CHRONIC IDIOPATHIC CONSTIPATION IN CHILDREN GUIDELINE

Appendix J - EVIDENCE TABLES

Key Components of the History Taking and the Physical Examination in Children with chronic constipation

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------|----------------------|-------------------------------|----------------------------|---------------------------------------|---|
| | level | prevalence | | standard | | |
| Borowitz et al. | Study type: | 220 children | 220 children | <u>Test</u> | Degree of difficulty with toilet | Additional information from study |
| Precipitants of | Case-control | | | History of events | training (mean ± SD) | Constipation defined as passage of < 3 |
| constipation | | <u>Inclusion</u> | -Patients | occurring in the 3 | (0=none, 4=extreme) | bowel movements each week for at |
| during early | Evidence | criteria: | n=125 | months prior to | | least 2 consecutive weeks |
| | <u>level</u> : | Aged 2y 0m to | mean age | onset of | Patients: 2.1±1.3 | |
| 2003. Journal of | Ш | 6y 11m, at least | (months): 44±13 | constipation: | Controls: 1.4±1.1 | 22 non-patient siblings matched as |
| the American | | average | 49% male | | p<0.001 | controls, an additional 73 non-sibling |
| Board of Family | Study aim: | intelligence | | -large/painful | | controls recruited from advertisements |
| Practice 16[3], | То | _ | -Controls | bowel movement | Degree of difficulty passing | |
| 213-218United | determine | - patients: First | n=95 | -toilet training | some bowel movements (% | Likert scale: 0 to 4. 0 being not at all |
| States. | the | time | mean age | -started day care | children) | difficult and 4 being extremely difficult |
| Borowitz, 2003 | precipitants | presentation to | (months): | -travelling | · | |
| | to | physician with | 46±18 | -liquid to solid | None: patients 3, controls 49 | Questionnaire for parents to fill out |
| | constipation | constipation | 54% male | foods | Mild: patients 86, controls 49 | describing children's bowel habits. |
| | in early | | | -breast to bottle | Moderate: patients 80, | - indication of how difficult toilet training |
| | childhood | - controls: no | | -family move | controls 10 | had been for bowel movements using |
| | | history of | Country: | -vomiting | Extreme: patients 76, controls | Likert scale |
| | | constipation | USA | /dehydration | 5 | - parents to indicate if any of 18 |
| | | | | -new medication | | different events occurred in the 3 |
| | | Exclusion | Setting: | -parental | p<0.001 (patients as compared | months preceding the onset of |
| | | criteria: | 26 primary care | separation | to controls in each category) | constipation, and which of these they |
| | | Underlying | facilities (15 | -birth of a sibling | | believed contributed to the onset of |
| | | medical | paediatricians, 11 | -tent camping | Degree of pain passing some | constipation |
| | | condition, | family medicine | -high fever | bowel movements (% children) | |
| | | medication that | centres) | -surgery | | Both groups comparable regarding age |
| | | could account | · | -extended bed | None: patients 5, controls 56 | and sex |
| | | for constipation | | | Mild: patients 82, controls 40 | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|--|---|
| | & Evidence | patients & | | Reference standard | | Reviewer comments Potential recall bias Source of funding: NIH grant RO1HD 28160 |
| | | | | | from breast to bottle and from liquid to solid diets having occurred more often before | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|------------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|--|------------------|
| | | | | | constipation in the younger children (30% vs. 0). Large or painful bowel movements were seen by far the most frequent precipitating event for both age groups. Toilet training was seen as more of a precipitant for older onset children (20% vs. 10%), whereas transition from breast to bottle and from liquid to solid foods was seen to be more of a problem for younger-onset children (25% vs. 0) | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---|---|--|---|---|--|---|
| The crying infant: Diagnostic testing and frequency of serious underlying disease. 2009. Pediatrics 123[3], 841-848United States. | Study type: Retrospectiv e case series Evidence level: III Study aim: To determine the proportion of children | 238 patients Inclusion criteria: - less than 12 months age - afebrile - presenting to ED during 9 month eligibility period with chief complaint of crying Exclusion criteria: Not stated | 238 patients Males 124 (52%) Median age 2.3 months (range 1.0 to 5.4) Country: Canada Setting: Tertiary care referral hospital | Tests Abdominal radiograph Abdominal ultrasound Reference Standard | constipation -History and examination were found to be the most important aspect in the evaluation of the crying infant. Investigations only helpful in 3% of sample in | cause 4) Neither Hx, PE nor investigations were diagnostic Required sample size calculated to yield stable estimates (±5%) of the primary outcome measure (proportion of infants who had potentially serious underlying aetiology). Estimated that |
| | | | | | this study | 10% sample would have underlying serious aetiologies. Minimum sample of 138 subjects required. Anticipated |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|---|
| | | | | | | follow-up telephone call response rate of only 75%. Final size after adjustment:: 245 |
| | | | | | | Reviewer comments No data on follow up care of accuracy of constipation cases |
| | | | | | | Minimum sample size required not achieved |
| | | | | | | Source of funding: Not stated |
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| | eudy type Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---|---|--|--|---|---|--|
| Diagnosing Hirschsprung's disease: increasing the odds of a positive rectal biopsy result. 2003. Journal of Pediatric Surgery 38[3], 412-416 Lewis et al., 2003 the phy exa and radi eva wou to a unn rect | trospective ohort dence el: dence el: test the pothesis of the history, ysical dence delication and dence delication allow devoid decessary of tall psies | Children presenting with constipation to | 315 children: -265 children who hade undergone rectal biopsy -50 children, concurrent selected cohort (cohort 2) Country: USA | Tests: Rectal biopsy | Clinical features in children with Hirschsprung's disease and idiopathic constipation (IC, n=40) -Onset of constipation <1 year old Delayed passage of meconium (%) HD: 65 IC: 13 P< 0.05 Abdominal distension (%) HD: 80 IC: 42 P< 0.05 Vomiting (%) HD: 72 IC: 21 P< 0.05 Faecal impaction requiring manual evacuation (%) HD: 6 IC: 30 P< 0.05 Enterocolitis (%) HD: 13 IC: 15 NS -Onset of constipation >1 year old Delayed passage of meconium | Additional information from study Questionnaires, telephone interviews and patients visits used to compile long- term data. In reporting features listed in the questionnaire only patients with definite information were included: the number of patients in each analysis varies to exclude those with missing data Delayed passage of meconium defined as failure to pass meconium in the first 48h of life. These data were available in 59% of cases Abdominal distension determined from parental response to questionnaire or data noted during patients visits Enterocolitis defined as diarrhoea associated with fever Reviewer comments: Data on clinical features not available for all children Unclear what kind of rectal biopsy was performed and how the diagnosis of HD was made Source of funding: Not stated |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|---|------------------|
| | | | | | HD: 81 IC: 1 P< 0.05 | |
| | | | | | Abdominal distension (%) HD: 53 IC: 7 P< 0.05 | |
| | | | | | Vomiting (%) HD: 23 IC: 0 P< 0.05 | |
| | | | | | Faecal impaction requiring manual evacuation (%) HD: 46 IC: 30 NS | |
| | | | | | Enterocolitis (%) HD: 13 IC: 14 NS | |
| | | | | | Age at onset of symptoms -Hirschsprung's (HD) (n=46) Mean: 8 months (range 1 day to 9 years) 1rst week of life: 60 % 1rst month of life: 70% 1rst year of life: 87% after 1 year of life: 13% | |
| | | | | | -Idiopathic constipation (IC) (n=40) Mean: 15 months (range 7 days to 16 years) | |

| 1rst week of life: 15% 1rst month of life: 55% 1rst year of life: 68% after 1 year of life: 32% At least 34% of HD patients had the classic triad (delayed passage of meconium + vomiting + abdominal distension). At least 1 feature of the triad noted in 98% of patients with HD. Only 60% of patients with IC had a history of delayed passage of meconium, vomiting or apdominal distancion, 100 % | Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|--|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|---|------------------|
| HD patients vs. 64% IC patients had 1 or more of the following: delayed passage of meconium, vomiting, abdominal distension and a transition zone on contrast enema. 36% of IC patients had none of these features. | | | | | | 1rst month of life: 55% 1rst year of life: 68% after 1 year of life: 32% At least 34% of HD patients had the classic triad (delayed passage of meconium + vomiting + abdominal distension). At least 1 feature of the triad noted in 98% of patients with HD. Only 60% of patients with IC had a history of delayed passage of meconium, vomiting or abdominal distension. 100 % HD patients vs. 64% IC patients had 1 or more of the following: delayed passage of meconium, vomiting, abdominal distension and a transition zone on contrast enema. 36% of IC patients had | |

Diagnostic Value of the Digital Rectal Examination (DRE) Children with Chronic Idiopathic Constipation

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|--|
| Beckmann et al. | Study type: | 251 children | Group 1: | Test: | Clinical variables (as a model) | Abdominal radiograph was either a |
| Accuracy of | Prospective | | 141 children with | Clinical variables | Sensitivity: 77% (+) | single flatplate or a flatplate with upright |
| clinical | case series | <u>Inclusion</u> | radiologically | | Specificity: 35% (-) | view, ordered by the ED attending |
| variables in the | | criteria: | proven | -History of | PPV 60% | physician based on customary |
| identification of | Evidence | Children aged | constipation | gastrointestinal | NPV: 55% | practices. The ED physicians ordering |
| radiographically | <u>level</u> : | 2-12 years old | | problems | | the radiographs were blinded to study |
| proven | III | who presented | Age: 7.9 +-3.1 | | Only the following clinical | objectives |
| constipation in | | to the | years | abdominal pain | variables were significantly | |
| children. 2001. | | Emergency | 63 (25%) male | -Stool habits | different between the two | 32% of the enrolled subjects did not |
| | Study aim: | Department | | -Straining on | groups: | undergo rectal exam |
| | to determine | (ED) of | | defecation | | |
| 100[1], 33- | whether | Children's | Group 2: | -Faecal consistency | History of normal/hard stool | A clinical diagnose previous to |
| 36United | clinical | Hospital of | 110 children with | | consistency: | radiology was made and reported. |
| States. | variables | Wisconsin with | no radiographic | -Medication | | However it was not clear how many of |
| | accurately | | evidence of | , | Group 1: | the clinical variables needed to be |
| | identify | and underwent | constipation | rebound, rigidity, | 74% (100/135) | present to diagnose constipation. |
| | children with | radiographic | | guarding, | | Furthermore, the physical exam was |
| | radiologically | evaluation. | Age: 7.4 +-3.0 | | Group 2: | completed by one of several paediatric |
| | proven | | years | _ | 61% (61/99) p: 0.016 | ED physicians and no assessment of |
| | constipation | <u>Exclusion</u> | 57 (23%) male | tenderness: diffuse, | | inter-rater reliability was performed. |
| | | criteria: | | each of four | Absence of rebound | |
| | | previous | Country: | quadrants, flank, | <u>tenderness</u> | Official radiologic diagnosis was |
| | | abdominal | USA | epigastric, | | provided by a single board certified |
| | | surgery, known | | | Group 1: | paediatric radiologist blinded to the |
| | | abdominal | | | 98% (138/141) | study. This was compared with the ED |
| | | pathology, | | bowel sounds, rectal | | physician interpretation of the |
| | | menarche or | | | Group 2: | radiograph and the patients were |
| | | sickle cell | | | 90% (99/110) p: 0.007 | divided into the two groups, but it is not |
| | | disease | | Clinical examination | | clear on the basis of what this decision |
| | | | | | Presence of left lower quadrant | |
| | | Setting: hospital | | exam) performed by | tenderness: | A data sheet with demographic-clinical |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|---|
| | | Emergency | | the ED physician | | data was required before an abdominal |
| | | Department | | | Group 1: | radiograph was ordered, but in 159 |
| | | | | Reference test: | 20% (19/96) | patients no data-sheet was submitted |
| | | | | Abdominal | | for various reasons. These patients |
| | | | | radiograph | Group 2: | were excluded from the study and the |
| | | | | | 9% (6/69) p: 0.0499 | lack of data makes impossible to tell |
| | | | | <u>Radiological</u> | | whether they differed from the group of |
| | | | | diagnose of | Stool present in rectal vault as | included patients |
| | | | | constipation (based | per rectal exam: | |
| | | | | on faecal loading | | Source of funding: |
| | | | | score originated and | | Not reported |
| | | | | | 69% (70/102) | |
| | | | | al and later revised | | |
| | | | | by Blethyn et al) | Group 2: | |
| | | | | | 43% (29/68) p: 0.008 | |
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| | | | | -Normal, grade 0: faeces in rectum and cecum only -Grade 1, mild constipation: faeces in rectum, cecum and discontinuous elsewhere -Grade 2, moderate constipation: faeces in rectum, cecum with continuous faeces affecting all segments but allowing for gas -Grade 3, severe constipation: continuous faeces with dilated colon and rectal impaction | 43% (29/68) p: 0.008 | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------|----------------------|-------------------------------|----------------------------|---------------------------------------|---|
| | level | prevalence | | standard | 4.14.11.1 | |
| Rockney et al. | Study type: | 60 encopretic | Age: 4-11 years | Test: | Values for rectal examination: | 78 encopretic children originally |
| The plain | Retrospectiv | children | old | Rectal examination | | enrolled but only 60 children for whom |
| abdominal | e case | | | | a) When the diagnosis of | Rx could be retrieved were included in |
| | series | Inclusion/ | Group 1 | Reference test: | retention by abdominal RX, | analysis. There were no significant |
| in the | | <u>exclusion</u> | 47 encopretic | Plain abdominal | systematic reading was agreed | differences between encopretic children |
| management of | | <u>criteria</u> | children with | roentgenogram | by at least two radiologists: | whose abdominal Rx were reviewed for |
| encopresis. | <u>level</u> : | Encopresis as | faecal retention | | | the study and those who did not have a |
| 1995. Archives | III | defined by the | by | Three radiologists, | (%) | Rx or whose Rx could not be retrieved. |
| of Pediatrics | | DSM Revised | roentgenogram | two paediatric and | Sensitivity: 88.6 | There were no significant differences in |
| | Study aim: | Third Edition: | criteria on | one general, at | Specificity: 41.6 | patients' characteristics at the two sites. |
| | to determine | "repeated | presentation | three separate | Positive predictive value: 84.8 | Not all data were available for every |
| 149[6], 623-627 | whether | involuntary (or, | | institutions, blind to | Negative predictive value: 50 | subject |
| | faecal | much more | Male sex: 74.5% | the identity of the | | |
| | retention in | rarely, | _ | | b) When the diagnosis of | Children with retention (as per Rx) were |
| | encopretic | intentional) | Group 2 | the plain abdominal | retention by abdominal RX, | significantly more likely to have stool in |
| | | passage of | 13 encopretic | Rx twice: a | systematic reading was agreed | the rectum on presentation (p 0.015) |
| | be assessed | | children without | "subjective" reading | by the three radiologists: | and were significantly less likely to have |
| | objectively | places not | faecal retention | assessed faecal | | parents report a difficult toilet training (p |
| | using the | appropriate for | by | content as markedly | | 0.018). There were no other significant |
| | plain | that purpose | roentgenogram | excessive, | Sensitivity: 91.7 | differences between the two groups |
| | abdominal | | | moderately | Specificity: 71.4 | regarding the rest of the variables |
| | | floor)the | presentation | | Positive predictive value: 94.3 | measured. |
| | am and | event must | | and a "systematic" | Negative predictive value: 62.5 | |
| | whether | occur at least | Male sex: 61.5 % | | | Each patient's medical record was |
| | | once a month | | | Not all data were available for | reviewed separately by one of the |
| | aphic | for at least 6 | Country: | record was | every subject | authors and a research assistant. When |
| | evidence of | months, the | USA | completed and a | | discrepancies existed charts were |
| | faecal | chronological | | score assigned (0- | | reviewed again conjointly and |
| | retention is | and mental age | | 25) reflecting the | | discrepancies resolved for both |
| | associated | of the child | | severity of faecal | | reviewers' satisfaction. |
| | with clinical | must be at least | | retention (score of | | The condition of the condition of the |
| | findings on | 4 years, and | | 10 or greater | | The reliability of the radiologists' |
| | | physical | | indicates faecal | | assessments was tested by two |
| | | disorders that | | retention, scale | | different procedures. |
| | children | can cause | | validated by Barr et | | Overall agree agree at agree agree the three |
| | | faecal | | al.) Final results | | Overall agreement among the three |
| | | incontinence, | | were taken from the | | radiologists was 77.8% for the |
| | | such as | | systematic reading | | subjective assessment, k=0.53 (z=7.04, |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|--|
| | ievei | aganglionic | | only. At least two | | p<0.0001). Agreement using the |
| | | megacolon, | | radiologists had to | | systematic assessment was 87.4%, |
| | | must be ruled | | agree in order to | | k=0.65 (z=7.2, p<0.0001). There were |
| | | out" | | classify | | no differences in interrater reliabilities |
| | | Children | | roentgenograms | | between pairs of radiologists. |
| | | younger than 4 | | either as in the | | bottion pane of radiologists. |
| | | years old and | | retention or | | The study from which the systematic |
| | | children who | | nonretention | | scoring system was derived has not |
| | | had a soiling | | category | | been replicated, and the cut-off point of |
| | | frequency of | | Presence of stool in | | 10, might not be valid for all populations |
| | | less than once | | rectal examination | | |
| | | a month or who | | was recorded in the | | Source of funding: |
| | | had recently | | patient records as | | Primary Care Faculty Development |
| | | stopped soiling | | "none", "small", | | Fellowship Programme at Michigan |
| | | were excluded | | "moderate" or | | State University, East Lansing. |
| | | | | "large" amount. | | |
| | | Setting: two | | Patients with | | |
| | | paediatric | | moderate or large | | |
| | | incontinence | | amounts of stool on | | |
| | | clinics, one | | rectal examination | | |
| | | located in the | | were classified as | | |
| | | ambulatory care | | having stool in the | | |
| | | facility of a | | rectum for | | |
| | | tertiary care | | subsequent | | |
| | | hospital and the | | analysis. | | |
| | | other at a | | | | |
| | | community | | The specific | | |
| | | hospital | | professional | | |
| | | | | qualification of the | | |
| | | | | person who | | |
| | | | | performed the rectal | | |
| | | | | examination was not | | |
| | | | | reported | | |

Prevalence of Coeliac Disease and Hypothyroidism in children with Chronic Idiopathic Constipation

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comment | |
|---------------------------|-----------------------------------|--------------------|-------------------------------|---------------------|--|--|--|
| Bonamico et al. | Study type: | 1202 patients | 1202 patients | Tests: | Signs/symptoms (%): | Additional information from study | |
| Prevalence and | Prospective | | | -Coeliac disease: | | Levels of IgA AGA were measured by | |
| clinical picture | cohort | <u>Inclusion</u> | 609 males | | -Group 1 (n=55): | enzymelinked immunosorbent assay by | |
| of celiac | | criteria: | | Revised | | the Alfa-gliatest (Eurospital, Trieste, | |
| disease in | <u>Evidence</u> | Down's | 1110 children | European | Growth failure 52.7 | Italy). Levels of EMA IgA were | |
| Italian down | level: | syndrome | age range: 15 | Society of | Diarrhoea 41.8 | evaluated by an indirect | |
| syndrome | 2+ | | months to 18 years | Paediatric | Vomiting 20 | immunofluorescence method | |
| patients: A | | Exclusion | | Gastroenterology, | Anorexia 18.2 | (Eurospital, Trieste, Italy). Sections | |
| multicenter | Study aim: | criteria: | 92 adults | Hepatology and | Constipation 29.1 | from the distal portion of monkey | |
| study. 2001. | To estimate | IgA deficiency | age range 18 to 46 | Nutrition | Distended abdomen 23.6 | oesophagus were used as a substrate, | |
| Journal of | the | | years | (ESPGHAN) | | and fluorescein-labeled goat antihuman | |
| Pediatric | prevalence | Setting: | | criteria Patients | -Group 2 (n=55): | IgA antibody was used as the second | |
| Gastroenterolog | of coeliac | Community | Country: | selected for | | antibody. The patients' serum was | |
| y and Nutrition | disease (CD) | | Italy | intestinal | Growth failure 10.9 | diluted 1:5 in phosphate buffer at pH | |
| 33[2], 139- | in patients | | | biopsy on the | Diarrhoea 1.8 | 7.2. The presence of a brilliant green | |
| 143United | with Down | | | basis of EMA | Vomiting 1.8 | network pattern under a fluorescence | |
| States. | syndrome | | -Group 1: 55 CD | positivity, AGA | Anorexia 1.8 | microscope was taken as a positive | |
| | and to define | | patients diagnosed | IgA positivity, | Constipation 14.5 | result. Intestinal biopsies performed by | |
| | the clinical | | by ESPGHAN | or both in children | Distended abdomen 14.5 | Watson capsule or by paediatric or | |
| | characteristi | | Criteria (36 males, | < 2 years of age | | adult endoscopes | |
| | cs of CD | | aged 4 to 46 years) | | -Group 3 (n=57): | | |
| | among | | | (AGA: antigliadin | | Patients selected for intestinal biopsy | |
| | Down | | -Group 2: 55 lgA | | Growth failure 7 | on the basis of both EMA positivity and | |
| | Syndrome | | AGA-positive EMA | antiendomysium | P < 0.001 | AGA IgA positivity in children < 2 years | |
| | patients | | negative DS | antibodies; IgA: | Diarrhoea 6.9 | of age, because in this age group, EMA | |
| | | | patients (33 males, | immunoglobulin | <i>P</i> < 0.001 | positivity may have a false-negative | |
| | | | aged 3 to 40 years) | A) | Vomiting 1.7 | result | |
| | | | | | P < 0.001 | | |
| | | | -Group 3: 57 IgA | -Down syndrome: | Anorexia 3.4 | A detailed questionnaire was completed | |
| | | | AGA-negative | confirmed by | P < 0.01 | to obtain information about familial | |
| | | | EMA-negative DS | cariotype in all | Constipation | gastroenterologic history with special | |
| | | | patients (34 males, | cases | 8.8 <i>P</i> < 0.05 | attention to feeding habits (breast milk | |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comment |
|---------------------------|-----------------------------------|--------------------|-------------------------------|------------------|--|---|
| | | | aged 4 to 38 years) | | | or formula, age of introduction of gluten-containing foods); gastrointestinal function, particularly the features of CD, such as chronic diarrhoea, vomiting, failure to thrive, and anorexia; presence of autoimmune or neoplastic conditions All patients were receiving a gluten-containing diet. Weight and height were evaluated using Down syndrome percentile charts (DSPC) The clinical features of 55 CD patients diagnosed by ESPGHAN Criteria (group 1) were compared with those observed in 55 IgA AGA-positive EMA negative DS patients (group 2) and in |
| | | | | | | 57 IgA AGA-negative EMA-negative DS patients (group 3). Group 2 and group 3 patients were selected randomly from among the screened patients to be age and gender matched to group 1. |
| | | | | | | 18 symptomatic patients belonging to group 2 underwent intestinal biopsy and showed normal small bowel mucosa |
| | | | | | | Parents of 8 EMA positive children and 2 EMA-positive adults did not give permission for intestinal biopsy to be performed and were not included among the 55 CD patients |
| | | | | | | Reviewer comments: It is unclear whether some patients had EMA and others had AGA IgA measured alternatively, or whether all patients had both EMA and AGA IgA |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comment |
|---------------------------|-----------------------------------|--------------------|-------------------------------|------------------|--|---|
| | level | | | | Effect Size | measured at the same. This considered it is also unclear why only IgA AGA-positive EMA-negative patients and IgA AGA-negative EMA-negative patients were chosen as control groups and there is no mention of the EMA-positive IgA AGA-negative group Source of funding: Not stated |
| | | | | | | |
| | | | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comment |
|--|--|---|--|--|--|---|
| Bingley et al. Undiagnosed coeliac disease at age seven: Population based prospective birth cohort study. 2004. British Medical Journal 328[7435], 322- 323United Kingdom. | Prospective cohort Evidence level: 2+ Study aim: to establish the prevalence of undiagnosed coeliac disease in the general population at age seven | based birth cohort study established in | 5470 children age: 7.5 years gender not reported Country: UK | radioimmunoassa y for antibodies to tissue transglutaminase (endomysial antigen) (tTG antibodies) 2. If positive to previous, serum IgA antiendomysial antibodies (IgA-EMA) by indirect | Any constipation reported at age 6.75 years (No, %): -tTG antibody negative controls (n=4285 questionnaires): 435 (10) -IgA-EMA positive (n=42 questionnaires): 6 (14) odds ratio (95% CI): 1.48 (0.62 to 3.52) Other symptoms reported at age 6.75 years (No, %): -tTG antibody negative controls (n=4285 questionnaires): any diarrhoea: 1450 (34) any vomiting: 1933 (45) any stomach pains: 2557 (60) ≥3 GI symptoms: 931 (22) -IgA-EMA positive (n=42 questionnaires): any diarrhoea: 21 (50) odds ratio (95% CI): 1.96 (1.06 to 3.59) any vomiting: 23 (55) odds ratio (95% CI): 1.47 (0.80 to 2.71) | Additional information from study Children with tTG antibodies < 97.5 th centile were defined as antibody negative Details of gastrointestinal symptoms and special diets collected by routine questionnaire at age 6.75 years Total tTG antibody negative controls (n=5333 children). Total IgA-EMA positive children (n=54) (1.0%; 95% confidence interval 0.8 to 1.4) 4324 children (79%) returned questionnaires An additional 137 children were tTG antibody positive, but Ig-EMA negative IgA-EMA were more common in girls (OR 2.12; 1.20 to 3.75). IgA-EMA positive children were shorter and weighted less than those who tested negative fro tTG antibody (p<0.0001 for all comparisons) Since ALPASC is an observational study based on analysis of anonymous samples, confirmatory biopsy was not possible Reviewer comments: Unclear how the symptom "constipation" was defined in the first place |
| | | | | | any stomach pains: 28 (66) | No data regarding clinical symptoms at |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comment |
|---------------------------|-----------------------------------|--------------------|-------------------------------|------------------|--|--|
| | | | | | odds ratio (95% CI): 1.35 (0.71 to 2.57) ≥3 GI symptoms: 17 (40) odds ratio (95% CI): 2.45 (1.33 to 4.5) | 6.75 years for 21% of the total sample Sources of funding: Coeliac UK, Medical Research Counci Wellcome Trust, UK government departments, and various charitable organisations and commercial companies, ALSPAC is part of the WHO initiated European Longitudinal Study on Pregnancy and Childhood |

| Bibliographic Information | Study type & Evidence | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures | Reviewer comment |
|---------------------------|-----------------------|----------------------|-------------------------------|------------------------------------|---------------------------------|---|
| | level | | | | Effect Size | |
| Cataldo et al. | Study type: | 1917 children | Total: 1917 children | | Clinical pattern and presenting | Additional information from study |
| | Retrospectiv | | with CD | -coeliac disease: | symptoms at diagnosis (n=36) | Classical forms not clearly defined, but |
| | e case | <u>Inclusion</u> | | diagnosis based | | included the following symptoms: |
| | series | criteria: | 36 immigrant | on the revised | -Classical forms (25/36) | chronic diarrhoea, weight loss, |
| immigrant | | Italian and | children with CD | criteria of the | (69.4%): | abdominal distension and vomit |
| | <u>Evidence</u> | immigrant | 15 males | European Society | | |
| | <u>level</u> : | children | age range 6 | of Paediatric | No child with constipation | Atypical forms included: iron-deficiency |
| An Italian | 3 | consecutively | months to 15 years | Gastroenterology | reported | anaemia, short stature, delayed |
| multicentre | O | diagnosed as | (mean 7.3) | and Nutrition | 4 | puberty, recurrent oral aphtae |
| | Study aim: | having CD | 4004 1/ 1/ | (ESPGAN): | -Atypical forms (9/36) (25%): | |
| Digestive and | To evaluate | between | 1881 Italian | 4 =: " (() (| A | Silent forms included: serological |
| | the | January 1999 to | | | Abdominal pain with | screening of first degree relative, loss of |
| | prevalence | December 2001 | | small intestinal | constipation: | Kerckring folds at endoscopy |
| | of immigrant | | age range 6 | mucosa with the | 2/9 | |
| | | | months to 16 years | features of | 0:1 | Clinical patterns in Italian children were |
| | coeliac | criteria: | (mean 7.9) | hyperplastic | -Silent forms (2/36) (5.5%): | similar to those of immigrant children |
| | ` ' | Not stated | Carrate v | villous atrophy on | No shild with aspetingtion | Daviewer comments: |
| | in Italy, the | Cattin au | Country: | histological | No child with constipation | Reviewer comments: |
| | | Setting: Hospital | Italy | examination of a | reported | Unclear how the symptom |
| | findings in these | (multicentre) | | biopsy specimen, while the patient | | "constipation" was defined in the first |
| | patients and | (municemie) | | is eating | | piace |
| | the possible | | | adequate | | Presenting symptoms at diagnosis were |
| | relationship | | | amounts of gluten | | not reported for Italian children |
| | between | | | amounts of gluten | | Thot reported for italian children |
| | immigration, | | | 2. Clear cut | | Source of funding: |
| | dietary | | | clinical remission | | Study supported by grants of Ministero |
| | habits and | | | on a strict gluten | | dell'Universita e della Ricerca |
| | CD in | | | free diet with | | Scientifica e Tecnologica (MURST) |
| | childhood | | | relief of all | | 60% di F.C. |
| | Ciliariood | | | symptoms of the | | 0070 011.0. |
| | | | | disease. This | | |
| | | | | response should | | |
| | | | | be reasonably | | |
| | | | | rapid occurring | | |
| | | | | within a matter of | | |
| | | | | weeks rather than | | |
| | | | | many months | | |

| tudy type Number of Evidence patients level | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comment |
|---|-------------------------------|--|--|------------------|
| | | 3. The finding of circulating antibodies (IgA gliadin, antireticulin, and antiendomysiun) at time of diagnosis and their disappearance when the patient is taking a gluten free diet add weight to the diagnosis | | |

| Bibliographic Information | Study type & Evidence | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures | Reviewer comment |
|--|-----------------------|---|--|---|---|--|
| | level | | | | Effect Size | |
| Egan-Mitchell et al. Constipation in childhood | level | patients 112 children Inclusion criteria: Coeliac disease Exclusion criteria: Not stated Setting: Regional and university hospitals | Characteristics 112 children 12 children with constipation: 6 males, age range 6 to 102 months Country: Ireland | normal, or the clinical observation of | Effect Size Incidence of constipation: 12 children constipated at some stage before diagnoses: -9 of those children presented with constipation and faecal impaction, of these 5 had intermittent diarrhoea and constipation but 4 never had diarrhoea. Of these 4, 3 children presented at around 1 year of age with anorexia, failure to thrive and faecal impaction -the 3 children who did not | Additional information from study Growth retardation assessed on the graphs of Tanner and Whitehouse (1959) and subsequently confirmed by catch-up growth following treatment with gluten-free diet Mucosal damage according to authors' classification (normal mucosa grade 0; mild non-specific change grade 1; grade 2 and 3 correspond to moderate and severe villous atrophy) Reviewer comments: Unclear whether authors' classification system for jejunal mucosa damage has been validated Source of funding: The main author was receiving a grant from the Medical Research Council of Ireland |
| | | | | impaction of abnormal amounts of hard | | |
| | | | | (usually pale) faeces in colon and rectum | | |

Diagnostic Value of the Anorectal Manometry in Children with Chronic Idiopathic Constipation

| Bibliographic | Study type | Number of | Population | Type of test (s) | Follow-up & Outcome | Reviewer comments |
|------------------|-----------------|------------------|--------------------|--------------------|--------------------------------|---|
| Information | & Evidence | patients | Characteristics | , , | Measures | |
| | level | | | | Effect Size | |
| Jarvi et al. | Study type: | 81 patients | 81 patients | Tests: | Rectoanal inhibitory reflex | Additional information from study |
| Anorectal | Retrospectiv | | 49 male | -Anorectal | (RAIR) and histology results | Records of all patients who met the |
| manometry with | e case | <u>Inclusion</u> | | manometry: | | inclusion criteria were reviewed |
| reference to | series | criteria: | median age at time | | -RAIR present (N=40) | |
| operative rectal | | Patients under | of ARM and biopsy: | Performed using | | In each case ARM was performed |
| biopsy for the | <u>Evidence</u> | 1 year of age | 2 months (range | a 4-cm long rectal | HD: no children | under ketamine anaesthesia by a |
| diagnosis/exclu | level: | who presented | 0.1 to 11 months) | balloon inflated | Normal histology: 39 children | consultant paediatric surgeon, and |
| sion of | III | with delayed | | incrementally with | Hypoganglionosis: 1 child | operative rectal biopsy was taken |
| Hirschprung's | | passage of | Country: | 5 to 50 mL of air | | simultaneously |
| disease in | Study aim: | meconium, | Finland | | -RAIR absent (N=41) | |
| children under 1 | To report on | abdominal | | -Operative rectal | | RAIR defined as greater than 25% drop |
| year of age. | the value of | distension and | | biopsy: | HD: 33 children | in the anal sphincter pressure for at |
| 2009. | anorectal | vomiting or | | | Normal histology: 8 children | least 5 seconds |
| International | manometry | constipation | | Taken 3 cm | | |
| Journal of | (ARM) with | who underwent | | above the dentate | Diagnostic variables for ARM | Patients who had HD were significantly |
| Colorectal | reference to | ARM | | in the posterior | and operative rectal biopsy in | younger at the time of investigation than |
| Disease 24[4], | operative | | | midline, | HD (%): | those who did not |
| 451- | rectal biopsy | <u>Exclusion</u> | | consisting of a | | |
| 454Germany. | in the | criteria: | | generous, | -Biopsy: | In the case of patients diagnosed with |
| | diagnosis/ex | Other | | Iongitudinal | | HD histology from bowel resected at |
| | clusion of | congenital | | specimen | Sensitivity: 100 | pull-through operation was consistent |
| | | gastrointestinal | | extending to the | Specificity: 100 | with pre-operative diagnosis in all cases |
| | g's disease | malformations | | submucosa | Positive predictive value: 100 | |
| | in children | such as | | | Negative predictive value: 100 | Operative rectal biopsy was adequate |
| | | anorectal | | | | and diagnostic in all cases. There was |
| | of age and | anomaly, funnel | | | -ARM: | one case of rectal bleeding following |
| | on the | anus or | | | | biopsy which required suturing in |
| | prognostic | gastroschisis | | | Sensitivity: 100 | theatre |
| | significance | | | | Specificity: 83 | |
| | of a normal | | | | Positive predictive value: 80 | Reviewer comments: |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---------------------------|-----------------------------------|--------------------|-------------------------------|------------------|--|--|
| | RAIR in these patients | | | | Regative predictive value: 100 | Unclear how the reviewing process was conducted Unclear how the biopsy specimens were processed and analysed Source of funding: Not stated |
| | | | | | | |

| Bibliographic Information | Study type & Evidence | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures | Reviewer comments |
|---------------------------|--------------------------|-----------------------------|-------------------------------|--------------------------|---|--|
| | level | | | | Effect Size | |
| Lee et al. | Study type: | 105 children | 105 children | Tests: | Rectoanal inhibitory reflex | Additional information from study |
| | Retrospectiv | | 61 boys | -Anorectal | (RAIR) and histology results | Severe abdominal distension defined as |
| and abdominal | e case | <u>Inclusion</u> | | manometry: | | an abdominal wall that protruded, was |
| distention | series | criteria: | Mean age: 2.1 ± | | -RAIR absent (N=48) | shiny and tense upon palpation |
| mimicking | | Infants < 6 | 0.9 months | Performed by | | |
| Hirschsprung's | <u>Evidence</u> | months of age | | paediatricians | HD: 34 | Reviewer comments: |
| disease in | <u>level</u> : | with severe | Country: | using a silicon | Normal histology: 10 | Unclear how the reviewing process was |
| infants. 2007. | III | abdominal | Korea | rubber catheter | AP: 2 | conducted |
| Acta | | distension that | | with an array of 8 | IND: 2 | |
| Paediatrica, | Study aim: | mimicked HD | | channels of | | Unclear what was the order in which |
| International | To evaluate | referred to | | sensors. Sedation | | investigations were carried out |
| Journal of | the | department of | | with chloral | -RAIR present (N=57) | |
| | incidence | paediatrics and | | hydrate for the | | Source of funding: |
| 96[12], 1784- | and clinical | division of | | procedure was | HD: 5 | Not stated |
| 1789United | aspects of | paediatric | | used | Normal histology: 43 | |
| Kingdom. | allergic | surgery and | | | AP: 5 | |
| | proctitis (AP) | underwent all | | -Suction rectal | IND: 4 | |
| | in patients | triple tests | | biopsy: | D | |
| | with | including | | | Diagnostic variables for ARM | |
| | symptoms | barium enema, | | Taken from 4 | and rectal suction biopsy in HD | |
| | that mimic | anorectal | | different sites | <u>(%):</u> | |
| | Hirschsprun | manometry and | | using a rectal | Diaman. | |
| | g's disease | rectal suction | | suction biopsy | -Biopsy: | |
| | (HD). In | biopsy. Some | | tube. Biopsy sites | Concitivity | |
| | addition authors | patients had associated | | were 3cm and 5 | Sensitivity: | |
| | determined | | | cm for anal verge. | , | |
| | the | symptoms like constipation, | | When ganglion cells were | Specificity: 100 % (94.50 to 100.00) | |
| | sensitivity | poor oral intake, | | observed to be | Positive predictive value 100% | |
| | and | vomiting, poor | | present with | Negative predictive value: | |
| | specificity of | weight gain and | | normal | 95.65% | |
| | anorectal | diarrhoea | | appearance on | 30.0070 | |
| | manometry | diaminoda | | haematoxylin- | -ARM: | |
| | and | Exclusion | | eosin staining HD | 7 11 11 11 | |
| | suction | criteria: | | was excluded. | Sensitivity: | |
| | rectal biopsy | Coeliac disease | | HD was finally | 87.18% (CI: 73.29 to 94.90) | |
| | used for | and cystic | | diagnosed with | Specificity: | |
| | evaluation of | | | full thickness | 78.79% (CI: 67.49 to 86.92) | |

| Bibliographic Information | Study type & Evidence level | | Population Characteristics | Type of test (s) | Measures Effect Size | | | Reviewer comments |
|---------------------------|-----------------------------------|--|-------------------------------|------------------|------------------------------|------------|--------|-------------------|
| | HD | considered in the differential | | biopsy | Positive 70.83% | predictive | value: | |
| | | the differential diagnosis because are extremely rare in Korea | | | 70.83% Negative 91.23% | predictive | value: | |
| | | | | | | | | |
| | | | | | | | | |

| Bibliographic Information | Study type & Evidence | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures | Reviewer comments |
|---|---|--|---|--|--|--|
| Kong et al. Screening Hirschsprung's disease by anorectal manometry. 1993. Chinese Journal of Gastroenterolog y 10[1], 29- 32Taiwan, Province of China. | Study type: Retrospective case series Evidence level: III Study aim: To evaluate the possibility of using anorectal manometry (ARM) for screening Hirschsprung's disease (HD) | 39 patients Inclusion criteria: Children with constipation or suspected HD Exclusion criteria: Systemic diseases like hypothyroidism or neurologic disorders | 39 patients age range: 3 days to 9 years (no other details provided) Country: Taiwan | Tests: -Anorectal manometry: Double lumen stainless steel manometric probes with internal diameter of 6 mm used. Entire system closed and water filled. Multiple-channel recorder used for recording results. No previous bowel preparation. Stimulus balloon placed from 3 to 5 cm from anal verge, depending on size of patients. For uncooperative patients intramuscular injection with mixture of chlorpromazine, promethazine and meperidine with or without intravenous diazepam was given | Sensitivity: 100 Specificity: 86 PPV: 83 NPV: 100 | Additional information from study A normal reflex (RAIR) was present when rythmicity of internal sphincter contractility was totally inhibited by rectal distension accompanied by a simultaneous drop of internal sphincteric pressure fro 5mmHg or more. A positive rectoanal response consisted of 3 successive pressure falls, each immediately following upon rectal distension by balloon. When rythmicity and internal sphincter pressure remained unchanged following rectal distension, the amount of air was increased gradually to 10 cc fro neonates and 50 cc for children. If RAIR was absent, a negative response was recorded The final diagnosis of HD was made by patient's clinical history, barium enema and rectal suction biopsy Inconclusive results with manometry due to poor tracing of internal sphincter contraction as a result of oversedation (n=2) and to anal stenosis (n=1) Reviewer comments: No definition of constipation given Insufficient details on how HD was diagnosed It is not completely clear whether or not all patients underwent rectal biopsy but it looks as this was probably the case |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---------------------------|-----------------------------------|--------------------|-------------------------------|---|--|--|
| | level | | | - Rectal suction biopsy (no other details provided) | Effect Size | The 3 children in whom manometry was inconclusive were not included in the calculation of the diagnostic variables and this introduces bias Source of funding: Not stated |
| | | | | | | |

| Bibliographic Information | Study type & Evidence | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures | Reviewer comments |
|---------------------------|--------------------------|--------------------|-------------------------------|--------------------|---------------------------------|--|
| | level | | | | Effect Size | |
| | Study type: | 261 patients | 261 patients | Tests: | Rectoanal inhibitory reflex | Additional information from study |
| Pitfalls and | Prospective | | | -Anorectal | (RAIR) and histology results | In no case the result of a rectal biopsy |
| limitations of | case series | <u>Inclusion</u> | -gender not | manometry | | was known at the time of manometry |
| testing the | | <u>criteria:</u> | reported for all | | -RAIR equivocal result | |
| rectoanal | <u>Evidence</u> | Patients | patients | No special bowel | (absent?): | RAIR considered to be present if the |
| inhibitory reflex | <u>level</u> : | referred for | | preparation given. | 9 children | anal pressure decreased on rectal |
| | III | anorectal | -Age: | if a considerable | | distension followed by recovery of the |
| hirschsprung's | | manometry in | < 6 months: 94 | amount of faecal | HD: 4 | basal tone. RAIR was also considered |
| disease. 1990. | Study aim: | order to confirm | (36%) | impaction was | Normal histology: 5 | to be present if the typical anal pressure |
| Pediatric | To better | or exclude | 6 month to 6 years: | found, patients | | waves were clearly abolished |
| Surgery | | Hirschsprung's | 106 (41%) | were sent back | -RAIR equivocal result | |
| International | traps and | disease. All | 6 to 15 years: 47 | for evacuating | (present?): | Confident interpretation of the RAIR |
| 5[4], 260- | | patients had | (18%) | enema (s) and | 8 children | was made in 232/261 patients (89%): |
| 265Germany. | testing the | presented with | 2 adolescents and | reexamination | | RAIR present in 207 cases and absent |
| | rectoanal | constipation | 12 adults (5%) | planned for the | HD: 2 | in 25. The result of this first manometric |
| | inhibitory | varying from | | next day. | Normal histology: 6 | evaluation was verified either by biopsy |
| | reflex | slight to | Country: | Children not | | or by repeated manometry in 54 cases. |
| | | intractable, with | Belgium | sedated. Entire | -RAIR confident interpretation: | In other cases the clinical evolution did |
| | frequently | highly differing | | system filled with | 232 children | not warrant further investigation. |
| | they occur | durations | | degassed water. | | |
| | and the | ranging from | | Multiple-channel | RAIR+: 207 | Manometrically the following factors |
| | possible | neonatal ileus | | | RAIR-: 25 | prevented examiners from reaching a |
| | • | to chronic | | recording results | | definite conclusion: low anal tone (n=8), |
| | | constipation in | | | Of the previous 54 children | restlessness of patient (n=7), reflex |
| | or false | adults | | - Superficial | underwent either biopsy or | external sphincter contraction partially |
| | results | | | biopsy of rectal | repeated manometry. Only | or completely masking possible RAIR |
| | | <u>Exclusion</u> | | mucosa and | false results reported: | (n=4), presence of megarectum (n=3), |
| | | criteria: | | submucosa taken | | artifacts (n=1), unstable RAIR (n=6) |
| | | Not stated | | with a laryngeal | -RAIR present and HD: 2 | |
| | | | | biopsy forceps. | children | Reviewer comments: |
| | | | | Frozen section | | Not all children underwent both |
| | | | | biopsies stained | -RAIR absent and normal | manometry and biopsy: 261 patients |
| | | | | for | histology: 4 children | underwent manometry and only 24 |
| | | | | acetylcholinestera | | underwent biopsy |
| | | | | se and | Incidence of false results and | |
| | | | | nicotinamide | age of patients at first | Details of both the manometry and |
| | | | | adenine | manometry | biopsy results were reported only in |
| | | | | dinucleotide- | | cases where the RAIR was equivocal in |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---------------------------|-----------------------------------|--------------------|-------------------------------|---|---|---|
| | | | | reduced diaphorase. Aganglionosis with hypertrophic bundles was diagnostic for HD | -In <1 month old: 5/22 (22.7.8%) -In > 1 month old: 4/239 (1.7%) Incidence of equivocal results and age of patients at first manometry -In <1 month old: 4/22 (18.2%) -In > 1 month old: 25/239 (10.4%) | the first manometry and in those children where the result proved to be false (either negative or positive). Considering this it is not possible to calculate the sensitivity, specificity, positive and negative predictive values of the anorectal manometry The incidence of false results in manometry performed by different examiners is reported in the paper, but there are missing data not accounted for and therefore we do not report it here Source of funding: Not stated |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---------------------------|-----------------------------------|--------------------|-------------------------------|----------------------------------|--|---|
| Low et al. | Study type: | 50 children | 50 children | Tests: | Rectoanal inhibitory reflex | Additional information from study |
| | Prospective | | | -Anorectal | (RAIR) and histology results | 5 children (10%) required repeat full- |
| | case series | Inclusion | (data available for | manometry | | thickness biopsy for inadequate |
| manometry in | | criteria: | 45 children) | | -RAIR absent (N=16) | sampling |
| the diagnosis of | Evidence | Children | 31 male | Performed as a | , , | |
| Hirschsprung's | level: | referred | 14 female | side-room | HD: 15 | All children underwent both manometry |
| disease. 1989. | III | consecutively to | | procedure. All | Normal histology: 1 | and biopsy. |
| Journal of | | one of the | Age range birth to | children under 4 | | |
| | Study aim: | authors for | 11 months | years of age who | -RAIR present (N= 34) | Biopsy specimens prepared in paraffin |
| Gastroenterolog | | anorectal | | were unable to | | sections and stained with haematoxylin |
| | accuracy of | manometric | Country: | cooperate were | HD: 4 | and eosin. Up to 60 6-µm-thick serial |
| L 3' | anorectal | studies | Singapore | tested after oral | Normal histology: 30 | sections of each specimen were |
| | manometry | | | sedation with | | examined histologically by pathologist |
| | in the | Exclusion | | chloral hydrate | Diagnostic variables for ARM, | for ganglion cells and hypertrophied |
| | diagnosis of | criteria: | | | total sample N=50 (%): | nerve bundles. Specimens not including |
| | Hirschsprun | Not stated | | | | the submucosal layer were considered |
| | g's disease | | | -Suction | Accuracy: 90 | inadequate and repeat full-thickness |
| | (HD) using | | | rectal biopsy | Sensitivity: 79 | operative rectal biopsies were taken |
| | histological | | | | Specificity: 97 | |
| | aganglionosi | | | Suction rectal | Positive predictive value: 94 | A normal reflex was present when |
| | s as the | | | biopsies obtained | Negative predictive value: 88 | rythmicity of internal sphincter |
| | reference | | | without | D | contractility was totally inhibited by |
| | point for final | | | anaesthesia by | Diagnostic variables for ARM, | rectal distension accompanied by |
| | diagnosis | | | paediatric | neonates N=10 (%): | simultaneous drop in internal |
| | | | | surgeon on | A | sphincteric pressure. Rythmicity and |
| | | | | outpatient basis. | Accuracy: 90 | tone recovered when rectal distension |
| | | | | Biopsies taken at 4 cms from the | Specificity: 100 | was removed. When rythmicity and |
| | | | | | Positive predictive value: 100 | internal sphincter pressure remained virtually unchanged after rectal |
| | | | | | Negative predictive value: 75 | distension a negative response was |
| | | | | biopsy set. | Negative predictive value. 75 | recorded |
| | | | | niopsy set. | Diagnostic variables for ARM, | lecolueu |
| | | | | | infants N=18 (%): | No complications encountered with |
| | | | | | <u> </u> | manometry in all 50 children studied |
| | | | | | Accuracy: 94.4 | manomoti y in all oo omidren stadied |
| | | | | | Sensitivity: 90 | Reviewer comments: |
| | | | | | Specificity: 100 | No definition of constipation/idiopathic |
| | | | | | Positive predictive value: 100 | constipation given |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---------------------------|-----------------------------------|--------------------|-------------------------------|------------------|--|---|
| Information | | patients | Characteristics | | | Unclear what "infant" meant for authors Source of funding: Research grant (RP53/81) from the National University of Singapore, Singapore |
| | | | | | | |

Diagnostic Value of the Plain Abdominal Radiography in Children with Chronic Idiopathic Constipation

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|---------------------------------------|---|
| Reuchlin- | Study type: | 6 studies (3 | Otherwise healthy | Test and | Diagnostic value: | MEDLINE searched from inception to |
| Vroklage et al. | Systematic | case series, 2 | children aged from | Reference | | April 2004, search terms reported and |
| Diagnostic | Review | case-control | 1 to 18 years old | Standard (studies | (LR: Likelihood ratio) | comprehensive. Results of this search |
| value of | | studies, | with signs and | could treat either | | combined with search strategy specific |
| abdominal | <u>Evidence</u> | 1retrospective | symptoms related | test as the | -Ability of the abdominal | to identify diagnostic studies. |
| radiography in | <u>level</u> : | re-examination | to constipation. | reference | radiography to discriminate | References lists of reviews articles and |
| constipated | 1+ | of abdominal | Some studies | standard) | between clinically constipated | included studies checked for further |
| children: a | | radiographs | included children | | and non constipated children | relevant articles. Experts in the field |
| systematic | Study aim: | | with soiling or | -Faecal loading | (4 studies): | contacted and asked to identify |
| review. 2005. | to evaluate | <u>Inclusion</u> | encopresis, while | on plain | | published and unpublished studies. No |
| Archives of | the | criteria: | others exclude this | abdominal | 1. | language restrictions applied |
| Pediatrics and | additional | Controlled, | group | radiography | Sensitivity: 76 (95% CI: 58 to | |
| Adolescent | diagnostic | observational | | according to a | 89) | Two reviewers independently screened |
| Medicine | value of the | studies | Country: | predefined | Specificity: 75 (95% CI: 63 to | the titles and abstracts f studies |
| 159[7], 671-678 | plain | investigating the | The Netherlands | scoring system | 85) | identified by the searches for eligibility. |
| | abdominal | relationship | | (reference test in | LR: 3.0 (95% CI: 1.6 to 4.3) | All potentially relevant studies were |
| | radiography | between faecal | | 3 studies) | | retrieved as full papers and |
| | in the | loading on plain | | | 2. | independently screened by two |
| | diagnosis of | abdominal | | -Clinical | Sensitivity: 60 (95% CI: 46 to | reviewers. Any disagreements were |
| | constipation | radiography and | | diagnosis of | 72) | resolved through consensus or by |
| | in children | symptoms and | | constipation | Specificity: 43 (95% CI: 18 to | arbitration of a third reviewer |
| | | signs related to | | according | 71) | |
| | | constipation in | | to the presence | LR: 1.0 (95% CI: 0.5 to 1.6) | Methodological quality of studies |
| | | otherwise | | or absence of | | assessed using the QUADAS tool. An |
| | | healthy children | | predefined | 3. | overall methodological quality value |
| | | aged from 1 to | | symptoms and | Sensitivity: 80 (95% CI: 65 to | was assigned to studies by calculating |
| | | 18 years old | | signs (reference | 90) | the number of positive scores |
| | | | | test in 3 studies) | Specificity: 90 (95% CI: 74 to | (maximum value 14). Studies with |
| | | Exclusion | | l | 98) | scores of 9 or higher >60%) were |
| | | criteria: | | In the 6 studies | LR: 8.0 (95% CI: 0.7 to 17.1) | arbitrarily regarded as being of "high" |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------|--|----------------------------|--|--|--|
| Illiormation | level | prevalence | Characteristics | standard | allu NFV | |
| | icvei . | Lack of control group, no data on diagnostic value presented, symptoms of constipation not related to the outcomes of a plain abdominal radiography Setting: all 6 studies hospital based | | included, 3 different scoring systems for assessing impaction on abdominal radiography were used: 3 studies: Barr- score | 84) Specificity: 35 (95% CI: 27 to 44) LR: 1.2 (95% CI: 1.0 to 1.4) -Association between a history of hard stool and faecal impaction on radiography: LR: 1.2 (95% CI, 1.0 to 1.4) -Association between a finding of absence of rebound tenderness and faecal impaction on radiography: LR: 1.1 (95% CI, 1.0 to 1.2) -Association between stool present on rectal examination and faecal impaction on abdominal radiography: LR: 1.6 (95% CI, 1.2 to 2.0) | methodological quality. Two reviewers independently assessed the methodological quality of the independent studies. Any disagreements were resolved by consensus or through consultation with third reviewer. Reviewers scored 84 items and agreed on 65 item (77.4%, k=0.54) Structured data extraction performed independently by two reviewers and any disagreement resolved by consensus Source of funding: Not reported |
| | | | | | abdominal radiography: | |
| | | | | | LR: 1.5 (95% CI, 0.8 to 2.3) | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|--|------------------|
| | | | | | Interobserver reliability: 5 studies: moderate to excellent (k range, 0.63 to 0.95) 1 study: poor to moderate (k=0.28 to 0.060) Intraobserver reliability: Evaluated in 3 studies, ranged from moderate (k=0.52) to excellent (k≥0.85) | |
| | | | | | | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|--------------------------|--------------------------------------|-------------------------------|----------------------------|---------------------------------------|---|
| de Lorijn et al. | level Study type: | prevalence 89 non selected | 89 children | standard Test: | Mean Leech score (using the | Children with clinical characteristics of |
| The Leech | | consecutive | os ciliuleii | | first score): | FAP and FNRFI were classified as the |
| method for | Case control | | Median age: 9.8 | diagnose | <u>llist score).</u> | control group, because according the |
| diagnosing | Case control | Ciliuleii | vears | constipation in | -Group 1 (constipation): 10.1 | authors they have "little or no faecal |
| | Evidence | Inclusion | years | plain abdominal | -Group 2 (controls): 8.5 | loading on an abdominal radiograph" |
| intra- and | level: | criteria: patients | Group 1 | radiography | -Group 2 (controls). 8.3 | loading on an abdominal radiograph |
| interobserver | III | referred for the | (constipation): | radiography | p=0.002 | Treatment with oral/rectal laxatives was |
| variability and | | evaluation of | n=52 (28 boys) | Reference test: | p=0.002 | discontinued in each patient for at least |
| | Study aim: | abdominal pain, | 11=32 (20 bbys) | | Mean CTT: | 4 days. Thereafter the patient ingested |
| Pediatric | to assess | constipation or | Group 2 (controls): | Time (CTT) | -Group 1 (constipation): 92 h | one capsule with 10 small radiograph |
| | intra- and | faecal | N=37 (24 boys) | Tillie (CTT) | -Group 2 (controls): 37 h | opaque markers on 6 consecutive days, |
| | interobserver | | 1N=37 (24 DUy3) | | -Gloup 2 (controls): 37 11 | in order to determine the CTT. |
| 43-49 | variability | Diagnosis of | 31: FNRFI | | p<0.0001 | Subsequently, a plain abdominal |
| | and | | 6: FAP | | | radiograph was taken on day 7. this |
| | determine | least two of the | 0.17.1 | Leech scoring | Diagnostic accuracy of Leech | radiograph was both used in the Leech |
| | diagnostic | following was | | method: | method vs. CTT method: | method and for CTT measurement |
| | | present: | Diagnosis of | Colon divided into | metriod vs. OTT metriod. | inclined and for OTT measurement |
| | the Leech | -defecation | functional non- | three segments: | -Leech method: | Three scorers independently scored the |
| | | frequency less | retentive faecal | right, left and | (cut-off point as per study | same radiography twice (4 weeks apart) |
| | identifying | than 3 | incontinence | recto sigmoid | comparable to 9 as per | using the Leech method, which was |
| | children with | | (FNRFI) based on: | Each segment | literature) | discussed amongst the three scorers |
| | functional | -2/more | 1) two/more faecal | provided with a | Sensitivity: 75% | previous to both readings |
| | | episodes of | incontinence | score from 0-5 | Specificity: 59% | provided to bear readings |
| | conoupation | faecal | episodes/week with | 0:no faeces | Specimenty: 5075 | Scorers were three experienced doctors |
| | | incontinence | no signs of | visible | (cut-off point 9 as per | (a 5 th year radiology resident, a |
| | | per week | constipation 2) | | literature) | paediatric radiologist and a senior |
| | | -production of | defecation | visible | Positive Predictive Value: 72% | paediatric gastroenterologist). No |
| | | large amounts | frequency 3/more | | Negative Predictive Value: | clinical information was about the |
| | | of stool once | times/week 3) no | loading | 63% | patients was made available to them. |
| | | over a period of | periodic passage of | 3: moderate | | panono nao mado aranadio to mom |
| | | 7-30 days | very large amounts | faecal loading | -сст: | A Leech score of 9 or more was |
| | | -presence of | of stool at least | 4: severe faecal | (cut-off point 54h as per study) | considered as suggestive of |
| | | palpable | once during a | loading | Sensitivity: 79% | constipation. |
| | | abdominal or | period of 7-30 days | 5: severe faecal | Specificity: 92% | |
| | | rectal mass | 4) no palpable | loading with | | CTT were assessed once by a single |
| | | | abdominal or rectal | bowel dilatation | (cut-off point 62h as per | scorer. It was assumed that the |
| | | (control children | mass on physical | | literature) | counting of radiopaque markers would |
| | | fulfilled criteria | examination fro a | Colonic transit | Sensitivity: 71% | not lead to intra- or interobserver |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|--------------------------|----------------------|-------------------------------|----------------------------|---------------------------------------|--|
| | level | prevalence | | standard | 0 15 15 0 504 | |
| | | for functional | period of at least 1 | time: | Specificity: 95% | variability |
| | | | week during the | Determined by | Positive Predictive Value: 69% | L. 50/ -f the Leesh |
| | | (FAP) and for | preceding 12 | the method of | Negative Predictive Value: | In 5% of cases the Leech scores of the |
| | | | weeks. Faecal | Bouchoucha. | 97% | same patient produced by different |
| | | | incontinence | Radiography on | B00 I : | scorers could differ by 4 points or more |
| | | incontinence | defined as the | day 7 used to | ROC analysis | |
| | | (FNRFI)) | voluntary/involuntar | count the number | | Source of funding: not stated |
| | | | y loss of loose | of markers in the | -AUC (Leech method): | |
| | | Exclusion | stools in the | | 0.68 (95% CI 0.58-0.80) | |
| | | criteria: not | | markers x 2 | -AUC (CTT method): | |
| | | reported | age of 4 years | produced total | 0.90 (95% CI 0.83-0.96) | |
| | | | Functional | CTT in hours. | | |
| | | | abdominal pain | Localization of | p=0.00015 | |
| | | 0 | (FAP) defined as | | AUC=Area Under the ROC | |
| | | y outpatients | abdominal pain of | calculated | curve | |
| | | clinic | at least 12 weeks | according to | ROC=Receiving Operator | |
| | | | duration 1)that was | previously | Characteristic | |
| | | | continuous or | described | | |
| | | | nearly | formula. Normal | Intraobserver variability (Leech | |
| | | | discontinuous in a | range fro total | score) | |
| | | | school-aged child | transit time based | | |
| | | | or adolescent 2) | on the upper | a. Systematic difference | |
| | | | that had no or only | limits (mean ± | (Mean, 95% CI): | |
| | | | an occasional | 2xSD) from a | -Scorer 1 | |
| | | | relationship with | study in healthy | 0.7 (0.2-1.2) | |
| | | | physiological | children. Based | P=0.89 | |
| | | | events 3) that was | on this study a | | |
| | | | accompanied by | CTT > 62 h was | -Scorer 2 | |
| | | | some loss of daily | considered | 0.03 (-0.4-0.5) | |
| | | | functioning 4) that | delayed. | P=0.0005 | |
| | | | was not feigned | | | |
| | | | and) for which | | -Scorer 3 | |
| | | | there were | | -1.6 (-2.0-1.3) | |
| | | | insufficient criteria | | P<0.0001 | |
| | | | to indicate the | | | |
| | | | presence of | | b. Variability (SD) | |
| | | | another functional | | -Scorer 1: | |
| | | | gastrointestinal | | 2.2 | |

| Bibliographic Study to Information Evidential Study to St | nce patients & | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|--|----------------|--|-------------------------------------|--|------------------|
| | | disorder <u>Country:</u> The Netherlands | | Limits of agreement: -6.0-5.0 -Scorer 2: 2.2 Limits of agreement: -7.0-7.0 -Scorer 3: 1.5 Limits of agreement: -5.0-3.0 Interobserver variability (using the first score): -Scorer 3 vs. scorer 1: Mean of differences 2.7 p<0.0001 -Scorer 3 vs. scorer 2: Mean of differences 2.9 p<0.0001 - Scorer 2 vs. scorer 1: no systematic differences found | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|--|--|--|---|--|---|---|
| van den Bosch et al. Systematic assessment of constipation on plain abdominal radiographs in children. 2006. Pediatric | Ievel Study type: Diagnostic retrospective case series Evidence level: III Study aim: To assess the reproducibilit y of there scoring systems (Barr, Leech and Blethyn) for plain abdominal radiography, in order to determine which one is most useful in clinical practice | prevalence 40 patients Inclusion criteria: consecutive patients referred to hospital for assessment of constipation. Patients complained of | 40 patients Mean age 7 years (range 3-12) 55% boys Country: The Netherlands | standard Test and Reference Standard (all tests compared to each other) -Barr scoring system -Leech scoring system -Blethyn scoring system: Quantifies the amount of faeces in four different bowel segments (ascending colon, transverse colon, descending colon and rectum) and also the consistency of the faces i.e. granular or rocky stools Constipation defined as Barr score>10 Blethyn system: Rough scoring system used to assess amount of faeces in large | Intraobserver variability (k values) -Observer 1: Barr: 0.75 Blethyn: 0.61 Leech: 0.88 -Observer 2: Barr: 0.66 Blethyn: 0.65 Leech: 1.00 Interobserver variability (k values) -Period 1 Barr: 0.45 Blethyn: 0.43 Leech: 0.91 -Period 2 Barr:0.71 Blethyn: 0.31 Leech: 0.84 All k values are statistically significant (p<0.05) Kappa (k) coefficients (level of agreement): <0.20: poor 021-0.40: fair 0.41-0.60: moderate | Masked abdominal radiographs of the children were independently evaluated by two observers, both experienced paediatric radiologists. Observers assessed each radiograph on two separate occasions, 6 weeks apart. Each abdominal radiograph was scored according to the three different scoring systems Intraobserver variability was determined for each scoring system by comparing data from the same observer at two different reading sessions. Interobserver reproducibility was determined by comparing data from the two observers on one occasion. Thus two intraobserver and two interobserver variabilities could be derived for each parameter. Kappa coefficients were calculated as indicators of intra- and interobserver variability. |
| | | | | bowel -Normal, grade 0: | | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|--------------------------|----------------------|-------------------------------|--------------------------------|---------------------------------------|------------------|
| | level | prevalence | | standard | | |
| | | | | faeces in rectum | | |
| | | | | and cecum only | | |
| | | | | -Grade 1, mild | | |
| | | | | constipation: | | |
| | | | | faeces in rectum, | | |
| | | | | cecum and | | |
| | | | | discontinuous | | |
| | | | | elsewhere | | |
| | | | | -Grade 2, | | |
| | | | | moderate | | |
| | | | | constipation: | | |
| | | | | faeces in rectum, | | |
| | | | | cecum with | | |
| | | | | continuous | | |
| | | | | faeces affecting | | |
| | | | | all segments | | |
| | | | | -Grade 3, severe | | |
| | | | | constipation: | | |
| | | | | faeces in rectum | | |
| | | | | and caecum, | | |
| | | | | continuous | | |
| | | | | elsewhere with | | |
| | | | | dilated colon and | | |
| | | | | rectal impaction | | |
| | | | | | | |
| | | | | Leech method: | | |
| | | | | The colon is | | |
| | | | | divided into there | | |
| | | | | segments: | | |
| | | | | 1.ascending and | | |
| | | | | proximal | | |
| | | | | transverse colon | | |
| | | | | 2.distal | | |
| | | | | transverse and | | |
| | | | | descending colon | | |
| | | | | rectosigmoid | | |
| | | | | Amount of faces | | |
| | | | | in each segment | | |

| Bibliographic Study t & Evide leve | nce patients & | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|------------------------------------|----------------|-------------------------------|--|---------------------------------------|------------------|
| | | | scored from 0 to 5. O indicates no faeces and 5 severe faecal loading and bowel dilatation. With a possible score of 0-15, > 8 considered to indicate constipation | | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|----------------------------|---------------------------|---------------------------|-------------------------------|----------------------------|---------------------------------------|--|
| Information | level | prevalence | Characteristics | standard | allu NPV | |
| | Study type: | 133 children | 133 children | Test and | | Authors defined constipation in the past |
| | Diagnostic | | 35 males | <u>Reference</u> | of constipation and faecal load | as "at least 2 weeks of hard, rock-like |
| of constipation | case control | Inclusion | Mean age: 5.6 | <u>Standard</u> | on abdominal X-ray: | stools passed less than 3 times/week |
| with childhood | | criteria: Cases: | years | (not clear which | | without evidence of structural, |
| urinary tract | Evidence | | (range: from | one was what) | Correlation coefficient=0.08 | endocrine or metabolic disease, other |
| infections. | <u>level</u> : III | history of UTIs | newborn to 14 | Ala ala sastra al | | useful association include: abnormally |
| 2005. Journal of | | who were | years) | -Abdominal | | large stools, and difficult or painful |
| Pediatric Urology 1[4], | Study aim: To evaluate | already undergoing a | Group 1 (history of | radiograph (KUB) | | defecation, associated with stools accidents or faecal smearing in |
| 273-278United | the | VCUG(voiding | UTI | -Clinical | | undergarments |
| Kingdom. | relationship | cystourethrogra | | variables: | | undergannents |
| Kingdom. | between a | m), who were | 11-100 | variables. | | Abdominal X-rays reviewed blindly by |
| | history of | | Group 2 (no history | Number of bowel | | three physicians: two paediatric |
| | constipation, | for the | of UTI) | movements/week | | radiologists an one paediatric urologist |
| | faecal | treatment of | n= 33 | | | and score for faecal loading based on a |
| | loading on | constipation | | Stools | | previously validated scoring system |
| | X-rays and a | | Country: | consistency | | (Leech) |
| | history of | Children | USA | | | |
| | UTIs in an | undergoing a | | | | Data collected prospectively on several |
| | office | plain film of the | | | | historical questions about constipation |
| | practice | abdomen for | | | | shortly after the X-ray was performed, |
| | | reasons that did | | | | but before they were reviewed with the |
| | | not include | | | | family. An interviewer filled out the |
| | | constipation/ | | | | history questionnaire using consensus |
| | | UTIs (e.g. renal | | | | of the child's and parents' responses. |
| | | calculi, | | | | Data were also obtained regarding a history of UTI. No data on the |
| | | gastroesophage al reflux) | | | | interviewer are reported |
| | | ai reliux) | | | | interviewer are reported |
| | | Exclusion | | | | Constipation history responses were |
| | | criteria: | | | | scored from 1 to 3 and a total history |
| | | Neurological | | | | score was obtained scored were |
| | | bowel and/or | | | | grouped as: |
| | | bladder | | | | 1-none or mild, 2-moderate, 3-severe |
| | | dysfunction or | | | | |
| | | lower | | | | Data derived from scores on faecal |
| | | gastrointestinal | | | | loading were averaged for each patient |
| | | problems. | | | | and the scores then grouped in the |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|---|
| | | - | | | | same way as previous. Questionnaire not piloted previous to the study As it was thought that children beyond toilet-training age would be more likely to have developed constipation related to overall elimination dysfunction and therefore UTIs as well, the data for children > 3 years were analysed separately |
| | | | | | | |

Diagnostic Value of the Rectal Biopsy in children with Chronic Idiopathic Constipation

| Bibliographic | Study type | Number of | Population | Type of test (s) | Follow-up & Outcome | Reviewer comments |
|------------------------------------|---------------------------|----------------------------------|---------------------------------------|------------------|----------------------------------|--|
| Information | & Evidence level | patients | Characteristics | | Measures Effect Size | |
| Lewis et al. | Study type: | 315 children | 315 children: | Tests: | Clinical features in children | Additional information from study |
| Diagnosing | Retrospectiv | | | Rectal biopsy | with Hirschsprung's disease | Questionnaires, telephone interviews |
| Hirschsprung's | e cohort | <u>Inclusion</u> | -265 children who | | and idiopathic constipation (IC, | and patients visits used to compile long- |
| disease: | | <u>criteria:</u> | hade undergone | | <u>n=40)</u> | term data. In reporting features listed in |
| increasing the | Evidence | -Cohort 1: | rectal biopsy | | | the questionnaire only patients with |
| odds of a | <u>level</u> : | Children | 50 abildese | | -Onset of constipation <1 year | definite information were included: the |
| positive rectal | 2+ | presenting with | -50 children, | | old | number of patients in each analysis |
| biopsy result. 2003. Journal of | Study aim: | constipation to diagnose | concurrent selected cohort (cohort 2) | | Delayed passage of meconium (%) | varies to exclude those with missing data |
| Pediatric | To test the | Hirschsprung's | Conon (conon 2) | | HD: 65 | luata |
| Surgery 38[3], | hypothesis | disease (HD) | Country: | | IC: 13 | Delayed passage of meconium defined |
| 412-416 | that key | (i.2) | USA | | P< 0.05 | as failure to pass meconium in the first |
| | features in | -Cohort 2: | | | | 48h of life. These data were available in |
| | the history, | idiopathic | | | Abdominal distension (%) | 59% of cases |
| | physical | constipation | | | HD: 80 | |
| | examination | | | | IC: 42 | Abdominal distension determined from |
| | and | Exclusion | | | P< 0.05 | parental response to questionnaire or |
| | radiographic | criteria: | | | \\ iti (0/) | data noted during patients visits |
| | evaluation would allow | Patients | | | Vomiting (%) HD: 72 | Enterocolitis defined as diarrhoea |
| | to avoid | undergoing re- evaluation fro | | | IC: 21 | associated with fever |
| | unnecessary | constipation | | | P< 0.05 | associated with level |
| | rectal | after pull- | | | 1 0.00 | Reviewer comments: |
| | biopsies | through | | | Faecal impaction requiring | Data on clinical features not available |
| | | procedure for | | | manual evacuation (%) | for all children |
| | | HD | | | HD: 6 | |
| | | | | | IC: 30 | Unclear what kind of rectal biopsy was |
| | | | | | P< 0.05 | performed and how the diagnosis of HD |
| | | | | | E | was made |
| | | | | | Enterocolitis (%) | Source of fundings |
| | | | | | HD: 13 IC: 15 | Source of funding: Not stated |
| | | | | | NS | Not stated |
| | | | | | | |
| | | | | | -Onset of constipation >1 year | |
| | | | | | old | |
| | | | | | Delayed passage of meconium | |
| | | | | | (%) | |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---------------------------|-----------------------------------|--------------------|-------------------------------|------------------|---|-------------------|
| | | | | | HD: 81 IC: 1 P< 0.05 | |
| | | | | | Abdominal distension (%) HD: 53 IC: 7 P< 0.05 | |
| | | | | | Vomiting (%) HD: 23 IC: 0 P< 0.05 | |
| | | | | | Faecal impaction requiring manual evacuation (%) HD: 46 IC: 30 NS | |
| | | | | | Enterocolitis (%) HD: 13 IC: 14 NS | |
| | | | | | Age at onset of symptoms -Hirschsprung's (HD) (n=46) Mean: 8 months (range 1 day to 9 years) 1rst week of life: 60 % 1rst month of life: 70% 1rst year of life: 87% after 1 year of life: 13% | |
| | | | | | -Idiopathic constipation (IC) (n=40) Mean: 15 months (range 7 days to 16 years) | |

| Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|--------------------|-------------------------------|------------------|--|-------------------|
| | | | 1rst week of life: 15% 1rst month of life: 55% 1rst year of life: 68% after 1 year of life: 32% At least 34% of HD patients had the classic triad (delayed passage of meconium + vomiting + abdominal distension). At least 1 feature of the triad noted in 98% of patients with HD. Only 60% of patients with IC had a history of delayed passage of meconium, vomiting or abdominal distension. 100 % HD patients vs. 64% IC patients had 1 or more of the following: delayed passage of meconium, vomiting, abdominal distension and a transition zone on contrast enema. 36% of IC patients had none of these features. | |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---------------------------|-----------------------------------|--------------------|-------------------------------|---------------------|--|---|
| Pini-Prato et al. | Study type: | 141 patients | 141 patients | Tests: | Clinical variables | Additional information from study |
| Rectal suction | Retrospectiv | - | median age: 20 | -Rectal suction | a. Meconium passage (%) | Total number of biopsies: 1118 |
| biopsy in the | e cohort | <u>Inclusion</u> | months | biopsy | -Failure/delay | performed on 429 patients (mean of 2.6 |
| workup of | | criteria: | mean 44 months ± | | FC (n=45): 7 | each). In 63 patients (14.7%) biopsies |
| childhood | Evidence | Patients with | 67 | -Clinical variables | HD (n=47): 87 | inadequate for a reliable diagnosis |
| chronic | level: | intestinal | | : | IND (49): 22.5 | absence of submucosal layer) 143 |
| constipation: | 2+ | dysganglinonos | Country: | | | patients (33.3%) received a diagnosis |
| indications and | | es (ID) | Italy | a. Meconium | FC vs. HD p<0.001 | of ID. 96/143 fulfilled inclusion criteria, |
| diagnostic | Study aim: | (Hirschsprung's | - | passage | | being 49 IND and 47 HD. 45 |
| value. 2007. | To describe | disease (HD) | | b. Symptoms | -Normal | consecutive patients with a diagnosis of |
| Pediatric | the clinical | and intestinal | | onset | FC (n=45): 93 | FC (out of the remaining 286 patients) |
| Surgery | features of a | neuronal | | c. Intestinal | HD (n=47): 13 | fulfilled inclusion criteria and were |
| International | group | dysplasia (IND)) | | obstruction | IND (49): 77.5 | consequently included, for a total |
| 23[2], 117-122 | patients with | who were | | d. Abdominal | | sample of 141 |
| | intestinal | diagnosed in | | distension | FC vs. HD p<0.001 | |
| | dysganglino | the period | | e. Reported | | Rectal suction biopsies (RSB) |
| | () | between | | enterocolitis | b. Symptoms onset (%) | performed with the instrument Solo- |
| | (Hirschsprun | February 2000 | | | - at < 1year old | RBT ©. Each patient underwent 2 to 4 |
| | g's disease | and July 2005 | | g. Palpable faecal | FC (n=45): 80 | biopsies 2 to 10 cms from the pectinate |
| | (HD) and | | | masses | HD (n=47): 96 | line. Various histochemical staining |
| | intestinal | <u>Exclusion</u> | | h. Soiling | IND (49): 94 | (AChE, LDH, ANE, NADPH-diaphorase |
| | neuronal | criteria: | | | | and Toluidine Blue) were used to |
| | dysplasia | Not stated | | | FC vs. HD p<0.02 | diagnose HD and IND. All biopsies |
| | (IND)) along | | | | | were evaluated by a single, senior and |
| | with a group | | | | - at > 1 year old | experienced pathologist. |
| | of | | | | FC (n=45): 20 | |
| | consecutive | | | | HD (n=47): 4 | HD diagnosed by demonstrating: |
| | patients with | | | | IND (49): 6 | a dramatic increased in AChE- |
| | functional | | | | | positive nerve fibres in the |
| | constipation | | | | FC vs. HD p<0.02 | lamina propia and muscularis |
| | (FC), to | | | | | mucosae |
| | compare | | | | c. Intestinal obstruction (%) | - thick nerve trunks |
| | them and to | | | | FC (n=45): 0 | absent ganglion cells in |
| | find out if the | | | | HD (n=47): 49 | submucosal |
| | clinical | | | | IND (49): 26.5 | |
| | criteria to | | | | | |
| | indicate | | | | FC vs. HD p<0.001 | In case on negative RSB functional |
| | rectal | | | | | constipation diagnosed according to |

| Bibliographic Study Information & Evidence Iev | dence patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---|----------------|-------------------------------|------------------|---|---|
| suction biopsy constip children exist | in pated | | | d. Abdominal distension (%) FC (n=45): 20 HD (n=47): 85 IND (49): 26.5 FC vs. HD p<0.001 e. Reported enterocolitis (%) FC (n=45): 9 HD (n=47): 10.5 IND (49): 20.5 FC vs. HD, NS f. Failure to thrive (%) FC (n=45): 11 HD (n=47): 27.5 IND (49): 22.5 FC vs. HD p<0.045 g. Palpable faecal masses (%) FC (n=45): 22 HD (n=47): 17 IND (49): 20.5 FC vs. HD, NS h. Soiling (%) FC (n=45): 46.5 HD (n=47): 4 IND (49): 4 FC vs. HD p<0.001 | Rome II criteria: At least 2 weeks of: -scybalous, pebble like, hard stools fro a majority of stools -firm stools 2 o less times/week absence of any organic cause of constipation (IND, HD, anorectal malformations, spinal dysraphism, metabolic disorders) Clinical variables retrospectively extracted from patients' notes Reviewer comments: Unclear how the reviewing process was conducted Source of funding: Not stated |

| Bibliographic Information | Study type & Evidence | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures | Reviewer comments |
|---------------------------|-----------------------|--------------------|-------------------------------|-----------------------------------|------------------------------------|---|
| Khan et al. The | Study type: | 182 patients | 182 patients | Tests: | -Total number of patients | Additional information from study |
| constipated | Retrospectiv | 102 pationto | 102 pationto | -Suction rectal | diagnosed with HD: 25 (14%) | Clinical details, laboratory investigations |
| child: how likely | е | Inclusion | 118 males | biopsy (SRB) and | a.a.gcca | and histopathological reports reviewed |
| is | case series | criteria: | | full-thickness | -mean age of patients | retrospectively |
| Hirschsprung's | | Patients who | Mean age 2.9 years | rectal biopsy | diagnosed with HD: 3.64 | , |
| disease? 2003. | Evidence | presented with | (range 2 days to 16 | , , | months (range 2 days to 4 | The Great Ormond Street (GOS) |
| Pediatric | level: | chronic | years) | -Clinical | years) | suction instrument (modified Nobblet) |
| Surgery | 3 | constipation or | , | variables: | , | was used. 2 of 4 specimens were |
| International | | intestinal | Country: | | Clinical symptoms in children | obtained at 2, 3 and 4 cm above the |
| 19[6], 439-442 | Study aim: | obstruction and | UK | a. Meconium | with HD (number of children): | dentate line, in the ward or theatre |
| | To review | had rectal | | passage | | without anaesthesias. All suction biopsy |
| | author's | biopsy to | | b. Constipation | Meconium passed> 48 h: | specimens were examined by routine |
| | experience | exclude HD in | | since birth | -In total sample: | fixation with HE staining and AChE |
| | of rectal | the University | | c. Intestinal | < 1 year old: 35 | histochemistry. All full thickness |
| | biopsy to | Hospital of | | obstruction | >1 year old: 6 | biopsies were done under general |
| | exclude | Wales, Cardiff | | d. Failure to thrive | | anaesthesia and examined by routine |
| | Hirschsprun | | | e. Chronic | % of clinical feature to HD: 39 | fixation with HE staining. The |
| | g's disease | <u>Exclusion</u> | | abdominal | | histochemical criteria used for the |
| | (HD) by | criteria: | | distension | Meconium passed< 24 h: | diagnosis of HD were those of Meier- |
| | haematoxyli | Not stated | | | -In total sample: | Ruge in 1972 i.e. the combination of an |
| | n-eosin (HE) | | | | < 1 year old: 40 | absence of submucosal ganglion cells |
| | staining and | | | | >1 year old: 74 | and an increased AChE activity with |
| | acetylcholine | | | | -In HD children: 6 | parasympathetic fibres of the |
| | sterase | | | | % of clinical feature to HD: 5 | muscularis mucosae and lamina propia |
| | (AChE) | | | | | mucosae. At least 60 sections were |
| | stains, and | | | | Passage of meconium | examined from each block to find the |
| | author's | | | | unknown: | submucosal ganglion cells |
| | clinical | | | | -In total sample: | |
| | criteria to | | | | < 1 year old: 29 | Suction biopsy accepted as adequate |
| | perform | | | | >1 year old: 17 | even if only 1 out of 2 to 4 specimens |
| | rectal biopsy | | | | -In HD children: 3 | contained mucosa and sub-mucosa |
| | in these | | | | % of clinical feature to HD: 11 | |
| | children | | | | On and in a diam of a large bloods | 182 patents who had rectal biopsies |
| | | | | | Constipation since birth: | provided355 specimens in which 79% |
| | | | | | -In total sample: | of suctions biopsies and 97% of full- |
| | | | | | < 1 year old: 33 | thickness biopsies were adequate. |
| | | | | | >1 year old: 20 | Adequate biopsies include rectal |
| | | | | | -In HD children: 17 | mucosa and submucosal according to |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---------------------------|-----------------------------------|--------------------|-------------------------------|------------------|--|---|
| | | patients | | | | Noblett. In 20 children with HD the diagnosis was made at the first attempt by suction rectal biopsy. Repeat biopsies performed on 14 (8%) of 182 patients because of inadequate initial biopsy, clarification of atypical inervation and confirmation of false negative results. 19/104 patients who underwent SRB were > 1 year old. Because 5 children (12 specimens) who were older than 1 year had inadequate suction biopsies at beginning of series, it was decided that SRB was not suitable fro children >1 year old. 3 patients with HD (aged 6 days, 12 days and 6 weeks) has false negative AChE staining. In these the diagnosis were later established from repeated biopsies: 1 full thickness biopsy, 1 laparotomy and 1 suction biopsy Reviewer comments: Unclear how the reviewing process was conducted No definition of constipation or other clinical symptoms given Authors explained that patients may have had more than one symptom, but these figures were not reported in the paper Source of funding: |
| | | | | | | Not stated |

| | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|---|---|--|--|---|---|--|
| Rectal biopsy in the investigation of constipation. 1998. Archives of Disease in Childhood 79[3], 266-268 St Tc cr wc re cc id. ch Hi g': (H th av tra ex ur re bic. | vidence evel: tudy aim: o develop riteria that rould eliably and onsistently dentify hildren with lirschsprun 's disease HD) and | Inclusion criteria: All children who had rectal biopsy to exclude Hirschsprung's disease between January 1, 1993 and December 31, 1995 at Southampton General Hospital Exclusion criteria: Not stated | 141 children age at biopsy: 1 day to 13 years gender not reported Country: UK | biopsy in children younger than 1 year Open transanal rectal biopsy under general anaesthesia performed at least 1cm above pectinate line, in older children or following repeated failure | history of delayed passage of meconium (>48h after birth): 10 (58.8%) age of onset of constipation: all 17 children: < 4 weeks bleeding per rectum: 0 anal fissures:0 sever behavioural/emotional problems: 0 soiling: 0 enterocolitis: 8 (47%) -No Hirschsprung's (n=124) age at biopsy: 1 day to 13 years 20 children: < 4 weeks 12 children: 4 to 12 weeks | Additional information from study Histological diagnosis usually made on haematoxylin and eosin staining with at least 100 serial sections looked at in detail. Acetylcholinesterase used occasionally but not as the main method of diagnosis Constipation defined as a decreased frequency of bowel movement s(<3/week), or a difficulty in defection which is perceived by the parents as a problem, requiring medication (oral or rectal) or manual intervention by the parents. This included anal stimulation with cotton bud, holding the buttocks apart and manual evacuation History of onset of constipation was available in 136 of the 141 children (96%). The 5 children in whom this history could not be obtained from the notes were all older than 1 year (3 teenagers) and none had HD A total of 186 biopsies performed, with 22% failures. (Suction: total 74, 35% failures; Open: total 100, 14% failures, operative total 12, no failures) Reviewer comments: Unclear how the reviewing process was conducted Source of funding: Not stated |

| Bibliographic Information | Study type & Evidence level | Number of patients | Population Characteristics | Type of test (s) | Follow-up & Outcome Measures Effect Size | Reviewer comments |
|------------------------------|-----------------------------------|--------------------|-------------------------------|------------------|---|-------------------|
| | | | | | age of onset of constipation: | |
| | | | | | 40 children: < 4 weeks | |
| | | | | | 32 children: 4 to 12 weeks 22 children: 12 weeks to 1 year | |
| | | | | | 25 children: > 1 year | |
| | | | | | 25 Children. > 1 year | |
| | | | | | bleeding per rectum: 37 (30%) | |
| | | | | | anal fissures: 14 (11%) | |
| | | | | | sever behavioural/emotional | |
| | | | | | problems: 10 (8%) | |
| | | | | | soiling: 16 (13%) | |
| | | | | | enterocolitis: 0 | |
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Diagnostic Value of the Abdominal Ultrasound in Children with Chronic Idiopathic Constipation

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------------------|--|--|---|---|---|--|
| tool for | Study type: Diagnostic. Case control Evidence level: | 49 patients Inclusion criteria: Positive diagnosis of | 49 patients aged between 5-13 years Group 1: 23 patient s with | Test: lower abdominal ultrasound of rectum Reference | Rectal diameter (cm) (Mean, standard deviation, 95% CI) -Group 1 (constipated, n=23): | Ultrasound done with the patient supine. 7.5 MHz probe applied on abdominal skin approximately 2cm above the symphysis. Measurement performed with moderate (30-70 % capacity of for age) filled bladder at an |
| | Study aim: to prove the accuracy of | constipation, made by patient history and physical examination | positive history of voiding dysfunction and constipation Group 2: | Standard : None reported | 4.9 (1.01; 4.4 to 5.3) -Group 2 (control, n=26) 2.1 (0.64; 1.8 to 2.4) | angle of about 15 degrees downward from the transverse plane. The diameter of the rectum, behind the bladder was measured twice. |
| Urology 172[5 Pt 1], 1986- 1988 | the transverse diameter of the rectum on ultrasonogra | when the patient had at least 2 positive signs, including: -2 or fewer bowel | 26 urological patients without lower urinary tract dysfunction and a normal defecation pattern, diagnosed | | p<0.001 | If stools had been passed in the last two hours or patients had an urge to defecate during the investigation the were not included in the study, but this situation did not occur |
| | phy as an additional parameter for diagnosing constipation | | with undescended testicle, periodic control for upper urinary tract dilatation, etc. | | | In all patients it was possible to obtain a reliable and repeatable measurement of the rectum if at least some bladder filling was present It was not reported who performed the |
| | in children with lower urinary tract dysfunction | weekly -periodic passage of a large amount of | Country: UK | | | ultrasound, or whether this person was blinded No significant difference in age between |
| | | stool once every 7 to 30 days -palpable abdominal and/or rectal | | | | the two groups (p=0.20) or in period between the last time a stool was passed prior to the rectal measurement (p=0.16) In all patients with voiding dysfunction |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|--|-------------------------------|---|---------------------------------------|--|
| | | mass Exclusion criteria: laxative therapy, constipation due to neurological disease, disease of the gastrointestinal tract based on endocrinological , metabolic, genetic or toxic disease, or connective tissue disease Setting: hospital | | | | and faecal constipation (Group 1) rectal examination confirmed stool in the rectum, but there are no data reported on this variable for the control group, probably for ethical reasons Source of funding: Not stated |

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|-----------------------------------|-----------------------|----------------------------|------------------------------------|-------------------|--------------------------------|--|
| Information | & Evidence | patients & | Characteristics | Reference | and NPV | |
| | level | prevalence | | standard | | |
| Singh et al. Use | | 177 children | 177 children | Test: | Median rectal crescent (cm) | A portable US machine with a 5-MHz |
| | Diagnostic. | | | Pelvic ultrasound | | probe (falcon 2101 Ultrasound scanner |
| ultrasound in | Case control | | Group 1: | D () | Group 1 (healthy children): | with a transducer type 8803 [3.0-5.0 |
| the diagnosis of | - · · | criteria: | 82 children (median | | 0.4./ | MHz], B-K Medical, Copenhagen, |
| | <u>Evidence</u> | | age 5.5 years, | none reported | 2.4 (range 1.3 to 4.2; IQR | Denmark) was used. |
| children with | <u>level</u> : III | referred after | range 0.30-15.30) | | 0.72) | |
| constipation. 2005. Journal of | | failing to | with no history of | | Group 2 (children with | The same individual performed all the US scans, but not other data on this |
| | Study aim: | respond to medical | constipation or other anorectal or | | constipation): | were reported (as blinding, individual's |
| | to establish | | gastrointestinal | | <u>constipation).</u> | experience in radiology, etc) |
| | normal | | problems and no | | 3.4 (range 2.10 to 7.0; IQR | experience in radiology, etc) |
| | values for | constipation | previous anorectal | | 1.0) | All children had a full or partially full |
| | the rectal | • | surgery | | 1.0) | bladder at the time of measurement. In |
| | crescent in | child had 2 or | ou.gory | | p<0.001 | cases where the child was initially |
| | healthy | more of the | Group 2: | | F 101001 | scanned and the bladder was noted to |
| | children, | following: | 95 children (median | | IQR= interquartile range | be empty, the US was abandoned and |
| | compare | -less than 3 | age 6.5 years, | | | the child was offered liberal fluids orally. |
| | them with | bowel | range 0.40-16.40) | | Receiver operating | The scan was repeated within an hour |
| | the rectal | movements/we | with a history of | | characteristic analysis: | and in all cases, by then, the child had |
| | crescent in | ek | constipation of at | | | a full or partially full bladder |
| | children with | -periodic | least 6 months | | -Area under the curve: | |
| | constipation | | duration, referred to | | 0.847 | The US probe was applied on the |
| | and explore | | a tertiary referral | | 95% CI: 0.791 to 0.904 | anterior abdominal wall in the midline, |
| | whether | discomfort or | centre | | | approximately 1-2 cm above the |
| | pelvic | pain | _ | | | symphysis at a 90 degrees angle to the |
| | ultrasound | -a palpable | Country: | | Cut-off point for establishing | abdominal wall. This showed the |
| | can hep in | abdominal | UK | | the diagnosis of megarectum: | impression of the rectum behind the |
| | establishing | mass on | | | | urinary bladder as a crescent which |
| | a diagnosis | physical | | | 3.0 cm | was measured in centimetres |
| | of | examination | | | | The same series and sittle series and series are series and series and series and series and series are series and series and series and series are series and series and series and series are series are series and series are seri |
| | megarectum | -faecal soiling in | | | | There were no significant differences |
| | | the presence of any of the | | | | between the two groups in terms of age, weight and height (p values 0.114, |
| | | any or the above | | | | 0.198 and 0.131 respectively) |
| | | above | | | | 0.130 and 0.131 respectively) |
| | | Exclusion | | | | Results were adjusted for confounders |
| | | criteria: | | | | (age, height and weight) |
| | | Previous | | | | |

| Bibliographic Study Information & Evid | lence patients & | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|--|--|-------------------------------|-------------------------------------|---------------------------------------|---|
| | anorectal surgery (e.g. pull-through procedures for Hirschsprung's disease or anorectal myectomy) Setting: tertiary referral centre | | Standard | | Age and rectal diameter were significantly related (p<0.0001): the older the child the bigger the rectal diameter Time to last evacuation was not ascertained and authors acknowledged this may influence the size of the rectal crescent Source of funding: not stated |

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|-------------------------------|-----------------------|------------------------|---|--------------------------------------|-----------------------------------|---|
| Information | & Evidence | patients & | Characteristics | Reference | and NPV | |
| Diine et el The | level | prevalence | 005 | standard | Diameters of sectal assessible by | 110 |
| Bijos et al. The | Study type: | 225 children | 225 children | Test: | Diameters of rectal ampulla by | US assessment of stool retention and |
| usefulness of | Diagnostic | la alcaia a | Craum 4. | Abdominal | US (mm, mean ± SD) | colonic enlargement involved |
| ultrasound | Case control | Inclusion criteria: | Group 1: 120 children with | ultrasound | Age (years) | measurement of the transverse |
| examination of the bowel as a | Evidonos | | | Deference tests | Croup 1 (constinuted): | diameter of the rectal ampulla (by US) |
| method of | Evidence level: | because of | chronic constipation (72 boys, mean age | Reference tests: Proctoscopy (for | -Group 1 (constipated): All ages: | and pelvic width (externally using a measuring tape) Pelvic width was |
| assessment of | <u>lever</u> . III | chronic | 6.25 years, range | diagnosing faecal | 43. 06 ± 9.68 (range 30 to 82) | defined as the distance between the |
| functional | 111 | constipation, | 1.6 to 17.9) | 0 0 | 45. 06 ± 9.66 (range 50 to 62) | external margins of the anterior |
| chronic | Study aim: | based on | 1.0 (0 17.9) | impaction) | ≤3: 38.35 ± 8.65 | superior iliac spines. The ratio between |
| | to determine | history and | Group 2: | | 3.1 to 6: 41.16 ± 8.72 | the transverse diameter of the rectal |
| children, 2007. | whether a | physical | 105 children with | | 6.1 to 12: 46.15 ± 9.56 | ampulla and transverse diameter of the |
| Pediatric | new method | examination: | normal defecation | | >12 years: 49.09 ± 10.19 | pelvis was calculated to give the |
| Radiology | of ultrasound | | pattern (mean age | | 712 years. 49.03 ± 10.19 | rectopelvic ratio. |
| 37[12], 1247- | (US) | | 8.25 years) | | -Group 2 (control): | Totopolvio ratio. |
| 1252 | assessment | persisting | 0.20 youro, | Transit times | All ages: | US was performed using a Philips HDI |
| 1202 | of stool | longer than 6 | Country: | (hours, upper limit | | 4000 US unit (Philips, Best, The |
| | retention | months, all | Poland | of 66 based on | on so is in the second | Netherlands) equipped with three |
| | could be | patients fulfilled | | literature) | ≤3: 27.07 ± 8.00 | electronic transducers with various |
| | used as a | Rome II criteria | | | 3.1 to 6: 29.25 ± 6.86 | frequencies from 2-14 MHz. children |
| | method of | for defecation | | ≤66: normal- | 6.1 to 12: 32.85 ± 8.73 | were examined before food and had a |
| | identifying | disorders | | transit | >12 years: 35.15 ± 7.18 | slightly filled bladder. Patients who |
| | children with | (frequency of | | constipation | | passed stool on the day of the |
| | functional | bowel | | | p<0.001 for every age group | examination were temporarily excluded |
| | chronic | movements less | | 66-100: slow- | | from the study until they became |
| | constipation, | than twice a | | transit | Mean rectopelvic ratios for all | constipated again. |
| | and to | week, | | constipation | ages (mean ± SD) | |
| | determine | consistency and | | | (Cut-off value to diagnose | Rectal ampulla width was measured |
| | whether | size of stool | | >100: very | megarectum: 0.189) | with the probe applied to the anterior |
| | children with | caused pain | | delayed slow- | | abdomen above the symphysis. |
| | an enlarged | during | | transit | -Group 1 (constipated): | Measurement was performed on |
| | rectum and | defecation, | | constipation | All ages: | oblique transaxial scanning plane to |
| | colon (as | withholding | | | 0. 22 ± 0.05 | obtain transverse diameter of the |
| | seen on US) | behaviour) | | | | ampulla. Measurement was taken |
| | should be | | | | ≤3: 0.24 ± 0.060 | several times and the highest one |
| | referred for | Exclusion | | | 3.1 to 6: 0.23 ± 0.05 | recorded taken as the final |
| | further | criteria: | | | 6.1 to 12: 0.22 ± 0.05 | measurement |
| | procedures | anatomic | | | >12 years: 0.19 ± 0.04 | Total and as supportal aslamic type: -!! |
| | such as | abnormality | | | | Total and segmental colonic transit |

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|---------------|---------------|-----------------------|-----------------|------------------|------------------------------------|---|
| Information | & Evidence | patients & | Characteristics | Reference | and NPV | |
| | level | prevalence | | standard | | |
| | proctoscopy | (Hirschsprung's | | | -Group 2 (control): | time measured by the modified sixth |
| | and | disease, | | | All ages: | day Hinton method. Total and |
| | assessment | congenital | | | 0.15 ± 0.04 | segmental time obtained by multiplying |
| | of colonic | abnormalities of | | | | the number of radiopaque markers |
| | transit time. | the anorectal | | | ≤3: 0.17 ± 0.05 | seen on the radiograph by 1.2 (time in |
| | | region) | | | 3.1 to 6: 0.16 ± 0.04 | hours/number of markers swallowed by |
| | | neurological | | | 1 to 12: 0.15 ± 0.05 | the patient) |
| | | and psychiatric | | | >12 years: 0.14 ± 0.03 | |
| | | conditions | | | | The same individual performed all the |
| | | (cerebral palsy, | | | p<0.001 for age groups (years): | US scans, but not other data on this |
| | | spina bifida, | | | ≤3; | were reported (as blinding, individual's |
| | | mental | | | 3.1 to 6; 6.1 to 12 | experience in radiology, etc) |
| | | retardation, | | | p=0.002 for >12 years | |
| | | anorexia | | | | It is not clear what number of children |
| | | nervosa) | | | US vs. proctoscopy in the | underwent each of the tests |
| | | ,metabolic | | | diagnosis of faecal impaction | |
| | | conditions | | | 0 ''' '' 00 00' | It is not clear how the authors |
| | | (diabetes | | | -Sensitivity: 88.3% | calculated the sensitivity of the US vs., |
| | | mellitus/insipidu | | | | proctoscopy to diagnose faecal |
| | | s) endocrine | | | Mean colonic transit times: | impaction, as the results of |
| | | disorders | | | Children with faecal impaction | proctoscopy are not reported |
| | | (hypothyroidism | | | (as per US) had significantly | 1. 1969 |
| | |), previous | | | longer average segmental | It is difficult to know exactly how many |
| | | thoracic or | | | transit time for the rectum, | children were diagnosed with faecal |
| | | abdominal | | | sigmoid and left colon | impaction by US, as these data are |
| | | surgery | | | (p<0.001, p=0.0015 and | reported only in the form of a bar |
| | | (| | | p=0.0104 respectively) there | graph. Data on number of children |
| | | (control | | | was not statistically significant | diagnosed with "overfilled colon" are |
| | | patients: normal | | | difference for the right side of | not reported at all. |
| | | defecation | | | the colon. Children with an | It is not also rubother "externed" |
| | | patterns, | | | I | It is not clear whether "enlarged" and |
| | | treated for | | | had a significantly longer transit | |
| | | various | | | | the authors, as no measurements of |
| | | symptoms like chronic | | | (p=0.0029) | "enlarged" colon are reported. |
| | | abdominal pain, | | | Definitions of: | Children apparently underwent DRE |
| | | food allergies) | | | Deminions or. | but no results are reported |
| | | ioou allergies) | | | -Faecal impaction (as per US in | |
| | 1 | | | 1 | r accai impaction (as per OS III | |

| Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|-----------------------------------|--|-------------------------------|-------------------------------------|---|------------------|
| | Setting: gastroenterolog y outpatient clinic | | | sagital plane): when pelvic structures were covered by stool masses and were not even partially visible. -Overfilled colon (as per US): Overfilled bowel at the splenic flexure: when it was impossible to visualise the entire length of the left kidney due to the lack of visibility of the lower pole of the kidney because of bowel contents. Probe applied to the long axis of the spleen. Overfilling of the transverse colon: when the superior mesenteric artery was not visible with the probe applied in the sagital plane over the aorta | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------|------------------------------|-------------------------------|----------------------------|---------------------------------------|--|
| imormation | level | prevalence | Onaracteristics | standard | and H | |
| | Study type: | 51 children | 51 children, aged 4- | Test: | Rectal diameter (mm) (mean ± | For transabdominal measurements of |
| | Diagnostic. | | 12 years | Transabdominal | 2SD) | rectal diameter: a 7.5 MHz probe |
| ultrasound of | Case control | | | ultrasound of | | applied to the abdomen approximately |
| rectum as a | | Children | Group 1: | rectum | -Children with rectal impaction | 2cm above the symphysis at 10 to15- |
| | <u>Evidence</u> | referred to | 27 children (mean | | as per DRE (n=22, 20 | degree downward angle. Diameter of |
| in childhood | <u>level</u> : | outpatient clinic | age 7.0±1.8 years) | Reference test: | constipated, 2 healthy): | the rectum measured in traverse plane. |
| constipation. | III | with either | diagnosed with | Digital rectal | | At each session (n=3) diameters were |
| 2008. Journal of | | constipation or | chronic constipation | | 40.5 ± 7.9 | measured three times and mean value |
| | Study aim: | faecal | by Rome III criteria | (DRE) | | was calculated. All children had a |
| 1997-2002 | To look into | incontinence, | _ | | -Children without rectal | partially full bladder range (28 to 450 |
| | a possible | with or without | Group 2: | | impaction as per DRE (n=26, 7 | ml) corresponding to 20-155% of |
| | correlation | urinary | 24 healthy children | | constipated, 19 healthy): | expected bladder capacity for age at |
| | between a | incontinence | (mean age 9.1±2.7 | | | the time of the measurement. In case |
| | dilated | | years) | | 21.0 ± 4.2 | of empty bladder fluid was offered |
| | rectum | UTI. Patients | | | | orally and scanning was repeated. If |
| | | fulfilled Rome III | | | p<0.001 | the child had a bowel movement within |
| | ultrasound | criteria, had at | the Netherlands | | | 3 hours before the investigation or had |
| | and a faecal | least 2 of the | | | Cut-off value for the presence | an urge to defecate, the result was |
| | mass | following | | | of rectal impaction (average | excluded. All investigations were |
| | detected by | characteristics: | | | rectal diameter of children | performed by the same observer (a |
| | digital rectal | -fewer than 3 | | | without impaction plus 2SD): | paediatric intern, who had no prior |
| | examination. | | | | | radiological experience) This observer |
| | To evaluate | movements/we | | | 29.4 mm | was not reported blinded to the study |
| | whether this | ek | | | | objectives and patient's characteristics |
| | method | -more than 1 | | | Rectal diameter (mm) (mean ± | |
| | could | episode of | | | <u>2SD)</u> | There was no significant difference in |
| | diagnose | faecal | | | Defense two etmos into | height and weight distribution between |
| | constipation | incontinence | | | Before treatment: | the 2 groups, but the healthy children |
| | according to | weekly | | | -Group 1 (Constipated, n=27): | were significantly older than the |
| | Rome III | -large stools in | | | 20.0 . 0.0 | constipated children |
| | criteria | rectum by DRE or palpable on | | | 39.6 ± 8.2 | Constipated children received 3 days of |
| | | abdominal | | | -Group 2 (Healthy): | disimpaction followed by 4 weeks of |
| | | palpation | | | -Group 2 (Fleatiffy). | laxative treatment with polyethylene |
| | | -occasional | | | 21.4 ± 6.00 | glycol and behavioural therapy. No |
| | | passage of | | | Z1.7 ± 0.00 | other details reported |
| | | large stools | | | p<0.001 | other details reported |
| | | -display of | | | | No significant correlation between |
| | | -uispiay Ui | | | | TWO SIGNINGATIL CONTENATION DELIVERN |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---|--|
| | 10701 | retentive | | Staridard | After treatment | bladder volume at the time of |
| | | posturing and | | | -Group 1 (Constipated, | measurement and rectal diameter |
| | | withholding | | | responded to treatment, n=15): | (r=0.04) |
| | | behaviour | | | | |
| | | painful | | | 26.9 ± 5.6 | There are missing data not accounted |
| | | defecation | | | | for |
| | | | | | p<0.01 (as compared to same | |
| | | (healthy control | | | group before) | Apparently healthy children diagnosed |
| | | children were | | | p<0.05 (as compared to group | with faecal impaction did not receive |
| | | recruited form | | | 2) | any laxative treatment, which is |
| | | employees of | | | | worrying from an ethical point of view |
| | | the Paediatrics | | | 11 children did not respond to | |
| | | Department at | | | treatment and no significant | Authors acknowledged the abdominal |
| | | the hospital) | | | differences were observed in | ultrasound technique might bear |
| | | F l i | | | their rectal diameter as | technical limitations related to artefacts |
| | | Exclusion | | | compared to pre-treatment | like: acoustic enhancement, speed |
| | | criteria: known | | | Intro ob o or you you in hility | error, and refraction artefacts although |
| | | organic causes of constipation, | | | Intraobserver variability: -coefficient of variation of the 3 | their possible influence on their results is unclear |
| | | including | | | consecutive measurements: | is unclear |
| | | Hirschsprung's | | | consecutive measurements. | No correlation was found between the |
| | | disease, spinal | | | 5.8% ± 4.3% | rectal diameter and age or sex of the |
| | | and anal | | | 3.0 70 ± 4.3 70 | children in either group |
| | | congenital | | | 7 of the constipated children | Crimarer in entiler group |
| | | abnormalities, | | | (26%) had a rectal diameter | Source of funding: |
| | | previous | | | smaller than the established | Supported by Karen Elise Jensen |
| | | surgery on the | | | cut-off point for rectal | Foundation |
| | | colon, | | | impaction, despite the fact that | |
| | | inflammatory | | | they fulfilled the Rome III | |
| | | bowel disease, | | | criteria for constipation. 2 | |
| | | allergy, | | | healthy children with rectal | |
| | | metabolic and | | | impaction had a markedly | |
| | | endocrine | | | larger rectal diameter (38 and | |
| | | diseases, | | | 31 mm) than the other healthy | |
| | | children | | | controls. | |
| | | receiving drugs | | | | |
| | | know to affect | | | | |
| | | bowel function | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---|-------------------------------|---|---------------------------------------|------------------|
| | | during a 2-mont period before initiation (not specified which) | | | | |
| | | Setting: outpatient clinic | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|----------------------------|----------------------|-------------------------------|----------------------------|---------------------------------------|--|
| | level | prevalence | | standard | | |
| | Study type: | 500 children | 500 children | Test: | Correlation between SSS and | Additional information from study |
| nan et al. A new | | | | Pelvic ultrasound | <u>US score</u> | -US scoring sheet (this score can be |
| ultrasound | prospective | Inclusion: | 317 male | | | used even with an empty bladder) |
| J , | case series | All children, | | Both transverse | -first visit (n=500) | |
| for assessing | | both new | median age: 8 | and longitudinal | | Stool height (x): (bladder effect (y)): |
| the severity of | Evidence | referrals and | years (age range 8 | planes | Mean SSS: 23.5 (SD 11.6) | |
| constipation in | <u>level</u> : | follow-up, | months to 18 years) | | 100.00 | No stool: 1 (empty bladder: 0 |
| | III | attending a | 0 | | Mean US total score: 4.02 (SD | Retro bladder: 2 (n compression: 0) |
| Pediatric | O. 1 . | constipation | Country: | same clinician | 2.8) | Just above bladder: 3 |
| Surgery | Study aim: To asses the | outpatient clinic | UK | after very brief | Pearson's correlation: 0.39 | Nearly umbilicus: 4 (indented bladder: |
| International | correlation | Exclusion | | training | P<0.001 | To umbilicus: 5 (Flattened bladder: 2) |
| 24[12], 1379- 1384 | between | criteria: | | | P<0.001 | Beyond umbilicus: 6 (displaced |
| 1304 | severity of | Children not | | Reference test: | -second visit (n=226) | bladder: 3) |
| | constipation | compliant to | | Clinical | -second visit (n=220) | Can't see upper edge: 7 |
| | and | have | | assessment: | Mean SSS: 19.9 (SD 12.6) | Uncooperative: 99 |
| | ultrasound | assessment | | assessificit. | Wear 000. 19.9 (0D 12.0) | Not available: 0 |
| | (US) | done by US, | | Standard | Mean US total score: 3.49 (SD | TVOT available. 0 |
| | findings, the | cases when the | | symptoms | 2.6) | total =x+y |
| | correlation | US machine | | severity scoring | | total =x+y |
| | between | was not | | sheet (SSS), | Pearson's correlation: 0.49 | -Symptom severity scoring sheet: |
| | clinical | available | | completed by | P<0.001 | January Committee of the Committee of th |
| | examination | | | parent or child if | | Filled in by parent, or child if old |
| | and US | Setting: | | old enough | -third visit (n=62) | enough. |
| | findings and | Constipation | | | , , | Q1 About the soiling problem (faecal |
| | the | outpatient clinic | | Clinical | Mean SSS: 23.02 (SD 13.7) | incontinence/mess in underclothes): |
| | correlation | | | assessment done | | - none (0) |
| | between | | | by detailed | Mean US total score: 3.66 (SD | |
| | findings at | | | history taking and | 2.6) | - occasionally (2) |
| | serial out- | | | abdominal | | - only is bowel loaded (5) |
| | patient | | | examination | Pearson's correlation: 0.26 | - continuous day only (8) |
| | follow-up | | | | P=0.04 | - continuous day and night (10) |
| | visits to | | | | | |
| | assess | | | | -fourth visit (n=12) | Q2 About the delay from passing one |
| | clinical | | | | | complete stool to the next: |
| | improvement | | | | Mean SSS: 28.5 (SD 16.8) | - daily stool (0) |
| | s and US | | | | | - every 2 or 3 days (1) |
| | findings | | | | Mean US total score: 4.9 (SD | - every 3-5 days (2) |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|---------------------------------------|---|
| | | | | | 3.2) | - every 5-10 days (5), |
| | | | | | | - greater than 10 (8) |
| | | | | | Pearson's correlation: 0.70 P=0.01 | - never (10) |
| | | | | | | Q3 About pain and difficulty with |
| | | | | | Pearson's correlation | passing stools: |
| | | | | | between US score and clinical | - none (0) |
| | | | | | examination of palpable faeces | - occasionally (1) |
| | | | | | <u>per abdomen</u> | - often (2) |
| | | | | | | - with most stools (4) |
| | | | | | -first visit (n=500) | - with every stool (5) |
| | | | | | Mean palpable faeces score: | Q4 About the amount and types of |
| | | | | | 1.42 (SD 1.6) | medicine needed regularly over the last month: |
| | | | | | Mean US total score: 4.02 (SD | - none (0) |
| | | | | | 2.8) | - softeners only e.g.: lactulose or Docusate or daily Movicol or methyl |
| | | | | | Pearson's correlation: 0.89 | cellulose (1) |
| | | | | | P<0.001 | - softeners and daily stimulants e.g.: |
| | | | | | 1 40.001 | Senokot or picosulphate (2) |
| | | | | | -second visit (n=226) | - softeners and daily stimulants and |
| | | | | | | weekend extra picosulphate or Movicol |
| | | | | | Mean palpable faeces score: | (4) |
| | | | | | 1.10 (SD 1.6) | - medicines as well as extra weekend klenprep or high dose Movicol (8) |
| | | | | | Mean US total score: 3.49 (SD | - medicines as well as regular enemas |
| | | | | | 2.6) | or suppositories (10) |
| | | | | | Pearson's correlation: 0.845 | Q5 About how your child's general |
| | | | | | P<0.001 | health has been affected by the bowel |
| | | | | | -third visit (n=62) | problem over the last month: - well (0) |
| | | | | | -tilita visit (11=02) | - well (0) - occasionally ill (2) |
| | | | | | Mean palpable faeces score: | - often ill (3) |
| | | | | | 1.10 (SD 1.6) | - ill most days (4) |
| | | | | | 1.10 (35 1.0) | - never well (5) |
| | | | | | Mean US total score: 3.66 (SD | (5) |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|--|---|
| | | | Citatacteristics | standard | 2.6) Pearson's correlation: 0.77 P<0.001 -fourth visit (n=12) Mean palpable faeces score: 1.92 (SD 1.7) Mean US total score: 4.9 (3.2) Pearson's correlation: 0.91 P<0.001 | Q6 About behavior related to the bowel problem: - cooperative OK (0) - needs reminding to use the lavatory/pot (2) - refuses the lavatory or pot (3) - also refuses medicines (4) - also generally difficult behavior (5) Q7 overall, which best describes how the problems are now compared with the last time seen at hospital: - nearly completely OK (0) - much better (1) - some improvement (4) - still as difficult (8) - getting worse (12) Filled in by practitioner Amount of stool detected on clinical examination of abdomen score: - None palpable: 0 - Little: 1 - Suprapubic only: 2 - To umbilicus: 3 - Beyond umbilicus: 5 - Reaching ribs: 8 Reviewers comments No control/comparison group Very small sample size at the fourth visit |
| | | | | | | Source of funding: Not stated |

Diagnostic Value of Transit Studies in Children with Chronic Idiopathic Constipation

Radiopaque Markers

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|------------------|--------------------|------------------|-----------------|---------------------|---------------------------------|--|
| Information | & Evidence | patients & | Characteristics | Reference | and NPV | |
| | level | prevalence | | standard | | |
| de Lorijn et al. | Study type: | 169 consecutive | 169 consecutive | Test: | Total and segmental transit | Additional information from study: |
| Prognosis of | Diagnostic | patients | patients | Colonic transit | times (hours), (median, 25 | Significant baseline differences |
| constipation: | prospective | | 65% boys | time (CTT) with | to75 th centiles) | between boys and girls: median |
| clinical factors | case series | <u>Inclusion</u> | Median age 8.4 | radiopaque | | defecation frequency at intake lower in |
| and colonic | | criteria: | years | markers | a. Boys (n=109) | girls than boys (1.0 vs. 2.0 times/week; |
| transit time. | Evidence | All referred | | | -total colon: 60 (38 to 103) | p=0.03); encopresis frequency more |
| 2004. Archives | <u>level</u> : III | patients ≥ 5 | Country: | Reference: | -delayed >62 h: 49% | than twice weekly reported more often |
| of Disease in | | years old, at | the Netherlands | Clinical variables: | -ascending colon: 10 (5 to 16) | in boys (94% vs. 73%; p=0.0002). More |
| Childhood | Study aim: | least two of the | | | -delayed >18 h: 23% | girls than boys reported no encopresis |
| 89[8], 723-727 | To | following: | | -defecation | -descending colon: 11 (4 to 18) | at all (20% vs. 6% p<0.05) |
| | investigate | 1) defecation | | frequency | -delayed >20 h: 21% | |
| | the relation | <3/week 2) | | -encopresis | -rectosigmoid: 37 (19 to 68) | At entry all children underwent CCT. |
| | between | encopresis | | frequency | -delayed >34h: 53% | Treatment with oral/rectal laxatives |
| | symptoms of | episodes | | -night-time | | discontinued for at least 4 days before |
| | chronic | >1/week 3) | | encopresis | b. Girls (n=60) | the test; during this period they took |
| | constipation | passing of very | | -rectal mass | -total colon: 53 (37 to 74) | one sachet of fibre (Volcolon, 6g) each |
| | and colonic | large stools | | | -delayed >62 h: 43% | day. Then they ingested a capsule |
| | transit time | every7-30 days | | | -ascending colon: 11 (5 to 15) | containing 20 radiopaque markers on 3 |
| | (CTT). To | 4)a palpable | | | -delayed >18 h: 18% | consecutive mornings. Abdominal X ray |
| | evaluate the | abdominal or | | | -descending colon : 8 (5 to 18) | performed on days 4 and 7 in morning. |
| | possible | rectal faecal | | | -delayed >20 h: 23% | Additional abdominal x ray performed |
| | relation | mass | | | -rectosigmoid: 31 (17 to 47) | on days 10, 13 and 16 if more than 20% |
| | between | | | | -delayed >34h: 38% | of markers remained on previous film. X |
| | symptoms | <u>Exclusion</u> | | | | ray localisation of markers based |
| | and CTT | criteria: | | | c. Total group (n=169) | identification of bony landmarks and |
| | and the | Hirschsprung's | | | -total colon: 58 (37 to 92) | gaseous outlines. Markers counted in |
| | outcome | disease, spinal | | | -delayed >62 h: 47% | right, left and rectosigmoid region and |
| | after one | and anal | | | -ascending colon: 10 (5 to 16) | mean segmental transit time calculated |
| | year of | abnormalities, | | | -delayed >18 h: 21% | according to previously described |
| | follow up | mental | | | -descending colon: 10 (5 to 18) | formula. |
| | | retardation, use | | | -delayed >20 h: 22% | |

| udy type Number Evidence patients level prevaler | & Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|--|--------------------|-------------------------------------|---|--|
| of drugs of than laxative setting: gastrointes outpatient of the setting setti | ner es tinal | | -rectosigmoid: 32 (18 to 63) -delayed >34h: 48% (no significant differences between boys and girls in the CTT and rectosigmoid transit time) Correlation between clinical parameters and transit time (hours) (RSTT: rectosigmoid transit time (hours) (RSTT: rectosigmoid transit time) 1. Defection frequency: a. 0 to1/week (n=79) CTT (median): 74 RSTT (median): 38 b. >1 to 3/week (n=55) CTT (median): 30 c. ≥ 3/week (n=35) CTT (median): 49 RSTT (median): 28 CTT: p=0.001 a. vs. b and a vs. c RSTT: p= 0.009 a. vs. b and a vs. c 2. Encopresis frequency (day and night) a. no encopresis (n=18) CTT (median): 49 RSTT (median): 24 | Normal ranges for total and segmental transit times based on upper limits (mean ± 2 SD) from a study in healthy children: CTT > 62 h considered delayed. Upper limits for right colon, left colon ad rectosigmoid transit time were 18, 20 and 34 hours respectively Reviewers' comments: Researchers not blinded No definition of encopresis given No control group Source of funding: not stated |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|--|------------------|
| | | | | | b. <1/day (n=24) CTT (median): 52 RSTT (median): 31 | |
| | | | | | c. 1 to 2/day (n=48) CTT (median): 50 RSTT (median): 30 | |
| | | | | | <u>d. ≥2/day (n=79)</u> CTT (median): 70 RSTT (median): 38 | |
| | | | | | CTT: p= 0.003 d vs. c, d vs. b, and d vs. a RSTT: p= 0.03 d vs. c, d vs. b, and d vs. a | |
| | | | | | 3. Night time encopresis: a. not present (n=106) CTT (median): 47 RSTT (median): 28 | |
| | | | | | b. present (n=63) CTT (median): 74 RSTT (median): 46 | |
| | | | | | CTT: p< 0.0001 RSTT: p< 0.0001 | |
| | | | | | 4. Rectal mass: a. not present (n=118) CTT (median): 48 RSTT (median): 28 b. present (n=51) CTT (median): 86 | |
| | | | | | RSTT (median): 64 CTT: p< 0.0001 | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|-----------------|------------------|
| | | | | | RSTT: p< 0.0001 | |
| | | | | | | |
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| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------|----------------------|-------------------------------|----------------------------|---------------------------------------|--|
| | level | prevalence | | standard | | |
| | Study type: | 211 children | 211 children | Test: | Total CTT (hours, mean and | Additional information from study: |
| Defaecation | diagnostic | | 0 4 (50) | Colonic transit | range): | Significant differences in the study |
| disorders in | case control | Inclusion | Group 1 (PC) | time (CTT) with | 0 4 (50 400) | population regarding clinical variables: |
| children, colonic | - · · | criteria: | N=129 | radiopaque | -Group 1 (PC, n=129): | more PC children reported large |
| | Evidence | | 64% boys | markers | 79.3 (2.4 to 384) | amount of stools, a palpable abdominal |
| | <u>level</u> : III | | Median age: 8 | 5 (| 0 0 0 1 1 1 1 5 0 5 1 0 | mass and rectal mass as compared to |
| score. 1995. | O. 1 . | defection, | years (5-14) | Reference test: | -Group 2 (isolated ES, n=54): | RAP children (p<0.001). More PC |
| European | Study aim: | soling, | 0 0 1 1 1 | Plain abdominal | 41.4 (16.6 to 104.4) | children reported abdominal pain and |
| Journal of | to | encopresis or | Group 2 (isolated | radiography (read | | no rectal sensation as compared to ES |
| | objectivate | recurrent | ES) | using the Barr | -Group 3 (RAP, n=23): | children (p<0.05) |
| 154[4], 277-284 | • | | N=54 | score) | 32.5 (4.8 to 69.6) | |
| | or absence | | 81% boys | | | Two experienced paediatric radiologists |
| | of faecal | | Median age: 9 | | -Healthy controls (n=23, mean | familiar with the Barr criteria and |
| | | patients who | years (5-17) | Barr scoring | + 2SD) (Arhan et al.) | without any knowledge of the clinical |
| | | met at least 2 of | | system: | 29.0 (62) | condition of the patient, independently |
| | _ | | Group 3 (RAP) | Quantifies the | | analysed in random order the first (day |
| | and compare | | N=23 | | p=0.03 group 2 vs. group 3 | 4) and second (day 7) plain abdominal |
| | these | constipation | 39% boys | in four different | | radiographs of the markers studies of |
| | findings to | | Median age: 9 | | Segmental CTT (hours, mean | the initial 101 consecutive patients. Barr |
| | | frequency less | years (5-16) | | and range): | scores were assessed in the different |
| | score | than 3 | _ | (0,1, or 2 points);, | -Right colon: | segments and total scores calculated. A |
| | | | Country: | | Group 1 (PC, n=129): | radiograph was considered positive if |
| | | or more | the Netherlands | (0,3, 4 or 5 | 13.2 (<1.2 to 60) | Barr score>10 |
| | | soling/encopresi | | points) | | |
| | | S | | | Group 2 (isolated ES, n=54): | Normal range for segmental and total |
| | | episodes/week | | (0,3, 4 or 5 | 7.9 (<1.2 to 26.4) | CTT taken from upper limits obtained in |
| | | 3) periodic | | points) | | healthy controls (mean ± 2SD), as |
| | | passage of very | | | Group 3 (RAP, n=23): | described by Arhan et al. |
| | | large amounts | | or 5 points) and | 7.7 (1.2 to 21.6) | Total CTT > 62h: delayed |
| | | of stools once | | also the | | Total CTT > 100h: slow transit |
| | | every 7-30 days | | | -Healthy controls (n=23, mean | constipation (based on study by |
| | | 4) a palpable | | faces i.e. scybala | + 2SD) (Arhan et al.) | Corazziari et al.) |
| | | abdominal or | | (0,1,2 or 3 | 7.7 (18) | Normal limits for segmental transit |
| | | rectal mass | | points); granular | | times (h): right colon (18), left colon |
| | | -Group 2: only | | (0,2, 4 or 5 | p<0.01 group 1 vs. group 2 | (20), rectosigmoid (34) |
| | | encopresis | | points) | and group 1 vs. group 3 | |
| | | and/or soiling | | | | Colonic transit time assessment |
| | | (ES), without | | | -Left colon: | method: Metcalf |

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|---------------|------------|--------------------------------|-----------------|------------------|--|--|
| Information | & Evidence | patients & | Characteristics | Reference | and NPV | Trovious Commont |
| | level | prevalence | | standard | | |
| | | any of the other | | | Group 1 (PC, n=129): | |
| | | criteria for PC. | | | 16.1 (<1.2 to 110.4) | Measurements of CTT performed with |
| | | Soiling defined | | | , | patients on their habitual diet. |
| | | as the loss of | | | Group 2 (isolated ES, n=54): | Treatment with laxatives ([ills or |
| | | loose stools. | | | 6.8 (<1.2 to 25.2) | enemas) discontinued for at least 4 |
| | | Encopresis | | | | days before the CTT study |
| | | defined as | | | Group 3 (RAP, n=23): | |
| | | (in)voluntary | | | 7.0 (1.2 to 25.2) | 5 patients excluded from study: 4 not |
| | | passage of a | | | | able to swallow capsule, 1 had |
| | | normal bowel | | | -Healthy controls (n=23, mean | "uninterpretable" abdominal X-ray |
| | | movement in | | | + 2SD) (Arhan et al.) | |
| | | the underpants | | | 8.7 (20) | Comparison of the Barr-score with the |
| | | or another | | | | marker method performed using the |
| | | unorthodox | | | p<0.01 group 1 vs. group 2 | mean Barr-score of the two observers |
| | | location with a | | | and group 1 vs. group 3 | obtained on radiograph I. Similar |
| | | frequency of 2 | | | Destesionesid | analysis using radiograph II revealed no |
| | | or more | | | -Rectosigmoid | differences compared to radiograph I, |
| | | times/week after the age of | | | Group 1 (PC, n=129): 49.7 (<1.2 to 226.8) | therefore only results with radiograph I are presented in detail |
| | | 4 in the | | | 49.7 (<1.2 to 226.6) | are presented in detail |
| | | absence of any | | | Group 2 (isolated ES, n=54): | According to authors the radiopaque |
| | | organic cause | | | 26.7 (4.8 to 93.6) | markers were no hindrance for the 2 |
| | | -Group 3: RAP | | | 20.7 (4.0 to 95.0) | observers in assessing the Barr-scores |
| | | defined as at | | | Group 3 (RAP, n=23): | observers in assessing the barr-scores |
| | | least 3 | | | 18.9 (1.2 to 49.2) | Reviewers' comments; |
| | | episodes/week | | | 10.0 (1.2 to 10.2) | There are missing data not accounted |
| | | of non specified | | | -Healthy controls (n=23, mean | for: only 101 abdominal radiographs |
| | | RAP, severe | | | + 2SD) (Arhan et al.) | were available for analysis, but there is |
| | | enough to | | | 12.4 (34) | no clear explanation for this |
| | | interfere with | | | | ' |
| | | day-to day | | | p<0.01 group 1 vs. group 2 | Source of funding: not stated |
| | | activities over at | | | and group 1 vs. group 3 | |
| | | least a 3-month | | | | |
| | | period, without | | | p=0.05 group 2 vs. group 3 | |
| | | any of the other | | | | |
| | | symptom of PC | | | CCT | |
| | | | | | -Interobserver agreement: | |
| | | Exclusion | | | | |

| Bibliographic Study type & Evidence level | | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---|--|-------------------------------|-------------------------------------|--|------------------|
| | criteria: Hirschsprung's disease, spinal/ anal anomalies, prior surgery of colon, metabolic diseases, mental retardation, use of drugs other than laxatives Setting: gastroenterolog y outpatients clinic | | | Radiograph 1 (n=101): Perfect agreement: 62% Difference of one marker: 25% Radiograph 2 (n=101): Perfect agreement: 92% Difference of one marker: 6% Barr scores (n=101) (mean of two observers) -Group 1 (PC, n=57) Radiograph 1: ≥10: 60% Radiograph 2: ≥10: 63% -Group 2 (isolated ES, n=30) Radiograph 1: ≥10: 47% Radiograph 2: ≥10: 60% -Group 3 (RAP, n=14) Radiograph 1: ≥10: 47% Radiograph 2: ≥10: 63% -Interobserver agreement (agreement between the 2 observers for the different segments on the same radiograph): k from 0.28 (fair) to 0.60 (moderate) -Intraobserver agreement (difference in quantity and quality of stool between radiograph I and II as scored | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|---|------------------|
| | | | | | by same radiologist): k from 0.05 (poor) to 0.47 (moderate) for both observers | |
| | | | | | -Intraobserver agreement (agreement on the existence of constipation as measured by a Barr-score of 10 or more points between radiographs I and II): fair for both observers, k= 0.22 and 0.25 respectively | |
| | | | | | Correlation of the Barr-score with Metcalf's makers method: Correlation between positive Barr score (≥10) and delayed total CTT (>62h): k=0.22 (fair) for all children. | |
| | | | | | K values by group: -PC group: 0.20 -ES group: 0.02 -RAP group: 0.46 | |
| | | | | | Abnormal Barr scores found in at least 46% of patients with normal transit times. Positive Barr scores correlated only with total CTT exceeding 100 h | |

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|----------------------------|------------------|----------------------------|--------------------|--------------------|---|---|
| Information | & Evidence level | patients & prevalence | Characteristics | Reference standard | and NPV | |
| Gutierrez et al. | Study type: | 68 children | 68 children | Test: | Total transit time (hours) | Additional information from study: |
| Total and | Diagnostic | | aged 2 to 14 years | Colonic transit | (mean ± SD, ranges) | Two children from patients group did |
| segmental | case control | <u>Inclusion</u> | | time (CTT) with | | not complete study: one refused to |
| colonic transit | | criteria: | Patients (n=38) | radiopaque | Patients (n=38) | swallow the capsules; one did not |
| time and | <u>Evidence</u> | Patients: history | | markers | 49.57 ± 25.38 (15.6 to 122.4) | comply (not clear exactly with what) |
| anorectal | level: | of chronic | Controls (n=30) | | | |
| manometry in | III | idiopathic | | Reference: | Controls (n=30) | No significant differences observed in |
| children with | | constipation > 6 | | Frequency of | 29.08 ± 8.30 (14.4 to 50) | mean daily fibre intake and calorie |
| chronic | Study aim: | months, | Spain | defecation | | consumption between the 2 groups |
| idiopathic | to evaluate | with/without | | | p<0.001 | Management was do not its abildon |
| constipation. | the use of a | secondary | | | Commontal transit times (hours) | Measurements made while children |
| 2002. Journal of Pediatric | motility study | encopresis, refractory to | | | Segmental transit time (hours) (mean ± SD, ranges) | maintained their usual diets. Laxative treatment discontinued 1 week before |
| Gastroenterolog | | conventional | | | (mean ± SD, ranges) | the test and a cleansing enema |
| | applied in | treatment of | | | -RC: | administered on the day before the test |
| 35[1], 31-38 | daily clinical | disimpaction, | | | Patients (n=38) | administered on the day before the test |
| 00[1], 01 00 | practice to | re-education of | | | 9.53 ± 9.07 (2.4 to 36) | No differences observed in CTT in |
| | more clearly | defecatory | | | 0.00 ± 0.07 (2.1 to 00) | relation to either se or age. Statistically |
| | define | habits, | | | Controls (n=30) | significant inverse correlation observed |
| | patients with | measures to | | | 7.52 ± 5.75 (2.4 to 15.6) | between total CTT and number of |
| | this disorder | increase dietary | | | , | weekly defecations (correlation |
| | and to | fibre content | | | p value NS | coefficient, r=0.68, p<0.001) |
| | improve | and | | | | |
| | therapy and | administration | | | -LC: | Reviewer comments: |
| | follow-up | of mineral oil or | | | Patients (n=38) | Researchers not blinded |
| | | osmotic-type | | | 15.41 ± 13.13 (2.4 to 32) | |
| | | laxatives | | | | Source of funding: Janssen |
| | | (lactulose or | | | Controls (n=30) | Pharmaceutical contributed the material |
| | | Lactinol). | | | 6.60 ± 6.20 (2.4 to 24) | required to determine the colonic transit |
| | | Encopresis defined as non- | | | p=0.01 | time. No further details provided |
| | | voluntary | | | p=0.01 | |
| | | defecation with | | | -RS: | |
| | | a frequency of | | | Patients (n=38) | |
| | | more than twice | | | 24.20 ± 16.77 (4.8 to 69.6) | |
| | | weekly in | | | (| |
| | | children older | | | Controls (n=30) | |
| | | than 4 years in | | | 14.96 ± 8.70 (2.4 to 19.2) | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|--------------------------|----------------------|-------------------------------|----------------------------|---------------------------------------|------------------|
| | level | prevalence | | standard | | |
| | | the absence of | | | | |
| | | any underlying | | | p=0.01 | |
| | | organic cause | | | | |
| | | Controls: | | | Clinical characteristic of the | |
| | | normal bowel | | | patients' group as a function of | |
| | | habits (between | | | colonic transit time: | |
| | | 3 defecations | | | | |
| | | daily and 3 | | | a) Age at onset of constipation | |
| | | weekly, without | | | (y, mean, SD): | |
| | | staring at stool, | | | -Total CTT within reference | |
| | | and faces of | | | values (n=19): 2.54 (1.18) | |
| | | normal | | | -Prolonged total CTT (n=19): | |
| | | consistency for | | | 1.77 (0.88) | |
| | | at least 12 | | | p<0.05 | |
| | | months before | | | | |
| | | the study, no | | | b) Family history of | |
| | | history of | | | constipation: | |
| | | previous | | | -Total CTT within reference | |
| | | abdominal/majo | | | values (n=19): 21% | |
| | | r extra- | | | -Prolonged total CTT (n=19): | |
| | | abdominal | | | 79% | |
| | | surgery, not on | | | p<0.01 | |
| | | medication with | | | | |
| | | effects on | | | c) Abdominal mass | |
| | | digestive tract, | | | -Total CTT within reference | |
| | | normal diet, and | | | values (n=19): 60% | |
| | | underwent | | | -Prolonged total CTT (n=19): | |
| | | abdominal | | | 93.8% | |
| | | radiography as | | | p<0.05 | |
| | | part of clinical | | | [| |
| | | study with | | | d) Encopresis episodes/night | |
| | | normal results | | | (mean, SD) | |
| | | | | | -Total CTT within reference | |
| | | Exclusion | | | values (n=19): 0.10 (0.44) | |
| | | criteria: | | | -Prolonged total CTT (n=19): | |
| | | Hirschsprung's | | | 0.60 (0.91) | |
| | | disease, spinal/ | | | p<0.05 | |
| | | anal | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|---------------------------|-----------------------------------|--|-------------------------------|---|---|------------------|
| | | malformations, prior surgery of colon, metabolic diseases, mental retardation | | | No significant differences found for age at diagnosis, sex, defecations/week, pain at defecation, enuresis, anal fissure, rectal mass or encopresis episodes/day | |
| | | Setting: gastroenterolog y outpatients clinic | | | | |
| | | | | | | |
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| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------|------------------------------|-------------------------------|----------------------------|---------------------------------------|--|
| | level | prevalence | | standard | | |
| Papadopoulou | Study type: | 52 children | 52 children | Test: | Patterns of transit time (n=52): | Additional information from study: |
| et al. The | Diagnostic | | Median age: 8 | Colonic transit | -normal transit: 21 (40%) | -To assess reliability of test |
| | prospective | <u>Inclusion</u> | years (range 2-13.5 | | -mild delay: 4 (8%) | interobserver error between 2 |
| | case series | criteria: | years) | radiopaque | -moderate delay: 9 (17%) | observers was measured: each |
| transit studies in | | Constipation | | markers | -severe delay: 18 (35%) | independently assessing 30 abdominal |
| childhood | <u>Evidence</u> | | Sex distribution not | | | X-rays and interobserver error by |
| constipation | level: III | One patient had | reported | Reference: | Patterns of marker distribution: | carrying out duplicate estimations by |
| and soiling. | | neurological | _ | Frequency of | -pancolonic transit delay: 15 | the same observer on the same 30 |
| | | problems due to | | bowel | (29%) | days |
| | assess the | ganglioneuroma | UK | movements and | -segmental transit delay: 5 | |
| | acceptability, | | | soiling | (10%) | -Assessment criteria of severity of |
| 153[8], 560-564 | the reliability | Constipation | | | -outlet obstruction: 11 (21%) | transit delay: |
| | of | defined as less | | | | a. normal transit: < 12 markers in colon |
| | interpretation | | | | Correlation between transit | (<40% of given markers) |
| | and the | movements/we | | | delay and clinical symptoms: | b. mild delay: 12-18 markers in colon |
| | clinical value | | | | | (41-60% of given markers) |
| | of solid | defined as | | | a) Fewer than 2 bowel | c. moderate delay: 19-24 markers in |
| | marker | involuntary | | | movements/week (%): | colon (61-80% of given markers) |
| | transit | passage of fluid | | | Children with acyers delay | d. severe delay: >24 markers in colon |
| | studies in children with | or semi-solid stools into | | | -Children with severe delay (n=18): | (>80% of given markers) |
| | soiling and | clothing 2/more | | | (11–16). 187 | -Assessment criteria of different |
| | spurious | times/week | | | -Children with normal transit | patterns of marker distribution: |
| | diarrhoea | tillies/ week | | | (n=21): 27 | a. pancolonic transit delay: no single |
| | (otherwise | Exclusion | | | (11—21). 21 | segment contains >75% of markers |
| | known as | criteria: | | | p<0.001 | remaining in colon |
| | overflow | Hirschsprung's | | | F 101001 | b. segmental transit delay: >75% of |
| | incontinence | disease | | | b) More than 3 soiling | markers remaining in colon clustered in |
| |)* | | | | episodes/week (%): | one segment |
| | ' | Setting: | | | | c. outlet obstruction: >60% of given |
| | | hospital | | | -Children with severe delay | markers clustered in rectosigmoid |
| | | 1 | | | (n=18): | |
| | | | | | 92 | -In 6 patients the transit studies were |
| | | | | | | repeated after colonic washout. |
| | | | | | -Children with normal transit | Significant improvements in transit |
| | | | | | (n=21): 35 | found after colonic emptying (p<0.05) |
| | | | | | | (exact number not reported in text, just |
| | | | | | p<0.005 | a bar graph) |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|---|---|
| | | | | | No correlation found between duration of symptoms and severity of delay Correlation between marker distribution and transit -Children with severe delay (n=18): Outlet obstruction: 39% Pancolonic transit delay: 56% Segmental transit delay (in descending colon): 5% -Children with mild delay (n=4): Pancolonic transit delay: 25% Segmental transit delay (in rectosigmoid): 75% P<0.005 Correlation between marker distribution and symptoms: -Fewer than 2 bowel movements/week (%): a. Outlet obstruction: 100% b. Pancolonic transit delay: 83% c. Segmental transit delay: 33% a vs. c and b vs. c: p<0.05 -More than 3 soiling | Laxative treatment not interrupted previous to measurements (97% were on laxatives) Reviewers' comments: Researchers not blinded No data on the type of diet children were on when measurements were made No data reported on the correlation between transit delay and clinical symptoms for children with mild/moderate delay Source of funding: not stated |
| | | | | | episodes/week (%): a. Outlet obstruction: 100% b. Pancolonic transit delay: | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|--|------------------|
| | | | | | 57% | |
| | | | | | c. Segmental transit delay: 0% | |
| | | | | | a vs. c and b vs. c: p<0.05 | |
| | | | | | Observer errors: | |
| | | | | | (coefficient of variation): -interobserver: 2.1 % | |
| | | | | | -intraobserver: 3.1 % | |
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| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|-----------------------------------|-------------------------------|-------------------------------------|--|--|
| | Study type: | 141 children | 141 children | Test: | Total gastrointestinal transit | Additional information from study: |
| Gastrointestinal | Diagnostic | | | Total | time (TGITT) (hours, mean ± | No patients receiving laxatives during |
| transit time, | case control | <u>Inclusion</u> | Patients: | gastrointestinal | SD, range) | investigation |
| frequency of | | criteria: | N=63 | transit time | -healthy controls (n=78) | |
| defecation, and | <u>Evidence</u> | | 40 boys | (TGITT) ¹ | 25.0 ± 3.7 (19 to 33) | Retention of contents in a given large |
| anorectal | level: III | | Mean age 5.4 ± 4.1 | | | bowel segment considered abnormally |
| manometry in | | constipation, | years (2 months to | Reference: | -patients with TGITT>33h | prolonged when transit index ≤60 (i/e |
| healthy and | | | 4 years) | -Frequency of | (n=53) | when on average, ≥ 30% of markers |
| constipated | quantify | reduced bowel | | defecation | 81.4% | were retained in that given segment at |
| | bowel | frequency | Controls: | | | least 33 h after ingestion of radiopaque |
| | function in | | N=78 | | -patients with TGITT<33h | pellets). Transit index of 60 chosen |
| | healthy | straining at | 37 boys | | (n=10) | because the lower confidence limit (?) |
| 106[3], 379-382 | | | Mean age 5.5 ± 3.2 | | 18.6% | of a normal adult population did not |
| | regard to | presence of | years (2 months to | | | exceed this value |
| | frequency of | | 12 years) | | Segmental transit time | |
| | , | blood on faeces | _ | | N=39 (out of 53 children with | Reviewers' comments: |
| | | | Country: | | prolonged transit time) | Not clear what type of diet patients |
| | nal transit | of mild laxatives | Italy | | | were following during investigation |
| | time and | -Controls: | | | Colon: lowest in 3 patients | |
| | | healthy children | | | | Segmental colonic transit times (right |
| | | free of bowel | | | Rectum: lowest in 24 patients | and left colon and rectosigmoid) |
| | cs of the | complaints | | | | measured but results not reported |
| | anorectal | | | | Colon and rectum: lowest in 12 | |
| | | Exclusion | | | patients | Accurate figures for CTT in patients not |
| | compare | criteria: | | | | reported |
| | variables of | secondary | | | Frequency of defecation | |
| | bowel | constipation | | | (times/week): | Segmental transit time not measured in |
| | function in | excluded after | | | -healthy controls (n=78) | controls |
| | | clinical interview | | | 6.3 ± 1.3 (range 4 to 9) | Describe was auto-d-fourth a baselthy, as atracks |
| | chronic | and | | | notion to with TOITT: 22h | Results reported for the healthy controls |
| | | examination, barium enema, | | | -patients with TGITT>33h | are not clearly stated in the paper that |
| | the normal | anorectal | | | (n=53) 2.5 ± 0.9 (range not reported) | there actually belong to this group, but as results for the patients group are |
| | | | | | 2.5 ± 0.9 (range not reported) | explicitly related to them, it was |
| | population | motility studies, rectosigmoidosc | | | -patients with TGITT<33h | assumed the others belonged to the |
| | | I ectosigniolausc | | 1 | -pauerns with TGHT (SSH | assumed the others belonged to the |

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¹ Italian papers included in this review (Corazziari, Cucchiara, Staiano) measured "total gastrointestinal transit time (TGITT)". Because of the similarity in the figures with the other studies' CTTs we assumed that TGITT is the name by which CTT known in Italy.

| Information & Evi | y type Number of dence patients & vel prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|-------------------|---|-------------------------------|-------------------------------------|--|--|
| | opy, rectal biopsy. Metabolic and endocrinologic abnormalities. Setting: unclear | | | (n=10) 5.1 ± 0.73 (range not reported) Stool frequency and TGITT significantly correlated in patients with prolonged transit time (r=0.75; p<0.001) and in healthy controls (r=0.78; p<0.001) In 7 of 53 patients with TGITT>33 h, the bowel frequency overlapped the range observed in the controls | controls Researchers not blinded Source of funding: not stated |

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|-----------------|------------------|-----------------------|---------------------|---------------------|---------------------------------|--|
| Information | & Evidence level | patients & prevalence | Characteristics | Reference standard | and NPV | |
| Benninga et al. | Study type: | 148 children | 148 children | Test: | Total transit time (hours, | Additional information from study: |
| Colonic transit | Diagnostic | | | Colonic transit | median, range) | Total and segmental CTT done as |
| time in | case control | <u>Inclusion</u> | -Patients (n=94): | time (CTT) with | -PSTC (n=24) | described by Metcalf |
| constipated | | criteria: | | radiopaque | 189 (104.4 to 380.4) | |
| children: does | <u>Evidence</u> | -Patients: | a. PSTC (paediatric | markers | | Based on upper limit (mean + 2SD) of |
| pediatric slow- | level: | otherwise | slow transit | | -NDTC (n=70) | previous study in 63 constipated |
| transit | III | | constipation): | Reference | 46.8 (3.6 to 99.6) | children (Corazziari, 1985), children in |
| constipation | | | 24 children | -Clinical variables | | current study arbitrarily separated in 2 |
| exist? 1996. | Study aim: | | 17 boys | | Segmental transit time (hours, | groups: |
| Journal of | To | with/without | Mean age 8 years | | median, range) | 1. CTT>100 h: paediatric slow transit |
| Pediatric | investigate | encopresis, | (range 5-14) | | | constipation (PSTC) |
| Gastroenterolog | | | | | Right colon: | 2. CTT<100 h: normal- or delayed- |
| 1 - | of slow | alone or | b. NDTC (normal | | -PSTC (n=24) | transit constipation (NDTC) (normal |
| 23[3], 241-251 | colonic | | delayed transit | | 27.0 (3.6 to 60) | transit ser at < 63h) |
| | transit in | | constipation) | | | Further analysis of the NDTC group |
| | children with | | 70 children | | -NDTC (n=70) | after separation into a group with total |
| | constipation | least 2 of the | 46 boys | | 8.4 (0 to 32.4) | CTT<63h and one with total CTT |
| | using | | Mean age 8 years | | | between 63 and 100h showed same |
| | radiopaque | for paediatric | (range 5-14) | | Left colon: | significant differences compared with |
| | markers | constipation: a) | | | -PSTC (n=24) | PSTC children as did the total PSTC |
| | | 2/fewer bowel | -Controls (n=54): | | 37.2 (0 to 110.4) | group allowing the merge of these |
| | | movements/we | 15 children (for | | | children |
| | | ek b)2/more | rectal manometry) | | -NDTC (n=70) | 0 |
| | | soiling or | 10 boys | | 7.2 (0 to 36.0) | CTT performed on patients taking their |
| | | | Mean age 11 years | | . | normal diet, any treatment with |
| | | episodes/week | (range 7-15) | | Rectosigmoid: | laxatives discontinued at least 4 days |
| | | c) passage of | | | -PSTC (n=24) | prior o test. No enemas given before |
| | | very large | Country: | | 116.4 (49.2 to 226.8) | transit studies. |
| | | amounts of | the Netherlands | | NIDTO (70) | D. i |
| | | stool once | | | -NDTC (n=70) | Reviewers' comments: |
| | | every 7-30 days | | | 27.0 (0 to 90.0) | Researchers not blinded |
| | | d) a palpable | | | Clinian Lynniah Is = : | Values for both total and a surrent d |
| | | abdominal | | | Clinical variables: | Values for both total and segmental |
| | | mass or rectal | | | -Daytime soiling (yes/no) (no., | transit times expressed as medians in |
| | | mass -Controls: | | | %) -PSTC (n=24) | the text and the heading of a table, and as means in the table itself. We have |
| | | | | | | |
| | | healthy | | | 22 (92) | chosen to report them as median |
| | | children. | | | -NDTC (n=70) | values because authors stated in the |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|--|-------------------------------|-------------------------------------|---------------------------------------|--|
| | & Evidence | | | Reference | | statistical analysis section that results were expressed as median and range for continuous variables Source of funding: major grant from the Stitching Kinderpostzegels Nederland and from an endowment from Zyma Nederland (Importal) |
| | | Setting: outpatient clinic of tertiary academic | | | -NDTC (n=70) 33 (49) p=0.03 | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|--|------------------|
| | | teaching hospital | | | -Pain during defecation (no., %) -PSTC (n=24) 8 (33) -NDTC (n=70) 28 (60) p=0.01 | |
| | | | | | -No rectal sensation (no., %) -PSTC (n=24) 8 (33) | |
| | | | | | -NDTC (n=70) 10 (14) p=0.03 | |
| | | | | | -Palpable abdominal mass (no., %) -PSTC (n=24) 17 (71) -NDTC (n=70) 27 (39) p=0.02 | |
| | | | | | -Palpable rectal mass (no., %) -PSTC (n=24) 17 (71) | |
| | | | | | -NDTC (n=70) 9 (13) p<0.01 | |
| | | | | | No significant differences regarding: sex, age, toilet training statue, age at which toilet training started, bowel movements/week, large amounts of stools very 7-30 | |

| Bibliographic Information | & Evidence Number patients level prevalen | & Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|------------------------------|---|-------------------|-------------------------------------|---|------------------|
| | ievei prevalen | | Standard | days, encopresis episodes/week, abdominal pain, poor appetite, daytime or nightime urinary incontinence Proportion of children with PSTC and rectal palpable mass, night time soiling or both: 0.34, 0.39 and 0.82 respectively. (multivariate analysis) only 7% of children without any of these characteristics had PSTC | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|---------------------------------------|---------------------------------------|
| Yang et al. | Study type: | 96 children | 96 children | Tests: | Total transit time (hours, mean | Reviewers' comments: |
| | Diagnostic | | | Colonic transit | <u>± SD)</u> | Researchers not blinded |
| | case control | <u>Inclusion</u> | -Patients (n=28): | time (CTT) with | | |
| gastrointestinal | | criteria: | 38 boys | radiopaque | -Patients (n=28) | No data available on diet, use of |
| | <u>Evidence</u> | | Mean age: 6 years | markers | 59.9 ± 2.3 | laxatives previous to the measurement |
| functional | <u>level:</u> III | confirmed | (range 3 to 14) | | | of CTT |
| constipation in | | functional | | Reference: | -Controls (n=68) | |
| | Study aim: to | | | none | 14.8 ± 0.8 | Source of funding: |
| Chinese Journal | | (FC). Two of | -Controls (n=68) | | | not stated |
| | the | the following for | | | p<0.01 | |
| | difference of | more than 3 | Mean age: 6 years | | | |
| | | months: | (range 3 to 13) | | Segmental transit time (hours, | |
| | nal transit | Evacuation less | _ | | mean ± SD) | |
| | time (GTT) | | Country: | | | |
| | between | evacuating | China | | Right colon: | |
| | constipated | pains, faecal | | | -Patients (n=28) | |
| | and normal | soiling every | | | 20.3 ± 1.2 | |
| | healthy | week or | | | | |
| | controls to | incontinence | | | -Controls (n=68) | |
| | elicit its | more 2 | | | 7.3 ± 1.1 | |
| | significance | times/week in | | | | |
| | in assessing | over 5 years | | | p<0.01 | |
| | the | old, touchable | | | Latteralani | |
| | • | stool by | | | Left colon: | |
| | the whole | abdominal or | | | -Patients (n=28) | |
| | gastro- | anal | | | 12.8 ± 1.7 | |
| | intestine and each | examination, | | | Controlo (n. 60) | |
| | | excessive defecation at | | | -Controls (n=68) 3.4 ± 0.8 | |
| | segment | interval of 7 to | | | 3.4 ± 0.6 | |
| | | 30 days. No | | | p<0.01 | |
| | | administration | | | p<0.01 | |
| | | of | | | Rectosigmoid: | |
| | | gastrointestinal | | | -Patients (n=28) | |
| | | dynamic and | | | 26.8 ± 1.4 | |
| | | evacuation | | | 20.0 ± 1.7 | |
| | | drugs for 2 | | | -Controls (n=68) | |
| | | weeks | | | 4.1 ± 1.2 | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|--|-------------------------------|-------------------------------------|---------------------------------------|------------------|
| | | -Controls: normal height and weight, normal frequency and character of evacuation fro 3 months without administration of any gastrointestinal dynamic and evacuation drugs | | | p<0.01 | |
| | | Exclusion criteria: organic ailment in alimentary tract and other organs ailment that would affect gastrointestinal function Setting: general hospital | | | | |

| level prevalence standard | |
|--|---|
| Cucchiara et al. Gastrointestinal transit time and anorectal manometry in children with fecal soiling. 1984. Journal of Pediatric Gastroenterolog y and Nutrition 3[4], 545-550 [All, 545-550] [All type: Diagnostic case-control anorectal manometry in children with fecal soiling. 1984. Journal of gastrointestinal total gastrointestinal transit time (TGITT) [Additional inform Controls matche but not sex with children with fecal soiling. 1984. Journal of the measure total gastrointestinal transit time (TGITT) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls matche but not sex with children with soiling (n=46) [Additional inform Controls matche but not sex with children with soiling (n=32) [Additional inform Controls (n=46) [Additional inform controls with soiling (n=32) [Additional inform in a pastrointestinal transit time (norm in a pastroin | ed for age and weight the constipated ments performed with heir usual diet ments: f constipation/soiling t blinded of laxatives previous to arium enema, without ing of the colon and ctosigmoid was monstrate the presence garectum or disease sit times not measured |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|----------------------------------|---|---------------------------------------|--|
| Martelli et al. | Study type: | 1182 children | 1182 children | Test: | Total transit time (hours, | Additional information from study: |
| Can functional | Diagnostic | | 63% boys | Colonic transit | median, range) | Patients classified into 4 groups: |
| constipation | retrospective | <u>Inclusion</u> | | time (CTT) with | | -"Normal" transit time |
| begin at birth? | case series | criteria: | Group 1: | radiopaque | -C+E patients (n=168): | -"Pancolic" constipation: delay in the 3 |
| 1998. | | Constipation | constipated | markers | 67.2 (2 to 168) | sites |
| Gastroenterolog | | with/without | children without | | | -"Terminal" constipation: delay in the |
| y International | <u>level:</u> III | encopresis | encopresis (C | Reference: | -C+4 patients (n=112): | rectosigmoid with/without delay in right |
| 11[1], 1-11Italy. | | Constipation | patients) | none | 54.6 (9 to 168) | or left colon |
| | | defined as less | | | | -"Non terminal" constipation: right |
| | analyse | than 3 | N=855 | | -C-4 patients (n=77) | and/or left delay but normal |
| | epidemiologi | spontaneous | 59%boys | | 49.6 (8 to 161) | rectosigmoid transit time |
| | С, | stools/week | | | | |
| | manometric | without any | 65% < 4 years old | | -Controls (n=21) Arhan et al. | Reviewers' comments: |
| | and | laxative or | (C-4 patients) | | 1983 | Researchers not blinded |
| | radiologic | motility- | 35% > 4 years old | | 22.8 (9.4 to 56.4) | l |
| | data in a | influencing | (C+4 patients) | | | Not all children underwent CTT |
| | large | drug. | | | p<0.0001 C+4/C-4/C+E | N 1 / 1 / 1 / 1 |
| | | Encopresis | Median age at first | | patients vs. controls | No data on diet or use of laxatives |
| | young | defined (in | evaluation: | | p<0.05 C+E patients vs. C+4 | previous to CTT measurement |
| | patients | France) as | C-4: 11 months | | patients | On the state of the state of |
| | presenting in | incontinent | (range 4 to 15 | | Commontal transit times (bours | Source of funding: not stated |
| | a paediatric | associated with | years) | | Segmental transit time (hours, | |
| | tertiary care hospital in | faecal | C+4: 7.7 years (range 4 to 15 | | median, range) | |
| | order to | | years) | | 1-Right colon: | |
| | classify | 3 years. Faecal | years) | | -Controls (n=21): Arhan et al. | |
| | different | impaction | Group 2: | | 1983 | |
| | types of | considered to | constipated | | 7.2 (0.6 to 19.2) | |
| | idiopathic | be present | children with | | -C-4 patients (n=77): | |
| | constipation | when | encopresis (C+E | | 14.8 (0 to 96) | |
| | according to | consistency of | patients) | | -C+4 patients (n=168): | |
| | age of onset, | faeces | pationto | | 12 (0 to 48) | |
| | sex and | | N=327 | | -C+E patients (n=112): | |
| | pelvic floor | rectum more | 78% boys | | 14 (0 to 144) | |
| | function | solid than that | Median age at first | | | |
| | | of stools | evaluation: 8.5 | | p<0.0005 C+4/C-4 patients vs. | |
| | | spontaneously | years (range 4 to | | controls | |
| | | emitted | 15 years) | | p<0.0001 C+E patients vs. | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|------------------|
| | ievei | prevalence | Country | Statiuaru | controls | |
| | | Exclusion | Country: France | | Controls | |
| | | criteria: | i iance | | 2-Left colon: | |
| | | children aged < | | | -Controls (n=21): Arhan et al. | |
| | | 48 months. | | | 1983 | |
| | | Local/general | | | 7.4 (1.2 to 22.8) | |
| | | causes of | | | -C-4 patients (n=77): | |
| | | constipation: | | | 12.4 (0 to 72) | |
| | | anal lesions | | | -C+4 patients (n=168): | |
| | | (anal fissures, | | | 12 (0 to 96) | |
| | | anal | | | -C+E patients (n=112): | |
| | | malposition), | | | 13.6 (0 to 96) | |
| | | neurogenic | | | | |
| | | constipation | | | p<0.0005 C-4 patients vs. | |
| | | (Hirschsprung's | | | controls | |
| | | disease, | | | p<0.005 C+4/C+E patients vs. | |
| | | neurointestinal | | | controls | |
| | | dysplasia, | | | | |
| | | spinal cord | | | 3-Rectosigmoid: Arhan et al. | |
| | | disorders, | | | 1983 | |
| | | chronic | | | -Controls (n=21): | |
| | | intestinal | | | 10.4 (1.21 to 34.2) | |
| | | pseudobstructio | | | -C-4 patients (n=77): | |
| | | n),endocrine | | | 18.4 (0 to 106) | |
| | | (hypothyroidism | | | -C+4 patients (n=168): | |
| | |), metabolic | | | 26.4 (0 to 108) | |
| | | disorders | | | -C+E patients (n=112): | |
| | | (diabetes | | | 30.2 (0 to 142) | |
| | | mellitus, renal | | | | |
| | | acidosis, | | | p<0.005 C-4 patients vs. | |
| | | hypercalcemia), | | | controls | |
| | | still breast-fed | | | p<0.0001 C+4/C+E patients | |
| | | patients with not | | | vs. controls | |
| | | symptoms other | | | Olassification of accepting ti | |
| | | than fewer than | | | Classification of constipation | |
| | | 3 stools/week | | | according to segmental colonic | |
| | | Cotting | | | transit times (n, %): | |
| | | Setting: | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|-----------------------------------|-------------------------------|---|--|------------------|
| | ievei | paediatric tertiary care hospital | | Standard | 1.Normal transit: -C-4 patients (n=77): 33 (43) -C+4 patients (n=168): 34 (30.5) -C+E patients (n=112): 38 (22.5) -Total (n=357): 105 (29) p<0.001 C+E vs. C-4 patients 2.Non terminal constipation: -C-4 patients (n=77): 18 (23) -C+4 patients (n=168): 26 (23) -C+E patients (n=112): 37 (22) -Total (n=357): 81 (23) 3.Terminal constipation: -C-4 patients (n=77): 17 (22) -C+4 patients (n=168): 42 (37.5) -C+E patients (n=112): 70 (41.5) -Total (n=357): 129 (36) p<0.05 C+4 vs. C-4 patients p<0.005 C+6 vs. C-4 patients p<0.005 C+7 vs. C-4 patients 14. Pancolic constipation: -C-4 patients (n=77): 9 (12) -C+4 patients (n=168): 10 (9) -C+E patients (n=112): 23 (14) -Total (n=357): 42 (12): 42 (12) | |
| | | | | | (p values not reported were not significant) | |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-------------------------|---------------------------|-------------------------------|----------------------------|---------------------------------------|--|
| | level | prevalence | | standard | | |
| Arhan et al. | Study type: | 176 patients | 176 patients | Test: | Segmental transit time of one | Additional information from study: |
| Idiopathic | Diagnostic | | aged 2 to 15 years | Colonic transit | radiopaque marker (hours, | Markers ingested 24h after beginning a |
| disorders of | case control | <u>Inclusion</u> | 64% boys | time (CTT) with | min; mean ± SD) | diet containing 0.5g/kg of crude fibres |
| fecal continence | | criteria: | | radiopaque | | |
| in children. | <u>Evidence</u> | -Patients: one | Controls: | markers | 1. Ascending colon: | Functional studies performed when |
| 1983. Pediatrics | | _ | 23 children (no | | -normal children (n= 23): | rectum free of stool either |
| 71[5], 774-779 | III | 1) history of | further data | -Reference: none | 7:10 ± 1:4 | spontaneously or as a result of |
| | | less than 3 | reported) | | -constipated children | cleansing enemas |
| | Study aim: to | | | | (with/without spina bifida | |
| | | stools/week 2) | Country: | | occulta) (n=176): | Reviewers' comments: |
| | clinical | evidence of | France | | 13:24 ± 1:5 | No clear definition of constipation given |
| | | faecaloma | | | - 0.05 | Danasan kana matikibada d |
| | of children with | (stools of harder | | | p<0.05 | Researchers not blinded |
| | | consistency than those | | | 2. Descending colon | Not along how many shildren underwant |
| | idiopathic disorders of | passed | | | -normal children (n= 23): | Not clear how many children underwent CTT |
| | faecal | spontaneously) | | | 7:37 ± 1:3 | |
| | continence | at rectal | | | -constipated children | Total transit time not measured |
| | and to | examination 3) | | | (with/without spina bifida | Total transit time not measured |
| | demonstrate | presence of | | | occulta) (n=176): | As no data are reported on the |
| | that they | faecal material | | | 13:49 ± 1:37 | characteristics of the control group it is |
| | have | in the entire | | | | not possible to tell whether they could |
| | functional | descending | | | p<0.05 | be significantly different from the |
| | abnormalitie | colon or | | | ľ | patients |
| | s of large- | faecaloma in | | | 3. Rectum | |
| | bowel | the | | | -normal children (n= 23): | Source of funding: partially by the |
| | motility | rectosigmoid | | | 11:4 ± 1:5 | Institut national de la Sante et de la |
| | | area diagnosed | | | -constipated children | Recherche Medicale (INSERM), CRL |
| | | radiologically | | | (with/without spina bifida | No.80-7002, grant MT-3511 from the |
| | | -Controls: | | | occulta) (n=176): | CRM, and by the French Canadian sub |
| | | children with no | | | 30:22 ± 2:42 | commission for health matters |
| | | intestinal | | | | |
| | | abnormalities | | | p<0.05 | |
| | | who had to | | | | |
| | | undergo a | | | No significant differences | |
| | | radiography of | | | between children with and | |
| | | the abdomen | | | without spina bifida occulta | |
| | | for medical | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|-------------------------------------|---------------------------------------|------------------|
| | | reasons | | | | |
| | | <u>Exclusion</u> | | | | |
| | | criteria: none | | | | |
| | | stated | | | | |
| | | Setting: hospital | | | | |
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| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|-----------------|-----------------|--------------------------|---------------------|------------------|-------------------------------------|---|
| Information | & Evidence | patients & | Characteristics | Reference | and NPV | |
| | level | prevalence | | standard | | |
| Staiano et al. | Study type: | 42 children | 42 children | Test: | Total gastrointestinal transit | Additional information from study: |
| Colonic transit | diagnostic | | | -Total | time (TGTT) (hours, mean ± | Severe brain damage: spastic |
| and anorectal | case control | <u>Inclusion</u> | Group1: children | gastrointestinal | <u>SD):</u> | tetraparesis/diplegia, generalised |
| manometry in | study | criteria: | with brain damage | transit time | | hypotonia |
| children with | | -patients: | N=16 | (TGITT) | -children with brain damage: | |
| severe brain | <u>Evidence</u> | children with | 10 boys | | 106.4 ± 6.1 | Children off all laxatives and/or |
| damage. 1994. | level: III | brain damage | Mean age 5.1 ± 3.5 | | | suppositories during the measurement |
| Pediatrics 94[2 | | referred for | years (range 1.5 to | segmental | -children with functional faecal | of total and segmental transit times |
| Pt 1], 169-173 | | gastroenterologi | 12 years) | gastrointestinal | retention (FFR): | |
| | | c evaluation of | | transit time | 98.6 ± 5.1 | Tracing coded and analysed by one of |
| | transit and | constipation | Group 2: children | (SGTT) | | the authors unaware of the clinical |
| | anorectal | -Controls: | with functional | | p value N.S | status of the child (not clear whether |
| | motility in | 1. functional | faecal retention | Reference | | this is CTT or manometry) |
| | | faecal retention: | | standard: | Segmental gastrointestinal | |
| | severe brain | | N=15 | None | transit time (SGTT): (mean, | Reviewers' comments: |
| | damage, | | 9 boys | | SEM) | 29 of the children originally undergoing |
| | looking for | normal | Mean age 6.0 ± 2.9 | | | evaluation for severe brain damage |
| | differences | frequency of | years (range 2 to | | Left colon: | were found to have constipation, but |
| | from | defecation and | 11 years) | | total number of markers at 48 | only 16 were included in the study. It is |
| | asymptomati | | | | h | not clear why the other 13 were |
| | c children | | Group 3: children | | -brain damaged: | excluded |
| | and from | 0 | with no | | 7.3 ± 1.3 | |
| | | disease | gastrointestinal | | | Functional faecal retention not defined |
| | functional | | <u>problems</u> | | -functional faecal retention | |
| | faecal | Exclusion | N=11 | | (FFR): | Exact values for all segmental transit |
| | retention and | | 7 boys | | 3.0 ± 1.0 | times in the 2 groups not reported |
| | normal | secondary | Mean age 5.6 ± 3.9 | | 0.05 | |
| | neurologic | constipation | years (range 2 to | | p< 0.05 | Source of funding: not stated |
| | development | | 12 years) | | | |
| | • | clinical | | | total number of markers at 72 | |
| | | interview, | Country: | | h: | |
| | | physical | Italy | | -brain damaged: | |
| | | examination, | | | 3.3 ± 0.8 | |
| | | barium enema, | | | functional faceal retention | |
| | | and anorectal | | | -functional faecal retention (FFR): | |
| | | manometry studies and/or | | | 0.5 ± 0.3 | |
| | | multiple suction | | | U.5 ± U.5 | |
| | | muniple suction | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|--|------------------|
| | | rectal biopsies | | | p<0.01 | |
| | | Setting: hospital | | | Distribution of markers in right colon and rectum not significantly different between the two groups | |
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| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|---|
| Zaslavsky et al. | Study type: | 61 adolescents | 61 adolescents | Test: | Colonic transit times patterns | Additional information from study: |
| Chronic | Diagnostic | | | Colonic transit | (N, %): | Radiographs interpreted by 2 of the |
| functional | case control | <u>Inclusion</u> | -Patients (n=48) | time (CTT) with | | authors (no further data provided) |
| constipation in | | <u>criteria:</u> | Mean age: 14 years | | Normal colonic transit: 8 (17) | |
| adolescents: | <u>Evidence</u> | -patients: aged | (range 12 to 18) | markers | Slow colonic transit: 29 (60) | Adolescents told to keep their usual diet |
| clinical findings | <u>level:</u> | 12 to 18 years, | 13 boys | | Pelvic floor dysfunction: 6 (13) | during examination and to discontinue |
| and motility | III | both sexes, | | Reference: | Slow colonic transit and pelvic | use of laxatives 7 days before |
| studies. 2004. | | normal sexual | -Controls (n=13) | Clinical variables | floor dysfunction: 5 (10) | examination |
| Journal of | | | 9 boys | | | |
| Adolescent | evaluate | | age not reported | | Total transit time (hours, mean | Patients underwent plain abdominal |
| Health 34[6], | symptoms | staging), <3 | | | ± SD, median and range) | radiography as per Metcalf method |
| 517-522 | and clinical | evacuations/we | | | | |
| | | | Brazil | | Constipated: | -Slow CTT: delay of total CTT and |
| | prospective | straining, | | | 62.9 ± 12.6 | delay of markers in the right and/or left |
| | series of | complaints for 1 | | | 69 (62.9 to 12.6) | colon |
| | adolescents | year or longer | | | | -Pelvic floor dysfunction: delay in the |
| | with | -controls: no | | | Non constipated: | rectosigmoid |
| | functional | digestive | | | 30.2 ± 13.2 | -Slow CTT associated with pelvic floor |
| | constipation | complaints, | | | 27.5 (10.8 to 50.4) | dysfunction: delay in the colon and |
| | and to identify | more than 3 bowel | | | p<0.001 | rectosigmoid together with delay in the total CTT |
| | colonic | movements/we | | | Segmental transit time (hours, | |
| | disorders by | ek | | | mean ± SD, range) | Cut-off points for measurements: mean |
| | measuring | (participated in | | | | value plus two SDs. Right colon (>14 |
| | total and | previous study | | | -Right colon: | h); left colon (>24h), rectosigmoid (>>36 |
| | segmental | by authors) | | | Constipated: | h) and total (>51 h) |
| | colonic | | | | 18.6 ± 15 | |
| | transit times | Exclusion | | | 13.2 (12 to 54) | Reviewers' comments: |
| | with | criteria: | | | · | Researchers not blinded |
| | radiopaque | neurologic/meta | | | Non constipated: | |
| | markers | bolic diseases, | | | 6.7 ± 3.9 | Cut-off points for total and segmental |
| | | Hirschsprung's | | | 4.8 (1.2 to 12) | transit times apparently taken from |
| | | disease (barium | | | P=0.001 | previous 1998 study by the authors |
| | | enema), spinal | | | | |
| | | disease, | | | -Left colon: | Source of funding: not stated |
| | | anorectal | | | Constipated: | - |
| | | anomalies, | | | 24.3 ± 13.7 | |
| | | surgery of the | | | 22.8 (2.4 to 51.6) | |

| Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|-----------------------------------|---|-------------------------------|-------------------------------------|--|------------------|
| | colon, mental retardation, use of drugs that act on digestive motility, no clinical evidence of bowel /systemic disease that could cause constipation Setting: hospital gastroenterolog y outpatients clinic | | | Non constipated: 7.9 ± 7.8 7.2 (0-28.8) P<0.001 -Rectosigmoid: Constipated: 20 ± 15.7 18 (0 to 54) Non constipated: 15.6 ± 10.7 12 (3.6 to 36) NS Interval between evacuations: -Slow colonic transit (n=29): 7.7 ± 6.6 days -Pelvic floor dysfunction (n=6): 3.7 ± 2.4 days p<0.003 Faecal mass palpable at initial examination statistically associated with slow colonic transit (p=0.03) Other clinical variables not statistically associated with delay in colon or rectosigmoid transit: onset of constipation, scybalous faeces, large volume, faecaloma, anal bleeding, soiling, previous use of | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---|------------------|
| | | | | | laxative/suppositories/enemas, history of constipation in family, anal fissure, daily ingestion of fibre, sex, age, skin colour | |
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| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|--------------------------|-----------------------------|-------------------------------|----------------------------|---------------------------------------|---|
| | level | prevalence | | standard | | |
| Koletzko et al. | Study type: | 48 children | 48 children | Test: | Total transit time (hours, mean | Additional information from study: |
| Is histological | Case series | | 25 boys | Colonic transit | <u>± SD)</u> | Hirschsprung's disease diagnosed in 9 |
| diagnosis of | (multicentre) | <u>Inclusion</u> | Mean age: 6.4 ± | time (CTT) with | | children excluded from further analysis |
| neuronal | | criteria: | 5.2 years | radiopaque | -Children with normal histology | |
| intestinal | <u>Evidence</u> | Initial symptoms | 0 | markers | (n=15): | Abortive neuronal intestinal dysplasia |
| dysplasia | <u>level:</u> III | of chronic | Country: Switzerland | Deference | 70.0 . 42.6 | (NID) and classic NID diagnosed in 17 |
| related to clinical and | Study aim: to | constipation or soiling, or | Switzeriand | Reference: | 70.0 ± 42.6 | and 6 patients respectively. |
| manometric | investigate | obstructive | | none | | Mean colonic transit times measured |
| findings in | the | symptoms in | | | | using the Metcalf method, in only 30 |
| constipated | relationship | early life | | | | children of the total population |
| children? | of clinical, | suggestive of | | | | ormanon or and total population |
| Results of a | manometric, | Hirschsprung's | | | | Reviewers' comments: |
| pilot study. | and | disease | | | | CTT results for children diagnosed with |
| 1993. Journal of | | | | | | abortive and classic NID not reported |
| Pediatric | findings in a | <u>Exclusion</u> | | | | for the purposes of this review as they |
| Gastroenterolog | | criteria: | | | | are considered organic causes of |
| , | | Anorectal | | | | constipation |
| 17[1], 59-65 | chronic | malformation or | | | | No data remente dan diat was af |
| | constipation in order to | mielomengonce le | | | | No data reported on diet, use of laxatives previous to the investigations |
| | evaluate the | ie . | | | | laxatives previous to the investigations |
| | role of | Setting: hospital | | | | Segmental transit times results not |
| | anorectal | Coung. Hoopital | | | | reported, and not clear whether they |
| | manometry | | | | | were measured |
| | in the | | | | | |
| | diagnosis of | | | | | Researchers not blinded |
| | neuronal | | | | | |
| | intestinal | | | | | Source of funding: not stated |
| | dysplasia | | | | | |
| | (NID) and | | | | | |
| | the | | | | | |
| | relationship of | | | | | |
| | histological | | | | | |
| | and | | | | | |
| | manometric | | | | | |
| | findings to | | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|--|---------------------------------|-------------------------------|---|---------------------------------------|------------------|
| | clinical severity of constipation and outcome | | | | | |
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| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|------------------|------------------|-----------------------|--------------------|--------------------|---------------------------------|---|
| Information | & Evidence level | patients & prevalence | Characteristics | Reference standard | and NPV | |
| Zaslavsky et al. | Study type: | 26 adolescents | 26 adolescents | Test: | Total transit time (hours, mean | Additional information from study: |
| Total and | Diagnostic | | aged 12-18 years | Colonic transit | ± SD, range) | No significant statistical differences |
| segmental | case control | Inclusion | Constipated (n=13) | time (CTT) with | -Constipated | between two groups regarding age, |
| colonic transit | | criteria: | Nonconstipated | radiopaque | 58.25 ± 17.46 | weight and height |
| time with radio- | Evidence | -patients: hard | (n=13) | markers | 68.4 (27.6 to 72) | |
| opaque markers | level: III | | 9 boys in each | | , | Total and segmental CTT measured |
| in adolescents | | in evacuating, | group | Reference: | -Non constipated | using Metcalf technique |
| with functional | Study aim: | less than 3 | | Clinical variables | 30.18 ± 13.15 | |
| constipation. | To measure | bowel | Country: | | 27.5 (10.8 to 50.4) | On the days the measurements were |
| 1998. Journal of | total and | movements/we | Brazil | | , , | performed adolescents were advised |
| Pediatric | segmental | ek, no evidence | | | P<0.001 | not to alter their diets and not to ingest |
| Gastroenterolog | colonic | of palpable | | | | food that might alter bowel motility. |
| y and Nutrition | transit time | rectal mass, | | | Segmental transit time (hours, | Fibre intake standardised at 15g/day |
| 27[2], 138-142 | in | history of | | | mean ± SD, range) | but due to poor compliance, test was |
| | constipated | constipation of | | | | performed on their normal diet. Any |
| | adolescents | at least one | | | -Right colon: | treatment with laxatives discontinued at |
| | and | year of duration | | | Constipated | least 7 days before test |
| | compared | -controls: no | | | 15.97 ± 12.48 | |
| | the results | digestive | | | 13.7 (2.4 to 43.2) | All radiographs interpreted by the same |
| | with those in | complaints, | | | | radiologist who did not know whether |
| | non | more than 3 | | | Non constipated | the patient was constipated |
| | constipated | bowel | | | 6.74 ± 3.91 | |
| | children | movements/we | | | 7.2 (1.2 to 12) | Patients with constipation considered to |
| | | ek | | | P=0.03 | have slow colonic transit when delay in |
| | | | | | | transit through the right colon, the left |
| | | <u>Exclusion</u> | | | -Left colon: | colon or both. They were considered to |
| | | criteria: | | | Constipated | have distal obstruction when the delay |
| | | neurologic/meta | | | 24.74 ± 13.39 | occurred in the rectosigmoid. |
| | | bolic diseases, | | | 25.7 (7.2 to 51.6) | |
| | | Hirschsprung's | | | | Normal values for total and segmental |
| | | disease, | | | Non constipated | transit times taken from the 95 th |
| | | spinal/anal | | | 7.94 ± 7.82 | percentile of adolescents without |
| | | anomalies, | | | 7.2 (0 to 28.8) | constipation |
| | | surgery of the | | | P<0.001 | |
| | | colon, mental | | | | Reviewers' comments: |
| | | retardation, | | | -Rectosigmoid: | Small sample size |
| | | history of drug | | | Constipated | |
| | | abuse | | | 17.60 ± 16.25 | Source of funding: not stated |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---|------------------|
| | | Setting: hospital | | | Non constipated 15.58 ± 10.69 12 (3.6 to 36) NS Interval between stools: -Constipated: 5.8 ± 2.3 days -Nonconstipated: Daily P<0.01 No significant differences between the 2 groups regarding: bulky or small stools, encopresis, rectal mass, intense use of laxatives, bowel movements/week and mean daily intake of fibres | |
| | | | | | mean daily intake of fibres | |

| Level Provalence Study type 225 children Lest: Abdominal Lest: Lest: Lest: Lest: Lest: Lest: Lest: Lest: | Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|--|---------------------------|-----------------------|----------------------|-------------------------------|----------------------------|---------------------------------------|--|
| usefulness of ultrasound (Case control Inclusion Care in Evidence method of assessment of constipation in chronic constipation in children 2007. Pediatric Radiology 37(12), 1247-1252 252 252 252 252 253 254 255 255 | | level | prevalence | | standard | | |
| lutrasound examination of the bowel as a method of assessment of functional chronic constitation in children. 2014. Study aim: Study aim: of stidior examination of the bowel as a method of assessment of functional chronic constitation in children. 2014. Pediatric Radiology 37[12], 1247-1252 1252 1252 1252 1252 1252 1252 1252 1252 1252 1252 1252 1252 1252 1252 1254 1255 1252 1254 1255 1252 1254 1252 | Bijos et al. The | Study type: | 225 children | 225 children | Test: | Mean colonic transit times: | |
| Exidence Evidence | | | | | | | |
| the bowel as a method of amethod of assessment of functional chronic constipation of functional children. 2007. Pediatric Radiology 37(12), 1247-1252 Table A sessment of functional children with a method of identifying children with functional chronic constipation, and to determine whether children with functional chronic constipation, and to determine whether children with functional chronic constipation, and to defect ation and a left colon (group 2: Group 2: Gro | ultrasound | Case control | | | ultrasound | | |
| method of sassesment of functional Scription in children 2007. Pediatric Radiology 37[12], 1247- 1252 Reseasement of inchildren with of stool retentine on could be used as a method of identifying children with functional schronic constipation, and to determine whether expective in the determine whether expective in the consistency and size of stool con (as gen on US) should be referred for further expective in the constipation in children with entition in children with an overfilled speak as a method of identifying children with an ocolon (as gen used to fix a session of the colon (as per use): (72 boys, mean age (a Colonic transit time (CTT) with radiopaque markers on US) that radiopaque markers on US (asoders assessment of sitool retention could be used as a method of identifying children with an ocolon (as session and to determine whether expension in could be used as a method of identifying children with an ocolon (as session and to determine whether expension identified the properties of the colon (as per us): (15) | | | | | | | |
| assessment of functional chronic functional chronic functional chronic constipation in constipation in constipation in constipation in children. 2014 Pediatric Radiology 37[12], 1247-1252 1247-1252 1 | | | | | | | even partially visible. |
| constipation in children. 2007. Pediatric whether a new method with readilology 37[12], 1247-1252 1252 1252 1252 | | | | | | | |
| Chronic constipation in children. 2007. Pediatric Radiology 37[12], 1247- 1252 1252 1252 1252 1252 1255 125 | | III | | | | | -Overfilled colon (as per US): |
| constipation in children. 2007. Pediatric new method Radiology 37/12J, 1247- 1252 | | | | 1.6 to 17.9) | | | |
| hildren. 2007. Pediatric Radiology 37[12], 1247- 1252 | | | | | | | |
| Pediatric Radiology 37[12], 1247- 1252 | | | | | | | |
| Radiology 37[12], 1247- 1252 Transit times disorders persisting of stool retention could be used as a method of identifying children with functional chronic constipation, and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further of the colon (as seen on US) should be referred for further of the colon (as sees and to for solor of stool retention of stool retention of stool retention of disorders disorders and the colon (as seen on US) should be referred for further of the colon (p=0.0029) Transit times (hours, upper limit (hours, upper limit transit transit transit transit transit transit transit transit transit constipation and to determine whether colon (p=0.0029) Transit times (hours, upper limit (hours, upper limit transit transit transit transit transit transit constipation and to determine whether colon (p=0.0029) Transit times (hours, upper limit transit transit transit transit transit transit transit constipation on US: 466: normal transit constipation on US: 661-100: slow-transit or constipation on US: 67 Total CTT (mean values are estimates taken from a bar chart) US: children examined before food and a slightly filled bladder. Patients without faecal impaction on US: 42 Patients with faecal impaction on US: 42 Patients without faecal impaction on US: 42 Patients without faecal impaction on US: 42 Patients without faecal impaction on US: 42 Po.001 Po.0 | | | | | | | |
| 37[12], 1247- (US) assessment of stool retention could be used as a method of identifying children with functional and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further | | | | | | | |
| assesment of stool retention could be used as a method of identifying children with functional chronic and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further of the first and to five the first and the colon (p=0.0029) Total CTT (mean values are estimates taken from a bar chart) Total CTT (mean values are estimates taken from a bar chart) US: children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 67 Patients without faecal impaction on US: 42 Total and segmental colonic transit time massurement the colon (p=0.0029) Total CTT (mean values are estimates taken from a bar chart) Total CTT (mean values are estimates taken from a bar chart) Total CTT (mean values are estimates taken from a bar chart) Total CTT (mean values are estimates taken from a bar chart) Total CTT (mean values are estimates taken from a bar chart) Total CTT (mean values are estimates taken from a bar chart) Total CTT (mean values are estimates taken from a bar chart) Total CTT (mean values a | | | | | | | |
| of stool retention could be patients fulfilled used as a method of identifying children with functional chronic and to determine whether size of stool children with an enlarged rectum and colon (as seen on US) should be referred for further of the first of the stool of identifying children with the free firered for the fre | | ` ' | | 8.25 years) | | | |
| retention could be used as a method of identifying children with functional chronic constipation, and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further of the function of the fu | 1252 | | | _ | | | spleen. |
| could be used as a method of identifying children with an enlarged recturn and colon (as seen on US) should be referred for further could be used as a method of identifying children with an enlarged recturn and colon (as seen on US) should be referred for further could be used as a method of identifying children with an enlarged recturn and to used as a method of identifying children with an enlarged referred for further could be used as a method of identifying children with an enlarged referred for further could be used as a method of identifying children with an enlarged referred for further could be used as a method of identifying children with an enlarged referred for further could be used as a method of identifying children with an enlarged referred for further could be used as a method of identifying children with an enlarged referred for further could be used as a method of identifying children with an enlarged referred for further could be used as a method of identifying children with an enlarged referred for further could be used as a method of identifying children with an enlarged referred for further could be used as a method of identifying the surprise taken from a bar chart): constipation (mean values are estimates taken from a bar chart): constipation (mean values are estimates taken from a bar chart): constipation (mean values are estimates taken from a bar chart): constipation (mean values are estimates) taken from a bar chart): constipation (mean values are estimates) taken from a bar chart): constipation (mean values are estimates) taken from a bar chart): constipation (mean values are estimates) taken from a bar chart): constipation (mean values are estimates) taken from a bar chart): constipation (mean values are estimates) taken from a bar chart): constipation (mean values are estimates) taken from a bar chart): constipation (mean values are estimates) taken from a bar chart): constipation (mean values are estimates) taken | | | | | <u>literature)</u> | the colon (p=0.0029) | |
| used as a method of identifying children with functional chronic constipation, and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further US children with functional chronic constipation (frequency of bowel transit constipation disorders taken from a bar chart): Total CTT (mean values are estimates taken from a bar chart): US children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 67 VS children examined before food and had a slightly filled bladder. Patients who passed stool on the day of the examination were temporarily excluded from the study until they became delayed slow-transit constipation VS children examined before food and had a slightly filled bladder. Patients with out faecal impaction on US: 67 VS children examined before food and had a slightly filled bladder. Patients with out faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with out faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients with faecal impaction on US: 42 VS children examined before food and had a slightly filled bladder. Patients wit | | | | Poland | | | |
| method of identifying children with functional chronic constipation, and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further of the constipation identifying children with functional chronic constipation (alternative children with functional chronic chronic constipation) and to determine with determine children with an enlarged rectum and colon (as seen on US) should be referred for further on the constipation identifying children with identification (alternative consistency and size of stool caused pain during defectation, withholding behaviour) and the children with the constipation on the day of the examination were temporarily excluded from the study until they became constipation on US: 42 constipated again. Measurement was taken several times and the highest one constipation on US: 42 constipated again. Measurement was taken several times and the highest one constipation on US: 42 constipated again. Measurement was taken several times and the highest one constipation on US: 42 constipated again. Measurement was taken several times and the highest one constipation on US: 42 constipated again. Measurement was taken from a bar chart) Segmental CTT (mean values are estimates taken from a bar chart) Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental immediate part of radiopaque markers seen on the constipation on US: 67 Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental colonic transit time obtained by multiplying the number of radiopaque markers seen on the constipation on US: 67 Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental colonic transit time obtained by multiplying the number of radiopaque markers seen on the constipation on US: 67 Total colon the day of the examination on US: 67 Total and segmental colonic transit time measured by the modified sixth day Hinton | | | • | | | | |
| identifying children with functional chronic movements less constipation, and to determine children with an enlarged rectum and colon (as seen on US) should be referred for further identifying children with functional chronic (frequency of bowel transit (chronic movements less constipation), and to week, determine whether consistency and whether seen on US) should be referred for further identifying children with functional bowel (frequency of bowel transit (constipation on US: 67 -Patients with faecal impaction on US: 42 -Patients without faecal from the study until they became examination were temporarily excluded from the study until they became delayed slow-transit constipation on US: 42 constipation on US: 42 constipated again. Measurement was taken several times and the highest one recorded taken as the final measurement Segmental CTT (mean values are estimates taken from a bar chart) Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental time obtained by multiplying the number of radiopaque markers seen on the | | | | | | | |
| children with functional chronic chronic chronic constipation, and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further Children with functional functional chronic chron | | | | | | , | the sagital plane over the aorta |
| functional chronic constipation, and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further functional chronic constipation and to determine children with an enlarged refurther further functional chronic constipation and to determine week, consistency and size of stool caused pain during defectation, withholding behaviour) further furth | | | | | | taken from a bar chart): | l., .,, |
| chronic constipation, and to determine determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further chronic constipation, and to week, consistency and size of stool cand to determine determine was than twice a week, consistency and size of stool cand to week, consistency and size of stool caused pain defecation, withholding behaviour) colon (as seen on US) should be referred for further children with an enlarged rectum and colon (as seen on US) should be referred for further children with an enlarged rectam and colon (as seen on US) should be referred for further children with an enlarged rectam and colon (as seen on US) should be referred for further constipation con US: 67 who passed stool on the day of the examination were temporarily excluded from the study until they became constipated again. Measurement was taken several times and the highest one recorded taken as the final measurement Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental time obtained by multiplying the number of radiopaque markers seen on the | | | | | | | |
| constipation, and to determine week, consistency and whether children with an enlarged rectum and colon (as seen on US) should be referred for further criteria: Constipation, and to week, consistency and size of stool caused pain defectation, withholding behaviour) Constipation | | | | | | | |
| and to determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further and to determine determine whether children with an enlarged rectum and colon (as seen on US) should be referred for further and to determine week, consistency and size of stool caused pain during defecation, withholding behaviour) - Patients without faecal impaction on US: 42 - Pa | | | | | constipation | on US: 67 | |
| determine whether size of stool caused pain an enlarged rectum and colon (as seen on US) should be referred for further citeria: delayed slow-transit constipated again. Measurement was taken several times and the highest one recorded taken as the final measurement delayed slow-transit constipated again. Measurement was taken several times and the highest one recorded taken as the final measurement Segmental CTT (mean values are estimates taken from a bar chart) Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental time obtained by multiplying the number of radiopaque markers seen on the | | | | | . 400 | Detients with sort for sel | |
| whether children with an enlarged rectum and colon (as seen on US) should be referred for further or iteria: whether children with an enlarged rectum and colon (as seen on US) should be referred for further children with an enlarged rectum and colon (as seen on US) should be referred for further children with an enlarged rectum and colon (as seen on US) should be referred for further children with an enlarged rectum and colon (as seen on US) should be referred for further children with an enlarged rectum and colon (as withholding behaviour) transit constipation p<0.001 Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental time obtained by multiplying the number of radiopaque markers seen on the | | | , | | | | |
| children with an enlarged rectum and colon (as seen on US) should be referred for further criteria: children with an enlarged rectum and colon (as seen on US) should be referred for further children with an enlarged rectum and colon (as seen on US) should be referred for further colon (as seen on US) should be referred for further colon (as withholding behaviour) constipation p<0.001 recorded taken as the final measurement Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental 1. Right colon -Patients with faecal impaction of radiopaque markers seen on the | | | , | | | impaction on US: 42 | |
| an enlarged rectum and colon (as seen on US) should be referred for further further defecation, colon (as rectum and colon (as seen on US) and segmental colonic transit time seen on the segmental colonic transit time (mean values are estimates (mean values are estimates) taken from a bar chart) measured by the modified sixth day Hinton method. Total and segmental time obtained by multiplying the number of radiopaque markers seen on the | | | | | | n =0 001 | |
| rectum and colon (as seen on US) should be referred for further criteria: Total and segmental colonic transit time taken from a bar chart) Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental time obtained by multiplying the number of radiopaque markers seen on the | | | | | Consupation | p<0.001 | |
| colon (as seen on US) behaviour) should be referred for further colon (as seen on US) behaviour) (mean values are estimates taken from a bar chart) (mean values are estimates taken from a bar chart) Total and segmental colonic transit time measured by the modified sixth day Hinton method. Total and segmental 1. Right colon time obtained by multiplying the number of radiopaque markers seen on the | | | | | | Commental CTT | measurement |
| seen on US) behaviour) should be referred for further seen on US) behaviour) taken from a bar chart) taken from a bar chart) measured by the modified sixth day Hinton method. Total and segmental time obtained by multiplying the number -Patients with faecal impaction of radiopaque markers seen on the | | | | | | | Total and cogmental colonic transit time |
| should be referred for further criteria: Hinton method. Total and segmental time obtained by multiplying the number -Patients with faecal impaction of radiopaque markers seen on the | | ` | | | | , | |
| referred for further criteria: 1. Right colon time obtained by multiplying the number -Patients with faecal impaction of radiopaque markers seen on the | | | Dellavioul) | | | lancii IIOIII a Dal Cilait) | |
| further criteria: -Patients with faecal impaction of radiopaque markers seen on the | | | Evolusion | | | 1 Right colon | |
| | | | | | | | |
| | | | | | | | |
| such as abnormality hours/number of markers swallowed by | | ļ • | | | | 00.0 | |

| Bibliographic Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---|---------------------------------|-------------------------------|---|--|--|
| | | | | -Patients without faecal impaction on US: 8 N.S 2. Left colon -Patients with faecal impaction on US: 18 -Patients without faecal impaction on US: 9 p=0.0104 3. Rectosigmoid: -Patients with faecal impaction on US: 32 -Patients without faecal impaction on US: 16 p=0.0015 | the patient) Reviewer's comments: No data on diet or use of laxatives previous to the measurement of CTT The same individual performed all the US scans, but not other data on this were reported (as blinding, individual's experience in radiology, etc) It is not clear what number of children underwent each of the tests It is not clear whether "enlarged" and "overfilled" colon mean the same for the authors, as no measurements of "enlarged" colon are reported. Data on number of children diagnosed with "overfilled colon" are not reported. It is not clear how many children were diagnosed with faecal impaction by US Children apparently underwent DRE but no results are reported Control group did not differ from patients regarding gender, the comparison regarding age is not clearly reported Source of funding: Not stated |
| | food allergies) | | | | |

| Bibliographic Information | Study type & Evidence level | prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---|-------------------------------|-------------------------------------|---------------------------------------|------------------|
| | | Setting: gastroenterolog y outpatient clinic | | | | |
| | | | | | | |
| | | | | | | |
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| | | | | | | |

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|-----------------|------------------------------|--------------------------|---------------------|--------------------|------------------------------------|---|
| Information | & Evidence level | patients & prevalence | Characteristics | Reference standard | and NPV | |
| | Study type: | 89 non selected | 89 children | Test: | Mean Leech score (using the | Additional information from study |
| | Diagnostic. | consecutive | | Plain abdominal | first score): | Diagnosis of functional non-retentive |
| method for | Case control | children | Median age: 9.8 | radiography (read | | faecal incontinence (FNRFI) based on: |
| diagnosing | | | years | using the Leech | -Group 1 (constipation): 10.1 | two/more faecal incontinence |
| constipation: | <u>Evidence</u> | <u>Inclusion</u> | _ | method) | -Group 2 (controls): 8.5 | episodes/week with no signs of |
| intra- and | <u>level</u> : | criteria: patients | | | | constipation 2) defecation frequency |
| interobserver | III | referred for the | (constipation): | Reference test: | p=0.002 | 3/more times/week 3) no periodic |
| variability and | | evaluation of | n=52 (28 boys) | Colonic transit | | passage of very large amounts of stool |
| | Study aim: | abdominal pain, | | time (CTT) with | Mean CTT: | at least once during a period of 7-30 |
| | to assess | | Group 2 (controls): | radiopaque | -Group 1 (constipation): 92 h | days 4) no palpable abdominal or rectal |
| 0, 1, | intra- and | | N=37 (24 boys) | markers | -Group 2 (controls): 37 h | mass on physical examination fro a |
| 43-49 | | incontinence. | 04 511551 | | 0.0004 | period of at least 1 week during the |
| | variability | 0 | 31: FNRFI | | p<0.0001 | preceding 12 weeks. Faecal |
| | and | | 6: FAP | | S | incontinence defined as the |
| | determine | least two of the | | | Diagnostic accuracy of Leech | voluntary/involuntary loss of loose |
| | diagnostic | following was | Country: | | method vs. CTT method: | stools in the underwear after the age of |
| | accuracy of | present: | the Netherlands | | l and a mark and | 4 years |
| | the Leech | -defecation | | | -Leech method: | Functional abdominal pain (FAP) |
| | method in | frequency less than 3 | | | (cut-off point as per study | defined as abdominal pain of at least 12 |
| | identifying children with | times/week | | | comparable to 9 as per literature) | weeks duration 1)that was continuous or nearly discontinuous in a school- |
| | functional | -2/more | | | Sensitivity: 75% | aged child or adolescent 2) that had no |
| | constipation | episodes of | | | Specificity: 59% | or only an occasional relationship with |
| | Constipation | faecal | | | Specificity : 59 % | physiological events 3) that was |
| | | incontinence | | | (cut-off point 9 as per | accompanied by some loss of daily |
| | | per week | | | literature) | functioning 4) that was not feigned and |
| | | -production of | | | Positive Predictive Value: 72% |) for which there were insufficient |
| | | large amounts | | | Negative Predictive Value: | criteria to indicate the presence of |
| | | of stool once | | | 63% | another functional gastrointestinal |
| | | over a period of | | | 00 70 | disorder |
| | | 7-30 days | | | -CCT: | Children with clinical characteristics of |
| | | -presence of | | | (cut-off point 54h as per study) | FAP and FNRFI classified as the |
| | | palpable | | | Sensitivity: 79% | control group: according to authors they |
| | | abdominal or | | | Specificity: 92% | have "little or no faecal loading on an |
| | | rectal mass | | | | abdominal radiograph" |
| | | | | | (cut-off point 62h as per | |
| | | (control children | | | literature) | Treatment with oral/rectal laxatives |
| | | fulfilled criteria | | | Sensitivity: 71% | discontinued in each patient for at least |

| Information & Ev | dy type Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|------------------|---|-------------------------------|-------------------------------------|---|---|
| | for functional abdominal pain (FAP) and for functional non-retentive faecal incontinence (FNRFI)) Exclusion criteria: not reported Setting: tertiary gastroenterolog y outpatients clinic | | | Specificity: 95% Positive Predictive Value: 69% Negative Predictive Value: 97% ROC analysis -AUC (Leech method): 0.68 (95% CI 0.58-0.80) -AUC (CTT method): 0.90 (95% CI 0.83-0.96) p=0.00015 AUC=Area Under the ROC curve ROC=Receiving Operator Characteristic | 4 days. Thereafter the patient ingested one capsule with 10 small radiograph opaque markers on 6 consecutive days, in order to determine the CTT. Subsequently, a plain abdominal radiograph was taken on day 7. this radiograph was both used in the Leech method and for CTT measurement CTT determined by the method of Bouchoucha. Radiography on day 7 used to count the number of markers in the colon. Number of markers x 2 produced total CTT in hours. Localization of markers and CTT calculated according to previously described formula. Normal range for total transit time based on the upper limits (mean ± 2xSD) from a study in healthy children. Based on this study a CTT > 62 h was considered delayed 3 scorers independently scored the same radiography twice (4 weeks apart) using the Leech method, discussed amongst the 3 scorers previous to both readings CTT assessed once by single scorer. Assumed the counting of radiopaque markers would not lead to intra- or interobserver variability Leech scoring method: Colon divided into three segments: right, left and recto sigmoid Each segment provided with a score from 0-5 0:no faeces visible |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|---|
| | | | | | | 1:scanty faeces visible |
| | | | | | | 2: mild faecal loading |
| | | | | | | 3: moderate faecal loading |
| | | | | | | 4: severe faecal loading 5: severe faecal loading with bowel |
| | | | | | | dilatation |
| | | | | | | Leech score of 9 or more: suggestive or constipation |
| | | | | | | Scorers: 3 experienced doctors (a 5 th year radiology resident, a paediatric radiologist and a senior paediatric |
| | | | | | | gastroenterologist). No clinical information about the patients was made available to them. |
| | | | | | | In 5% of cases the Leech scores of the same patient produced by different scorers could differ by 4 points or more |
| | | | | | | Reviewer's comments: |
| | | | | | | No data reported on type of diet given prior to the measurement of CTT |
| | | | | | | Source of funding: not stated |

Radioisotopes Markers

| | & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|-----------------------|------------|----------------------------------|-------------------------------|--------------------------------|---|--|
| | level | prevalence | | standard | | |
| | | 101 consecutive | 101 children | Test: | Mean sum of GC for the 4 | Additional information from study: |
| | | nuclear transit | | Colonic transit | imaging periods (mean ± SD, | Four imaging periods: 6, 24, 30 and 48h |
| transit to assess ref | | | 62 boys | time (CTT) with | range) | |
| | | on children with | | radioisotopes | | Intake of laxatives stopped 5 days |
| large bowel | | | Mean age 7.3 ± | | 1-Normal transit time (n=24): | before the transit time and patients |
| | | • | 3.7 years | <u>Reference</u> | | fasted for 4 h before start of test. Rectal |
| | | over a 2-year | | Standard: | 15.7±3.3 (7.3-19.1) | disimpaction not carried out before |
| chronic | | | Country: | None stated | | study in any patient. |
| | Study aim: | | Australia | | 2-SCT (n=50): | Radiopharmaceutical technetium 99m- |
| | | <u>Inclusion</u> | | Three categories | | calcium phytate colloid, suspended in |
| 2005. Journal of the | | <u>criteria:</u> | | of colonic transit | 11.2±1.9 (7.5-16.3) | 20mL of milk was administered by |
| | | All patients | | according to | | mouth. |
| | | seen by the | | visual | p<0.001 as compared to | |
| | | senior author or | | assessment | normal transit time and FFR | A nuclear medicine radiologist from the |
| | | a | | -Normal transit | groups | hospital performed qualitative visual |
| | | gastroenterologi | | time: tracer | | assessment of the images acquired at |
| _ | | st paediatrician. | | reached the | 3-FFR (n=22): | each time interval. Colonic transit times |
| | | All had | | caecum by 6 | 45 4 4 5 (40 7 40 0) | was estimated by analysis of the |
| | | symptoms of . | | hours, passed | 15.1±1.5 (12.7-18.2) | images acquired between 6 and 48 |
| | | severe chronic | | through the colon | 45 4 5 | hours |
| | | constipation | | | 4-Borderline (n=5) | |
| | | and/or | | excreted by 6 | not reported | Geometric centre (GC): six regions of |
| | | encopresis that | | hours | | interest were defined: |
| | | had not | | Class salania | GC at each of the 4 imaging | 1-precolonic region |
| | | responded to at least six months | | -Slow colonic transit time | periods (mean ± SD, range) | 2-caecum and ascending colon as far |
| | - 3 | of medical | | | 1 Normal transit time (n. 24). | as the hepatic flexure 3-transverse colon from hepatic to |
| | | | | (SCT): when the tracer reached | 1-Normal transit time (n=24): | splenic flexure |
| | | therapy with laxatives, | | | 6h: 2.0±0.5 (1-3.5) 24h: 3.9±1.1 (1-5.9) | 4- descending colon from splenic |
| CII | | dietary | | hours but most | 30h: 4.6±1.2 (2-5.9) | flexure to start of sigmoid |
| | | alterations and | | | 48h: 5.2±0.9 (2.3-6) | 5-sigmoid colon |
| | | behaviour | | retained in the | 46 . 5.2±0.9 (2.3-6) | 6-faeces |
| | | modification | | | 2-SCT (n=50): | GC refers to the median point of the |
| | | mounication | | | 6h: 1.8±0.3 (1-2.5) | distribution of activity within the colon. It |
| | | Exclusion | | | 24h: 2.6±0.5 (1.9-4.4) | was calculated by multiplying the |
| | | criteria: | | | 30h: 3.1±0.6 (1.8-4.5) | fraction of the administered activity in a |
| | | Obviously | | retention/outlet | 48h: 3.7±0.9 (1.9-5.7) | region, by a region number and the 6 |
| | | palpable | | obstruction | 1011. 0.7 ±0.0 (1.0 0.7) | numbers for each image episode were |

| Bibliographic | Study type | Number of | Population | Type of test and | Sensitivity, Specificity, PPV | Reviewer comment |
|---------------|------------|-----------------|-----------------|-------------------|----------------------------------|--|
| Information | & Evidence | patients & | Characteristics | Reference | and NPV | |
| | level | prevalence | | standard | | |
| | | faecaloma in | | | p<0.05 at 6h and p<0.001 at | added |
| | | rectum or | | | 24, 30 and 48 h, as compared | |
| | | sigmoid colon. | | | to normal transit and FFR | Reviewers' comments: |
| | | Anorectal | | 24 to 30 h but | groups | No control group, or comparison with a |
| | | malformation, | | was not passed | | reference test |
| | | spinal | | | 3-FFR (n=22): | |
| | | deformity, | | | 6h: 2.0±0.4 (1.2-3) | Not clear definition of constipation |
| | | Hirschsprung's | | | 24h: 3.6±0.7 (2.5-5) | reported |
| | | disease, bowel | | according to | 30h: 4.4±0.5 (3.5-5.4) | |
| | | washout or | | | 48h: 5.1±0.3 (4.4-5.7) | No diagnosis prior to the application of |
| | | enema in the | | functional | | the test was made |
| | | week before | | | 4-Borderline (n=5) | |
| | | study to remove | | | not reported | Researchers not reported blinded |
| | | faecaloma | | trough the colon" | | |
| | | _ | | | No significant difference in the | Source of funding: Not stated |
| | | Setting: | | | GC at any imaging time when | |
| | | continence | | | comparing patients with normal | |
| | | clinic | | | transit with those with FFR. | |
| | | | | | | |
| | | | | | Two of the 101 children (not | |
| | | | | | clear in which group) had a GC | |
| | | | | | of 1.0 at 6 h indicating that | |
| | | | | | 100% of the tracer was located | |
| | | | | | in the small bowel, suggesting | |
| | | | | | impairment. | |
| | | | | | | |
| | | | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|--|--|---|--|--|--|---|
| Vattimo et al. Total and segmental colon transit time in constipated children assessed by scintigraphy with 111In- DTPA given orally. 1993. Journal of Nuclear Biology and Medicine 37[4], 218-222 | who underwent total and segmental | Inclusion criteria: Constipation defined as 2 or fewer bowels motions/week or straining for more than 25% of the defecating time Exclusion criteria: Normal children (no other details given) Setting: unclear, but children were outpatients | 39 children 23 females Age range: 2-13 years Country: Italy | Test: Colonic transit time (CTT) with radioisotopes Reference test: none reported | Total transit time (hours, mean ± SD) -Normal transit time (n=13) 27.79 ± 4.10 -Mainly rectosigmoid retention (n=5) 53.36 ± 29.66 -Prolonged transit time in all segments (n=14) 62.09 ± 7.23 -More prolonged transit time in rectosigmoid tract (n=7) 92.36 ± 24.16 Segmental transit time (hours, mean ± SD) -Normal transit time (n=13) Right colon: 9.11 ± 2.53 Left colon: 9.80 ± 3.50 Rectosigmoid: 8.88 ± 4.09 -Mainly rectosigmoid retention (n=5) Right colon: 10.38 ± 2.34 Left colon: 10.40 ± 4.00 Rectosigmoid: 32.58 ± 29.64 -Prolonged transit time in all | Additional information from study: -RC: right colon from caecum to midtransverse -LF: left colon from mid-transverse to descending colon-sigmoid junction -RS: rectosigmoid from the sigmoid junction to rectum From the point of view of radiation dosimetry the most heavily irradiated organs were the lower large intestine and the ovaries and the level of radiation burden depended on the colon transit time Reviewers' comments: No data reported on diet or use of laxatives previous to the measurement of CTT It is unclear whether the children suffered from severe/intractable constipation. Otherwise if might be difficult to justify this study No data on the researchers or their performance was reported Results for children with dolichocolon (n=7) not reported as this would be secondary constipation Source of funding: not stated |
| | | | | | segments (n=14) | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|--|------------------|
| | | | | | Right colon: 21.81 ± 5.29 | |
| | | | | | Left colon: 23.32 ± 6.14 | |
| | | | | | Rectosigmoid: 16.95 ± 4.52 | |
| | | | | | -More prolonged transit time in rectosigmoid tract (n=7) Right colon: 19.78 ± 9.03 | |
| | | | | | Left colon: 21.05 ± 5.70 Rectosigmoid: 51.53 ± 17.82 | |
| | | | | | Interval between defecations: (hours, mean ± SD) | |
| | | | | | -Normal transit time (n=13) 23.38 ± 5.42 | |
| | | | | | -Mainly rectosigmoid retention (n=5) 35.60 ± 14.54 | |
| | | | | | -Prolonged transit time in all segments (n=14) 53.00 ± 15.97 | |
| | | | | | -More prolonged transit time in rectosigmoid tract (n=7) 85.71 ± 32.25 | |
| | | | | | | |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|---------------------------------------|---|
| Shin et al. Signs | Study type: | 180 children | 180 children | Test: | FFR (n=19) | Additional information from study: |
| and symptoms | Retrospectiv | | 92 boys | Colonic transit | STC (n=161) | Clinical stories reviewed retrospectively |
| of slow-transit | e case | Inclusion | Mean ages: 10.5 | time (CTT) with | , , | and augmented by interview or |
| constipation | series | criteria: | years (STC); 6 | radioisotopes | FFR vs. SCT | questionnaire |
| versus | | Severe, | years (FFR) | | (Clinical variables (%)) | |
| functional | Evidence | intractable | , , | Reference: | | No gender differences between both |
| retention. 2002. | level: III | constipation | Country: | -Clinical variables | -Constipation: 89 vs.91 | groups |
| Journal of | | which did not | Korea & Australia | -Stool | -Soling: 42 vs.64 | |
| Pediatric | | respond to at | | characteristics | -Bloating: 26 vs. 46 | Normal CCT defined as the presence of |
| Surgery 37[12], | Study aim: | least 6 months | | | -Abdominal pain: 42 v. 51 | tracer in the caecum by 6 h, in the |
| 1762-1765 | to correlate | of medical | | | -Anal pain: 16 vs. 19 | rectosigmoid by 30 h and passed in the |
| | symptoms, | therapy | | | -Vomiting: 7 vs. 16 | faces by 48h. Slow CCT defined as |
| | signs, transit | instituted by a | | | -Failed toilet training: | global colonic delay with hold-up of |
| | times and | general | | | -Poor appetite: 42 vs. 22 | tracer proximal to the rectosigmoid at |
| | immunohisto | practitioner or | | | -Behavioural problems: 21 vs. | 30 and 48 h (with no rectal faecaloma). |
| | chemistry to | paediatrician | | | 22 | FFR identified by hold-up of tracer |
| | determine | | | | -Prematurity: 6 vs. 5 | proximal to the rectosigmoid at 48 h |
| | the | <u>Exclusion</u> | | | -Meconium passage > 24 after | preceded by normal transit |
| | diagnostic | criteria: | | | birth: 41 vs. 33 (35% unknown) | |
| | differences | None reported | | | -Family history of constipation: | Visual inspection of collected |
| | between | | | | 61 vs. 52 | radiographic images augmented by use |
| | slow transit | Setting: unclear | | | -Constipation present at birth: | of a "colonic transit index" (sum of the |
| | constipation (STC) and | | | | 11 vs. 26 (p=0.17) | geometric centres of radioactivity at 6, 24, 30 and 48 h) |
| | functional | | | | (p values not reported are not | |
| | faecal | | | | significant) | Normal values for CTT derived from |
| | retention | | | | | several studies of transit time in healthy |
| | (FFR) | | | | FFR vs. SCT | children |
| | | | | | (Stool characteristics (%)) | |
| | | | | | | Slow-transit constipation, STC: slow |
| | | | | | -Volume: | transit through the colon |
| | | | | | Small moderate: 68 vs. 47 | FFR: chronic constipation caused by |
| | | | | | Large: 26 vs. 52 | delay of anorectal release |
| | | | | | Not known: 5 vs. 2 | |
| | | | | | | Reviewers' comments: |
| | | | | | -Consistency: | Exclusion criteria not reported |
| | | | | | Hard/firm: 78 vs. 58 | |
| | | | | | Soft/variably soft: 16 vs. 39 | Questionnaires not piloted. No data on |

| Bibliographic Information | Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|-----------------------------------|---------------------------------|-------------------------------|---|--|--|
| | | | | | (p<0.001) Not known: 5 vs. 3 -Frequency: >1 week: 56 vs. 40 1/week: 26 vs. 22 <1 week: 11 vs. 28 Not known: 5 vs. 10 | intrarater/interrater reliability No data on diet or use of laxatives previous to CTT No data of individual(s) performing readings: blinding, etc. Actual figures for CTT not reported Source of funding: Not reported |

| Bibliographic Information | Study type & Evidence | Number of patients & | Population Characteristics | Type of test and Reference | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|---------------------------|---------------------------|----------------------------------|-------------------------------|----------------------------|---|---|
| Chitkara et al. | level | prevalence 67 adolescents | 67 adolescents | standard | Colonia transit time (n. 44) | Additional information from study |
| The role of | Study type: Diagnostic | or addlescents | Mean age: 14.7± | Test: Colonic transit | Colonic transit time (n=41) (FC=12; FFR=8; C-IBS=21) | Additional information from study: Patients were classified in three groups |
| pelvic floor | retrospective | <u>Inclusion</u> | 3.3 yr | time (CTT) with | (FC=12, FFR=6, C-1B3=21) | according to paediatric Rome II criteria |
| | case series | criteria: | 67% female | radioisotopes | -Geometric centre at 24 h | based on the symptoms and diagnoses |
| slow colonic | 0430 301103 | -constipation | 07 70 ICITIAIC | radioisotopes | Total: 2.03 ± 0.99 | provided by the clinician who evaluated |
| transit in | Evidence | unresponsive to | Group 1: | | FC: 1.73 ± 0.29 | the patient prior to the ARM and BET |
| adolescents | level: III | first line, | (n=16) Functional | Reference tests: | FFR: 2.04 ± 0.38 | |
| with refractory | | symptomatic | constipation (FC) | -Clinical variables | | Patients instructed to discontinue all |
| constipation. | Study aim: | treatments | (1, | (nausea, | -Slow colonic transit (%) | medications known to affect intestinal |
| 2004. American | to examine | - completion of | Group 2: | vomiting, | Total: 30 | motility 48 h prior to study. Patients |
| Journal of | the | clinically | (n=18) Functional | bloating, weight | FC: 42 | given the radioisotope after overnight |
| Gastroenterolog | | indicated ARM | faecal retention | loss and | FFR: 14 | fast |
| | and pelvic | and BET for | (FFR) | incomplete rectal | | |
| 1584 | floor function | the evaluation | | evacuation) | -Fast colonic transit (%) | A geometric centre at 24h of ≤ 1.6 was |
| | by anorectal | of constipation | Groups 3: | | Total: 7.5 | classified as slow colonic transit and > |
| | manometry | -age ≤ 18 yr | (n=33) | | FC: 0 | 3.8 considered fast colonic transit. |
| | (ARM) and | -able and willing | | | FFR: 0 | D. t |
| | balloon | to follow | predominant | | No significant appointing of | Reviewers' comments: |
| | expulsion | instructions in the balloon | irritable bowel | | No significant association of abnormal GC at 24h (fast or | Methodology poorly described. Researchers not reported blinded. |
| | test (BET) in adolescents | expulsion study | syndrome IBS(C-IBS) | | slow) and individual | Intrarater/interrater reliability |
| | ≤ 18 years of | | 163) | | gastrointestinal symptoms (no | measurements not reported |
| | age referred | experienced | Country: | | further details reported) | lineasurements not reported |
| | to a tertiary | test operator | USA | | luttiei details reported) | Only 61% of total sample underwent |
| | care centre | -presence of | 00/1 | | | colonic transit time, but not clear |
| | for | gastrointestinal | | | | explanation for this |
| | symptoms of | complaints in | | | | ' |
| | refractory | the absence of: | | | | Not clear on what basis the cut off |
| | constipation, | | | | | points for the geometric centre were |
| | and to | <u>Exclusion</u> | | | | determined |
| | describe the | criteria: | | | | |
| | results of | colonic | | | | Insufficient data to allow calculation of |
| | scintigraphic | resection or | | | | other parameters of diagnostic value of |
| | colonic | systemic | | | | CTT (Sensitivity, Specificity, PPV and |
| | transit | organic disease | | | | NPV) |
| | measuremen | | | | | Depute for CIDC noticets and remarks |
| | ts in the | mellitus, | | | | Results for C-IBS patients not reported, |
| | patients wno | hypothyroidism, | |] | | as population outside the remit of this |

| Bibliographic Study type & Evidence level | Number of patients & prevalence | Population Characteristics | Type of test and Reference standard | Sensitivity, Specificity, PPV and NPV | Reviewer comment |
|--|--|-------------------------------|---|---------------------------------------|---|
| also munderwent el this test re lo de H di | nielomeningoc ele, mental etardation/deve opmental lelay, dirschsprung's lisease) Setting: tertiary are centre | | | | guideline Source of funding: In part by the GlaxoSmithKline Institute of Digestive General Research Award to D. Chiktara ad NIH grants to M. Camilleri |

Pharmacological and Surgical Interventions for Disimpaction in Children with Chronic Idiopathic Constipation

| Candy et al. Treatment of faecal impaction with polyethelene glycol plus electrolytes (PGE + E) followed by a double-blind Candy et al. Study Type: Prospective case series (phase 1 of the study)* Evidence followed by a double-blind Study Type: Study Type: Prospective case series (phase 1 of the study)* Study Type: Prospective case series (phase 1 of the stu | nents |
|--|-------------------|
| Treatment of faecal impaction with polyethelene glycol plus electrolytes (PGE + E) followed by a faecal impaction with faecal impaction with polyethelene glycol plus electrolytes (PGE) followed by a faecal impaction of faecal impaction with polyethelene glycol as series (phase 1 of the study)* Treatment of faecal impactive case series (phase 1 of the study)* Constipation Mean age: 5.7 years(56% g powder dissolved in at least 125 ml water per sachet) plus electrolytes (PEG) followed by a followed by a followed by a faccal impaction was procedural: Children were glycol 3350 (13.8 g powder dissolved in at least 125 ml water per sachet) plus electrolytes (PEG) followed by a followed by a followed by a faccal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours faecal impaction was procedural: Children were would, in the normal cours | |
| faecal impaction with polyethelene glycol plus electrolytes (PGE + E) followed by a followed by a faccal impaction with gibble and the study) and the study | |
| with polyethelene glycol plus electrolytes (PGE + E) followed by a follo | |
| polyethelene glycol plus electrolytes (PGE + E) followed by a followed b | |
| glycol plus electrolytes (PGE + E) (solution) followed by a 2 to 11 years with (PGE + E) (solution) followed by a 2 to 11 years (per sachet) plus electrolytes (PEG to 11 years) (per sachet) plus electrolytes (PEG to 12 ml water per sachet) plus electrolytes (PEG to 12 ml water per sachet) plus electrolytes (PEG to 13 ml water per sachet) plus electrolytes (PEG to 14 (n=28)) (per sachet) plus electrolytes (PEG to 14 (n=28)) (per sachet) plus electrolytes (PEG to 14 (n=28)) (per sachet) plus electrolytes (PEG to 15 ml water per sachet) plus e | |
| electrolytes (PGE + E) (PGE + E) (Followed by a solution of the study plane of the study | reated for |
| (PGE + E) | |
| followed by a 3 constipation + E; Movicol® disimpaction no: 3 (11) noncomparative because | _ |
| | |
| Idouble blind that had failed Country: LIK Ladministered without any laurescent rate obtained at it | |
| | |
| comparison of Study aim: to to respond to orally in hospital additional -age 5 to 11 (n=35) experience in treating imp | |
| PEG + E versus assess the conventional according to an intervention yes: 33 (94) with PEG + E in the author | ors' unit: it was |
| lactulose as efficacy of treatment and escalating dosing no: 2 (6) considered unethical to ra | indomise the |
| maintenance polyethylene would require regime until 2. Time to children to an alternative t | treatment |
| therapy. 2006. glycol 3350 hospital disimpaction was disimpaction 2. Time to | |
| Journal of plus admission for achieved (up to 7 (primary disimpaction (days) Sample size: intended to r | recruit 60 |
| Pediatric electrolytes disimpaction days) efficacy (mean, SD; median, children to obtain approximately days) | mately 45 |
| Gastroenterolog (PEG + E; (otherwise endpoint) range): children continuing to end | of phase 2 |
| y and Nutrition Movicol ®) as been admitted -PEG + E dosing -total (n=63) | |
| 43[1], 65-70 oral for enemas, regime 3. Maximum 5.7 ± 1.2 Successful disimpaction in | ndicated by the |
| monotherapy manual dose required 6.0 (3 to 7) passage of watery stools. | - |
| in the removal or No. PEG + E to achieve | |
| treatment of intestinal sachets: disimpaction -age 2 to 4 (n=28) Dose regime chosen because | ause it had |
| faecal lavage with 5.8 ± 1.2 shown to be effective in a | |
| impaction in PEG + E a. 2 to 4 years 4. Safety 6.0 (3 to 7) from the same unit | · |
| children and solutions) Day 1: 1 | |
| to compare Day 2: 2 -age 5 to 11 (n=35) After disimpaction children | n continued to |
| PEG + E with Exclusion Day 3: 2 5.6 ± 1.1 received PEG + E at the control of the | |
| lactulose as criteria: any Day 4: 3 6.0 (3 to 7) achieved disimpaction for | |
| maintenance Condition Day 5: 3 | - |
| therapy in a contraindicati Day 6: 4 3. Maximum dose the bowel had occurred | раскоп от |
| randomised ng the use of Day 7: 4 | |

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^{*} Study comprised two phases. Outcomes for the second phase (RCT) regarding maintenance therapy will be presented at the next review

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------------|---------------------------|---------------------------|---------------------|-----------------------------------|--|
| | Level | | s | · | Measures | | |
| | trial | PEG+E or | | | | (sachets/day): | Use of additional interventions |
| | | lactulose, | | b. 5 to 11 years | | -total (n=63): 6 | necessary to achieve disimpaction |
| | | including | | Day 1: 2 | | -age 2 to 4 (n=28): 4 | (laxatives, suppositories, enemas, |
| | | intestinal | | Day 2: 3 | | | washouts or manual removal) necessary |
| | | perforation or | | Day 3: 4 | | | to achieve disimpaction was also |
| | | obstruction, | | Day 4: 5 | | 4. Mean number (SD) | recorded |
| | | allergy to any | | Day 5: 6 | | of sachets required to | |
| | | of the | | Day 6: 6 | | | 3 children withdrew before receiving any |
| | | ingredients of | | Day 7: 6 | | -total (n=63): 19.6 | study medication and 2 children failed to |
| | | the trial | | _ | | (7.5) | disimpact within the time allowed, but |
| | | products, | | Comparison: none | | | they were included in results |
| | | paralytic ileus, | | | | -age 2 to 4 (n=28): | |
| | | toxic | | | | 14.3 (4.5) | Reviewer comments: |
| | | megacolon, | | | | | No explicit definition of "watery stools" |
| | | Hirschsprung' | | | | -age 5 to 11 (n=35): | given |
| | | s disease , | | | | 23.6 (6.8) | 1 |
| | | severe | | | | No significant | It is not clear who assessed the |
| | | inflammatory | | | | No significant | outcome "passage of watery stools", |
| | | bowel | | | | differences between | although it looks like it was probably the researchers |
| | | disease, uncontrolled | | | | the two age groups for any of the | researchers |
| | | renal/hepatic/ | | | | outcomes measured | Individual assessing outcomes not |
| | | cardiac | | | | outcomes measured | reported blinded to study objectives |
| | | disease. | | | | The 2 children who | reported billided to study objectives |
| | | uncontrolled | | | | failed to disimpact in | Not reported whether there were any |
| | | endocrine | | | | the 7 days specified | differences between the children who |
| | | disorder or | | | | in the study protocol | withdrew before receiving any |
| | | any | | | | were continued on | medication, those who failed to |
| | | neuromuscula | | | | PEG+E | disimpact and the ones who completed |
| | | r condition | | | | administration and | the study and disimpacted during the |
| | | affecting the | | | | eventually | time allowed |
| | | bowel | | | | disimpacted | |
| | | | | | | | Not clear whether vomiting affected the |
| | | Setting: | | | | 4. Safety: | dose required to achieve disimpaction |
| | | hospital | | | | -Number of children | or whether children receive any |
| | | | | | | experiencing adverse | medication to prevent / stop vomiting |
| | | | | | | effects: 39 (62%). | |
| | | | | | | (non of these judged | Source of funding: supported by Norgine |

| Bibliographic Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|--------------------|--------------------------------|---------------------------|------------------------------------|--|----------------------|
| | | | | | by investigator to be serious) Most commonly reported events: gastrointestinal (51% children) (abdominal pain, nausea, pruritus, ani / proctalgia and vomiting) No differences in the overall incidence of adverse effects or of gastrointestinal effects for the two age groups, except for vomiting (32% of age 2 to 4 children vs. 9% of aged 5 to 11 children)) results showed a direct correlation between incidence of vomiting and day of dosing | Pharmaceuticals Ltd. |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|----------------------------------|-----------------------|-----------------------|---------------------------|---------------------------------|---------------------------------|----------------------|--|
| | Level | 44 1 11 1 | S | | Measures | | |
| Youssef et al. | Study Type: | | 41 children | Intervention: | Follow-up | Clearance of faecal | Additional information from study: |
| Dose response | RCT | | 27 male | Polyethylene | period: 5 days | impaction (number of | Functional faecal retention: difficulty |
| of PEG 3350 for | | | median age 7.5 | glycol PEG 3350 | after starting | patients, %) | passing stools >3 months (straining, |
| the treatment of childhood fecal | <u>Evidence</u> | | years (3.,3 to | Comporisons (4 | treatment (48 | -Achieved | grunting, stool "getting stick") and |
| impaction. | level: | functional | 13.1) | Comparisons (4 arms): | hour after their last drug use) | total: 30 (75) | passage of stools <3 times/week |
| 2002. Journal of | - | | Country: USA | <u>aiiiis).</u> | last drug use) | lotal. 30 (73) | Planned to enrol 10 children in each |
| Pediatrics | Study aim: to | retention as | Country. USA | 1) 0.25 g/kg per | Outcome | (Values for each | group |
| | investigate | defined by | | day | Measures: | group are estimates | group |
| 141[0], 410-414 | the efficacy | Rome criteria, | | 2) 0.5 g/kg per | a. Primary | taken from a Bar | All medications for constipation |
| | | aged 3 to 18, | | day | outcome: | chart.): | discontinued 7 days before baseline |
| | 4 different | male or | | 3) 1.0 g/kg per | | onara). | examination and also during the |
| | | female, with | | day | -clearance of | a) 0.25 g/kg per day | duration of study |
| | | evidence of | | 4) 1.5 g/kg per | faecal | (n=10): 5 | |
| | glycol (PEG) | faecal | | day | impaction | | Faecal impaction: a palpable mass in |
| | 3350 in the | impaction at | | | ' | b) 0.5 g/kg per day | the left abdomen and/or a dilated rectum |
| | treatment of | physical | | Each of them to | b. Secondary | (n=10): 4 | filled with a large amount of hard stool |
| | childhood | examination | | be taken for 3 | outcomes: | | on rectal examination |
| | faecal | | | consecutive days, | | c) 1.0 g/kg per day | |
| | disimpaction | <u>Exclusion</u> | | premixed with a | -number of | (n=10): 9 | Presence or absence of faecal |
| | | criteria: | | solution flavoured | | | impaction assessed by abdominal and |
| | | previous | | | movements | d) 1.5 g/kg per day | rectal examination. Physical |
| | | gastrointestin | | Light (Kraft Food, | l | (n=9): 10 | examinations performed by 2 examiners |
| | | al surgery, no | | Inc) in the | -characteristics | | to confirm presence of faecal impaction |
| | | allergy | | morning with | of stools | p<0.05 c and d (95%) | |
| | | /sensitivity to | | breakfast at a | | vs. a and b (55%) | Investigators blinded to randomisation |
| | | PEG solution | | dose of | -safety | Niveshau of haveal | allocation sequence and concealment |
| | | Or phoophotoo | | 10mL/kg/day. If volume exceeded | | Number of bowel | maintained until patients enrolled |
| | | phosphates, signs and | | 240 ml, the | | movements in 5 days: | completed |
| | | signs and symptoms | | remaining daily | | >3 bowel movements | All medications dispensed to families in |
| | | suggestive of | | dose was equally | | during the 5-day | a clear container labelled with only a |
| | | obstruction | | divided | | study: 33 (83%) of | random sequence number generated by |
| | | (vomiting, | | throughout the | | total sample | manufacturer. All containers initially |
| | | abdominal | | remaining meals. | | istar odrijero | contained PEG 3350: 50g, 100g, 200g |
| | | distension | | Maximum dose | | (Values for each | or 300g. Each container was then |
| | | and | | 100 g daily | | group are estimates | constituted to a 2000 ml solution for |
| | | abdominal | | | | taken from a Bar | respective four doses |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-------------------------|--------------------------------|---------------------------|------------------------------------|--------------------------|--|
| | | mass that | | | | chart. Baseline value | |
| | | extended | | | | is less than 2 for all | Characteristics of stools measured by |
| | | beyond the level of the | | | | groups): | diaries provided to parents. Diaries had visual analog scales marked from 0 to |
| | | umbilicus) | | | | a) 0.25 g/kg per day | 10, each mark evenly spaced 1 cm |
| | | diffibilious) | | | | (n=10): 6 | apart, 0 minimum and 10 maximum. |
| | | | | | | (11–10). 0 | Children and parents asked to report |
| | | | | | | b) 0.5 g/kg per day | each defecation and its associated |
| | | | | | | (n=10): 8 | straining (0, very easy and no pushing; 1 |
| | | | | | | | to 10, very difficult and much effort), |
| | | | | | | c) 1.0 g/kg per day | consistency of stool (0, too loose and |
| | | | | | | (n=10): 11 | watery; 1 to 10 very hard), amount of |
| | | | | | | | stools per defecation (0, very little; 1 to |
| | | | | | | d) 1.5 g/kg per day | 10, a lot) associated gas (0, none; 1 to |
| | | | | | | (n=9): 12 | 10 too much) and cramping (0, none; 1 to 10 very painful) |
| | | | | | | p<0.005 for each | 45 |
| | | | | | | group compared to | 5 th day after initiation of treatment |
| | | | | | | the others | chosen for follow-up visit because of |
| | | | | | | | author's previous clinical experience |
| | | | | | | -time of first bowel | with PEG 3350 showed initial effect |
| | | | | | | movement after | between 1 and 2 days after beginning |
| | | | | | | initiation of treatment | use of medication |
| | | | | | | (mean ± SD) 1.89 ± | Clearance of faceal impaction defined |
| | | | | | | 0.46 days (total sample) | Clearance of faecal impaction defined as rectal vault that was either empty or |
| | | | | | | Sample) | had a small amount of soft stools. In |
| | | | | | | Characteristics of | those with abdominal examination |
| | | | | | | stools and symptoms | findings, resolution of the left lower |
| | | | | | | during treatment | quadrant mass in addition to an empty |
| | | | | | | acig troutilont | rectal vault was defined as successful |
| | | | | | | No significant | disimpaction. Clearance of faecal |
| | | | | | | differences in any of | impaction confirmed by 2 examiners |
| | | | | | | the following | |
| | | | | | | parameters among | Success of disimpaction not significantly |
| | | | | | | the 4 groups: | related to the independent factors of |
| | | | | | | straining, | age, duration of constipation, current |
| | | | | | | consistency, stool | use of medication for constipation and |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|---|
| | | | | | | amount, gas and cramping (copy actual | baseline constipation score |
| | | | | | | results) | One child receiving 1.5 g/kg/day did not show up at follow-up visit |
| | | | | | | Adverse effects: -Nausea (5%) | Reviewer comments: |
| | | | | | | -Vomiting (5%) | Small sample, no sample size |
| | | | | | | -Bloating/flatulence: 18% | calculation |
| | | | | | | -Pain/cramping: 5% -Loose stools (13%) -Diarrhoea: higher | Methods of randomisation and allocation concealment not described |
| | | | | | | | Examiners performing physical examination not clearly reported blinded. |
| | | | | | | (2/20); p<0.02 Acceptability of study | Unclear whether the two examiners who confirmed clearance of faecal impaction |
| | | | | | | medication by | were the same who assessed children |
| | | | | | | <u>children:</u> 95% of children took | at baseline Unclear who prepared the 2000 ml |
| | | | | | | | solution for respective four doses |
| | | | | | | · | Source of funding: supported by |
| | | | | | | Mean daily volumes required to take the | Braintree Laboratories Incorporated, General Clinical Research Centre, |
| | | | | | | appropriate study | Children's Hospital of Pittsburgh, |
| | | | | | | dose: no significant differences between | Pennsylvania |
| | | | | | | groups | |
| | | | | | | All children said they | |
| | | | | | | would repeat a 3-day regimen of PEG3350 | |
| | | | | | | to help treat future faecal impaction | |
| | | | | | | Duration of constipation at | |
| | | | | | | baseline significantly | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | longer for the group receiving 1.5 g/kg per day as compared to the group receiving 0.5 g/kg per day (p<0.03) | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|------------------------|---|
| | Level | | S | | Measures | | |
| Tolia et al. A | Study Type: | 48 children | 48 children | Intervention: | Follow-up | Frequencies (%) | Additional information from study: |
| prospective | RCT | | | 2-8 tablespoons | period: 2 days | (total sample for all | Constipation defined as the passage of |
| randomized | | | Data available | of mineral oil in 2 | | outcomes, n=36) | infrequent, large sized, firm to hard |
| study with | <u>Evidence</u> | | for 36 patients | divided doses for | | 1.History: | stools with or without associated rectal |
| mineral oil and | level: | | who completed | 2 days. Dose | Measures: | | pain or bleeding |
| oral lavage | 1- | > 2 years with | study: | empirically | , | a. number of bowel | |
| solution for | | constipation, | | determined (30 | | movements after | Randomisation performed by a |
| treatment of | Study aim: | normal growth | | ml/10 kg of body | | | computer-generated table |
| | to compare | | (mineral oil): | weight) | | none): | |
| in children. | the efficacy | | 11 males | | movements | -Group I (mineral oil, | Significantly more patients in the lavage |
| 1993. | and | | Mean age: 6.88 | If parents had | after treatment | n=17): 2/10/5 | group gave a history of previous |
| Alimentary | acceptability | | ± 3.26 years | difficulty in | -vomiting | | treatment with mineral oil (p<0.05). No |
| Pharmacology | of the | s disease | | administering the | -compliance | -Group II (lavage | significant differences at baseline |
| and | treatment of | excluded on | -Group II | oil they were | | solution, n=19): 9/8/2 | between 2 groups regarding: duration of |
| Therapeutics | | | (flavoured | | cramps/bloating | p<0.005 | constipation, frequency of stooling, |
| 7[5], 523-529 | impaction | , | lavage | it by blending it | -first bowel | | associated encopresis, rectal bleeding, |
| | using either | | solution): | with 120-180 ml of | | | previous treatments with enemas/fibre |
| | mineral oil or | | 6.44 ± 2.36 | orange juice | treatment | (none/occasional/a | diet, palpable abdominal masses, |
| | pineapple | , | years | | consider same | lot): | abdominal distension, anal fissure, |
| | isotonic | presence of | | Comparison: | treatment | -Group I (mineral oil, | perineal soiling, sphincter tone and |
| | intestinal | | Country: USA | pineapple | | n=17):17/0/0 | consistency of stool. |
| | lavage | faecal | | | 2.Physical | | |
| | solution | impaction in | | balanced oral | examination: | -Group II (lavage | Parents kept diaries assessing: |
| | | the anal canal | | lavage solution | | solution, n=19): | compliance of child with medication, |
| | polyethylene | and rectal | | containing | -palpable | 12/6/1 | time of first bowel movement after |
| | glycol-3350 | ampulla on an | | polyethylene | abdominal | p<0.005 | treatment, number of bowel movements |
| | (Colyte) | otherwise | | glycol-3350 | masses | | on each day, consistency of bowel |
| | | normal; | | (Colyte) | | c. compliance | movements, abdominal distension, |
| | | complete | | (sweetened with | distension) | (good/fair/poor): | cramps, nausea and vomiting, and |
| | | physical | | Nutra-Sweet) to | -consistency of | -Group I (mineral oil, | willingness to repeat the same treatment |
| | | examination | | drink in the dose | stool | n=17): 14/3/0 | in the future if impaction recurred |
| | | | | of 20 ml/kg/h for 4 | -anal fissure | | |
| | | Exclusion | | h once daily on 2 | -anal sphincter | -Group II (lavage | After treatment patients re-evaluated by |
| | | criteria: | | consecutive days. | | solution, n=19): 6/7/6 | the same physician who repeated the |
| | | medical | | Maximum | -perineal soiling | p<0.01 | abdominal and rectal examination in the |
| | | history of | | amount/hour: 1 | | , , , , , | same way as before |
| | | recurrent | | litre | | d. cramps/bloating | 40 |
| | | vomiting | | | | (none/ a few/a lot): | 12 patients failed to return for |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-------------------------|---------------------------|----------------------------------|---------------------|--------------------------------------|--|
| | Level | | S | | Measures | | |
| | | and/or aspiration, | | In addition patients received | | -Group I (mineral oil, n=17): 13/4/0 | reassessment in two days |
| | | central | | a single oral dose | | 11-11). 16/ 1/6 | Post-treatment history and physical |
| | | nervous | | of | | -Group II (lavage | examination further analysed after |
| | | system | | metoclopramide | | solution, n=19): | stratifying for previous use of mineral |
| | | problems or | | (0.1 mg/kg) before | | 10/8/1 | oils and stratified results did not differ |
| | | known history | | dinking the lavage | | N.S | significantly from unstratified analysis. |
| | | of liver, kidney and | | solution on both days to prevent | | e. first bowel movement after | Results presented are unstratified |
| | | heart disease | | nausea and | | | Reviewer comments: |
| | | | | vomiting | | day/none): | Small sample size. No sample |
| | | | | | | -Group I (mineral oil, n=17): 6/6/5 | calculation made |
| | | | | | | | Method of allocation concealment not |
| | | | | | | -Group II (lavage | described |
| | | | | | | solution, n=19): | |
| | | | | | | 14/3/2 | Physician-researchers not reported |
| | | | | | | p<0.01 | blinded |
| | | | | | | f. consider same treatment | Intention to treat analysis not performed |
| | | | | | | (yes/maybe/no): | Unclear how descriptive outcomes |
| | | | | | | -Group I (mineral oil, n=17): 12/3/2 | converted to numerical before analysis |
| | | | | | | | Source of funding: Block Drug |
| | | | | | | -Group II (lavage | Company, Inc. (Jersey City, NJ, USA) |
| | | | | | | solution, n=19): | provided the supplies for the study |
| | | | | | | 11/6/2 | |
| | | | | | | N.S | |
| | | | | | | 2.Physical | |
| | | | | | | examination: | |
| | | | | | | -palpable abdominal | |
| | | | | | | masses (none/a | |
| | | | | | | few/many): | |
| | | | | | | -Group I (mineral oil, | |
| | | | | | | n=17): 10/4/3 | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | -Group II (lavage solution, n=19): 17/1/1 p<0.005 | |
| | | | | | | -abdominal distension (none/some): -Group I (mineral oil, n=17): 11/6 | |
| | | | | | | -Group II (lavage solution, n=19): 11/8 N.S | |
| | | | | | | -consistency of stool (soft/firm/hard): -Group I (mineral oil, n=17): 12/3/2 | |
| | | | | | | -Group II (lavage solution, n=19): 14/3/2 N.S | |
| | | | | | | -anal fissure (none/healing): -Group I (mineral oil, n=17): 15/2 | |
| | | | | | | -Group II (lavage solution, n=19): 15/4 N.S | |
| | | | | | | -anal sphincter tone (normal/decreased): -Group I (mineral oil, n=17): 14/3 | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | -Group II (lavage solution, n=19): 15/4 N.S -perineal soiling (absent/present): -Group I (mineral oil, n=17): 10/7 -Group II (lavage solution, n=19): 13/6 N.S | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|------------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|-----------------------|---|
| Guest et al. | Study Type: | 224 children | 224 children | Intervention: | Follow-up | Percentage of | Additional information from study: |
| Clinical and | Multicentre | | aged 2 to 11 | macrogol 3350 | period: | patients disimpacted | Clinical data contained in patients' case |
| economic | retrospective | | vears | plus electrolytes | 12 weeks | within 5 days (%, | notes transcribed onto case report forms |
| impact of using | cohort | Inclusion | , | | (including | Confidence limit) | designed specifically for this study by |
| macrogol 3350 | | criteria: aged | 5 centres in | Comparison 1: | maintance | | one independent nurse, who examined |
| plus electrolytes | Evidence | between 2 | England and | enemas and | treatment) ² | -macrogol 3350 plus | the case notes of all patients at all |
| | level: | and 11 years, | Wales | suppositories | , | electrolytes (n=5 | centres |
| setting | 2- | suffering from | | | Outcome | centres): 97% (94%, | |
| compared to | | intractable | | Comparison 2: | Measures: | 100%) | Patients stratified according to centre |
| enemas and | Study aim: to | constipation | | manual | -Percentage of | 10070 | and initial treatment for disimpaction. |
| suppositories | estimate the | and initially | -macrogol 3350 | evacuation of the | patients | -enemas and | Individual clinical outcomes quantified |
| and manual | clinical and | disimpacted | plus electrolytes | | disimpacted | suppositories (n=5 | for each treatment at each centre. |
| evacuation to | economic | between | | anaesthesia | within 5 days | centres): 73% (58%, | Clinical centre was the unit of analysis |
| | | 01/01/01 and | n=5 centres | | | 89%) | |
| faecal impaction | • | 31/01/06 | | | -Time to initial | 3373) | Reviewer comments: |
| | macrogol | | -enemas and | | disimpaction | -manual evacuation | No clear definition of "intractable |
| clinical practice | 3350 plus | Exclusion | suppositories | | | of the bowel under | constipation" given |
| in England and | electrolytes | criteria: not | n=101 children | | -time for | anaesthesia (n=2 | green |
| Wales, 2007. | (macrogol | initially | n=5 centres | | disimpaction for | centres): 89% (67%, | Very small sample size for the manual |
| Current Medical | 3350; | disimpacted | | | those who did | 100%) | evacuation of the bowel |
| | Movicol, | between | -manual | | not disimpact | 10070 | |
| | Movicol | previous | evacuation of | | within 5 days | p<0.001 | Not reported which enemas and |
| 2213-2225 | Paediatric | dates or had | the bowel under | | | | suppositories children were treated with |
| | Plain) in an | any condition | anaesthesia | | -reported | Time to initial | for disimpaction |
| | outpatient | contraindicati | n=11 children | | adverse effects | disimpaction and time | , |
| | setting | ng the use of | n= 2 centres | | | for disimpaction for | Having another nurse (or other |
| | compared to | macrogol | | | | those who did not | professional) independently examining |
| | enemas and | 3350 | Country: UK | | | disimpact within 5 | the case notes or reviewing the |
| | suppositories | | | | | days: | transcriptions might have decreased the |
| | and manual | | | | | | risk of potential bias |
| | evacuation to | | | | | No significant | |
| | treat | | | | | differences amongst | According to the reported results it is |
| | paediatric | | | | | the 3 groups | unclear that clinical centre was the unit |
| | faecal | | | | | | of analysis |
| | impaction | | | | | Doses required for | , |
| | | | | | | successful | Source of funding: sponsored financially |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|-------------------------|--|
| | Level | | s | • | Measures | | |
| | | | | | | disimpaction within 5 | by Norgine Pharmaceuticals Ltd, |
| | | | | | | | Harefiled, UK, manufactures of Movicol |
| | | | | | | | (macrogol 3350 plus electrolytes) |
| | | | | | | -macrogol 3350 plus | |
| | | | | | | electrolytes (sachets): | |
| | | | | | | 29 (13 to 44) | |
| | | | | | | | |
| | | | | | | -enemas (units): | |
| | | | | | | 2 (1 to 3) | |
| | | | | | | -suppositories (units): | |
| | | | | | | 1 (1 to 2) | |
| | | | | | | | |
| | | | | | | Percentage of | |
| | | | | | | patients on different | |
| | | | | | | treatments during the | |
| | | | | | | week before initial | |
| | | | | | | treatment: | |
| | | | | | | Significantly more | |
| | | | | | | children disimpacted | |
| | | | | | | with manual | |
| | | | | | | evacuation were | |
| | | | | | | taking lactulose and | |
| | | | | | | senna compared with | |
| | | | | | | other 2 groups | |
| | | | | | | (p<0.001) | |
| | | | | | | (10.00.) | |
| | | | | | | Significantly more | |
| | | | | | | children disimpacted | |
| | | | | | | with Macrogol were | |
| | | | | | | taking picosulphate | |
| | | | | | | compared with other | |
| | | | | | | 2 groups (p<0.01) | |
| | | | | | | | |
| | | | | | | Significantly more | |
| | | | | | | children disimpacted | |
| | | | | | | with enemas and | |
| | | | | | | suppositories were | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | taking lactulose and other combinations (p<0.01), other laxatives (p<0.001) or were not treated ((p<0.001) when compared with other 2 groups | |
| | | | | | | No significant differences between the 3 groups for patients taking lactulose only or those taking Senna Adverse effects: a. Vomiting (%): -macrogol 3350 plus electrolytes (n=112 | |
| | | | | | | patients): 2 -enemas and suppositories (n=101 patients): 2 -manual evacuation of the bowel under anaesthesia (n=11 patients): 18 | |
| | | | | | | p<0.01 No significant differences among 3 groups for: urinary tract infection, dermatitis around anus, thrush and gastric illness | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|-----------------------|--|
| | Level | | S | | Measures | | |
| Pashankar et al. | Study Type: | 24 children | (data available | Intervention: | Follow-up | Soiling frequency | Additional information from study: |
| | Prospective | | for only 20 | PEG solution, | period: 8 weeks | (n=9) (mean ± SEM) : | Diagnosis of constipation based on |
| optimal dose of | case series | | children who | initial dose ~1g/kg | | | symptoms of at least 3 months' duration |
| daily | | | completed | body weight per | | before treatment: | including at least 2 of: hard stools, |
| polyethylene | Evidence | | study) | day (14 ml/kg/d | | 10.0 ± 2.4 | painful defection, withholding of stools, |
| glycol 3350 for | level: 3 | children | 0.1 | solution) given in | | during treatment: | faecal soiling, palpable faecal mass and |
| treatment of | | between ages | | 2 divided doses | | 1.3 ± 0.7 | fewer than 3 bowel movements/week |
| constipation | | | aged 18 months | for 8 weeks | frequency | p= 0.003 | |
| | | | to 11 years | | , | | Administration of all other medications |
| | efficacy and | | Mean age 6.09 | Parents instructed | | Total resolution of | for constipation stopped on enrolment. |
| 2001. Journal of | | | ± 4.2 years | to dissolve 17 g of | | soiling: 4 patients | No enemas or cathartics given either. |
| | PEG in | criteria: | 44 1911 | PEG powder in | faecal mass | (44.4%) | Initial doses of PEG prescribed based |
| 139[3], 428-432 | | | 11 children: | each 240 ml (8 | | | on authors' previous experience with |
| | constipation | | constipation | ounces) of water, | | Presence of | this agent |
| | | | alone | juice or other | faecal rectal | abdominal faecal | Ota al a su sista u su sa sa sa al har hista u s |
| | | anorectal | 0 1311 | clear-liquid | impaction | mass (n=18) | Stool consistency assessed by history |
| | | malformations | | beverage, families | -1:1-414-1 | h - f - u - t u t t | on a scale of 1 to 5 as follows: 1, hard; |
| | | | constipation + | allowed free | | before treatment: | 2, firm; 3, soft; 4, loose and 5, watery |
| | | 0 , | soiling | choice of clear | | 44% | Detients accessed as an alleger to det |
| | | any systemic | O | liquid beverage. | | | Patients examined on enrolment and at |
| | | | Country: USA | For determination | -painful | p<0.0029 | the end of 8 weeks of therapy for the |
| | | could lead to | | of best dose for | defecation | D | presence or absence of a palpable |
| | | constipation | | each child, | t | Presence of faecal | faecal mass, faecal impaction and rectal |
| | | | | parents asked to | -fear of | rectal impaction | dilatation |
| | | | | increase or | defecation | <u>(n=18)</u> | Children of annuantiate developmental |
| | | | | decrease volume | /stool | h - f - u - t u t t | Children of appropriate developmental |
| | | | | of PEG solution | | before treatment: | status advised to sit on toilet for 5 |
| | | | | by 20% every 3 | | 83% | minutes after each meal |
| | | | | days as required | | during treatment: | Deticate howel hobits hofers DEC |
| | | | | to yield 2 soft-to- | | 22% | Patients bowel habits before PEG |
| | | | | loose stools | | p<0.0006 | treatment compared with those recorded |
| | | | | (consistency | | Dilatad rootal varilt | on diary forms during the last 2 weeks |
| | | | | score of 3 to 4) | | Dilated rectal vault | (weeks 7 and 8) of treatment |
| | | | | per day | | <u>(n=18)</u> | 4 aubicate drapped from study because |
| | | | | Comparison: page | | before treatment: | 4 subjects dropped from study because of failure to return required symptoms |
| | | | | Comparison: none | | 78% | diaries: 2 of these had an excellent |
| | | | | | | | |
| | | | | | | during treatment: | response to therapy by parent report |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|--|
| | Level | | S | | Measures | p<0.0001 Fear of defecation /stool withholding (N=20) before treatment: 70% during treatment: 5% p<0.0001 Final effective dose during last 2 weeks of treatment (mean ± SEM) (g/kg/day): 0.84 ± 0.27 (range 0.27 to 1.42) Palatability: all children reported willingness to take PEG and found it highly palatable (to prepare PEG patients used sweeteners, fruit juices, water and cow's milk) | and two were lost to follow up Reviewer comments: Small sample size, no sample size calculation No data reported on who performed physical examination on enrolment and at the end of 8 weeks of therapy Not clear why data on physical examination available for only 18 children Source of funding: not stated |
| | | | | | | Adverse effects: no significant except for | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | diarrhoea during adjustment of dose. Flatulence (n=2) Abdominal pain (n=10) | |
| | | | | | | | |
| | | | | | | | |
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| | | | | | | | |

Pharmacological Interventions for Ongoing Treatment/ Maintenance in Children with Chronic Idiopathic Constipation

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|------------------|-----------------|---------------------|-----------------|--------------------|-----------------|----------------------|---|
| Information | Evidence | Patients | Characteristic | Comparison | Outcome | 2.11001 0.120 | Trovious Comments |
| in or manon | Level | . anomo | S | - Companicon | Measures | | |
| Candy et al. | Study Type: | 65 children | -Phase 1: | Intervention: | Duration of | Number of successful | Additional information from study: |
| Treatment of | Double-blind | | 65 children | Polyethylene | treatment | defecations/week | Sample size: intended to recruit 60 |
| faecal impaction | RCT** | <u>Inclusion</u> | | glycol 3350 (13.8 | 12 weeks | (last on-treatment | children to obtain approximately 45 |
| with | | criteria: | -Phase 2: | g powder | | <u>value)</u> | children continuing to end of phase 2 |
| polyethelene | <u>Evidence</u> | children aged | 58 children | dissolved in at | Assessment | Mean, SD, range | |
| glycol plus | level: | 2 to 11 years | | least 125 ml water | | | Children and investigators blinded to |
| electrolytes | 1+ | with | 67% boys | per sachet) plus | Immediately | -PEG+E (n=27): | medication which was dispensed |
| (PGE + E) | | intractable | | electrolytes (PEG | after treatment | | according to randomisation list |
| followed by a | Study aim: | | Mean age: 5.7 ± | + E; Movicol ®) | finished | 9.4 (4.56; 2 to 24) | generated by the study sponsor |
| double-blind | to assess the | that had failed | • | | | | |
| | efficacy of | to respond to | (range 2 to 11 | Comparison: | Follow-up | -Lactulose (n=26): | Blindness reasonably maintained as |
| PEG + E versus | polyethylene | conventional | years) | Lactulose (10 g | period: | | appearance of 2 products very similar |
| | glycol 3350 | treatment and | | | No follow-up | 5.9 (4.29; 2 to 23) | and both packed in sachets of an |
| | plus | | Country: UK | | made after | | identical size |
| | electrolytes | hospital | | water) | treatment | Difference in means: | |
| Journal of | (PEG + E; | admission for | | | finished | 3.5 | 5 children did not complete phase 1: 3 |
| | Movicol ®) as | disimpaction | | For both | | 95% CI: 1.0 to 6.0 | children withdrew before receiving any |
| 9 | oral | (otherwise | | medications | <u>Outcome</u> | p=0.007 | study medication and 2 children failed to |
| | monotherapy | been | | children received | Measures: | | disimpact within the time allowed |
| 43[1], 65-70 | in the | admitted for | | oral maintenance | | Reimpaction rate (n, | |
| | treatment of | enemas, | | doses | 1. Primary | % children): | 58 children entered phase 2. 5 were |
| | faecal | manual _. | | | efficacy | -PEG+E (n=27): 0 | excluded from the ITT population as |
| | impaction in | removal or | | ½ of the numbers | endpoint: | | they did not provide any on-treatment |
| | children and | intestinal | | of sachets | -number of | -Lactulose (n=26): 7 | efficacy data. |
| | to compare | lavage with | | required for | successful | (23%) | 10 children (17%) did not complete |
| | PEG + E with | PEG + E | | disimpaction/day | defecations/we | 0.044 | phase 2: 7 on lactulose reimpacted, 2 on |
| | lactulose as | solutions) | | D | ek | p=0.011 | lactulose did not want to continue, 1 on |
| | maintenance | | | Disimpaction | | | PEG+E did not complete the diary card |
| | therapy in a | Exclusion | | regime (n | | Number of sachets | |
| | randomised | criteria: | | sachets): | 2.Secondary | used each day: | No significant differences at baseline |
| | trial | any condition | | . 0 4- 4 | efficacy | -PEG+E (n=27): 0.91 | between 2 groups regarding: age, sex, |
| | | contraindicati | | a. 2 to 4 years | endpoints: | (0.41) | height and weight |
| | | ng the use of | | Day 1: 1 | | Lastulass (n. OC): | No abildran with draw farms the attention |
| | | PEG+E or | | Day 2: 2 | -reimpaction | -Lactulose (n=26): | No children withdrew form the study for |
| | | lactulose, | | Day 3: 2 | rate | 2.41 (0.91) | safety reasons |

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^{**} This is phase 2 of the study. Phase 1 was a prospective case series already discussed in the review for disimpaction

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|---------------|--------------|--------------------------|----------------|--------------------------------|------------------|-----------------------|---|
| Information | Evidence | Patients | Characteristic | Comparison | Outcome | | |
| | Level | | S | | Measures | | |
| | | including | | Day 4: 3 | | | |
| | | intestinal | | Day 5: 3 | -number of | Use of senna as | Reviewer comments: |
| | | perforation or | | Day 6: 4 | sachets used | rescue medication | No clear definition of constipation given |
| | | obstruction, | | Day 7: 4 | each day | -PEG+E (n=27): 0 | |
| | | allergy to any | | | | | Method of allocation concealment not |
| | | of the | | b. 5 to 11 years | -use of senna | -Lactulose (n=26): 8 | described |
| | | ingredients of | | Day 1: 2 | as rescue | (31%) | |
| | | the trial | | Day 2: 3 | medication | p=0.002 | Results not controlled for confounders |
| | | products, | | Day 3: 4 | | | |
| | | paralytic | | Day 4: 5 | -amount of stool | | Missing data on 2 children who did not |
| | | ileus, toxic | | Day 5: 6 | | differences in mean | enter phase 2 of the study |
| | | megacolon, | | Day 6: 6 | -predominant | values per patient | |
| | | Hirschsprung' | | Day 7: 6 | | between 2 groups | Source of funding: |
| | | s disease , | | | | with respect to: | supported by Norgine Pharmaceuticals |
| | | severe | | | | amount of stool, | Ltd. |
| | | inflammatory | | Additional laxative | -pain | predominant bowel | |
| | | bowel | | treatment with | | movement form, pain, | |
| | | disease, | | senna allowed as | -straining | straining, rectal | |
| | | uncontrolled | | rescue medication | | bleeding, abdominal | |
| | | renal/hepatic/ | | if the response to | -rectal bleeding | pain, soiling and | |
| | | cardiac | | a single agent | | overall assessment of | |
| | | disease, uncontrolled | | alone was judged inadequate by | -abdominal pain | treatment | |
| | | endocrine | | investigator | -soiling | Safety (% children) | |
| | | disorder or | | gato. | 9 | (n=58): | |
| | | any | | | -overall | -PEG+E: 64 | |
| | | neuromuscula | | | assessment of | -Lactulose: 83 | |
| | | r condition | | | treatment | | |
| | | affecting the | | | | Similar incidence in | |
| | | bowel | | | | each age group. Most | |
| | | | | | 3. Safety | commonly reported | |
| | | | | | | events | |
| | | | | | | gastrointestinal and | |
| | | | | | | resolved during the | |
| | | | | | | study. No clinically | |
| | | | | | | significant abnormal | |
| | | | | | | values observed in | |
| | | | | | | urine and plasma | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | electrolytes after 12 weeks of maintenance therapy | |
| | | | | | | | |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|-------------------------------------|-------------------------|--|
| | Level | | S | | Measures | | |
| Voskuijl et al. | Study Type: | 100 children | 91 children | Run-in phase (1 | Duration: | Defecation | Additional information from study: |
| PEG 3350 | RCT | | 49 male | week before | 8 weeks (RCT) | frequency/week | Childhood constipation defined as |
| (Transipeg) | | <u>Inclusion</u> | | treatment): | 18 weeks (case | | having at least 2 to 4 of the following |
| versus lactulose | Evidence | criteria: | age range: 6 | No laxatives | series) | -PEG 3350: 7.12 | symptoms for the last 3 months: less |
| in the treatment | level: | children aged | months to 15 | allowed. | | (5.14) | than 3 bowel movements/week, |
| of childhood | 1+ | 6 months to | years | At the end all | Assessment | -Lactulose: 6.43 | encopresis more than once/week, large |
| functional | | 15 years with | | patients received | point (s): | (5.18) | amounts of stool every 7 to 30 days |
| constipation: a | Study aim: | constipation | Age (y) (mean | | | N.S | (large enough to clog the toilet) and |
| double blind, | to compare | | (SD)) PEG | 3 days: | weeks after | | palpable abdominal or rectal mass on |
| randomised, | the clinical | Exclusion | 3350 6.5 (3.2) | -Children ≤ 6 | starting | <u>Encopresis</u> | physical examination |
| controlled, | efficacy and | criteria: | | | treatment | frequency/week: | |
| multicentre trial. | safety of PEG | organic | Lactulose 6.5 | (sodium | | | Estimated that a total sample of 90 |
| 2004. Gut | 3350 | causes for | (3.4) | | Follow-up | -PEG 3350: 3.11 | patients would be adequate to show a |
| 53[11], 1590- | (Transipeg; | defecation | | ate and sorbitol) | period: | (5.41) | difference of at least 30% more success |
| 1594 | polyethylene | disorders, | Country: the | -Children > 6 | 26 weeks after | -Lactulose: 2.84 | at 8 weeks using PEG 3350 compared |
| | glycol with | including | Netherlands | years: 120 ml Klyx | | (3.59) | to lactulose, with a 2 tailed alpha level of |
| | | Hirschsprung' | | | series phase | N.S | 0.05 with a power of 80% |
| | | s' disease, | | 1. Initial phase: | | | |
| | | spina bifida | | | <u>Outcome</u> | Success percentages | Unlabelled number boxes with |
| | constipation | occulta or | | Intervention: | Measures: | (95% CI) | unlabelled sachets prepared by the |
| | | hypothyroidis | | PEG 3350 | | | AMC pharmacy and handed out to |
| | | m | | | | PEG 3350: 56 (39 to | patients after randomisation. The box |
| | | | | -children aged 6 | | 70) | contained 180 sachets containing either |
| | | | | months to 6 years | -frequency of | | lactulose 6g/sachet or PEG 3350 2.95g |
| | | | | (inclusive): one | stools | Lactulose: 29 (16 to | per sachet. |
| | | | | sachet (2.95g) per | | 44) | |
| | | | | day | -frequency of | | Toilet training advised after each meal |
| | | | | | encopresis | P=0.02 | (5 minutes) and small gifts and praise |
| | | | | -children older | | | used to enhance compliance |
| | | | | than 6 years: 2 | -overall | Overall treatment | |
| | | | | sachets (5.9g) per | | | No significant differences at baseline |
| | | | | day | | | between the 2 groups with respect to: |
| | | | | | | | age, sex, defecation frequency, |
| | | | | Comparison: | 2. Safety | laxatives for more | encopresis, large amounts of stool and |
| | | | | Lactulose | | than 1 year prior to | faecal impaction |
| | | | | | | the start of the study. | |
| | | | | -children aged 6 | · · · · · · · · · · · · · · · · · · | | 9 dropouts: 4 on PEG 3350, 5 on |
| | | | | months to 6 years | gastrointestinal | less than 1 year a | lactulose. 2/each group lost to follow-up, |

| omments |
|---|
| Inknown. 2 on acter positive, 1 latability of study |
| ess defined 3 or t/week and 1 less every 2 |
| on and allocation |
| ribed comes not se of this review |
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| Bibliographic Study Evid Le | ence Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|-----------------------------|---------------|--------------------------------|------------------------------|------------------------------------|--|-------------------|
| | | | | | significant differences between 2 groups regarding: bloating, diarrhoea, flatulence, nausea, hard stool consistency and vomiting. Significantly more children complained of bad palatability of PEG compared to lactulose and this caused the premature withdrawal of 1 patient. | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|---------------------|---|
| | Level | | S | | Measures | | |
| Loening-Baucke | Study Type: | 79 children | 79 children | General: | Duration of | | Additional information from study: |
| et al. A | RCT | | 65 boys | disimpacted with | treatment: | -at 12 months: | Functional constipation defined by |
| randomized, | | <u>Inclusion</u> | age range: 4 to | 1 or 2 phosphate | 12 months | PEG (n=34): 62 | duration of ≥ 8 weeks and ≥ 2 of the |
| prospective, | <u>Evidence</u> | | 16.2 years | enemas in the | | MOM (n=21): 43 | following: frequency of bowel |
| comparison | level: | 4 years and | (median 7.4; | clinic on the day | <u>Assessment</u> | | movements <3 stools/week, >1 episode |
| study of | 1- | presence of | mean 8.1 ± 3.0) | of the visit, if | | NS | of faecal incontinence/week, large stools |
| polyethylene | | functional | | necessary and | 1, 3, 6 and 12 | | noted in rectum or felt during abdominal |
| glycol 3350 | Study aim: to | constipation | Country: USA | started laxative | | Recovery rate (%) | examination, passing of stools so large |
| without | compare the | with faecal | | therapy that | initiating | -at 12 months: | that they obstructed the toilet |
| | efficacy, | incontinence | | evening | | PEG (n=34): 33 | |
| milk of | safety and | | | | | MOM (n=21): 23 | Randomisation performed by children |
| magnesia for | patient | Exclusion | | Intervention: | Follow-up | | drawing a sealed envelope with and |
| | | criteria: stool | | polyethylene | | NS | enclosed assignment |
| | polyethylene | toileting | | glycol (PEG) 3350 | | | |
| | glycol (PEG) | refusal, faecal | | without added | | Bowel movement | Investigators, children and their parents |
| incontinence. | 3350 without | incontinence | | electrolytes 0.7 | treatment | frequency (mean ± | aware of the study group assignment |
| 2006. Pediatrics | | but no | | | finished | <u>SD,</u> | |
| 118[2], 528-535 | | constipation, | | daily for 12 | | episodes/week) | Estimated that 38 subjects required in |
| | | previous | | months | <u>Outcome</u> | -Baseline: | each group to be able to detect a |
| | magnesia | refusal of one | | | | (/ | difference in failure rates between the 2 |
| | (MOM) over | of study | | capful of PEG (17 | | MOM (n=40): 3.5 ± 6 | groups of 30% in 12 months (40% vs. |
| | 12 months | medications, | | g) mixed in 8 oz of | | | 10%), at the .05 significance level with |
| | | children who | | | outcomes: | -at 12 months: | .80 power. Authors hypothesized that |
| | | came from far | | Kool-Aid, Crystal | | | PEG would be as successful as MOM in |
| | | away for a | | Light or water) | | MOM (n=21): 8.2 ± | treating chronic constipation and faecal |
| | | second | | making a solution | | 3.9 | incontinence. Authors' previous study |
| | | opinion, | | of ~2g/30 mL | -recovery | | showed that 33% of children refused to |
| | | Hirschsprung' | | | | P<0.005 for both | take MOM during the first 12 months of |
| | | s disease, | | Comparison: | | groups compared to | treatment. |
| | | chronic | | | outcomes: | baseline | |
| | | intestinal | | (MOM) 2mL/kg | | <u>_</u> | Children treated with minimal effective |
| | | pseudobstruct | | body weight daily | | Faecal Incontinence | dosage of PEG or MOM, allowing for a |
| | | ion, previous | | for 12 months | stool frequency | frequency (mean ± | daily stool and preventing abdominal |
| | | surgery | | | per week | <u>SD,</u> | pain and faecal incontinence. Parents |
| | | involving | | plain MOM could | | episodes/week) | instructed to aim for 1 or 2 stools of |
| | | colon or anus | | be mixed into | -improvement in | | milkshake consistency each day. |
| | | | | apple sauce or | | PEG (n=39): 12.2 ± | Parents asked to increase dosage if |
| | | | | milkshakes, or | faecal | 13 | stools too hard or not frequent enough |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|---------------------|--------------------------------|---|
| | Level | | S | | Measures | | |
| | | | | chocolate and | incontinence | MOM (n=40): 13.5 ± | and to decrease the dosage if stools |
| | | | | other flavouring | per week | 15.5 | watery or too numerous. Small changes, |
| | | | | could be added | | | such as 2 oz of PEG or 0.5 tbsp of MOM |
| | | | | | -resolution of | -at 12 months: | every 3 days, were recommended. |
| | | | | Large doses of | | | Regular stool sittings for 5 minutes after |
| | | | | both medications | | MOM (n=21): 0.5 ± | each meal required initially. Toilet sitting |
| | | | | could be divided | | 1.6 | frequency reduced after children |
| | | | | into 2 doses | -safety profile | P<0.005 for both | recognized urge to defecate and initiated toilet use themselves. |
| | | | | | | groups compared to | initiated tollet use themselves. |
| | | | | | | baseline | No significant differences at baseline |
| | | | | | compliance | bacomic | between the 2 groups regarding: age, |
| | | | | | | Abdominal pain (%) | sex, primary faecal incontinence, |
| | | | | | | -Baseline: | previous treatment with laxatives, history |
| | | | | | | PEG (n=39): 71.8 | of retentive posturing, frequency of |
| | | | | | | MOM (n=40): 52.5 | bowel movements, bowel movements |
| | | | | | | | obstructing the toilet, frequency of faecal |
| | | | | | | -at 12 months: | incontinence, presence of abdominal |
| | | | | | | PEG (n=34): 3 | pain, presence of abdominal faecal |
| | | | | | | MOM (n=21): 0 | mass and presence of rectal faecal mass |
| | | | | | | P<0.005 for both | |
| | | | | | | groups compared to | By 12 months a total of 27 dropouts/lost |
| | | | | | | baseline | to follow-up. PEG: 2 children lost to |
| | | | | | | | follow-up monitoring, 2 (5%) had refused |
| | | | | | | At 12-month frequency of bowel | PEG, 1 child allergic to PEG, 2 children were receiving senna. These 7 children |
| | | | | | | movements, | counted as not improved and not |
| | | | | | | frequency of | recovered. MOM: 2 |
| | | | | | | episodes of faecal | Children lost to follow-up monitoring, 3 |
| | | | | | | incontinence, and | children had discontinued study |
| | | | | | | percentage of | participation, 14 children (35%) had |
| | | | | | | children | refused to take MOM, and 1 child was |
| | | | | | | with abdominal pain | receiving senna |
| | | | | | | not significantly | |
| | | | | | | different | Efficacy analyses performed with |
| | | | | | | | intention to treat population, other |
| | | | | | | MOM group | outcomes calculated from available |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|------------------------|--|
| | Level | | S | | Measures | | |
| | | | | | | | follow-up data |
| | | | | | | Patient Acceptance | |
| | | | | | | Several children | Reviewer comments: |
| | | | | | | complained about | Results not controlled for potential |
| | | | | | | taste of PEG and MOM. | confounders High drop-out / lost to follow-up rate: |
| | | | | | | 2 children (5%) | 30.4% |
| | | | | | | continued to refuse | 00.470 |
| | | | | | | PEG vs. 14 children | Source of funding: Braintree |
| | | | | | | (35%) continued to | Laboratories (Braintree, MA) supported |
| | | | | | | refuse MOM during | study with an unrestricted research |
| | | | | | | the 12 months of the | grant. According to authors, the funding |
| | | | | | | study (P < .001 | source had no involvement in the study design, collection, analysis, |
| | | | | | | Treatment doses | interpretation of data, writing of the |
| | | | | | | (mean ± SD): | report or decision to submit the article |
| | | | | | | 1 | for publication |
| | | | | | | -PEG (g/kg body | |
| | | | | | | weight) | |
| | | | | | | 1 month: 0.7 ± 0.2 | |
| | | | | | | 3 months: 0.6 ± 0.3 | |
| | | | | | | additional senna at | |
| | | | | | | some point: 3 children | |
| | | | | | | -MOM (mL/kg body | |
| | | | | | | weight) | |
| | | | | | | 1 month: 1.2 ± 0.7 | |
| | | | | | | 3 months: 1.2 ± 0.8 | |
| | | | | | | additional senna at | |
| | | | | | | some point: 1 child | |
| | | | | | | mean doses similar in | |
| | | | | | | children who | |
| | | | | | | improved and who | |
| | | | | | | did not improve for | |
| | | | | | | both treatments | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | safety profiles PEG: 1 child allergic No other significant clinical effects for either medication, apart from transient diarrhoea disappearing with dose reduction -Laboratory tests: PEG: 1 child with elevated platelets before and after treatment, 1 child with decreased sodium levels at 6 months, but normal at 12 months MOM: 1 child high platelet count, 1 low serum sodium level, elevated AST, 1 elevated ALT | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|-----------------------------|---------------------------|------------------------|-----------------------|--|
| | Level | . anomo | S | Companicon | Measures | | |
| Dupont et al. | Study Type: | 96 children | 96 children | Intervention: | Duration of | Stool frequency | Additional information from study: |
| Double-blind | RCT | | 51 male | PEG 4000 | treatment: | (number of stools/wk, | Constipation defined as less than 1 |
| randomized | | <u>Inclusion</u> | | | 3 months | median (interquartile | stool/day for > 1 month in children 6 to |
| evaluation of | <u>Evidence</u> | criteria: | | -Starting dose: | | range) | 12 months old and less than 3 |
| clinical and | level: | children with | -Age (months) | | <u>Assessment</u> | | stools/week for > 3 months in children |
| biological | 1+ | constipation | median (25 th to | 1 placebo to be | point (s): | -D42 | aged 13 months to 3 years |
| tolerance of | | despite their | 75th | taken at breakfast | | NS in babies | |
| polyethylene | Study aim: to | usual dietary | percentiles) | | and day 84 | Toddlers: | PEG 4000 and lactulose packaged in a |
| glycol 4000 | | treatment for | | Comparison: | (D84) after | PEG 4000 (n=51): | double-blind and double-dummy design, |
| versus lactulose | safety of a | at least 1 | PEG 4000: | Lactulose | starting | 8 (6–10) | by means of coupled sachets, according |
| in constipated | polyethylene | month, aged | 28 (19.5–33.7) | | treatment | Lactulose (45): | to a randomisation list. Double dummy |
| children. 2005. | | 6 months to 3 | | -Starting dose: | | 6 (5–7) | design required because of the |
| Journal of | 4000 laxative | years, | Lactulose: | 1 sachet (3.33g) | Follow-up | (P=0.013). | difference of taste between the drugs. |
| Pediatric | without | ambulatory | 25.8 (12.3–33) | and 1 placebo to | period: | | Numbered boxes provided to |
| Gastroenterolog | additional | | | be taken at | No follow-up | -D84 | investigators at each site in equal |
| y and Nutrition | salts in | <u>Exclusion</u> | Country: | breakfast | performed after | NS in babies or | numbers. Investigators randomly |
| 41[5], 625-633 | paediatric | <u>criteria:</u> | France | | treatment | toddlers | allocated either PEG 4000 or lactulose |
| | patients | history of | | | finished | | to the children for a 3-month period, with |
| | | intractable | | For both drugs, | | Frequency of hard | the same strategy for dose adaptation |
| | | faecaloma, | | dose could be | <u>Outcome</u> | <u>stools</u> | |
| | | Hirschsprung' | | doubled if | Measures: | | 3 children not included because of a |
| | | s disease, | | ineffective in | | -D42 | baseline laboratory value ONR (out of |
| | | neurologic, | | children aged 13 | -Efficacy: | PEG 4000: 9% | normal range) before amendment |
| | | endocrine or | | months to 3 years | | (4 of 46) | applied. 2 children in PEG 4000 group |
| | | metabolic | | If maximum | stool frequency | Lactulose (45): 34% | dropped out before any study drug |
| | | disorders, | | authorised dose | frequency of | (14 of 41) | intake, so the intention to treat |
| | | allergic | | unsuccessful, one | | P = 0.003 | population included 51 children (10 |
| | | disease or | | micro-enema of | enema use | | babies and 41 toddlers) in the PEG |
| | | allergies | | glycerol per day | faecal | -D84 | 4000 group and 45 (12 babies and 33 |
| | | | | could be | impaction | PEG 4000 (n=51): | toddlers) in the lactulose group. 76 of |
| | | | | prescribed for a | • | 6% (3 of 47) | these children included in the per |
| | | | | maximum of 3 | appetite | Lactulose (45): 28% | protocol analysis and 20 excluded by |
| | | | | consecutive days. | 5 | (11 of 40) | the independent scientific committee for |
| | | | | If child not | -Biological | P = 0.008 | at least one major deviation, 11 in the |
| | | | | produced stools | tolerance: | _ | PEG 4000 group and 9 in the lactulose |
| | | | | after treatment 2 | l. | Enema use | group. Reasons for exclusion were no |
| | | | | enemas could be | ion | D. 40 | laboratory test at D84, one or more one |
| | | | | administered at a | electrolytes | -D42: | missing laboratory results at D84, |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|--|---|---|---|
| | | | | liquid stools for >1 day or > 2 or 3 stools/day depending on age, dose could | total protein albumin vitamin A vitamin D folates | of 48) Lactulose: 43% (19 of 44) -D84: PEG 4000: 17% (8 of 48) | delayed laboratory test at D84 (n = 12), inadequately long exposure to the study drug (n = 2), personal reasons (n = 5) and unauthorized concomitant treatment (n = 1) No clinically relevant differences between 2 treatment groups at baseline for clinical or biologic parameters Stool frequency, abdominal pain, vomiting, and nausea recorded on Self-Diary Evaluation Booklet Reviewer comments: Methods of randomisation and allocation concealment not clearly described No sample calculation performed Results not controlled for potential confounders Source of funding: not stated |
| | | | | | | P<1.00 | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | Appetite score improvement | |
| | | | | | | PEG 4000 (n=51): +19% Lactulose (45): -4% | |
| | | | | | | p<0.003 | |
| | | | | | | Clinical tolerance (ITT population) | |
| | | | | | | -6 adverse effects (all non serious): 5 diarrhoea (5 episodes in 2 children in both treatment groups) 1 anorexia (on lactulose) | |
| | | | | | | -median (interquartile range) duration of either new onset or worsened flatulence (days): | |
| | | | | | | PEG 4000: 3 (1 to 4.5) Lactulose: 5 (3 to 19.5) P=0.005 | |
| | | | | | | -median (interquartile range) duration of either new onset or | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | worsened vomiting episodes (days): | |
| | | | | | | PEG 4000: 1 (1 to 2) Lactulose: 2 (1 to 6) P<0.05 | |
| | | | | | | -anal irritation: 5% (2 out of 40 children, both on lactulose) | |
| | | | | | | -no difference between PEG 4000 and lactulose groups with regards to other digestive tolerance outcomes | |
| | | | | | | -Body height and body weight unaffected during the 3-monht treatment for | |
| | | | | | | both boys and girls Biological tolerance (ITT population): No significant difference between | |
| | | | | | | treatment groups for the % of children with ONR values on D84 compared to baseline | |
| | | | | | | status. No treatment- related changes found in serum iron, electrolytes, total protein, albumin | |
| | | | | | | and vitamins A, D and folates | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | Dose used (sachets/day) (median (interquartile range)) -Babies: 1 (0.9 to 1) PEG 1 (1 to 1.3) lactulose | |
| | | | | | | P = 0.67 -Toddlers 1 (1 to 1.3) PEG 1.1 (0.9 to 1.5) lactulose P = 0.58 | |
| | | | | | | Treatment stopped in 1 child because of lack of efficacy (lactulose group). | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|-----------------------|---|
| | Level | | S | | Measures | | |
| Perkin. | Study Type: | 21 children | 21 children | Intervention: | <u>Duration:</u> | Number of patients | Additional information from study: |
| Constipation in | RCT | | | Senna syrup | 1 week each | passing stools of any | Patients given either treatment |
| childhood: a | (crossover) | <u>Inclusion</u> | Country: UK | 10 to 20 ml daily | period with 1 | kind each day: | according to a code-list of random |
| controlled | | criteria: | | for 1 week | week no | Lactulose vs. Senna | numbers, placed in a series of sealed |
| comparison | <u>Evidence</u> | children aged | | | | N.S | envelopes, one of which was opened |
| between | level: | <15 years | | Comparison: | between | | each time a child entered the trial |
| lactulose and | 1- | with a history | | Lactulose | | Number of patients | |
| standardized | | of | | | <u>Assessment</u> | passing normal stools | 1 dropout: 1 patient on senna at the |
| senna. 1977. | Study aim: | constipation | | for 1 weeks | point (s): | each day (mean) | beginning of study failed to attend at the |
| Current Medical | | treated at | | | immediately | 1 1 10 1 | end of 1 st week |
| Research and | effectiveness | home for 3 | | Each preparation | after treatment | -Lactulose: 13.4 | |
| Opinion 4[8], | and side | months or | | | completed | -Senna: 8.43 | No written or oral indication of any |
| 540-543 | effects | more | | the appropriate | | p <0.01 | medical preference for other preparation |
| | between a | | | treatment week in | | | given and patients presented with single |
| | standardised | Exclusion | | a daily dose | period: | Adverse effects (n | bottle of one or other of the preparations |
| | senna syrup | criteria: | | varied according | No follow up | patients): | according to the coded instruction at |
| | | any cause of | | to the age of the | | a- senna week: | start of trial. On 3 rd week a bottle of |
| | in the | constipation | | patient | | 12 (8 colic, 1 | alternative preparation was given |
| | treatment of | requiring | | | finished | diarrhoea, 2 colic+ | |
| | childhood | surgical or | | 1 intermediate | | diarrhoea, 1 colic + | Outcomes recorded by parents in written |
| | constipation | medical | | week with not | <u>Outcome</u> | distension) | diaries |
| | | correction in | | treatment | Measures: | | |
| | | addition to | | | | | 4-point scale of stool consistency: loose, |
| | | laxation | | | | 4 (3 colic, 1 colic + | normal, hard, none |
| | | | | | consistency | distension) | |
| | | | | | | | Reviewer comments: |
| | | | | | | c- lactulose week | No clear definition of constipation given |
| | | | | | | 1 (colic) | Very small sample size, no sample size |
| | | | | | each day | | calculation |
| | | | | | | p<0.001 (a vs. c) | Inadequate method of allocation |
| | | | | | -adverse effects | NS (b vs. c) | concealment |
| | | | | | | | Patients' baseline characteristics not |
| | | | | | | | reported |
| | | | | | | | Study not reported as blinded |
| | | | | | | | Results not controlled for confounders |
| | | | | | | | Very short treatment period |
| | | | | | | | According to authors the number of |
| | | | | | | | stools passed each day was recorded, |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|-------------|---------------------|
| | | | | | | | but is not reported |
| | | | | | | | Source of funding: |
| | | | | | | | not stated |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------------|---------------------------|------------------------------------|---------------------|-----------------------|--|
| | Level | | s | | Measures | | |
| Farahmand. A | Study Type: | 247 children | 247 children | General: | Duration of | Stool frequency | Additional information from study: |
| randomised trial | RCT | | | 1 or 2 enemas | treatment: | (mean ± SD) | Diagnosis of chronic functional |
| of liquid paraffin | | <u>Inclusion</u> | 127 male | daily for 2 days to | 8 weeks | | constipation based on having at least 2 |
| versus lactulose | <u>Evidence</u> | criteria: | | clear any rectal | | week): | of the following symptoms for the last 3 |
| in the treatment | <u>level:</u> | chronic | aged 2 to 12 | | | Liquid paraffin | months: less than 3 bowel |
| of chronic | 1- | functional | years old (mean | cc/10 kg of | | (n=127) | movements/week, faecal soiling more |
| functional | | constipation | 4.1± 2.1 years) | paraffin oil) | | 1.6 ± 1 | than once/week, large amounts of stool |
| constipation in | Study aim: | | | | | Lactulose (n=120) | every 7 to 30 days and palpable |
| children. 2007. | | <u>Exclusion</u> | Country: Iran | Intervention: | | 1.8 ± 1.2 | abdominal or faecal mass on physical |
| Acta Medica | the clinical, | criteria: | | Liquid paraffin | | p=0.155 | examination |
| | | organic | | orally, 1 to 2 | Follow-up | | |
| | safety of liquid | | | ml/kg, twice daily | period: | -during first 4 weeks | Apart form laxative treatment, parents |
| | | defecation | | for 8 weeks | 12 weeks after | (per week): | given instructions to increase their daily |
| | | disorders | | | treatment | Liquid paraffin | fibre intake to an amount of grams equal |
| | | including | | Comparison: | finished | (n=127) | to their age plus 10. Toilet training after |
| | | Hirschsprung' | | Lactulose orally, 1 | 0.1 | 12.1 ± 3.2 | each meal advised to enhance |
| | | s' disease, | | | <u>Outcome</u> | Lactulose (n=120) | compliance |
| | | spina bifida | | daily for 8 weeks | | 9.2 ± 2.1 | T |
| | | occulta, | | | | p<0.001 | Treatment success defined as 3 or more |
| | | hypothyroidis | | | -stool frequency | duning last 4 was les | bowel movements/week and encopresis |
| | | m, cystic | | | | -during last 4 weeks | episodes less than 2/week |
| | | fibrosis, | | Can data main atian | -encopresis | (per week): | No significant baseline differences |
| | | neurological abnormalities. | | For determination of best dose for | rrequency | Liquid paraffin | No significant baseline differences between the 2 treatment groups |
| | | intestinal | | | augaga rata | (n=127) 13.1 ± 2.3 | regarding: age, sex, duration of |
| | | pseudo | | child, parents asked to increase | | Lactulose (n=120) | constipation, defection frequency, |
| | | obstruction | | the volume of | -optimal dose of | | number of patients with history of |
| | | ODSTRUCTION | | each drug by 25% | | p<0.001 | encopresis, large amount of stool, faecal |
| | | | | every 3 days as | urug | p<0.001 | impaction in rectum, rectal bleeding, lost |
| | | | | | -side effects | Encopresis frequency | to follow-up after 8 weeks, bad |
| | | | | or 2, firm-loose | -side ellecis | (mean ± SD) | palatability of study medication |
| | | | | stools | | -Before treatment | paratability of Study Medication |
| | | | | 310013 | | (per week): | Reviewer comments: |
| | | | | | | Liquid paraffin | Method of randomisation and allocation |
| | | | | | | (n=127) | concealment not described |
| | | | | | | | Non blinded study |
| | | | | | | | No sample calculation performed |
| | | | | | | | No withdrawals/dropouts reported |

| Information Evidence Patients Characteristic Comparison Outcome Level s Measures | ct Size Reviewer Comments |
|--|--|
| p=0.1 -during firs (per week Liquid para (n=127) 1 ± 4.3 Lactulose 2 ± 4.6 p=0.07 -during las (per week Liquid para (n=127) 0 ± 0 Lactulose 3 ± 4.1 p<0.001 Success rigs%) -during firs Liquid para (n=127) 90 Lactulose 52 p<0.001 -at end of Liquid para (n=127) 85 Lactulose 29 | Source of funding: not stated, but authors reported "no conflicts of interests" (n=120) at 4 weeks affin (n=120) ate (%, CI at 4 weeks: affin (n=120) 8 weeks: affin |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|------------------------|-------------------|
| | | | | | | Optimal dose of drug | |
| | | | | | | -Final effective dose | |
| | | | | | | (mean, ml/kg/day): | |
| | | | | | | Liquid paraffin | |
| | | | | | | (n=127) | |
| | | | | | | 1.72 ± 0.13 | |
| | | | | | | Lactulose (n=120) | |
| | | | | | | 2.08 ± 0.21 | |
| | | | | | | p<0.001 | |
| | | | | | | Side effects (during 4 | |
| | | | | | | to 12 week) (not clear | |
| | | | | | | whether, n or %, but | |
| | | | | | | probably %) | |
| | | | | | | (estimates taken from | |
| | | | | | | bar chart, outcomes | |
| | | | | | | not reported in text): | |
| | | | | | | Lactulose (n=120) | |
| | | | | | | Abdominal pain: 10 | |
| | | | | | | Bad palatability: 15 | |
| | | | | | | Pain at defecation: 10 | |
| | | | | | | Bloating: 10 | |
| | | | | | | Diarrhoea: 10 | |
| | | | | | | Anal oil leakage: 20 | |
| | | | | | | Flatulence: 10 | |
| | | | | | | Nausea: 10 | |
| | | | | | | Hard stool: 20 | |
| | | | | | | Vomiting: 0 | |
| | | | | | | Liquid paraffin | |
| | | | | | | (n=127) | |
| | | | | | | Abdominal pain: 50 | |
| | | | | | | Bad palatability: 40 | |
| | | | | | | Pain at defecation: 50 | |
| | | | | | | Bloating: 20 | |
| | | | | | | Diarrhoea: 30 | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | Anal oil leakage: 40 Flatulence: 20 Nausea: 5 Hard stool: 6 Vomiting: 0 | |
| | | | | | | | |
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| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|-----------------------|---|
| Gremse et al. | Study Type: | 44 children | 44 children | Intervention: | Duration of | Mean number of | Additional information from study: |
| Comparison of | RCT | | | PEG 3350 without | treatment: | bowel movements | 7 patients withdrew during the first 2- |
| polyethylene | (crossover) | <u>Inclusion</u> | Age range: 2 to | electrolytes | 2 weeks each | | week treatment period due to lack of |
| glycol 3350 and | | criteria: | 16 years (mean | (MiraLax) | period | -PEG 3350 (n=37): | efficacy of the assigned intervention: 6 |
| lactulose for | <u>Evidence</u> | patients aged | 7.8 ± 3.7) | 10g/m2/d orally | | 14.8 ± 1.4 | patients taking lactulose at time of |
| treatment of | level: | 2 to 16 years, | | for 2 weeks | <u>Assessment</u> | | withdrawal |
| chronic | 1- | referred for | Country: USA | | point (s): | -Lactulose (n=37): | |
| constipation in | | subspecialty | | Mean weight | | 13.5 ± 1.5 | Stool form scoring: 0 hard, 1 firm, 2 soft, |
| children. 2002. | Study aim: | evaluation of | | adjusted dose: 0.3 | after each | | 3 loose, 4 watery |
| Clinical | | constipation | | | treatment | Stool form (mean | |
| Pediatrics 41[4], | the efficacy of | | | to 0.5) | period | sum of scores) | Stool passage scoring: 0 hard, 1 difficult, |
| 225-229 | PEG 3350 | Exclusion | | | | | 2 easy, 3 urgency, 4, no control |
| | and lactulose | criteria: | | Comparison: | Follow-up | -PEG 3350 (n=37): | |
| | in the | organic | | Lactulose 1.3 | period: | 25.9 ± 3.0 | Stool frequency, form and easy of |
| | treatment of | disease of the | | g/kg/d orally for 2 | No follow-up | | passage recorded by parent or guardian |
| | chronic | large or small | | weeks | made after | -Lactulose (n=37): | in symptom diary |
| | constipation in | | | | | 27.9 ± 1.5 | |
| | | known allergy | | | completed | | Reviewer comments: |
| | | to PEG or | | (no washout | | Stools passage | No definition of constipation given |
| | | lactulose, | | period) | <u>Outcome</u> | | Baseline characteristics between groups |
| | | previous | | | Measures: | -PEG 3350 (n=37): | not compared |
| | | gastrointestin | | | | 28.5 ± 4.2 | Method of randomisation and allocation |
| | | al surgery, | | | -Stool | | concealment not described |
| | | renal; or heart | | | frequency | -Lactulose (n=37): | Non blinded study |
| | | failure, bowel | | | | 26.2 ± 5.1 | Small sample size, no sample size |
| | | obstruction, | | | -Stool form | | calculation |
| | | ileus, | | | | Effectiveness (% | No follow-up period |
| | | pregnancy, | | | -Easy of | effective) | Intention to treat analysis not performed |
| | | lactation, | | | passage | -PEG 3350 (n=37): | 15.9 % dropout rate |
| | | galactosemia, | | | | 84 | Results not controlled for potential |
| | | diabetes | | | -Effectiveness | | confounders |
| | | mellitus | | | (global | -Lactulose (n=37): 46 | |
| | | | | | assessment, as | p=0.002 | Source of funding: |
| | | | | | reported by | | not stated |
| | | | | | parent or | Laxative preference | |
| | | | | | guardian) | (% preferred): | |
| | | | | | | -PEG 3350 (n=37): | |
| | | | | | -Laxative | 73 | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|---|--------------------------|-------------------|
| | | | | | preference (based on efficacy, ease of administration and side effects) | -Lactulose (n=37): 27 | |
| | | | | | | | |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|------------------------|---|
| | Level | | S | | Measures | | |
| Wald et al. | Study Type: | 50 children | 50 children | Intervention: | Duration of | Children in remission | Additional information from study: |
| Evaluation of | RCT | | | Biofeedback , one | | | At baseline 2 groups comparable |
| biofeedback in | | <u>Inclusion</u> | 40 boys | 25 to 30-minute | 12 weeks | <u>(%)</u> | respect to age, sex, duration and |
| childhood | <u>Evidence</u> | criteria: | Age range 6 to | session | | | severity of soiling, anorectal motility |
| encopresis. | level: | encopresis of | 15 years (mean | | <u>Assessment</u> | | parameters and expulsion patterns |
| 1987. Journal of | 1- | at least 6 | 8.4) | Children with | point (s): | as exact figures not | |
| Pediatric | | months of | | abnormal | Immediately | reported in text) | Single blinded design |
| Gastroenterolog | | duration | Country: USA | expulsion pattern | after treatment | | |
| y and Nutrition | to evaluate | | | taught a | completed | -3 months: | Initial and follow-up office visits at 2, 4 |
| 6[4], 554-558 | the efficacy of | | | technique to | | biofeedback (n=24): | and 8 weeks similar in duration for both |
| | biofeedback | criteria: | | normalise their | Follow-up | 54 | groups. All outcomes recorded by |
| | | not stated | | patterns and they | period: | | parents in written calendar. Follow-up |
| | encopresis | | | and children with | 6 and 12 | mineral oil (n=26): 54 | interviews by telephone performed at 3, |
| | | | | normal expulsion | months after | | 6 and 12 months by investigator |
| | | | | pattern told to use | | -6 months: | unaware of treatment or results of |
| | | | | the technique | finished | biofeedback (n=24): | anorectal studies |
| | | | | whenever they | | 50 | |
| | | | | attempted to | <u>Outcome</u> | | Based on outcomes, children placed in |
| | | | | defecate | Measures: | | groups at each assessment: 1-some |
| | | | | | -frequency of | | improvement, 2-some improvement, but |
| | | | | Reinforcement | defecation | -12 months: | major soiling (<1/week), 3-marked |
| | | | | sessions at 2, 4 | | biofeedback (n=24): | improvement (rare major soiling |
| | | | | and 8 weeks | -frequency of | 50 | <1/week or minor soiling) 4-complete |
| | | | | | gross | | remission |
| | | | | Comparison: | incontinence | mineral oil (n=26): 59 | |
| | | | | Mineral oil orally | | | 2 dropouts at 3 months (1 from each |
| | | | | in graded | -frequency of | | group), 3 additional dropouts at 6 |
| | | | | amounts (range 1 | staining or | period | months (2 biofeedback) and 5 lost to |
| | | | | to 4 | minor soiling | | follow-up at 12 months (3 biofeedback). |
| | | | | tablespoons/day), | | No significant | All dropouts designated as treatment |
| | | | | designed to | -parental | differences in | failures for each subsequent |
| | | | | induce a soft | perception of | | assessment point |
| | | | | bowel movement | clinical status | with abnormal | |
| | | | | daily | and overall | | Reviewer comments: |
| | | | | | satisfaction | | No clear definition of encopresis given |
| | | | | | | expulsion patterns | Method of randomisation and allocation |
| | | | | | | | concealment not described |
| | | | | | | | No sample size calculation. ITT analysis |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|-------------|---|
| | | | | | | | apparently performed Unclear how the 4 outcomes groups were defined from the clinical variables |
| | | | | | | | Source of funding: not stated |
| | | | | | | | |
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| | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|------------------|-------------------|------------------|------------------|--------------------|-----------------------------|-----------------------|--|
| Information | Evidence Level | Patients | Characteristic s | Comparison | Outcome Measures | | |
| Thomson et al. | Study Type: | 51 children | 51 children | Intervention: | Duration of | Number of complete | Additional information from study: |
| Polyethylene | RCT (cross | | 29 girls | PEG + E (6.9 g | treatment: | defecations per week | Chronic constipation defined according |
| glycol 3350 plus | over, | <u>Inclusion</u> | mean age 5.4 | powder/sachet) | 2 weeks each | (Mean (SD), range) | to Rome criteria as fewer than 3 |
| electrolytes for | multicentre) | criteria: | years (range: | | treatment | (data do not include | complete bowel movements/week, and |
| chronic | | chronic | 24 months to 11 | Comparison: | period | washout period) | at least 1 of the following: pain on |
| constipation in | <u>Evidence</u> | constipation | years) | Placebo (6.9 g | separated by a | | defecation on at least 25% of days; at |
| children: a | level: | for at least 3 | | powder/sachet) | | a. ITT population | least 25% of bowel movements with |
| double blind, | 1+ | months | Country: UK | | washout | | straining, and at least 25% of bowel |
| placebo | | | | | | -PEG+E (n = 47): | movements with hard or lumpy stools |
| controlled, | Study aim: to | Exclusion | | Washout period in | | 3.12 (2.050) | |
| crossover | assess the | criteria: | | between: 2 weeks | | 0.00-8.87 | Random sequence group computer |
| study.[erratum | efficacy and | current or | | | immediately | | generated before start of recruitment |
| | safety of | previous | | Dosing regime for | | -Placebo (n = 48) | using block size of 4 patients and study |
| | polyethylene | faecal | | both PEG + E and | treatment | 1.45 (1.202) | medication labelled accordingly. |
| Jan;93(1):93]. | glycol 3350 | impaction | | | period, | 0.00-3.73 | Random blocks (with numbers stored in |
| | plus | decided by | | sachets/day): | including | | sealed code-break envelopes) sent to |
| | electrolytes | either | | | washout | | investigator sites as required. As |
| Childhood | | physical | | -children aged 2 | | 1.64 | children enrolled, sites allocated |
| - L 3/ | | examination | | to 6 years | Follow-up | | treatment supplies sequentially, started |
| 1000 | of chronic | or abdominal | | days 1-2: 1 | period: | p Value (95% CI) | with lowest possible number. Both the |
| | constipation in | • . | | days 3-4: 2 (taken | | <0.001 (0.99 to 2.28) | children (and their parents/guardians) |
| | children | previous | | together) | made after | | and those administering treatment were |
| | | intestinal | | days 5-6: 3 (2 | treatment | b. PP population | blinded to allocation schedule |
| | | perforation/ob | | morning, 1 | completed | | |
| | | struction, | | evening) | | -PEG+E (n = 36): | A sample size of 50 children was |
| | | paralytic | | days 7-8: 4 (2 | <u>Outcome</u> | 3.63 (1.980) | planned to achieve 40 evaluable |
| | | ileus, | | morning, 2 | Measures: | 0.00–8.87 | children, giving 90% power to detect a |
| | | Hirschsprung' | | evening) | | | true treatment difference of 0.3 bowel |
| | | s disease, | | | 1. Primary | -Placebo (n = 36): | movements/week using a two-tailed |
| | | severe | | -children aged 7 | efficacy | 1.63 (1.229) | significance test at the 5% level. As |
| | | inflammatory | | to 11 years | endpoint: | 0.00-3.73 | dropout rate was higher than originally |
| | | conditions of | | days 1-2: 2 (taken | | | estimated, recruitment target was |
| | | the intestinal | | together) | -number of | | increased to 60 children |
| | | tract, severe | | days 3-4: 2 (taken | • | 1.96 | |
| | | gastroesopha | | together) | defecations per | | At baseline, clinically significant |
| | | geal reflux, | | days 5-6: 5 (2 | week | <0.001 (1.19 to 2.72) | abnormalities |
| | | diabetes, | | morning, 3 | | | on physical examination (mainly |
| | | receiving | | evening) | Secondary | (95% CI, 95% | associated with faecal loading but not |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|------------------------------|---------------------------|-----------------------------------|---------------------|--|--|
| | Level | | s | • | Measures | | |
| | | doses of | | days 7-8: 6 (3 | efficacy | confidence interval; | impaction) recorded for 8 children (5/27 |
| | | stimulant | | morning, 3 | outcomes: | | in the PEG+E/placebo group, 3/24 in the |
| | | laxatives | | evening) | | PP per protocol) | placebo/PEG+E group). Before |
| | | considered by | | | -total number of | | randomisation, 47 children taking other |
| | | local | | For both groups if | defecations | Secondary efficacy | laxatives (most frequently lactulose) |
| | | observers to be at higher | | diarrhoea, doses was decreased by | noin on | outcomes, ITT population (mean, | 13/51 children (7/27 in the |
| | | end of their | | | | SD) | PEG+E/placebo |
| | | own doses | | a day. If loose | delecation | <u>3D)</u> | group, 6/24 in the placebo/PEG+E |
| | | spectrum | | stools doses | -straining on | a. Total number of | group) recorded at least one deviation |
| | | | | | | defaecations | from the study protocol (1 child recorded |
| | | | | sachet | | | 2 protocol deviations). Main reason for |
| | | | | | | | deviation was non-compliance with |
| | | | | | consistency | (2.771) | study medication (7/51 children), |
| | | | | | | Placebo* (n = 47): | followed by failure to supply sufficient |
| | | | | | | 4.10 (2.503) | bowel movement data (4/51 children), |
| | | | | | hard stools | Treatment difference: | and taking concomitant non-study |
| | | | | | | 1.58 p Value (95% CI)= | laxative medication after randomisation (3/51 children). |
| | | | | | | 0.003 (0.55 to 2.60) | (3/31 children). |
| | | | | | on delecation | 0.003 (0.33 to 2.00) | Reviewer comments: |
| | | | | | -faecal | b. Pain on | Blinding procedures not clearly |
| | | | | | | defaecation | described |
| | | | | | | PEG+E (n = 47): 0.49 | Unclear whether outcomes assessors |
| | | | | | | (0.727) | were also blinded to treatment allocation |
| | | | | | | Placebo (n = 47): | Study not controlled for potential |
| | | | | | | 0.77 (0.863) | confounders |
| | | | | | | Treatment difference: | 0 (() |
| | | | | | | -0.28 | Source of funding: |
| | | | | | | p Value (95% CI): 0.041 (–0.52 to – | Norgine Ltd. One of the authors was an employee of Norgine Ltd at the time the |
| | | | | | | 0.041 (-0.52 to - | study was written. The others declared |
| | | | | | | 0.01) | that they had nothing to declare |
| | | | | | | c. Straining on | and any ridd fielding to dooldro |
| | | | | | | defaecation | |
| | | | | | | PEG+E (n = 47): 0.72 | |
| | | | | | | (0.789) | |
| | | | | | | Placebo (n = 47): | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | 1.37 (1.041) Treatment difference: -0.65 p Value (95% CI): 0.001 (-0.97 to - 0.33) | |
| | | | | | | d. Stool consistency PEG+E (n = 47): 1.73 (0.497) Placebo (n = 47): 2.21 (0.556) Treatment difference: -0.48 p Value (95% CI): 0.001 (-0.68 to - 0.27) | |
| | | | | | | e. Percentage hard stools PEG+E (n = 47): 14.64 (26.041) Placebo (n = 47): 38.19 (39.508) Treatment difference: -23.55 p Value (95% CI): <0.001 | |
| | | | | | | f. Abdominal pain on defaecation PEG+E (n = 47): 0.67 (0.789) Placebo (n = 47): 0.79 (0.903) Treatment difference: 20.12 p Value (95% CI) | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | NS | |
| | | | | | | | |
| | | | | | | g. Faecal incontinence | |
| | | | | | | PEG+E (n = 47): 4.70 | |
| | | | | | | (6.344) | |
| | | | | | | Placebo (n = 47): | |
| | | | | | | 4.85 (7.863) | |
| | | | | | | Treatment difference: | |
| | | | | | | 20.15 | |
| | | | | | | p Value (95% CI) | |
| | | | | | | NS | |
| | | | | | | Maan offactive door | |
| | | | | | | Mean effective dose of PEG 3350 | |
| | | | | | | (g/kg/day): | |
| | | | | | | 0.6 (2 to 6-year-old) | |
| | | | | | | 0.7 (7 to 11-year-old) | |
| | | | | | | , | |
| | | | | | | Adverse events: | |
| | | | | | | PEG+E (31/49, 63%) | |
| | | | | | | Placebo (28/49, 57%) | |
| | | | | | | during periods I and | |
| | | | | | | III. None serious, | |
| | | | | | | most judged by | |
| | | | | | | investigator to be | |
| | | | | | | moderate or mild in | |
| | | | | | | severity | |
| | | | | | | 20 children (41%) on | |
| | | | | | | PEG+E: 41 events | |
| | | | | | | 22 children (45%) on | |
| | | | | | | placebo: 45 events, | |
| | | | | | | judged by investigator | |
| | | | | | | to be at least possibly | |
| | | | | | | related to the study | |
| | | | | | | treatment. Most | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|-------------------------|-------------------|
| | Level | | s | • | Measures | | |
| | | | | | | gastro-intestinal | |
| | | | | | | disorders (particularly | |
| | | | | | | abdominal pain), | |
| | | | | | | PEG+E (39%, 39 | |
| | | | | | | events); placebo | |
| | | | | | | (45%, 41 events). 1 | |
| | | | | | | child in | |
| | | | | | | placebo/PEG+E | |
| | | | | | | group withdrawn at | |
| | | | | | | week 3 because of | |
| | | | | | | abdominal pain, | |
| | | | | | | assessed by | |
| | | | | | | investigator as being | |
| | | | | | | related to treatment, | |
| | | | | | | this child was taking | |
| | | | | | | placebo at the time of | |
| | | | | | | withdrawal. New | |
| | | | | | | clinically significant | |
| | | | | | | abnormalities on | |
| | | | | | | physical examination | |
| | | | | | | (mainly associated | |
| | | | | | | with faecal loading): | |
| | | | | | | 13 children (8/27 in | |
| | | | | | | the PEG+E/placebo | |
| | | | | | | group, 5/24 in the | |
| | | | | | | placebo/ | |
| | | | | | | PEG+E group). When | |
| | | | | | | analysed for what | |
| | | | | | | these children were | |
| | | | | | | taking for the 2 weeks | |
| | | | | | | before the physical | |
| | | | | | | examination, 23 out | |
| | | | | | | of the 24 reports | |
| | | | | | | (95.8%) occurred | |
| | | | | | | when child taking | |
| | | | | | | placebo. Only 1 | |
| | | | | | | report of an abnormal | |
| | | | | | | abdominal | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|--|-------------------|
| | | | | | | examination while patient on PEG+E | |
| | | | | | | Mean weight similar before and after treatment, no significant difference found between the 2 groups for change in weight while on treatment (p=0.357) | |
| | | | | | | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|-------------------------|---|
| Sondheimer et | Level Study Type: | 37 children | s 37 children | General: | Measures Duration: | Daily bowel | Additional information from study: |
| al. Lubricant | RCT | 37 Gillaren | 37 Gillaten | 5-day course of | Unclear, | movement (% | Diagnosis of chronic functional |
| versus laxative | KCI | <u>Inclusion</u> | 26 male | oral bisacodyl | probably 6 | patients) | constipation made on basis of historical |
| in the treatment | Evidence | criteria: | 20 maie | (most patients) | months | <u>patierits)</u> | features and physical exam |
| of chronic | level: | patients | age range: 3 to | and daily enema | monuis | at 1 month: N.S | demonstrating dilated rectum, excessive |
| functional | 1- | treated for | 12 years | for 3-5 days in | Assessment | at Thonas N.S | retained stool directly within anal verge |
| constipation of | • | chronic | 12 years | addition (a | point (s): | at 3 months: | and in most cases, evidence of perianal |
| children: a | | functional | Country: | minority) | 1, 3 and 6 | at 3 months. | soiling |
| comparative | | constipation | USA | Tillionty) | months after | -Mineral oil (n=18): | Soming |
| study. 1982. | | in specialist | OOA | Intervention: | initiating | 100 | Children assigned to 1 of 2 treatment |
| Journal of | mineral oil | clinic | | | treatment | -Senokot (n=18): 72 | groups according to the last digit of their |
| Pediatric | and | Cili liC | | twice daily in | licalificit | p<0.05 | hospital number. All patients seen by |
| Gastroenterolog | | Exclusion | | doses sufficient to | Follow-up | p<0.03 | same physician. Parents informed that 1 |
| y and Nutrition | senna | criteria: | | induce loose | period: | latest follow-up: | of 2 acceptable medications would be |
| 1[2], 223-226 | | neurological | | stools and | -Mineral oil | latest follow up. | used to accomplish the discussed |
| 1[2], 220 220 | | impairment, | | | group, mean | -Mineral oil (n=18): 89 | objectives |
| | | faecal soiling | | | 10.1 months | -Senokot (n=18): 50 | objectives |
| | constipation in | 0 | | week of | | p<0.05 | No significant baseline differences |
| | children | absence of | | treatment, dose | -Senokot group, | P 10:00 | between 2 groups regarding mean age, |
| | ormaron | retained stool | | reduced until | mean 10.5 | Daily soiling (% | median age at onset of symptoms and |
| | | rotaliloa otool | | leakage ceased. | months | patients) | percent of patients who had received |
| | | | | This dose (range | THO THE TO | <u>patierney</u> | prior treatment with constipation, sex |
| | | | | 1.5 to 5.0 | Outcome | at 1 month: | ratio, faecal soiling, overt retentive |
| | | | | cc/kg/day) | Measures: | at i monan. | behaviour, enuresis, "difficult" toilet |
| | | | | maintained for | <u>modouroo.</u> | -Mineral oil (n=18): 11 | training and primary failure of toilet |
| | | | | minimum 3 | -daily bowel | -Senokot (n=18): 39 | training. |
| | | | | months. | movements | p<0.05 | g. |
| | | | | | | F 10.00 | Patients allowed to discontinue |
| | | | | Comparison: | -daily soiling | at 3months: | medications after 3 months if symptom |
| | | | | Senokot (tablet or | adii, coiii.g | | control unsatisfactory |
| | | | | syrup), doses | -compliance | -Mineral oil (n=18): 11 | ` ansaustasts. |
| | | | | sufficient to | with medication | -Senokot (n=18): 50 | 1 patient on mineral oil lost o follow-up |
| | | | | induce at least 1 | | p<0.05 | after 3-month visit and not considered in |
| | | | | bowel movement | | | results. No dropouts/lost to follow-up in |
| | | | | daily during first 2 | | latest follow-up: | other group |
| | | | | weeks of | | - T | 3 - 1 |
| | | | | treatment. This | | -Mineral oil (n=18): 6 | During 1rst month patients/parents kept |
| | | | | dose maintained | | | records of medication, stool frequency |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---|------------------------------------|---|--|
| | | | | for 3 months. Tapering accomplished by changing from daily to every other day and then every 3 rd day medication | | Compliance with medication (% reliably compliant) -Mineral oil (n=19): 68 -Senokot (n=18): 78 % successfully discontinued regular medication at latest follow-up: -Mineral oil (n=18): 55 -Senokot (n=18): 22 an additional 33% discontinued Senokot because of unacceptable | and faecal soiling. From then on outcomes measured by telephone interviews and during consultations Reviewer comments: Study inadequately randomised. Allocation concealment not described Clinicians/researchers not blinded. Blinding procedures for parents/patients not clearly described No sample size calculation performed Results not controlled for potential confounders Definition of "reliably compliant" not given Source of funding: not stated |
| | | | | | | F .0.0 ! | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|-----------------------------|--|
| | Level | | S | | Measures | | |
| Bu et al. | Study Type: | 45 children | 45 children | Intervention: | Duration of | <u>Defecation frequency</u> | Additional information from study: |
| Lactobacillus | RCT | | 23 male | MgO 50 mg/kg | treatment: | (times/day) | Chronic constipation defined as a stool |
| casei | | <u>Inclusion</u> | | per day, twice a | 4 weeks | -MgO (n=18) | frequency of <3 times/week for >2 |
| rhamnosus | <u>Evidence</u> | <u>criteria:</u> | | day | | 0.55 ± 0.13 | months and at least 1 of the following |
| Lcr35 in | <u>level:</u> | children | Age (months, | | <u>Assessment</u> | | minor criteria: anal fissures with |
| children with | 1+ | | mean, SD) | Comparison 1: | point (s): | -probiotic (n=18) | bleeding due to constipation, faecal |
| chronic | | years old with | | Lcr35 8 X 10^8 | Immediately | 0.57 ± 0.17 | soiling or passage of large and hard |
| constipation. | | | MgO group | c.f.u/day | after treatment | | stool |
| 2007. Pediatrics | | constipation | | (Antiobiophilus | completed | -placebo (n=9) | |
| International | the effect of | | Probiotic group | 250 mg, 2 | | 0.37 ± 0.10 | Children randomly assigned into the 3 |
| 49[4], 485-490 | Probiotics | <u>Exclusion</u> | | capsules, twice a | Follow-up | | groups according to a computer - |
| | (Lactobacillus | criteria: | Placebo group | day) | period: | MgO vs. probiotic NS | generated randomisation list |
| | case | organic | | | No follow up | Placebo vs. probiotic | |
| | rhamnosus, | causes of | | Comparison 2: | made after | P=0.006 | Blinding achieved by the use of 3 |
| | Lcr35) alone | constipation | Country: | Placebo (starch in | treatment | MgO vs. placebo | interventions with similar appearances |
| | in the | like | Taiwan | content) | finished | p=0.01 | and placed into identical capsules, |
| | | Hirschsprung' | | | | | which were either swallowed o as a |
| | chronic | s disease, | | | <u>Outcome</u> | Hard stool (%) | whole or opened and the contents of the |
| | constipation in | spina bifida | | | Measures: | -MgO (n=18) | capsule administered in milk or fluid |
| | children and | (occulta), | | | | 23.5 ± 7.9 | |
| | to compare | hypothyroidis | | Lactulose use | -frequency of | | Throughout the duration of study all |
| | the effect with | m, or other | | (1mL/kg/day) | defecation | -probiotic (n=18) | investigators, participants and data |
| | magnesium | metabolic/ren | | allowed when no | | 22.4 ± 14.7 | analysts were blinded to the assigned |
| | oxide (MgO) | al | | stool passage | -consistency of | | treatment |
| | and placebo, | abnormalities, | | noted for 3 days. | stools | -placebo (n=9) | |
| | respectively | drugs | | Glycerin enema | | 75.5 ± 6.1 | Sample size determined by doing |
| | | influencing | | used only when | -episodes of | | primary trial with 9 patients using non- |
| | | gastrointestin | | no defecation for | soiling | MgO vs. probiotic NS | inferiority to test. Equivalent margin |
| | | al function | | >5days or | _ | Placebo vs. probiotic | chosen with reference to effect of active |
| | | other than | | abdominal pain | -episodes of | p=0.02 | control in the data of preliminary trial. |
| | | laxatives | | suffered due to | abdominal pain | MgO vs. placebo | Unbalance design of allocation number |
| | | (calcium | | stool impaction | | p=0.03 | used for more interest in the new drug |
| | | channel | | | -use of | | (Lcr35): allocation rate set at 2:2:1. One |
| | | blockers, | | | lactulose or | Abdominal pain | sided significance level set at 0.05 and |
| | | antidysrythmi | | | enema | (times) | power was 80%. Under these |
| | | c agents, | | | | -MgO (n=18) | assumptions the smallest sample size |
| | | anticonvulsiva | | | | 4.8 ± 3.7 | was 45 and the sample size of MgO, |
| | | nts, | | | | | Lcr35 and placebo was 18, 18 and 9 |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|--------------------------------|---|
| | Level | | S | | Measures | | |
| | | antidepressan | | | | -probiotic (n=18) | respectively |
| | | ts, | | | | 1.9 ± 1.6 | |
| | | anticholinergi | | | | | No significant differences at baseline |
| | | c agents) | | | | -placebo (n=9) | amongst the 3 group regarding: sex, age |
| | | | | | | 6.7 ± 3.3 MgO vs. probiotic | of enrolment, age of onset of constipation, duration of constipation, |
| | | | | | | p=0.04 | previous treatment, defecation period, |
| | | | | | | Placebo vs. probiotic | stool consistency, abdominal pain, |
| | | | | | | p=0.01 | faecal soiling, bleeding during |
| | | | | | | MgO vs. placebo NS | defecation, use of enema, taking fruit or |
| | | | | | | Ingo vo. placobo ivo | vegetable daily |
| | | | | | | Use of glycerine | 1.19 |
| | | | | | | enema (times) | Patients asked to discontinue any |
| | | | | | | -MgO (n=18) | laxatives previously prescribed 3 days |
| | | | | | | 1.3 ± 1.9 | before entering protocol, and also asked |
| | | | | | | | to avoid any other probiotics, yogurt or |
| | | | | | | -probiotic (n=18) | beverage containing probiotics for at |
| | | | | | | 1.6 ± 1.9 | least 2 weeks before treatment and during therapy |
| | | | | | | -placebo (n=9) | during thorapy |
| | | | | | | 4.0 ± 2.1 | All outcomes measures recorded by |
| | | | | | | | parents in a stool diary |
| | | | | | | MgO vs. probiotic NS | , |
| | | | | | | Placebo vs. probiotic | 4 patients discontinued medication |
| | | | | | | p=0.04 | during study period: 2 in MgO, 1 in |
| | | | | | | MgO vs. placebo | probiotic, 1 in placebo group (2 patients |
| | | | | | | p=0.03 | suffered from acute gastroenteritis and |
| | | | | | | | 2 patients lost to follow-up) |
| | | | | | | No significant | |
| | | | | | | differences regarding | Reviewer comments: |
| | | | | | | use of lactulose, | Allocation concealment not described |
| | | | | | | faecal soiling and | Not clear whether the 2 patients who |
| | | | | | | change of appetite | suffered from acute gastroenteritis had it |
| | | | | | | amongst 3 groups | as consequence of the study medication Study not controlled for potential |
| | | | | | | Patients with | confounders |
| | | | | | | treatment success | Comounders |
| | | | | | | (%) | Source of funding: not stated |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|---|-------------------|
| | Level | | S | | Measures | -MgO (n=18): 72.2 -probiotic (n=18): 77.8 -placebo (n=9): 11.1 MgO vs. probiotic NS Placebo vs. probiotic p=0.01 MgO vs. placebo p=0.01 no adverse effects noted in probiotic and placebo groups, only 1 patient in the MgO group suffered from mild diarrhoea | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|-----------------------|--|
| | Level | | S | | Measures | | |
| Loening- | Study Type: | 49 children | -Miralax group: | Intervention: | Duration of | Bowel movement | Additional information from study: |
| Baucke. | Prospective | | 28 children | MiraLax | treatment: | frequency (mean, | Initial dose of Miralax 0.5 g/kg daily |
| Polyethylene | cohort | <u>Inclusion</u> | 20 boys | 17 dissolved | 12 months | results are estimates | suggested for children whose rectums |
| glycol without | | criteria: | Mean age ± SD: | in 240 mL of a | | taken form bar chart | were loaded with stool but who had no |
| electrolytes for | <u>Evidence</u> | children ≥4 | 8.7 ± 3.6 years | | <u>Assessment</u> | as not reported in | fecal abdominal masses at the initial |
| children with | <u>level:</u> | years of age | Range 4.1 to | juice or Kool-Aid | point (s): | text) | physical examination and no history of |
| constipation | 2+ | referred for | 17.5 years | initial dose: | 1, 3, 6, and 12 | | long intervals between huge bowel |
| and encopresis. | | functional | | 0.5 to 1 g/kg/daily | months after | -baseline: | movements. Those with |
| 2002. Journal of | | constipation | -MOM group: | | initiating | PEG: 3.2 | palpable abdominal fecal masses or |
| Pediatric | to determine | and | 21 children | Comparison: | treatment | MOM: 2.5 | history of infrequent huge bowel |
| Gastroenterolog | the efficiency, | encopresis | 17 boys | MOM | | | movements started on 1 g/kg daily |
| y and Nutrition | acceptability, | Functional | Mean ± SD: 7.3 | Initial dose: | Follow-up | -1 month | |
| 34[4], 372- | and treatment | constipation | ± 3.0 years | 1 to 2.5 mL/kg | period: | PEG: 9.0 | Milk of Magnesia given if family could |
| 377United | dosage of | defined as | Range: 4.0 to | | No follow-up | MOM: 6.5 | afford only the use of a cheaper laxative |
| States. | MiraLax | delay/difficulty | 13.9 years | | made after | | or if child had previously received MOM |
| | (polyethylene | in defecation | - | | treatment | -3 months | without refusal. For these children, MOM |
| | glycol 3350 | and | Country: | | finished | PEG: 9.5 | reintroduced or adjusted to adequate |
| | without | encopresis | USA | | | MOM: 7.0 | dosage. Parents told how to improve the |
| | electrolytes) | (≥1/week) for | | | Outcome | | taste by mixing the child's preferred |
| | during a 12- | more than 1 | | Large laxative | Measures: | -6 months | flavoring with plain MOM. Initial daily |
| | month | year | | dosages divided | | PEG: 8.8 | dosage of 1 mL/kg body weight |
| | treatment | | | into 2 daily doses. | -bowel | MOM: 6.3 | suggested for children with rectal fecal |
| | period in | Exclusion | | Parents told to | movement | | masses only at initial evaluation and if |
| | children with | criteria: | | adjust the dose of | frequency | -12 months | no history of infrequent large bowel |
| | functional | Children <4 | | medication by 30 | | PEG: 6.8 | movements. Dosage of 2.5 mL/kg |
| | constipation | years of age; | | mL for MiraLax | -consistency of | MOM: 7.2 | prescribed for those with fecal |
| | and | children who | | and by 7.5 mL | stools | | abdominal masses at the initial |
| | encopresis | refused the | | (one-half | | P<0.01 when | evaluation or history of huge, infrequent |
| | • | toilet for | | tablespoon) for | -soiling | comparing values at | bowel movements. |
| | | stooling but | | MOM every 3 | frequency | every assessment | |
| | | who had no | | days to a dosage | | point to baseline for | Regular stool sittings for 5 minutes after |
| | | constipation, | | that resulted in 1 | -abdominal pain | both treatments | each meal required for initial months |
| | | Hirschsprung' | | to 2 soft bowel | frequency | | |
| | | s disease, | | movements/day | ' ' | Soiling frequency | Patients and parents provided with diary |
| | | chronic | | and prevented | -medication | (mean, results are | sheets to record each outcome |
| | | intestinal | | soiling and | dosage | estimates taken form | measured |
| | | pseudo- | | abdominal pain. | | bar chart as not | |
| | | obstruction, | | If child retained | -clinically | reported in text) | Doing well defined as 3 or more bowel |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|---------------------|---------------------------------------|--|
| | Level | | s | | Measures | | |
| | | or | | stools despite | significant side | | movements/week and 2 or fewer soiling |
| | | previous | | compliance with | effects | -baseline: | episodes / month. Improved defined as |
| | | surgery of the | | assigned laxative, | | PEG: 12.0 | 3 or more bowel movements / week and |
| | | colon or anus | | daily senna added | | MOM: 8.5 | more than 75% decrease in soiling but |
| | | | | to treatment. | with medication | | not more than 1 soiling / week. Not |
| | | | | | | -1 month | doing well defined as fewer than 3 bowel |
| | | | | | | PEG: 3.0 | movements / week, less than 75% |
| | | | | | | MOM: 0.5 | decrease in soiling frequency, use of senna, or refusal to take the assigned |
| | | | | | | -3 months | laxative. Recovered defined as 3 or |
| | | | | | | PEG: 1.8 | more bowel movements / week and 2 or |
| | | | | | | MOM: 0.2 | fewer soiling episodes / month while not taking laxatives. |
| | | | | | | -6 months | |
| | | | | | | PEG: 1.0 | No significant baseline differences |
| | | | | | | MOM: 0.8 | between 2 groups |
| | | | | | | -12 months | Reviewer comments: |
| | | | | | | PEG: 0.9 | No sample size calculation performed |
| | | | | | | MOM: 0.1 | |
| | | | | | | | Outcomes for consistency of stools not |
| | | | | | | P<0.01 when | reported |
| | | | | | | comparing values at | Nick according to the chinical background |
| | | | | | | every assessment | Not reporting on the clinically significant |
| | | | | | | point to baseline for both treatments | side effects (or lack of them) for MOM |
| | | | | | | P<0.01 when | Source of funding: |
| | | | | | | comparing values | Dr. Loening-Baucke recipient of grant |
| | | | | | | between 2 groups at | support from Braintree Pharmaceuticals, |
| | | | | | | 1 and 12 months | Braintree, MA, U.S.A., for continuing |
| | | | | | | | studies on childhood constipation |
| | | | | | | Children with | |
| | | | | | | abdominal pain (%): | |
| | | | | | | -baseline: | |
| | | | | | | PEG: 61 | |
| | | | | | | MOM: 81 | |
| | | | | | | -1 month | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|------------------------|---|-------------------|
| | Level | | S | | Measures | DE0 44 | |
| | | | | | | PEG: 14 | |
| | | | | | | MOM: 14 | |
| | | | | | | O ma a matha a | |
| | | | | | | -3 months | |
| | | | | | | PEG: 13 MOM: 5 | |
| | | | | | | IVIOIVI. 5 | |
| | | | | | | -6 months | |
| | | | | | | PEG: 8 | |
| | | | | | | MOM: 11 | |
| | | | | | | IVIOIVI. I I | |
| | | | | | | -12 months | |
| | | | | | | PEG: 4 | |
| | | | | | | MOM: 0 | |
| | | | | | | IVIOWI. 0 | |
| | | | | | | P<0.01 when | |
| | | | | | | comparing values at | |
| | | | | | | every assessment | |
| | | | | | | point to baseline for | |
| | | | | | | both treatments | |
| | | | | | | | |
| | | | | | | Medication dosage | |
| | | | | | | (Mean doses and | |
| | | | | | | range for children | |
| | | | | | | who were doing well | |
| | | | | | | or improved) (PEG, | |
| | | | | | | g/kg; MOM, mL/kg) | |
| | | | | | | | |
| | | | | | | 1 month | |
| | | | | | | PEG: | |
| | | | | | | $0.6 \pm 0.2 \ (0.3 \ \text{to} \ 1.1)$ | |
| | | | | | | MOM: | |
| | | | | | | 1.4 ± 0.6 (0.6 to 2.6) | |
| | | | | | | | |
| | | | | | | 3 months | |
| | | | | | | PEG: | |
| | | | | | | $0.6 \pm 0.3 (0.3 \text{ to } 1.4)$ | |
| | | | | | | MOM: | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | 1.2 ± 0.5 (0.6 to 2.4) | |
| | | | | | | 12 months | |
| | | | | | | PEG: | |
| | | | | | | $0.4 \pm 0.1(0.1 \text{ to } 0.7)$ | |
| | | | | | | MOM: | |
| | | | | | | only 2 children still | |
| | | | | | | required MOM. Their | |
| | | | | | | dosages were 0.4 | |
| | | | | | | and 1.6 mL/kg, both | |
| | | | | | | less than the initial treatment dosage. | |
| | | | | | | liteatifierit dosage. | |
| | | | | | | mean doses for both | |
| | | | | | | treatments at 12 | |
| | | | | | | months | |
| | | | | | | did not differ | |
| | | | | | | significantly between | |
| | | | | | | children with or | |
| | | | | | | without initial | |
| | | | | | | palpable abdominal | |
| | | | | | | faecal masses. None | |
| | | | | | | of the patients required an increased | |
| | | | | | | dosage of either | |
| | | | | | | medication over time | |
| | | | | | | | |
| | | | | | | 5 children received a | |
| | | | | | | stimulant laxative in | |
| | | | | | | addition to PEG and | |
| | | | | | | 1 child received a | |
| | | | | | | stimulant laxative | |
| | | | | | | in addition to MOM (P | |
| | | | | | | > 0.2) | |
| | | | | | | Clinically significant | |
| | | | | | | side effects | |
| | | | | | | <u> </u> | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | PEG: no significant clinical side effects. Some children had diarrhea. None of the children in the PEG group became dehydrated. Children receiving PEG and | |
| | | | | | | their parents did not report increased flatus, abdominal distention, or new onset of abdominal pain Compliance with | |
| | | | | | | medication: -PEG: No children reported disliking the taste, no parents reported that child refused to take it in juice or Kool-Aid | |
| | | | | | | Parental noncompliance with administering the laxative and supervising toilet use: 14% children | |
| | | | | | | -MOM: 33% children refused to take it Parental noncompliance with | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | administering the laxative and supervising toilet use: 4% children | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|------------------------|--------------------------------|-----------------------------|------------------------------------|-------------------------|---|
| Urganci et al. A | Study Type: | 40 patients | 40 patients | Intervention: | Duration of | Stool consistency | Additional information from study: |
| comparative | RCT | | 22 male | Liquid paraffin | treatment: | (mean ± SD) | Diagnosis of constipation based on |
| study: the | | <u>Inclusion</u> | mean age 3.7 ± | | 8 weeks | | symptoms of ay least 3 months duration |
| efficacy of liquid | | criteria: | 2.7 years | Comparison: | | -first 4 weeks: | including at least 2 of the following: hard |
| paraffin and | <u>level:</u> | children 2 to | | Lactulose | | Liquid paraffin (n=20): | stool, painful defecation, rectal bleeding, |
| lactulose in | 1- | 12 years old | | | | 2.17 ± 0.5 | encopresis and < 3 bowel |
| management of | | referred for | | | | Lactulose (n=20): | movements/week |
| chronic | Study aim: | evaluation of | Country: | Medication | | 1.71 ± 0.5 | |
| functional | | constipation | Turkey | administered | after initiation of | p<0.01 | Open-label randomised study |
| constipation. | and compare | with evidence | | orally as a | treatment | | |
| | efficacy, | of faecal | | suspension at 1 | | -last 4 weeks: | Children also met with a nutritionist, |
| International | safety and | impaction | | mL/kg, twice daily | | | were given instructions to increase daily |
| 47[1], 15-19 | optimal dose | | | for each drug | | 2.29 ± 0.2 | fibre intake to amount of gram equal to |
| | of liquid | <u>Exclusion</u> | | | | Lactulose (n=20): | their age plus 10, parent asked to have |
| | paraffin and | criteria: | | | | 2.21 ± 0.4 | children sit on the toilet 4 times daily |
| | lactulose in | Hirschsprung' | | of best dose for | | N.S | after meals |
| | | s disease, | | , | finished | | |
| | | hypothyroidis | | parents asked to | | Stool frequency | Stool frequency and stool consistency |
| | | m, mental | | increase or | <u>Outcome</u> | (mean ± SD) (per | recorded by parents in daily diary forms. |
| | constipation | deficiency, chronic | | decrease the volume of each | Measures: | week) | Stool consistency scoring: 1, hard; 2, firm; 3, loose |
| | | debilitating | | drug by 25% | -stool | -first 4 weeks: | |
| | | diseases, | | every 3 days as | consistency | | No significant baseline differences |
| | | neurological | | required, to yield | | 13.3 ± 4.2 | between 2 groups |
| | | abnormalities, | | 2 firm-loose stools | -stool frequency | Lactulose (n=20): | |
| | | previous | | per day. | | 10.2 ± 4.4 | Effective treatment defined as clearance |
| | | surgery of | | Maximum dose | -optimal dose of | p<0.05 | of impaction: more than 3 bowel |
| | | colon | | | drugs | | movements/week and improvement in |
| | | | | the study: 3 mL/kg | | -last 4 weeks: | stool consistency |
| | | | | per day for each | -compliance | Liquid paraffin (n=20): | |
| | | | | drug | rate | 16.1 ± 2.2 | Patients considered compliant if ≥ 80% |
| | | | | | | Lactulose (n=20): | of prescribed dose taken correctly. |
| | | | | | | 12.3 ± 6.6 | Patients instructed to take both empty |
| | | | | | | p<0.05 | and full containers to calculate amount |
| | | | | | | | of medication taken |
| | | | | | | Optimal dose of drugs | |
| | | | | | | (mean ± SD) | Reviewer comments: |
| | | | | | | (mL/kg/day) | Randomisation method not described |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|--|
| | Evidence | | Characteristic | | Outcome | -data reported in table, assumed that for the whole study period: Liquid paraffin (n=20): 1.88 ± 0.27 Lactulose (n=20): 2.08 ± 0.27 N.S | No sample size calculation performed No clear definition of "evidence of faecal impaction" given Apparently no children dropped out the study/were lost to follow-up Study not controlled for potential confounders Source of funding: not stated |
| | | | | | | -first 4 weeks: Liquid paraffin (n=20): 95 Lactulose (n=20): 90 N.S -end of 8 weeks: Liquid paraffin (n=20): 90 Lactulose (n=20): 60 p=0.02 Adverse effects: | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|-------------------------|-------------------|
| | | | | | | No patient stopped | |
| | | | | | | treatment because of | |
| | | | | | | adverse effects | |
| | | | | | | (adverse effects not | |
| | | | | | | reported). During first | |
| | | | | | | 4 weeks, taste | |
| | | | | | | aversion in 1 child on | |
| | | | | | | liquid paraffin and | |
| | | | | | | abdominal distension | |
| | | | | | | in 2 patients on | |
| | | | | | | lactulose influenced | |
| | | | | | | compliance. During | |
| | | | | | | last 4 weeks, poor | |
| | | | | | | symptom control in 5 | |
| | | | | | | patients, side-effects | |
| | | | | | | (abdominal distension | |
| | | | | | | and cramping) in 3 | |
| | | | | | | on lactulose, and | |
| | | | | | | watery stools in 2 on | |
| | | | | | | liquid paraffin | |
| | | | | | | influenced | |
| | | | | | | compliance | |
| | | | | | | | |
| | | | | | | | |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|----------------------|---|
| | Level | | S | | Measures | | |
| Berg et al. A | Study Type: | 44 children | 40 children | General: | Duration of | Severity of soiling: | Additional information from study: |
| controlled trial | Quasi RCT | | | Behavioural | treatment: | | Children randomly allocated to 1 of 3 |
| of 'Senokot' in | | <u>Inclusion</u> | mean age: 7.9 | treatment, | 3 months | -At 3 months: | treatment groups, A, B and virtually in a |
| faecal soiling | <u>Evidence</u> | criteria: | years (S.D. = | focusing on use of | | Senokot (n=14) | random fashion |
| treated by | <u>level:</u> | children who | 2.3) | the toilet and | Assessment | Placebo (n=11) | |
| behavioural | 1- | had soiling as | | freedom from | point (s): | No tablets (n=15) | No significant baseline differences |
| methods. 1983. | | a main | gender not | soiling | 3 months after | | between the 3 groups |
| Journal of Child | Study aim: | complaint | reported | | starting | NS between the 3 | |
| Psychology and | | and | | Intervention: | treatment | groups (outcomes | Psychiatrist and psychologists did not |
| Psychiatry and | | uncomplicate | Country: UK | Senokot | | not reported by | know which tablets actually contained |
| Allied | | d functional | | | Follow-up | group) | the laxative. Tablets made up in packs |
| Disciplines | therapy would | | | Comparison 1: | period: | | labelled A and B. |
| 24[4], 543-549 | | incontinence | | placebo tablets in | 6 months to 1 | Number of soiling- | |
| | own in the | after an initial | | similar dosage to | - | free children | Methods used in behavioural treatment: |
| | treatment of | assessment | | Senokot | entering trial | | identifying targets, discussing use of |
| | | and physical | | | (but after 3 | -Relieved (less than | rewards, star charting, reinforcement of |
| | persistent | examination | | Comparison 2: | months the | once/week or not at | using the toilet appropriately and staying |
| | faecal soiling | | | No medication | study was a | all) | clean, mainly by Mothers advised to |
| | | Exclusion | | | case series for | | avoid castigating children. Initially, |
| | | criteria: | | Children started | Senokot only, | Senokot (n=14): 5 | children taken to toilet 3 times a day, |
| | | not clearly | | on 1 tablet at | therefore not | (35%) | then prompted to go unaccompanied, |
| | laxative | stated | | night. On the next | reported here) | Placebo (n=11): 2 | then expected to go on own initiative |
| | as well | | | visit to the clinic, if | | (18%) | |
| | | | | no improvement | <u>Outcome</u> | | 4 children dropped out after only 1 or 2 |
| | | | | in 'use of the | Measures: | -Not relieved | visits |
| | | | | toilet' and 'being | | | |
| | | | | clean' on the | -severity of | Senokot (n=14): 9 | Severity of soiling rating: 0 = none, 1 = |
| | | | | charts dosage | soiling | Placebo (n=11): 9 | less than once a week, 2 = at least once |
| | | | | increased to 2 | | | a week but less than daily, 3 = daily |
| | | | | tablets. Number of | | NS between the 3 | |
| | | | | tablets increased | soiling-free | groups | Reviewer comments: |
| | | | | to 3 on following | children | | No definitions of soiling/functional faecal |
| | | | | visit if | | | incontinence given |
| | | | | improvement had | | | Inadequate randomisation |
| | | | | still not occurred. | | | Allocation concealment not described |
| | | | | When soiling | | | Soiling frequently apparently assessed |
| | | | | getting better and | | | by interviewing parent at time of |
| | | | | child using toilet | | | consultation |

| umber of Patient Patients Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|--|---|------------------------------------|-------------|---|
| | dosage kept the same. Once child going regularly to toilet and not soiling tablets stopped altogether | | | No sample size calculation performed Not clear whether the 4 children who dropped out had already received any study medication There is a mistake in the paper regarding outcomes for the "no tablets" groups, therefore not reported here Results not controlled for potential confounders Source of funding: Messrs Reckitt and Coleman provided the medication and gave their support in carrying out this trial |

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|-------------------|-------------------|------------------|------------------|---------------------|---------------------|------------------------|--|
| Information | Evidence Level | Patients | Characteristic s | Comparison | Outcome Measures | | |
| Nurko et al. | Study Type: | 103 children | 103 children | General: | Duration of | Proportion of children | Additional information from study: |
| PEG3350 in the | RCT | | | Behavioural | treatment: | who responded to | Chronic constipation diagnosed when |
| treatment of | (multicentre) | Inclusion | 69 boys | treatment: | 3 weeks | treatment (% | for at least 3 months there was a history |
| childhood | , | criteria: | J | instructions to sit | | children) | of <3 spontaneous bowel |
| constipation: a | Evidence | Children aged | mean age: 8.5 ± | on toilet for 10 | Assessment | Group 1 (n=26): 77 | movements/week and ≥ 1 associated |
| multicenter, | level: | | 3 years | minutes twice | point (s): | , , , | symptoms including: straining, hard |
| double-blinded, | 1+ | with chronic | | after meals, | 7 and 14 days | Group 2 (n=27): 74 | stools sensation of incomplete |
| placebo- | | constipation. | Country: | positive | after medication | , | evacuation, production of large bowel |
| controlled trial. | Study aim: | Patients | USA | reinforcement | started | Group 3 (n=26): 73 | movements that may obstruct the toilet |
| 2008. Journal of | To establish | taking other | | using age- | | | or painful defecation |
| Pediatrics | the efficacy | laxatives only | | appropriate | Follow-up | Placebo (n=24): 42 | |
| 153[2], 254-261 | and best | included if | | printed calendars | period: | , , | Faecal impaction defined as presence of |
| Nurko et al., | starting dose | they had <3 | | and special | N.A | P<0.04 each group | faecal hypogastric mass palpable on |
| 2008 | of | bowel | | stickers for days | | vs. placebo | abdominal examination and presence of |
| | polyethylene | movements/w | | without episodes | Outcome | P=0.026 all | hard stool on rectal examination. |
| | glycol (PEG) | eek while | | of faecal | Measures: | treatments groups vs. | diagnosis of faecal impaction made by 2 |
| | 3350 in the | taking the | | incontinence and | | placebo | independent observers, no |
| | short-term | laxative | | others with bowel | Efficacy: | NS between | disagreement found in the assessment |
| | treatment of | | | movements | - | treatment groups | of any patient |
| | children with | Exclusion | | | -primary | | |
| | functional | criteria: | | Intervention | outcome: | Weekly number of | Sample size calculation performed |
| | constipation | Taking a | | (Group 1): | | bowel movements | |
| | - | stable dose of | | Polyethylene | proportion of | (BM) | Patient randomly assigned in blinded |
| | | PEG3350, | | glycol (PEG) 3350 | children who | Group 1 (n=26): | fashion in a 1:1:1:1 ratio within each |
| | | evidence of | | Miralax): | responded to | Before 1.7±0.9 | participant site. Randomisation schedule |
| | | faecal | | 0.2g/kg per day- | treatment | | at each site constructed by using |
| | | impaction, | | single dose | | Group 2 (n=27): | random blocks of 20 patients, which |
| | | guiac- | | Maximum: 8.5 g | -secondary | Before 1.5±1.0 | provided balanced treatment |
| | | positive stool, | | per day | outcomes: | | assignments in order to ensure the |
| | | anorectal | | | | Group 3 (n=26): | specified treatment ratio |
| | | malformations | | Comparison 1 | weekly number | Before 1.5±0.5 | · |
| | | , | | (Group 2): | of bowel | | Miralax and placebo provided as a |
| | | Hirschsprung' | | Polyethylene | movements | Placebo (n=24): | powder containing flavouring in |
| | | s disease, | | glycol (PEG) 3350 | | Before 1.6±0.7 | identically labelled bottles reconstituted |
| | | myelomening | | Miralax): | weekly number | | with water to 4000 mL by study |
| | | ocele, | | 0.4g/kg per day- | of faecal | Overall difference | personnel in the pharmacy. Dosing |
| | | hypothyroidis | | single dose | incontinence | between treatment | calculated by pharmacy staff and water |
| | | m or other | | Maximum: 17 g | episodes | groups and placebo | added. All dose calculated to be given |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|---------------------------|---------------------------|------------------------------------|---------------------------------|--|
| | Levei | organic | S | per day | | p=0.017 | on a 10-mL/kg basis by pharmacy staff. |
| | | causes of | | | changes in | P=0.015 dose- | The blinded research team received the |
| | | constipation | | Comparison 2 | stool | response trend | reconstituted identical jugs, which were |
| | | | | (Group 3): | consistency | Maakk mumbanaf | distributed to patient's |
| | | | | Polyethylene | atrainin a | Weekly number of | parents/caregivers. No difference in |
| | | | | glycol (PEG) 3350 | straining | faecal incontinence | colour, appearance r taste amongst |
| | | | | Miralax): | | episodes mean ± SD) | different doses. Patients took single |
| | | | | | proportion of children who | Croup 1 (p. 26). | dose per day. No adjustment of study |
| | | | | single dose | | Group 1 (n=26): | medication allowed during study. No |
| | | | | Maximum: 34 g | responded to | Before 3.8±4.8 After 3.0±4.6 | other laxatives allowed during study |
| | | | | per day | treatment in the second week | Aller 3.0±4.6 | Comilian completed deily diamy that |
| | | | | Comparison 3: | Second week | Group 2 (n=27): | Families completed daily diary that included number and characteristics of |
| | | | | Placebo | Safety: | Before 3.5±4.9 | bowel movements an documentation of |
| | | | | riacebo | Salety. | After 1.8±2.6 | episodes of faecal incontinence |
| | | | | | -incidence and | Aiter 1.0±2.0 | episodes of faecal incontinence |
| | | | | | severity of | Group 3 (n=26): | Response to treatment defined as ≥3 |
| | | | | | adverse effects | Before 7.2±18.7 | bowel movements during the second |
| | | | | | adverse effects | After 3.5±7.8 | week of treatment. Patients considered |
| | | | | | | 7 (IIC) 0.0±1.0 | failures and withdrawn from study if they |
| | | | | | | Placebo (n=24): | had no bowel movements (BM) for 7 |
| | | | | | | Before 2.4±3.8 | days or developed faecal impaction at |
| | | | | | | After 1.4±3.7 | any point. |
| | | | | | | Alter 1.4±0.7 | arry point. |
| | | | | | | NS amongst different | No significant differences in baseline |
| | | | | | | groups | characteristics between the 4 groups |
| | | | | | | groupo | onaraciencies servicenturo i greape |
| | | | | | | Changes in stool | 14 patients did not complete the 2-week |
| | | | | | | consistency (mean ± | treatment: |
| | | | | | | SD) | -8 because of treatment failure (5 with |
| | | | | | | | impaction (2 Group 1, 3 Group 2), and |
| | | | | | | Group 1 (n=26): | 3 with > 7 days without a BM) (2 Group |
| | | | | | | Before 2.8±0.8 | 1, 1 Group 3)] |
| | | | | | | After 2.1±0.7 | - 3 because of adverse events (1 |
| | | | | | | | increased abdominal pain (placebo), 1 |
| | | | | | | Group 2 (n=27): | fever, malaise, headache (placebo), 1 |
| | | | | | | Before 2.6±0.9 | exacerbation bipolar (placebo)) |
| | | | | | | After 1.7±0.6 | - 1 withdrawal (lack of response |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|---|
| | | | | | | Group 3 (n=26): Before 2.9±0.7 After 1.5±0.7 | (placebo)) - 2 non compliance (1 Group 2, 1 Group 3) |
| | | | | | | Placebo (n=24): Before 3.0±0.8 After 2.4±0.9 | - 3 serious adverse events occurred requiring hospitalisation (2 cases impaction, 1 case of exacerbation of bipolar/depression) |
| | | | | | | P<0.003 each group vs. placebo | IIT analysis performed |
| | | | | | | | There were no significant predictors of success by controlling for age, duration of constipation, prior laxative use, presence of stool in rectum, sex and presence of faecal incontinence at |
| | | | | | | Straining scores (mean ± SD) | baseline |
| | | | | | | Group 1 (n=26): Before 2.3±1.1 After 1.4±0.9 | Source of funding: Supported in part by Braintree Laboratories Inc. |
| | | | | | | Group 2 (n=27): Before 1.9±1.2 After 1.0±1.0 | |
| | | | | | | Group 3 (n=26): Before 2.0±1.0 After 0.9±0.6 | |
| | | | | | | Placebo (n=24): Before 2.7±1.2 After 1.5±1.2 | |
| | | | | | | P<0.003 each group vs. placebo P<0.003 test for trend P<0.003 overall | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|--------------------|--------------------------------|------------------------------|------------------------------------|------------------------------|-------------------|
| | | | | | | difference between | |
| | | | | | | treatment groups | |
| | | | | | | Proportion of children | |
| | | | | | | who responded to | |
| | | | | | | treatment in the second week | |
| | | | | | | Group 1 (n=26): 58% | |
| | | | | | | (with no faecal | |
| | | | | | | incontinence 31%) | |
| | | | | | | Group 2 (n=27): 48% | |
| | | | | | | (with no faecal | |
| | | | | | | incontinence 26%) | |
| | | | | | | Group 3 (n=26): 62% | |
| | | | | | | (with no faecal | |
| | | | | | | incontinence 31%) | |
| | | | | | | Placebo (n=24): 29% | |
| | | | | | | (with no faecal | |
| | | | | | | incontinence 8%) | |
| | | | | | | P<0.27 group 3 vs. | |
| | | | | | | placebo | |
| | | | | | | Incidence and | |
| | | | | | | severity of adverse | |
| | | | | | | <u>effects</u> | |
| | | | | | | Group 1 (n=26): 9 (34.6%) | |
| | | | | | | | |
| | | | | | | Group 2 (n=27): 16 | |
| | | | | | | (59.3%) | |
| | | | | | | Group 3 (n=26): 17 | |
| | | | | | | (65.4%) | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | Placebo (n=24): 14 (58.3%) | |
| | | | | | | NS difference amongst groups | |
| | | | | | | No differences in the type of non- | |
| | | | | | | gastrointestinal related events, most common was | |
| | | | | | | headache. Higher incidence of GI-related events in | |
| | | | | | | patients receiving PEG vs. placebo. As dose of PEG | |
| | | | | | | increased, it also increased incidence of flatulence, | |
| | | | | | | abdominal pain, nausea and diarrhoea. | |
| | | | | | | No electrolyte abnormalities or differences in | |
| | | | | | | laboratory values amongst groups | |
| | | | | | | Treatment Failures Group 1 (n=26): 6 (4 BM frequency criteria, | |
| | | | | | | 2 with stool impaction) | |
| | | | | | | Group 2 (n=27): 7(3 BM frequency criteria, 4 with stool | |

| Bibliographic Study Type 8 Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|--------------------|--------------------------------|------------------------------|------------------------------------|---|-------------------|
| Level | | 5 | | Wedsures | impaction) Group 3 (n=26): 7 (6 BM frequency criteria, 1 with stool impaction) Placebo (n=24): 14 (all related to BM frequency criteria) | |

Adverse Effects of medium- to long-term use of Laxatives in Children with Chronic Idiopathic Constipation

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|--|---|--|--|---|---|---|---|
| Erickson et al. Polyethylene glycol 3350 for | Study Type: Retrospective case series | | s 46 children 35 girls | Intervention: Polyethylene glycol 3350 | Measures Duration of treatment | Side effects: -Diarrhoea: 9/46 | Additional information from study: Diagnosis of constipation based on history of |
| constipation in children with dysfunctional elimination. 2003. Journal of Urology 170[4 Pt 2], 1518- | Evidence level: 3 Study aim: To review the | Children diagnosed with dysfunctional voiding and | mean age: 7.7 years (range 4.5 to 11.2 years) 11 boys mean age: 7.6 | without electrolytes (MiraLax) 17 gm (1 capful) mixed with 8 ounces of fluid of | Mean: 194.3 days (SD 133.5) Assessment points | children, all female age at start of PEG (mean ± SD, years): patients with diarrhoea (n=9): | Infrequent bowel movements (less than very other day) and/or hard, large or painful bowel movements. Most children also had confirmatory abdominal x-ray demonstrating accumulation of stool in the rectum and throughout the colon |
| 1520 | for the treatment of constipation in children with dysfunctional elimination | who received polyethylene glycol 3350 between January 2000 and July 2002 Exclusion criteria: | years (range 4.4 to 11.1 years) | Starting dose: 8 ounces of mixture each day with instructions to adjust the amount consumed by 1 to 2 ounces every 3 | Not clear Outcome Measures: side effects | 6.8 ± 1.1 patients without diarrhoea (n=37): 8.2 ± 1.8 p=0.04 duration of follow-up | 25 patients also underwent biofeedback, and 8 patients began anticholinergic medication during the course of PEG treatment Reviewer comments: Not clear how side effects measured in the first place |
| | and asses bladder function following treatment | Known neurological impairments | | days to achieve the goal of 1 to 2 soft bowel movements per day Final dose normalised to patient weight Average final | | (mean ± SD, days): patients with diarrhoea (n=9): 336 ± 153 patients without diarrhoea (n=37): 108 ± 11 | Not clear how the reviewing process was conducted Source of funding: not stated |
| | | | | dose: 0.63 gm/kg (reported in abstract) 0.59 gm/kg (reported in text) Comparison: | | p=0.0028 1 child stopped taking PEG because of side effects | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|-------------|-------------------|
| | | | | None | | | |
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| Loening-Bauck et al. Rudy Type: et al. Rudy Type: (ase series glycol 3350 without electrolytes onstipation in infants and todilers. 2004. Journal of foundtional constipation in infants and todilers. 2004. Journal of gastroenterology and Nutrition and SIGS 5. Safe-539 | Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|--|--|---|---|--|---|---|--|
| | et al. Polyethylene glycol 3350 without electrolytes for the treatment of functional constipation in infants and toddlers. 2004. Journal of Pediatric Gastroenterolog y and Nutrition 39[5], 536-539 | Study Type: Retrospective case series Evidence level: 3 Study aim: to evaluate the safety and efficacy of PEG 3350 without electrolytes for the treatment of constipation in children < 2 years of age | Inclusion criteria: Children with constipation <2 years of age at start of PEG therapy Exclusion criteria: Hirschsprung's disease, chronic intestinal pseudo-obstruction, previous surgery of colon/anus, disease states that place limitations on the act of defecation such as hypotonia, cerebral palsy and severe mental | 75 children 36 boys mean age 17 months (range 1 to 21 months) | PEG 3350 without electrolytes (MiraLax) Starting average dose 1g/kg body weight/day Parents asked to adjust dose to yield 1 to 2 soft painless stools/day Comparison: | Duration of treatment (months, mean ± SD) -short term: 2.3 ± 1.3 (range: 1 to 4) -long term: 10.6 ± 8.1 (range 6 to 37) Assessment points -short term: ≤ 4 months (mean 2 months) -long term: ≥ 6 months (mean 11 months) Outcome Measures: | a. ≤ 4 months (n=71) 5 children (7%): runny stools (Dose of PEG (g/kg body weight/day): Range 0.4 to 2.3 Mean 1.1 ± 1.2 Median (0.82) b. ≥ 6 months (n=47) 1 child (2%): watery stools (he was only brought by his mother for a 6-month follow-up). The diarrhoea disappeared after lowering the dose of PEG. (Dose of PEG (g/kg body weight/day): Range 0.3 to 2.1 Mean 0.8 ± 0.4 Median (0.67) Parents did not report increased flatus, abdominal distension, vomiting or new onset abdominal pain. None stopped PEG | Constipation defined according to NASPGHAN criteria Reviewer comments: Authors reviewed charts from their own clinics. Not clear how the reviewing process was conducted Not completely clear how side effects were measured in the first place, it seems that parents were asked about the at the time of consultation Source of funding: not stated |

| Bibliographic Study Evido | nce Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|--------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | Complete blood counts (in 24 children), electrolytes (in 9 children), renal functions (in 8 children) and liver functions (in 8 children) occasionally done in children on long-term PEG treatment, and all were within normal limits. | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|---|--|--|---|---|--|---|
| | Evidence Level Study Type: Retrospective case series Evidence level: 3 Study aim: to determine | | | Intervention: PEG 3350 administered orally, mixed in a ratio of 17 g to 240 mL of fluid, as recommended by | Outcome Measures Duration of treatment Mean 6.2 ± 5 months (range, 3 weeks to 21 months) Assessment points at initial visit | Side effects: Total: 5 (17.9%) of patients 1 (3.6%) infant experienced increased passage of gas per rectum 4 (14.3%) infants | Additional information from study: Diagnostic criteria for functional constipation in infants and preschool children adapted from Rasquin-Weber and included: 2 weeks of hard stools (the majority of stools), or firm stools 2 or fewer times a week in the absence of structural, endocrine, or metabolic disease No patient placed on a clean-out protocol using any other drug |
| 39[2], 197-199 | optimal dose of polyethylene glycol powder for treatment of | Exclusion criteria: organic aetiology for constipation: Hirschsprung's disease, anorectal malformation, bowel obstruction, or systemic | gender not reported <u>Country:</u> USA | the sole diet. After initial dose, families asked to titrate the dose to obtain at least one nonformed bowel movement daily. Change in dose permitted within 24 hours, if necessary Mean initial | | resolved after dose adjustment | Duration of therapy and side effects retrieved from the patient's chart. Information not available in the chart was obtained by telephone interview. Only 1 family needed to be contacted by telephone Reviewer comments: Authors reviewed charts from their own clinics. Not clear how the reviewing process was conducted |
| | | illness (hypothyroidis m, cystic fibrosis, or lead poisoning associated with constipation. Taking medication that could potentially | | Dose: 0.88 g/kg/day (range, 0.26–2.14 g/kg/day) Mean effective maintenance dose: 0.78 g/kg/day (range, 0.26–1.26 g/kg/day) Comparison: | | | Source of funding: not stated |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|-------------|-------------------|
| | | change the | | none | | | |
| | | frequency or | | | | | |
| | | consistency of bowel | | | | | |
| | | movements | | | | | |
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| Pasharkar et al. Lung-term (Propective cohort (Propective cohort efficacy of polyethylene glycol 3350 for the treatment of chronic constipation in children with and without encopresis. 2030. Slinical Pediatrics 42[9], 815-819 **Refice (Propective cohort (Propective cohort (Propection)) (Propert efficacy of chronic constipation in children with constipation and children |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|-------------------------|---|
| | Level | 00 1 11 1 | <u>S</u> | 1 | Measures | | |
| Pashankar et al. | | 83 children | 83 children | Intervention: | Duration of | Clinical adverse | Additional information from study: |
| Safety of | Prospective | | | PEG 3350 without | | effects Minor and | Diagnosis of chronic constipation based |
| polyethylene | case series | | Male/female: | electrolytes | | acceptable over | on symptoms of at least 3 months' |
| glycol 3350 for | | | 48/35 | (MiraLax) | | mean duration of | duration, including at least 2 of the |
| | <u>Evidence</u> | Children > | 7.4 | 1 ''' 1 1 0 0 | | therapy | following: hard stools, painful defecation, |
| chronic | <u>level:</u> | | Mean age 7.4 | Initial dose: 0.8 | months | | encopresis, or fewer than 3 bowel |
| constipation in | 3 | | years (range | g/kg per day | | 8 patients (10%): | movements per week |
| children. 2003. | | | 2.0 to 16.9 | According to | | frequent watery stools | |
| Archives of | Study aim: | | years) | manufacturer's | | sometime during | All other laxative treatments stopped |
| Pediatrics and | to assess the | who were | | directions, parents | | therapy. Diarrhoea | before starting PEG |
| | | | Country: | instructed to | <u>Outcome</u> | disappeared with | |
| Medicine | and | | USA | dissolve 17 g of | Measures: | reduction of dose | Parents interviewed using structured |
| | | >3 months | | PEG powder in | | | questionnaire and asked about dose of |
| | profile of long- | | | | Adverse effects: | | PEG given, medication compliance, any |
| | | Exclusion | | or other beverage | | bloating or flatulence | possible adverse effects of PEG, and |
| | 3350 | <u>criteria</u> : | | and to give | -clinical | | particularly about excessively loose or |
| | treatment in a | | | prepared solution | | 2 children (2%): | frequent stools, abdominal pain, |
| | large cohort of | | | in 2 divided | -laboratory | abdominal pain | flatulence, bloating, and nausea. |
| | children and | s disease, | | doses. Families | | | Parents asked about overall |
| | also | anorectal | | allowed choice of | | 1 patient each (1%): | improvement in bowel movement |
| | paediatric | malformations | | beverage to suit | | thirst, fatigue, and | pattern regarding stool frequency and |
| | patient | , or any | | child's preference. | | nausea after | consistency with PEG therapy. |
| | | systemic | | Parents asked to | | receiving PEG | Following interview and physical |
| | 3 | illness | | adjust dose of | | solution on | examination, 4 mL of blood obtained for |
| | PEG therapy | potentially | | PEG solution as | | an empty stomach | measurement of different parameters |
| | | leading to | | required to yield 2 | | | |
| | | constipation | | soft painless | | None of the patients | Results of blood tests considered |
| | | | | stools per day. | | stopped treatment | abnormal if outside (even by 1 point) the |
| | | | | Over time, | | due to adverse | age- and sex appropriate reference |
| | | | | parents instructed | | effects and all were to | range established in authors' hospital. If |
| | | | | to gradually | | continue PEG | results abnormal, blood tests repeated |
| | | | | decrease | | therapy. | within 8 weeks while patient continued |
| | | | | dose of PEG if | | | to receive therapy |
| | | | | symptoms of | | General physical | |
| | | | | constipation and | | examination findings | |
| | | | | encopresis | | revealed no new | Source of funding: |
| | | | | showed | | significant | Study financially assisted by Braintree |
| | | | | improvement | | abnormalities | Laboratories |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | Comparison: | | compared with the pre-treatment | |
| | | | | None | | Laboratory evaluation results: | |
| | | | | | | Haemoglobin, haematocrit, serum | |
| | | | | | | electrolytes, blood urea nitrogen, serum | |
| | | | | | | creatinine, serum albumin, and | |
| | | | | | | osmolality, normal in all patients (10 patients did not have | |
| | | | | | | serum osmolality measured) | |
| | | | | | | 9 patients (11%) had slightly elevated ALT | |
| | | | | | | level (<1.5 times the upper limit of normal; | |
| | | | | | | range, 31 to 45 U/L). 8 of these patients | |
| | | | | | | had ALT levels remeasured within 8 weeks, 7 of whom still | |
| | | | | | | receiving PEG therapy. 7 of these 8 | |
| | | | | | | patients had values in the reference range, 1 had slightly | |
| | | | | | | elevated ALT level (<1.2 times normal; | |
| | | | | | | 28 U/L). 3 patients (4%) had | |
| | | | | | | an elevated aspartate aminotransferase | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|---|-------------------|
| | | | | | | level (<1.5 times normal; range, 42-52 U/L), and all had normal values when remeasured while still receiving PEG therapy Dose and duration of PEG therapy not significantly different in patients with abnormal values compared with those with laboratory values in the reference range | |
| | | | | | | | |

| Clark et al. Serum beta-carotene, Prospective case series retinol, and alpha- tocopherol levels during mineral oil therapy for constipation. 1987. American Journal of Diseases of Children receiving 141[11], 1210-Diseases of 1212 Clark et al. Setudy Type: Prospective case series retinol, and alpha- tocopherol levels during mineral oil throughout the early phase of treatment of treatment with mineral oil throughout the early phase of treatment tocopherol levels during mineral oil throughout the early phase of treatment tocopherol levels during mineral oil throughout the early phase of treatment tocopherol levels during alpha- tocopherol level level (isimpaction (micrograms/dL) (so. 55.7 ± 26.0) (micrograms/dL) (so. 55.7 ± 26.0) (micrograms | Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|--|---|---|---|--|---|---|--|--|
| Month 1: 4.0 ± 1.4 Month 2: 2.9 ± 1.2 Month 3: 2.1 ± 0.5 Month 4: 1.4 ± 0.4 Comparison: none Retinol: NS as compared to baseline -Month 3 (n=10): Serum beta-carotene: Baseline: 1.1 ± 0.6 (60.4 ± 30.0) Treatment: 0.6 ± 0.2 (34.7 ± 12.3) | Information Clark et al. Serum beta- carotene, retinol, and alpha- tocopherol levels during mineral oil therapy for constipation. 1987. American Journal of Diseases of Children 141[11], 1210- 1212 | Evidence Level Study Type: Prospective case series Evidence level: 3 Study aim: to prospectively monitor children receiving large doses of mineral oil throughout the early phase of | Patients 25 children Inclusion criteria: Children with encopresis, over 1 year old with no previous treatment with mineral oil Exclusion criteria: | Characteristic s 25 children mean age: 7.83 years (range 1.75 to 14.27 years) gender not reported Country: | Intervention: Following initial disimpaction (not reported with what), mineral oil, 45 mL twice daily between meals Dose gradually decreased on monthly basis (usually 30 mL/mo) depending on patient's reported performance and results of serial rectal examinations -Mean ± SEM: Month 1: 4.0 ± 1.4 Month 2: 2.9 ± 1.2 Month 3: 2.1 ± 0.5 Month 4: 1.4 ± 0.4 Comparison: | Outcome Measures Duration of treatment 4 months Assessment points 1, 2, 3 and 4 months Outcome Measures: Serum beta-carotene level Retinol level Alfa tocopherol level | Serum levels (micromols/L (micrograms/dL) (mean ± SEM): -Month 1 (n=25): Serum beta-carotene: Baseline: 1.0 ± 0.5 (55.7 ± 26.0) Treatment: 0.7± 0.4 (35.9 ± 22.1) P<0.01 Retinol: NS as compared to baseline -Month 2 (n=17): Serum beta-carotene: Baseline: 1.1 ± 0.6 (59.5 ± 30.6) Treatment: 0.7 ± 0.5 (38.2 ± 28.4) P<0.05 Retinol: NS as compared to baseline -Month 3 (n=10): Serum beta-carotene: Baseline: 1.1 ± 0.6 (60.4 ± 30.0) Treatment: 0.6 ± 0.2 | Additional information from study: Vitamin supplementation not prescribed Normal serum values for authors' laboratory: -Serum beta-Carotene: >0.6 micromols/L (>30 micrograms/dL) -Retinol: 0.70 micromols/L (20 micrograms/dL) -Alfa tocopherol: >9 micromols/L (>0.4 micrograms/dL) Since number of patients returning for subsequent visits gradually decreased, basal levels were recalculated for each month of treatment using the remaining patients as their own controls |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|---|-------------------|
| | | | | | inousures - | Treatment: 2.22 ± 0.77 (63.5 ± 22.1) P<0.01 -Month 4 (n=5): Serum beta-carotene: NS as compared to baseline Retinol: NS as compared to baseline Serum alfa tocopherol levels remained relatively unchanged throughout study. No statistical significant difference between baseline levels and those obtained throughout the 4 months of therapy | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|-------------------------|---|
| Hardikar et al. | | 81 children | 77 children | Intervention: | Duration of | Mean numbers of | Additional information from study: |
| Macrogol 3350 | Prospective | | | Macrogol 3350 | treatment | sachets/day during | Chronic constipation defined as fewer |
| | case series | <u>Inclusion</u> | 44% boys | plus electrolytes | | | than 3 complete bowel movements per |
| for chronic | | | mean age: | (Movicol) | | 1.3 (6.9 g) | week over previous14 days in |
| constipation in | Evidence | | 4.9 ± 2.6 years | (/ | Assessment | - (3) | association with either straining or |
| children: a | level: | 24 months to | · , , | Each sachet | points | Adverse effects | passage of hard stools in at least a |
| single-centre, | 3 | 11 years with | | (6.563 g | Adverse effects | (n=78) | guarter of bowel movements |
| open-label | | | Country: | Macrogol) | | 72 children (92%) | |
| study. 2007. | Study aim: | | Australia | dissolved 62.5 mL | throughout the | reported a total of | If investigator considered it to be |
| Journal of | | for at least 6 | | of water | | 318 events | clinically necessary patients could be |
| Paediatrics and | the safety and | months, | | | samples for | | given another laxative provided they had |
| Child Health | efficacy of a | which was | | Number of | | 241 (76%) assessed | failed to respond to the maximum dose |
| 43[7-8], 527- | macrogol | either | | sachets first 5 | | as unrelated to study | for 3 days |
| 531 | 3350-based | untreated or | | | | treatment | , |
| | electrolyte | inadequately | | -Children aged 2 | days. Vital | | No other therapeutic interventions, |
| | containing | treated by | | to 6 years: | signs measured | 262 (82%): mild | including an increase in oral fluids or |
| | preparation in | laxatives | | Days 1 & 2: 1/day | | 302 (95%): resolved | dietary fibre were instituted |
| | the treatment | | | Days 3 & 4: 1 | 84 days | by end of study | - |
| | of chronic | Exclusion | | twice a day | - | | Any child who developed faecal |
| | constipation in | criteria: | | Day 5: 1 three | Outcome | 6 serious adverse | impaction (faecal loading) which |
| | children | children | | times/day | Measures: | events in 4 children: 4 | required treatment was withdrawn from |
| | | treated for | | | | affected | study and classified as treatment failure |
| | | faecal | | -Children aged 7 | | gastrointestinal | |
| | | impaction with | | to 11 years | | system. All assessed | 78 (96%) patients included in safety |
| | | bowel | | Day 1 & 2: | | by investigator as | analysis. |
| | | washouts | | 1 twice a day | | unrelated or unlikely | 65 (80%) patients completed study. 16 |
| | | during the | | | laboratory tests | to be related to study | patients withdrew prematurely: 6 unable |
| | | previous 2 | | 2 twice a day | | medication and | or refused to take medication, 4 protocol |
| | | months, or | | | | resolved at end of | deviation, 3 poor compliance, 1 failed to |
| | | had a past | | Thereafter and | signs | study. 1 serious | return for final visit, 1 parent refused to |
| | | history of | | until end of study | | adverse event (faecal | give medication, 1 serious adverse |
| | | intestinal | | dosage titrated | | impaction) led to | effect |
| | | perforation/ob | | according to | | patient's premature | |
| | | struction, | | faecal form. This | | withdrawal from study | |
| | | Hirschsprung' | | dose increased by | | | 6 serious adverse events in 4 children: 4 |
| | | s disease, | | 1 sachet/day in | | - | affected gastrointestinal system, |
| | | paralytic | | the event of | | washout | remaining 2 not reported |
| | | ileum, toxic | | continued hard | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|--|--------------------------------|--|------------------------------------|---|--|
| | | megacolon, severe inflammation of the intestinal tract, urinary tract infection,, uncontrolled renal, hepatic or cardiac diseases, endocrine disorders, or any other severe unstable coexisting disease during he previous 30 days | | stools/no bowel movements, and decreased by 1 to 2 sachets/day in the event of loose stools or diarrhoea Comparison: None | | Changes in vital signs: No clinically significant changes as result of study medication | Not clear how clinical adverse effects were asked for Source of funding: Movicol sachets supplied by Norgine Ltd. Uxbridge, UK. Study supported by a research grant from Norgine Ltd. Uxbridge, UK and Norgine PTY, Sydney, Australia |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|---------------------------|---------------------------|---------------------------|---------------------|-----------------------------|--|
| | Level | | S | | Measures | | |
| Urganci et al. A | Study Type: | | 40 patients | Intervention: | Duration of | | Additional information from study: |
| comparative | RCT | | 22 male | Liquid paraffin | treatment: | (mean ± SD) | Diagnosis of constipation based on |
| study: the | - · · | | mean age 3.7 ± | | 8 weeks | (mL/kg/day) | symptoms of ay least 3 months duration |
| | <u>Evidence</u> | | 2.7 years | Comparison: | A | data mananta diin | including at least 2 of the following: hard |
| paraffin and | <u>level:</u> 1- | children 2 to | | Lactulose | Assessment | -data reported in | stool, painful defecation, rectal bleeding, |
| lactulose in | 1- | 12 years old referred for | | | | table, assumed that | encopresis and fewer |
| management of chronic | Study aim: | | Country: | Medication | | for the whole study period: | Open-label randomised study |
| functional | | | Turkey | administered | after initiation of | репоа. | Open-laber randomised study |
| constipation. | and compare | with evidence | Turkey | orally as a | treatment | Liquid paraffin (n=20): | Children also met with a nutritionist, |
| | efficacy, | of faecal | | suspension at 1 | liealinent | 1.88 ± 0.27 | were given instructions to increase daily |
| | safety and | impaction | | mL/kg, twice daily | Outcome | Lactulose (n=20): | fibre intake to amount of grams equal to |
| | optimal dose | Impaction | | for each drug. | | 2.08 ± 0.27 | their age plus 10, parent asked to have |
| | of liquid | Exclusion | | .o. odo d.dg. | | N.S | children sit on the toilet 4 times daily |
| | paraffin and | criteria: | | For determination | -optimal dose of | | after meals |
| | lactulose in | Hirschsprung' | | of best dose for | drugs | -data reported in text | |
| | | s disease, | | each child, | | | Stool frequency and stool consistency |
| | | hypothyroidis | | parents asked to | | treatment: | recorded by parents in daily diary forms. |
| | functional | m, mental | | increase or | rate | | Stool consistency scoring: 1, hard; 2, |
| | constipation | deficiency, | | decrease the | | | firm; 3, loose |
| | | chronic | | volume of each | | 1.72 ± 0.18 | |
| | | debilitating | | drug by 25% | | Lactulose (n=20): | No significant baseline differences |
| | | diseases, | | every 3 days as | | 1.82 ± 0.57 | between 2 groups |
| | | neurological | | required, to yield | | | |
| | | abnormalities, | | 2 firm-loose stools | | Compliance rate (%) | Patients considered compliant if ≥ 80% |
| | | previous | | per day. | | | of prescribed dose taken correctly. |
| | | surgery of | | Maximum dose | | | Patients instructed to take both empty |
| | | colon | | used throughout | | ' ' ' ' | and full containers to calculate amount |
| | | | | the study: 3 mL/kg | | 95 | of medication taken |
| | | | | per day for each | | Lactulose (n=20): | D |
| | | | | drug | | 90 N.S | Reviewer comments: |
| | | | | | | C.VI | Randomisation method not described |
| | | | | | | -end of 8 weeks: | No sample size calculation performed No clear definition of "evidence of faecal |
| | | | | | | | impaction" given |
| | | | | | | 190 | Apparently no children dropped out the |
| | | | | | | Lactulose (n=20): | study/were lost to follow-up |
| | | | | | | 60 | Study not controlled for potential |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|--|--|
| | Level | | S | | Measures | p=0.02 No patient stopped treatment because of adverse effects (adverse effects not reported). During first 4 weeks, taste aversion in 1 child on liquid paraffin and abdominal distension in 2 patients on lactulose influenced compliance. During last 4 weeks, poor symptom control in 5 patients, side-effects (abdominal distension and cramping) in 3 on lactulose, and watery stools in 2 on liquid paraffin influenced compliance | confounders Source of funding: not stated |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|-------------------------|--|
| | Level | | s | • | Measures | | |
| Dupont et al. | Study Type: | 96 children | 96 children | Intervention: | Duration of | Clinical tolerance | Additional information from study: |
| Double-blind | RCT | | 51 male | PEG 4000 | treatment: | (ITT population) | Constipation defined as <1 stool/day for |
| randomized | | <u>Inclusion</u> | | | 3 months | -6 adverse effects (all | >1 month in children 6 to 12 months old |
| evaluation of | <u>Evidence</u> | criteria: | Age (months) | -Starting dose: | | non serious): | and <3 stools/week for > 3 months in |
| clinical and | level: | ambulatory | (median, (25th- | 1 sachet (4g) and | <u>Assessment</u> | 5 diarrhoea (5 | children aged 13 months to 3 years |
| biological | 1+ | | 75th | 1 placebo to be | point (s): | episodes in 2 children | |
| tolerance of | | | percentiles) | taken at breakfast | | in both treatment | PEG 4000 and lactulose packaged in a |
| polyethylene | Study aim: | despite their | | | | groups) | double-blind and double-dummy design, |
| glycol 4000 | to assess the | usual dietary | -PEG 4000: | Comparison: | (D84) after | 1 anorexia (on | by means of coupled sachets, according |
| versus lactulose | safety of a | | 28 (19.5–33.7) | Lactulose | starting | lactulose) | to a randomisation list. Double dummy |
| in constipated | polyethylene | at least 1 | | | treatment | | design required because of the |
| children. 2005. | glycol (PEG) | month, aged | -Lactulose: | -Starting dose: | | -median (interquartile | difference of taste between the drugs. |
| Journal of | | | 25.8 (12.3–33) | 1 sachet (3.33g) | <u>Outcome</u> | range) duration of | Numbered boxes provided to |
| Pediatric | without | years | | and 1 placebo to | Measures: | either new onset or | investigators at each site in equal |
| Gastroenterolog | additional | | Country: France | | | worsened flatulence | numbers. Investigators randomly |
| y and Nutrition | salts in | <u>Exclusion</u> | | breakfast | -Biological | (days): | allocated either PEG 4000 or lactulose |
| 41[5], 625-633 | paediatric | criteria: | | | tolerance: | | to the children for a 3-month period, with |
| | patients | history of | | | | PEG 4000: 3 (1 to | the same strategy for dose adaptation |
| | | intractable | | For both drugs, | | 4.5) | |
| | | faecaloma, | | dose could be | electrolytes | Lactulose: 5 (3 to | 3 children not included because of a |
| | | Hirschsprung' | | doubled if | | 19.5) | baseline laboratory value ONR (out of |
| | | s disease, | | ineffective in | | P=0.005 | normal range) before the amendment |
| | | neurologic, | | children aged 13 | vitamin A | | was applied. 2 children in PEG 4000 |
| | | endocrine or | | months to 3 years | vitamin D | -median (interquartile | group dropped out before any study |
| | | metabolic | | If maximum | folates | range) duration of | drug intake, so the intention to treat |
| | | disorders, | | authorised dose | | either new onset or | (ITT) population included 51 children (10 |
| | | allergic | | unsuccessful, one | | worsened vomiting | babies and 41 toddlers) in the PEG |
| | | disease or | | micro-enema of | tolerance: | episodes (days): | 4000 group and 45 (12 babies and 33 |
| | | allergies | | glycerol per day | | | toddlers) in the lactulose group. 76 of |
| | | | | | , , | PEG 4000: 1 (1 to 2) | these children included in the per |
| | | | | | body weight | Lactulose: 2 (1 to 6) | protocol analysis and 20 excluded by |
| | | | | maximum of 3 | adverse effects | P<0.05 | the independent scientific committee for |
| | | | | consecutive days. | | | at least 1 major deviation, 11 in the PEG |
| | | | | If child not | | -anal irritation: 5% (2 | 4000 group and 9 in the lactulose group. |
| | | | | produced stools | | out of 40 children, | Reasons for exclusion were no |
| | | | | after treatment 2 | | both on lactulose) | laboratory test at D84, 1 or more one |
| | | | | enemas could be | | d:#fa.sa.s | missing laboratory results at D84, |
| | | | | administered at a | | -no difference | delayed laboratory test at D84 (n = 12), |

| Bibliographic Study Type & Number of Information Evidence Level Patients Study Type & Patient Characteristic Comparison | Follow-up & Effect Size Reviewer Comments Outcome Measures |
|--|---|
| 48-h interval. The procedure was only allowed two during the study of the child produce liquid stools for more than 1 days of the child produce liquid stools for more than 2 days of the child and or more than 2 days of the child and or more than 3 stools/day depending on age, dose could be decreased by the child and the ch | and lactulose groups with regards to other digestive tolerance outcomes -Body height and body weight unaffected during the 3-month treatment for both boys and girls -Biological tolerance (ITT population): No significant drug (n = 2), personal reasons (n = 5) and unauthorized concomitant treatment (n = 1). There were no clinically relevant differences between the 2 treatment groups at baseline for clinical or biologic parameters. Stool frequency, abdominal pain, vomiting, and nausea recorded by parents on Self-Diary Evaluation Booklet Reviewer comments: Methods of randomisation and allocation concealment not clearly described |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|---|-------------------|
| | | | | | | 1 (1 to 1.3) PEG 1.1 (0.9 to 1.5) lactulose P = 0.58 | |
| | | | | | | Treatment stopped in 1 child because of lack of efficacy (lactulose group) | |
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| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|--|---|--|--|--|---|---|
| Perkin. Constipation in childhood: a controlled comparison between lactulose and standardized senna. 1977. Current Medical Research and Opinion 4[8], 540-543 | Study Type: RCT (crossover) Evidence level: 1- Study aim: to compare effectiveness and side effects between a standardised senna syrup and lactulose in the treatment of childhood constipation | Inclusion criteria: children aged | 21 children (age and gender not reported) Country: UK | Intervention: Senna syrup 10 to 20 ml daily for 1 week Comparison: Lactulose 10 to 15 ml daily for 1 weeks Each preparation given throughout the appropriate treatment week in a daily dose varied according to the age of the patient 1 intermediate week with not treatment | week no treatment in between Assessment point (s): immediately after treatment completed Outcome Measures: | 4 (3 colic, 1 colic + distension) c- lactulose week 1 (colic) p<0.001 (a vs. c) | Additional information from study: Patients given either treatment according to a code-list of random numbers, placed in a series of sealed envelopes, one of which was opened each time a child entered the trial 1 dropout: 1 patient on senna at the beginning of study failed to attend at the end of 1 st week No written or oral indication of any medical preference for other preparation given and patients presented with single bottle of one or other of the preparations according to the coded instruction at start of trial. On 3 rd week a bottle of alternative preparation was given Outcomes recorded by parents in written diaries 4-point scale of stool consistency: loose, normal, hard, none Reviewer comments: Very small sample size, no sample size calculation Inadequate method of allocation concealment Patients' baseline characteristics not reported Study probably non blinded Results not controlled for confounders Very short treatment period According to authors the number of stools passed each day was recorded, but is not reported |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|-------------|-------------------------------|
| | | | | | | | Source of funding: not stated |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|------------------------|-------------------------|---|
| iniormation | Level | i dilonio | S | Companicon | Measures | | |
| Thomson et al. | Study Type: | 51 children | 51 children | Intervention: | Duration of | Mean effective dose | Additional information from study: |
| Polyethylene | RCT (cross | | 29 girls | PEG + E (6.9 g | treatment: | of PEG 3350 | Chronic constipation defined according |
| glycol 3350 plus | over, | | mean age 5.4 | powder/sachet) | 2 weeks each | (g/kg/day): | to Rome criteria as < 3 complete bowel |
| electrolytes for | multicentre) | criteria: | years (range: | | treatment | 0.6 (2 to 6-year-old) | movements/week, and at least 1 of the |
| chronic | | chronic | 24 months to 11 | Comparison: | | 0.7 (7 to 11-year-old) | following: pain on defecation on at least |
| constipation in | <u>Evidence</u> | constipation | years) | Placebo (6.9 g | separated by a | | 25% of days; at least 25% of bowel |
| children: a | level: | for at least 3 | | powder/sachet) | | Adverse events: | movements with straining, and at least |
| double blind, | 1+ | months | Country: UK | | washout | | 25% of bowel movements with hard or |
| placebo | | | | | | PEG+E (31/49, 63%) | lumpy stools |
| controlled, | Study aim: | <u>Exclusion</u> | | Washout period in | | Placebo (28/49, 57%) | |
| crossover | to assess the | criteria: | | between: 2 weeks | point (s): | during periods I and | Random sequence group computer |
| study.[erratum | efficacy and | current or | | | immediately | III. None serious, | generated before start of recruitment |
| | safety of | previous | | Dosing regime for | | most judged by | using block size of 4 patients and study |
| | polyethylene | faecal | | both PEG + E and | | investigator to be | medication labelled accordingly. |
| | glycol 3350 | impaction | | | | moderate or mild in | Random blocks (with numbers stored in |
| | plus | decided by | | sachets/day): | including | severity | sealed code-break envelopes) sent to |
| | electrolytes | either | | | washout | | investigator sites as required. As |
| Childhood | | physical | | -children aged 2 | | 20 children (41%) on | children enrolled, sites allocated |
| 92[11], 996- | | examination | | to 6 years | | PEG+E: 41 events | treatment supplies sequentially, started |
| 1000 | of chronic | or abdominal | | days 1-2: 1 | | 22 children (45%) on | with lowest possible number. Both the |
| | constipation in | • . | | days 3-4: 2 (taken | | placebo: 45 events, | children (and their parents/guardians) |
| | children | previous | | | | | |
| | | intestinal | | days 5-6: 3 (2 | | | blinded to allocation schedule |
| | | perforation/ob | | morning, 1 | | related to the study | |
| | | struction, | | evening) | | treatment. Most | A sample size of 50 children was |
| | | paralytic ileus, | | days 7-8: 4 (2 | | gastro-intestinal | planned to achieve 40 evaluable |
| | | Hirschsprung' | | morning, 2 | | disorders (particularly | children, giving 90% power to detect a |
| | | s disease, | | evening) | | abdominal pain), | true treatment difference of 0.3 bowel |
| | | severe | | | | PEG+E (39%, 39 | movements/week using a two-tailed |
| | | inflammatory | | -children aged 7 | | events); placebo | significance test at the 5% level. As |
| | | conditions of | | to 11 years | | (45%, 41 events). 1 | dropout rate was higher than originally |
| | | the intestinal | | days 1-2: 2 (taken | | child in | estimated, recruitment target was |
| | | tract, severe | | together) | | placebo/PEG+E | increased to 60 children |
| | | gastroesopha | | days 3-4: 2 (taken | | group withdrawn at | |
| | | geal reflux, | | together) | | week 3 because of | At baseline, clinically significant |
| | | diabetes, | | days 5-6: 5 (2 | | abdominal pain, | abnormalities |
| | | receiving | | morning, 3 | | assessed by | on physical examination (mainly |
| | | doses of | | evening) | | investigator as being | associated with faecal loading but not |

| Information Ev | dy Type & Number of vidence Patients Level | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|----------------|---|--------------------------------|---|------------------------------------|---|--|
| | stimulant laxatives considered by local observers to be at higher end of their own doses spectrum | | days 7-8: 6 (3 morning, 3 evening) For both groups if diarrhoea, dose was decreased by 2 sachets or miss a day. If loose stools dose decreased by 1 sachet | | placebo at the time of withdrawal. New clinically significant abnormalities on physical examination (mainly associated with faecal loading): 13 children (8/27 in the PEG+E/placebo group, 5/24 in the placebo/PEG+E group). When analysed for what these children were taking for the 2 weeks before the physical examination, 23 out of the 24 reports (95.8%) occurred when child taking placebo. Only 1 | impaction) recorded for 8 children (5/27 in the PEG+E/placebo group, 3/24 in the placebo/PEG+E group). Before randomisation, 47 children taking other laxatives (most frequently lactulose) 13/51 children (7/27 in the PEG+E/placebo group, 6/24 in the placebo/PEG+E group) recorded at least 1 deviation from the study protocol (1 child recorded 2 protocol deviations). Main reason for deviation was noncompliance with study medication (7/51 children), followed by failure to supply sufficient bowel movement data (4/51 children), and taking concomitant nonstudy laxative medication after randomisation (3/51 children) Safety monitored by adverse events recording, physical examination findings, and weight changes Reviewer comments: Blinding procedures not clearly described Unclear whether outcomes assessors were also blinded to treatment allocation Study not controlled for potential confounders Source of funding: Norgine Ltd. One of the authors was an employee of Norgine Ltd. At the time the study was written. The others declared that they had nothing to declare |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|------------------------|------------------------|---|
| | Level | | S | | Measures | | |
| Farahmand. A | Study Type: | 247 children | 247 children | General: | Duration of | Optimal dose of drug | Additional information from study: |
| randomised trial | RCT | | | 1 or 2 enemas | treatment: | -Final effective dose | Diagnosis of chronic functional |
| of liquid paraffin | | <u>Inclusion</u> | 127 male | , | 8 weeks | (mean, ml/kg/day): | constipation based on having at least 2 |
| versus lactulose | | <u>criteria:</u> | | clear any rectal | | Liquid paraffin | of the following symptoms for the last 3 |
| in the treatment | | | aged 2 to 12 | impaction (30 | <u>Assessment</u> | (n=127) | months: <3 bowel movements/week, |
| of chronic | 1- | | years old (mean | | | 1.72 ± 0.13 | faecal soiling >once/week, large |
| functional | | constipation | 4.1± 2.1 years) | paraffin oil) | | Lactulose (n=120) | amounts of stool every 7 to 30 days and |
| constipation in | Study aim: | | | | | 2.08 ± 0.21 | palpable abdominal or faecal mass on |
| | to compare | | Country: Iran | Intervention: | started | p<0.001 | physical examination |
| Acta Medica | the clinical, | criteria: | | Liquid paraffin | | | |
| Iranica 45[3], | efficacy and | organic | | orally, 1 to 2 | <u>Outcome</u> | | Apart from laxative treatment, parents |
| 183-188Iran, | safety of liquid | | | ml/kg, twice daily | Measures: | | given instructions to increase their daily |
| Islamic | paraffin and | defecation | | for 8 weeks | | | fibre intake to an amount of grams equal |
| Republic of. | lactulose in | disorders | | | • | | to their age plus 10. Toilet training after |
| | | including | | Comparison: | drug | | each meal advised to enhance |
| | | Hirschsprung' | | Lactulose orally, 1 | | bar chart, outcomes | compliance |
| | childhood | s' disease, | | to 2 ml/kg, twice | -side effects | not reported in text): | |
| | constipation | spina bifida | | daily for 8 weeks | | Lactulose (n=120) | Treatment success defined as 3 or more |
| | | occulta, | | | | | bowel movements/week and encopresis |
| | | hypothyroidis | | | | Abdominal pain: 10 | episodes < 2/week |
| | | m, cystic | | | | Bad palatability: 15 | |
| | | fibrosis, | | | | | No significant baseline differences |
| | | neurological | | For determination | | Bloating: 10 | between the 2 treatment groups |
| | | abnormalities, | | of best dose for | | Diarrhoea: 10 | regarding: age, sex, duration of |
| | | intestinal | | child, parents | | Anal oil leakage: 20 | constipation, defection frequency, |
| | | pseudo | | asked to increase | | Flatulence: 10 | number of patients with history of |
| | | obstruction | | the volume of | | Nausea: 10 | encopresis, large amount of stool, faecal |
| | | | | each drug by 25% | | Hard stool: 20 | impaction in rectum, rectal bleeding, lost |
| | | | | every 3 days as | | Vomiting: 0 | to follow-up after 8 weeks, bad |
| | | | | required to yield 1 | | | palatability of study medication |
| | | | | or 2, firm-loose | | Liquid paraffin | |
| | | | | stools | | (n=127) | Parents received chart to record side effects |
| | | | | | | Abdominal pain: 50 | |
| | | | | | | Bad palatability: 40 | Reviewer comments: |
| | | | | | | Pain at defecation: 50 | Method of randomisation and allocation |
| | | | | | | Bloating: 20 | concealment not described |
| | | | | | | Diarrhoea: 30 | Non blinded study |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|---|--|
| | Level | | 5 | | | Anal oil leakage: 40 Flatulence: 20 Nausea: 5 Hard stool: 6 Vomiting: 0 | No sample calculation performed No withdrawals/dropouts reported Results not controlled for confounders Source of funding: not stated, but authors reported "no conflicts of interests" |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|-----------------------------|---------------------------|------------------------|-------------------------|--|
| Loening- | Level Study Type: | 49 children | s -Miralax group: | Intervention: | Measures Duration of | Medication dosage | Additional information from study: |
| Baucke. | Prospective | | 28 children | MiraLax | treatment: | (Mean doses and | Initial dose of Miralax 0.5 g/kg daily |
| | cohort | | 20 boys | 17 dissolved | 12 months | range for children | suggested for children whose rectums |
| glycol without | COHOIT | | Mean age ± | in 240 mL of a | 12 1110111113 | who were doing well | were loaded with stool but who had no |
| | <u>Evidence</u> | | SD: | beverage such as | Assassment | or improved) (PEG, | fecal abdominal masses at the initial |
| children with | level: | | 8.7 ± 3.6 years | juice or Kool-Aid | point (s): | g/kg; MOM, mL/kg) | physical examination and no history of |
| | 2 + | | Range 4.1 to | initial dose: 0.5 to | 1, 3, 6, and 12 | g/kg, MOM, IIIL/kg) | long intervals between huge bowel |
| and encopresis. | 2 1 | functional | 17.5 years | 1 g/kg/daily | months after | 1 month | movements. Those with |
| 2002. Journal of | Study aim: | constipation | 17.5 years | i g/kg/dally | initiating | PEG: | palpable abdominal fecal masses or |
| | to determine | and | -MOM group: | Comparison: | treatment | 0.6 ± 0.2 (0.3 to 1.1) | history of infrequent huge bowel |
| | | | 21 children | MOM | licalificiti | MOM: | movements started on 1 g/kg daily |
| | acceptability, | | 17 boys | Initial dose 1 to | Outcome | 1.4 ± 0.6 (0.6 to 2.6) | Intovernents started on 1 g/kg daily |
| | and treatment | | Mean ± SD: 7.3 | | Measures: | 1.4 ± 0.0 (0.0 to 2.0) | Milk of Magnesia given if family could |
| | dosage of | | ± 3.0 years | 2.0 IIIL/Ng | <u>ivicasarcs.</u> | 3 months | afford only the use of a cheaper laxative |
| | MiraLax | delay/difficulty | | | -medication | PEG: | or if child had previously received MOM |
| Glates. | | | 13.9 years | | dosage | 0.6 ± 0.3 (0.3 to 1.4) | without refusal. For these children, MOM |
| | glycol 3350 | and | 10.5 years | | dosage | MOM: | reintroduced or adjusted to an adequate |
| | | | Country: | | -clinically | 1.2 ± 0.5 (0.6 to 2.4) | dosage. Parents told how to improve the |
| | | (≥1/week) for | | | significant side | 1.2 ± 0.0 (0.0 to 2.4) | taste by mixing the child's preferred |
| | during a 12- | more than 1 | 00/1 | | effects | 12 months | flavoring with plain MOM. Initial daily |
| | month | vear | | Large laxative | enecis | PEG: | dosage of 1 mL/kg body weight |
| | treatment | you | | dosages divided | -compliance | 0.4 ± 0.1(0.1 to 0.7) | suggested for children with rectal fecal |
| | period in | Exclusion | | into 2 daily doses. | | MOM: | masses only at initial evaluation and if |
| | children with | criteria: | | Parents told to | With medication | only 2 children still | they had no history of infrequent large |
| | functional | Children <4 | | adjust the dose of | | required MOM. Their | bowel movements. Dosage of 2.5 mL/kg |
| | constipation | years of age; | | medication by 30 | | dosages were 0.4 | prescribed for those with fecal |
| | and | children who | | mL for MiraLax | | and 1.6 mL/kg, both | abdominal masses at the initial |
| | encopresis | refused the | | and by 7.5 mL | | less than the initial | evaluation or history of huge, infrequent |
| | encopiesis | toilet for | | (one-half | | treatment dosage | bowel movements |
| | | stooling but | | tablespoon) for | | licatificht dosage | bower movements |
| | | who had no | | MOM every 3 | | mean doses for both | Regular stool sittings for 5 minutes after |
| | | constipation, | | days to a dosage | | treatments at 12 | each meal required for initial months. |
| | | Hirschsprung' | | that resulted in 1 | | months | Caon mean required for initial months. |
| | | s disease, | | to 2 soft bowel | | did not differ | Patients and parents provided with diary |
| | | chronic | | movements/day | | significantly between | sheets to record each outcome |
| | | intestinal | | and prevented | | children with or | measured |
| | | pseudo- | | soiling and | | without initial | |
| | | obstruction, or | | abdominal pain. | | palpable abdominal | Global assessment of whether child was |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|--------------------------------|------------------------|-----------------------------------|--|
| IIIIOIIIIatioii | Level | ratients | S | Companison | Measures | | |
| | | previous | | If child retained | | faecal masses. None | "doing well," "improved," or "not doing |
| | | surgery of the | | stools despite | | of the patients | well" was recorded. Doing well defined |
| | | colon/anus | | compliance with | | | as 3 or more bowel movements/week |
| | | | | assigned laxative, | | dosage of either | and 2 or fewer soiling episodes / month. |
| | | | | daily senna added to treatment | | medication over time | Improved defined as 3 or more bowel movements / week and a more than |
| | | | | | | 5 children received a | 75% decrease in soiling but not more |
| | | | | | | stimulant laxative in | than 1 soiling / week. Not doing well was |
| | | | | | | addition to PEG and | defined as fewer than 3 bowel |
| | | | | | | 1 child received a | movements / week, a less than 75% |
| | | | | | | stimulant laxative | decrease in soiling frequency, use of |
| | | | | | | in addition to MOM (P | senna, or refusal to take the assigned |
| | | | | | | > 0.2) | laxative. Recovered defined as 3 or |
| | | | | | | | more bowel movements / week and 2 or |
| | | | | | | Clinically significant | fewer soiling episodes / month while not |
| | | | | | | side effects | taking laxatives. |
| | | | | | | PEG: no significant | No significant baseline differences |
| | | | | | | clinical side effects. | between 2 groups |
| | | | | | | Some children had | |
| | | | | | | | Reviewer comments: |
| | | | | | | children in the PEG | No sample size calculation performed |
| | | | | | | group became dehydrated. Children | Outcomes for consistency of stools not |
| | | | | | | receiving PEG and | reported |
| | | | | | | their parents did not | reported |
| | | | | | | report increased | Not reporting on the clinically significant |
| | | | | | | flatus, abdominal | side effects (or lack of them) for MOM |
| | | | | | | distention, or new | Side chects (or lack of them) for Mow |
| | | | | | | onset of abdominal | Source of funding: |
| | | | | | | pain | Dr. Loening-Baucke recipient of grant |
| | | | | | | | support from Braintree Pharmaceuticals, |
| | | | | | | Compliance with | Braintree, MA, U.S.A., for continuing |
| | | | | | | medication: | studies on childhood constipation |
| | | | | | | -PEG: No children | |
| | | | | | | reported disliking the | |
| | | | | | | taste, no parents | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|--|-------------------|
| | | | | | | reported that child refused to take it in juice or Kool-Aid Parental noncompliance with administering the laxative and | |
| | | | | | | supervising toilet use: 14% children -MOM: 33% children refused to take it Parental noncompliance with administering the | |
| | | | | | | laxative and supervising toilet use: 4% children | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|------------------------|-------------------------|---------------------------|---------------------------|------------------------|------------------------|---|
| | Level | | S | | Measures | | |
| Loening-Baucke | Study Type: | 79 children | 79 children | General: | Duration of | Patient Acceptance | Additional information from study: |
| et al. A | RCT | | 65 boys | disimpacted with | treatment: | Several children | Functional constipation defined by |
| randomized, | | <u>Inclusion</u> | age range: 4 to | 1 or 2 phosphate | 12 months | complained about | duration of ≥ 8 weeks and ≥ 2 of the |
| prospective, | <u>Evidence</u> | criteria: | 16.2 years | enemas in the | | taste of PEG and | following: frequency of bowel |
| comparison | <u>level:</u> | age ≥ 4 years | (median 7.4; | clinic on the day | <u>Assessment</u> | MOM. | movements <3 stools/week, >1 episode |
| study of | 1- | and presence | mean 8.1 ± 3.0) | | point (s): | 2 children (5%) | of faecal incontinence/week, large stools |
| polyethylene | | of functional | | necessary and | 1, 3, 6 and 12 | continued to refuse | noted in rectum or felt during abdominal |
| glycol 3350 | Study aim: | constipation | Country: USA | started laxative | months after | PEG vs. 14 children | examination, passing of stools so large |
| without | to compare | with faecal | | therapy that | initiating | (35%) continued to | that they obstructed the toilet |
| | the efficacy, | incontinence | | evening | treatment | refuse MOM during | |
| milk of | safety and | | | | | the 12 months of the | Randomisation performed by children |
| magnesia for | patient | Exclusion | | Intervention: | <u>Outcome</u> | study | drawing a sealed envelope with and |
| children with | acceptance of | criteria: | | polyethylene | Measures: | (P < 0.001) | enclosed assignment |
| | polyethylene | stool toileting | | glycol (PEG) 3350 | | | |
| | glycol (PEG) | refusal, faecal | | without added | -safety profile | Treatment doses | Investigators, children and their parents |
| incontinence. | 3350 without | incontinence | | electrolytes 0.7 | | (mean ± SD): | aware of the study group assignment |
| | added | but no | | g/kg body weight | -patient's | DEC /a/lea book | It was a stime at a dith at 20 ambig ata ways |
| 118[2], 528-535 | | constipation, | | daily for 12 | acceptance and | ίο ο | It was estimated that 38 subjects were |
| | | previous refusal of one | | months | compliance | weight) | required in each group to be able to detect a difference in failure rates |
| | magnesia (MOM) over | of study | | capful of PEG (17 | | 1 month: 0.7 ± 0.2 | between the 2 groups of 30% in 12 |
| | 12 months | medications, | | g) mixed in 8 oz of | | 3 months: 0.6 ± 0.3 | months (40% vs. 10%), at the 0.05 |
| | 12 1110111115 | children who | | beverage (juice, | | additional senna at | significance level with 0.80 power. |
| | | came from far | | Kool-Aid, Crystal | | | Authors hypothesized that PEG would |
| | | away for a | | Light or water) | | Some point. S children | be as successful as MOM in treating |
| | | second | | making a solution | | -MOM (mL/kg body | chronic constipation and faecal |
| | | opinion, | | of ~2g/30 mL | | weight) | incontinence. Authors' previous study |
| | | Hirschsprung' | | or EgroomE | | Worght, | showed that 33% of children refused to |
| | | s disease, | | Comparison: | | 1 month: 1.2 ± 0.7 | take MOM during the first 12 months of |
| | | chronic | | milk of magnesia | | 3 months: 1.2 ± 0.8 | treatment. |
| | | intestinal | | (MOM) 2mL/kg | | additional senna at | |
| | | pseudobstruct | | body weight daily | | some point: 1 child | Children treated with minimal effective |
| | | ion, previous | | for 12 months | | | dosage of PEG or MOM, allowing for a |
| | | surgery | | | | mean doses similar in | daily stool and preventing abdominal |
| | | involving | | plain MOM could | | children who | pain and faecal incontinence. Parents |
| | | colon or anus | | be mixed into | | improved and who | instructed to aim for 1 or 2 stools of |
| | | | | apple sauce or | | did not improve for | milkshake consistency each day. |
| | | | | milkshakes, or | | both treatments | Parents asked to increase dosage if |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|--|------------------------------------|---|--|
| | | | | chocolate and other flavouring could be added Large doses of both medications could be divided into 2 doses | | safety profiles PEG: 1 child allergic No other significant clinical effects for either medication, apart from transient diarrhoea disappearing with dose reduction -Laboratory tests: PEG: 1 child with elevated platelets before and after treatment, 1 child with decreased sodium levels at 6 months, but normal at 12 months MOM: 1 child high platelet count, 1 low serum sodium level, elevated AST, 1 elevated ALT | stools too hard or not frequent enough and to decrease the dosage if stools watery or too numerous. Small changes, such as 2 oz of PEG or 0.5 tbsp of MOM every 3 days, were recommended. Regular stool sittings for 5 minutes after each meal required initially. Toilet sitting frequency reduced after children recognized urge to defecate and initiated toilet use themselves. No significant differences at baseline between the 2 groups regarding: age, sex, primary faecal incontinence, previous treatment with laxatives, history of retentive posturing, frequency of bowel movements, bowel movements obstructing the toilet, frequency of faecal incontinence, presence of abdominal pain, presence of abdominal faecal mass and presence of rectal faecal mass By 12 months a total of 27 dropouts/lost to follow-up monitoring, 2 (5%) had refused PEG, 1 child allergic to PEG, 2 children were receiving senna. These 7 children counted as not improved and not recovered. MOM: 2 Children lost to follow-up monitoring, 3 children had discontinued study participation, 14 children (35%) had refused to take MOM, and 1 child was receiving senna Efficacy analyses performed with |
| | | | | | | | intention to treat population, other |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|-------------|---|
| | | | | | | | outcomes calculated from available follow-up data |
| | | | | | | | Patients and parents questioned with respect to side effects during each visit |
| | | | | | | | Reviewer comments: Results not controlled for potential confounders High drop-out / lost to follow-up rate: 30.4% |
| | | | | | | | Source of funding: Braintree Laboratories (Braintree, MA) supported study with an unrestricted research grant. According to authors, the funding source had no involvement in the study design, collection, analysis, interpretation of data, writing of the report or decision to submit the article for publication |
| | | | | | | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|-----------------------|---|
| intormation | Level | - ationio | S | Companicon | Measures | | |
| Adler. Effective | Study Type: | 134 patients | 134 patients | Intervention: | Duration of | Mean dose at end of | Reviewer's' comments |
| Treatment of | Prospective | | 88 males | Movicol (macrogol | treatment: | observational period | It is difficult to asses the quality criteria |
| Constipation | Case series | <u>Inclusion</u> | age not clearly | 3350 with | Mean: 50 | | and to make comments on this study |
| and Encopresis | | <u>criteria:</u> | reported | electrolytes,13.8g | weeks (SD ±50 | Age 2 to 6: 0.42 | because we have only been able to |
| with Movicol | <u>Evidence</u> | children | | sachets) | weeks; range 1 | sachets | review the abstract. This abstract was |
| (Macrogol 3350 | <u>level:</u> | referred with | Country: | | to 211 weeks) | | included because it provides some |
| with | 3 | constipation | Sweden | -Mean starting | | Age 7 to 11: 0.49 | evidence on long-term treatment with |
| Electrolytes) in | | and/or | | dose: | <u>Assessment</u> | sachets | Movicol |
| Children and | | encopresis to | | Age 2 to 6: 0.58 | point (s): | | |
| Adolescents. | to assess the | The Queen | | sachets | unclear | -overall mean | |
| 2005. Gut | effectiveness | Silvia | | | | change: 0.553 to | Source of funding: |
| 54[Suppl VII], | | Children's | | Age 7 to 11: 0.51 | <u>Outcome</u> | 0.477 sachets/day | Not stated |
| A217 | | Hospital, | | sachets | Measures: | | |
| Adler, 2005 | 3350 with | Sweden | | | | Side-effects were | |
| | electrolytes), | | | | -final treatment | reported in 10 (7.5%) | |
| | | Exclusion | | Doses adjusted in | dose | patients and these | |
| | course of long | | | each patient to | | were generally mild | |
| | term | Not stated | | achieve symptom | -side effects | and transient | |
| | treatment in | | | relief with the | | | |
| | children with | | | minimally effective | | | |
| | constipation | | | dosage | | | |
| | | | | 0 | | | |
| | | | | Comparison: | | | |
| | | | | None | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |

Effectiveness of Diet and Lifestyle modifications in Children with Chronic Idiopathic Constipation

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|------------------------|---------------------------|---------------------------|------------------------|----------------------------|--|
| | Level | | S | | Measures | | |
| Bongers et al. | Study Type: | 38 children | 38 children | Intervention: | | Clinical efficacy | Additional information from study: |
| The clinical | Double-blind | | | Nutrilon Omneo | | after period 1 | Constipation defined as the presence of |
| | | <u>Inclusion</u> | 19 boys | (new formula, NF) | | Defecation frequency | at least 1 of the following symptoms: 1) |
| infant formula in | over) | criteria: | median age: 1.7 | | weeks each | (mean ± SD) | frequency of defecation < 3/week; 2) |
| term infants | | Otherwise | months | -Nutrients per 100 | | | painful defecation (crying); 3) abdominal |
| with | <u>Evidence</u> | healthy, term | | : | <u>Assessment</u> | SF (n = 15): 4.9 ± 2.5 | or rectal palpable mass |
| constipation: a | <u>level:</u> | infants with | | ml: | | NF (n = 20): 5.6 ± 2.8 | |
| double-blind, | 1+ | constipation, | Country: | | After period 1 | | Infants randomised by a computer |
| randomized | | between 3 to | The | Energy (kcal) 70 | and period 2 | Difference of means | program to either NF or SF in period 1 |
| cross-over trial. | Study aim: | 20 weeks of | Netherlands | | | (95% CI): | and crossed-over after 3 weeks to |
| 2007. Nutrition | | age, who | | Protein (g) 1.7 | | 0.7 (-0.8 to 2.3) | treatment period 2 |
| Journal 6, 8 | hypothesis | received at | | Casein - | | N.S | |
| | that Nutrilon | least 2 bottles | | Intact whey | No follow-up | | In order to mimic the taste of Nutrilon |
| | Omneo (new | of milk-based | | protein - | conducted after | Improvement of hard | Omneo, the whey-based control formula |
| | | formula per | | Whey protein | treatment | to soft stools (n) | was partly mixed with a formula based |
| | will have a | day | | hydrolysate 1.7 | finished | 0- (| on hydrolyzed whey protein (mixture of |
| | positive effect | | | | | SF (n = 15): 50% | 75% Nutrilon 1 and 25% Aptamil HA I). |
| | | Exclusion | | | <u>Outcome</u> | (5/10) | Formula cans were labelled with codes |
| | characteristics | | | (g) 3.3 | | NF (n = 20): 90% | to mask identity of the study feedings. |
| | | Hirschsprung' | | Palmitic acid 0.6 | | (9/10) | Neither the parents nor the physicians |
| | children | s disease, | | - at the sn-2 | Primary | DD (050(OI) | were aware of the composition of the |
| | | spinal or anal | | position (%) 41.0 | | RR (95% CI): | formula until the entire study was |
| | | anomalies, | | Linoleic acid 0.4 | | 1.8 (0.9 to 3.5) | completed |
| | | previous | | α-linolenic acid | | N.S | Drients start of the study somewhat sine |
| | | colonic | | 0.08 | 1) defecation | No pointul defending | Prior to start of the study, sample size, |
| | | surgery, metabolic, | | Carbabydratas (a) | frequency | No painful defecation | based on a cross-over design, was calculated to allow detection of a 30% |
| | | cerebral and | | Carbohydrates (g) 8.4 | > 3/WEEK | <u>(n)</u> | difference in improvement between NF |
| | | renal | | - | 2) normalization | SF (n = 15): 33% | and SF. Under the assumption of a |
| | | abnormalities, | | Maltodextrin 4.0 | of stool | (5/15) | significance level of 0.05 with |
| | | children who | | Starch 1.5 | | NF (n = 20): 35% | a power of 0.80, and 2-sided hypothesis |
| | | were treated | | 0.0.011 1.0 | Consistency | (7/20) | testing, a minimal sample size of 34 with |
| | | with laxatives | | Fibre (g) 0.8 | 3) no more | (1,20) | 17 children in each group was |
| | | at enrollment | | | | RR (95% CI): | determined |
| | | | | (90% GOS, 10% | | 1.0 (0.4–2.7) | |
| | | | | IcFOS) 0.8 | | N.S | Only 24 children (63%) completed the |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|------------------|---------------------------------------|---------------------|------------------------|---|
| Information | Level | Patients | Characteristic s | Comparison | Outcome Measures | | |
| | | | - | | Secondary | | cross-over study. In period 1, 3 SF |
| | | | | Minerals and | outcome: | Clinical efficacy | patients dropped out; 2 patients stopped |
| | | | | trace elements | | after cross-over | because of severe constipation; 1 |
| | | | | (mg) | -safety | (period 1 and 2) | patient switched to hypoallergenic |
| | | | | Calcium 53 | | Defecation frequency | feeding, because of suspected cow's |
| | | | | Phosphorus 29 | | (mean) | milk protein allergy. Parents of 1 patient |
| | | | | Sodium 23 | | | decided that they did not want to cross- |
| | | | | Potassium 82 | | SF (n =12): 5.9/week | over because she was free of symptoms |
| | | | | Chloride 44 | | NF (n =12): 5.5/week | and they started openly with NF instead. |
| | | | | Iron 0.5 | | | 3 patients dropped out after switching to |
| | | | | Zinc 0.5 | | Difference of means | NF; 2 patients stopped after less than 1 |
| | | | | | | (95% CI): | week because of recurrence of |
| | | | | Comparison: | | - 0.5 (-1.6 to 0.6) | constipation symptoms. 1 patient was |
| | | | | Standard formula | | N.S | lost to follow-up. 7 patients dropped out |
| | | | | (SF, mixture of | | _ , , | after switching to SF; 6 patients stopped |
| | | | | 75% Nutrilon I | | Frequency of soft | after 1 week because of recurrence of |
| | | | | and 25% Aptamil | | stools: | constipation symptoms. 1 patient was |
| | | | | HA I) | | | lost to follow-up |
| | | | | Energy (keel) 67 | | had soft stools when | Data analysis based on the group of 25 |
| | | | | Energy (kcal) 67 | | stools with SF, | Data analysis based on the group of 35 patients that completed period 1 and a |
| | | | | Protein (g) 1.5 | | | subgroup analysis of 24 patients who |
| | | | | Casein 0.5 | | with soft stools when | completed the cross-over |
| | | | | Intact whey | | receiving SF and no | Completed the cross-over |
| | | | | protein 0.6 | | | No significant differences in baseline |
| | | | | Whey protein | | with NF (p = 0.046) | characteristics between 2 groups |
| | | | | hydrolysate 0.4 | | (p = 0.0 10) | groupe |
| | | | | l l l l l l l l l l l l l l l l l l l | | Painful defecation | During both periods parents asked to |
| | | | | Fat (triglycerides) | | not significantly | daily record in a diary details on formula |
| | | | | (g) 3.5 3.3 | | different between the | intake, formula tolerance (vomiting, |
| | | | | Palmitic acid 0.6 | | periods | flatulence, colic, rash), passage of stools |
| | | | | - at the sn-2 | | on NF and SF | and stool |
| | | | | position (%) 11.5 | | | consistency compared to 4 validated |
| | | | | Linoleic acid 0.4 | | <u>Safety</u> | photographs of runny, mushy soft, |
| | | | | α-linolenic acid | | Throughout the study | formed soft and hard stools |
| | | | | 0.07 | | there were no serious | |
| | | | | | | adverse effects in | Reviewer comments: |
| | | | | Carbohydrates (g) | | either group. Both | Allocation concealment method not |

| T.3 Lactose 7.2 Maltodextrin - Starch - Fibre (g) - Oligosaccharides (90% GOS, 10% IcFOS) - Minerals and trace elements (mg) Calcium 53 Phosphorus 29 Sodium 22 Potassium 69 Chloride 42 Iron 0.5 Tinc 0.5 Phosphorus 29 Tinc 0.5 Tinc 0.5 | Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|--|---------------------------|-----------------------------|-----------------------|--------------------------------|--|------------------------------------|-------------|---|
| Feeding patterns not described | | | | | Lactose 7.2 Maltodextrin - Starch - Fibre (g) - Oligosaccharides (90% GOS, 10% IcFOS) - Minerals and trace elements (mg) Calcium 53 Phosphorus 29 Sodium 22 Potassium 69 Chloride 42 Iron 0.5 Zinc 0.5 | | | Study not controlled for potential confounders Source of funding: study supported by a grant of Nutricia Nederland BV, Zoetermeer, The |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|----------------------|---|
| | Level | | S | | Measures | | |
| Chao et al. | Study Type: | 93 children | 93 children | Intervention: | Duration of | Improved (number | Additional information from study: |
| Therapeutic | Open label | | 47 boys | Magnesium- | treatment | and % of children) | Study non-blinded, according to authors |
| effect of | RCT | <u>Inclusion</u> | mean age 3.8 ± | enriched infant | 2 months | -At 2 weeks: | this was not possible because all infants |
| Novalac-IT in | | criteria: | 1.7 months | formula, Novalac- | | Novalac-IT (n=47): | were included in 1 centre |
| infants with | <u>Evidence</u> | Children aged | | IT | <u>Assessment</u> | 31 (66) | |
| constipation. | <u>level:</u> | 2 to 6 months | Country: | | point (s): | | Randomisation performed applying an |
| 2007. Nutrition | 1- | referred to | Taiwan | Composition per | At 2 weeks, 1 | Strengthened formula | envelope drawing system |
| 23[6], 469-473 | | paediatric | | 100 mL: | month and 2 | (n=46): 23 (50) | |
| | Study aim: | gastroenterol | | | months | N.S | Assigned nurse educated the family to |
| | | ogy clinic at | | Energy (cal/100 | | | prepare the 20% strengthened formula |
| | commercialise | | | mL): 70.7 | Follow-up | -At 1 month: | (20% extra formula) (regular |
| | d formula, | centre with | | | period: | Novalac-IT (n=47): | concentration of the formula is 13%) |
| | | constipation ≥ | | Protein (g): 1.70 | No follow-up | 39 (83) | |
| | (Intestinal | 2 weeks, fed | | Whey/casein: | conducted after | | No significant differences in baseline |
| | | exclusively | | 60/40 | treatment | | characteristics (clinical or demographic) |
| | France) | with formula. | | | finished | (n=46): 23 (50) | between the 2 groups |
| | against a | Participation | | Fat (g): 3.54 | | P=0.002 | |
| | "strengthened | in trial | | | <u>Outcome</u> | | Intake of formula and clinical parameters |
| | regular | proposed | | Carbohydrates | Measures: | -At 2 months: | regarding constipation and weight and |
| | formula", the | before a | | (g): 8.06 | Remission / | Novalac-IT (n=47): | all relevant information recorded by |
| | traditional | more | | 100 % Lactose | improvement / | 42 (89) | family daily in a diary during the entire |
| | approach in | complete | | | failure | | intervention period |
| | infants with | diagnostic | | Major minerals | according to | Strengthened formula | |
| | digestive | workup for | | (mg) | severity scoring | (n=46): 25 (54) | Severity scoring system developed and |
| | problems in | cow's milk | | Sodium 17.46 | system based | P<0.001 | evaluated in pilot study: |
| | Taiwan | protein | | Potassium 61.58 | on stool | | Hard stool: 0, no hard stool; 1, hard and |
| | | allergy, | | Chloride 43.40 | consistency, | Good response | long form, 2; |
| | | Hirschsprung' | | Calcium 60.87 | frequency and | (number and % of | Difficulties with defecation: 0, no |
| | | s disease and | | Phosphate 31.46 | volume of | <u>children)</u> | difficulties; 1, irritability; 2, crying |
| | | others | | Magnesium 9.12 | stools and | -At 2 weeks: | Frequency of defecation: 0, >3 |
| | | | | | difficulties in | Novalac-IT (n=47): | times/week; 1, 1 to 3 times/week; 2, <1 |
| | | Exclusion Page 1 | | Osmolality: 300 | | 17 (36) | time/week |
| | | <u>criteria</u> : | | | 3 mild | | Stool weight (g//kg/week): 1, >35; 2, 20 |
| | | unclear | | Comparison: | constipation; 4 | | to 35; 3, <20 |
| | | | | 20% strengthened | | (n=46): 13 (28) | |
| | | | | Novalac regular | 7 or 8 severe) | | Reviewer comments: |
| | | | | infant formula | | -At 1 month: | No sample size calculation performed |
| | | | | | -Remission: | Novalac-IT (n=47): | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|--|---|--|--|
| | | | | Composition per 100 mL Energy (cal/100 mL): 78 Protein (g): 1.89 Whey/casein: 50/50 Fat (g): 3.96 Carbohydrates (g): 8.69 70% Lactose, 30% Maltodextrin Major minerals (mg) Sodium 21.24 Potassium 70.20 Chloride 46.80 Calcium 70.20 Phosphate 42.12 Magnesium 7.02 Osmolality: 300 | asymptomatic -Good response: decrease in severity of ≥ 2 -Fair response: decrease in severity of 1 to 3 | 22 (47) Strengthened formula (n=46): 11 (24) Fair response (number and % of children) -At 2 weeks: Novalac-IT (n=47): 14 (30) Strengthened formula (n=46): 10 (22) -At 1 month: Novalac-IT (n=47): 17 (36) Strengthened formula (n=46): 23 (50) Not improved (number and % of children) -At 2 weeks: Novalac-IT (n=47): 16 (34) Strengthened formula (n=46): 23 (50) -At 1 month: Novalac-IT (n=47): 8 (17) Strengthened formula | Irrelevant reason given for non-blinding the study Unclear how both formulas were administered No dropouts/lost to follow-up reported Study not controlled for potential confounders Source of funding: Not stated Intestinal Transit provided free samples of Novalac-IT formula. According to authors there was no other grant from the company, which was neither involved in the design of the study |
| | | | | | | Strengthened formula (n=46): 23 (50) | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | Symptoms free (number and % of children) -At 2 weeks: Novalac-IT (n=47): 18 (38) Strengthened formula (n=46): 12 (26) N.S -At 1 month: Novalac-IT (n=47): 28 (60) Strengthened formula (n=46): 16 (35) P=0.029 -At 2 months: Novalac-IT (n=47): 35 (75) Strengthened formula (n=46): 18 (39) P<0.001 | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-------------------------|-----------------------------|---------------------------|------------------------|--------------------------------|--|
| Covine et al | Level | 400 abildras | s 95 children | latam rantian. | Measures | Charl fra avvanav | Additional information from attacks |
| Savino et al. | Study Type: | 123 children | | Intervention: | Duration of | Stool frequency | Additional information from study: |
| | | la alvaia a | 50 boys | New formula (NF) | treatment | (number/day) (mean | Constipation defined as a stool |
| management of | RCI | Inclusion criteria: | aga at atudu | Composition per | 14 days | ± SD) | frequency of less than 1 stool a day |
| digestive problems during | <u>Evidence</u> | Enteria. Formula-fed | age at study entry (months) | 100 ml (Omneo / | Assassment | -at study entry | Parents given a structured questionnaire |
| the first months | level: | healthy term | entry (months) | Conformil): | Assessment point (s): | NF group (n=55): 0.53 ± 0.5 | in order to monitor frequency of |
| of life. 2005. | 1- | | -intervention | Comornin). | On days 1, 7 | 0.55 ± 0.5 | symptoms, feeding volume and side |
| Acta | • | months of | | Energy: 70 kcal | and 14 | SF group (40): | effects |
| Paediatrica | | age with | group: 1.55 ± 0.88 | Protein equivalent | | 0.60 ± 0.5 | enecis |
| 94[SUPP 449], | | constipation | 1.55 ± 0.66 | (g): 1.7 | Follow-up | N.S | No significant differences in baseline |
| 120- | the efficacy | Consupation | -control group: | Casein: whey: | period: | IN.S | characteristics between the 2 groups |
| 124Norway. | | Exclusion | 1.28 ± 0.66 | 100% whey | No follow-up | -on day 7 | characteristics between the 2 groups |
| 124INOIWay. | | criteria: | 1.20 ± 0.00 | hydrolysate | conducted after | NF group (n=55): | When an infant eligible to study came to |
| | formula based | | Country: | liyuloiysale | treatment | 1.79 ± 0.96 | the doctor, child was randomly assigned |
| | | problems | Italy | Carbohydrate (g): | finished | 1.79 ± 0.90 | to the study or the control group, the |
| | | and/or any | italy | 8.4 | IIIIISIIGU | SF group (40): | next infant with the same symptoms was |
| | predominantly | | | Lactose:2.9 | Outcome | 1.31 ± 0.89 | matched to the previous infant and |
| | | any kind of | | Maltodextrine: 4.0 | | 1.51 ± 0.09 | assigned to the other group |
| | the β-position, | | | Starch: 1.5 | <u>ivicasures.</u> | difference: | assigned to the other group |
| | oligosaccharid | | | Glaren. 1.5 | -stool | 0.48 (CI 95%: 0.09; | 28 children excluded after randomisation |
| | | before the | | Prebiotic | characteristics: | 0.87) | because at entry they had more than 1 |
| | , | beginning of | | | frequency and | p=0.02 | evacuation |
| | , | the study and | | (g): 0.8 | consistency | p=0.02 | Cvacation |
| | | during the | | (9). 0.0 | Consistency | -on day 14 | Reviewer comments: |
| | | study period | | Fat (g): 3.3 | | NF group (n=55): | Sample size calculation not performed |
| | hydrolysed | olday polica | | Palmitic acid:0.60 | | 2.04 ± 1.04 | Campie 6/20 Galcalation flot portofflied |
| | protein, low | | | T ammilio aora.o.oo | | 2.0121.01 | Inadequate randomisation |
| | lactose | | | Minerals (mg) | | SF group (40): | madequate randomication |
| | content and | | | Sodium:23 | | 1.64 ± 0.99 | Allocation concealment not described |
| | higher density | | | Potassium: 66 | | 1.0.1 = 0.00 | |
| | ingilor delicity | | | Chloride: 50 | | difference: | Study not reported as blinded |
| | | | | Calcium: 53 | | 0.40 (CI 95%: -0.03; | otaa, roportoa ao omiaoa |
| | | | | Phosphorus: 31 | | 0.83) | Stool consistency post- treatment not |
| | | | | Iron: 0.5 | | p=0.07 | reported |
| | | | | Zinc: 0.5 | | r | - F |
| | | | | | | Mean difference in | No dropouts/lost to follow-up children |
| | | | | Comparison: | | stool frequency | reported |
| | | | | Standard formula | | between the 2 groups | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---|------------------------------------|---|-------------------------------|
| | | | | (SF) (composition not reported in paper) Feeding volume based on a feeding ad libitum procedure. Feeding frequency decided by the parents and not influenced by the study protocol | | adjusted for gender, age at entry, maternal instruction, parity, birth weight, number of feedings/day and stool frequency at entry -Days 0 to 7: 0.60 (CI 95%: 0.19; 1.01) p=0.004 -Days 0 to 14: 0.53 (CI 95%: 0.11; 0.90) p=0.015 | Source of funding: not stated |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|------------------------------|----------------------------|---------------------------|---------------------------|------------------------|--|---|
| | Level | | S | | Measures | | |
| Savino et al. | Study Type: | 604 children | 604 children | Intervention: | Duration of | Stool frequency | Additional information from study: |
| "Minor" feeding | Prospective | | (232 with | New formula (NF) | treatment | 232 infants with | Constipation defined as a stool |
| problems during | case series | <u>Inclusion</u> | constipation) | | 14 days | constipation | frequency of less than 1 stool a day |
| the first months | | criteria: | | Composition per | | | |
| | <u>Evidence</u> | Formula-fed | age at entry | 100 ml | <u>Assessment</u> | -increase in number | Parents given a questionnaire in order to |
| a partially | level: | healthy term | (months, total | | point (s): | of stools per day | monitor frequency of symptoms, feeding |
| hydrolysed milk | 3 | infants up to 3 | | Energy: 70 kcal | On days 1, 7 | during study period: | volume and side effects. Number of |
| formula | | months of | 1.35 ± 0.77 | Protein equivalent | and 14 | 147 infants (63.4%) | stools were recorded daily |
| containing | Study aim: | age seen by | | (g): 1.7 | | | |
| fructo- and | To investigate | | gender not | Casein: whey: | Follow-up | -average increase: | A total of 932 infants enrolled: 604 |
| galacto- | whether a | because of | reported | 100% whey | period: | | |
| | new infant | colic and/or | 0 | hydrolysate | No follow-up | 0.27; p<0.005) | 358 infants excluded from study: 154 |
| s. 2003. Acta | formula | constipation | Country: | Carla alas salvadas (as). | conducted after | | completed only the first step and did not |
| Paediatrica | commercially | and/or | Italy | Carbohydrate (g): | treatment | -average increase | return for the visit on day 14, 131 infants |
| | available in | regurgitation. | | 8.4 Lactose:2.9 | finished | between day 1 and | excluded because of incomplete data. |
| | Italy is useful as a dietary | Normal birth weight (>2500 | | Maltodextrine: 4.0 | Outcomo | day 7: 0.41 (CI 95%: 0.51 to 0.23; p<0.05) | 73 infants required medication during the 1rst week of study and were |
| 90Norway. | | g), normal | | Starch: 1.5 | Measures: | 0.51 to 0.25, p<0.05) | therefore excluded |
| | infants with | weight gain (≥ | | Starti. 1.5 | ivicasures. | -average increase | linererore excluded |
| | minor feeding | 150g/week) | | Prebiotic | -stool frequency | between day 7 and | Reviewer comments: |
| | problems | and normal | | oligosaccharides | -3tool frequency | day 14: 0.04 (NS) | No description of the scoring system |
| | problemo | physical | | (g): 0.8 | -parents' | day 11. 0.01 (1 10) | used to evaluate parent's satisfaction |
| | | examination | | (9). 0.0 | evaluation of | -no improvement of | was provided |
| | | Ondirini dilori | | Fat (g): 3.3 | formula | symptoms: 85 infants | mas provided |
| | | Exclusion | | Palmitic acid:0.60 | | (26.6%) | Source of funding: |
| | | criteria: | | | | () | Not stated |
| | | Neonatal | | Minerals (mg) | | Mean parent | |
| | | problems, use | | Sodium: 23 | | evaluation of formula | |
| | | of any kind of | | Potassium: 66 | | | |
| | | medication | | Chloride: 50 | | 7.9 ± 1.8 | |
| | | the week | | Calcium: 53 | | | |
| | | before the | | Phosphorus: 31 | | 550 parents (91%) | |
| | | beginning of | | Iron: 0.5 | | gave a positive | |
| | | the study or | | Zinc: 0.5 | | judgement (score 6 to | |
| | | during the | | <u>_</u> | | 10) | |
| | | study period | | Feeding volume | | | |
| | | | | based on a | | | |
| | | | | feeding ad libitum | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|--|------------------------------------|-------------|-------------------|
| | | | | procedure. Feeding frequency decided by the parents and not influenced by the study protocol | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|-----------------------|--|
| Pina et al. | Study Type: | 3487 children | 604 children | Intervention: | | 91.6% of cases of | Additional information from study: |
| Prevalence and | Prospective | (total | (with | Novalac Anti- | treatment | constipation resolved | Study on effectiveness included 2069 |
| dietetic | case series | population) | constipation) | Constipation: | 30 days | within 7 days | infants with MGDs. Effectiveness was |
| management of | | | | formula with | | | evaluated among 1441 infants who |
| | <u>Evidence</u> | Inclusion | 52.2% boys (of | adapted | Assessment | Number of daily | completed follow-up. Premature study |
| gastrointestinal | level: | criteria: | the total | concentration of | point (s): | stools (mean ± SD) | termination due to adverse events in |
| disorders in | 3 | Infants up to | population) | magnesium and | Immediately | Baseline: 0.6 ± 0.7 | 2.7% cases, parent decision in 6.9%, |
| milk-fed infants. | | 4 months of | age at | lactose | after treatment | At 30 days: 1.7 ± 0.8 | loss to follow-up in 1.64%, protocol |
| 2008. World | Study aim: | age fed with | consultation: 1 | | was completed | | violations in 2.46% and non-specified |
| Journal of | To assess the | artificial milk | week to 17 | No other details | | Type of stools (% | reasons in 16.62% |
| Gastroenterolog | prevalence of | formulas, | weeks (total | regarding feeding | Follow-up | children) | |
| y 14[2], 248- | mild | presence of | population) | volume/frequency | period: | | A questionnaire addressing the different |
| 254China. | gastrointestin | MGDs, | | were provided | No follow-up | -Normal: | symptoms and their intensity was |
| | al disorders | possibility of | Country: | | made after | Baseline: 33.40 | designed for each disorder |
| | (MGDs) in | feeding | Spain | Comparison: | treatment | At 30 days: 95.60 | _ |
| | milk-fed | infants with | | N.A | finished | - | Satisfaction of parents/tutors with the |
| | infants in | some product | | | | -Hard | formulas assessed on final visit by |
| | paediatric | of the | | | | Baseline: 66.60 | means of a Likert-type scale with 5 |
| | practice and | Novalac line | | | Measures: | At 30 days: 4.40 | possible answers: from very satisfied to |
| | to evaluate | of formulas, | | | | - | very dissatisfied |
| | the | continuation | | | -type of stools | Presence of pain or | - |
| | effectiveness | of these | | | | discomfort (% | Reviewer comments: |
| | and | formula on an | | | -presence of | children) | No definition of constipation given |
| | satisfaction | exclusive | | | pain or | | |
| | with dietetic | basis for at | | | discomfort | -Yes: | Not completely clear how outcomes |
| | treatment: | least 30days | | | | Baseline: 90.00 | were measured and who measured |
| | specifically | with no | | | -external help | At 30 days: 10.40 | them |
| | elaborated | incorporation | | | needed for | - | |
| | formulas | of other foods | | | defecation | -No: | No definition of "resolved case" given |
| | belonging to | to the diet | | | | Baseline: 10.00 | |
| | the Novalac | | | | -satisfaction of | At 30 days: 89.60 | Source of funding: |
| | line of | Exclusion | | | parents/tutors | | Not stated |
| | products | criteria: | | | | External help needed | |
| | | Not clearly | | | -adverse events | for defecation | |
| | | stated | | | | -Yes: | |
| | | | | | | Baseline: 76.10 | |
| | | | | | | At 30 days: 8.80 | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|---|-------------------|
| | | | | | | -No: | |
| | | | | | | Baseline: 23.90 | |
| | | | | | | At 30 days: 91.20 | |
| | | | | | | -satisfaction of | |
| | | | | | | parents/tutors: | |
| | | | | | | 90.0% of parents | |
| | | | | | | satisfied with | |
| | | | | | | treatment | |
| | | | | | | treatment | |
| | | | | | | Adverse events (for | |
| | | | | | | all formulas, no | |
| | | | | | | subgroup analysis): | |
| | | | | | | Reported in 3.9% | |
| | | | | | | infants of total | |
| | | | | | | population. Most | |
| | | | | | | frequent affected | |
| | | | | | | digestive tract (1.4%), | |
| | | | | | | including diarrhoea | |
| | | | | | | and constipation, and | |
| | | | | | | respiratory apparatus | |
| | | | | | | | |
| | | | | | | (0.7%) (E.g. bronchiolitis and | |
| | | | | | | | |
| | | | | | | bronchitis). 10 infants (0.5%) required | |
| | | | | | | | |
| | | | | | | hospital admission for | |
| | | | | | | septicaemia (n=1), | |
| | | | | | | dehydration (n=2), | |
| | | | | | | vomiting (n=1), hernia | |
| | | | | | | (n=1) and bronchitis or bronchiolitis (n=2) | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|---------------------------|-----------------------|---------------------------|------------------------------|------------------------------|----------------------------|--|
| | Level | | S | | Measures | | |
| Kokke et al. A | Study Type: | 135 children | 97 children | Intervention: | Duration of | <u>Defecation</u> | Additional information from study: |
| dietary fiber | Double-blind | | | Yogurt drink with | treatment | frequency/week(| Randomisation performed by use of |
| mixture versus | RCT | <u>Inclusion</u> | fibre mix group | mixed dietary fibre | | mean) | sequential numbers allocated to patients |
| lactulose in the | | criteria: | (n=42): | (10g/125mL) | intervention | -At 8 weeks: | at study entry and coordinated by the |
| treatment of | <u>Evidence</u> | Constipated | 20 boys | | period | Fibre (n=42): 7 | logistic manager of Numico Research |
| childhood | <u>level:</u> | children | median age: 5.5 | | | | using a block design |
| constipation: a | 1+ | referred to | years (1 to 12 | (per 100mL): | 4-week | Lactulose (n=55): 6 | |
| double-blind | | hospital | years) | 3.0 g | weaning period | N.S | Bottles with yogurt prepared and packed |
| randomized | Study aim: | outpatient | | transgalacto- | | | by Numico Research and transported to |
| controlled trial. | | clinic for | lactulose group | oligosacharides | Assessment | Number of patients | hospital. Treatment products could not |
| 2008. Journal of | | constipation | (n=55): | 3.0 g inulin | point (s): | with ≥ 1 faecal | be distinguished from each other with |
| Pediatric | efficacy and | who fulfilled | 23 boys | 1.6 g soy fibre | At 3, 8 and 12 | <u>incontinence</u> | respect to colour, taste or consistency |
| Gastroenterolog | | | median age 5.0 | 0.33g resistant | weeks | episodes/week | |
| y and Nutrition | dietary fibre | criteria for | years (1 to 12 | starch 3 | | -At 8 weeks: | Sample size based on primary outcome |
| 47[5], 592-597 | mixture and | constipation: | years) | 0 | Follow-up | Fibre (n=42): 9 | variable, defecation frequency. It was |
| | compare it | stool | 0 | Comparison: | period: | | calculated that a random allocation of |
| | | frequency <3 | Country: The | Yogurt drink | No follow-up conducted after | Lactulose (n=55): 5 N.S | 150 children would allow for the |
| | in the | times/week, | Netherlands | containing lactulose(10g/125 | treatment | N.S | detection of a mean difference in defecation of 1.0/week between the 2 |
| | treatment of childhood | faecal incontinence | inemenanus | mL) (Duphalac | finished | Ctool consistency | |
| | constipation | ≥ 2 | | Lactulose) | linisnea | Stool consistency (mean) | groups |
| | Consupation | times/week. | | Lacidiose) | Outcome | -At 3 weeks: | No significant differences found in |
| | | periodic | | | | Fibre (n=42): 3.5 | baseline characteristics between the 2 |
| | | passage of | | | <u>ivicasures.</u> | | groups with a power of 80% and |
| | | large | | Both products | 1. primary | Lactulose (n=55): 4.5 | alfa=0.05 |
| | | amounts of | | taken at breakfast | | P<0.01 | ana-0.00 |
| | | stool at least | | and in case of ≥ 2 | outoonio. | 1 40.01 | Defecation noted on a daily basis during |
| | | once very 7 to | | bottles also at | -defecation | -At 8 weeks: | treatment period. Faecal incontinence |
| | | 30 days, or a | | lunch | | Fibre (n=42): 3.6 | each day assessed "yes" or "no", stool |
| | | palpable | | | o quioo,, oo | | consistency according to Bristol Stool |
| | | abdominal or | | Amount of | 2. secondary | Lactulose (n=55): 4.0 | Form Scale. Data recorded daily in |
| | | rectal mass | | fibre/fluid intake | | P=0.01 | bowel diary by parents or patients. |
| | | | | daily depended on | | | , , , , , |
| | | Exclusion | | patient's body | -faecal | Number of patients | Adverse effects defined as any adverse |
| | | criteria: | | weight: | incontinence | using step-up | change from baseline (pre-treatment) |
| | | Organic | | | each day | medication | condition, which occurred during the |
| | | causes of | | Intervention | _ | -At 3 weeks: | course of the study after treatment |
| | | defecation | | period: | -stool | Fibre (n=42): 13 | started, whether it was considered to be |

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|---------------|--------------|--------------------------|----------------|--------------------------------|------------------|--------------------------|---|
| Information | Evidence | Patients | Characteristic | Comparison | Outcome | | |
| | Level | | S | | Measures | | |
| | | disorders | | <15 kg: 1 bottle | consistency | | related to treatment |
| | | including | | (125 mL, 10g | | Lactulose (n=55): 7 | |
| | | Hirschsprung' | | fibres) | -use of step-up | P=0.028 | 33 patients dropped-out during study |
| | | s disease, | | | medication | | period: 22 in fibre group after 1 to 56 |
| | | spina bifida, | | 15 to 20kg: 2 | | -At 8 weeks: | days (median 7) and 11 in lactulose |
| | | hypothyroidis | | bottles (250 mL, | -adverse effects | Fibre (n=42): 20 | group after 1 to 51 days (median 8) |
| | | m or other | | 20g) | | | (p=0.020). Those patients refused to |
| | | metabolic/ren | | 001 01 11 | | Lactulose (n=55): 21 | drink the yogurt. 3 patients lost to follow- |
| | | al | | >20 kg: 3 bottles | | N.S | up: 1 fibre, 2 lactulose. 2 exclusions |
| | | abnormalities, | | (375 mL, 30g) | | A4 40 also | after randomisation in lactulose group: 1 |
| | | mental | | 14/a a mim ay m a mia ali | | -At 12 weeks: | coeliac disease, 1 spina bifida occulta |
| | | retardation, | | Weaning period: <15 kg: 0.5 | | Fibre (n=42): 21 | Paviouer comments: |
| | | use of drugs influencing | | bottle/day (week 9 | | Lactulose (n=55): 26 | Reviewer comments: Method of allocation concealment not |
| | | gastrointestin | | & 10); 0.5 every | | N.S | described |
| | | al function | | other day (week | | IN.S | described |
| | | other than | | 11 &12) | | Adverse effects | Study not controlled for potential |
| | | laxatives, use | | 11 (412) | | No serious or | confounders |
| | | of lactulose, | | 15 to 20kg: 1 | | significant side effects | Comoundoro |
| | | other | | bottle/day (week 9 | | recorded | Unclear how adverse effects were |
| | | laxatives, | | & 10); 1 every | | | recorded. |
| | | prebiotics, | | other day (week | | Fibre (n=42): 1 dose- | |
| | | probiotics or | | 11 &12) | | related persistent | ITT analysis not performed |
| | | antibiotics in | | , | | diarrhoea | , , |
| | | the previous 4 | | >20 kg: 2 | | | Source of funding: |
| | | weeks before | | bottles/day (week | | Lactulose (n=55): 2 | The Scientific Research Foundation |
| | | the first visit | | 9 & 10); 1 | | dose-related | project SW) 2001. |
| | | | | bottle/day (week | | persistent diarrhoea | One author received financial support |
| | | | | 11 &12) | | | throught project no.9.001, which is a |
| | | | | | | | subproject of Business aimed |
| | | | | If persistent | | | Technological Cooperation project |
| | | | | diarrhoea | | | 00176. 2 authors were researchers and |
| | | | | reported, original | | | employees of Danone Research BV |
| | | | | dose reduced by | | | (formerly Numico Research BV) |
| | | | | 50% | | | |
| | | | | If aliminal | | | |
| | | | | If clinical | | | |
| | | | | parameters | 1 | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|--|------------------------------------|-------------|-------------------|
| | | | | compared to baseline did not improve 3 weeks after start of intervention period, step-up medication (Macrogol 3350) given per protocol | medsures | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|------------------------|---|
| | Level | | S | | Measures | | |
| Loening-Baucke | | 31 children | 31 children | <u>General</u> | Duration of | Children with <3 | Additional information from study: |
| et al. Fiber | Double-blind | | | Disimpaction with | treatment | BMs/week (%) | Constipation defined as a delay or |
| , , | RCT (cross- | <u>Inclusion</u> | 16 boys | | 2 treatment | | difficulty in |
| | over) | criteria: | | | periods of 4 | Fibre (n= 31): 19% | defecation, present for >2 weeks, and |
| the treatment of | | Otherwise | age: 4.5 to 11.7 | | weeks each | P<0.05 | sufficient to cause significant distress to |
| childhood | <u>Evidence</u> | healthy | years (mean: | during rectal | | | child |
| constipation. | level: | | 7.1 ± 2.0 years) | examination | <u>Assessment</u> | Stool consistency | |
| 2004. Pediatrics | 1+ | than 4 years | | (58% of patients | point (s): | Initial (n= 31): 0.3 ± | Encopresis defined as the involuntary |
| 113[3 Pt 1], | | who had | Countries: | continued with | At 4 and 8 | 0.9 | loss of formed, semiformed, or liquid |
| e259-e264 | Study aim: | chronic | USA & Italy | their | weeks | Placebo (n= 31): 1.2 | stool into the child's underwear in the |
| | | functional | | preevaluation | | ± 0.9 | presence of functional constipation after |
| | | constipation | | laxative during | Follow-up | Fibre (n= 31): 1.5 | the child |
| | supplementati | | | whole study | period: | ±0.9 | has reached the age of 4 years |
| | on with | with or | | period) | No follow-up | P<0.05 as compared | |
| | glucomannan | without | | | conducted after | to initial data | It had been previously calculated that at |
| | | encopresis | | Intervention: | treatment | | α=0.05; 26 subjects would allow a power |
| | the treatment | | | | finished | Children with | of approximately 0.95 to detect a |
| | of children | Exclusion | | capsule | | <u>encopresis</u> | difference of 0.7 versus 0.2 in achieving |
| | | criteria: | | containing | <u>Outcome</u> | Initial (n= 31): 58% | normal bowel patterns in the crossover |
| | constipation | Hirschsprung' | | glucomannan, a | Measures: | | design |
| | | S | | polysaccharide of | -efficacy: | Fibre (n= 31): 42% | |
| | | disease, | | d-glucose and d- | | | Patients randomized by envelope into 1 |
| | | hypothyroidis | | | changes in | Frequency of soiling | of 2 treatment arms. Blinding done by |
| | | m, mental | | | frequency of | episodes/wk (n=18) | having the medication labelled |
| | | deficiency, | | alimentary fibre. | bowel | Initial (n=18):_9.9 ± | glucomannan A and glucomannan B |
| | | chronic | | | movements | 12.3 | with the code kept by the company until |
| | | debilitating | | Comparison: | (BMs) | Placebo (n= 18): 4.2 | study was completed and analyzed. |
| | | diseases, | | Glucomannan A: | | ± 4.8 | Glucomannan A was a capsule |
| | | neurological | | capsule | soiling | Fibre (n= 18): 4.0 ± | containing maltodextrins as placebo. |
| | | abnormalities, | | containing | frequency | 6.3 | Glucomannan B was a capsule |
| | | previous | | maltodextrins as | | P<0.05 as compared | containing glucomannan, a |
| | | surgery of the | | placebo. | -successful | to initial data | polysaccharide of d-glucose and d- |
| | | colon or anus | | | treatment | | mannose, =450 mg of alimentary fibre |
| | | | | Group 1: placebo | | Successful treatment | |
| | | | | first and then | -parents' global | Placebo (n= 31): 13% | Patients and their parents kept diary |
| | | | | glucomannan | assessment | Fibre (n= 31): 45% | sheets during the 8 weeks of study. |
| | | | | | | P<0.05 as compared | They recorded daily each BM, soiling |
| | | | | Group 2: | -overall | to placebo treatment | episode, abdominal pain episode, and |

| | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|-------------|-------------------|-----------|------------------|---------------------------------------|---------------------|--|---|
| Information | Evidence Level | Patients | Characteristic s | Comparison | Outcome Measures | | |
| | 2010. | | | glucomannan first | | | medication used and reported at the end |
| | | | | and then placebo | palatability | Improved (parent | of each treatment period the associated |
| | | | | | | <u>rating)</u> | subjective symptoms such as stool |
| | | | | -Placebo and | -safety: side | | consistency, new occurrence of |
| | | | | glucomannan | effects | Fibre (n= 31): 68% | abdominal pain, bloating, abdominal |
| | | | | doses: | | P<0.05 as compared | distension, excessive gas, or diarrhoea. |
| | | | | 100 mg/kg body | | to placebo treatment | Stool consistency was assessed rating |
| | | | | weight daily | | | the stool consistency as hard like rocks, |
| | | | | (maximal 5 | | Outcomes controlled | pellets= 0, firm = 1, soft like banana = 2, |
| | | | | g/day), rounded to the nearest 500 | | for confounders -successful treatment | loose like milkshake = 3, and watery = 4 |
| | | | | mg, because each | | | Successful treatment rated by physician |
| | | | | capsule contained | | | and defined as ≥ 3 bowel movements |
| | | | | 500 mg. Each | | | per week and ≤ 1 soiling episode in the |
| | | | | capsule either | | low or acceptable | last 3 weeks with no abdominal pain. |
| | | | | opened and | | fibre intake (P>0.6) | Parents' global assessments: whether |
| | | | | sprinkled on food | | | they believed that the child was better |
| | | | | given with 50 mL | | - more children with | during the first or second treatment |
| | | | | of fluid per | | encopresis in the | period |
| | | | | capsule; given as | | laxative group (78% | |
| | | | | a solution, | | vs. 31%; P<0 .02), | No significant differences in baseline |
| | | | | whereby the | | | characteristics between the 2 groups |
| | | | | content of each | | children in the | |
| | | | | 500-mg capsule | | laxative group were | 46 children originally recruited. 13 |
| | | | | was mixed with 50 | | treated successfully | children did not show up for the 4-week |
| | | | | mL of fluid of the | | | follow-up: 7 children randomized to |
| | | | | child's choice; or | | placebo | placebo first and 6 children randomized |
| | | | | swallowed as a | | (P <0 .01) | to fibre first. 2 constipated girls |
| | | | | capsule with 50 mL of fluid for | | - Children with | completed the first 4 weeks of the study |
| | | | | | | | only: 1 received placebo and 1 received |
| | | | | each capsule. | | constipation only were significantly | fibre; both recovered from chronic constipation and abdominal pain during |
| | | | | In addition. | | more likely to be | the first 4 weeks of treatment and did |
| | | | | parents instructed | | treated successfully | not return for the 8-week visit. Data from |
| | | | | to have the child | | with fibre | the 13 children who entered the study |
| | | | | sit on the toilet 4 | | | and were randomized but did not come |
| | | | | times daily after | | constipation and | for follow-up and the 2 children who did |
| | | | | meals and to keep | | encopresis (28%; | not complete the study were excluded |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---|------------------------------------|------------------------------------|---|
| | | | | a stool diary. No enemas given during each treatment period, unless rectal disimpaction felt during rectal examination at assessment visits | | onset of abdominal pain, bloating, | from the analysis. Initial data of these 15 children not significantly different from the data of the 31 children who completed the study, except soiling frequency per week was significantly less (4.0 ± 1.4; P<0.001). Data analysis includes 31 children with functional constipation with or without encopresis Reviewer comments: No definition of soiling given. Unclear how different this would be from the authors' definition of encopresis High dropout rate: 28%. ITT analysis not performed Source of funding: DicoFarm (Rome, Italy) provided research support and the medications for the study |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|------------------------|------------------------------|--|
| | Level | . anomo | S | Companicon | Measures | | |
| Castillejo et al. | Study Type: | 56 children | 56 children | Intervention: | Duration of | No. of bowel | Additional information from study: |
| A controlled, | Double-blind | | | cocoa husk | treatment | movements per week | Chronic functional constipation defined |
| randomized, | RCT (pilot | <u>Inclusion</u> | 22 boys | supplement rich | 4 weeks | (mean ± SD) | in accordance with Rome II diagnostic |
| double-blind | study) | | Mean age 6.3 ± | in dietary fibre + | | Difference (95% CI): | criteria, by the presence, for at least 12 |
| trial to evaluate | | Children aged | 2.2 years | standardized toilet | | | (not necessarily consecutive) weeks in |
| the effect of a | <u>Evidence</u> | 3 to 10 years | | training | point (s): | 0.67 (-0.76 to 2.10) | the preceding 12 months, of at least 2 of |
| supplement of | level: | referred to | Country: | procedures | Immediately | p=0.780 | the following symptoms: straining in |
| cocoa husk that | 1+ | pediatric | Spain | | after treatment | | >25% of defecations; lumpy or hard |
| is rich in dietary | | gastroenterol | | 1 sachet (5.2 g): | finished | -Cocoa husk group | stools in >25% of defecations; a |
| fiber on colonic | Study aim: | ogy | | 4 g cocoa husk + | | Basal (n=24): 3.86 | sensation of incomplete |
| transit in | to evaluate | outpatients' | | 1 g | Follow-up | ±2.05 | evacuation in >25% of defecations; a |
| constipated | the effect of a | clinic between | | betafructosans | period: | Final (<i>n</i> =24): 6.16 | sensation of anorectal |
| pediatric | palatable | January 2004 | | | | ±3.35 | obstruction/blockage in >25% of |
| patients. 2006. | cocoa husk | and April | | (53.2 g of fibre | made after | Difference (95% CI): | defecations; a need for manual |
| Pediatrics | supplement | 2005 with | | (39.6 g of total | | 2.40±3.16 | maneuvres to facilitate >25% of |
| 118[3], e641- | that is rich in | chronic | | fibre and 13.6 g of | finished | | defecations (e.g., digital evacuation, |
| e648 | fibre on | constipation | | betafructosans) | | -Placebo group | support of the pelvic floor); and <3 |
| | intestinal | • | | per 100 g of | Outcome | Basal (<i>n</i> =24): 3.18± | defecations per week |
| | transit time | Exclusion | | product. Insoluble | Measures: | 1.93 | · |
| | and other | criteria: | | fibre 37.2% and | -number of | Final (<i>n</i> =24): 5.08 | Treatment was blinded to both patients |
| | indices of | presence | | soluble fibre 2.4% | bowel | ±2.10 | and investigator until the study was |
| | constipation | of fecal | | of total fibre | movements per | Difference (95% CI): | completed and analyzed. Patients |
| | in children | impaction that | | Cellulose and | week | 1.73 ±1.73 | randomly assigned to treatment 1 or 2 in |
| | with idiopathic | required | | uronic acids the | | | a ratio of 1:1. A randomization list was |
| | chronic | enema in the | | main type of | -stool | Hard stool | designed by the manufacturers of the |
| | constipation | 7 days | | insoluble fibre and | consistency | consistency (% | supplement and the placebo (Madaus |
| | - | before the | | soluble fibre, | - | children) | SA) using a computer random-number |
| | | start of the | | respectively) | -pain with | -Cocoa husk group | generator in 20 blocks of 4 patients |
| | | study, | | | defecation | Basal (n=24): 95.8 | each. The details of the randomization |
| | | treatment with | | Comparison: | | Final (n=24): 41.7 | codes were kept in sealed envelopes |
| | | dietary fibre, | | placebo + | -safety | | away from the investigators. Only in |
| | | bulk-forming | | standardized toilet | , | -Placebo group | cases of the utmost necessity (eg, |
| | | agents, or | | training | | Basal (<i>n</i> =24): 95.8 | serious adverse events) did the |
| | | laxatives in | | procedures | | Final (n=24): 75.0 | coordinator of the study allow the |
| | | the 2 weeks | | | | P=0.017 | investigator to know the treatment |
| | | before the | | 1 sachet (5.2 g): | | | assigned to the patient |
| | | start of the | | glucose, cocoa | | Subjective | |
| | | study, | | flavouring, and | | improvement in stool | Because of lack of previous studies and |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--|---------------------------|---------------------------|------------------------|--|--|
| imormation | Level | latients | S | Comparison | Measures | | |
| | | constipation | | excipients | | consistency (n | likelihood of methodological difficulties |
| | | attributable | | | | children) | (in the evaluation of the main |
| | | to organic or | | -doses for both | | P=0.039 | parameters) |
| | | anatomic | | products: | | | in carrying out a study on this kind of |
| | | causes | | | | Cocoa husk group | population, authors designed a pilot |
| | | (Hirschsprung | | Children aged 3 to | | (<i>n</i> =24) | study with a minimum sample from the |
| | | 's disease, | | 6 years: 1 sachet | | Improvement : 14 | statistical point of view |
| | | hypothyroidis | | before lunch and | | No Improvement: 10 | |
| | | m, mental | | 1 sachet before | | | Fibre supplement and placebo |
| | | deficiency, | | dinner | | Placebo group (<i>n</i> =24) | administered as a soluble powder in |
| | | psychiatric | | | | Improvement : 6 | sachets of identical weight (5.2 g) and |
| | | illnesses, | | Children aged 7 to | | No Improvement: 18 | presentation |
| | | chronic | | 10 years: 2 | | <u>Subjective</u> | |
| | | debilitating | | sachets | | improvement in pain | At baseline and after 4 weeks of |
| | | diseases, | | before lunch and | | P=0.109 | treatment, investigators evaluated bowel |
| | | neurologic | | dinner | | Cocoa husk group | movement habits and stool consistency |
| | | abnormalities, | | | | (n=24) | using a diary completed by patients' |
| | | or previous | | Parents instructed | | Improvement : 16 | parents; and received a subjective |
| | | surgery of the | | to dissolve | | No Improvement: 8 | evaluation from the parents regarding |
| | | colon or | | content of the | | Placebo group | the efficacy of the treatment. Adherence |
| | | anus), renal | | sachets in 200 mL | | (n=24) | to the intervention evaluated by the |
| | | insufficiency, | | of whole milk | | Improvement : 11 | same investigator using a visual |
| | | hypocalcemia | | before ingestion | | No Improvement: 13 | analogical scale (in the case of |
| | | , | | | | 0-1-1 | standardized toilet training procedures) |
| | | hyperkalemia, | | | | Safety No significant | and counting the empty sachets that were returned |
| | | or any other metabolic | | | | | were returned |
| | | | | | | adverse effects, such | No significant differences in baseline |
| | | | | | | | |
| | | | | | | | characteristics between the 2 groups |
| | | J . | | | | | 9 shildren withdraw from study before its |
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| | | diseases at the start of the study; long-term use of drugs that affect gastrointestin al motility (eg, imipramine, iron or calcium | | | | as a new onset of abdominal pain, bloating, abdominal distension, excessive gas, diarrhoea, or anaphylactic symptoms, reported during the 4-week period with either treatment No significant | No significant differences in baseline characteristics between the 2 groups 8 children withdrew from study before a completion (5 children discontinued study because of the difficulty of the protocol, and 3 were excluded because of the presence of positive antigliadin and antiendomysium antibodies). Data refer only to 48 participants who completed the study |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|--|--------------------------------|------------------------------|------------------------------------|---|--|
| | | supplements, anticonvulsan ts), inability to adhere to the study's medications or procedures | | | | changes between groups in relation to hemoglobin concentrations; hematocrit; serum ferritin; or plasma levels of zinc, iron, or calcium | Reviewer comments: Study not controlled for potential confounders ITT analysis not performed Source of funding: Study supported by Madaus, SA, and by grants from the Instituto de Salud Carlot III, Red de Centros RCMN (C03/08), and Red de Grupos (G03/140), Madrid Spain. One author had received consulting or lecture fees from Madaus Laboratories and another one belonged to Madaus Laboratory |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|--|--|--|---|---|---|--|---|
| MAFFIA. Treatment of functional constipation with prune-malt. 1955. Archives of Pediatrics 72[10], 341-346 | Study Type: Open label non-RCT Evidence level: 1- Study aim: To evaluate the effectiveness in the treatment of functional constipation in infants and children of a palatable | 3 months to 8 years with functional constipation Exclusion criteria: Organic constipation | 200 children age range: 3 months to 8 years gender not reported Country: USA | Intervention: Prune-Malt ® added to diet -Infants 3 weeks to 1 year old: 2 tablespoonfuls daily added to milk or juice -children 1 to 4 years: 3 tablespoonfuls daily added to milk or food -children 4 to 8 years: 4 tablespoonfuls daily added to milk or food (no changes made in usual diet, no drugs given) Comparison: No intervention | Duration of treatment 3 weeks Assessment point (s): Immediately after treatment completed Follow-up period: No follow-up made after treatment finished Outcome Measures: | Returned to normality (number of children) Prune-Malt ®: 28 Controls: 16 Improved (number of children) Prune-Malt ®: 51 Controls: 25 Not improved (number of children) Prune-Malt ®: 21 Controls: 59 Acceptability (number of parents) Good: 132 Fair: 47 Poor: 21 | Additional information from study: Diagnosis of constipation made on the following: 1) decreases in frequency of stools as compared to the child's usual bowel habits, 2) passage of hard, dry stools Wherever possible, cases of equal severity and ages were equally divided between the 2 groups All mothers given a card to record daily number and description of stools, all associated findings if any and acceptability of Prune Malt by the child Reviewer comments: No sample size calculation performed No comparison made between baseline characteristics No definitions/scoring system given for: "improvement", "no improvement", "return to normality", "good", "fair" and "poor" No dropouts/lost to follow-up children reported Study not controlled for potential confounders Source of funding: |
| | | | | | | | Prune-Malt provided by the Benson- Nuen Laboratories Inc., New York No other details provided |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
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| | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|--|--|---|---|--|-----------------------------|---|---|
| Information | Evidence Level | Patients | Characteristic s | Comparison | Outcome Measures | | |
| Dietary fibre intake and constipation in children with severe developmental disabilities. 2000. Journal of Paediatrics and Child Health 36[3], 236-239Australia. | Prospective case series pilot study) Evidence evel: Study aim: To evaluate ibre intake of severe developmenta ly disabled | 20 children Inclusion criteria: severe developmenta Ily disabled children able to take oral feeding and medically stable Exclusion criteria: Not stated | age range 3 to 17 years gender not reported Country: Hong Kong | Intervention: Fibre supplementation: wheat bran (All Bran ® , Kellogg) added in breakfast -Stage 1: 15 g added to each serving of breakfast (total fibre intake, 17g) -Stage 2: 19 g added to each serving of breakfast (total fibre intake, 21g) Comparison: N.A | supplementatio n:10 days | Number of laxatives per week -at baseline: 1.22 (SD 0.36) -at end of stage 1: 0.9 (SD 0.75) p<0.05 as compared to baseline -at end of stage 2: 0.7 (SD 0.40) p<0.01 as compared to baseline N.S comparing stage 1 and 2 | Additional information from study: Definition of constipation: in the centre where the study was conducted if a child does not have a spontaneous bowel movement for 2 consecutive days a laxative is administered. Those who need more than 1 laxative per week are defined as having constipation Baseline fibre intake around 2g/day Reviewer comments: Unclear who measured study outcomes and how Outcomes for bowel movements not reported in paper Source of funding: Study sponsored by the Society for Relief of Disabled Children, Pokfulam, Hong Kong. 'All Bran' ® sponsored by Kellogg's Asia Ltd Wanchai, Hong Kong |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------------------|---------------------------|----------------------|-----------------------|--|
| Bu et al. | Level Study Type: | 45 children | s 45 children | Intervention: | Measures Duration of | Defecation frequency | Additional information from study: |
| Lactobacillus | double-blind | 40 official | 23 male | MgO 50 mg/kg | treatment: | (times/day) | Chronic constipation defined as a stool |
| casei | RCT | Inclusion | 20 maio | per day, twice a | 4 weeks | -MgO (n=18) | frequency of <3 times/week for >2 |
| rhamnosus | I.O.I | criteria: | | day | | 0.55 ± 0.13 | months and at least 1 of the following |
| Lcr35 in | <u>Evidence</u> | children | Age (months, | day | Assessment | 0.00 ± 0.10 | minor criteria: anal fissures with |
| children with | level: | under 10 | mean, SD) | Comparison 1: | point (s): | -probiotic (n=18) | bleeding due to constipation, faecal |
| chronic | 1+ | years old with | , , , , , , , , , , , , , , , , , , , | Lcr35 8 X 10^8 | <u> </u> | 0.57 ± 0.17 | soiling or passage of large and hard |
| constipation. | | chronic | -MgO group | c.f.u/day | after treatment | 0.07 = 0.11 | stool |
| | Study aim: | constipation | 32.4 ± 13.9 | (Antiobiophilus | completed | -placebo (n=9) | |
| International | to investigate | | | 250 mg, 2 | | 0.37 ± 0.10 | Children randomly assigned into the 3 |
| 49[4], 485-490 | the effect of | Exclusion | -Probiotic group | capsules, twice a | Follow-up | | groups according to a computer - |
| 1 1 | Probiotics | criteria: | 36.7 ± 14.5 | day) | period: | MgO vs. probiotic NS | generated randomisation list |
| | (Lactobacillus | organic | | ,, | | Placebo vs. probiotic | |
| | case | causes of | -Placebo group | Comparison 2: | made after | P=0.006 | Blinding achieved by the use of 3 |
| | rhamnosus, | constipation | 35 ± 14.7 | Placebo (starch in | treatment | MgO vs. placebo | interventions with similar appearances |
| | Lcr35) alone | like | | content) | finished | p=0.01 | and placed into identical capsules, |
| | in the | Hirschsprung' | Country: | | | | which were either swallowed o as a |
| | treatment of | s disease, | Taiwan | | Outcome | Hard stool (%) | whole or opened and the contents of the |
| | chronic | spina bifida | | | Measures: | -MgO (n=18) | capsule administered in milk or fluid |
| | constipation in | | | Lactulose use | | 23.5 ± 7.9 | |
| | children and | hypothyroidis | | (1mL/kg/day) | -frequency of | | Throughout the duration of study all |
| | to compare | m, or other | | | defecation | -probiotic (n=18) | investigators, participants and data |
| | the effect with | metabolic/ren | | stool passage | | 22.4 ± 14.7 | analysts were blinded to the assigned |
| | magnesium | al | | noted for 3 days. | -consistency of | | treatment |
| | oxide (MgO) | abnormalities, | | Glycerin enema | stools | -placebo (n=9) | |
| | and placebo, | drugs | | used only when | | 75.5 ± 6.1 | Sample size determined by doing |
| | respectively | influencing | | no defecation for | -episodes of | | primary trial with 9 patients using non- |
| | | gastrointestin | | >5days or | | | inferiority to test. Equivalent margin |
| | | al function | | abdominal pain | | Placebo vs. probiotic | chosen with reference to effect of active |
| | | other than | | suffered due to | | p=0.02 | control in the data of preliminary trial. |
| | | laxatives | | stool impaction | | MgO vs. placebo | Unbalance design of allocation number |
| | | (calcium | | | | p=0.03 | used for more interest in the new drug |
| | | channel | | | -use of | l | (Lcr35): allocation rate set at 2:2:1. One |
| | | blockers, | | | lactulose or | Abdominal pain | sided significance level set at 0.05 and |
| | | antidysrythmi | | | enema | (times) | power was 80%. Under these |
| | | c agents, | | | | -MgO (n=18) | assumptions the smallest sample size |
| | | anticonvulsiva | | | | 4.8 ± 3.7 | was 45 and the sample size of MgO, |
| | | nts, | | | | | Lcr35 and placebo was 18, 18 and 9 |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--------------------------------|---|
| | Level | antidepressan | 3 | | Weasures | -probiotic (n=18) | respectively |
| | | ts, | | | | 1.9 ± 1.6 | Soperation |
| | | anticholinergi | | | | | No significant differences at baseline |
| | | c agents) | | | | -placebo (n=9) | amongst the 3 group regarding: sex, age |
| | | | | | | 6.7 ± 3.3 | of enrolment, age of onset of |
| | | | | | | MgO vs. probiotic | constipation, duration of constipation, |
| | | | | | | p=0.04 | previous treatment, defecation period, |
| | | | | | | Placebo vs. probiotic p=0.01 | stool consistency, abdominal pain, |
| | | | | | | MgO vs. placebo NS | faecal soiling, bleeding during defecation, use of enema, taking fruit or |
| | | | | | | lvigo vs. piacebo ivo | vegetable daily |
| | | | | | | Use of glycerine | , and the same of |
| | | | | | | enema (times) | Patients asked to discontinue any |
| | | | | | | -MgO (n=18) | laxatives previously prescribed 3 days |
| | | | | | | 1.3 ± 1.9 | before entering protocol, and also asked |
| | | | | | | | to avoid any other probiotics, yogurt or |
| | | | | | | -probiotic (n=18) 1.6 ± 1.9 | beverage containing probiotics for at least 2 weeks before treatment and |
| | | | | | | 1.0 ± 1.9 | during therapy |
| | | | | | | -placebo (n=9) | daming unorapy |
| | | | | | | 4.0 ± 2.1 | All outcomes measures recorded by |
| | | | | | | | parents in a stool diary |
| | | | | | | MgO vs. probiotic NS | |
| | | | | | | Placebo vs. probiotic | 4 patients discontinued medication |
| | | | | | | p=0.04 | during study period: 2 in MgO, 1 in |
| | | | | | | MgO vs. placebo | probiotic, 1 in placebo group (2 patients |
| | | | | | | p=0.03 | suffered from acute gastroenteritis and 2 patients lost to follow-up) |
| | | | | | | No significant | 2 panerio iosi to ionow-up) |
| | | | | | | differences regarding | Reviewer comments: |
| | | | | | | use of lactulose and | Allocation concealment not described |
| | | | | | | faecal soiling | |
| | | | | | | amongst 3 groups | Not clear whether the 2 patients who |
| | | | | | | | suffered from acute gastroenteritis had it |
| | | | | | | Patients with | as consequence of the study medication |
| | | | | | | treatment success | Church a mat a material land for a material land |
| | | | | | | (%) MaO (n=19): 72.2 | Study not controlled for potential |
| | | | | | | -MgO (n=18): 72.2 | confounders |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|--|----------------------------------|
| | | | | | | 77.8 -placebo (n=9): 11.1 MgO vs. probiotic NS | Source of funding: not stated |
| | | | | | | Placebo vs. probiotic p=0.01 MgO vs. placebo p=0.01 no adverse effects noted in probiotic and | |
| | | | | | | placebo groups, only 1 patient in the MgO group suffered from mild diarrhoea | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|------------------------|--------------------|--|
| IIIIOIIIIatioii | Level | ratients | S | Companison | Measures | | |
| Banaszkiewicz | Study Type: | 84 children | 84 children | General: | Duration of | Treatment success | Additional information from study: |
| et al. | Triple-blind | | | Rectal | treatment | <u>(%)</u> | Allocation sequence and randomisation |
| Ineffectiveness | RCT | <u>Inclusion</u> | mean age | disimpaction | 12 weeks | -At 12 weeks: | list computer generated by investigators |
| of Lactobacillus | | criteria: | (months) | with phosphate | | LGG (n=43): 72 | |
| GG as an | Evidence | Children aged | -lactulose + | and saline enema | (from weeks 13 | Placebo (n=41): 68 | Blinding achieved by the use of study |
| adjunct to | level: | 2 to 16 years | LGG group | in all patients | to | N.S | products with similar appearances and |
| lactulose for the | 1+ | with | 79 ± 47 | before study | 24, patients | | tastes, packed identically and |
| treatment of | | constipation | | treatment | instructed to | -At 24 weeks: | indistinguishable from each other. |
| constipation in | Study aim: | defined as < 3 | | | continue the | LGG (n=43): 64 | Throughout duration of study all |
| children: a | To assess the | bowel | placebo group | Intervention: | use of lactulose | Placebo (n=41): 65 | investigators, participants, outcomes |
| double-blind, | effectiveness | movements | 65 ± 36 | Lactulose 70%, 1 | or other | N.S | assessors and data analysts were |
| placebo- | of | per week for | | mL/kg/day (in 2 | laxatives as | | blinded to the assigned treatment |
| controlled | lactobacillus | at least 12 | gender not | divided doses) + | needed | Spontaneous bowel | |
| randomized | rhamnosus | weeks | reported | 10^9 colony | | | No significant differences in baseline |
| trial. 2005. | GG (LGG) as | | | forming units | <u>Assessment</u> | (mean± SD) | characteristics between the 2 groups |
| Journal of | and adjunct to | <u>Exclusion</u> | Country: | (CFU) of | point (s): | -At 4 weeks | |
| Pediatrics | lactulose in | <u>criteria</u> : | Poland | lactobacillus | At 4, 8, 12 | LGG (n=43): | All patients received stool diaries to |
| 146[3], 364-369 | the treatment | Constipation | | rhamnosus GG | | 5.9 ± 2.3 | record frequency of daily bowel |
| | of | caused by | | (LGG) | Follow-up | Placebo (n=41): | movements, faecal soling, straining, |
| | constipation in | | | | period: | 7.7 ± 5.4 | stool consistency as well as any |
| | children | r, anatomic or | | Comparison: | At 24 weeks | N.S | symptoms they consider important (e.g. |
| | | metabolic | | Lactulose 70%, 1 | after study | | abdominal pain, bloating, diarrhoea) |
| | | diseases (as | | mL/kg/day (in 2 | treatment | -At 8 weeks | |
| | | established | | , | finished | LGG (n=43): | Treatment success defined as ≥3 |
| | | by medical | | placebo | | 6.1 ± 2.3 | spontaneous bowel movements per |
| | | history , an | | | <u>Outcome</u> | Placebo (n=41): | week with no episodes of faecal soiling |
| | | abnormal | | | Measures: | 7.2 ± 3.8 | |
| | | thyroid | | | | N.S | 5 children in LGG group discontinued |
| | | hormone level | | | -primary | | intervention (4 clinical improvement, 1 |
| | | or prior | | | outcome: | -At 12 weeks | abdominal pain) vs. 3 patients in |
| | | anorectal | | | treatment | LGG (n=43): | placebo group (2 refused to participate, |
| | | manometry, | | | success | 6.1 ± 1.8 | 1 provided other reason) |
| | | barium or | | | , | Placebo (n=41): | · · · · · · |
| | | ionogram | | | -secondary | 6.8 ± 3.1 | Reviewer comments: |
| | | examination) | | | outcomes: | N.S | Sample size calculation not performed |
| | | | | | number of | Episodes of faecal | Study not controlled for potential |
| | | | | | bowel | soiling per week | confounders |

| Bibliographic Study Type Information Evidence Level | & Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|----------------------|--------------------------------|---------------------------|--|---|--|
| Level | | S | | movements per week number of episodes of faecal soiling per week stool consistency straining frequency per week percentage of patients using laxatives adverse events | (mean± SD) -At 4 weeks LGG (n=43): 0.9 ± 2.1 Placebo (n=41): 0.7 ± 1.5 N.S -At 8 weeks LGG (n=43): 0.8 ± 2.2 Placebo (n=41): 0.3 ± 0.8 N.S -At 12 weeks LGG (n=43): 0.8 ± 1.8 Placebo (n=41): 0.3 ± 0.9 N.S Straining frequency per week (mean± SD) -At 4 weeks LGG (n=43): 1.6 ± 1.9 Placebo (n=41): 1.4 ± 1.9 N.S -At 8 weeks LGG (n=43): 1.4 ± 1.7 Placebo (n=41): 1.4 ± 1.8 N.S | Outcomes for stool consistency not reported Source of funding: Not stated |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|----------------------------|-------------------|
| | | | | | | -At 12 weeks | |
| | | | | | | LGG (n=43): | |
| | | | | | | 1.3 ± 1.5 | |
| | | | | | | Placebo (n=41): | |
| | | | | | | 1.6 ± 1.8 | |
| | | | | | | N.S | |
| | | | | | | Patients using | |
| | | | | | | laxatives (%) | |
| | | | | | | -At 24 weeks: | |
| | | | | | | LGG (n=43): 44 | |
| | | | | | | Placebo (n=41): 43 | |
| | | | | | | N.S | |
| | | | | | | Advarage offeets (0/ | |
| | | | | | | Adverse effects (% | |
| | | | | | | patients) LGG (n=43): 9 | |
| | | | | | | Placebo (n=41): 14.6 | |
| | | | | | | N.S | |
| | | | | | | N.S | |
| | | | | | | LGG well tolerated. | |
| | | | | | | Side effects profile of | |
| | | | | | | LGG similar to that of | |
| | | | | | | placebo: 3 patients in | |
| | | | | | | LGG group vs. 5 | |
| | | | | | | patients in placebo | |
| | | | | | | group developed | |
| | | | | | | abdominal pain. 1 | |
| | | | | | | patients in LGG | |
| | | | | | | group developed | |
| | | | | | | vomiting and 1 in the | |
| | | | | | | placebo group | |
| | | | | | | experienced | |
| | | | | | 1 | headache | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|----------------------|---|
| | Level | | S | | Measures | | |
| Bekkali et al. | Study Type: | 20 children | 20 children | General: | Duration of | Frequency of bowel | Additional information from study: |
| The role of a | Prospective | | | Disimpaction: | treatment | movements (BMs) | Constipation defined by Rome III criteria |
| probiotics | case series | <u>Inclusion</u> | 10 boys | rectal enema | 4 weeks | per week, total | as having at least 2 out of 6 of the |
| mixture in the | (pilot study) | | Median age: 8 | (Klyx: sodium- | | sample | following |
| treatment of | | Children | years (4 to 16) | | Assessment | -Baseline: | symptoms: bowel movements <3 |
| childhood | Evidence | between 4 to | | ate and sorbitol) | point (s): | 2.0 (1.0 to 5.0) | times/week; faecal incontinence >2 |
| constipation: a | level: | 16 years of | Country: | once daily for 3 | At 2 and 4 | | times/week; large amounts of stools |
| pilot study. | 3 | 3 | The | days | weeks | -Week 2: | obstructing the toilet once in 10 days; |
| 2007. Nutrition | | | Netherlands | | | 4.2 (0.0 to 16.0) | painful defecation; withholding |
| Journal 6, 17 | Study aim: | clinic with | | Intervention: | Follow-up | p = 0.10 | behaviour; palpable abdominal or rectal |
| | to determine | constipation | | Daily probiotics | period: | | mass on physical examination |
| | the | | | mixture of | No follow-up | -Week 4: | |
| | therapeutic | <u>Exclusion</u> | | 4 × 109 colony | conducted after | and 3.8 (2.1 to 7.0) | 7 days prior to baseline assessment and |
| | effect of a | criteria: | | forming units | treatment | p = 0.13 | during treatment period all children |
| | combination | use of any | | \ // | finished | | recorded frequency of bowel |
| | of probiotics | oral laxative < | | Bifidobacteria | | Frequency of bowel | movements, number of faecal |
| | strains, | 4 weeks | | (B.) bifidum, B. | <u>Outcome</u> | movements (BMs) | incontinence episodes, stool |
| | containing the | before intake, | | infantis, B. | Measures: | per week in 12 | consistency, abdominal pain, flatulence |
| | bifidobacteria | mental | | longum, | | children presenting | and pain during defecation as well as |
| | B. bifidus, B. | retardation, | | Lactobacilli (L.) | Primary | with <3 BMs per | adverse effects such as vomiting and |
| | infantis and B. | | | casei, L. | outcomes: | week at baseline: | diarrhoea in a standardized bowel diary |
| | longum and | disease, | | plantarum and L. | | -Baseline: | |
| | the lactobacilli | functional | | rhamnosus | -frequency of | 1.0 (0.0 to 2.0) | Stool consistency rated by patients as |
| | L. casei, L. | non-retentive | | | bowel | | hard, normal or watery |
| | plantarum and | incontinence, | | Comparison: | movements per | -Week 2: | |
| | L. rhamnosus, | and a history | | N.A | week | 3.0 (0.0 to 7.0) | During treatment period children |
| | on childhood | of gastro- | | | | p = 0.01 | instructed to start toilet training. Toilet |
| | constipation | intestinal | | | -stool | | training consisted of sitting on the toilet |
| | | surgery | | | consistency | -Week 4: | 3 times per day for 5 minutes after each |
| | | | | | | 3.0 (0.0 to 10.0) | meal with the intention of trying to |
| | | | | | Secondary | p = 0.009 | defecate. Use of laxatives not allowed |
| | | | | | outcomes: | | during treatment period |
| | | | | | | Stool consistency | · . |
| | | | | | -number of | Hard stools (n | Reviewer comments: |
| | | | | | faecal | children): | No dropouts/lost to follow-up children |
| | | | | | incontinence | -Baseline: 7 | were reported |
| | | | | | episodes per | | |
| | | | | | week | -Week 2 : 4 | Source of funding: |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| Bibliographic Information | Evidence | | Characteristic | | Outcome | p = 0.23 -Week 4: 6 p = 1.00 At week 4, hard stools appeared in 5 children who also had hard stools at baseline. 1 child with normal stools at baseline, reported hard stools only at the end of the study. 2 of the 7 children who presented with hard stools, reported normal stools at the end of the study Number of faecal incontinence episodes per week Baseline: 4.0 (0.0 to 35.0) | Not stated |
| | | | | | | Week 2: 1.5 (0.0 to 14.0) p = 0.007 Week 4: 0.3 (0.0 to 7.0) p = 0.001 <u>Side effects</u> | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | effects such as vomiting, bloating and increased flatulence during the study period | |
| | | | | | | | |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|---------------------|--|
| | Level | | S | | Measures | | |
| Young et al. | Study Type: | 108 children | 90 children | Intervention: | Duration of | Stool frequency | Additional information from study: |
| Increasing oral | Open label | | | Increased water | <u>treatment</u> | (mean) | Constipation Assessment Score based |
| fluids in chronic | RCT | <u>Inclusion</u> | 31 boys | | 2 weeks | H2O (water) | on 8 variables assessed during the past |
| constipation in | | criteria: | (47.46%) | instructed to | | HiOsm (high | 3 days: abdominal distension or |
| children. 1998. | <u>Evidence</u> | Prepubertal | | increase water | <u>Assessment</u> | osmolality) | bloating, change in amount of gas |
| Gastroenterolog | <u>level:</u> | children with | mean age 7.5 | intake by 50% on | point (s): | | passed rectally, less frequent bowel |
| y Nursing 21[4], | 1- | moderate to | years (range | the basis of total | At week 2 and 3 | | movements, oozing liquid stools, rectal |
| 156-161 | | severe simple | | measured oral | | Control: 3.45 | fullness or pressure, rectal pain with |
| | | constipation | years) | liquid intake | | H2O: 3.52 | bowel movement, smaller stool size, |
| | | as assessed | | during1st baseline | | HiOsm: 3.75 | urge but inability to pass stool. Each |
| | whether or not | | Country: | week | No follow-up | | variable scored as 0, no problem; 1, |
| | increasing | Constipation | USA | | made after | -week 2: | some problem and 2, severe problem. |
| | | Assessment | | Comparison 1: | | Control: 4.05 | |
| | by either | Score | | 71 | finished | H2O: 3.57 | A gift certificate to a toy store was used |
| | excess water | | | liquids: group | | HiOsm: 4.31 | as incentive to return data collection |
| | | Exclusion | | administered | <u>Outcome</u> | | forms |
| | excess | <u>criteria</u> : | | supplemental | Measures: | -week 3: | |
| | , · | Post pubertal | | liquid in the form | | Control: 3.40 | The concentration of 600 mOsm/L |
| | • | children, | | of Kool-Aid, juice, | -stool frequency | H2O: 3.70 | chosen because it was considered to be |
| | | hypercalcemi | | soda pop or other | | HiOsm: 3.44 | a level above which a significant osmotic |
| | significantly | a, | | liquids know to | -stool | _ | load in the small bowel would result in |
| | | Hirschsprung' | | contain more than | consistency | Stool consistency | significant plasma to lumen flux. The |
| | | s disease, | | 600 mOsm/L | | (mean) | 50% increase arbitrarily chosen as being |
| | | hypothyroidis | | | -difficulty of | -baseline: | feasible , >50% considered potentially |
| | | m, cardiac or | | Comparison 2: | stool passage | Control: 6.30 | burdensome for children/caregiver and |
| | children | renal | | Control group: no | | H2O: 6.13 | probably not therapeutically obtainable |
| | | disorders, | | intervention | | | under normal situations |
| | | children | | | | -week 2: | |
| | | receiving | | | | Control: 6.33 | Stool frequency, consistency and |
| | | specialised | | | | H2O: 5.99 | difficulty with passage assessed daily by |
| | | diets, | | | | | parents using a simple form. The Stool |
| | | malnourished | | | | -week 3: | Consistency Continuum previously |
| | | children | | | | Control: 6.30 | developed by Bergstrom chosen to |
| | | already | | | | H2O: 5.79 | evaluate stool form. Difficulty of passage |
| | | receiving | | | | Difficulty of stool | scored as: 0, no problem; 1, some |
| | | stool | | | | Difficulty of stool | problem; 2 severe problem |
| | | softeners or | | | | passage (mean) | |
| | | laxative | | | | -baseline: | A second round of analysis excluded all |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|------------------------------|--------------------------------|---------------------------|------------------------------------|--------------------------------------|--|
| | | preparations, | | | | Control: 0.96 | subjects who failed to comply with at |
| | | children who | | | | H2O: 0.78 | least 75% of assigned intervention, and |
| | | were | | | | HiOsm: 0.77 | this did not change the study outcomes |
| | | physically or intellectually | | | | -week 2: | Reviewer comments: |
| | | challenging | | | | Control: 0.95 | Sample size calculated on the basis of |
| | | (?) or who | | | | H2O: 0.84 | preliminary power analysis but no details |
| | | had an | | | | HiOsm: 0.74 | provided. Non probability convenience |
| | | underlying | | | | | sample was used |
| | | central | | | | -week 3: | |
| | | nervous | | | | | No comparison made of baseline |
| | | system | | | | H2O: 0.87 | characteristics |
| | | disease | | | | HiOsm: 0.62 | |
| | | | | | | | Methods of randomisation and allocation |
| | | | | | | | concealment not described |
| | | | | | | water intake nor | |
| | | | | | | increasing | 108 children originally included, but only |
| | | | | | | | 90 completed the entire study as |
| | | | | | | | assigned. 18 children failed to comply |
| | | | | | | | with 75% of the intervention, but there |
| | | | | | | | are no clear explanations as to why that |
| | | | | | | | happened |
| | | | | | | consistency or | |
| | | | | | | _ | Outcomes for stool consistency in the |
| | | | | | | | HiOsm group not reported |
| | | | | | | groups when | Ctudy not controlled for notantic! |
| | | | | | | | Study not controlled for potential |
| | | | | | | made with previous weeks, or between | |
| | | | | | | , | |
| | | | | | | the 3 groups during | |
| | | | | | | the same week | Source of funding: Not clearly stated |
| | 1 | 1 | 1 | | | i | INULUICAIIY SLALCU |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|----------------------|---|
| 0 | Level | 00 131 | S | | Measures | | |
| Staiano et al. | | 20 children | 20 children | General: | Duration of | Number of stools per | Additional information from study: |
| Effect of the | Case series | | | Disimpaction with | treatment | week (mean ± SD) | Children fed by mouth with semi-liquid |
| dietary fiber | F · · | Inclusion | 14 boys | enemas for 2 or 3 | 12 weeks | -at 4 weeks | diet including formula and pureed food |
| glucomannan | <u>Evidence</u> | criteria: | mean age 5.7 ± | days (not clear | | Glucomannan (n=9): | |
| on chronic | | Severe | 4.2 years | what medication | Assessment | 4.0 ± 1.3 | No significant differences in baseline |
| constipation in | 3 | neurologic | | used) | point (s): | Placebo (n=10): | characteristics between 2 groups |
| neurologically | | damage, | Country: | | At 4, 8 and 12 | 1.1 ± 0.2 | |
| impaired | | constipation | Italy | Intervention: | weeks | | 1 patient receiving glucomannan |
| children. 2000. | | of at least 12 | | Glucomannan | | -at 8 weeks | withdrawn from study after 3 weeks of |
| Journal of | the efficacy of | | | | Follow-up | Glucomannan (n=9): | treatment because of concomitant |
| Pediatrics | | most patients | | a day | period: | 3.3 ± 1.0 | increase in seizure frequency |
| 136[1], 41-45 | as a treatment | | | | No follow-up | Placebo (n=10): | associated with blood level of |
| | | not possible | | Comparison: | | 2.5 ± 1.2 | Phenobarbital below the therapeutic |
| | constipation in | | | Placebo | treatment | | range |
| | | enema. All | | 100mg/kg 2 times | finished | -at 12 weeks | |
| | severe | patients had | | a day | | Glucomannan (n=9): | During study period a daily diary card |
| | | severe | | | <u>Outcome</u> | 3.8 ± 0.9 | was completed for recording symptoms, |
| | damage | /profound | | | Measures: | Placebo (n=10): | dietary fibre intake, number of bowel |
| | | mental | | Both | | 2.0 ± 0.6 | movements per week, stool consistency, |
| | | retardation | | glucomannan and | Stool frequency | | presence of painful defecation and use |
| | | (IQ level < 35) | | placebo consisted | | p<0.01 for | of laxative (lactulose 1g/kg/dose) or |
| | | and exhibited | | of a 500-mg | Stool | glucomannan group | glycerol suppository. Arbitrary scoring |
| | | severe clinical | | capsule. Oral | consistency | at all periods as | system used for assessment of |
| | | manifestation | | dose given by | _ | compared to baseline | symptoms: |
| | | s of brain | | mixing the | Presence of | | -stool consistency: 1, pellets; 2, hard; 3, |
| | | damage | | contents of one | painful | Stool consistency | soft; 4, loose; 5, liquid |
| | | etiologically | | capsule with 100 | defecation | score (mean ± SD) | -presence of painful defecation: 1, often; |
| | | related to | | mL of water | | -at 4 weeks | 2, occasionally; 3, none |
| | | prenatal or | | | Laxative use | Glucomannan (n=9): | , , , |
| | | perinatal | | | | 2.4 ± 0.5 | Reviewer comments |
| | | hypoxia: 12 | | | | Placebo (n=10): | No definition of constipation given |
| | | patients had | | | | 1.3 ± 0.6 | g |
| | | classical | | | | | Very small sample size. Sample size |
| | | tetraplegia, 6 | | | | -at 8 weeks | calculation not performed |
| | | severe | | | | Glucomannan (n=9): | F |
| | | spastic | | | | 2.8 ± 0.7 | Randomisation and allocation |
| | | diplegia, 2 | | | | Placebo (n=10): | concealment |
| | | persistent | | | | 1.3 ± 0.5 | methods not described |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|--|--------------------------------|---------------------------|------------------------------------|---|---|
| | Level | hypotonia Exclusion criteria: unclear | S | | Measures | -at 12 weeks Glucomannan (n=9): 2.7 ± 0.7 Placebo (n=10): 1.4 ± 0.7 p<0.01 for glucomannan group at all periods as compared to baseline Painful defecation score(mean ± SD) -at 4 weeks Glucomannan (n=9): 1.4 ± 1.1 (N.S as compared to baseline) Placebo (n=10): 0.9 ± 0.8 -at 8 weeks | Blinding procedures poorly described Unclear who measured study outcomes Study not controlled for potential confounders Source of funding: One of the authors supported by a grant from Dicofarm, Italy. No other details provided |
| | | | | | | Glucomannan (n=9): 1.7 ± 1.4 (N.S as compared to baseline) Placebo (n=10): 1.2 ± 0.8 -at 12 weeks Glucomannan (n=9): 1.9 ± 1.2 Placebo (n=10): 1.2 ± 0.9 p<0.01 for glucomannan group as compared to | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---------------------------------------|-------------------|
| | | | | | | baseline | |
| | | | | | | Laxative use (number | |
| | | | | | | per week, (mean ± SD) | |
| | | | | | | -at 4 weeks | |
| | | | | | | Glucomannan (n=9): 0.3 ± 0.8 | |
| | | | | | | Placebo (n=10): 2.0 ± 0.6 | |
| | | | | | | | |
| | | | | | | p<0.01 for glucomannan group | |
| | | | | | | as compared to | |
| | | | | | | baseline | |
| | | | | | | -at 8 weeks Glucomannan (n=9): | |
| | | | | | | 0.5 ± 0.8 (N.S as | |
| | | | | | | compared to baseline) | |
| | | | | | | Placebo (n=10): | |
| | | | | | | 1.8 ± 1.6 | |
| | | | | | | -at 12 weeks Glucomannan (n=9): | |
| | | | | | | 0.3 ± 0.5 | |
| | | | | | | Placebo (n=10): 2.1 ± 0.4 | |
| | | | | | | | |
| | | | | | | p<0.01 for glucomannan group | |
| | | | | | | as compared to | |
| | | | | | | baseline | |
| | | | | | | All outcomes for placebo group at all | |
| | | | | | | points were N.S as | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|----------------------|-------------------|
| | | | | | | compared to baseline | |
| | | | | | | | |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|----------------------|---------------------------|---------------------------|------------------------|-----------------------|--|
| | Level | | s | | Measures | | |
| Eisenberg et al. | Study Type: | 22 children | 22 children | Intervention: | Duration of | Prevalence of | Additional information from study: |
| Contribution of | Non-RCT | | | Trial of David Hart | | constipation (number, | Intervention and control children |
| stepping while | | <u>Inclusion</u> | Intervention | Walker (HW) | 6 months | % of children) | matched for age and sex |
| standing to | <u>Evidence</u> | criteria: | group (n=11): | device (to | | | |
| function and | level: | Aged | 6 males | | <u>Assessment</u> | -At entry: | HW device – The David Hart Walker |
| secondary | 1- | | mean age (yr) | stepping while | point (s): | | (HW) Orthosis, a hands free walker |
| conditions | | | 6.1±2.1 | standing) in | at 6 months | HW: 6 (54.5) | provides weight-bearing support and leg |
| | Study aim: | at first visit, | | addition to | after treatment | SF: 6 (54.5) | alignment while allowing upper extremity |
| with cerebral | To explore the | | Controls (n=11) | physical therapy | initiated | | freedom, aiming to allow the action of |
| palsy. 2009. | feasibility and | | : | sessions. | | NS | stepping while standing |
| Pediatric | efficacy of | | 6 males | Beginning with 30 | | | |
| Physical | stepping while | | mean age (yr) | minute sessions 4 | | -at 6 months: | Constipation defined as 2 bowel |
| Therapy 21[1], | | function | 6.7±1.6 | times a week, | None after | | movements per week, or 2 of the |
| 79-85 | | classification | _ | parents and | intervention | HW: 1 (9.1) | following on more than 1 of 4 occasions: |
| Einsberg et al. | | system | Country: | | period finished | SF: 6 (54.5) | straining, hard stools and a feeling of |
| 2009 | | (GMFCS) | Israel | encouraged to | | | incomplete evacuation |
| | prevalence of | level 4 or 5, | | use device at | <u>Outcome</u> | p = 0.02 | |
| | secondary | inability to | | home | Measures: | | Diary of bowel function kept by parent |
| | conditions | stand and | | | Prevalence of | | and/or the physical therapist and |
| | among | walk with | | Comparison: | constipation | | maintained throughout follow-up period |
| | | traditional | | Program in | | | used to assess for constipation |
| | severe | walker/rollator | | standing frame | | | |
| | | due to | | (SF) (passive | | | At baseline children in the HW group |
| | (CP) | insufficient | | standing) as part | | | had higher significant mean scores in |
| | | upper | | of physical | | | the self-care and social function domain |
| | | extremity | | therapy session. | | | of the Paediatric Evaluation of Disability |
| | | control, | | 30-minute | | | Inventory (PEDI) score than children in |
| | | attempts | | sessions 4 times | | | the SF group |
| | | steps when in | | a week, parents | | | Daviewer comments |
| | | a supported | | and children | | | Reviewer comments |
| | | standing | | encouraged to | | | Vancarall atual manufation |
| | | position, flexion | | use SF at home | | | Very small study population |
| | | contractures | | | | | No dropouts/loss to follow-up reported |
| | | of the hips | | | | | ino diopodis/ioss to follow-up reported |
| | | and knees of | | | | | PEDI scores may have confounded the |
| | | less than 30° | | | | | effects of the intervention and this was |
| | | 1699 111911 90 | | | | | not accounted for |
| | | | | | | | HOL ACCOUNTED TO |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|--------------------------------------|--------------------------------|------------------------------|------------------------------------|-------------|----------------------------------|
| | | Exclusion criteria: Not stated | | | | | Source of funding: Not stated |
| | | Not stated | | | | | Not Stated |
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Effectiveness of excluding Cow's Milk from the Diet in Children with Chronic Idiopathic Constipation

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--------------------|--|
| lacono et al. | Study Type: | 65 patients | Age (mo) | Intervention: | Follow-up | Observation period | The order of treatment was randomly |
| Intolerance of | Cross over | | 34.6+-17.1 | Excluding cow's | period: Mean: | <u>(n=65)</u> | assigned by a computer-generated |
| cow's milk and | randomised | 33 patients | | milk and its | 10 months | | method with the individual patient as the |
| chronic | controlled trial | | Sex M/F 29/36 | derivatives from | | Number of bowel | unit of randomisation. The researchers |
| constipation in | | cow's milk | | the diet of children | | movements: 4 | were unaware of the order of the |
| children. 1998. | <u>Evidence</u> | and 32 soy | | with constipation | | Median: 3-5 | treatment. |
| New England | level: 1+ | milk during | | | | 25th to 75th | |
| Journal of | | the fist study | | Comparison: | | percentile | At baseline and end of two study periods |
| Medicine | | period | | Cow's milk vs. soy | bowels | | children were examined by a researcher |
| 339[16], 1100- | | | | milk | | Qualitative faecal | who was unaware of laboratory test results |
| 1104United | | 32 patients | | | | score | and histological findings |
| States. | | received | | Weeks 1-2: | | 1: 0 | |
| | | cow's milk | | observation | | 2: 0 | To ensure that children did not receive any |
| | | and 33 soy | | period | | 3: 65 | other kind of milk-containing food during |
| | | milk during | | all medication | during a | | the study periods parents were given a list |
| | | the second | | stopped | | Weeks 3-4 and 6-7 | of most common milk-containing food to |
| | | study period | | | period were | -Cow's milk group: | be avoided |
| | | | | Weeks 3-4: one | considered to | | |
| | | <u>Inclusion</u> | | group received | | Number of bowel | 6 patients were withdrawn from the study |
| | | <u>criteria:</u> | | cow's milk and | | movements: | during the cow's-milk study period (on |
| | | consecutive | | unrestricted diet | | Median: 4 | days 9-12) because of the reappearance |
| | | children | | and the other had | | 25th to 75th | of constipation and other related disorders. |
| | | referred by | | cow's milk and its | | percentile: 3-5 | For these children the number of bowel |
| | | family | | derivatives | 1: mushy or | | movements per period was prorated. |
| | | paediatricians | | excluded from diet | | Qualitative faecal | Intention to treat analysis was used |
| | | to a paediatric | | and received soy | | score | |
| | | gastroenterol | | milk instead | | 1: 0 | The mean (±SD) daily consumption was |
| | | ogy clinic | | | | 2: 0 | 450±120 ml of soy milk and 470±135 ml of |
| | | diagnosed | | Week 5: washout | | 3: 65 | cow's milk. Analysis of the main |
| | | with chronic | | period for both | and difficulty | | constituents of the diet (proteins, |
| | | constipation. | | groups, | and pain on | -Soy milk group: | carbohydrates and fibres) did not show |
| | | Chronic | | unrestricted diet | passing stools | | any qualitative or quantitative variation |
| | | constipation | | and intake of soy | | Number of bowel | during the study period (data not shown). |
| | | defined as | | or cow's milk and | Both Number of | | |
| | | chronic faecal | | its derivatives | | Median: 10 | Patients were highly selected and this |
| | | retention (one | | | movements and | 25th to 75th | might have led to overestimate the |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|--------------------------|--------------------------------|---------------------------|------------------------------------|-------------------------|--|
| | | bowel | | Week 6-7: | qualitative | percentile: 4-12 | frequency of cow's milk intolerance as a |
| | | movement | | patients switched | faecal score | | cause of constipation. Paediatricians who |
| | | every 3 to 15 | | to the other type | were recorded | Qualitative faecal | referred the patients may have preselected |
| | | days) often | | of milk | by parents | score | them having being the centre where the |
| | | associated | | | | 1: 2 | study was conducted experience in the |
| | | with | | Total amount of | | 2: 42 | treatment of food allergies. The inclusion |
| | | abdominal | | milk given to the | | 3: 21 | of patients with no response to laxatives |
| | | symptoms | | patient during the | | | may have also contributed to this issue. |
| | | (abdominal | | two weeks: 5-10 | | p values were < | |
| | | pain, painful defecation | | litres | | 0.001 for all variables | |
| | | and so forth) | | Bottles coded A or | | Challenge with cow's | |
| | | , | | B by hospital | | milk (n=44) | |
| | | Exclusion | | dispensary | | | |
| | | criteria: | | Infants < 15 | | -Placebo group (soy | |
| | | Anatomical | | months age: | | milk): 0 clinical | |
| | | causes | | formula based on | | reactions | |
| | | (Hirschsprung | | cow's milk | | | |
| | | 's disease, | | (Transilat, | | -Cow's milk group: 0 | |
| | | spinal | | Plasmon, Milan, | | acute reaction, but in | |
| | | disease) | | Italy) or formula | | all patients | |
| | | another | | based on soy | | constipation | |
| | | disorder | | (Plasmonsoy, | | associated with hard | |
| | | (hypothyroidis | | Plasmon). | | stools and discomfort | |
| | | m, | | Children > 15 | | on defecation | |
| | | psychomotor | | months age: | | reappeared after 5-10 | |
| | | retardation), | | commercially | | days on the diet. | |
| | | prior anal | | available whole | | Cow's-milk-free diet | |
| | | surgery, | | cow's milk or soy | | was recommenced, | |
| | | medication | | milk | | with a consequent | |
| | | that can | | | | normalisation of | |
| | | cause | | After the two | | bowel movements in | |
| | | constipation | | study periods | | all patients | |
| | | (chlorpromazi | | children with a | | | |
| | | ne) and | | response to | | Follow-up: | |
| | | referral for | | cow's-milk free | | | |
| | | other reasons | | diet were given | | 0 children with | |
| | | | | the soy-milk diet | | response had | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|--|------------------------------------|--|-------------------|
| | | | | for another month and then underwent a 2-week double-blind challenge with cow's milk at hospital. Children were randomly assigned to receive cow's milk or a placebo containing soy milk. If no clinical reactions were observed within 12 hours, patients were discharged and the challenge continued at home with bottles coded A or B by the hospital dispensary. Challenge was stopped when a clinical reaction occurred, in particular when there were not bowel movements for 72 hours and the patient had abdominal pain, perianal lessons or both. | | constipation Cow's milk reintroduced into the diets of 15 children after 8-12 of cow's milk-free diet and in all cases constipation returned within 5-10 days Children with no response to soy-milk diet were treated with high doses of laxatives, with subsequent improvement in stool frequency. In all cases symptoms returned once treatment with laxatives was stopped | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|----------------------------------|---------------------|---------------------|--|
| | Level | | s | • | Measures | | |
| Carroccio et al. | Study Type: | 52 | Age (months): | Intervention: | Follow-up | Observation period: | Qualitative faecal score previously |
| Chronic | Case series | consecutive | 51.2±18 | Cow's milk-free | period: | | validated according to authors |
| constipation | and | infants and | | diet, with the | None reported | -Patients with food | |
| and food | embedded | children with | Sex (M/F): | exclusion of cow's | | intolerance (n=30) | Randomisation method used during the |
| intolerance: A | randomised | chronic | 22/30 | milk and all its | <u>Outcome</u> | | cow's milk challenge not described |
| model of | controlled | constipation | | derivatives | Measures: | Number of bowel | |
| proctitis causing | challenge | unresponsive | | | Number of | movements/week: | To ensure that children did not receive any |
| constipation. | | to previous | | Comparisons: | bowels | Median: 1.5 | other kind of milk-containing food during |
| 2005. | <u>Evidence</u> | treatments | | 1. Cow's milk-free | movements/we | 25th to 75th | the study periods parents were given a list |
| Scandinavian | <u>level</u> : | examined at | | diet vs. soy milk | ek | percentile: 1-2 | of most common milk-containing food to |
| Journal of | 3 | the | | Cow's milk vs. | | | be avoided. Furthermore, they were asked |
| Gastroenterolog | | outpatients | | ass's milk | Qualitative | Qualitative faecal | to record the amount and type of food their |
| y 40[1], 33- | | clinic of a | | | faecal score | score | children had eaten every day. Frequent |
| 42Norway. | | hospital. | | | , | 1: 0 | telephone contacts helped to ensure |
| | | Chronic | | 1. Cow's milk-free | liquid stool | 2: 0 | adherence to the diet |
| | | constipation | | diet vs. soy milk | 2: soft faeces | 3: 30 | |
| | | defined as | | | and no pain in | | Patients with chronic constipation caused |
| | | chronic faecal | | -2 weeks | passing stools | - Patients with | by food intolerance showed at baseline a |
| | | retention (one | | observation | 3: hard faeces | constipation | higher frequency of a personal history of |
| | | bowel | | period: all | and difficulty | unrelated to food | previous food intolerance (p=0.02) and |
| | | movement | | medications | and pain on | intolerance (n=22): | concomitant signs of food intolerance |
| | | every 3 days | | stopped | passing stools | | (bronchospasm five cases, rhinitis four |
| | | or more) with | | and at the end a | | Number of bowel | cases, dermatitis two cases) (p=0.03) than |
| | | painful | | clean-out with | both number of | movements/week: | patients with constipation unrelated to food |
| | | elimination of | | single dose of | bowels | Median: 1.5 | intolerance. |
| | | hard stools | | polyethylene | | 25th to 75th | No difference was observed between the |
| | | associated | | glycol 4000 | ek | percentile: 1-2 | 24 patients with CM intolerance and the 6 |
| | | with | | (0.75g/kg). | and qualitative | | patients with multiple food intolerance for |
| | | abdominal | | Normal diet, no | faecal score | Qualitative faecal | outcome measures considered (number of |
| | | pain | | restrictions | were recorded | score | bowel movements and qualitative faecal |
| | | l | | | by parents | 1: 0 | score), either at baseline or on elimination |
| | | <u>Inclusion</u> | | -4 weeks of cow's | | 2: 0 | diet. However in comparison with patients |
| | | criteria: | | milk free diet | Children with | 3: 22 | intolerant to CM alone, patients suffering |
| | | -a history of | | (without cow's | eight or more | | from multiple food intolerance were older |
| | | chronic | | milk derivatives | bowel | Elimination diet | (p=0/04) and had a higher frequency of |
| | | constipation | | too) | movements | period: | family history of atopic disease (p=0.03) |
| | | lasting at | | | during a | | |
| | | least 6 | | Infants < 15 | treatment | -Patients with food | Analysis of the main constituents of the |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|-----------------------|--|
| | Level | months | 5 | months age: soy- | period were | intolerance (n=30) | diet (proteins, carbohydrates and fibres) |
| | | -lack of | | based (Nutrilon- | considered to | , , | did not show any qualitative or quantitative |
| | | response to a | | soya, Nutricia, | have a | Number of bowel | variation during the study period (data not |
| | | previous | | Milan, Italy) | response | movements/week: | shown) |
| | | increase in | | | | Median: 5 | , |
| | | dietary fibre | | Children > 15 | Normalised | 25th to 75th | Patients with food intolerance (to CM only |
| | | intake and/or | | months age: | stools habits: | percentile: 4-7 | or multiple) were treated as a group for the |
| | | to laxative | | commercially | bowel | | purpose of analysing the data, therefore it |
| | | treatment | | available soy milk | frequency of at | Qualitative faecal | is not possible to offer specific data for the |
| | | (milk of | | | least five | score | CM group only |
| | | magnesia 1-2 | | Patients | evacuations/we | | |
| | | ml per kg | | unresponsive to | ek with the | 2: 28 | The high frequency of chronic constipation |
| | | bodyweight, | | CM-free diet | | 3: 0 | owing to food intolerance likely due to a |
| | | polyethylene | | placed on | soft stools, | | selection bias, as mainly food-intolerant |
| | | glycol 4000 | | oligoantigenic diet | without painful | - Patients with | patients are treated at the centre where |
| | | mean dose | | 4 weeks (also | defecation | constipation | study was conducted. |
| | | 0.75 g per kg | | excluding cow's | | unrelated to food | |
| | | daily) | | milk): exclusively | | intolerance (n=22): | Funding source: partly supported by a |
| | | attempted for | | rice, lamb, | | | grant from MURST and from the MiPAF |
| | | at least one | | carrots, ass's | | Number of bowel | (progetto "ALICE", D.D. n 86 dated |
| | | month | | milk, olive oil and | | movements/week: | 30.01.2002) |
| | | -regular | | sugar | | Median: 1.5 | |
| | | dietary intake | | | | 25th to 75th | |
| | | of cow's milk | | 2. Cow's milk vs. | | percentile: 1-2 | |
| | | and | | ass's milk | | | |
| | | derivatives | | | | Qualitative faecal | |
| | | <u>Exclusion</u> | | Double-blind | | score | |
| | | criteria: | | placebo-controlled | | 1: 0 | |
| | | -prior anal | | challenge with | | 2: 0 | |
| | | surgery | | cow's milk, after | | 3: 22 | |
| | | -use of | | 12 weeks, to all | | | |
| | | medication | | patients cured on | | Cow's milk challenge: | |
| | | that can | | CM-free or | | No specific data are | |
| | | cause | | oligoantigenic | | reported apart from | |
| | | constipation | | diet. | | saying that in all | |
| | | -referral for | | Placebo: ass's | | cases cow's milk | |
| | | reasons other | | milk | | readministration | |
| | | than chronic | | If no clinical | | caused the | |

| udy Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------------|---|--------------------------------|---|------------------------------------|---|-------------------|
| | constipation -anatomical /neurological causes of constipation (Hirschsprung 's disease, spinal disease, psychomotor retardation) -another disease causing constipation (hypothyroidis m, coeliac disease) | | reactions after 12 hours, patients were discharged and challenge continued at home with bottles coded A or B. Challenge was stopped when a clinical reaction occurred | | reappearance of constipation within 5 days after commencing the challenge (median 2 days, range 1-5 days) | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---|------------------------|---|
| | Level | | S | | Measures | | |
| lacono et al. | Study Type: | 27 infants | 15 boys | Intervention: | Follow-up | Mean number (+-SD) | Analysis of the patient's dietary diaries did |
| Chronic | Case series | | | Excluding cow's | | of stools per day | not show any significant variations in daily |
| constipation as | | <u>Inclusion</u> | | milk and its | for a mean | during unrestricted | fibre and liquid intake during the various |
| a symptom of | Evidence | criteria: | | derivatives from | period of 18 | | study periods |
| cow milk | level: 3 | referred to a | | the diet of children | | CMP-free diet | |
| allergy. 1995. | | paediatric | months) | with chronic | 10 to 30 | | It is not reported whether any medication |
| Journal of | | gastroenterol | | constipation | months) | Stools from patients | was stopped at the beginning of the study |
| Pediatrics | | ogy clinic | | | | on CMP-free diet | |
| 126[1], 34-39 | | during the 12 | | Comparisons: | <u>Outcome</u> | -Cured (n=21) | Funding: not reported |
| | | months | | 1. Cow's milk-free | Measures: | | |
| | | preceding the | | diet vs. soy milk/ | | a. UD: 0.24+-0.10 | |
| | | study and | | ass's milk | | b. 1rst CMP-free diet: | |
| | | considered to | | 2. Cow's milk vs. | | 1.04+-0.12 | |
| | | have | | ass's milk | | c. 1rst CMP | |
| | | idiopathic | | | | challenge: 031+-0.14 | |
| | | constipation. | | 1. Cow's milk-free | | d. 2nd CMP-free diet: | |
| | | Diagnosis of | | diet vs. soy milk/ | - · · · · · · · · · · · · · · · · · · · | 1.05+-0.11 | |
| | | constipation | | ass's milk | passing them = | | |
| | | made on the | | | | Significance: b and d | |
| | | basis of a | | -First 7 days: All | score | vs. a and c, p<0.0005 | |
| | | history of | | patients were | | | |
| | | reduced | | being fed the | Qualitative | -Unimproved (n=6) | |
| | | frequency of | | same diet as at | score: | | |
| | | stools (one | | the time of | | UD: 0.18+-0.12 | |
| | | evacuation | | diagnosis: various | | 1rst CMP-free diet: | |
| | | every 3 to 7 | | form of | | 0.20+-0.13 | |
| | | days- and on | | commercial | | CMP challenge: - | |
| | | pain in the | | formula derived | , | CMP-free diet: - | |
| | | passage of | | from cow milk or | no pain | | |
| | | hard stools. In | | whole cow milk | , | Qualitative score: | |
| | | all patients | | and its derivatives | liquid stool | | |
| | | the frequency | | | During the | -Cured (n=21) | |
| | | of stools per | | -For the next | various study | | |
| | | day was lower | | month: all patients | 1 (| a. UD: 2.85+-0.05 | |
| | | than the 3rd | | started a cow's | | b. 1rst CMP-free diet: | |
| | | percentile of | | milk protein-free | , | 1.90+-0.08 | |
| | | the values | | diet. Three | | c. 1rst CMP | |
| | | observed in a | | patients aged < | | challenge: 2.75+-0.11 | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|------------------------|------------------------|-------------------|
| miormation | Level | - ationto | S | Companicon | Measures | | |
| | | large | | 12 months were | | d. 2nd CMP-free diet: | |
| | | population of | | fed a formula | | 1.85+-0.10 | |
| | | healthy | | containing soy | | | |
| | | subject | | protein and the | | (p<0.001) | |
| | | participating | | others received | | | |
| | | in an Italian | | soy milk or ass's | | -Unimproved (n=6) | |
| | | multicentre | | milk (eight cases) | | | |
| | | study | | and all milk | | UD: 3 | |
| | | <u>Exclusion</u> | | derivatives were | | 1rst CMP-free diet: 3 | |
| | | criteria: | | excluded | | CMP challenge: - | |
| | | Hirschsprung' | | | | CMP-free diet: - | |
| | | s disease, | | After a month: | | | |
| | | mental | | -Patients whose | | Difficulty in passing | |
| | | retardation | | symptoms abated: | | stools: | |
| | | | | cow milk | | | |
| | | | | challenge. Cow | | -Cured (n=21) | |
| | | | | milk given for a | | | |
| | | | | maximum of 10 | | a. UD: | |
| | | | | days, again an | | B. 1rst CMP-free diet: | |
| | | | | exclusion diet for | | none had difficulty | |
| | | | | 1 month and then | | c. 1rst CMP | |
| | | | | a second cow milk | | challenge: Painful | |
| | | | | challenge. All | | d. 2nd CMP-free diet: | |
| | | | | challenges were | | none had difficulty | |
| | | | | performed in | | D | |
| | | | | hospital. Before | | During the second | |
| | | | | the challenge a | | challenge symptoms | |
| | | | | prick test was | | reappeared within 24 | |
| | | | | performed with | | to 48 h: all 21 | |
| | | | | CMP. In patients | | patients had painful | |
| | | | | with a negative | | passage of stools and | |
| | | | | result, the | | for this reason | |
| | | | | challenge was | | challenge was | |
| | | | | performed by | | suspended on the | |
| | | | | giving whole cow | | third day | |
| | | | | milk in a singles | | Unimproved (n=6) | |
| | | | | feeding; if there | | -Unimproved (n=6) | |
| | | 1 | | were no clinical | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|------------------------|-------------------|
| | | | | reactions, the | | Control: ? | |
| | | | | same food was | | 1rst CMP-free diet: no | |
| | | | | given the | | changes | |
| | | | | following days. In | | CMP challenge: - | |
| | | | | patients with a | | CMP-free diet: - | |
| | | | | positive test | | | |
| | | | | result, the | | Follow-up period: | |
| | | | | challenge was | | Reintroduction of cow | |
| | | | | performed by | | milk was cautiously | |
| | | | | giving a formula | | attempted in 16 | |
| | | | | containing CMP, | | children 6-9 months | |
| | | | | beginning with an | | after the diagnosis of | |
| | | | | initial quantity of | | CMPA-dependant | |
| | | | | 10 ml and | | constipation. In eight | |
| | | | | gradually | | children CMP did not | |
| | | | | increasing the | | cause the onset of | |
| | | | | amount to reach | | any problems and it | |
| | | | | the dose | | was reintroduced on | |
| | | | | equivalent to a full | | a permanent basis; in | |
| | | | | feeding after 48 | | eight patients CMP | |
| | | | | hours. No other | | led to the | |
| | | | | change in diet | | reappearance of | |
| | | | | was made. | | constipation within 2 | |
| | | | | | | to 3 days after | |
| | | | | Reintroduction of | | introduction, and | |
| | | | | cow milk | | these infants were | |
| | | | | cautiously | | still following CMP- | |
| | | | | attempted in 16 | | free diet at the time | |
| | | | | children 6-9 | | the paper was written. | |
| | | | | months after the | | | |
| | | | | diagnosis of | | | |
| | | | | CMPA-dependant | | | |
| | | | | constipation | | | |
| | | | | -Patients with no | | | |
| | | | | abatement in | | | |
| | | | | symptoms: | | | |
| | | | 1 | permanently given | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---|------------------------------------|-------------|-------------------|
| | | | | an unrestricted diet, except for one infant who had episodes of recurrent bronchospasm related to ingestion of cow milk | | | |
| | | | | | | | |
| | | | | | | | |

| Level S Measures S Measures S Sudy Types 36 Cas series infants and chronic constipation manometry and chronice manometry and of Hepatology and 19[2], 143-150 S S S S S S S S S | Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|--|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|---------------------|---|
| Case series and enhoderonic constipation: manometry and enhoded constipation: and chronic constipation: and chronic constipation and original of Gastroenterolog y and Hepatology 18(2), 143-150 8(2), 143-150 8(2), 143-150 8(2), 143-150 8(3), 143-150 8(4), 143-150 8(4), 143-150 8(5), 143-150 8(5), 143-150 8(6), 143-150 8(7), 143-150 8(7), 143-150 8(8), 1 | | | | | | | | |
| intolerance and chronic controlled constituation: manmetry and histology study. 2006. Europea Journal of Sastroenterolog yand Hepatology 18[2], 143-150 Appearance of the sum of the s | | | 36 | 20 females | | Follow-up | Observation period: | |
| ektronic constipation anometry and character manometry and character of the deasures: Evidence level: 3 | | | | | | | | |
| constipation: manometry and manometry and histology study. 2006. European Journal of Castroenterolog y and Hepatology 18[2], 143-150 Facility of Chronic Constipation histology study. 2006. European Journal of Gastroenterolog y and Hepatology 18[2], 143-150 Facility of Chronic Constipation histology study. 2006. European Journal of Gastroenterolog y and Hepatology 18[2], 143-150 Facility of Chronic Constipation defined as less than 3 bowel on hard stools of Chronic Constipation defined as less than 3 bowel on hard stools on history of chronic Constipation lasting at least 3 months old a commercially available soy milk. Comparisons: 1. Cow's milk and all its derivatives (Manusculation of histology study. 2006. European Journal of Castroenterology by death of the study to evaluate adherence to the diet movements/week. Number of bowel movements/week. 2. Cow's milk vs. ass's milk v | intolerance and | and | infants and | Aged 9 months | diet, with the | Not reported | -Patients with food | asked to record the amount and type of |
| manometry and chistology study. 2006. European Journal of Gastroenterolog y and Hepatology 18[2], 143-150 Federal Pacification of Hepa | chronic | | | | | | intolerance (n=17) | |
| histology study. 2006. European Journal of Gastroenterolog y and Hepatology 18[2], 143-150 Evidence or y and the quantity of milk consumed bowle movements/we kexamined at the outpatient the uptatient of district of a hospital Paediatric Gastroenterol ogy Division. Chronic constipation defined as less than 3 bowle movements/r week with painful elimination of hard stools and points of chronic constipation lasting at least 3 months old a commercially aleast 3 months old a commercially available soy milk. | | | | | | | | |
| Journal of Gastroenterolog y and Hepatology 18[2], 143-150 Fig. 2 | | | | years) | derivatives | | | |
| Journal of Gastroenterology and Hepatology 18[2], 143-150 Septiments Septiment | | challenge | | | | | | and the quantity of milk consumed |
| Gastroenterolog y and Hepatology 18[2], 143-150 Sevel | | | | | | | | |
| y and Hepatology 18[2], 143-150 Secondary 18[2], 143-15 | | | | | | | | |
| Hepatology 18[2], 143-150 A | _ | | | | | ek | percentile: 1-2 | |
| hospital Paediatric Gastroenterol ogy Division. Chronic constipation defined as less than 3 bowel movements/r week with painful painful painful painful chronic criteria: on a history of chronic constipation at gastro of the mouths of the diet vs. soy milk: of difficulty in passing stools and difficulty and pain on passing stools and difficulty and pain on a history of chronic constipation lasting at least 3 months lea | | | | | | | | contained asses' or cows' milk. |
| Paediatric Gastroenterol ogy Division. Chronic constipation defined as less than 3 bowel movements/r week with painful elimination of hard stools elimination of hard stools on a history of chronic constipation as time at least 3 months least 3 months lasting at least 6 months old are more mere lasting at least 6 months old a commercially available soy milk. 1. Cow's milk-free diditiculty in difficulty | | 3 | | | ass's milk | 1 1 | | |
| Gastroenterol ogy Division. Chronic constipation defined as less than 3 bowel movements/r week with painful elimination of hard stools 1 Inclusion criteria: constipation defined as less than 3 bowel stopped 1 Inclusion criteria: constipation defined as less than 3 bowel movements/r week with painful elimination of hard stools 1 Inclusion criteria: constipation al least 3 months -lack of response to a | 18[2], 143-150 | | | | | | | |
| ogy Division. Chronic Chronic constipation defined as less than 3 bowel movements/r week with painful elimination of hard stools oriteria: - a history of chronic constipation lasting at least 3 months - lack of response to a commercially available soy milk constipation observation of efaced score: - Patients with constipation unrelated to food intolerance (n=19): Specific data related to number of bowel movements and qualitative faecal score were not reported for the challenge period. Number of bowel movements/week: and no pain in patients on cow's milk challenge not described cow's milk challenge not described on turnelated to food intolerance (n=19): Number of bowel movements/week: and no pain in patients on cow's milk challenge not described covs's milk challenge not described on worements with constipation unrelated to food intolerance (n=19): Number of bowel movements/week: and no pain in patients on cow's milk challenge not described covs's milk challenge not described covs's milk challenge not described on worements with constipation unrelated to food intolerance (n=19): Number of bowel movements/week: and no pain in patients on cow's milk challenge not described covs's milk challenge not described on worements and qualitative faecal score were not reported for the challenge period. Number of bowel movements/week: and no pain in patients on cow's milk challenge not described on tonelated to food intolerance (n=19): Number of bowel movements/week: and no pain in passing stools and difficulty and pain on passing stools Qualitative faecal score were not reported for the challenge period. Number of bowel movements/week: and no pain | | | | | | | | faecal score had been previously validated |
| Chronic constipation defined as less than 3 bowel movements/r week with painful elimination of hard stools criteria: - a history of chronic constipation anoths - lack of response to a commercially available soy milk. Chronic constipation observation defined as less than 3 bowel movements/r week with painful elems than 3 bowel movements and qualitative faecal score were not reported for the challenge period. Number of bowel movements and qualitative faecal score were not reported for the challenge period. Analysis of the main constituents of the diet (proteins, carbohydrates and fibres) did not show any qualitative or quantitative variation during the study period (data not shown) passing stools (All outcomes measures were recorded by parents) (All outcomes measures were recorded by pare | | | | | diet vs. soy milk: | | | |
| constipation defined as less than 3 bowel stopped stopped stopped similation of hard stools elimination of hard stools or criteria: constipation attempts of chronic constipation lasting at least 3 months old a response to a less than 3 bowel a less than 3 medications period: all medications and period: all movements and qualitative faecal score were not reported for the challenge period. Analysis of the main constituents of the movements/week: all movements/week: diet (proteins, carbohydrates and fibres) and difficulty and pain on passing stools shown) and pain on passing stools (All outcomes movements and qualitative faecal score were not reported for the challenge period. Analysis of the main constituents of the movements/week: diet (proteins, carbohydrates and fibres) and difficulty and pain on passing stools (All outcomes score or multiple) were treated as a group for the period: and outring the study period (data not shown) and pain on passing stools (All outcomes measures were not reported for the challenge period. Qualitative faecal score were not reported for the challenge period. Qualitative faecal score diet (proteins, carbohydrates and fibres) and difficulty and pain on passing stools (All outcomes measures were not reported for the challenge period. Qualitative faecal score movements/week: diet (proteins, carbohydrates and fibres) and pa | | | | | | | 3: 17 | |
| defined as less than 3 bowel movements/r week with painful elimination of hard stools - a history of chronic constipation lasting at least 3 months lasting at least 3 months lasting at least 3 months lasting at least 6 movements (less than 3 bowel movements/r week with painful eless than 3 bowel movements/r week with painful eless than 3 bowel movements/r week with painful elimination of hard stools and difficulty and pain on painsing stools and difficulty and pain on passing stools and difficulty and pain on passing stools and difficulty and pain on passing stools are ceived a formula based on soy (Nutrilon-soya, Nutricia, months old a months old a months old a months least 3 months old a months of response to a less than 3 bowel medications stopped unrelated to food intolerance (n=19): were not reported for the challenge period. Number of bowel movements/week: diet (proteins, carbohydrates and fibres) did not show any qualitative or quantitative variation during the study period (data not show any qualitative faecal score were not reported for the challenge period. Number of bowel movements/week: diet (proteins, carbohydrates and fibres) did not show any qualitative or quantitative faecal score were not reported for the challenge period. Analysis of the main constituents of the movements/week: did not show any qualitative or quantitative faecal score were not reported for the challenge period. Analysis of the main constituents of the movements/week: did not show any qualitative or quantitative faecal score were not reported for the challenge period. Analysis of the main constituents of the movements/week: did not show any qualitative or quantitative faecal special period (proteins, carbohydrates and fibres) did not show any qualitative or quantitative faecal special period (proteins, carbohydrates and fibres | | | | | | | | cow's milk challenge not described |
| less than 3 bowel movements/r week with painful elimination of hard stools Inclusion criteria: - a history of chronic constipation lasting at least 3 months -lack of response to a eliminate of bowel months old a commercially available soy milk. Ideast 3 months -lack of response to a elimination showel movements/r week with painful patients on cow's stopped stooped movements/r week with patients on cow's milk free diet. Patients on cow's milk into the passing stools and no pain in patients on cow's milk free diet. Patients with food intolerance (n=19): Inclusion or the diet (proteins, carbohydrates and fibres) and difficulty and pain on passing stools formula based on soy (Nutrilon-soy, Nutricia, measures were recorded by parents) Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, months old a commercially available soy milk. Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, milaty), children>15 months old a commercially available soy milk. Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, measures were recorded by parents) Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, milaty), children>15 months old a commercially available soy milk. Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, measures were recorded by parents) Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, measures were recorded by parents) Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, measures were recorded by parents) Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, measures were recorded by parents) Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, measures were recorded by parents) Inclusion oriteria: formula based on soy (Nutrilon-soy, Nutricia, measures were recorded by parents) Inclusion oriteria: formula to food intolerance (n=19): Number of bowel movements/week: Analysis of the m | | | | | | faecal score: | | |
| bowel movements/r week with painful elimination of hard stools Inclusion criteria: formula based on chronic constipation lasting at least 3 months eleast 3 m | | | | | 1 | | | |
| movements/r week with painful elimination of hard stools elimination of chronic constipation lasting at least 3 months old response to a elimination of elements on cow's hard stools elimination of more elimination of elimination of elimination of elimination eliminati | | | | | | | | |
| week with painful elimination of hard stools Inclusion criteria: | | | | | stopped | | intolerance (n=19): | were not reported for the challenge period. |
| painful elimination of hard stools Inclusion criteria: - a history of chronic constipation lasting at least 3 months old a months -lack of response to a patients on cow's milk free diet. Infants < 15 molths old a commercially available soy milk. patients on cow's milk free diet. Infants < 15 milk free diet. Infants < 15 movements/week: Median: 1.5 (and difficulty and pain on passing stools) 3. Hard stools Median: 1.5 (25th to 75th percentile: 1-2 shown) Patients with food intolerance (to CM only or multiple) were treated as a group for the purpose of analysing the data, therefore it is not possible to offer specific data for the stool habits: a bowel Normalised stool habits: a bowel Milur and MiPAF: project "Alimetazione e celiachia (ALICE)", D.D. n 86 dated | | | | | | | | |
| elimination of hard stools Millan, Italy), constipation lasting at least 3 months old aronsponse to a Millan, Italy Lack of response to a Millan, Italy available soy milk. Median: 1.5 | | | | | | | | |
| hard stools Infants < 15 months old received a formula based on criteria: - a history of chronic constipation lasting at least 3 months -lack of response to a Infants < 15 months old received a formula based on soy (Nutrilon- soya, Nutricia, constipation lasting at least 3 months -lack of response to a Infants < 15 months old received a formula based on soy (Nutrilon- soya, Nutricia, children>15 months old a commercially available soy milk. Infants < 15 months old received a formula based on soy (Nutrilon- soya, Nutricia, children>15 months old a commercially available soy milk. Infants < 15 months old received a formula based on soy (Nutrilon- soya, Nutricia, measures were recorded by parents) Infants < 15 months old passing stools Qualitative faecal score 1: 0 2: 0 3: 19 CM group only CM group only CM group only Funding: partly supported by a grant from MIUR and MiPAF: project "Alimetazione e celiachia (ALICE)", D.D. n 86 dated | | | | | | | | |
| Inclusion criteria: - a history of chronic constipation lasting at least 3 months -lack of response to a months old received a formula based on soy (Nutrilon- soya, Nutricia, Milan, Italy), children>15 months old received a formula based on soy (Nutrilon- soya, Nutricia, Milan, Italy), children>15 months old a commercially available soy milk. months old received a formula based on soy (Nutrilon- soya, Nutricia, Milan, Italy), children>15 months old a commercially available soy milk. months old received a formula based on soy (Nutrilon- soya, Nutricia, Milan, Italy), recorded by parents) least 3 months -lack of response to a months old received a formula based on soy (Nutrilon- soya, Nutricia, Milan, Italy), recorded by parents) least 3 months -lack of response to a months old received a formula based on soy (Nutrilon- soya, Nutricia, Milan, Italy), recorded by parents) least 3 months -lack of response to a months old soore leach of received a formula based on soy (All outcomes measures were recorded by parents) leach of received a formula based on soy (Nutrilon- soya, Nutricia, Milan, Italy), parents) leach of leach of received a formula based on soy (All outcomes measures were recorded by parents) leach of leach | | | | | | | | |
| Inclusion criteria: - a history of chronic constipation lasting at least 3 months - lack of response to a received a formula based on soy (Nutrilon-soya, Nutricia, months old a criteria: - a history of chronic constipation lasting at least 3 months old a received a formula based on soy (Nutrilon-soya, Nutricia, Milan, Italy), children>15 months old a commercially available soy milk. Normalised stool habits: a bowel Stool habits: a bowel | | | nard stools | | | - | | · · · · · · · · · · · · · · · · · · · |
| criteria: - a history of chronic constipation lasting at least 3 months -lack of response to a lasting at least of the chronic constipation lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response | | | la alcaia a | | | | percentile: 1-2 | snown) |
| - a history of chronic soya, Nutricia, Milan, Italy), children>15 months old a months - lack of response to a soy (Nutrilon-soya, Nutricia, Soya, Nutricia, Milan, Italy), children>15 months old a soy milk. | | | | | | passing stools | Ouglitative forced | Datients with food intelerence (to CM only |
| chronic constipation lasting at least 3 months -lack of response to a lasting at least 3 and lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months -lack of response to a lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at least 3 months old a commercially available soy milk. Stool habits: a labowel lasting at lasting | | | | | | (All autaana | | |
| constipation lasting at least 3 months -lack of response to a lasting at least 3 and response to a lasting at least 3 months -lack of response to a lasting at least 3 months old a commercially available soy milk. In the control of the children is not possible to offer specific data for the condition is not possible to offer specific data for the condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group only condition is not possible to offer specific data for the CM group | | | , | | • ` | | | |
| lasting at least 3 months old a commercially available soy milk. In the parents of response to a children>15 months old a commercially available soy milk. In the parents of p | | | | | | | | |
| least 3 months old a commercially available soy milk. Size of response to a months old a commercially available soy milk. Size old habits: a bowel Elimination diet period: Elimination diet period: MIUR and MiPAF: project "Alimetazione e celiachia (ALICE)", D.D. n 86 dated | | | | | | , | | |
| months -lack of response to a commercially available soy milk. Normalised stool habits: a bowel Elimination diet period: Funding: partly supported by a grant from MIUR and MiPAF: project "Alimetazione e celiachia (ALICE)", D.D. n 86 dated | | | J | | | parents) | 3. 18 | Civi group orily |
| -lack of response to a available soy milk. stool habits: a bowel period: MIUR and MiPAF: project "Alimetazione e celiachia (ALICE)", D.D. n 86 dated | | | | | | Normalised | Elimination diet | Funding: partly supported by a great from |
| response to a bowel celiachia (ALICE)", D.D. n 86 dated | | | | | | | | |
| | | | | | available suy IIIIK. | | penou. | |
| intevious I reguents ittenuency of at 1-eattents with food 180 01 2007 | | | previous | | Patients | frequency of at | -Patients with food | 30.01.2002) |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------|---------------------|-----------------------|-------------------|
| information | Level | Patients | S | Comparison | Measures | | |
| | | increase in | | unresponsive to | least three | intolerance (n=17) | |
| | | dietary fibre | | CM-free diet | evacuations per | | |
| | | intake or to | | placed on | week, with the | Number of bowel | |
| | | laxative | | oligoantigenic diet | elimination of | movements/week: | |
| | | treatment | | 4 weeks (also | soft stools, | Median: 5 | |
| | | (milk of | | excluding cow's | without painful | 25th to 75th | |
| | | magnesia 1-2 | | milk): exclusively | defecation | percentile: 3-7 | |
| | | ml/ kg of body | | rice, lamb, | | | |
| | | weight) | | carrots, ass's | | Qualitative faecal | |
| | | -a regular | | milk, olive oil and | | score | |
| | | dietary intake | | sugar | | 1: 1 | |
| | | of cow's milk | | | | 2: 16 | |
| | | and | | 2. Cow's milk vs. | | 3: 0 | |
| | | derivatives | | ass's milk: | | | |
| | | | | | | - Patients with | |
| | | <u>Exclusion</u> | | After 12 weeks: | | constipation | |
| | | criteria: | | patients cured on | | unrelated to food | |
| | | -previous | | cow's milk-free | | intolerance (n=19): | |
| | | evaluation for | | diet and | | | |
| | | chronic | | oligoantigenic | | Number of bowel | |
| | | constipation | | underwent a 2- | | movements/week: | |
| | | -anatomical | | week double-blind | | Median: 1.5 | |
| | | /neurological | | placebo-controlled | | 25th to 75th | |
| | | causes | | challenge with | | percentile: 1-2 | |
| | | (Hirschsprung | | cow's milk. Asses' | | | |
| | | 's disease, | | milk was used as | | Qualitative faecal | |
| | | psychomotor | | placebo. If no | | score | |
| | | retardation) | | clinical reactions | | 1: 0 | |
| | | -another | | after 12 hours, | | 2: 0 | |
| | | disease | | patients were | | 3: 19 | |
| | | (coeliac | | discharged and | | | |
| | | disease, | | challenge | | Cow's milk challenge | |
| | | hypothyroidis | | continued at | | <u>period</u> | |
| | | m) | | home with bottles | | | |
| | | -previous anal | | coded A or B. | | Reappearance of | |
| | | surgery | | Challenge was | | constipation in all | |
| | | -use of | | stopped when a | | cases (n=17), very | |
| | | medication | | clinical reaction | | often associated with | |

| Bibliographic Information | Evidence Level | Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-------------------|---|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | that causes constipation -referral for reasons other than constipation | | occurred | | painful defecation, within 5 days after the commencement of the challenge (median 2 days, range 1-5 days). | |
| | | | | | | | |
| | | | | | | | |

Psychological/Behavioural Interventions for Ongoing Treatment/Maintenance in Children with Chronic Idiopathic Constipation

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|------------------------|--|
| D | Level | 101 111 | S | | Measures | IDD : :: | |
| van Dijk et al. | Study Type: | 134 children | 134 children | General: | <u>Intervention</u> | IRR: incidence rate | Additional information from study: |
| Behavioral | Parallel-RCT | | 70.1 | -Disimpaction: | | ratio | At entry, patients had to meet at least |
| therapy for | Fridalesses | Inclusion | 76 boys | daily Klyx enemas | | RR: relative risk | 2 of 4 criteria: defecation frequency< 3 |
| childhood | Evidence | criteria: | 4. | (sodium- | | | times per week, faecal incontinence ≥ 2 |
| constipation: a | <u>level:</u> | | age range: 4 to | | | BT (n=67) | times per week, passage of large |
| randomized, | 1+ | functional | 18 years | ate and sorbitol; | weeks with | | amounts of stool at least once every 7 to |
| controlled trial. | | constipation | | 60 mL/day for | similar intervals | Defecation frequency | 30 days (large |
| 2008. Pediatrics | | aged 4 to 18 | -mean age: | · | between | | enough to clog the toilet), or a palpable |
| 121[5], e1334- | To evaluate | years referred | | of age; 120 | treatment | <u>CI)</u> | abdominal or rectal faecal mass |
| e1341 | the clinical | | CT group: 6.5 | mL/day for | sessions | | |
| | effectiveness | | (2.1) | children > 6 years | | -Post-treatment | After baseline measurement and if |
| | of | al outpatient | | of age) for 3 | <u>Assessment</u> | | written informed consent was given, a |
| | behavioural | | BT group: 6.9 | | | CT: 7.2 (6.1 to 8.5) | research assistant performed a |
| | therapy with | Emma | (2.5) | was prescribed by | follow-up | BT: 5.4 (4.3 to 6.7) | telephone call to a randomization centre |
| | laxatives | Children's | | paediatric | period: | | and revealed the allocation to parents |
| | compared | Hospital | Country: | gastroenterologist | | -Follow-up | immediately. A computer-based system |
| | with | | The | s before starting | At the last visit | | used to generate a sequence of random |
| | conventional | 2002 and | Netherlands | treatment | (posttreatment | CT: 6.6 (5.0 to 8.8) | group assignment for consecutive |
| | treatment in | August 2004 | | | time point) and | BT: 5.3 (4.4–6.3) | patients. Random assignment stratified |
| | treating | | | -Maintenance: | 6 months after | | by age (4 to 8 years or ≥8 years) and |
| | functional | Exclusion | | polyethylene | the 22-week | Group (main effect of | gender. Within 2 weeks after random |
| | constipation in | criteria: | | glycol 3350, 1 | treatment | BT): | assignment, patients received their 1rst |
| | childhood | Having | | sachet (10 g) per | ended (follow- | | treatment session |
| | | received a | | day, and if | up). | IRR=0.75 (0.59 to | |
| | | comprehensiv | | treatment | Time between | 0.96) p=0.021 | Sample size calculated to allow |
| | | e BT in the | | considered to | baseline | | detection of a 25% difference in the |
| | | previous 12 | | have insufficient | assessment | Group x time | proportion of success between BT and |
| | | months, use | | effect dose | and follow-up: | (interaction effect of | CT. It was estimated that CT reached |
| | | of drugs | | increased by 1 | ~1 year | BT with measurement | success in 35% of the children at follow- |
| | | influencing | | sachet. If | | at follow up): | up. Under the additional assumption of a |
| | | gastrointestin | | spontaneous | Outcome | ., | significance level of .05, a power of .80, |
| | | al function | | defecation | | IRR= 1.06 (0.75 to | and 2-sided hypothesis testing, a |
| | | other than | | delayed for >3 | | 1.50) p=0.758 | minimal sample size of 124 with 62 |
| | | laxatives, | | days, parents | -Primary | , , , , , , , , | children in each group was determined |
| | | organic | | advised to give an | | Faecal incontinence | 3 11, 11 11 11 11 |
| | | causes for | | enema or | | | During treatment 2 (3.1%) of 64 in the |

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|---------------|-------------------|----------------|------------------|---------------------|---------------------|-------------------------------------|--|
| Information | Evidence Level | Patients | Characteristic s | Comparison | Outcome Measures | | |
| | | defecation | | bisacodyl | a. defecation | CI) | CT group and 9 (13.8%) of 65 in the BT |
| | | disorders, e.g | | suppository of 5 | frequency | | group discontinued intervention |
| | | Hirschsprung' | | mg. In BT | per week | -Post-treatment | (P=0.054). At follow-up, 4 patients |
| | | s disease, | | preferred to give | | | dropped out in CT. There was 1 loss of |
| | | spina bifida | | oral bisacodyl | b. faecal | CT: 2.1 (0.8 to 5.8) | contact, and 3 children were referred for |
| | | occulta, | | tablets of 5 mg | incontinence | BT: 5.0 (2.1 to 12.0) | BT directly after CT, making them |
| | | hypothyroidis | | instead of rectal | frequency per | | unsuitable for follow-up measurements. |
| | | m, or other | | laxatives. During | week | -Follow-up | Questionnaires were not returned by 3 |
| | | metabolic | | BT, paediatric | | | patients in both intervention arms at |
| | | or renal | | psychologists | c. successful | CT: 6.4 (3.5 to 11.7) | posttreatment and by 9 patients (CT: 6; |
| | | abnormalities | | adjusted laxative | treatment | BT: 8.6 (4.0 to 18.3) | BT: 3) at follow-up |
| | | | | dose and | 1 | | |
| | | | | consulted | -Secondary | | Except for painful defecation (65.0% CT |
| | | | | paediatric | outcomes: | BT): | vs. 43.1% BT, P=0 .014), no significant |
| | | | | gastroenterologist | | IDD 0.00 (0.77.) | differences between the 2 groups in |
| | | | | | a. stool | IRR=2.36 (0.77 to | baseline sociodemographic factors or for |
| | | | | | withholding | 7.31) p=0.135 | clinical characteristics |
| | | | | groups, patients | behaviour | Cuarra ve tima a | Intent to treat analysis and stand |
| | | | | kept a bowel diary | | Group x time (interaction effect of | Intent-to-treat analyses conducted. |
| | | | | Intervention: | | | Because of withdrawal before treatment start, dropouts during the study, failure |
| | | | | Protocolised | | at follow up): | to fill out questionnaires, or research |
| | | | | behavioural | | at follow up). | procedure violations, missing data |
| | | | | therapy (BT) | | IRR= 0.57 (0.12 to | occurred. Imputation of missing values |
| | | | | пстару (Вт) | | 2.61) p=0.467 | used to make intent-to-treat analyses |
| | | | | -developed by | | 2.01) p=0.401 | feasible |
| | | | | paediatric | | Success, % (95% CI) | Todololo |
| | | | | psychologists of | | | Treatment considered successful if |
| | | | | the psychosocial | | -Post-treatment | patients achieved a defecation |
| | | | | department of our | | CT: 62.3 (51.1 to | frequency of ≥3 |
| | | | | hospital. Basic | | 76.1) BT: 51.5 (39.7 | times per week and a faecal |
| | | | | assumption that | | to 66.9) | incontinence frequency of ≤1 times per |
| | | | | phobic reactions | | | 2 weeks, irrespective of laxative use |
| | | | | related to | | RR= 0.83 (0.60 to | |
| | | | | defecation | | 1.14) p=0.249 | Reviewer comments: |
| | | | | can be reduced | | | Insufficient details on how outcomes |
| | | | | and that adequate | | -Follow-up | were measured |
| | | | | toileting behaviour | | CT: 57.3 (46.6 to | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|--|---------------------|--|---|
| | | | | and appropriate defecation straining can be (re)acquired by teaching parents behavioural procedures and by behavioural play therapy with the child in presence of his or her parents. The protocol consists of 2 age-related modules: a module for children aged 4 to 8 years and a module for children aged ≥8 years. Learning process for child and parents: 5 sequential steps (know, dare can, | Outcome Measures | 70.4) BT: 42.3 (31.8 to 56.4) RR= 0.74 (0.52 to 1.05) p=0.095 Stool withholding behaviour at follow-up (% children with behaviour) CT: 13.8 BT: 10.6 NS | Results controlled for confounders Source of funding: funded in part by the Dutch Digestive Disease Foundation (SWO 02-16) |
| | | | | will, and do). This approach is derived from a multidisciplinary BT to treat children with defecation disorders. For all involved psychologists, a detailed manual for both agerelated modules | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|-------------------------------------|------------------------------------|-------------|-------------------|
| | | | | available to | | | |
| | | | | ensure a standard | | | |
| | | | | delivery of | | | |
| | | | | therapy. Visits | | | |
| | | | | lasted ~45 | | | |
| | | | | minutes | | | |
| | | | | Comparison: | | | |
| | | | | Conventional | | | |
| | | | | treatment (CT) | | | |
| | | | | -conducted by | | | |
| | | | | paediatric | | | |
| | | | | gastroenterologist | | | |
| | | | | s, visits lasted ~20 | | | |
| | | | | to 30 minutes, | | | |
| | | | | laxative treatment | | | |
| | | | | and bowel diary | | | |
| | | | | discussed. | | | |
| | | | | Patients and their parents received | | | |
| | | | | education to | | | |
| | | | | explain that | | | |
| | | | | symptoms are not | | | |
| | | | | harmful and are | | | |
| | | | | common in | | | |
| | | | | children with | | | |
| | | | | functional | | | |
| | | | | constipation and | | | |
| | | | | that a positive, | | | |
| | | | | non-accusatory | | | |
| | | | | approach is | | | |
| | | | | essential. | | | |
| | | | | Children | | | |
| | | | | instructed not to | | | |
| | | | | withhold stool | | | |
| | | | | when they feel | | | |
| | | | | urge to defecate. | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|--|------------------------------------|-------------|-------------------|
| | | | | Motivation enhanced by praise and small gifts from the paediatric gastroenterologist s | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|---|---------------------------|------------------------------------|---------------------------|--|
| Ritterband et al. | Study Type: | 24 children | s 24 children | Intervention: | | Percentage change | Additional information from study: |
| An Internet | Parallel-RCT | 24 Gillaren | 24 Ciliulen | Laxatives + Web | | from pre- to post- | Computer and internet access provided |
| intervention as | (multicentre) | Inclusion | 19 boys | intervention | | assessment | to all families who contacted the |
| adjunctive | (mandochilo) | /exclusion | 10 boys | intorvention | o weeks | uooooomon. | research centre and met the inclusion |
| | <u>Evidence</u> | | mean age: 8.46 | Comparison: | Assessment | Number of faecal | criteria |
| pediatric | level: | | years (SD1.81) | Laxatives only | point (s): | accidents per week | ontona |
| encopresis. | 1+ | between 6 | , | | 3 weeks after | (mean, SD) | Participants received a \$25 gift |
| 2003. Journal of | | | -Web group: 12 | Laxatives: all | | -Web group: 0.50 | certificate to a local toy sore for |
| | Study aim: | | children (10 | children instructed | | (.85) | completing the pre-treatment |
| Clinical | To examine | | boys) | to start with a | Follow-up | , | assessment and another \$25 gift |
| Psychology | the utility and | and have no | , | basic regime of | period: | -No-Web group: 8.27 | certificate for completing the post- |
| | effectiveness | medical | -No-Web group: | one square of Ex- | None | (13.83) | treatment assessment |
| | of an Internet- | diagnosis | 12 children (9 | Lax (senna), twice | | | |
| | based version | other than | boys) | a day | <u>Outcome</u> | | Information regarding BM assessed by |
| | of enhanced | constipation | | | Measures: | | parent report on the Child Information |
| | toilet training | that could | | -The Web site: | | passed in the toilet | Form. Question regarding child's bowel |
| | | explain their | Country: | Web-based | | | habits included such as number of BMs |
| | | faecal | USA | | per week | -Web group: +152% | in toilet and use of toilet with / without |
| | | incontinence | | treatment of | | | parental prompts. Questions regarding |
| | | | | paediatric | -number of | | use of internet programme also included |
| | | | | encopresis (U- | | | in post-treatment form for the |
| | | | | CAN-POOP-TOO | movements | | intervention group. The Virginia |
| | | | | | | | Encopresis/Constipation Apperception |
| | | | | | | prompts | Test (VECAT) also administered. It |
| | | | | programme, | week | -Web group: +109% | assesses bowel specific problems |
| | | | | targets primarily 5 | 1 4 | A1 147 I | related to the process of encopresis, |
| | | | | to 10 years old | - bathroom use | -No-Web group: - | such as avoidance of the toilet, non |
| | | | | | | 37% | responsiveness to rectal distension cues |
| | | | | designed to be | | p=0.021 | and fear of defecation pain. A generic |
| | | | | used by child and | | Detharman | subscale included as a comparison |
| | | | | parent (s) | | Bathroom use with prompts | measure, addresses problem |
| | | | | together | -internet use | | behaviours not related to bowel issues. |
| | | | | 3 core modules | /most/least | -Web group: +47% | The VECAT consists of 18 pairs of drawings (9 pairs bowel-specific and 9 |
| | | | | take 60 to 90 | (| -No-Web group: -45% | parallel generic events) and child selects |
| | | | | | the programme; | | the picture in each pair that best |
| | | | | | preference | 140 | describes him/herself |
| | | | | users instructed to | <u>.</u> | Internet use (Web | docomboo mini/nordon |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|---------------------|---|---|
| | Level | | S | | Measures | | |
| | | | | review them | regarding | group only) | No significant differences in baseline |
| | | | | during the first | individual cores | | characteristics between the 2 groups |
| | | | | week: | an modules) | Most useful aspect | (age, gender, race, stage of bowel |
| | | | | 1. The body | | of the programme: | movement training, length of current |
| | | | | (anatomy, | | -the step by step | laxative regime or any of the outcomes |
| | | | | physiology and | | program to get the | measured) |
| | | | | pathophysiology | | child regulated | |
| 1 | | | | of digestion) | | -understanding why | CM1: anatomy and pathophisiology |
| 1 | | | | 2. How to poop | | | CM2: medication (enemas/laxatives) |
| | | | | (behavioural | | needs to do | CM3: behavioural intervention |
| | | | | techniques for | | everyday-and what | |
| | | | | treatment of | | happens when he | Reviewer comments: |
| | | | | encopresis) | | doesn't have a BM | No definition of constipation / soling |
| | | | | 3. Medication | | and health | given |
| | | | | (clean-out and | | consequencesinfor | Small sample size, no sample size |
| | | | | laxative | | mation was | calculation |
| 1 | | | | treatment) | | | Randomisation and allocation concealment method not described |
| | | | | New modules | | -developing a feeling that he can control | |
| | | | | assigned each | | his own body | No dropouts/lost to follow up reported |
| 1 | | | | week based on a | | | Results not controlled for potential |
| | | | | follow-up | | the only child with this | |
| 1 | | | | assessment the | | problemthat was | Contounders |
| | | | | user completes | | reassuring | Source of funding: |
| | | | | about their child's | | | National Institutes of Health Grant RO1 |
| | | | | status. Not all | | 2. Least useful aspect | |
| | | | | modules | | of the programme | 11520100 |
| | | | | necessarily used | | or the programme | |
| 1 | | | | by all users, only | | -difficulty with | |
| | | | | those modules | | connections | |
| | | | | identified as | | -modules regarding | |
| | | | | relevant are | | fear of toilet and | |
| | | | | assigned and | | "monsters" | |
| | | | | reviewed. | | -art work of the body | |
| | | | | However all | | did not print out | |
| | | | | modules can be | | -Miralax should have | |
| I | | | | viewed by all | | been included (as a | |
| | | | | users. Follow-up | | choice of laxative) | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------------|---------------------|--|-------------------|
| | Level | | S | | Measures | | |
| | | | | comprised of 17 | | -nutrition portion was | |
| | | | | to 20 questions, | | too limited | |
| | | | | depending on the | | latamant com a discussion | |
| | | | | week. System contains a total o | | Internet experience: parents' views / | |
| | | | | 22 modules, each | | satisfaction | |
| | | | | takes 5 to 10 | | | |
| | | | | minutes to review | | -found material | |
| | | | | | | understandable | |
| | | | | | | (mean 5.00, SD 0.00, | |
| | | | | | | N = 20) | |
| | | | | | | | |
| | | | | | | -found it easy to use | |
| | | | | | | (mean 4.62, SD 0.74, N = 21) | |
| | | | | | | 14 – 21) | |
| | | | | | | -believed their child | |
| | | | | | | liked the program | |
| | | | | | | (mean 4.05, SD 1.28, | |
| | | | | | | N = 21) | |
| | | | | | | | |
| | | | | | | - believed their child | |
| | | | | | | found it | |
| | | | | | | understandable (mean 4.32, SD 0.89, | |
| | | | | | | N = 19 | |
| | | | | | | | |
| | | | | | | - believed their child | |
| | | | | | | found it easy to use | |
| | | | | | | (mean 4.47, SD 0.77, | |
| | | | | | | N = 19) | |
| | | | | | | | |
| | | | | | | 2 Droforonoo | |
| | | | | | | 3. Preference regarding cores | |
| | | | | | | modules (CM) (mean, | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | SD) (score 0 to 4) | |
| | | | | | | a. How useful: CM1: 3.84 (0.38) CM2: 3.94 (0.24) CM3: 4.00 (0.00) | |
| | | | | | | b. How well did you understand the material CM1: 3.89 (0.32) CM2: 3.89 (0.32) | |
| | | | | | | c. how well did your child understand the material CM1: 3.53 (0.61) | |
| | | | | | | CM2: 3.28 (1.07) CM3: 3.54 (1.13) | |
| | | | | | | d. How much did you enjoy using the module CM1: 3.68 (0.48) CM2: 3.67 (0.49) CM3: 3.69 (0.48) | |
| | | | | | | e. How much did your child enjoy using the module CM1: 3.63 (0.76) CM2: 3.61 (0.98) CM3: 3.46 (1.13) | |
| | | | | | | 0.40 (1.10) | |

| | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|-----------------|----------------------|----------------|-------------------------|---------------------|---|----------------------|--|
| Information | Evidence | Patients | Characteristic | Comparison | Outcome | | |
| Borowitz et al. | Level Study Type: | 87 children | s 87 children | Intervention: | Measures Duration of | Soling | Additional information from study: |
| Treatment of | Parallel-RCT | or cilialen | or cilialen | Intensive medical | treatment | frequency(mean, SD) | Using a random number generator, |
| childhood | l alaliel-IVO1 | Inclusion | 72 boys | therapy (IMT) | Unclear | -at 3 months: | blocks of six consecutive children were |
| encopresis: A | Evidence | criteria: | 12 boys | linerapy (livir) | Unclear | IMT: 0.54 (0.68) | randomly assigned to one of 3 treatment |
| | level: | | Mean age at | 1 of 2 paediatric | Assessment | 11011. 0.54 (0.00) | groups |
| comparing three | | aged between | | gastroenterologist | | ETT: 0.22 (0.21) | groups |
| treatment | 17 | - | enrollment: 8.6 | s directed | follow-up period | L11. 0.22 (0.21) | All data were collected using the |
| | Study aim: | vears | ± 2.0 years | treatment: colonic | lollow-up period | BF: 0.34 (0.51) | Automated Patient Symptom Monitor |
| Journal of | | of age who | (range, 5 to 13 | | When subjects | DI . 0.34 (0.31) | system, a computerized voice-mail |
| | short- and | had | years) | a series of | had been | -at 6 months: | system that telephones the families |
| Gastroenterolog | | experienced | years) | enemas followed | enrolled in the | IMT:0.44 (0.52) | each day. With each telephone call, the |
| v and Nutrition | effectiveness | encopresis for | Country | by sufficient | study, data | 11011.0.44 (0.52) | computer asked parents the same 8 pre- |
| 34[4], 378- | of | a minimum of | | laxative therapy to | | ETT: 0.38 (0.45) | recorded questions relating to bowel |
| | | 6 months. | 00/1 | produce at least 1 | | L11. 0.00 (0.40) | habits during the previous 24 hours. |
| States. | treatment | defined as at | | soft stool each | were collected | BF:0.20 (0.26) | After parents had answered all |
| | protocols in | least weekly | | day without | for 14 | DI :0:20 (0:20) | questions, the computer checked |
| | children | episodes of | | associated pain. | consecutive | -at 12 months: | responses to ensure all items were |
| | experiencing | faecal soiling | | Laxatives | days before | IMT:0.33 (0.48) | answered and that responses were |
| | | for at least 6 | | prescribed: Milk of | | | within acceptable ranges. If the |
| | encopresis | months | | | | ETT: 0.36 (0.53; 95% | computer detected an error, the |
| | опоортоою | montrio | | senna (Senokot, | visit, and again | confidence interval, | questionnaire was repeated |
| | | Exclusion | | Ex-Lax, or | at 3 months, | 0.05 to 0.47) | queenemane mae repeateu |
| | | criteria: | | | 6 months, and | 0.00 10 0.17) | No significant differences in baseline |
| | | any chronic | | Castoria). | | BF:0.27 (0.37) | clinical or demographics characteristics |
| | | underlying | | Laxative dosages | initiation of | 2 (0.0.) | between the 3 groups |
| | | medical | | | therapy | NS among the 3 | groups |
| | | conditions or | | to produce 1 to 3 | ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, | groups at any time | Treatment considered successful if the |
| | | developmenta | | soft bowel | Outcome | groupe and any and | child experienced no episodes of faecal |
| | | l disabilities | | movements daily. | Measures: | Improvement rate (% | soiling during the 2-week assessment |
| | | | | An enema or | -soling | children) | 12 months after initiation of therapy |
| | | | | suppository | frequency | -at 2 weeks: | |
| | | | | administered if | | IMT: 41 | Reviewer comments: |
| | | | | child had not | -improvement | | No definition of constipation given |
| | | | | produced a bowel | | ETT: 48 | |
| | | | | movement during | | | No sample size calculation performed |
| | | | | a 48-hour period. | -cure rate | BF: 62 | |
| | | | | No specific dietary | | | Method of allocation concealment not |
| | | | | recommendations | -number of | NS between 3 groups | reported |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|------------------------------------|---------------------|------------------------------------|---|
| | Level | | S | | Measures | | |
| | | | | or manipulations | bowel | | |
| | | | | undertaken. | movements | | No drop outs/lost to follow up children |
| | | | | Families received | passed in the | IMT: 45 | reported |
| | | | | specific | toilet each day | FTT 05 | 0 (() |
| | | | | instructions and | | ETT: 85 | Source of funding: |
| | | | | written brochure | -self-initiated | DE: 04 | supported by National Institutes of |
| | | | | detailing | toileting each | BF: 61 | Health grant RO1 HD 28160 |
| | | | | treatment protocol | day | at C magnetha. | |
| | | | | and need for children to attend | lovetive use | -at 6 months: IMT: 41 | |
| | | | | the toilet at least | -laxative use | 11011.41 | |
| | | | | twice dally, | | ETT: 74 | |
| | | | | preferably after | | L11.74 | |
| | | | | breakfast and | | BF: 58 | |
| | | | | supper | | D1 . 00 | |
| | | | | очррог | | -at 12 months: | |
| | | | | Comparison 1: | | IMT: 41 | |
| | | | | Intensive medical | | | |
| | | | | therapy + | | ETT: 78 | |
| | | | | enhanced toilet | | | |
| | | | | training (ETT) | | BF: 61 | |
| | | | | | | | |
| | | | | Similar enema | | At 3 months, 6 | |
| | | | | and laxative | | months, and 12 | |
| | | | | therapy, with 1 | | months, the number | |
| | | | | clinical | | of children who | |
| | | | | psychologist | | responded in the ETT | |
| | | | | adjusting laxative | | group was | |
| | | | | dose. Only | | significantly greater | |
| | | | | difference from | | than in either the IMT | |
| | | | | previous therapy | | or the BF group (P < | |
| | | | | was that laxative | | 0.05), and these results were very | |
| | | | | therapy was decreased | | stable over time (P < | |
| | | | | gradually when | | 0.001). With all 3 | |
| | | | | children | | regimens, response | |
| | | | | demonstrated | | to treatment during | |
| | | | | stable bowel | | the first 2 weeks of | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|-------------------------------|---------------------|--------------------------------------|-------------------|
| Illiorniation | Level | ratients | S | Companison | Measures | | |
| | | | | frequency with no | | therapy strongly | |
| | | | | soiling episodes. | | correlated with | |
| | | | | As long as child | | response to treatment | |
| | | | | had daily bowel | | at 3, 6, and 12 | |
| | | | | movements of | | months ($r > 0.90, P <$ | |
| | | | | normal size for a | | 0.0001 in all cases). | |
| | | | | week, laxative | | Of those children who | |
| | | | | dose was | | had significant | |
| | | | | decrease by one | | improvement | |
| | | | | quarter. This | | after 2 weeks of | |
| | | | | process was | | therapy, 86 continued | |
| | | | | continued until | | to improve at 3 | |
| | | | | laxative therapy | | months, 83 at 6 | |
| | | | | was discontinued. | | months, and 81 at 12 | |
| | | | | If child did not | | months | |
| | | | | pass daily bowel movements of | | Cura rata (number of | |
| | | | | normal size, | | Cure rate (number of children cured) | |
| | | | | laxative dose was | | -at 12 months: | |
| | | | | increased. | | -at 12 months. | |
| | | | | Parents and child | | IMT: 10/29 (34.5%) | |
| | | | | instructed on the | | 11011. 10/29 (34.376) | |
| | | | | psychophysiology | | ETT: 12/27 (44.4%) | |
| | | | | of constipation | | LII. 12/21 (44.470) | |
| | | | | and encopresis, | | BF: 11/31 (35.5%) | |
| | | | | and how | | DI . 11/31 (33.370) | |
| | | | | responding to | | chisquare=0.9488 | |
| | | | | early rectal | | ornoquaro—oro roo | |
| | | | | distention cues | | p=0.7005 | |
| | | | | along with regular | | | |
| | | | | toileting was | | Number of bowel | |
| | | | | critical to avoid | | movements passed in | |
| | | | | reimpaction and | | the toilet each day | |
| | | | | to establish | | (mean, SD) | |
| | | | | regular bowel | | -at 3 months: | |
| | | | | habits. Various | | IMT:1.44 (0.57) | |
| | | | | incentive | | | |
| | | | | programs | | ETT: 1.21 (0.49) | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|--------------------------|-------------------|
| | Level | | S | | Measures | | |
| | | | | established, | | | |
| | | | | depending on the | | BF: 1.25 (0.64) | |
| | | | | developmental | | | |
| | | | | age and the | | -at 6 months: | |
| | | | | motivation of the | | IMT:1.36 (0.61) | |
| | | | | child. Target | | | |
| | | | | behaviours: | | ETT:1.31 (0.63) | |
| | | | | spontaneous trips | | | |
| | | | | to the toilet and | | BF:1.12 (0.60) | |
| | | | | clean pants. | | | |
| | | | | Toilet training was | | -at 12 months: | |
| | | | | "enhanced" | | IMT:1.30 (0.61) | |
| | | | | because | | ETT (04 (0 54) | |
| | | | | instructions were | | ETT:1.01 (0.51) | |
| | | | | given on the role | | DE 4.40 (0.07) | |
| | | | | of paradoxic | | BF:1.16 (0.67) | |
| | | | | constriction of the | | NC amage that 2 | |
| | | | | external anal | | NS among the 3 | |
| | | | | sphincter, and because | | groups at any time | |
| | | | | appropriate | | Self-initiated toileting | |
| | | | | defecation | | each day (times/day, | |
| | | | | straining was | | mean, SD) | |
| | | | | modeled. The | | -at 3 months: | |
| | | | | therapist sat on a | | IMT: 1.53 (0.77) | |
| | | | | portable toilet and | | 11011. 1.55 (0.77) | |
| | | | | demonstrated | | ETT: 1.62 (0.82) | |
| | | | | how to relax the | | 211.1.02 (0.02) | |
| | | | | legs and feet, how | | BF:1.40 (0.71) | |
| | | | | to take in a deep | | 2 | |
| | | | | breath and hold it | | -at 6 months: | |
| | | | | while sitting up | | IMT:1.49 (0.60) | |
| | | | | straight, and how | | _ (/ | |
| | | | | to push down with | | ETT:1.67 (0.95) | |
| | | | | the held breath | | , , | |
| | | | | and pull in from | | BF:1.34 (0.72) | |
| | | | | the lower | | , , | |
| | | | | abdomen (rectus | | -at 12 months: | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|---------------------------|------------------------------------|------------------------------------|---|-------------------|
| | Levei | | S | abdominous | ivieasures | IMT:1.40 (0.76) | |
| | | | | muscle) to propel | | 11011.1.40 (0.76) | |
| | | | | out a stool. The | | ETT:1.31 (0.83) | |
| | | | | child then | | E11.1.31 (0.63) | |
| | | | | replicated this | | BF:1.31 (0.69) | |
| | | | | while sitting on a | | ы .1.51 (0.09) | |
| | | | | portable toilet. | | NS among the 3 | |
| | | | | The child | | groups at any time | |
| | | | | received "hand | | groups at any time | |
| | | | | feedback" by | | Laxative use (number | |
| | | | | placing one hand | | of children using) | |
| | | | | on the abdomen | | -at 12 months: | |
| | | | | just below the | | IMT: 17/29 (58.6%) | |
| | | | | navel to feel the | | | |
| | | | | abdomen move | | ETT: 9/27 (33.3%) | |
| | | | | out when the | | (************************************** | |
| | | | | breath was | | BF: 17/31 (54.8%) | |
| | | | | pushed down, and | | (chi-square= 4.1414, | |
| | | | | placing the | | P= 0.1261) | |
| | | | | second hand just | | | |
| | | | | below the first to | | | |
| | | | | feel inward | | | |
| | | | | movement with | | | |
| | | | | contraction of the | | | |
| | | | | rectus | | | |
| | | | | abdominous. | | | |
| | | | | Parents instructed | | | |
| | | | | to prompt these | | | |
| | | | | behaviours at | | | |
| | | | | home. | | | |
| | | | | Additionally, 8 to | | | |
| | | | | 12 minutes of "toilet time" was | | | |
| | | | | scheduled daily, | | | |
| | | | | beginning 15 to | | | |
| | | | | 30 minutes after | | | |
| | | | | the same two | | | |
| | | | | meals. | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|-------------|-------------------|
| | Level | | S | | Measures | | |
| | | | | During these | | | |
| | | | | times, children | | | |
| | | | | were instructed to | | | |
| | | | | practice tensing | | | |
| | | | | and relaxing the | | | |
| | | | | external anal | | | |
| | | | | sphincter for the | | | |
| | | | | first 4 minutes, | | | |
| | | | | with the objective | | | |
| | | | | of localizing | | | |
| | | | | control of and | | | |
| | | | | fatiguing the | | | |
| | | | | external anal | | | |
| | | | | sphincter, and to | | | |
| | | | | mechanically | | | |
| | | | | stimulate the | | | |
| | | | | rectum. To | | | |
| | | | | desensitize | | | |
| | | | | children to toilet | | | |
| | | | | sitting, the second | | | |
| | | | | 4 minutes were | | | |
| | | | | spent "having fun" | | | |
| | | | | while being read | | | |
| | | | | to or playing | | | |
| | | | | games. During the | | | |
| | | | | final 4 minutes, | | | |
| | | | | the child was to | | | |
| | | | | strain and attempt | | | |
| | | | | to have a bowel | | | |
| | | | | movement while | | | |
| | | | | relaxing his or her | | | |
| | | | | legs and feet. This | | | |
| | | | | routine toilet | | | |
| | | | | sitting was | | | |
| | | | | discontinued 2 | | | |
| | | | | weeks after the | | | |
| | | | | last scheduled | | | |
| | | | | treatment session | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---|------------------------------------|-------------|-------------------|
| | | | | Comparison 2: Intensive medical therapy + enhanced toilet training + anal sphincter biofeedback (BF) Same instructions that previous 2 groups and simultaneously received surface electromyographic biofeedback training. Same 2 psychologists who worked with the ETT group also worked with the BF group | | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--------------------------|--|
| Loening- | Study Type: | 43 children | 43 children | Intervention: | Duration of | Recovery rate | Additional information from study: |
| Baucke. | Parallel-RCT | | | Conventional | treatment | (number | Constipation and encopresis defined as |
| Modulation of | | Inclusion | 33 boys | treatment alone | 6-month | recovered, %) | having ≥ 2 soiling episodes/week and |
| abnormal | Evidence | criteria: | | (CT) | protocol. | | evidence of a huge amount of faecal |
| defecation | level: | Children 5 to | Mean age: 8.9 | (-) | | -at 7 months: | material in the rectal ampulla at rectal |
| dynamics by | 1+ | 16 years with | years (range 5 | CT: use of | Assessment | | examination. In many patients stool |
| biofeedback | | chronic | to 16) | laxatives, | point (s) and | CT (n=19): 1(5) | evacuation was incomplete as |
| treatment in | Study aim: | constipation | , | increase of dietary | | - (- / (- / | evidenced by periodic passage of very |
| chronically | To determine | and | Country: | fibre and | period: | BF (n=22): 12 (55) | large amounts of stools (every 7 to 30 |
| constipated | whether | encopresis | USA | scheduled | | , , , , , , | days), often clogging the toilet |
| children with | outcome in | and abnormal | | toileting | 7 & 12 months | P<0.001 | 33 3 1 1 1 |
| encopresis. | chronically | defecation | | J 3 3 3 | | | Abnormal defecation dynamics defined |
| 1990. Journal of | | dynamics | | Disimpaction with | Outcome | Recovery rates did | as abnormal contraction of the external |
| Pediatrics | and | 1 | | enemas (type and | | not differ between | anal sphincter and pelvic floor during |
| 116[2], 214-222 | encopretic | Exclusion | | dose not reported) | | boys and girls in | defecation attempts, as determined by |
| 1 1 | children with | criteria: | | , , | Recovery rate | general and within | anorectal manometry |
| | abnormal | Hirschsprung' | | Maintenance: milk | , | the biofeedback | , |
| | defecation | s disease, | | of magnesia ~ | | group in particular. | Sample size and calculation: 2 pairs of |
| | dynamics | hypothyroidis | | 2ml/kg body | | Prior unsuccessful | subjects would be needed per group to |
| | could be | m, mental | | weight daily to | | treatment no related | allow a power of approximately 0.9 to |
| | improved with | deficiency, | | induce at least 1 | | to treatment outcome | detect a difference of 0.7 vs. 0.2 in |
| | biofeedback | chronic | | bowel movement | | in either group | achieving normal bowel habits (recovery |
| | training | debilitating | | daily and prevent | | | from constipation and encopresis) |
| | | diseases, | | faecal retention. | | Patients with an initial | . , |
| | | neurologic | | Doses decrease | | abdominal faecal | Sealed envelopes with cards indicating |
| | | abnormalities, | | gradually to | | mass (severe | either conventional therapy alone or |
| | | previous | | maintain daily | | constipation) | conventional therapy with biofeedback |
| | | surgery of the | | bowel movement | | significantly more | training used for randomisation |
| | | colon | | and prevent | | likely to recover with | |
| | | | | faecal retention | | BF training than with | 1 boy in the conventional treatment |
| | | | | and soiling | | CT alone (46% vs. | group was lost to follow-up 1 month after |
| | | | | J | | 0%, p<0.02) | treatment began. At that visit he was |
| | | | | Patients | | , | taking milk of magnesia and his soiling |
| | | | | instructed to | | -at 12 months : | had resolved. 1 boy was lost to follow-up |
| | | | | discontinue | | | in the biofeedback group |
| | | | | laxative therapy at | | CT (n=19): 3 (16) | after the first biofeedback session |
| | | | | 6 ± 0.5 months | | | |
| | | | | after initiation of | | BF (n=22): 11 (50) | Baseline characteristics not significantly |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---|------------------------------------|---|---|
| | | | | therapy Comparison: Conventional treatment (CT) + biofeedback (BF) Up to 6 sessions of biofeedback therapy 7 +/- 2 days apart. 1 session included approximately 30 to 35 defecation trials and lasted approximately 45 minutes Patients instructed to discontinue laxative therapy at 6 ± 0.5 months after initiation of therapy | | P<0.05 A 14-yeor old boy in the BF group had a relapse. He had severe faecal impaction with enormous abdominal distension initially. Faecal impaction recurred 4 months after successful discontinuation of milk of magnesia. at time study was written he had no soiling but required intermittent treatment for constipation | different between both groups apart from gender: more girl in the BF group than in the CT group (41% vs. 5%, p<0.02). During initial evaluation the following significantly more frequent in girls than in boys: severe constipation (an abdominal faecal mass present) (90% vs. 48%, p<0.03), daytime urinary incontinence (70% vs. 23%, p<0.02) and a history of previous urinary tract infection (60% vs. 6%, p<0.001) Patients considered to have recovered if they had ≥3 bowel movements/week and soiling ≤ 2 episodes/month while not receiving laxatives for 4 weeks. Patients considered not to have recovered if they had <3 bowel movements/week or were soiling >2 times/month or had been started on a regime of laxatives again Re-evaluation of patients included review of last month's stool, soiling and medication dairy. Follow-up interview by questionnaire at 12 months Reviewer comments: Not completely clear who measured outcomes and how, and whether questionnaires were piloted ITT analysis not performed Source of funding: Supported by grant No. M01-RR-00059 from the General Clinical Research Centre Program,, Division of Research Resources, National Institute of Health; |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|-------------|--|
| | Level | | S | | Measures | | the Children's Miracle Network Telethon and the Spelman-Rockefeller Child and Parenting Seed Grant |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|---------------------|---|
| | Level | | s | | Measures | | |
| Sunic-Omejc et | Study Type: | 49 children | 49 children | Intervention: | Duration of | Therapeutic success | Additional information from study: |
| al. Efficiency of | Parallel-RCT | | | Conventional | <u>treatment</u> | (number of children | Treatment considered successful if a |
| biofeedback | | <u>Inclusion</u> | 27 male | treatment (CON) | 12 weeks | cured) | frequency of ≥ 3 stools /week and < 2 |
| therapy for | Evidence | criteria: | | | | | episodes of soling or encopresis per |
| chronic | level: | Children aged | | Per oral | Assessment | -CON: 15/24 (62.5%) | month were achieved without laxatives |
| constipation in | 1+ | >5 years who | | administration of | point (s): | | |
| children. 2002. | | met at least 2 | 94 ± 33 months | Portalak | At 12 weeks | -BFB: 21/25 (84%) | Therapeutic success evaluated by use |
| Collegium | | of the | | (lactulosis, 240 | | | of questionnaires distributed on weekly |
| Antropologicum | To asses the | following | Mean age | mg/day or 10 mL | Follow-up | P<0.05 | visits |
| 26 Suppl, 93- | success of | criteria fro | (BFB): | syrup) with dose | period: | | |
| 101 | biofeedback | chronic | 92 ± 35 months | titration for the | None | | No significant differences in baseline |
| | method vs. | constipation: | | patient to have at | | | characteristics between 2 groups |
| | conventional | defecation | Country: | least 3 | <u>Outcome</u> | | |
| | method in the | frequency < 3 | Croatia | stools/week. | Measures: | | All children completed treatment |
| | | times/week, ≥ | | When | | | |
| | | 2 episodes of | | spontaneous | Therapeutic | | Reviewer comments: |
| | constipation in | | | defecation failed | success | | Small sample size, no sample size |
| | | encopresis | | to occur for > 3 | | | calculation |
| | over a 12- | /week, | | days in spite of | | | |
| | | periodic | | appropriate | | | Randomisation and allocation |
| | | evacuation of | | therapy an enema | | | concealment methods not described |
| | up the effect | large volume | | was used. In | | | |
| | of | stools at least | | addition a fibre- | | | Insufficient details on who measured |
| | biofeedback | once every 7 | | rich diet and | | | outcomes and how |
| | | to 30 days | | attempting | | | |
| | defecation | and palpable | | defecation after | | | Results not controlled for potential |
| | dynamics and | abdominal or | | meal were | | | confounders |
| | other | faecal mass | | advised | | | |
| | anorectal | | | | | | Source of funding: |
| | manometric | <u>Exclusion</u> | | Comparison: | | | Not stated |
| | parameters | <u>criteria</u> : | | Conventional | | | |
| | | Hirschsprung' | | treatment (CON, | | | |
| | | s disease, | | as previous) + | | | |
| | | spina bifida, | | Biofeedback | | | |
| | | hypothyroidis | | (BFB) | | | |
| | | m, metabolic | | | | | |
| | | or renal | | Pressure | | | |
| | | disorders, | | technique. | | | |

| Bibliographic S Information | tudy Type & Evidence Level | Number of Patients | Patient Characteristic s | | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|-----------------------------|----------------------------------|--------------------------------------|--------------------------------|--|------------------------------------|-------------|-------------------|
| | | mental retardation, taking drugs for | | Child and parents instructed on how to perform Kegel exercises at home. Exercises include alternating 10-second contraction and relaxation of sphincter and pubo-rectal muscle, performed 5 times a day in 20 cycles | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|---------------------|---|
| IIIIOIIIIatioii | Level | ratients | S | Companison | Measures | | |
| van der Plas et | Study Type: | 192 children | 192 children | Intervention: | Duration of | Treatment success | Additional information from study: |
| al. Biofeedback | Parallel-RCT | | | Conventional | intervention | (number of children | A faecal mass defined as a large hard or |
| training in | | <u>Inclusion</u> | 126 boys | laxative treatment | 6 weeks | cured, %) | soft stool in the rectum which completely |
| treatment of | Evidence | criteria: | | (CT) | | | filled the rectal vault. Soiling defined as |
| childhood | level: | Patients with | -age range | | Assessment | -at 6 weeks | loss of loose stools in underwear. |
| constipation: a | 1+ | paediatric | (total | 5 outpatient | point (s) and | CT (n=94): | Encopresis defined as voluntary or |
| randomised | | constipation | population): 5 to | visits lasting | follow-up | 31/94 (33%) | involuntary passage of a quantitatively |
| controlled | Study aim: | who fulfilled | 16 years | approximately 30 | period: | | normal bowel movement in underwear in |
| study. 1996. | | at least 2 of | | min during which | after the last | CT+BF (n=98): | children over the age of |
| | | these 4 | -median age for | laxative | visit of the | 31/98 (32%) | 4, occurring on a regular basis without |
| | biofeedback | criteria: stool | both groups: 8 | treatment and | intervention | | any organic cause. A large amount of |
| 780 | | frequency <3 | years | information from a | | NS | stool was estimated to be twice the |
| | | per week, ≥2 | | | weeks, then at | | standard shown in a clay model |
| | | | Country: | | 6 months, 1 | -at 6 months | |
| | | encopresis | The | frequency and | year, and 1 1/2 | CT (n=94): | High percentage of non compliance |
| | dynamics and | episodes per | Netherlands | | years | 48/93 (52%) | reported by parents if the child was |
| | outcome in | week, | | soiling episodes | | | asked to attempt toilet training 15–30 |
| | chronically | periodic | | were discussed | <u>Outcome</u> | CT+BF (n=98): | min after the meal to profit from the |
| | ı ı | passage of | | | Measures: | 44/94 (47%) | gastro—colic reflex |
| | children | very large | | High-fibre diet | | | |
| | | amounts of | | advised but | Treatment | NS | Treatment was considered successful if |
| | | stool at least | | | success | | the patients achieved ≥3 bowel |
| | | once every 7- | | supplements | | -at 1 year | movements per week and < 2 soiling or |
| | | 30 days, or a | | not prescribed | | CT (n=94): | encopresis episodes per month while |
| | | palpable | | | | 54/92 (59%) | not receiving laxatives for 4 weeks |
| | | abdominal or | | Patients | | | |
| | | rectal mass. | | instructed to try to | | CT+BF (n=98): | It was estimated that a sample of 180 |
| | | Children | | defecate on the | | 46/92 (50%) | patients would be adequate to show a |
| | | needed to be | | toilet for 5 min | | | difference of at least 70% success at 6 |
| | | at least 5 | | immediately after | | NS | months for CT+BF compared to 45% |
| | | years | | each meal | | | success using CT with a two-tailed alfa |
| | | old to | | | | -at 1 ½ year | 2 of 0.05 with a power of 90% |
| | | understand | | During the first 3 | | CT (n=94): | |
| | | the | | days patients | | 52/92 (57%) | At baseline patients were comparable |
| | | manometric | | were to use daily | | OT DE () | for gender, age, and frequency of |
| | | procedures | | enemas (120 mL | | CT+BF (n=98): | gastrointestinal complaints, and urinary |
| | | and | | sodiumdioctylsulfo | | 44/92 (48%) | problems |
| | | instructions | | succinate, 1 mg | | | |

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|---------------|-------------------|--------------------|------------------|-------------------------------------|---------------------|-------------|--|
| Information | Evidence Level | Patients | Characteristic s | Comparison | Outcome Measures | | |
| | 2010. | and had to | | sorbitol, 250 mg | cubui co | | At 6 months, 5 patients were lost (4 |
| | | have had | | per mL, Klyx) at | | | patients in the CT+BF and 1 patient in |
| | | treatment with | | home. If, on day | | | the CT group), and at 1 year 8 patients |
| | | laxatives for a | | 3, enemas still | | | were lost to follow up (another 2 in the |
| | | minimum of 1 | | resulted in large | | | CT+BF and 1 in the CT group). Patients |
| | | month before | | amounts of stool, | | | lost to follow up were withdrawn from |
| | | randomisation | | enemas were | | | further analysis |
| | | | | continued for a | | | |
| | | Exclusion | | maximum of 7 | | | During the intervention period, 3 patients |
| | | criteria: | | days. After the | | | in the CT group refused manometry at |
| | | Hirschsprung' | | initial 3-day | | | the end of the treatment period: 1 |
| | | s disease, | | enema treatment, | | | patient was successfully treated and the |
| | | spina bifida | | patients started | | | parents refused permission for |
| | | occulta, | | oral laxatives with | | | manometry; 1 patient was |
| | | hypothyroidis | | Importal (lactitol | | | unsuccessfully treated and refused |
| | | m or other | | betagalactoside | | | manometry; and 1 patient was lost to |
| | | metabolic or renal | | sorbitol, 1 sachet | | | follow-up after two visits. 2 patients of |
| | | abnormalities, | | of 5 g/10 kg body weight per day | | | the CT+BF group discontinued treatment: one 5-year-old patient did not |
| | | mental | | divided in 2 | | | cooperate and another patient |
| | | retardation, | | doses). Enemas | | | discontinued treatment because his |
| | | and | | given whenever | | | parents could not afford the cost of |
| | | children using | | spontaneous | | | transport. |
| | | drugs | | defaecation was | | | transport. |
| | | influencing | | delayed for more | | | At the beginning and end of the 6-week |
| | | gastrointestin | | than three days. | | | treatment period, each patient had a |
| | | al function | | Motivation | | | detailed medical history, abdominal and |
| | | other | | enhanced by | | | rectal examination, and anorectal |
| | | than laxatives | | praise and small | | | manometry. The child and parents were |
| | | | | gifts | | | asked about bowel function, frequency |
| | | | | 3 | | | of defaecation soiling and/or encopresis, |
| | | | | Comparison: | | | consistency and size of stool, pain |
| | | | | | | | during defaecation, and associated |
| | | | | 5 outpatient visits, | | | symptoms such as abdominal pain, |
| | | | | including the | | | appetite, and enuresis. Follow up |
| | | | | same | | | done either during a clinical visit using a |
| | | | | conventional | | | standard questionnaire or by telephone |
| | | | | treatment as | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---|------------------------------------|-------------|--|
| | | | | described above, in combination with 5 biofeedback training sessions. As far as possible, both groups received equal attention. | | | Because other studies have selected patients for evaluation according to the presence of abnormal defaecation dynamics at the start of the study, authors compared defaecation dynamics at randomisation and after treatment, and found no correlation between achievement of normal defaecation dynamics and success. Analysis of all patients showed no relationship between post-treatment defaecation dynamics and success. Log-linear modelling showed significant relationships between pre-treatment and post-treatment defaecation dynamics (x2= 13·91, p<0·001) and between treatment and post-treatment defaecation dynamics (x2=28·38, p<0·001). There was no association between post-treatment defaecation dynamics and treatment success after 6 weeks (x2=2·41, p=0·12). The results at 6 months and 1 year were similar Reviewer comments: Randomisation and allocation concealment methods not reported Not completely clear who measured outcomes and how ITT analysis not performed Source of funding: Not stated |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|------------------------|-----------------------|--|
| intornation | Level | i dilonio | S | Companicon | Measures | | |
| Nolan et al. | Study Type: | 29 children | 29 children | Intervention: | Duration of | Treatment outcome | Additional information from study: |
| Randomised | Parallel-RCT | | | EMG biofeedback | | | Originally, it was planned to recruit 25 |
| controlled trial | | <u>Inclusion</u> | 24 boys | training and | CT: Unclear | -Full remission: | subjects into each group, which would |
| of biofeedback | <u>Evidence</u> | criteria: | | conventional | | BFT+CT (n=14): | mean that, at the alfa = 0.05 level (one |
| training in | <u>level:</u> | Children aged | age range: 4.8 | medical treatment | BFT: up to 4 | 2 (14%) | tailed), there would be 80% power to |
| persistent | 1+ | ≥4 years, | to 14.9 years | (BFT+CT) | weeks | | detect at least a 38% point advantage of |
| encopresis with | | judged to be | | | | CT (n=15): | biofeedback (32% against 70% or |
| anismus. 1998. | Study aim: | of adequate | -mean age | | <u>Assessment</u> | 2 (13%) | better) in the comparison group. An |
| Archives of | To determine | maturity to | (years) (SD): | at weekly intervals | point (s): | | interim analysis conducted when it |
| Disease in | whether | cooperate | | conducted for | 6 months | 95% CI on difference, | became clear that successful and |
| Childhood | surface | | BFT+CT: | each patient, each | | -24% to 26% | sustained biofeedback outcomes were |
| 79[2], 131- | electromyogra | | 9.2 (2.7) | session consisting | | | not occurring. A revised sample size |
| | | treatment and | | of ~ 30–35 | period: | -Improved: | calculation was based on argument that |
| Kingdom. | | | CT: | defecation | None | BFT+CT (n=14): | if no successful outcomes were to be |
| | training | | 8.4 (2.3) | attempts. Aim was | | 2 (14%) | achieved in 15 subjects |
| | produces | more of | | to achieve 10 | <u>Outcome</u> | | randomised to biofeedback, there would |
| | sustained | conventional | Country: | relaxations of the | Measures: | CT (n=15): | be a 95% confidence that the true rate |
| | faecal | multimodal | Australia | external anal | | 4 (27%) | of successful outcome could not be |
| | | therapy; had | | | Treatment | | greater than 18%. The precision of the |
| | medical | continuing | | visual feedback in | success | p = 0.7; 95%Cl on | final result was expressed in the |
| | treatment | soiling with or | | 2 successive | | difference, −46% to | confidence interval (CI) around the |
| | resistant | without | | sessions. | | 23% (for remission | difference in remission rates |
| | and/or | laxative | | If this occurred in | | and improvement | |
| | treatment | treatment | | less than 4 | | combined) | Procedure to determine whether |
| | | (more than | | sessions then | | | anismus was present involved the use of |
| | children with | once a | | biofeedback was | | -No improvement: | a balloon filled with 50 ml warm water. |
| | anismus | month) or had | | discontinued. | | BFT+CT (n=14): | After a tuition period to explain what was |
| | | achieved | | At completion of | | 10 (71%) | required to achieve correct straining and |
| | | remission | | training, subjects | | , | squeezing, patient asked to make 5 |
| | | from soiling | | followed at | | CT (n=15): | alternating attempts each to squeeze |
| | | but could not | | monthly intervals | | 9 (60%) | and strain. Normal strain response |
| | | sustain | | by | | | defined as a persistent decrease in |
| | | continence | | a single | | 3/14 patients in the | external anal sphincter activity |
| | | without | | paediatrician, who | | BFT group completed | (measured by a decrease in amplitude |
| | | continued | | gave verbal | | the training in 3 | of the electromyographic recording and |
| | | laxative | | reinforcement of | | sessions, and the | an increase in rectal pressure of at least |
| | | treatment; | | the skills learned | | | 50 mm Hg) in at least 3 of 5 attempts. A |
| | | and had | | during training | | 4 sessions. Only 1 | persistent increase in external anal |

| anismus on EMG during anorectal manometry anorectal anorectal manometry anorectal manometry anorectal manometry anorectal anor | Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|--|---------------------------|-----------------------------------|--|--------------------------------|---|------------------------------------|--|--|
| bisacodyl tablets. Medication use decreased to a level consistent with bisacodyl tablets. at least one level from baseline status, but without achieving full remission Presence or absence of continued soiling ascertained on the basis of parental report, assisted by daily diary | | | EMG during anorectal manometry Exclusion criteria: known structural congenital or postoperative anatomical defect (such as spina bifida or anorectal malformation), or Hirschsprung's disease (excluded by rectal biopsy only if clinically | | Conventional medical treatment alone (CT) -Laxative therapy in 2 phases: 1. Initial disimpaction phase: 3-day cycles of 5 mL 'Microlax' enemas (sodium citrate) on day 1, one 5 mg bisacodyl tablet after school and 1 in evening of day 2. Up to 4 cycles (12 days) undertaken. Further cycles prescribed if later evidence of stool reaccumulation 2. Maintenance phase: liquid paraffin 5 to 30 ml once or twice a day, senna granules and or bisacodyl tablets. Medication use decreased to a level consistent | | demonstrate relaxation of the external anal sphincter with attempted defecation. Only 1 patient (same one) was unable to defecate the biofeedback balloon by the time of their final session. All complied well with instructions and procedures involved in the training. 2 complained of transient discomfort when the biofeedback apparatus was inserted. No other adverse effects seen | increase in rectal pressure in at least four of five attempts were deemed as indicating anismus Randomisation carried out using a stratified, blocked schedule, with subjects stratified on the basis of whether they were soiling or were in laxative dependent remission. Each treatment allocation was recorded on a card in an opaque numbered and sealed envelope and stored sequentially. An individual not connected with the clinic or the study carried out the randomisation plan Full remission defined as no medication and no soiling for at least 4 weeks; full remission on medication was defined as on medication and no soiling for at least 4 weeks; partial remission defined as soiling no more than once a week, regardless of medication used. The use of medication was attempted by all those not in full remission, not only those who were worse or not improved. The remainder were those who were soiling more than once a week, regardless of medication use. Improvement defined as progression by at least one level from baseline status, but without achieving full remission Presence or absence of continued soiling ascertained on the basis of |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|-------------|---|
| | Level | | S | maintenance of | Measures | | record. Patient data recorded |
| | | | | continence as | | | prospectively in a relational database |
| | | | | monitored by | | | was also used for appointment |
| | | | | bowel | | | scheduling and data quality control |
| | | | | diary | | | Some daming and data quality control |
| | | | | ulary | | | At baseline there were slightly more |
| | | | | -Standard | | | subjects with primary encopresis in the |
| | | | | paediatric | | | biofeedback group than in the control |
| | | | | behaviour | | | group |
| | | | | modification: | | | 3 - 1 |
| | | | | clarification during | | | Reviewer comments: |
| | | | | joint parent-child | | | No definition of constipation given |
| | | | | interview of the | | | |
| | | | | postulates | | | Small sample size |
| | | | | underlying | | | · |
| | | | | physiological | | | Unclear how the use of medication was |
| | | | | basis for | | | measured |
| | | | | encopresis. Bowel | | | |
| | | | | training | | | No dropouts/lost to follow up reported |
| | | | | programme used | | | |
| | | | | positive | | | Results not controlled for potential |
| | | | | reinforcement for | | | confounders |
| | | | | successful | | | |
| | | | | defection in toilet | | | Source of funding: |
| | | | | and additional | | | grants from the National Health and |
| | | | | reinforcement for | | | Medical Research Council (grant |
| | | | | each 24h without | | | 910621) and the Royal Children's |
| | | | | soiling. | | | Hospital Research Foundation |
| | | | | Reinforcement | | | |
| | | | | consisted of | | | |
| | | | | parental praise | | | |
| | | | | and use of start- | | | |
| | | | | chart diary (fitness | | | |
| | | | | training card) to | | | |
| | | | | indicate soiling- | | | |
| | | | | free days. Regular | | | |
| | | | | sitting programme | | | |
| | | | | of 5 to 10 minutes | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---|------------------------------------|-------------|-------------------|
| | | | | toilet-time within 30 minutes of each meal was basis of the programme. -Dietary advice, general counselling and support provided by paediatrician. Psychiatric assessment or treatment initiated when indicated clinically | | | |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|---|---|--|---|--|--|---|
| Loening-Baucke. Biofeedback treatment for chronic constipation and encopresis in childhood: long-term outcome. 1995. Pediatrics 96[1 Pt 1], 105-110 | Study Type: Retrospective cohort Evidence level: 2+ Study aim: To evaluate if patients who received biofeedback treatment (BF) continued with improved outcome compared | 129 children Inclusion criteria: Children 5 to 18 years with chronic constipation and encopresis (≥1 soiling episode per week) Exclusion criteria: Hirschsprung's disease, hypothyroidis m, mental deficiency, chronic debilitating diseases, neurologic abnormalities, previous surgery of the colon | s 129 children 97 boys Mean age (years): -CT group Initial: 9.1 ± 3.3 Follow-up: 13.4 ± 3.3 -BF group Initial: 10.4 ± 3.2 Follow-up: 14.5 ± 3.3 Country: USA | to 6 weekly training sessions given. 1 session included approximately 30 to 35 defecation trials and lasted approximately 45 to 60 minutes. Number of training sessions given depended on how soon child learned to relax external sphincter. Sessions stopped after 10 relaxations of the external sphincter without visual feedback could be accomplished in each of 2 successive training sessions Comparison: Conventional | Duration of treatment BF: between 2 and 6 weeks CT: unclear Follow-up period: -CT group: 4.2 ± 2.5 years -BF group: 4.1 ± 2.4 years Outcome Measures: -stool frequency -presence of soiling -soiling frequency | Stool frequency/week (mean ± SD) BF (n=63): 5 ± 3 CT (n=66): 6 ± 3 N.S % of children soiling BF (n=63): 35 CT (n=66): 24 N.S Soiling frequency/week (mean ± SD) BF (n=63): 1 ± 2 CT (n=66): 1 ± 2 N.S Recovery rate (number of children, %) BF (n=63): 28 (44) CT (n=66): 41 (62) N.S Laxative use (% children using laxatives) BF (n=63): 25 DE (n=63): 25 | Additional information from study: Parents and children instructed to keep diary of bowel movements, faecal soiling and medication used Of 64 patients who originally received biofeedback 1 patient did not return after the first unsuccessful biofeedback session and was lost to follow-up. The 63 patients included in the biofeedback group were combined from 2 studies (clinical characteristics of both groups were similar): 21 patients from an RCT (included already in this review, see Loening-Baucke, 1990) and 42 patients who had not recovered after at least 6 months of conventional treatment. Patients were charged for this service. Because of cost, inability to return for weekly biofeedback training or parent's and children's satisfaction with the marked improvement of constipation and encopresis with conventional treatment. 23 patients have been originally included in the RCT but 1 boy was lost to follow-up after the first biofeedback session and a second patient received a central nervous system shunt during the follow-up period and was exclude from analysis In May 1993 parents requested by email |
| | | | | treatment alone (CT) CT: use of | | CT (n=66): 18 N.S | to fill out with the help of their children a structured questionnaire eliciting information on the presence of soiling and frequency and amount of soiling per |

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|---------------|-------------------|-----------|------------------|--------------------------------|---------------------|-------------|---|
| Information | Evidence Level | Patients | Characteristic s | Comparison | Outcome Measures | | |
| | | | - | laxatives, increase of dietary | | | week, the frequency and size of bowel movements per week and the use of |
| | | | | fibre and scheduled | | | laxatives. In December 1993 |
| | | | | toileting (child | | | questionnaires again were mailed to non responders and to those families |
| | | | | instructed to | | | evaluated between January and May |
| | | | | defecate fro 5 | | | 1993. non responders were contacted |
| | | | | minutes after each meal and | | | by telephone |
| | | | | after returning | | | Patients considered to have recovered if |
| | | | | from school for | | | they had ≥3 bowel movements/week |
| | | | | the initial months, and try to | | | and soiling ≤ 2 episodes/month while off laxatives for at least 1 month. Patients |
| | | | | defecate at least | | | considered not to have recovered if they |
| | | | | daily once they | | | had <3 bowel movements/week or were |
| | | | | could recognise | | | soiling >2 times/month or had been |
| | | | | the urge to defecate | | | started on a regime of laxatives again |
| | | | | derecate | | | Baseline characteristics were |
| | | | | Disimpaction with | | | comparable between both groups |
| | | | | enemas (type and | | | except for the presence of an abdominal |
| | | | | dose not reported) | | | faecal mass (number of children, BF: 60 vs. CT: 41; p<0.05) |
| | | | | Maintenance: milk | | | νοι στι τι, ρ τοισογ |
| | | | | of magnesia ~ | | | Age and follow-up age were not related |
| | | | | 2ml/kg body weight daily to | | | to outcome in either group. The length of follow-up was significantly related to |
| | | | | induce at least 1 | | | recovery for the biofeedback group |
| | | | | bowel movement | | | (p<0.02) and for all patients (p<0.01) but |
| | | | | daily and prevent | | | showed no relationship for the |
| | | | | faecal retention. | | | conventionally treated group |
| | | | | Doses decreased gradually to | | | Reviewer comments: |
| | | | | maintain daily | | | No clear definition of constipation given |
| | | | | bowel movement | | | |
| | | | | and prevent | | | Supported by great No. M01 BB 00050 |
| | | | | faecal retention and soiling. | | | Supported by grant No. M01-RR-00059 from the General Clinical Research |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|--|------------------------------------|-------------|---|
| | | | | Occasionally mineral oil or senna used instead of milk of magnesia | | | Centre Program,, Division of Research Resources, National Institute of Health; the Children's Miracle Network Telethon and the Spelman-Rockefeller Child and Parenting Seed Grant |
| | | | | | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|-------------------------------------|---------------------|----------------------|---|
| | Level | | S | | Measures | | |
| Silver et al. | Study Type: | 108 children | 108 children | Intervention: | Duration of | EXT (n=54) | Additional information from study: |
| Family therapy | Retrospective | and their | | Externalizing | treatment | OTH (n=54) | 162 sets of notes of all referrals for |
| and soiling: An | audit | families | 3 to 5 years: 45 | Treatment (EXT) | (mean, months) | | soiling over a four-year period were |
| audit of | | | >6 years: 63 | | -EXT: 7.8 | Not all children | audited |
| externalizing | <u>Evidence</u> | <u>Inclusion</u> | | Families were | -OTH: 6.6 | assessed for all | |
| and other | level: | | mean age | only included if | | outcomes | Some children clearly diagnosed in the |
| approaches. | 3 | Children | (years): | the approach | <u>Assessment</u> | | referral letter as 'constipated' or 'not |
| 1998. Journal of | | treated for | -EXT: 6.98 | included: | point (s) & | | constipated', but in some referral letters |
| Family Therapy | Study aim: | soiling | -OTH: 6.68 | | | | it was not stated whether the referring |
| 20[4], 413-422 | To asses the | problems. | | 1 Externalizing | At a minimum | parents) | doctor had checked for constipation |
| | effectiveness | Referrals | Country: | | of 6 months | -EXT: | |
| | of | included | UK | first interview with | | Helpful: 24 | The treatment given depended only on |
| | Externalizing | 'faecal | | the child and | months) after | Unhelpful: 5 | the current approach of the therapist |
| | Treatment | soiling', | | J (| treatment | | who received the referral. All the families |
| | EXT) as | 'encopresis', | | | Ended | -OTH: | had received either 'externalizing' or |
| | compared to | 'psychological | | White and Epston, | | Helpful: 10 | 'other treatments' |
| | traditional | soiling', 'failed | | 1990) | <u>Outcome</u> | Unhelpful: 20 | |
| | treatments in | toileting', | | | Measures: | | No significant differences between the |
| | children with | 'constipation | | 2 Developing a | -Parent | p = 0.0001 | groups on baseline variables |
| | soiling | with overflow' | | | assessment of | | |
| | problems | and | | child and family | | End of treatment | At a minimum of 6 months' follow-up |
| | | 'deliberate | | · · · · · · · · · · · · · · · · · · | treatment | outcome (from notes) | (mean 23 months), all parents (including |
| | | soiling'. | | see themselves | | -EXT: | those who dropped out) sent a |
| | | | | as capable, skilful | | No soiling/improved: | questionnaire |
| | | <u>Exclusion</u> | | and determined to | | 42 | with a letter from the secretary, |
| | | criteria: | | | /frequency | Soiling: 5 | explaining that we could learn a great |
| | | Families who | | lesson, outwit the | (parents' | | deal from their responses, whether |
| | | failed to | | poo or defeat the | assessment | -OTH: | negative or positive, with no names |
| | | attend or | | poo | | No soiling/improved: | being recorded. Parents asked whether |
| | | cancelled | | | | 30 | there had been any further soiling |
| | | their first | | 3 Not using | | Soiling: 13 | incidents since they were last seen and |
| | | appointment, | | , | notes | | frequency of these incidents in the past |
| | | the problem | | interpretation, | | p = 0.02 | month. Parents asked whether they had |
| | | had been | | confrontation or | -Number of | | found their treatment helpful or unhelpful |
| | | resolved, the | | paradoxical | appointments | GP follow-up | and what was helpful or unhelpful and to |
| | | children were | | interventions as | | -EXT: | offer other comments. Where children |
| | | put into care | | therapeutic | | No soiling: 29 | had returned for paediatric consultation, |
| | | or sent to | | manoeuvres. | | Soiling: 8 | frequency of soiling stated in paediatric |

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|---------------|--------------|---------------------------|----------------|---------------------------|-------------|-------------------------|--|
| Information | Evidence | Patients | Characteristic | Comparison | Outcome | | |
| | Level | | S | | Measures | | |
| | | boarding | | 4 Attempting to | | | notes was recorded even if parents did |
| | | school very | | see the whole | | -OTH: | not reply to the audit. GPs asked |
| | | early in | | family at least | | No soiling: 24 | whether they were aware of any further |
| | | treatment or | | once. | | Soiling: 18 | soiling after treatment had ended |
| | | the soiling | | | | | |
| | | had a medical | | Comparison: | | p = 0.045 | Reviewer comments: |
| | | cause | | Other Treatments | | Parent follow-up | No definition of constipation given |
| | | (Hirschsprung | | (OTH) | | -EXT: | |
| | | 's disease). | | | | No soiling/stains: 24 | Unclear exactly how many children |
| | | Children who | | Mixed group of | | Soiling: 14 | dropped out/ were lost to follow up |
| | | had full | | traditional | | OTU. | Course of fundings |
| | | control, but | | treatments with | | -OTH: | Source of funding: Not stated |
| | | would insist | | predominantly | | No soiling/stains: 13 | inot stated |
| | | on a nappy for a bowel | | (but not only) a | | Soiling: 22 | |
| | | movement. | | behavioural approach in a | | p = 0.026 | |
| | | 3 more | | family systems | | p = 0.026 | |
| | | families | | context. There | | Number of | |
| | | where a | | were no elements | | appointments (mean) | |
| | | therapist | | of externalizing in | | appointments (mean) | |
| | | who usually | | any | | -EXT: 8.2 | |
| | | used | | OTH sessions | | -OTH: 10 | |
| | | externalizing | | 0 0000.0 | | NS | |
| | | switched to a | | | | | |
| | | behavioural | | | | Externalizing proved | |
| | | approach in a | | | | to | |
| | | systems | | | | be superior for boys, | |
| | | context in the | | | | for children aged ≥ 6 | |
| | | belief that | | | | years, for those with | |
| | | externalizing | | | | frequent soiling at the | |
| | | would not | | | | outset, for those with | |
| | | work. Within | | | | over 2 years' | |
| | | the remaining | | | | continuous soiling | |
| | | families in the | | | | and those diagnosed | |
| | | audit there | | | | as constipated on | |
| | | was no | | | | referral | |
| | | known | | | | | |
| | | selection for a | | | | | |

| Bibliographic Information | Study Type & Evidence Level | Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|-------------|-------------------|
| | | particular therapy | | | | | |
| | | шетару | | | | | |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|------------------------|--|
| | Level | | S | | Measures | | |
| Taitz et al. | Study Type: | 47 children | 47 children | General | Duration of | Treatment success | Additional information from study: |
| Factors | Quasi-RCT | | | In cases where | | did not differed | One year after the beginning of |
| associated with | | <u>Inclusion</u> | 26 boys | constipation was | -BhM: 6 weekly | between both groups. | treatment parents sent a postal |
| outcome in | <u>Evidence</u> | criteria: | | severe with large | intervals for | | questionnaire, which sought to elicit the |
| management of | level: | children who | age not | faecal masses | | It is not possible to | response to treatment. This survey |
| defecation | 1+ | presented | reported | children initially | months | report the figures | included all patients who 'dropped out' of |
| disorders. 1986. | | with faecal | | admitted to the | and 1 year | here, as they were | this study at any stage. They were |
| Archives of | Study aim: | soiling, with | Country: | ward for | | only analysed by the | asked whether they considered the child |
| Disease in | To report our | or without | UK | defecation was | <u>Assessment</u> | authors according to | cured, improved, or unchanged and |
| Childhood | experience | constipation | | made impossible | point (s): | compliance with | asked how often the child defecated; |
| 61[5], 472-477 | with children | | | by severe | 1 year after | treatment and with | whether and how often soiling occurred; |
| | who | Exclusion | | | initiating | children social class, | and whether and how often laxatives |
| | presented | criteria: | | were then | | but not according to | were needed. These answers were |
| | with faecal | identified | | continued on | | treatment groups | made as objective as possible by |
| | soiling, with or | | | | Follow-up | | requesting parents to place ticks in |
| | without | disease or | | | period: | | appropriate boxes. This response was |
| | | neurological | | before referral. | None | | then graded into three categories-cured, |
| | who were | handicaps | | Where no laxative | | | improved, and no response, on the |
| | treated by | | | had previously | <u>Outcome</u> | | basis of the parents' answers to the |
| | incentive | | | been used the | Measures: | | questionnaire, compared with the clinical |
| | based | | | child was offered | | | assessment before allocation to |
| | behavioural | | | a twice | Treatment | | treatment groups. Assessment of results |
| | modification, | | | daily dose of | success | | were thus made by the parents at home |
| | plus or minus | | | lactulose. If no | | | and not by the professionals involved |
| | psychotherap | | | accumulation of | | | |
| | y, and | | | faeces no | | | Criteria for the classification of the |
| | consider | | | laxatives | | | results of treatment: |
| | factors that | | | prescribed. No | | | (1) Cured. At least 5 normal stools each |
| | might predict | | | other | | | week without soiling. Only occasional |
| | the outcome | | | laxatives used in | | | use of laxatives (less than once a week) |
| | for a non- | | | this study, and in | | | (2) Improved. At least three stools each |
| | intensive | | | general their | | | week and soiling less than once a week |
| | approach and | | | use was | | | (3) Non-responders. Less than three |
| | in particular, | | | minimised, with | | | stools each week or soiling more than |
| | to draw | | | the parents | | | once a week. These children were |
| | attention to | | | encouraged to | | | considered as failing to improve, despite |
| | social | | | stop the treatment | | | the fact that in most cases there was |
| | background | | | with laxatives as | | | less soiling than at the beginning of |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|------------------------|-------------|--|
| | Level | | S | | Measures | | |
| | as a | | | soon as a regular | | | treatment |
| | prognostic | | | bowel habit | | | |
| | indicator | | | established. In | | | 4 children dropped out from the study |
| | | | | none of the | | | and 13 failed to keep adequate 'star |
| | | | | children | | | charts'. The 'drop outs' occurred at 1, 2, |
| | | | | were | | | 3, and 4 months. 2 children were |
| | | | | suppositories | | | subsequently found to be cured |
| | | | | used at any time. | | | |
| | | | | All the children | | | Reviewer comments: |
| | | | | were encouraged | | | No definition of constipation given |
| | | | | to take a high | | | |
| | | | | residue diet and in | | | Small sample size, no sample size |
| | | | | particular were | | | calculation |
| | | | | asked to | | | |
| | | | | take bran with | | | Baseline characteristics not compared |
| | | | | their breakfast | | | |
| | | | | cereal | | | Randomisation and allocation |
| | | | | | | | concealment methods not reported |
| | | | | Intervention: | | | |
| | | | | Behaviour | | | ITT analysis not performed |
| | | | | modification | | | |
| | | | | (BhM) | | | Source of funding: |
| | | | | | | | Grants from the Hawley Trust, National |
| | | | | Carried out by | | | Health Service Locally Organised |
| | | | | paediatrician. All | | | Research Grant (Trent RHA) and |
| | | | | children placed on | | | CHRIS Fund, Children's Hospital |
| | | | | a star chart | | | |
| | | | | regimen. Children | | | |
| | | | | offered varying | | | |
| | | | | coloured stars for | | | |
| | | | | 'sitting on the | | | |
| | | | | toilet' and | | | |
| | | | | 'remaining | | | |
| | | | | unsoiled for a full | | | |
| | | | | day'. In | | | |
| | | | | some cases stars | | | |
| | | | | awarded to | | | |
| 1 | | | | encourage | | | |

| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
|---------------|--------------|-----------|----------------|---------------------------------|-------------|-------------|-------------------|
| Information | Evidence | Patients | Characteristic | Comparison | Outcome | | |
| | Level | | S | ala il alua sa | Measures | | |
| | | | | children | | | |
| | | | | who were | | | |
| | | | | reluctant to take | | | |
| | | | | bran in their diet. | | | |
| | | | | Contract | | | |
| | | | | negotiated between child and | | | |
| | | | | | | | |
| | | | | parent (usually | | | |
| | | | | father) for an award to be made | | | |
| | | | | at the discretion of | | | |
| | | | | the paediatrician. | | | |
| | | | | Child was to | | | |
| | | | | understand that | | | |
| | | | | the giving of the | | | |
| | | | | award would | | | |
| | | | | depend on | | | |
| | | | | response to | | | |
| | | | | treatment. | | | |
| | | | | 'Demystification', | | | |
| | | | | alleviation of guilt, | | | |
| | | | | and use of | | | |
| | | | | explanatory | | | |
| | | | | diagrams | | | |
| | | | | generally followed | | | |
| | | | | the lines | | | |
| | | | | recommended | | | |
| | | | | by Levine and | | | |
| | | | | Bakow. | | | |
| | | | | Children seen at 6 | | | |
| | | | | weekly intervals | | | |
| | | | | by paediatrician | | | |
| | | | | for between 3 | | | |
| | | | | months | | | |
| | | | | and 1 year and | | | |
| | | | | subjected to | | | |
| | | | | shows of affection | | | |
| | | | | and interest, | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------------|------------------------------------|-------------|-------------------|
| | | | | which included | | | |
| | | | | careful and | | | |
| | | | | serious inspection | | | |
| | | | | of the charts. | | | |
| | | | | Failure to keep a | | | |
| | | | | star chart on 2 | | | |
| | | | | successive visits | | | |
| | | | | resulted in firm statement of | | | |
| | | | | displeasure. 2 | | | |
| | | | | further failures at | | | |
| | | | | 6 | | | |
| | | | | week intervals led | | | |
| | | | | to the stopping of | | | |
| | | | | treatment and | | | |
| | | | | discharge with the | | | |
| | | | | option of | | | |
| | | | | psychiatric | | | |
| | | | | referral. | | | |
| | | | | Discharge of | | | |
| | | | | cured patients | | | |
| | | | | was at discretion | | | |
| | | | | of the parents | | | |
| | | | | Comparison: | | | |
| | | | | Behaviour | | | |
| | | | | modification (as | | | |
| | | | | previous) + | | | |
| | | | | psychotherapy | | | |
| | | | | (BhM +Psy) | | | |
| | | | | | | | |
| | | | | -Psychotherapy: | | | |
| | | | | children seen by | | | |
| | | | | the child | | | |
| | | | | psychiatrist at roughly monthly | | | |
| | | | | intervals for | | | |
| | | | | periods between | | | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|-------------|-------------------|
| | Level | | S | | Measures | | |
| | | | | two and 12 | | | |
| | | | | months. | | | |
| | | | | Treatment was | | | |
| | | | | organised along | | | |
| | | | | the following lines: | | | |
| | | | | (1) At each | | | |
| | | | | appointment | | | |
| | | | | mother (and also | | | |
| | | | | father in 4 cases) | | | |
| | | | | seen for 15-30 | | | |
| | | | | minutes to explore | | | |
| | | | | her feelings in | | | |
| | | | | respect of the | | | |
| | | | | child's bowel | | | |
| | | | | problem and its | | | |
| | | | | effect on the | | | |
| | | | | family and her | | | |
| | | | | own relationship | | | |
| | | | | with the child. | | | |
| | | | | Whenever | | | |
| | | | | possible mother's | | | |
| | | | | own history | | | |
| | | | | explored and | | | |
| | | | | other emotional | | | |
| | | | | problems | | | |
| | | | | discussed where | | | |
| | | | | relevant e.g. | | | |
| | | | | expressions of | | | |
| | | | | grief, anger, | | | |
| | | | | depression, etc. | | | |
| | | | | (2) Child seen for | | | |
| | | | | between 15-30 | | | |
| | | | | minutes for play, | | | |
| | | | | including picture | | | |
| | | | | drawing, games, | | | |
| | | | | and sharing of | | | |
| | | | | their own toys and | | | |
| | 1 | | | belongings. Their | | | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---|------------------------------------|-------------|-------------------|
| | | | | feelings concerning their problem also explored. Behavioural star chart also often brought, and reviewed and child praised and encouraged according to progress (3) Mother and child seen together sometimes early in treatment, sometimes later, depending on their relationship and success with management of the problems to assess to overall progress | | | |

Complementary Therapies for Ongoing Treatment/Maintenance in Children with Chronic Idiopathic Constipation

| Bibliographic Sinformation | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|---|--|--|--|--|---|---|
| Reflexology in the management of encopresis and chronic constipation. 2003. Paediatric Nursing 15[3], 20-21 | Prospective case series Evidence evel: Study aim: To investigate he efficacy of reating patients with | 50 children Inclusion criteria: Children diagnosed with encopresis / chronic constipation Exclusion criteria: Not stated | age range 3 to 14 years 64% boys Country: UK | Intervention: Reflexology: 6 sessions , 30 minutes each at weekly intervals (no other details provided) Comparison: N.A | point (s): Immediately after treatment was completed Follow-up period: No follow-up made after treatment finished | no soiling/week: 6 -After: at least daily: 20 1 to 3 times/week: 30 no soiling/week: 48 p<0.05 (unclear for which comparisons) Frequency of bowel movements (BM)(n=48) % children -Before: No BM/week: 36 1 to 4 BMs/week: 46 | Additional information from study: With the help of their parents, children completed questionnaires on bowel motions and soling patterns before, during and after treatment Parents completed questionnaires on their attitude towards reflexology Existing medications were unaltered 2 children only attended the first session Reviewer comments: No definition of constipation/encopresis given Questionnaire not reported as piloted Results not controlled for potential confounders Baseline outcomes for the 2 children who only attended the first session were reported but it is unclear whether they were included in the analysis Source of funding: Not stated |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | . Trought of | daily BMs: 24 p<0.05 (unclear for which comparisons) Parents' attitude towards reflexology 70% parents keen to try treatment, 72% satisfied with outcome | |

Surgical Interventions for Maintenance: Effectiveness of the ACE procedure in Children with Chronic Idiopathic Constipation

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|----------------------|--|
| King at al. The | Level | 50 abildes e | s 42 children | l-4 | Measures | AOE | Additional information forms at the |
| King et al. The | Study Type: | 56 children | 42 children | Intervention: | Follow-up period: | ACE usage | Additional information from study: |
| antegrade | Retrospective cohort | Inducion | 31 boys | appendicostomy (ACE): | Mean: 48 | a. ACE regimes | Independent investigator conducted confidential telephone interviews using a |
| continence enema | COHOIL | Inclusion criteria: | 31 boys | | months (median | a. ACE regimes | modified questionnaire |
| | Cvidonos | | maan aga at | laparoscopy or | | -median initial | modilled questionnaire |
| | <u>Evidence</u> | | mean age at | mini-laparotomy | 39, range 3 to | | Continuos consumo difical |
| | <u>level:</u> | | interview: 13.1 | C | 118) | regimes used (% | Continence score: modified |
| slow-transit | 2+ | | years (median | Comparison: none | | children): | Holschneider (maximum score 12). |
| constipation. | Ct. d. cias | | 12.4; range 6.9 | | <u>Outcome</u> | Cal. 4al. (70) | Modification required because the |
| 2005. Journal of | | • | to 25.0) | - | Measures: | Golytely (79) | criterion of "frequency of defecation" not |
| | to determine | formed | | Enemas: | A C = | Liquorice (12) | appropriate for the cohort |
| Surgery 40[12], | whether ACE | | mean age at | | -ACE usage | Water (2) | Overlite of life consequence difficult Terror letter |
| 1935-1940 | are successful | | procedure: 9.1 | -median initial | AOE -#: | Other (7) | Quality of life score: modified Templeton |
| | for idiopathic | | years (median | regimes used: | -ACE efficacy | | and Toogood |
| | paediatric | | 7.8, range 3.1 | O-1-4-1- (DEO | 405 | -outcome (% | |
| | slow transit | | to 18.5) | Golytely (PEG | -ACE | children): | Frequency score used for all frequency |
| | | criteria for | | 3350 and | complications | Excellent (29) | measures: daily=6, 3 to 6 d/wk=5, 1 to 2 |
| | (STC) | functional | -recurrent | electrolytes): 250 | | Good (36) | d/wk=4, 1 to 2 d/fortnight=3, 1 to 2 |
| | | | soiling: 29/42 | to 500 ml every | | Average (7) | d/mo=2, once every 2 to 3 months=1 |
| | | | (69%) | second day, | | Poor (28) | and never=0) |
| | | faecal | | infused over 20 to | | | |
| | | | -inability to | 30 mins for 1 to 3 | | -median regime at | Reviewer comments: |
| | | | adequately | months | | time of interview: | Originally 56 children met the inclusion |
| | | | pass stool: 7/42 | | | | criteria, but only 42 (75% of the families) |
| | | | (17%) | Liquorice , 250 to | | Golytely: (how many | were interviewed without a clear |
| | | period of | | 500 ml daily, | | , | explanation for that |
| | | | -recurrent | infused over 10 to | | occurred 20 to 30 | |
| | | | hospital | 20 mins infused | | mins after ACE | Source of funding: |
| | | • | admissions for | over 10 to 20 | | | Dr. King funded by scholarships from |
| | | | nasogatric | mins for 1 to 3 | | mins spent on toilet | the NHMRC (Australia) and the Royal |
| | | | washouts: 6/42 | months | | | Australian College of Surgeons |
| | | | (14%) | | | Majority of patients | |
| | | criteria: | | -median regime at | | (25/42, 60%) either | |
| | | not stated | | time of interview: | | using the initial | |
| | | | Country: | Golytely (PEG | | regime or had tried | |
| | | | Australia | 3350 and | | one regimen change. | |
| | | | | electrolytes): 500 | | No correlation | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---|------------------------------------|--|-------------------|
| | | | | to 750 ml every second day, infused over 10 to 20 mins with no need for disimpaction | | between numbers of ACE regimens tried, patient satisfaction or length of ACE usage. Many families believed regimes changes were a necessary response to increased tolerance to a particular ACE solution | |
| | | | | | | b. patient input into ACE regimen (n children) -completely independent: 7 (all older 10 years) -requiring supervision only: 5 -needing help setting up and cleaning up: 15 -completely dependent: 15 | |
| | | | | | | c. patients satisfaction with ACE (n children) -very satisfied or satisfied: 37 (88%) -families would recommend ACE to other children: 41 (98% | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | -families felt significant improvement in quality of child's life: 39 (93%) -mean optimal age for appendicostomy formation, as felt by families: 4.9 years | |
| | | | | | | (median 4, range 2 to 12) d. effectiveness | |
| | | | | | | -effective: 41 (98%) e. symptoms resolution (n patients) | |
| | | | | | | -ceased ACE: 15 (36%): in 7 symptoms resolved, in 4 a colostomy was formed, in 2 an ileostomy was formed and 2 patients returned to | |
| | | | | | | conservative management -successful ACE: 34 (81%) | |
| | | | | | | ACE efficacy (mean, median and range): -continence score: pre-ACE: 2.5 (2; 0 to 8) | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | | post-ACE:: 5.2 (5; 1 to 12) p<0.0001 | |
| | | | | | | -quality of life score: pre-ACE: 1.4 (1.5; 0.5 to 3.0) post-ACE: 2.2 (2.5; 0.5 to 3.0) p<0.0001 | |
| | | | | | | -soiling frequency score: pre-ACE: 5.7 (6; 0 to 6) post-ACE: 3.0 (3; 0 to 6) p<0.0001 | |
| | | | | | | -abdominal pain severity score: pre-ACE: 7.4 (8; 0 to 10) post-ACE: 3.0 (3; 0 to 8) p<0.0001 | |
| | | | | | | -abdominal pain frequency score: pre-ACE: 5 (6; 0-6 to 3-6 d/week) post-ACE: 2.5 (2.5; 0-6 to 1-2 d/month) p<0.0001 | |
| | | | | | | ACE complications: a. symptoms at some | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|--------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | stage of treatment: | |
| | | | | | | Total: 20/42 (710/) | |
| | | | | | | Total: 30/42 (71%) cramping: 18/30 | |
| | | | | | | nausea: 17/30 | |
| | | | | | | vomiting: 7/30 | |
| | | | | | | sweating: 14/30 | |
| | | | | | | dizziness: 10/30 | |
| | | | | | | pallor: 10/30 | |
| | | | | | | (3 or more symptoms | |
| | | | | | | present in 12/30 | |
| | | | | | | patients) | |
| | | | | | | | |
| | | | | | | b. Long-term | |
| | | | | | | complications (n, %), N=42: | |
| | | | | | | N=42. | |
| | | | | | | -granulation tissue: | |
| | | | | | | 33 (79), unresolved: | |
| | | | | | | 15% | |
| | | | | | | -anxiety about ACE: | |
| | | | | | | 21 (50), unresolved: 29% | |
| | | | | | | -stomal infection: 18 | |
| | | | | | | (43), unresolved: 11% | |
| | | | | | | -stomal leakage (ACE | |
| | | | | | | days): 16 (38), | |
| | | | | | | unresolved:13% | |
| | | | | | | -embarrassment | |
| | | | | | | about device: 16 (36), | |
| | | | | | | unresolved: 87% | |
| | | | | | | -dislikes device: 12 (29), unresolved: 58% | |
| | | | | | | -stomal leakage (non | |
| | | | | | | ACE days): 12 (29), | |
| | | | | | | unresolved: 8% | |
| | | | | | | -stomal pain: 11 (26), | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | unresolved: 45% -stomal stenosis: 8 (19), unresolved: 0 -new behavioural disturbance: 7 (17), unresolved: 72% -stomal prolapse: 6 (14), unresolved: 33% -stomal bleeding: 6 (14), unresolved: 0 -limited activity: 4 (10), unresolved: 75% -weight loss: 2 (5), unresolved: 0 -perforation: 2 (5), unresolved: 0 | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---|------------------------------------|-----------------------|--|
| Youssef et al. | Study Type: | 12 children | 12 children | Intervention: | Follow-up | Bowel | Additional information from study: |
| Management of | Retrospective | | 9 boys | Caecostomy | period: | movements/week | A questionnaire used to interview |
| intractable | case series | | mean age: 8.7 | (surgically and by | 13.5 ± 8.5 | before: 1.4 ± 0.7 | caregivers |
| constipation | | | ± 4.4 years | interventional | months | after: 7.1 ± 3.8 | 13.5 ± 8.5 months after caecostomy |
| with antegrade | <u>Evidence</u> | children | , | radiology) | | p<0.005 | placement. No caregiver refused to |
| enemas in | level: | | Country: USA | , | Outcome | F 101000 | participate in interview |
| neurologically | 3 | tertiary care | | Comparison: | Measures: | Soiling | |
| intact children. | | motility centre | | none | | episodes/week | Scoring for episodes of abdominal pain: |
| 2002. Journal of | Study aim: | for further | | | -Bowel | before: 4.7 ± 3.2 | 0 = none, 1=once or twice, 2=a few |
| Pediatric | | evaluation of | | | movements/we | after: 1.0 ± 1.4 | times, 3=fairly often, 4=very often, 5= |
| Gastroenterolog | benefit of | intractable | | Choice of | | p<0.01 | everyday |
| y and Nutrition | antegrade | constipation, | | irrigation solution | | ľ | |
| 34[4], 402-405 | colonic | who had | | used after | -Soiling | Number of | Scoring for overall health and |
| | enemas | undergone | | caecostomy | episodes/week | medications used for | emotional state: 1=poor, 2=fair, 3=good, |
| | through | caecostomy | | varied, based on | • | constipation | 4=very good, 5=excellent |
| | caecostomy | placement for | | preference of | -Number of | before: 4.0 ± 1.0 | |
| | catheters in | administration | | treating physician. | medications | after: 0.8 ± 0.6 | Reviewer comments: |
| | children with | of antegrade | | Most patients | | p<0.005 | Very small sample |
| | severe | enemas | | began with low | constipation | | · |
| | constipation | | | volume | · | Abdominal pain | Not clear who performed the review of |
| | who were | Exclusion | | infusions of | -Episodes of | score: | the clinical records |
| | referred to a | criteria: | | solution, which | abdominal | before: 2.9 ± 1.6 | |
| | tertiary care | neurologic | | were increased | pain/week | after: 0.9 ± 1.0 | Not clear who interviewed the parents |
| | centre | handicap and | | according | | p<0.005 | · |
| | | other organic | | to therapeutic | -Missed school | | Researchers not reported blinded |
| | | causes of | | response. 67% of | days/month | Missed school | · |
| | | constipation | | patients used 200 | | (days/month) | Questionnaire not reported |
| | | | | mL to 1,000 mL | -Emotional | before: 7.5 ± 6.9 | piloted/validated |
| | | | | (mean 478 mL ± | health | after: 1.5 ± 2.5 | |
| | | | | 262 mL) | | p<0.02 | Source of funding: |
| | | | | polyethylene | -Overall health | | not stated |
| | | | | glycol irrigation | | Emotional health | |
| | | | | solution, daily to | -Physician | <u>score</u> | |
| | | | | every other day. | office visits/year | before: 1.9 ± 0.8 | |
| | | | | 25% of patients | | after: 3.6 ±1.1 | |
| | | | | used a | | p<0.005 | |
| | | | | combination of | | | |
| | | | | saline and | | Overall health score: | |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|-----------------------|---------------------------|---------------------------------|---------------------|-------------------------|-------------------|
| | Level | | S | | Measures | | |
| | | | | glycerin, mixing | | before: 1.7 ± 0.9 | |
| | | | | 60 mL to 75 mL of | | after: 3.6 ± 0.9 | |
| | | | | glycerin in 240 mL | | p<0.005 | |
| | | | | to 300 mL of | | DI | |
| | | | | saline. 1 patient | | Physician office | |
| | | | | received 90 mL | | visits/year | |
| | | | | phosphate soda | | before: 24.0 ± 19.1 | |
| | | | | solution followed | | after: 9.2 ± 14.2 | |
| | | | | by 300 mL of saline. Evacuation | | p<0.05 | |
| | | | | occurred within 1 | | No acute adverse events | |
| | | | | hour | | Evento | |
| | | | | of enema | | Postoperative | |
| | | | | administration in 7 | | adverse events (n | |
| | | | | children and | | children): | |
| | | | | occurred within 3 | | -skin breakdown | |
| | | | | hours in the other | | and development of | |
| | | | | 5 children. | | granulation tissue: 1 | |
| | | | | o crinareri. | | -leakage of irrigation | |
| | | | | | | solution: 1 | |
| | | | | | | -accidental removal of | |
| | | | | | | the catheter with | |
| | | | | | | subsequent easy | |
| | | | | | | catheter replacement | |
| | | | | | | by the interventional | |
| | | | | | | radiologist: 2 | |
| | | | | | | radiologica = | |
| | | | | | | No adverse event led | |
| | | | | | | to discontinuation | |
| | | | | | | of antegrade enema | |
| | | | | | | use. | |
| | | | | | | No child has required | |
| | | | | | | admission to a | |
| | | | | | | hospital because of | |
| | | | | | | faecal impaction | |
| | | | | | | since starting | |
| | | | | | | antegrade enemas. | |
| | | | | | | 5 patients | |

| Information Ev | y Type & Number of idence Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|----------------|------------------------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | | | | | discontinued antegrade enemas with removal of the caecostomy at a mean of 14.6 ± 9.1 months after beginning treatment. None has redeveloped problems with constipation or faecal soiling. | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|----------------------------|------------------------------------|--|---|
| Cascio et al. | Study Type: | 49 children | 49 children | Intervention: | Follow-up | Soiling (n children in | Additional information from study: |
| MACE or | Retrospective | | 15 boys | Malone antegrade | period: | which stopped | One patient with CB and one with MACE |
| caecostomy | cohort | <u>Inclusion</u> | | enema (MACE) | Mean, 18 | completely) | moved to another region and were lost |
| button for | | criteria: | -MACE: | | months | | to follow-up |
| idiopathic | <u>Evidence</u> | children who | 37 children | Antegrade | | MACE (n=37): 30 | |
| constipation in | <u>level:</u> | underwent | 15 boys | enemas started | <u>Outcome</u> | (81%) | Success criteria: |
| children: a | 2+ | MACE or CB | | on the 4 th | Measures: | CB (12): 9 (75%) | -full: totally clean or minor or minor |
| comparison of | | between June | | postoperative day | | | rectal leakage on the night of the |
| complications | Study aim: | 1998 and | 12 children | and Foley | -Soiling | Occasional soiling still | |
| and outcomes. | to compare | August 2002 | 9 boys | catheter left in | | present in 1 child with | |
| 2004. Pediatric | the results | for intractable | | appendicostomy | -Failure | | rectal leakage, occasional major leak, |
| Surgery | | idiopathic | Country: UK | for 6 weeks | | 1 child with CB | still wearing protection but perceived by |
| International | and outcomes | | | | -Surgical | resumed regular | the child or parent to be an improvement |
| 20[7], 484-487 | | and faecal | | Comparison: | complications | activity and CB was | -failure: regular soiling or constipation |
| | antegrade | soiling that | | Caecostomy | | removed | persisted , no perceived improvements, |
| | enema | had failed | | button (CB) | | | procedure abandoned usually to a |
| | (MACE) with | conventional | | | | <u>Failure</u> | colostomy |
| | the | treatment | | Enemas started | | -MACE (n=37): 6 | |
| | caecostomy | | | on 4 th | | (16.2%) | Source of funding: |
| | button (CB) in | Exclusion | | postoperative day | | | not stated |
| | children with | criteria: not | | and MIC-KEY | | 4 patients' colonic | |
| | intractable | clearly stated, | | gastrostomy tube | | washouts ineffective. | |
| | constipation | but all rectal | | changed to | | 1 patient: colonic | |
| | | biopsies were | | standard | | washout associated | |
| | | aganglionic. | | gastrostomy | | with abdominal pain | |
| | | | | button after 6 | | during enema. 1 | |
| | | | | weeks | | patient required | |
| | | | | | | revision for | |
| | | | | _ | | perforation of | |
| | | | | Enemas | | appendicostomy and | |
| | | | | performed by | | the fibrotic-ischaemic | |
| | | | | administering | | appendix was | |
| | | | | saline (20ml/kg) to | | replaced with a CB | |
| | | | | empty the entire | | CD (42), 4 (0.20/) | |
| | | | | colon at a convenient time | | -CB (12): 1 (8.3%) Reason for failure | |
| | | | | | | | |
| | | | | for patient. | | was leaking faecal | |
| | | | | Children not | | content around the | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|--|------------------------------------|---|-------------------|
| | Level | | 5 | responding to saline wash-out | Wiedsures | button, converted to MACE after 20 | |
| | | | | used Klean-Prep. Frequency and | | months P >0.05 | |
| | | | | volume of enemas individualised to each patient to | | Surgical complications %): | |
| | | | | achieve cleanliness and | | a. requiring operative intervention | |
| | | | | stop soiling | | MACE (n=37) -total: 9 (24%) | |
| | | | | | | -stoma stenosis: 11% -iatrogenic perforation | |
| | | | | | | appendicostomy: 5% -difficult catheterization: 5% | |
| | | | | | | -adhesive obstruction: 3% | |
| | | | | | | CB (n=12) -total: 0 | |
| | | | | | | -adhesive obstruction: | |
| | | | | | | Others N.A | |
| | | | | | | P=0.009 for total | |
| | | | | | | b. not requiring operative intervention | |
| | | | | | | MACE (n=37) -total: 7 (19%) | |
| | | | | | | -pain/difficult catheterisation: 11% | |
| | | | | | | -stoma granulosa: 5% -stoma stenosis: 3% | |
| | | | | | | -faecal leakage: 0 -pain around button: | |

| Bibliographic Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|--------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | N.A CB (n=12) -total: 11 (92%) -pain/difficult catheterisation: N.A -stoma granulosa: (33%) -stoma stenosis: N.A -faecal leakage: 42% -pain around button: 92% p<0.001 for total | |

| Exclusion criteria: not clearly stated -failure: regular soiling or constipation persisted , no perceived improvements, procedure abandoned usually to a colostomy Reviewer comments: Retrospective study Low response rate to the proforma Results for patients with diagnoses other than constipation not reported here because they are outside the remit of this review. Main complications not related in paper | Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---|---|---|--|--------------------------------------|---|---|--|---|
| not reported here Source of funding: | MACE procedure: experience in the United Kingdom. 1999. Journal of Pediatric Surgery 34[2], | Retrospective survey Evidence level: 3 Study aim: to find out the current status of the Malone Antegrade Continence Enema | Inclusion criteria: MACE procedures performed by UK members of the British Association of Paediatric Surgeons (or their units) up to the end of 1996 Exclusion criteria: not clearly | Mean age: 12.3 years <u>Country:</u> | Malone Antegrade Continence Enema (MACE) Comparison: | period: Mean 2.4 years (range 0.3 to 6) Outcome Measures: -children diagnoses -success rate -complications | Including both full and partial): 79% Success rate based on diagnosis (%): Constipation (n=23) Full: 52 Partial: 10 Failure: 38 | Results included figures from authors' previous study, reported figures from one other UK centre and replies to proformas sent by authors to BAPS members 102 proformas sent, 58 returned Success criteria: -full: totally clean or minor or minor rectal leakage on the night of the washout; -partial: clean, but significant stomal or rectal leakage, occasional major leak, still wearing protection but perceived by the child or parent to be an improvement failure: regular soiling or constipation persisted, no perceived improvements, procedure abandoned usually to a colostomy Reviewer comments: Retrospective study Low response rate to the proforma Results for patients with diagnoses other than constipation not reported here because they are outside the remit of this review. Main complications not related in paper to the clinical diagnosis and therefore not reported here |

| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---------------------------|-----------------------|--------------------|---------------------------|---------------------------|------------------------|-----------------------|--|
| mormation | Level | rationts | S | Companison | Measures | | |
| Mousa et al. | Study Type: | 31 children | -total population | Intervention: | Duration of | (all values are | Additional information from study: |
| Cecostomy in | Retrospective | | 31 children | Caecostomy | study period: | median) | Standardised questionnaire used to |
| children with | cohort | <u>Inclusion</u> | 58% boys | performed | 4 years | | obtain data on outcomes measured |
| defecation | | criteria: | | percutaneously by | | Type of antegrade | |
| disorders. 2006. | <u>Evidence</u> | Children who | | interventional | Follow-up | enemas used | Frequency of bowel movements scored |
| Digestive | level: | received a | -9 children with | radiologist | period: | | as: 1, <5 bowel movements/week; 2, |
| Diseases and | 2+ | caecostomy | functional | | Median 11 | performed | 5/week to 3/day; 3, 3/day |
| Sciences 51[1], | | fro | constipation | Comparison: | months (range | | |
| 154-160 | | constipation, | | Caecostomy | 1 to 45) after | Bowel movement | Soling frequency scoring: 1, none; 2, |
| | | faecal soiling | median age at | performed by | caecostomy | frequency (n=9) | occasional, 3, few episodes/week; 4. |
| | authors' 4- | or a | time of | open surgical | | Pre: <5/week | few episodes/week to daily; 5, |
| | year | combination | caecostomy: 12 | approach | <u>Outcome</u> | Post: 5/week to 3/day | constantly |
| | | of both. | years old | | Measures: | P<0.01 | |
| | with 2 | Underlying | (range 3 to 16) | | | | Quality of life assessed by scoring |
| | different | conditions | | | -type of | Soiling frequency | limitations of activity (none, mild, |
| | techniques of | included | Country: | | antegrade | <u>(n=9)</u> | moderate and severe), global health |
| | the | functional | USA | | enemas used | Pre: constant | score, and global emotional score (poor, |
| | | constipation, | | | | Post: none | fair, good, very good and excellent) |
| | procedure | Hirschsprung' | | | -bowel | P=0.01 | |
| | and to | s disease, | | | movement | | Reviewer comments: |
| | compare the | imperforate | | | frequency | Number of | Not clear who interviewed the parents |
| | | anus, | | | | medications (n=9) | |
| | | imperforated | | | -soiling | Pre: 4 | Source of funding: |
| | caecostomy in | | | | frequency | Post: 1 | study supported in part by the Ter |
| | | combined | | | | P=0.01 | Meulen Fund, Royal Netherlands |
| | defection | with tethered | | | -number of | | Academy of Arts and Sciences |
| | disorders | spinal cord | | | medications | Number of physician | |
| | secondary to | syndrome and | | | | visits related to | |
| | functional | spinal | | | -number of | defecation problems | |
| | | abnormalities | | | physician visits | <u>(n=9)</u> | |
| | imperforate | | | | related to | Pre: 6 | |
| | | Exclusion | | | defecation | Post: 2 | |
| | spinal | criteria: | | | problems | P<0.01 | |
| | abnormalities | Not stated | | | 1 | | |
| | | | | | -number of | Number of hospital | |
| | | | | | hospital | admissions for | |
| | | | | | admissions for | disimpaction (n=9) | |
| | | | | | disimpaction | Pre: 4 | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|---|-------------------|
| | Evidence | | | Comparison | Outcome | Post: 0 P<0.01 Number of missed school days per month (n=9) NS Global health score (n=9) Pre: poor Post: good P=0.01 Global emotional score (n=9) Pre: poor Post: good P=0.01 Limitations of activity (n=9) Pre: moderate Post: mild P<0.01 Complications No major complications like perforation, stoma stenosis, or stoma prolapse. No | |
| | | | | | | difference found in occurrence of number of complications between different procedures/technique s | |

| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---------------------------|------------------------------------|--|-------------------|
| | | | | | | Other outcomes not reported here as no subgroup analysis performed | |
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| Bibliographic Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|---|--|---|--|--|---|--|--|
| Information Jaffray. What happens to children with idiopathic constipation who receive an antegrade continent enema?. An actuarial analysis of 80 consecutive cases. 2009. Journal of Pediatric Surgery 44[2], 404-407United | Evidence Level Study Type: Prospective case series Evidence level: 3 Study aim: to perform an actuarial analysis of the outcomes of antegrade continent enema (ACE) procedure in | 80 children Inclusion criteria: All children with idiopathic constipation undergoing ACE surgery by 1 surgeon. In all children symptoms had persisted despite medical management supervised by | Characteristic s 80 children 44 boys median age at | Intervention: Antegrade continent enema (ACE) procedure Children followed up in a nurse-led continence clinic Lavage regime was supervised by specialist nurses and used a solution of saline prepared by parents at a volume of | | 53 children: conventional ACE 27 children: laparoscopic ACE - ACE lavage failed in 12 children: 4 children were identified where the appendicostomy was not being used. Although these children could be lavaged, parent's had | Additional information from study: In the first 32 cases the diagnosis was confirmed by the use of marker studies using an established protocol. However because the marker studies did not alter treatment decisions and to avoid unnecessary radiation exposure, this practice was stopped Previous treatment was heterogeneous and had always included prolonged treatment with laxatives, usually with periods of in-patient administration of surgical bowel cleansing solutions, frequent manual disimpaction and often involvement of a clinical psychology |
| States. | have idiopathic constipation | paediatrician for at least 3 years Exclusion criteria: Hirschsprung's disease (excluded by rectal biopsy in all cases) | | 20mL/kg body weight Comparison: N.A | because colonic lavage has not been found to improve the child's bowel habit or the child's colon had not proved to be lavageable and symptoms had deteriorated -Cure: the appendicostom y was closed/reversed because the child achieved normal bowel | not found it to be of help in the child's bowel management and had ceased use In 8 children, deterioration of symptoms occurred despite ACE lavage and required alternative treatment of symptoms. These children could not be lavaged Kaplan Meier probability of an ACE failing: 0.3 at 8.5 | In calculating the Kaplan Meier probability of an ACE being reversed or failing, the following times were calculated: -ongoing lavage: length of follow up calculated as time from the date of formation of ACE to current date -time to failure calculated as the time from creation of the ACE to the clinic letter stating that the parents had ceased using the ACE, or the date of commencement of alternative treatment -cure: the date of the operation to |

| Bibliographic S Information | Study Type & Evidence | Number of Patients | Patient Characteristic | Intervention & Comparison | Follow-up & Outcome | Effect Size | Reviewer Comments |
|-----------------------------|-----------------------|-----------------------|---------------------------|---------------------------|---------------------|---|--|
| | | Patients | Characteristics | Comparison | | years; estimated mean failure time: 8.6 years (95% CI 7.9 to 9.2) -12 children had normal bowel habit, no longer performed colonic lavage and underwent closure of appendicostomy. The Kaplan Meier probability of an ACE being reversed was 0.2 at 6.2 years, estimated mean time to reversal (9.1 years (95% CI: 8.4 to 9.7) -56 children currently performing colonic lavage Colonic transit time (CTT), age at surgery and duration of follow-up were not significantly associated with ACE failure, but sex was (p=0.04) the higher failure rate amongst girls was significant (p=0.02) | reverse the ACE was used as the censoring time A minimum of 6 months follow-up judged to be appropriate because a decision regarding "cure" would take no less than 6 months to determine Children who could not be lavaged defined as those having failed too have a bowel evacuation despite an appropriate volume of lavage fluid. These children were assessed by performing continuous lavage though the appendicostomy over several days while in hospital. Typically such children accommodate very large volumes of fluid in their colon, often in excess of 10 L without bowel evacuation Criteria for ACE reversal: for at least the previous 6 months, child had stopped using their ACE, was stooling spontaneously at least every other day, was not requiring laxative therapy and was not soiling. ACE reversed by dissecting the appendix to the caecal wall and ligating and removing it No patient was discharged, and none was lost to follow up Source of funding: Not stated |
| | | | | | | CTT significant factor in predicting failure in | |

Information and Support for Children with Chronic Idiopathic Constipation and their families

| | | | | Clinic-based In | nterventions | | |
|-------------------|-----------------|------------------|------------------|-------------------------------|---------------------|--------------------------|--|
| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
| Information | Evidence | Patients | Characteristic | Comparison | Outcome | | |
| | Level | | S | | Measures | | |
| Burnett et al. | Study Type: | 102 children | 102 children | Intervention: | <u>Intervention</u> | Primary outcomes | Additional information from study: |
| Nurse | RCT | | 55 males | Nurse led clinic | period: | | Constipation defined as (1) decreased |
| management of | | <u>Inclusion</u> | | (NLC) | 30 months | Time to cure at last | frequency of bowel movements |
| intractable | <u>Evidence</u> | <u>criteria:</u> | median age at | | | | (that is, decreased from the individual's |
| functional | level: | All children | study entry: 4.6 | Comparison: | <u>Assessment</u> | by telephone | previous pattern); and/or (2) harder stool |
| constipation: a | 1+ | aged 1 | (NLC) and 4.8 | Consultant led | point (s): | | consistency; and/or (3) subjective |
| randomised | | to 15 years | years (PGC) | paediatric | Unclear | -Number cured, % | difficulty, including pain and distress |
| controlled trial. | Study aim: | presenting to | | gastroenterology | | NLC (n = 52): | associated |
| 2004. Archives | | | age range: 13 | clinic (PGC) | | 34 (65.4%) | with defecation |
| of Disease in | the | 3 | months to 14.7 | | | PGC (n = 50): | |
| Childhood | effectiveness | ogy service at | years | -Assessment: | | 25 (50.0%) | Interpretation of abdominal radiograph |
| 89[8], 717-722 | | the John | _ | Nurse led clinic | months for both | | obtained at the time of initial |
| | ` ' | Radcliffe | Country: | designed to be a | groups | -Time to event | assessment made though a validated |
| | compared | Hospital, | UK | follow up clinic for | | (median (95% CI, | scoring system (Leech) using scores |
| | with a | Oxford, UK | | children who had | <u>Outcome</u> | months) | ranging from 0 (no stool) to 5 (gross |
| | consultant led | with | | undergone a full | Measures: | | faecal loading with bowel dilatation) in |
| | paediatric | constipation | | and detailed | | NLC (n = 52): | three areas of the colon, giving a total |
| | gastroenterolo | | | medical | | 18.0 (8.5 to 27.5) | severity score ranging from 0 to 15. |
| | gy clinic | Exclusion | | assessment in the | | PGC (n = 50): | Using this system a radiographic score |
| | (PGC) in the | criteria: | | paediatric | | 23.2 (17.3 to 29.2) | of >9 has been shown to have a high |
| | management | Organic or | | gastroenterology | last visit or later | | specificity and sensitivity in the |
| | of chronic | neurological | | clinic leading to a | confirmed by | Hazard ratio(one | diagnosis of childhood constipation |
| | constipation | disease | | diagnosis of | telephone | sided 95% CI): | _ , , , , , , , , , , , , , , , , , , , |
| | | | | idiopathic | | 1.332 (0.860 to ∞) | The primary outcome of cure at last visit |
| | | | | functional | -Time to cure at | Times metic (one sided | or later confirmed by telephone used to |
| | | | | constipation | last visit. | ` | assess sample size. For non-inferiority |
| | | | | luva etimetia nev | | 95% CI): | to be concluded between NLC and |
| | | | | -Investigations: | | 0.816 (0 to 1.032) | PGC, 200 patients (100 per arm) would |
| | | | | Where it was | study | Time a to avera at least | be required for a power of 80% and a |
| | | | | clinically | termination. | Time to cure at last | one-sided significance level of 0.05, |
| | | | | appropriate, an | 2 Cocondor: | <u>visit</u> | assuming the success rate of the PGC |
| | | | | abdominal | 2. Secondary | Number oured 9/ | to be 50%. The range of clinical |
| | | | | radiograph obtained at the | outcomes: | -Number cured, % | equivalence was defined to be within |
| | | | | | | NLC (n = 52): | 15%, therefore non-inferiority was |
| | | | | time of initial | -number of | 27 (51.9%) | defined as the ruling out of a hazard |

| | | | Clinic-based Ir | nterventions | | |
|---|-----------------------|--------------------------------|---|--|--|---|
| Bibliographic Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| | | | assessment both as a diagnostic tool and as a semi-quantitative marker of the severity of constipation -Treatment: a standardised treatment algorithm (constructed for the study, similar to a number of published guidelines) provided the basis for management decisions in all consultations in both clinics -initial phases: involved child and parent education about diet (fibre and fluid), exercise, toilet training, and the actions of the laxatives prescribed. Laxative therapy comprised a combination of stool softeners | clinic visits -number requiring additional medication/in- patient procedures during the scheduled treatment period | PGC (n = 50): 22 (44.0%) -Time to event (median (95% CI, months) NLC (n = 52): 22.1 (15.1 to 29.2) PGC (n = 50): 25.1 (17.0 to 33.2) Hazard ratio(one sided 95% CI): 1.207 (0.749 to ∞) Time ratio (one sided 95% CI): 0.855 (0 to 1.112) Premature study termination -Number, % NLC (n = 52): 5 (9.6) (2 lost to follow-up, 3 withdrew) PGC (n = 50): 14 (28) (10 lost to follow-up, 4 withdrew) -Time to event (median (95% CI, months) NLC (n = 52): NA | ratio less than 0.85 on the basis of the lower limit of the one sided 95% confidence interval. Conversely, for an outcome where a reduction of events is preferable, non-inferiority is defined as the ruling out of a hazard ratio greater than 1.176 on the basis Allocation concealment facilitated by using sequentially numbered sealed envelopes produced by an external source for consecutive and eligible study patients. Randomisation performed using block randomisation with fixed blocks of size four Time to cure at last visit or later confirmed by telephone relates to all those children confirmed cured either at their last visit, or subsequently, confirmed over the telephone. Children who were close to achieving the definition of "cured" at their last visit but who were still being weaned off medication, were not required to attend for a further follow up appointment but received their follow up via the telephone. Time to cure at last visit relates to only those children confirmed cured at their last visit (a subset of the previous outcome). Premature study termination comprises those patients who were either lost to follow up or withdrawn for whatever reason Baseline demographic and clinical presentation characteristics as well as |

| | | | | Clinic-based In | iterventions | | |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|--|------------------------------------|--|---|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| | Level | | S | (for example, lactulose, docusate sodium) and stimulants. Stimulants of different potencies (senna, bisacodyl, sodium picosulphate) were prescribed according to the clinical response as indicated by the bowel diaries. If there was an inadequate clinical response to this initial phase, the patient moved on to an advanced treatment regime which might include, enemas, intestinal lavage, manual removal of faeces under general anaesthesia, or psychological referral as was appropriate in each case | Measures | PGC (n = 50): NA Hazard ratio(one sided 95% CI): 0.334 (0 to 0.788) Time ratio (one sided 95% CI): NA Secondary outcomes Number of clinic visits -Median number of visits in each clinic: 6.0 -Median number of inter-visit contacts: NLC: 6.0 (range 2 to 16) PGC: 0.0 (range 0.0 to 29) Number requiring additional medication/in-patient procedures during the scheduled treatment period No significant differences between both groups | previous laxative usage well balanced across clinics ITT analysis conducted for all outcomes. Survival analysis conducted for the primary time-to-event outcomes Reviewer comments: Unclear who measured outcomes Results not controlled for potential confounders Source of funding: Research grants from Norgine Ltd and from WellChild |
| | | | | -Monitoring /follow-up: Bowel | | both groups | |

| | | | | Clinic-based In | terventions | | |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---|------------------------------------|---|-------------------|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| | | | | diaries, which report the frequency, size, and consistency of stools, presence or absence of soiling, and a record of daily laxative medication, were used in both clinics to monitor progress and response to treatment. Dedicated case report forms were used for each study participant and, together with detailed clinical history (including a detailed dietetic history) and clinical findings on initial assessment, documented details of bowel habit and drug therapy at all subsequent outpatient visits. Any other contact with the families, e.g. on the | | 10 children (5 NLC, 5 PGC) completed study as per the protocol but were not cured (treatment failures): -8/10: formally referred for psychological / psychiatric management -9/10: had documented serious behavioural problems -3/10: also referred for surgical assessment and management A total of 15/102 children still undergoing follow up, as they are not cured. In this group, 7/15 children are followed up in the PGC and 8/15 in the NLC. 7/15 children had documented psychosocial problems associated with poor compliance in attending clinic appointments | |
| | | | | telephone or a | | | |

| | | | | Clinic-based In | terventions | | |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|--|------------------------------------|-------------|-------------------|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| | | | | home visit, was documented using inter-visit contact forms -Discharge: Child defined as having been "cured" of their constipation when, for a period of at least 1 month, they had been opening their bowels, producing a normal formed stool without difficulty at least 3 times per week and without any laxative therapy | | | |

| | | | | Clinic-based I | nterventions | | |
|---|-----------------------------------|---|---|--|---|---|--|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| Sullivan et al. Parent satisfaction in a nurse led clinic compared with a paediatric gastroenterolog y clinic for the management of intractable, functional constipation. 2006. Archives of Disease in Childhood 91[6], 499-501 | parent's satisfaction | gastroenterol ogy service at the John Radcliffe Hospital, Oxford, UK with constipation Exclusion criteria: Organic or neurological | 102 children 55 males median age at study entry: 4.6 (NLC) and 4.8 years (PGC) age range: 13 months to 14.7 years Country: UK | Intervention: Nurse led clinic (NLC) Comparison: Consultant led paediatric gastroenterology clinic (PGC) Intervention as described in previous study | Duration of treatment As previous RCT Assessment point (s): After 12 months' follow-up or before this if the child has been "cured" Outcome Measures: 1. Parent satisfaction, 6 domains: -provision of information -empathy with patient -technical quality and competence -attitude towards the patient | Provision of information scores (median) NLC: 8.7 PGC: 7.5 P<0.001 Empathy with patient scores (median) NLC: 9.0 PGC: 7.3 P<0.001 Technical quality and competence scores (median) NLC: 9.1 PGC: 8.0 P<0.001 Attitude towards the patient scores (median) NLC: 8.7 PGC: 7.3 P<0.001 Access to and continuity with the caregiver scores (median) NLC: 8.2 PGC: 6.7 P<0.001 | Additional information from study: Satisfaction with care defined as "the degree to which parents perceive the needs of their children are met" Parent satisfaction measured using a validated instrument based on the Leeds Satisfaction Questionnaire (LDQ), which has been shown to be easy and quick to complete sensitive to change, reliable and reproducible. Questions in the LDQ were pertinent to a rheumatology clinic and thus adapted for the purposes of this constipation clinic. Questionnaire covered 6 separate domains in 48 statements: provision of information, empathy with the patient, access to and continuity with the caregiver and overall satisfaction. The "overall satisfaction" component was added for the purposes of validation. 5 point Likert scales used fro responses ranging from "strongly agree" to "strongly disagree", stability of the instrument tested using the test-retest method An attempt was made to record all "inter-visit" contacts (by telephone or day ward attendances) made by parents outside their schedules outpatients appointment A total of 90 questionnaires returned from 107 families canvassed (84%); 40/51 (78%) from the PGC and 50/56 |
| | | | | | -access to and continuity with | Overall satisfaction scores (median) | (89%) from the NLC. Robustness and high reliability of the questionnaire |

| | | | | Clinic-based I | nterventions | | |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|--|--|--|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| | | | | | the caregiver -overall satisfaction 2. Number of inter-visit contacts | NLC: 8.7 PGC: 7.3 P<0.001 Number of inter-visit contacts (mean (SD)) NLC: 2.37 ± 4.17 PGC: 1.70 ± 4.79 NS | demonstrated by calculating the internal consistency for each domain; lowest Cronbach's alpha: 0.81 Reviewer comments: This study is an evaluation of the previous RCT ITT analysis performed for all outcomes Source of funding: Research grant form WellChild |

| | | | | Clinic-based In | nterventions | | |
|--|---|---|--|---|---|---|---|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| Information Poenaru et al. The Pediatric Bowel Management Clinic: initial results of a multidisciplinary approach to functional constipation in children. 1997. Journal of Pediatric Surgery 32[6], 843-848 | Evidence Level Study Type: Prospective case series Evidence level: | 114 patients Inclusion criteria: Children up to 19 years old referred to the clinic with constipation after a 3- month | s 114 patients Mean age: 5.4 ± 3.8 years (range 4 months to 19 | Intervention: Bowel Management Clinic -Clinic staff: a physician (rotating between 2 paediatricians, 1 paediatric gastroenterologist and 1 paediatric | Measures Duration of treatment Mean time span between first and last visit to clinic: 4.5 months Assessment point (s): 2 and 4 months after initial clinic visit Outcome Measures: -stool frequency per month -stool consistency -occurrence | Stool frequency per month, mean (n=26) 1rst visit: 11.73 last visit:: 29.77 p=0.00026 Stool consistency (n=55) (Unclear whether the following are number of children or %) -liquid 1rst visit: 0 last visit: 1 -soft 1rst visit: 4 last visit: 13 -formed 1rst visit: 16 last visit: 13 -hard 1rst visit: 10 last visit: 3 p=0.00004 Occurrence of | Additional information from study: Children considered constipated when they had persistent symptoms (soling, pain, bleeding, etc) related to bowel movements which tend to be infrequent Total number of visits was 257 with average of 6 patients per clinic. 62 patients seen more than once with a mean of 3.1 visits per patient and a mean time span between the first and the last visit to clinic of 4.5 months Sample size varies in each category of symptoms because of incomplete observations and stool frequencies were only included for non-soiling patients 13 children appeared to be lost to follow-up (no return to clinic in over 6 months) and 11 were discharged Among the discharges the mean number of clinics visits was 3.5 Patient data collected prospectively from the families and the clinic staff. Before initial clinic visit families filled out several mailed questionnaires covering medical, psychological and social issues surrounding the child's problem. These included a medical information |
| | | | | to establish components of individualised management. | with care, 5 scales: respectful and supportive care, | symptoms (%) -Soiling (n=42) 1rst visit: 57 last visit: 43 | questionnaire, a family information questionnaire, the Family Assessment Device (FAD), the Chronic Illness psychosocial Inventory (CI-PSI) and a |
| | | | | Further referral to other BMC staff | enabling and partnership, | NS | knowledge quiz. Parents also required to complete a "constipation/soiling diary" |

| | | | Clinic-based li | nterventions | | |
|--|--------------------|--------------------------------|---------------------------|--|---|--|
| Bibliographic Study Type of Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| | | | of organic cause | providing specific information, coordinated and comprehensive care | -Rectal pain (n=51) 1rst visit: 53 last visit: 22 p=0.0003 -Rectal bleeding (n=54) 1rst visit: 26 last visit: 4 p=0.00035 Frequency of symptoms per month Soiling (n=26) 1rst visit: 30.7 last visit: 12.8 p=0.015 Rectal pain (n=23) 1rst visit: 9.5 last visit: 2.0 N.S Rectal bleeding (n=11) 1rst visit: 0.6 last visit: 0.2 N.S Satisfaction with care Results only reported in a graph from which it is difficult to extract estimates Scores were normal | fro one week, detailing the child's stools and symptoms. At the first clinic visit a structured history/physical examination completed by physician. At each follow up families completed a short progress questionnaire and asked to continue diaries throughout. The FAD, CI-PSI questionnaires and knowledge quiz were repeated at 2 and 4 months after initial clinic visit. A Measure of Processes of Care (MOPC) questionnaire was also administered at the 4-month point. MPOC is a self report measure of the parents' perceptions of the extent to which 5 behaviours of health care professionals occur (respectful and supportive care, enabling and partnership, providing general information, providing specific information, coordinated and comprehensive care). The scores from the study group were compared with those from a normative group of 653 patients Source of funding: Educational grant from Janssen Pharmaceutica through Queen's GI Motility Education Centre |

| | Clinic-based Interventions | | | | | | | | |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|---|------------------------------------|--|-------------------|--|--|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments | | |
| | | | | taking effect. Choice of enemas are phosphate and tap water or saline. High colonic saline irrigations used in severe cases, suppositories not routinely employed. Choice of laxative based on compliance and nature of symptoms. Most patients treated with senna, Docusate sodium and mineral oil. Multiple laxatives avoided. Patient started on recommended dosages, then increased by 50% every 4 to 5 days until symptomatic improvement noted. Individualised dosage then maintained minimum 3 to 6 months, during which dietary and psychosocial issues are dealt | | or higher that the norm for: respectful and supportive care, enabling and partnership and coordinated and comprehensive care Scores were lower than the norm for providing general information and providing specific information | | | |

| | | | | Clinic-based In | terventions | | |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|--|------------------------------------|-------------|-------------------|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| | | | | with. Patient is then slowly weaned off medications -Follow-up: arranged by each health care professional as needed. Visits used to monitor progress and continue education process. Patients who show no progress are reassessed by physician and may become candidates for diagnostic testing -Discharge: when patient is asymptomatic and off medications. Patient referred back to the referring physician, with information for maintaining healthy bowel routine | | | |
| | | | | Comparison: N.A | | | |

Information and Support for Children with Chronic Idiopathic Constipation and their families

| | | | | Web-based In | terventions | | |
|-------------------|---------------------|-----------------------|----------------------|-----------------------------|-----------------------------|---|--|
| Bibliographic | Study Type & | Number of | Patient | Intervention & | Follow-up & | Effect Size | Reviewer Comments |
| Information | Evidence | Patients | Characteristic | Comparison | Outcome | | |
| | Level | | S | | Measures | | |
| Borowitz et al. | Study Type: | 1142 | 1142 | Intervention: | <u>Outcome</u> | The tutorial received | Additional information from study: |
| Using the | Online survey | participants | participants | Multimedia tutorial | | 157 326 successful | The tutorial also includes a one-page |
| Internet to teach | | | | | -clarity and | page requests from | feedback form comprised of 6 multiple- |
| parents and | <u>Evidence</u> | <u>Inclusion</u> | only 887 (78%) | Directed primarily | easiness of | 38 012 distinct hosts | choice questions and one open-ended |
| children about | <u>level:</u> | criteria: | answered the | at parents and | information | | comment field. Questions were |
| constipation | 4 | Children and | questions | older children. | presented in | Was the information | developed in consultation with the |
| and encopresis. | | parents who | categorising the | Includes | tutorial | presented in the | university division of survey research. All |
| 2001. Medical | Study aim: | accessed a | reader: | information about | | tutorial clear and easy | completed form were sent via email |
| Informatics and | | tutorial about | | differential | -usefulness of | to understand? | directly to the main author |
| the Internet in | the feedback | childhood | -789 (89%): | diagnosis, | tutorial: helping | (N=883) | |
| Medicine 26[4], | received | constipation | parents and | aetiology, | parents to | | Responses to multiple-choice questions |
| 283-295 | regarding a | and . | guardians of a | treatment and | understand why | -Very clear: 812 | were tabulated. One author reviewed all |
| | | encopresis, | child with | potential side | children | (92%) | free text comments and identified the |
| | tutorial about | developed | constipation or | , | develop | -Pretty clear: 71 (8% | central them of each comment. |
| | chronic | and installed | encopresis | follow-up | constipation | -Nobody chose "not | Comment were categorised as: |
| | childhood | on the web | 44 (50() | including regular | and/or . | very clear" or "not | -appreciation for making the information |
| | constipation | pages of the | -44 (5%): | monitoring, | encopresis, | clear at all" | available |
| | and | Children's | grandparent or | natural history | making parents | Did the tutorial hale | -question (s) about a particular child's |
| | | Medical centre at the | other family members | and prognosis and a list of | better able to take care of | Did the tutorial help | symptoms or treatment |
| | during 28 months | University of | members | references | their child | you to understand why children develop | -a general question not specific to any particular child |
| | between | Virginia, and | -30 (3%): | references | lineii ciilia | constipation and/or | -a referral request |
| | January 1998 | also | teachers | Comparison: | -usefulness of | encopresis? (N=696) | -a request for dietary recommendations |
| | and April | completed an | leadileis | N.A | tutorial as a | encopiesis: (IV=030) | -a request for additional online |
| | 2000 | online | -9 (1%): | IV.A | good way to | -Completely: 174 | information, such as online forum or a |
| | 2000 | feedback | physicians | | teach people | (25%) | frequently asked questions (FAQ) site |
| | | form. No | priyololario | | about health | -Somewhat: 174 | -specific recommendations as to how to |
| | | internal or | -35 (4%): other | | problems | (25%) | improve the tutorial |
| | | external | healthcare | | | -A little: 13 (2%) | in prove the tatemen |
| | | announceme | providers | | -questions or | -Not al all: 0 | Definition of constipation in the tutorial: a |
| | | nt made to | i. | | comments or | | child is constipated when he or she |
| | | communicate | Country: | | suggestions as | After completing the | passes bowel movements less than |
| | | the availability | | | to how to | tutorial, do you think | every other or every third day and when |
| | | of the tutorial, | | | improve the | you are better able to | he or she passes a bowel movement, it |
| | | but access to | | | tutorial | take care of a child | often is large and hard and perhaps |

| Bibliographic Information Study Type & Evidence Level The website was not limited in any way. These Study Type & Evidence Level Study Type & Evidence Comparison Study Type & Evidence Characteristic Scomparison Study Type & Evidence Comparison Study Type & Evidence Comparison Study Type & Evidence Comparison Suffering from Constipation and/or encopresis? (N=696) Reviewer Comments Suffering from Constipation and/or encopresis? (N=696) Not all participants answered as | |
|---|--|
| was not constipation and/or limited in any encopresis? (N=696) Reviewer comments: | |
| pages can be found by a link in the university homepage called "tutorials for families" about health problems? (N-691) Exclusion criteria: Not stated Source of funding: Not stated Source of unding: Not stated | |

| | | | | Web-based In | terventions | | |
|---------------------------|-----------------------------------|-----------------------|--------------------------------|------------------------------|------------------------------------|---|-------------------|
| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| | | | | | | symptoms or treatment: 167 (20%) -a general question not specific to any particular child: 96 (11%) -a referral request: 46 (5%) -a request for dietary recommendations: 34 (4%) -a request for additional online information, such as online forum or a frequently asked questions (FAQ) site: 21 (2%) -specific recommendations as to how to improve the tutorial: 38 (4%) | |

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| Bibliographic S Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| An Internet intervention as adjunctive therapy for pediatric encopresis. 2003. Journal of Consulting and Clinical Psychology 71[5], 910-917 | Evidence evel: 1+ Study aim: To examine the utility and effectiveness of an Internet- cased version of enhanced coilet training | between 6 and 12 years, soling at least once a week and have no medical diagnosis | 24 children 19 boys mean age: 8.46 years (SD1.81) -Web group: 12 children (10 boys) -No-Web group: 12 children (9 boys) Country: USA | Intervention: Web intervention Comparison: No-Web intervention -The Web site: Web-based program for the treatment of paediatric encopresis (U-CAN-POOP-TOO) (please refer to Ritterband, 2008 for a description of the program) | Duration of intervention: 3 weeks Assessment point (s): 3 weeks after initial home visit Follow-up period: None Outcome Measures: -number of faecal accidents per week -number of bowel movements (BM) passed in the toilet per week - bathroom use without prompts -bathroom use with prompts -internet use (most/least useful aspect of the programme; | p=0.001 | Additional information from study: Computer and internet access provided to all families who contacted the research centre and met the inclusion criteria Participants received a \$25 gift certificate to a local toy sore for completing the pre-treatment assessment and another \$25 gift certificate for completing the post-treatment assessment assessment Information regarding BM assessed by parent report on the Child Information Form. Question regarding child's bowel habits included such as number of BMs in toilet and use of toilet with / without parental prompts. Questions regarding use of internet programme also included in post-treatment form for the intervention group. The Virginia Encopresis/Constipation Apperception Test (VECAT) also administered. It assesses bowel specific problems related to the process of encopresis, such as avoidance of the toilet, non responsiveness to rectal distension cues and fear of defecation pain. A generic subscale included as a comparison measure, addresses problem behaviours not related to bowel issues. The VECAT consists of 18 pairs of drawings (9 pairs bowel-specific and 9 parallel generic events) and child selects the picture in each pair that best |

| | | Web- | based Interventions | | |
|----------|---|----------------------|--|---|--|
| . | umber of Patient Patients Characteristi | Characteristic Compa | | Effect Size | Reviewer Comments |
| | | | regarding individual cores an modules) | Internet use (Web group only) 1. Most useful aspect of the programme: -the step by step program to get the child regulated -understanding why his body does what it needs to do everyday-and what happens when he doesn't have a BM and health consequencesinfor mation was tremendously useful developing a feeling that he can control his own body realising that he's not the only child with this problemthat was | No significant differences in baseline characteristics between the 2 groups (age, gender, race, stage of bowel movement training, length of current laxative regime or any of the outcomes measured) CM1: anatomy and pathophisiology CM2: medication (enemas/laxatives) CM3: behavioural intervention Reviewer comments: No definition of constipation / soling given Small sample size, no sample size calculation Randomisation and allocation concealment method not described No dropouts/lost to follow up reported Results not controlled for potential confounders Source of funding: National Institutes of Health Grant RO1 HD28160 |

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| | Level | | s | | Measures | did not print out -Miralax should have been included (as a choice of laxative) -nutrition portion was too limited Internet experience: parents' views / satisfaction -found material understandable (mean 5.00, SD 0.00, N = 20) -found it easy to use (mean 4.62, SD 0.74, N = 21) -believed their child liked the program (mean 4.05, SD 1.28, N = 21) - believed their child found it understandable (mean 4.32, SD 0.89, N = 19) - believed their child found it easy to use (mean 4.47, SD 0.77, N = 19) 3. Preference regarding cores modules (CM) (mean, | |
| | | | | | | modules (CM) (mean, SD) (score 0 to 4) | |

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| | | | | | | a. How useful: CM1: 3.84 (0.38) CM2: 3.94 (0.24) CM3: 4.00 (0.00) b. How well did you understand the material CM1: 3.89 (0.32) CM2: 3.89 (0.32) CM3: 3.92 (0.28) c. how well did your child understand the material CM1: 3.53 (0.61) CM2: 3.28 (1.07) CM3: 3.54 (1.13) d. How much did you enjoy using the module CM1: 3.68 (0.48) CM2: 3.67 (0.49) CM3: 3.69 (0.48) e. How much did your child enjoy using the module CM1: 3.63 (0.76) CM2: 3.61 (0.98) CM3: 3.46 (1.13) | |

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| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| Ritterband et al. Using the internet to provide information prescriptions. 2005. Pediatrics | Level Study Type: RCT-Survey Evidence level: 1+ (RCT component) 3 (survey component) Study aim: To determine if families of children suffering from chronic constipation and/or encopresis will visit an educational Web site that is specifically prescribed by their physician and whether | 83 patients and their families Inclusion/exclusion criteria: Families with a child who was being seen for the first time in the paediatric gastroenterol ogy clinic at the University of Virginia with a chief complaint of chronic constipation and/or encopresis. To be eligible, families had to have access to the Internet in their home and have an | | Intervention: E-mail-prompt group (n=43) Comparison: No E-mail-prompt group (n=40) At the conclusion of the patient's clinic visit, 1 of the 2 attending gastroenterologist s provided a form with the Web-site address and a log-in identification number. The handout, signed by the physician, stated: "It is important to learn as much as you | Measures Duration of intervention 1 week Assessment point (s): 1 week Follow-up period: None Outcome Measures: -Number of families who visited the prescribed Web site within 1 week of their clinic visit -Perceived barriers to accessing the Web site | Number of families who visited the prescribed Web site within 1 week of their clinic visit (N=83) 54 (65%) Perceived barriers to accessing the Web site 18 interviewed subjects did not go to the Web site because (n, %): 1. Personal / family / behaviour: -just forgot: 11 (61) -didn't have much time: 11 (61 -lost flyer: 6 (33) -interrupted: 3 (17) -computer in use by another: 2 (11) -did not think it would be useful: 2 (11) -did not want to go: 1 (6) | Additional information from study: On the Web page, users read the following instructions: "We hope you find the information in this website to be helpful. Before you can begin, please enter the ID number you were given in the space below, and then click the button to begin." When the "submit" button was clicked, the 2-digit identification number and the date and time were logged in a database. The 2-digit identification number identified the family as a member of the e-mail-prompt group or no-prompt group. This was the only information captured in the database No significant differences between the 2 groups on type and speed of Internet connection, the number of times they reported checking their e-mail, or frequency of using the Internet There were no significant differences in the ages of the children between the 2 groups Approximately 1 week after the clinic visit, the study coordinator attempted to contact the primary caretaker of each patient by telephone or e-mail to ask about their experience accessing the |
| | the Web site. In addition, barriers to accessing the prescribed | account. | | this Web site and review the relevant material. This should be beneficial to your | | -did not like typing in URLs: 1 (6) -did not know how to type in URLs: 1 (6) -child not | Web site. Families who did not access the Web site were encouraged to identify barriers that they may have experienced in accessing the prescribed Web site. They were presented with a |

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| | Web site were identified | | | child's treatment." Families were assigned randomly into a "prompt" group or "noprompt" group or "noprompt" group. 2 business days after the clinic visit, an e-mail containing the Web-site address and a reminder to visit the Web site was sent to those in the "prompt" group -The Web site: an abbreviated version of a larger Web-based program for the treatment of paediatric encopresis (U-CAN-POOP-TOO) -3 modules: (1) "How to Strain": reviewed proper defecation dynamics, including proper positioning, straining, and | | cooperating: 0 -did not know how to use internet: 0 -family thought it was a bad idea: 0 2. Technical issues/obstacles -computer broken: 4 (22) -internet connection broken: 2 (11) -difficulty logging on: 1 (6) -too long to log on:1 (6) No significant differences in identified obstacles between the families who received the e-mail reminder and those who did not | list of potential barriers and were asked whether the item had been a barrier for them to accessing the Web site. Individuals were able to select multiple barriers, if applicable Of the 83 families, 67 (81%) were contacted by telephone (n= 57) or e-mail (n= 10) No significant differences were found in identified obstacles between the families who received the e-mail reminder and those who did not Reviewer comments: No definition of chronic constipation or encopresis given No sample size calculation performed Randomisation and allocation concealment methods not described Results controlled for potential confounders Source of funding: Partially supported by National Institutes of Health grant RO1 HD28160 |

| | Web-based Interventions | | | | | | | | | | |
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| | | | | muscle control/ strength-building exercises (2) "Giving and Getting Enemas": reviewed techniques for administering enemas (3) "The SuperCleanout game": An arcade-style game for children with a learning message. Parents and children were able to view as much of the site as they wanted and could come back as often as they liked | | | | | | | |

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| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| Ritterband et al. | Study Type: | 49 children | 49 children and | Intervention: | Duration of | Motivation scores | Additional information from study: |
| Examining the | Single sample | and their | their families | Modified modules | <u>intervention</u> | (lower score reflects | Families who agreed to participate |
| added value of | cross-over | families | | including audio, | Each module | more motivation) | received a |
| | RCT | | 32 boys | graphics and | with or without | | \$25 gift certificate forma a local toy store |
| and interactivity | Multicentre | <u>Inclusion</u> | | interactivity | each | -Audio | |
| in an internet | | criteria: | mean age: 7.98 | | component | | Parents asked to complete the |
| intervention for | (and these | Children aged | | Comparison: | presented once | Audio-computer | motivation and readiness to change |
| pediatric | are the results | 5 to 12 years | (SD=1.88) | Modules without | | | items from their child's perspective: |
| encopresis. | of 3 individual | who were | | audio, graphics or | <u>Assessment</u> | a. Child | |
| 2006. Children's | studies for | being see for | Country: | interactivity | point (s): | Pre: 6.00 | -Motivation: a 3-item parallel drawing |
| Health Care | each | encopresis at | USA | | Immediately | Post: 5.13 | selection measure was created in the |
| 35[1], 47- | component) | 2 paediatric | | | after each | P≤0.004 | same manner as the Virginia |
| 59United | | gastroenterol | | 2 modules of the | module was | | Encopresis-Constipation Apperception |
| States. | Evidence | ogy clinics | | original U-CAN- | presented | b. Parent | Test for both the enema and proper |
| | level: | | | POOP-TOO | | Pre: 7.56 | defecation dynamics modules. |
| | 1+ | Exclusion | | intervention were | Follow-up | Post: 6.25 | Respondents select the image in each |
| | | criteria: | | revised: | period: | P=0.06 | pair which they feel is closest to |
| | Study aim: | Not stated | | | None | | represent how they might act given the |
| | To determine | | | -"Giving and | | 2. Audio-person | scenario presented in the picture (e.g. |
| | the | | | Getting Enemas": | Outcome | · | child does not want an enema vs. child |
| | usefulness | | | reviewed | Measures: | a. Child | wants an enema, child feels urge to |
| | and user | | | techniques for | -motivation | Pre: 6.19 | poop but keeps on playing vs. go right |
| | preference for | | | administering | | Post: 5.63 | away to sit on toilet). Respondents are |
| | audio (use of | | | enemas | -readiness to | N.S | then asked whether he or she is "a lot |
| | sound), | | | "How to Strain": | change | | like or "a little like" the image selected. |
| | graphics (use | | | reviewed proper | | b. Parent | Pre-post reliability correlations on the |
| | of images) | | | defecation | | Pre: 8.75 | motivation scale for the enemas and |
| | and | | | dynamics, | | Post: 7.13 | dynamic modules were .66 and.83 |
| | interactivity | | | including proper | | P≤0.02 | respectively |
| | (triggering of | | | positioning, | | | |
| | events by the | | | straining, and | | -Graphics | -Readiness to change: a 1-item scale |
| | user causing | | | muscle control/ | | , | with4 response options was created to |
| | various | | | strength-building | | 1. Graphics + | identify the child's stage of change as |
| | actions, i.e. | | | exercises | | · | defined by Prochaska and DiClemente, |
| | clickable | | | | | a. Child | 1983) with respect to both receiving an |
| | buttons) in a | | | Design was | | Pre: 5.69 | enema and proper defecation dynamics |
| | paediatric | | | significantly | | Post: 5.19 | |

| Bibliographic Study Type & Number of Inferrention & Follow-up & Effect Size Characteristic Study Type & Patients Characteristic Study Type & Patients Characteristic Study Type & Patients Characteristic Study Type & Study Type | Reviewer Comments iewer comments: definition of chronic constipation oppresis given |
|---|--|
| Internet- improved with N.S Revi | definition of chronic constipation of |
| | |
| | opresis given |
| intervention given to graphical, b. Parent enco | |
| specifically animation and Pre: 7.13 | |
| designed for interactive Post: 6.06 No s | sample size calculation |
| patients with elements. For P≤0.03 | • |
| encopresis each of the 3 | eline characteristics not compare |
| studies 2. Graphics - | , , , , , , , , , , , , , , , , , , , |
| | domisation and allocation |
| | cealment methods not described |
| modified to either Pre: 5.75 | sourment methode het decembed |
| | dropouts/lost to follow up reporte |
| constructs of N.S | aropouto/lost to rollow up roporto |
| | ults controlled for potential |
| | ounders |
| interactivity) or Pre: 8.06 | ounders |
| | ree of fundings |
| | rce of funding: |
| | onal Institutes of Health grant RC |
| | 28160 |
| both modules -Interaction | |
| were created with | |
| and without 1. Interaction + | |
| sound. For the | |
| study examining a. Child | |
| graphics both Pre:6.00 | |
| modules were Post: 4.71 | |
| created with P=0.03 | |
| graphics and | |
| completely text b. Parent | |
| based; and for the Pre: 8.35 | |
| study examining Post: 6.88 | |
| interactivity both NS | |
| modules were | |
| created with 2. Interaction - | |
| interaction (use | |
| the mouse to click a. Child | |
| various aspects of Pre: 5.18 | |

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| | | | | the screen and navigation) and as a movie (where no interaction was necessary and the participant could just watch the module play from beginning to end | | Post: 4.41 P=0.02 b. Parent Pre: 7.76 Post: 7.29 NS | |
| | | | | | | Stage of change scores -Audio | |
| | | | | | | 1. Audio-computer a. Child Pre: 2.88 Post: 3.00 N.S | |
| | | | | | | b. Parent Pre: 2.19 Post: 2.69 N.S | |
| | | | | | | 2. Audio-person a. Child Pre: 2.69 Post: 2.63 N.S | |
| | | | | | | b. Parent Pre: 2.25 Post: 2.75 P=0.04 | |

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| | | | | | | -Graphics | |
| | | | | | | 1. Graphics + | |
| | | | | | | a. Child Pre: 3.38 Post: 3.31 NS | |
| | | | | | | b. Parent Pre: 2.44 Post: 2.88 P=0.01 | |
| | | | | | | 2. Graphics - | |
| | | | | | | a. Child Pre: 3.38 Post: 3.25 NS | |
| | | | | | | b. Parent Pre: 2.75 Post: 3.13 NS | |
| | | | | | | -Interaction | |
| | | | | | | 1. Interaction + | |
| | | | | | | a. Child Pre: 2.47 Post: 2.71 NS | |

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| | | | | | | b. Parent Pre: 2.18 Post: 1.94 NS 2. Interaction - a. Child Pre: 2.53 Post: 2.53 NS b. Parent Pre: 1.82 Post: 1.94 NS | |

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| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments |
| Ritterband et al. | Study Type: | 22 children | 22 children | Intervention: | Duration of | Number of faecal | Additional information from study: |
| Real world use | Prospective | | | Internet-based | intervention | accidents over a 2- | Of 46 patients originally provided with |
| of an Internet | case series | <u>Inclusion</u> | 13 males | intervention for | 2 weeks | week period (mean) | the Web-based information prescription |
| intervention for | | criteria: | | childhood | | -initial period: | 10 could not be reached by phone or |
| pediatric | Evidence | Children with | mean age: 8.10 | encopresis: U- | Assessment | 13.86 (SD 10.40, | email for interview, of the remaining 36 3 |
| encopresis. | level: | a | years (SD 2.3 | CAN-POOP-TOO | point (s) and | median 13.00) | did not provide consent, 3 stated that |
| 2008. Journal of | 3 | documented | years) range | | follow-up period | | they never received the initial email with |
| Medical Internet | | diagnosis of | 5.1 years to | Child-focused | | -follow-up period: | their personalised log-in information, 5 |
| Research 10[2], | Study aim: | encopresis as | 12.11 years | programme, | -initial period: 2 | 2.14 (SD 2.21, | never logged on and 3 logged but never |
| e16 | To examine | noted in their | - | targets primarily 5 | weeks before | median 1.00) | viewed any of the intervention material. |
| | the utility and | medical | Country: | to 10 years old | children were | P < .001 | No subsequent data was collected on |
| | impact of an | records and | USA | children but was | enrolled in the | | these patients |
| | Internet | their families, | | designed to be | program | Number of bowel | |
| | intervention | seen at the | | used by child and | | movements (BM) | Number of faecal accidents, number of |
| | for childhood | Paediatric | | parent (s) | -follow-up | passed in the toilet | bowel movements passed in the toilet |
| | encopresis as | Gastroenterol | | together | period: 2 weeks | over a 2-week period | and average amount of perianal pain |
| | part of | ogy Clinic at | | | immediately | (mean, SD) | experienced during defection were |
| | standard | the University | | 3 core modules | before phone | -initial period (n=21, | obtained from children's medical charts |
| | medical care | of Virginia | | take 60 to 90 | interview | missing data) | and though a phone interview with |
| | in a "real | Children's | | minutes to | | 14.62 (10.68) | parents. Interview also included open- |
| | world" setting | Hospital | | | <u>Outcome</u> | | ended questions about what the parents |
| | | between . | | users instructed to | | -follow-up period: | believed were the most helpful and least |
| | | all children | | review them | | 14. 82 (8.65) | helpful components of the programme. 3 |
| | | had been | | during the first | -number of | NS | structured questionnaire mostly |
| | | given access | | | faecal accidents | | developed for this interview were also |
| | | to the | | 1. The body | over a 2-week | Average amount of | completed: U-CAN-POOP-TOO Utility |
| | | paediatric | | (anatomy, | period | perianal pain | Questionnaire administered to all |
| | | encopresis | | physiology and | | experienced during | parents who had used the program |
| | | Internet | | pathophysiology | -number of | defection over a 2- | (extent to which the parent and child |
| | | intervention | | of digestion) | bowel | week period (mean, | found program useful, enjoyable, |
| | | as part of | | 2. How to poop | | <u>SD)</u> | understandable and easy to use); U- |
| | | their | | (behavioural | (BM) passed in | | CAN-POOP-TOO Impact Questionnaire |
| | | treatment | | techniques for | the toilet over a | -initial period: | administered to all parents who had |
| | | | | treatment of | 2-week period | 0.56 (0.78) (n=18, | used the program (parents to rate how |
| | | Exclusion | | encopresis) | | missing data) | much they perceived the programme |
| | | criteria: | | Medication | -average | | helped their child) and Internet |
| | | Not stated | | (clean-out and | amount of | -follow-up period: | Intervention Adherence Measure |

| | Web-based Interventions | | | | | | | | | |
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| Bibliographic Information | Study Type & Evidence Level | Number of Patients | Patient Characteristic s | Intervention & Comparison | Follow-up & Outcome Measures | Effect Size | Reviewer Comments | | | |
| | | | | laxative treatment) New modules assigned each week based on a follow-up assessment the user completes about their child's status. Not all modules necessarily used by all users, only those modules identified as relevant are assigned and reviewed. However all modules can be viewed by all users. Follow-up comprised of 17 to 20 questions, depending on the week. System contains a total o 22 modules, each takes 5 to 10 minutes to review Comparison: N.A | perianal pain experienced during defection over a 2-week period -utility and impact of the programme :parents' views/satisfacti on -adherence | N = 21) -believed their child liked the program (mean 4.05, SD 1.28, N = 21) - believed their child found it understandable (mean 4.32, SD 0.89, N = 19) - believed their child found it easy to use (mean 4.47, SD 0.77, N = 19) -most helpful components of the program: tutorials about anatomy and pathophysiology, | administered to patients who stopped using the programme for some reason other than that their problem was "resolved". Those who responded "not applicable" to items on the U-CAN-POOP-TOO Utility Questionnaire were not included in the analysis for that item (explaining the varying sample sizes) The U-CAN-POOP-TOO Impact Questionnaire was administered to examine how much the parents believed the program affected outcome. Those who responded "not applicable" were not included in the analysis for that item No significant correlations found between computer/Internet usage and the change from initial to follow-up period for accident frequency (<i>r</i> = .09, <i>P</i> < .69, N = 22), BMs passed in the toilet (<i>r</i> = .38, <i>P</i> < .09, N = 21), or amount of pain associated with defecation (<i>r</i> = .08, <i>P</i> < .76, N = 18). Internet comfort and connection speed were also not significantly correlated to changes in any of the bowel-related outcome variables (<i>r</i> values ranged from17 to .27; <i>P</i> values ranged from .25 to .59) Of the 22 patients who used U-CAN-POOP-TOO, 18 (82%) completed all three assigned cores (main treatment components). All 22 patients completed the Anatomy Core; 20 completed the Medication Core; and 18 completed the Behavior Core. A total of 12 patients | | | |

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| | | | | | | the child, but that it was comprehensive and non-judgemental least helpful components of the program: no clear themes emerged leaved the program helped them and their children: On average, 19/25 items (76%) rated at least "somewhat helpful," no item described as "not at all helpful." On the 1-to 5-point scale, average responses ranged from a low of 2.33 (the program helped reduce the number of times parents had to remind their child to use the bathroom) to a high of 4.2 (the program helped the child feel more comfortable using the toilet at home). Adherence 16/22 patients examined, stopped using the program for some reason other | Reviewer comments: Unclear how encopresis was defined/diagnosed Small sample size, no sample size calculation Unclear whether questionnaires were piloted Source of funding: Partially supported by NIH grant RO1 | | | | |

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| | | | | | | than that their problem was "resolved." -Obstacles to using the program (only 2 items with a mean score of 2 or greater (on a 1- to 3-point scale)): I just forgot [to go to the website]" (mean 2.00, SD 0.89) "I didn't have time in my schedule" (mean 2.06, SD 0.85) | |