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Sedation in children and young people

Foreword

Advances in medicine, over the last 20 years, have increased the demand for invasive investigations and procedures. The type of procedure can range from painless imaging that requires immobility to painful or uncomfortable minor surgery. Whereas adults can cope with these children often need more than simple reassurance and pain relief; they need either sedation or anaesthesia. The problem with sedation is its unpredictability. If managed well, it can be effective but sometimes it is not effective enough unless the doses are increased and this risks causing unconsciousness and suppression of vital protective reflexes leading to potentially dangerous hypoxia. If however, sedation is inadequate, the distress can be remembered for a lifetime and make any subsequent procedure much more difficult. There is a dilemma therefore between giving too much and too little. Anaesthesia, in comparison, is reliable but involves specialist skills and facilities, and may not always be an appropriate use of resources.

There is evidence that large numbers of children in the UK undergo single or repeated procedures and the perception is that there is considerable variation in the services that are provided. The common question asked is "What drugs are safe and effective?" and the Scottish Guideline Network guideline published in 2007 reviewed the evidence and drew useful conclusions. However, at the stakeholder meeting at the inception of this NICE guideline a different concern was raised — "Healthcare practitioners need to be trained to use sedation safely?". In other words, it became clear that the problem was less "What drugs?" but more "Who can administer them?". Indeed, if it can be agreed that a chosen drug technique is effective, people need to know who can use it safely.

In consequence we have had two broad aims. Our first was to review the evidence of efficacy and safety of common drug techniques, and our second was to form a consensus view on what resources are necessary, and this included not only the facilities, the equipment and staff, but also the training of staff to ensure that they have the knowledge, the skills and judgement.

Our guideline development group included doctors, nurses, dentists and a psychologist who were all expert and experienced in working with children. We are especially grateful to our dentists who have been pioneers in this field and to our parent representatives who have made sure we have considered the patient's perspective. In our discussions we soon realised that we would be unable to review and advise on all aspects of sedation and we decided to limit our searches for evidence that would help guide 90% of scenarios. Nevertheless we wanted to make clear statements of principle that are applicable and relevant to all situations.

We began by identifying key questions. We wanted to advise on how patients should be assessed, prepared and managed, and to specify the necessary resources. The psychological needs and behavioural management have also been considered. All these were tackled by consensus methods. Other questions related to whether sedation drugs are effective and safe,

1 and we hoped that these could be answered from published evidence. There is a long list of 2 potentially useful drugs but we decided to choose drugs that were in common use in the UK, 3 and those that could be applied to the "90% of scenarios". In particular we chose not to 4 review evidence for analgesia alone except for those that have a sedative component or 5 those that are commonly used in combination with another sedative. 6 When considering safety of sedation the concepts of "consciousness", "margin of safety" and 7 "target depth" are important. The ideal safe sedation technique is one that can be relied 8 upon to not cause sedation deeper than the target depth of moderate sedation (also known 9 as conscious sedation). At this level the patient responds to stimuli and vital reflexes are 10 active. Drugs with a wide margin of safety have a large difference between the doses that 11 cause moderate sedation and those that depress vital reflexes. 12 Propofol and sevoflurane are potent anaesthetic drugs that are being administered in small 13 doses to achieve short acting and controlled moderate sedation. It is debatable whether these 14 drugs can reliably sedate rather than stray unintentionally beyond the target depth into 15 anaesthesia. The truth probably depends upon the dose and the pain of the procedure, and 16 we decided to consider published evidence about these drugs provided the authors had the 17 intention of causing sedation. 18 Our technical team found surprisingly few high quality published reports and clinical trials. 19 This perhaps was due to the practical difficulties of enrolling sufficient numbers of children 20 into adequately controlled and blinded protocols. We have only considered efficacy data 21 from RCTs but used both cohort studies and RCTs for safety data. 22 Different procedures need different sedation techniques and we wanted to develop a 23 practical algorithm to facilitate effective and safe decisions. We limited ourselves to four 24 common scenarios that are short painful procedures in the emergency department, 25 gastrointestinal endoscopy, dental extractions and painless imaging, and we are confident 26 that guidance for these can be applied to the "90%". 27 The cost-effectiveness of sedation has to be compared with anaesthesia. The "quality of 28 patient experience" is rarely published in clinical trials and when it did it was difficult to 29 interpret. The cost was the more measurable factor and was the cost of the healthcare 30 practitioners involved. However there was disagreement about whether or not the data 31 described the true everyday situation. If sedation fails its cost must take into account the cost 32 of anaesthesia and therefore we wanted to estimate the failure rate that would make the 33 investment of an anaesthesia service worthwhile. 34 A change in sedation services to children has become necessary because demand has 35 increased. Moreover change is within our grasp if healthcare professionals work together to 36 improve standards. My GDG colleagues and I have been privileged to develop this guideline 37 and it is our sincerest hope that it will make a significant contribution to making diagnostic 38 and therapeutic procedures less distressing and safer for children and young people. 39 Mike Sury 40 Chair, Guideline Development Group

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Guideline Review Panel

The Guideline Review Panel is an independent panel that oversees the development of the guideline and takes responsibility for monitoring concordance to NICE guideline development processes. In particular, the panel ensures that stakeholder comments have been adequately considered and responded to. The panel includes members from the following perspectives.

1 2

Stakeholder Involvement

3 To be added after consultation

Abbreviations

A&E Accident and Emergency

AGREE Appraisal of Guidelines Research and Evaluation

ALS Advanced Life Support
ANCOVA Analysis of covariance

ASA American Society of Anesthiologists

BNF British National Formulary

BLS Basic Life Support

CCA Cost-consequences analysis
CEA Cost-effectiveness analysis

CI Confidence interval

CPR Cardiopulmonary Resuscitation
CT Computerised Tomography

CUA Cost-utility analysis

DH Department of Health

ED Emergnecy Department

GA General Anaesthesia

GDG Guideline Development Group

GI Gastrointestinal

GP General Practitioner

GRADE Grading of Recommendations Assessment, Development and Evaluation

GRP Guideline Review Panel

HRQL Health-related quality of life
HTA Health technology assessment
ICC Intraclass correlation coefficient

ICER Incremental cost-effectiveness ratio

ILS Intermediate Life Support

IM Intramuscular
IN Intranasal

INB Incremental net benefit

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Inh Inhaled

IQR Inter-quartile range
ITT Intention to treat

IV Intravenous
Los Length of Stay

LY Life-year

MHRA Medicines and Healthcare Products Regulatory Agency

MRI Magnetic Resonance Imaging
MTC Mixed-treatment comparisons

NCGC National Clinical Guidelines Centre

NHS National Health Service

NICE National Institute for Health and Clinical Excellence

NNT Number needed to treat

OGD Oesophago-Gastro Duodenoscopy

OR Odds ratio

PASA NHS Purchasing and Supply Agency

PICO Framework incorporating patients, interventions, comparison and

outcome

PPIP Patient and Public Involvement Programme

PSA Probabilistic sensitivity analysis

QALY Quality-adjusted life year

RCA Royal College of Anaesthetists

RCN Royal College of Nursing
RCT Randomised controlled trial

RR Relative risk

SD Standard deviation
SR Systematic review

VS. Versus

Glossary of Terms

Absolute risk reduction (Risk difference)

1

The difference in the risk of an event between two groups (one

subtracted from the other) in a comparative study.

Abstract Summary of a study, which may be published alone or as an

introduction to a full scientific paper.

Adherence The extent to which the patient's behaviour matches the prescriber's

recommendations. Adherence emphasises the need for agreement and that the patient is free to decide whether or not to adhere to the

doctor's recommendation. 105

Adjustment A statistical procedure in which the effects of differences in

composition of the populations being compared (or treatment given

at the same time) have been minimised by statistical methods.

Advanced Life Support

Advanced Life Support is the management of the child or young person who is deteriorating, in respiratory arrest or in cardiac arrest. Senior healthcare professionals (doctors, nurses, paramedics) work together in a structured team environment in managing the child or young person, with advanced skills in airway management and ventilation, chest compression, administration of life support drugs

and support to the child or young person's family/carers.

Algorithm (in guidelines)

A flow chart of the clinical decision pathway described in the guideline, where decision points are represented with boxes, linked

with arrows.

Allocation concealment

The process used to prevent advance knowledge of group assignment in a RCT. The allocation process should be impervious to any influence by the individual making the allocation, by being administered by someone who is not responsible for recruiting

participants.

Anaesthetic agent A drug used to cause general anaesthesia. Anaesthetic agents are

potent and reliably cause anaesthesia but they may be given in low or "sub-anaesthetic" doses to cause sedation. Sedation techniques using anaesthetic agents have been called "narrow margin of safety" techniques because the difference between the sedation dose and

the angesthesia dose is small.

Applicability The degree to which the results of an observation, study or review

are likely to hold true in a particular clinical practice setting.

Appraisal of An international collaboration of researchers and policy makers

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Guidelines Research and Evaluation, (AGREE) whose aim is to improve the quality and effectiveness of clinical practice guidelines (http://www.agreecollaboration.org). The AGREE instrument, developed by the group, is designed to assess the quality of clinical guidelines.

Arm (of a clinical study)

Sub-section of individuals within a study who receive one particular intervention, for example placebo arm

Association

Statistical relationship between two or more events, characteristics or other variables. The relationship may or may not be causal.

Audit See 'Clinical audit'.

Baseline The initial set of measurements at the beginning of a study (after run-

in period where applicable), with which subsequent results are

compared.

Basic Life Support Basic Life Support is the maintenance of a child or young person's

airway and support of breathing and the circulation without using equipment other than a simple airway device or pocket mask. A combination of expired air ventilation (rescue breathing) and chest compression is known as cardiopulmonary resuscitation (CPR).

Bias Systematic (as opposed to random) deviation of the results of a study

from the 'true' results that is caused by the way the study is designed

or conducted.

Blinding (masking) Keeping the study participants, caregivers, researchers and outcome

assessors unaware about the interventions to which the participants

have been allocated in a study.

Capital costs Costs of purchasing major capital assets (usually land, buildings or

equipment). Capital costs represent investments at one point in time.

Carer (caregiver) Someone other than a health professional who is involved in caring

for a person with a medical condition.

Case-control study Comparative observational study in which the investigator selects

individuals who have experienced an event (For example, developed a disease) and others who have not (controls), and then collects data

to determine previous exposure to a possible cause.

Case series Report of a number of cases of a given disease, usually covering the

course of the disease and the response to treatment. There is no

comparison (control) group of patients.

Clinical audit A quality improvement process that seeks to improve patient care

and outcomes through systematic review of care against explicit

criteria and the implementation of change.

Clinical efficacy The extent to which an intervention is active when studied under

controlled research conditions.

Clinical effectiveness The extent to which an intervention produces an overall health

benefit in routine clinical practice.

Clinical impact The effect that a guideline recommendation is likely to have on the

treatment or treatment outcomes, of the target population.

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Clinical question In guideline development, this term refers to the questions about

treatment and care that are formulated to guide the development of

evidence-based recommendations.

Clinician A healthcare professional providing direct patient care, for example

doctor, nurse or physiotherapist.

Cluster A closely grouped series of events or cases of a disease or other

related health phenomena with well-defined distribution patterns, in relation to time or place or both. Alternatively, a grouped unit for

randomisation.

Cochrane Library A regularly updated electronic collection of evidence-based

medicine databases including the Cochrane Database of Systematic

Reviews.

Cochrane Review A systematic review of the evidence from randomised controlled

trials relating to a particular health problem or healthcare intervention, produced by the Cochrane Collaboration. Available

electronically as part of the Cochrane Library.

Cohort study A retrospective or prospective follow-up study. Groups of individuals

to be followed up are defined on the basis of presence or absence of exposure to a suspected risk factor or intervention. A cohort study can be comparative, in which case two or more groups are selected on the basis of differences in their exposure to the agent of interest.

Co-morbidity Co-existence of more than one disease or an additional disease

(other than that being studied or treated) in an individual.

Comparability Similarity of the groups in characteristics likely to affect the study

results (such as health status or age).

Compliance The extent to which a person adheres to the health advice agreed

with healthcare professionals. May also be referred to as

'adherence' or 'concordance'.

Concordance This is a recent term whose meaning has changed. It was initially

applied to the consultation process in which doctor and patient agree therapeutic decisions that incorporate their respective views, but now includes patient support in medicine taking as well as prescribing communication. Concordance reflects social values but does not address medicine-taking and may not lead to improved adherence.

Conference proceedings

Compilation of papers presented at a conference.

Confidence interval

(CI)

A range of values for an unknown population parameter with a stated 'confidence' (conventionally 95%) that it contains the true value. The interval is calculated from sample data, and generally straddles the sample estimate. The 'confidence' value means that if the method used to calculate the interval is repeated many times, then that proportion of intervals will actually contain the true value.

Confounding

In a study, confounding occurs when the effect of an intervention on an outcome is distorted as a result of an association between the population or intervention or outcome and another factor (the 'confounding variable') that can influence the outcome independently of the intervention under study.

Consensus methods

Techniques that aim to reach an agreement on a particular issue. Formal consensus methods include Delphi and nominal group techniques, and consensus development conferences. In the development of clinical guidelines, consensus methods may be used where there is a lack of strong research evidence on a particular topic. Expert consensus methods will aim to reach agreement between experts in a particular field.

Control group

A group of patients recruited into a study that receives no treatment, a treatment of known effect, or a placebo (dummy treatment) - in order to provide a comparison for a group receiving an experimental treatment, such as a new drug.

Controlled clinical trial(CCT)

A study testing a specific drug or other treatment involving two (or more) groups of patients with the same disease. One (the experimental group) receives the treatment that is being tested, and the other (the comparison or control group) receives an alternative treatment, a placebo (dummy treatment) or no treatment. The two groups are followed up to compare differences in outcomes to see how effective the experimental treatment was. A CCT where patients are randomly allocated to treatment and comparison groups is called a randomised controlled trial.

Cost benefit analysis

A type of economic evaluation where both costs and benefits of healthcare treatment are measured in the same monetary units. If benefits exceed costs, the evaluation would recommend providing the treatment.

Cost-consequences analysis (CCA)

A type of economic evaluation where various health outcomes are reported in addition to cost for each intervention, but there is no overall measure of health gain.

Cost-effectiveness analysis (CEA)

An economic study design in which consequences of different interventions are measured using a single outcome, usually in 'natural' units (For example, life-years gained, deaths avoided, heart attacks avoided, cases detected). Alternative interventions are then compared in terms of cost per unit of effectiveness.

Cost-effectiveness model

An explicit mathematical framework, which is used to represent clinical decision problems and incorporate evidence from a variety of sources in order to estimate the costs and health outcomes.

Cost-utility analysis (CUA)

A form of cost-effectiveness analysis in which the units of effectiveness are quality-adjusted life-years (QALYs).

Credible interval

The Bayesian equivalent of a confidence interval.

Decision analysis An explicit quantitative approach to decision making under

uncertainty, based on evidence from research. This evidence is translated into probabilities, and then into diagrams or decision trees which direct the clinician through a succession of possible scenarios,

actions and outcomes.

Decision problem A clear specification of the interventions, patient populations and

outcome measures and perspective adopted in an evaluation, with an explicit justification, relating these to the decision which the analysis is

to inform.

Deep sedation Drug-induced depression of consciousness during which patients are

asleep and cannot be easily roused but do respond purposefully to repeated or painful stimulation. The ability to maintain ventilatory function independently may be impaired. Patients may require assistance in maintaining a patent airway. Spontaneous ventilation may be inadequate. Cardiovascular function is usually maintained.

DiscountingCosts and perhaps benefits incurred today have a higher value than

costs and benefits occurring in the future. Discounting health benefits reflects individual preference for benefits to be experienced in the present rather than the future. Discounting costs reflects individual preference for costs to be experienced in the future rather than the

present.

Dominance An intervention is said to be dominated if there is an alternative

intervention that is both less costly and more effective.

Dosage The prescribed amount of a drug to be taken, including the size and

timing of the doses.

Double

blind/masked study (investigator/clinician) is aware of which treatment nor intervention the subject is receiving. The purpose of blinding/masking is to protect

the subject is receiving. The purpose of blinding/masking is to protect

against bias.

Drop-out A participant who withdraws from a clinical trial before the end.

Economic evaluation Comparative analysis of alternative health strategies (interventions

or programmes) in terms of both their costs and consequences.

A study in which neither the subject (patient) nor the observer

Effect (as in effect measure, treatment effect, estimate of effect, effect size) The observed association between interventions and outcomes or a statistic to summarise the strength of the observed association.

Effectiveness See 'Clinical effectiveness'.

Efficacy See 'Clinical efficacy'.

Epidemiological

study

The study of a disease within a population, defining its incidence and

prevalence and examining the roles of external influences (For

example, infection, diet) and interventions.

Equity Fair distribution of resources or benefits.

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Evidence Information on which a decision or guidance is based. Evidence is

obtained from a range of sources including randomised controlled trials, observational studies, expert opinion (of clinical professionals

and/or patients).

Evidence table A table summarising the results of a collection of studies which, taken

together, represent the evidence supporting a particular recommendation or series of recommendations in a guideline.

Exclusion criteria (literature review)

Explicit standards used to decide which studies should be excluded

from consideration as potential sources of evidence.

Exclusion criteria (clinical study)

Criteria that define who is not eligible to participate in a clinical

study.

Expert consensus See 'Consensus methods'.

Extended dominance If Option A is both more clinically effective than Option B and has a

lower cost per unit of effect, when both are compared with a donothing alternative then Option A is said to have extended

dominance over Option B. Option A is therefore more efficient and

Extrapolation In data analysis, predicting the value of a parameter outside the

range of observed values.

Follow up Observation over a period of time of an individual, group or initially

should be preferred, other things remaining equal.

defined population whose appropriate characteristics have been assessed in order to observe changes in health status or health-

related variables.

General anaesthesia Drug-induced loss of consciousness during which patients are not

rousable, even by painful stimulation. Patients often require assistance in maintaining a patent airway. Ventilatory function is often impaired. Positive pressure ventilation may be required because of depressed spontaneous ventilation or drug-induced depression of neuromuscular function. Cardiovascular function may

be impaired.

Generalisability The extent to which the results of a study based on measurement in a

particular patient population and/or a specific context hold true for another population and/or in a different context. In this instance, this is the degree to which the guideline recommendation is applicable across both geographical and contextual settings. For instance, guidelines that suggest substituting one form of labour for another should acknowledge that these costs might vary across the country.

Gold standard See 'Reference standard'.

Goodness-of-fit How well a statistical model or distribution compares with the

observed data.

Grey literature Reports that are unpublished or have limited distribution, and are

not included in the common bibliographic retrieval systems.

Harms Adverse effects of an intervention.

Healthcare professional

For the purposes of this guideline the term 'healthcare professional' refers to a trained and registered individual involved the care of a sedated patient: and includes doctor, dentist or nurse.

Health economics

The study of the allocation of scarce resources among alternative healthcare treatments. Health economists are concerned with both increasing the average level of health in the population and improving the distribution of health.

Health-related quality of life

A combination of an individual's physical, mental and social well-being; not merely the absence of disease.

Heterogeneity

Or lack of homogeneity. The term is used in meta-analyses and systematic reviews when the results or estimates of effects of treatment from separate studies seem to be very different – in terms of the size of treatment effects or even to the extent that some indicate beneficial and others suggest adverse treatment effects. Such results may occur as a result of differences between studies in terms of the patient populations, outcome measures, definition of variables or duration of follow-up.

Homogeneity

This means that the results of studies included in a systematic review or meta-analysis are similar and there is no evidence of heterogeneity. Results are usually regarded as homogeneous when differences between studies could reasonably be expected to occur by chance.

Hypothesis

A supposition made as a starting point for further investigation.

Inclusion criteria (literature review)

Explicit criteria used to decide which studies should be considered as potential sources of evidence.

Incremental analysis

The analysis of additional costs and additional clinical outcomes with different interventions.

Incremental cost

The mean cost per patient associated with an intervention minus the mean cost per patient associated with a comparator intervention.

Incremental cost effectiveness ratio (ICER)

The difference in the mean costs in the population of interest divided by the differences in the mean outcomes in the population of interest for one treatment compared with another.

 $ICER = (Cost_A - Cost_B) / (Effectiveness_A - Effectiveness_B).$

Incremental net benefit (INB)

The value (usually in monetary terms) of an intervention net of its cost compared with a comparator intervention. The INB can be calculated for a given cost-effectiveness (willingness to pay) threshold. If the threshold is £20,000 per QALY gained then the INB is calculated as: $(£20,000 \times QALYs \text{ gained})$ – Incremental cost.

Index

In epidemiology and related sciences, this word usually means a rating scale, for example, a set of numbers derived from a series of observations of specified variables. Examples include the various health status indices, and scoring systems for severity or stage of cancer.

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Indication (specific) The defined use of a technology as licensed by the Medicines and

Healthcare products Regulatory Agency (MHRA).

Intention-to-treat analysis (ITT analysis) An analysis of the results of a clinical study in which the data are analysed for all study participants as if they had remained in the group to which they were randomised, regardless of whether or not they remained in the study until the end, crossed over to another

treatment or received an alternative intervention.

Intermediate Life Support Intermediate Life Support is the initiation of cardiopulmonary resuscitation in the clinical setting, including effective chest compressions and ventilation and early safe defibrillation. Those healthcare professionals with intermediate life support skills are able to utilise a wider range of life support adjuncts (such as the laryngeal mask) and should also recognise the child or young person who is at risk of deterioration, therefore preventing cardiac arrest.

Intermediate outcomes

Outcomes that are related to the outcome of interest but may be more easily assessed within the context of a clinical study: for example, intraocular pressure reduction is related to the risk of conversion to COAG or COAG progression.

Internal validity

The degree to which the results of a study are likely to approximate the 'truth' for the participants recruited in a study (that is, are the results free of bias?). It refers to the integrity of the design and is a prerequisite for applicability (external validity) of a study's findings. See 'External validity'.

Intervention

Healthcare action intended to benefit the patient, for example, drug treatment, surgical procedure, psychological therapy.

Intraoperative

The period of time during a surgical procedure.

Kappa statistic

An index which compares the agreement against that which might be expected by chance

Length of stay

The total number of days a participant stays in hospital.

Licence

See 'Product licence'.

Life-years gained

Mean average years of life gained per person as a result of the intervention compared with an alternative intervention.

Literature review

An article that summarises the evidence contained in a number of different individual studies and draws conclusions about their findings. It may or may not be systematically researched and developed.

Markov model

A method for estimating long term costs and effects for recurrent or chronic conditions, based on health states and the probability of transition between them within a given time period (cycle).

Medical devices

All products, except medicines, used in healthcare for the diagnosis, prevention, monitoring or treatment of illness or handicap.

Medicines and Healthcare Products Regulatory Agency (MHRA) The Executive Agency of the Department of Health protecting and promoting public health and patient safety by ensuring that medicines, healthcare products and medical equipment meet appropriate standards of safety, quality, performance and effectiveness, and are used safely.

Meta-analysis

A statistical technique for combining (pooling) the results of a number of studies that address the same question and report on the same outcomes to produce a summary result. The aim is to derive more precise and clear information from a large data pool. It is generally more reliably likely to confirm or refute a hypothesis than the individual trials.

Minimal sedation

A drug-induced state during which patients are awake and calm, and respond normally to verbal commands. Although cognitive function and coordination may be impaired, ventilatory and cardiovascular functions are unaffected.

Moderate sedation (also known as conscious sedation): Drug-induced depression of consciousness during which patients are sleepy but respond purposefully to verbal commands, either alone or accompanied by light tactile stimulation (reflex withdrawal from a painful stimulus is not a purposeful response). No interventions are required to maintain a patent airway. Spontaneous ventilation is adequate. Cardiovascular function is usually maintained.

Multivariate model

A statistical model for analysis of the relationship between two or more predictor (independent) variables and the outcome (dependent) variable.

Narrative summary

Summary of findings given as a written description.

Number needed to treat (NNT)

The number of patients that who on average must be treated to prevent a single occurrence of the outcome of interest.

Observational study

Retrospective or prospective study in which the investigator observes the natural course of events with or without control groups; for example, cohort studies and case—control studies.

Odds ratio

A measure of treatment effectiveness. The odds of an event happening in the treatment group, expressed as a proportion of the odds of it happening in the control group. The 'odds' is the ratio of events to non-events.

Off-label

A drug or device used treat a condition or disease for which it is not specifically licensed.

Older people

People over the age of 65 years.

Operating costs

Ongoing costs of carrying out an intervention, excluding capital costs.

Opportunity cost

The opportunity cost of investing in a healthcare intervention is the loss of other healthcare programmes that are displaced by its introduction. This may be best measured by the health benefits that could have been achieved had the money been spent on the next best alternative healthcare intervention.

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Outcome Measure of the possible results that may stem from exposure to a

preventive or therapeutic intervention. Outcome measures may be intermediate endpoints or they can be final endpoints. See

'Intermediate outcome'.

P value The probability that an observed difference could have occurred by

chance, assuming that there is in fact no underlying difference between the means of the observations. If the probability is less than 1 in 20, the P value is less than 0.05; a result with a P value of less than 0.05 is conventionally considered to be 'statistically significant'.

Peer review A process where research is scrutinised by experts that have not

been involved in the design or execution of the studies.

Perioperative The period from admission through surgery until discharge,

encompassing preoperative and post-operative periods.

Placebo An inactive and physically identical medication or procedure used as

a comparator in controlled clinical trials.

Placebo effect A beneficial (or adverse) effect produced by a placebo and not due

to any property of the placebo itself.

Postoperative Pertaining to the period after patients leave the operating theatre,

following surgery.

Preoperative Pertaining to the period before surgery commences.

Primary care Healthcare delivered to patients outside hospitals. Primary care

covers a range of services provided by GPs, nurses and other healthcare professionals, dentists, pharmacists and opticians.

Primary research Study generating original data rather than analysing data from

existing studies (which is called secondary research).

Product licence An authorisation from the MHRA to market a medicinal product.

Prognosis A probable course or outcome of a disease. Prognostic factors are

patient or disease characteristics that influence the course. Good prognosis is associated with low rate of undesirable outcomes; poor prognosis is associated with a high rate of undesirable outcomes.

Prospective study A study in which people are entered into the research and then

followed up over a period of time with future events recorded as they happen. This contrasts with studies that are *retrospective*.

Qualitative research Research concerned with subjective outcomes relating to social,

emotional and experiential phenomena in health and social care.

Quality of life See 'Health-related quality of life'.

Quality-adjusted life year (QALY)

An index of survival that is adjusted to account for the patient's quality of life during this time. QALYs have the advantage of incorporating changes in both quantity (longevity/mortality) and quality (morbidity, psychological, functional, social and other factors) of life. Used to measure benefits in cost-utility analysis. The QALYs gained are the mean QALYs associated with one treatment minus the mean QALYs associated with an alternative treatment.

Quantitative research

Research that generates numerical data or data that can be converted into numbers, for example clinical trials or the national Census which counts people and households.

Quick Reference Guide

An abridged version of NICE guidance, which presents the key priorities for implementation and summarises the recommendations for the core clinical audience.

Randomisation

Allocation of participants in a research study to two or more alternative groups using a chance procedure, such as computergenerated random numbers. This approach is used in an attempt to ensure there is an even distribution of participants with different characteristics between groups and thus reduce sources of bias.

Randomised controlled trial (RCT)

A comparative study in which participants are randomly allocated to intervention and control groups and followed up to examine differences in outcomes between the groups.

RCT See 'Randomised controlled trial'.

Reference standard

The test that is considered to be the best available method to establish the presence or absence of the outcome – this may not be the one that is routinely used in practice.

Relative risk (RR)

The number of times more likely or less likely an event is to happen in one group compared with another (calculated as the risk of the event in group A/the risk of the event in group B).

Remit

The brief given by the Department of Health and Welsh Assembly Government at the beginning of the guideline development process. This defines core areas of care that the guideline needs to address.

Resource implication

The likely impact in terms of finance, workforce or other NHS resources.

Retrospective study

A retrospective study deals with the present/ past and does not involve studying future events. This contrasts with studies that are prospective.

Secondary benefits

Benefits resulting from a treatment in addition to the primary, intended outcome.

Sedation

Sedation is a state of depressed consciousness. There are depths or levels of sedation that range from minor to major depression of consciousness. Whereas depression of consciousness is a continuum, with no clear boundaries between levels, three levels of sedation have been defined and are in common use: minimal, moderate and

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deep sedation; they are recommended internationally ^{1,6,44,194}. The target level of sedation is the level that is intended for the patient. The level of sedation can vary according to the drug, the dose, the patient and the stimulus of the procedure. The level of sedation varies over time due to two main factors: the change in the concentration of the sedation drug within the patient and the variation in the stimulation that opposes sedation.

Sedation nurse

A registered nurse trained to both deliver sedation and manage the sedated patient.

Sedationist

A healthcare professional who is trained to both deliver sedation and manage the sedated patient.

Sedative

A drug that causes minimal, moderate or deep sedation. All sedation drugs have a variable effect on conscious level. Some sedation drugs may either not be effective enough or cause sedation deeper than the intended target level. High or excessive doses of sedatives may cause unintended deep sedation or anaesthesia. Sedation drugs or techniques that are unlikely to cause anaesthesia have been called drugs with a "wide margin of safety" because they are unlikely to cause appreciable depression of airway reflexes or breathing.

Selection bias (also allocation bias)

A systematic bias in selecting participants for study groups, so that the groups have differences in prognosis and/or therapeutic sensitivities at baseline. Randomisation (with concealed allocation) of patients protects against this bias.

Selection criteria

Explicit standards used by guideline development groups to decide which studies should be included and excluded from consideration as potential sources of evidence.

Sensitivity

Sensitivity or recall rate is the proportion of true positives which are correctly identified as such. For example in diagnostic testing it is the proportion of true cases that the test detects.

See the related term 'Specificity'

Sensitivity analysis

A means of representing uncertainty in the results of economic evaluations. Uncertainty may arise from missing data, imprecise estimates or methodological controversy. Sensitivity analysis also allows for exploring the generalisability of results to other settings. The analysis is repeated using different assumptions to examine the effect on the results.

One-way simple sensitivity analysis (univariate analysis): each parameter is varied individually in order to isolate the consequences of each parameter on the results of the study.

Multi-way simple sensitivity analysis (scenario analysis): two or more parameters are varied at the same time and the overall effect on the results is evaluated.

Threshold sensitivity analysis: the critical value of parameters above or below which the conclusions of the study will change are identified.

Probabilistic sensitivity analysis: probability distributions are assigned to the uncertain parameters and are incorporated into evaluation models based on decision analytical techniques (For example, Monte Carlo simulation).

Specialist in Sedation

Healthcare professional experienced in delivering sedation in children with more complex medical conditions.

Specificity

The proportion of true negatives that a correctly identified as such. For example in diagnostic testing the specificity is the proportion of non-cases incorrectly diagnosed as cases.

See related term 'Sensitivity'.

In terms of literature searching a highly specific search is generally narrow and aimed at picking up the key papers in a field and avoiding a wide range of papers.

Stakeholder

Those with an interest in the use of the guideline. Stakeholders include manufacturers, sponsors, healthcare professionals, and patient and carer groups.

Statistical power

The ability to demonstrate an association when one exists. Power is related to sample size; the larger the sample size, the greater the power and the lower the risk that a possible association could be missed.

Synthesis of evidence

A generic term to describe methods used for summarising (comparing and contrasting) evidence into a clinically meaningful conclusion in order to answer a defined clinical question. This can include systematic review (with or without meta-analysis), qualitative and narrative summaries.

Systematic review

Research that summarises the evidence on a clearly formulated question according to a pre-defined protocol using systematic and explicit methods to identify, select and appraise relevant studies, and to extract, collate and report their findings. It may or may not use statistical meta-analysis.

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Time horizon The time span used in the NICE appraisal which reflects the period

over which the main differences between interventions in health effects and use of healthcare resources are expected to be experienced, and taking into account the limitations of supportive

evidence.

Treatment allocation Assigning a participant to a particular arm of the trial.

Treatment options The choices of intervention available.

Utility A measure of the strength of an individual's preference for a specific

health state in relation to alternative health states. The utility scale assigns numerical values on a scale from 0 (death) to 1 (optimal or 'perfect' health). Health states can be considered worse than death

and thus have a negative value.

1 Introduction

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Our clinical guidelines are recommendations for the care of individuals in specific clinical conditions or circumstances within the National Health Service (NHS) – from prevention and self-care through primary and secondary care to more specialised services. We base our clinical guidelines on the best available research evidence, with the aim of improving the quality of health care. We use predetermined and systematic methods to identify and evaluate the evidence relating to specific clinical questions.

- 9 Clinical guidelines can:
 - provide recommendations for the treatment and care of people by health professionals
 - be used to develop standards to assess the clinical practice of individual health professionals
 - be used in the education and training of health professionals
- help patients to make informed decisions
 - improve communication between patient and health professional
- While guidelines assist the practice of healthcare professionals, they do not replace their knowledge and skills.
- We produce our guidelines using the following steps:
 - Guideline topic is referred to the National Institute for Health and Clinical Excellence (NICE) from the Department of Health
 - Stakeholders register an interest in the guideline and are consulted throughout the development process.
- The scope is prepared by the National Clinical Guideline Centre (NCGC)
 - The NCGC establish a guideline development group

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1 2		 A draft guideline is produced after the group assesses the available evidence and makes recommendations
3		There is a consultation on the draft guideline.
4		The final guideline is produced.
5 6		The National Clinical Guideline Centre and NICE produce a number of versions of this guideline:
7 8		 the full guideline contains all the recommendations, plus details of the methods used and the underpinning evidence
9 10		 the NICE guideline presents the recommendations from the full version in a format suited to implementation by health professionals and NHS bodies
11 12		 the quick reference guide presents recommendations in a suitable format for health professionals
13 14		 information for the public ('understanding NICE guidance') is written using suitable language for people without specialist medical knowledge.
15 16		This version is the full version. The other versions are available from NICE www.NICE.org.uk.
17	1.2	The need for this guideline
18 19 20 21 22 23 24 25 26 27 28 29 30		Many children present to hospitals and dental clinics needing effective sedation or anaesthesia for painful or distressing diagnostic or therapeutic procedures. There are many sedation techniques available but there is insufficient guidance on which techniques are effective and what resources are required to deliver them safely. Sedation is not always effective enough and will occasionally require the procedure to be delayed until the child can be anaesthetised perhaps in another healthcare setting or on another day. Consequently sedation failure is both distressing for the child and has major NHS cost implications. Excessive doses of sedation can cause unintended loss of consciousness and dangerous hypoxia. Planned anaesthesia, in comparison, is effective but may have resource implications. The need for sedation or anaesthesia will depend upon the type of procedure. Some types of procedures are very common and healthcare providers and practitioners need to understand whether sedation or anaesthesia is the most cost effective method of managing them
31	1.3	The National Clinical Guideline Centre

This guideline was commissioned by NICE and developed by the NCGC. The NCGC is one of four national collaborating centres (Cancer, Women and Children's Health, Mental Health and the NCGC) funded by NICE and comprises a partnership between a variety of academic, professional and patient-based organisations. As a multidisciplinary centre we draw upon the expertise of the healthcare professions and academics and ensure the involvement of patients in our work.

1.4 Remit

- The following remit was received by the NCGC from the Department of Health in March 2008 as part of NICE's 18th wave programme of work.
- 4 The Department of Health asked NICE:
 - "To prepare a clinical guideline on sedation for diagnostic and therapeutic procedures in infants, children and young people up to the age of 19."

1.5 What the guideline covers

Clinical need for the guideline:

- In adults, many procedures can be undertaken with local anaesthesia and reassurance. In children and young people this is often not possible because the procedures are too frightening, too painful and need to be carried out in children who may be ill, or in pain or have behavioural problems. Therefore special consideration is necessary for children and young people undergoing distressing procedures.
- It is estimated that more than 2 million children and young people are taken to emergency departments each year following accidental injury. Many of these children and young people will undergo procedures that require sedation. For example, in 2005–6 there were 866 children aged 14 and younger who required a closed reduction of a dislocated joint. Sedation is also frequently used for invasive diagnostic procedures such as lumbar punctures, bone marrow biopsies and endoscopies. In 2005–6 there were 4700 gastroscopies, 9000 diagnostic spinal punctures and 2100 bone marrow biopsies carried out on children aged 14 and younger. Sedation is also commonly used in dental practice where the use of general anaesthesia is now restricted to the hospital setting.
- Sedation is only one of the management options available for children and young people undergoing therapeutic or diagnostic procedures. Non-pharmacological techniques may also be useful in reducing anxiety and managing behaviour, and analgesia may be used to provide pain control. These techniques may be used in combination with sedation or as an alternative to sedation. Another alternative to using sedation for diagnostic or therapeutic procedures is to carry out the procedure under general anaesthesia, in which case the usual standards of care for patients undergoing anaesthesia must be met.
- Sedation is a drug-induced depression of consciousness. The aims of sedation during diagnostic or therapeutic procedures may include reducing fear and anxiety, providing pain control and minimising movement. The importance of each of these aims will vary depending on the nature of the procedure and the characteristics of the patient. For example, in younger children sedation may be necessary to ensure that movement is minimised during non-painful procedures such as a magnetic resonance imaging (MRI) scan; in older children sedation may be necessary to minimise the physical and psychological consequences of a painful procedure such as a lumbar puncture.

the awake state, through progressively deeper levels of sedation to anaesthesia. Anaesthesia is an unresponsive state in which vital airway and breathing reflexes are likely to be suppressed. The American Society of Anesthesiologists (ASA) has published useful definitions of sedation levels, classifying them as 'minimal', 'moderate' and 'deep'. Minimal sedation equates to anxiolysis and has no appreciable effect on vital reflexes. In a state of moderate sedation the patient is able to breathe adequately without assistance and responds purposefully to verbal stimulus or tactile stimulation. This is often referred to as conscious sedation. During deep sedation, the patient cannot be roused easily but will respond purposefully to repeated or painful stimuli and may require assistance with their airway or breathing. The level of sedation that is appropriate will depend on the nature of the procedure and the needs of the individual. Deeper levels of sedation require more advanced management because the patient's protective reflexes are affected and they have the potential to progress to anaesthesia.

The effect of sedation drugs on consciousness level is a continuum ranging from

- The level of sedation achieved depends on the drug used and the dose at which
 it is given. When choosing between sedation techniques, healthcare professionals
 must consider the effectiveness of the drug in achieving the required level of
 sedation, the duration of that effect, and the margin of safety between the dose
 required to achieve sedation and the dose that is likely to cause anaesthesia.
- There may be serious adverse effects if the level of sedation is greater than
 intended. If breathing is unintentionally depressed and this complication is not
 recognised and managed appropriately, then this may lead to hypoxic brain
 injury or death. Sedation drugs may also have other unexpected adverse effects
 such as prolonged emergence, paradoxical excitement or post-sedation nausea
 and vomiting.
- If sedation is unsuccessful, this can result in a painful and traumatic experience for the child. It may be necessary to complete the procedure under general anaesthesia or the procedure may need to be abandoned and rescheduled. If the child becomes distressed due to a failure to provide adequate sedation, their parent or carer may chose to refuse consent for further procedures. A distressing experience may also have long-term psychological consequences for the patient, especially if they are required to undergo repeated procedures.
- There is significant variation in practice across the NHS, with sedation being carried out by a variety of healthcare professionals using a wide range of techniques, within different clinical settings. The Scottish Intercollegiate Guidelines Network (SIGN) published a guideline on this topic in 2004. This covered moderate sedation but not deep sedation, and the evidence base it considered has not been updated since 2002. The aim of this guideline is to provide recommendations to both improve the effectiveness and safety of all types of procedural sedation and to reduce current variations in standards of care.

Groups that will be covered:

1 Infants, children and young people (under 19 years) receiving sedation by any 2 technique for painful or non-painful diagnostic or therapeutic procedures. 3 The GDG will consider whether different recommendations are required for 4 different age groups in the population. 5 Healthcare setting: 6 Hospital settings, including inpatients, outpatients, radiology and emergency 7 departments. 8 Primary care, including dental and medical general practice settings. 9 Clinical management 10 Assessment of the patient to determine whether sedation is appropriate. 11 Clear communication, in a child-friendly manner, of information relating to the 12 preparation required for the procedure or investigation, and related sedation 13 technique. This will include the needs of the patient and their parents or carers, 14 ensuring that implications (sedation safety and efficacy) are clearly understood 15 by both the patient and their parent or carer prior to informed consent. 16 Preparation required for the procedure or investigation and related sedation 17 technique. 18 The clinical environment, including the availability of equipment, facilities and 19 20 Patient monitoring during and after sedation and criteria for discharge following 21 sedation. 22 The effectiveness, safety and limitations of sedation techniques. This will include 23 the use of sedation in combination with non-pharmacological techniques and in 24 combination with analgesia. Note that guideline recommendations will normally 25 fall within licensed indications. Where clearly supported by evidence, use outside 26 a licensed indication may be recommended. The guideline will assume that 27 prescribers will use a drug's summary of product characteristics and the 'British 28 National Formulary for Children' to inform their decisions for individual patients. 29 The Guideline Development Group (GDG) will take reasonable steps to identify 30 ineffective interventions and approaches to care. If robust and credible 31 recommendations for re-positioning the intervention for optimal use, or changing 32 the approach to care to make more efficient use of resources, can be made, they 33 will be clearly stated. If the resources released are substantial, consideration will 34 be given to listing such recommendations in the 'Key priorities for implementation' 35 section of the guideline. 36

1		Training and competence:
2 3 4		 Training for practitioners involved in procedural sedation, irrespective of specialty background, that will be relevant to the sedation techniques and the clinical environment.
5 6 7 8		 Training that enables practitioners to be competent in the practical aspects of effective and safe delivery of sedation techniques relevant to the clinical situation, and the management of adverse events (for example, airway management skill in the inadvertently anaesthetised patient).
9	1.6	What the guideline does not cover
10		Groups that will not be covered
11 12		 Patients requiring sedation for purposes other than for diagnostic or therapeutic procedures including:
13		o sedation in critically ill patients requiring mechanical ventilation
14		o sedation in palliative care
15		o sedation in the treatment of mental health conditions
16 17		 sedation given as premedication for general anaesthesia or as postoperative analgesia
18		o night sedation
19		• Patients having diagnostic or therapeutic procedures under general anaesthesia.
20	1.7	Who developed this guideline?
21 22 23		A multidisciplinary GDG comprising professional group members and consumer representatives of the main stakeholders developed this guideline (see section on Guideline Development Group Membership and acknowledgements).
24 25 26		NICE funds the NCGC and thus supported the development of this guideline. The GDG was convened by the NCGC and chaired by Dr Mike Sury in accordance with guidance from NICE.
27 28 29 30 31		The group met every 6-8 weeks during the development of the guideline. At the start of the guideline development process all GDG members declared interests including consultancies, fee-paid work, share-holdings, fellowships and support from the healthcare industry. At all subsequent GDG meetings, members declared arising conflicts of interest, which were also recorded (Appendix B).
32 33 34		Members are either required to withdraw completely or for part of the discussion if their declared interest makes it appropriate, however this was not deemed necessary for any aroup members on this guideline.

2 Methodology

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This guideline was commissioned by NICE and developed in accordance with the guideline development process outlined in 'The guidelines manual' (NICE 2009)¹⁷⁰.

4 2.1 Developing the clinical questions

Clinical questions were developed to guide the literature searching process and to facilitate the development of recommendations by the guideline development group (GDG). They were drafted by the review team and refined and validated by the GDG. The questions were based on the scope (Appendix A).

The full list of clinical questions addressed by the guideline is summarised in table 2.1 below

11 Table 1: full list of clinical questions

Question	Relevant chapter	Method used to formulate recommendations
Pre-sedation assessment, communication, patient information and consent		
For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques what factors should be assessed to justify the use of sedation rather than no sedation or general anaesthesia?	4	Consensus*
For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques what validated tools should be used to support assessment?	4	Consensus (as no relevant papers were identified for review)
For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques who should make the assessment and how should the assessment be recorded?	4	Consensus*

^{*} Questions denoted with * were agreed with NICE as consensus style questions *a priori*. These questions were based upon stakeholder desire to include these aspects even though routine care. The GDG felt that there would be limited evidence in these areas and as such they were background questions that were not congruent with the style of a full and systematic evidence based approach Sedation in children and young people: full guideline DRAFT (May 2010) Page 32 of 371

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques how should consent be obtained for	4	Consensus*
sedation?		
Fasting		
In children and young people under the age of 19 undergoing sedation techniques, should fasting versus no fasting be implemented to prevent adverse outcomes?	4	Evidence based (literature review)
Psychological preparation		
For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques what standard psychological preparation, coping skills and strategies should be used?	4	Evidence based (literature review)
Personnel and training		
For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation what generic and specific skills are required for different team members and for different levels of sedation?	4	Consensus*
For personnel involved in delivering sedation to children and young people under the age of 19 undergoing diagnostic and therapeutic procedures what training and competences are required?	4	Consensus*
Clinical environment and monitoring		
For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under moderate or deep sedation techniques what monitoring and equipment is required to reduce the risk of complications?	4	Consensus*
When should monitoring stop for children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques?	4	Consensus*
Discharge criteria		
For children and young people under the age of 19 after diagnostic and therapeutic procedures under moderate or deep sedation techniques what discharge criteria are required?	4	Consensus*
Efficacy and safety of midazolam		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is midazolam (with or without: analgesia, another drug or psychological techniques) effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?	6	Evidence based (literature review)
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is	6	Evidence based (literature review)

^{*} Questions denoted with * were agreed with NICE as consensus style questions a priori. These questions were based upon stakeholder desire to include these aspects even though routine care. The GDG felt that there would be limited evidence in these areas and as such they were background questions that were not congruent with the style of a full and systematic evidence based approach.

psychological techniques) safe for sedation (at mild, moderate, and deep levels) in different settings? Efficacy and safety of ketamine		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is ketamine (with or without: analgesia, another drug or psychological techniques) effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?	6	Evidence based (literature review)
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is ketamine (with or without: analgesia, another drug or psychological techniques) safe for sedation (at mild, moderate, and deep levels) in different settings?	6	Evidence based (literature review)
Efficacy and safety of chloral hydrate		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is chloral hydrate (with or without: analgesia, another drug or psychological techniques) effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?	6	Evidence based (literature review)
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is chloral hydrate (with or without: analgesia, another drug or psychological techniques) safe for sedation (at mild, moderate, and deep levels) in different settings?	6	Evidence based (literature review)
Efficacy and safety of nitrous oxide		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is nitrous oxide (with or without: analgesia, another drug or psychological techniques) effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?	6	Evidence based (literature review)
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is nitrous oxide (with or without: analgesia, another drug or psychological techniques) safe for sedation (at mild, moderate, and deep levels) in different settings?	6	Evidence based (literature review)
Efficacy and safety of opioids		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, are opioids (with or without: analgesia, another drug or psychological techniques) effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?	6	Evidence based (literature review)
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, are opioids (with or without: analgesia, another drug or psychological techniques) safe for sedation (at mild,	6	Evidence based (literature review)

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moderate, and deep levels) in different settings?		
Efficacy and safety of propofol		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is propofol (with or without: analgesia, another drug or psychological techniques) effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?	6	Evidence based (literature review)
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is propofol (with or without: analgesia, another drug or psychological techniques) safe for sedation (at mild, moderate, and deep levels) in different settings?	6	Evidence based (literature review)
Efficacy and safety of sevoflurane		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is sevoflurane (with or without: analgesia, another drug or psychological techniques) effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?	6	Evidence based (literature review)
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is sevoflurane (with or without: analgesia, another drug or psychological techniques) safe for sedation (at mild, moderate, and deep levels) in different settings?	6	Evidence based (literature review)
Efficacy and safety of triclofos sodium		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is triclofos sodium (with or without: analgesia, another drug or psychological techniques) effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?	6	Evidence based (literature review)
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is triclofos sodium (with or without: analgesia, another drug or psychological techniques) safe for sedation (at mild, moderate, and deep levels) in different settings?	6	Evidence based (literature review)
Sedation sparing		
For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, does a combination of psychological techniques and sedation drugs lead to sedation sparing?	6	Evidence based (literature review)

From these clinical questions, the technical team produced review questions and protocols to address these questions. The protocols are reported in appendix H.

3 2.2 Searching the literature

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4 2.2.1 Clinical literature search

The search strategies and the databases searched are presented in detail in Appendix C. All searches were conducted on the following databases with no date restrictions.

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Database	Interface	Date searched from
Medline	OVID	1950
Embase	OVID	1980
Cinahl	EBSCO	1982
The Cochrane Library (to 2009 Issue 4)	www.thecochranelibrary.com	All dates searched: 1996 for Cochrane Reviews 1995 for DARE 1898 for CENTRAL 1904 for Methods Studies 1995 for HTA and NHSEED

Databases were searched using relevant subject headings and free-text terms. Where appropriate, study design filters were applied. Non-English language studies and abstracts were not reviewed.

All searches were updated to 18th January 2010. Hand-searching was not undertaken following NICE advice that exhaustive searching on every guideline review topic is not practical or efficient¹⁷⁰. Reference lists of articles were checked for studies of potential relevance.

2.2.2 Sifting process

Once the search had been completed, the following sifting process took place:

- 1st sift: one reviewer sifted the title/abstract for articles that potentially met the eligibility criteria; this was checked where necessary by a second reviewer.
- 2nd sift: full papers were ordered that appeared relevant and eligible or where relevance/eligibility was not clear from the abstract.
- 3rd sift: full papers were appraised that meet eligibility criteria. Generally, one
 reviewer appraised the papers using an inclusion criteria form, and this was
 checked where necessary by a second reviewer.

Once individual papers were retrieved, the articles were checked for methodological rigour (see section 2.4), applicability to the UK and clinical significance. Assessment of study quality concentrated on dimensions of internal validity and external validity. At this stage, some studies were excluded if the interventions were not licensed for use in the UK or they were not regularly used in the UK. Studies in which the interventions were obsolete were also excluded.

2.2.3 Economic literature search

Economic evidence was obtained from systematic searches of the following databases in accordance with the NICE Guidelines Manual: Medline, Embase, the Health Technology Appraisals (HTA) database and the NHS Economic Evaluations Database (NHSEED. The latter two databases were searched via The Cochrane Library. Health economics searches were restricted by date on Medline and Embase to studies published since 2006.

Detailed search strategies can be found in Appendix C.

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2.3	Clinical	effectiveness	review	methods

This section describes the methods of reviewing that are common to all reviews of intervention studies. Further specific details are given in the individual reviews and in Appendix H. Details on consensus chapters are given in section 2.4.4

References identified by the systematic literature search were screened for appropriateness by title and abstract by an information scientist and systematic reviewer. Studies were selected that reported one or more of the outcomes listed in section 2.3.2. Selected studies were ordered and assessed in full by the NCGC team using agreed inclusion/exclusion criteria specific to the guideline topic, and using NICE methodology quality assessment checklists appropriate to the study design. Further references suggested by the GDG were assessed in the same way. Not enough data was available from RCTs for serious adverse events related to pharmacological interventions. Consequently, an additional literature review of observational data was performed to supplement the RCT evidence.

2.3.1 Patients covered by this guideline

Infants, children and young people (under 19 years) receiving sedation by any technique for painful or non-painful diagnostic or therapeutic procedures including dental surgery and minor operations carried out under local anaesthesia.

This guideline will not cover:

- Patients requiring sedation for purposes other than for diagnostic or therapeutic procedures including:
 - o sedation in critically ill patients requiring mechanical ventilation
- 23 o sedation in palliative care
- 24 o sedation in the treatment of mental health conditions
- o sedation given as premedication for general anaesthesia or as postoperative analgesia
- o night sedation.
- Patients having diagnostic or therapeutic procedures under general anaesthesia.

2.3.2 Outcome measures

The following outcomes were considered.

31 **Primary outcome:**

- Successful completion of diagnostic or therapeutic procedure
- o measured as the number of patients for whom the diagnostic or therapeutic procedure was carried out and completed.

35 **Secondary outcomes:**

1	•	Behavioural ratings including:
2 3 4		 pain as assessed by the patient or parent or other observer using validated pain scales e.g. Visual Analogue Scale (VAS), Children's Hospital of Eastern Ontario Pain Scale (CHEOPS), Faces Pain Scale (FPS).
5 6 7		 procedural distress and/or anxiety as assessed by the patient or parent or other observer using validated scales e.g. Visual Analogue Scale (VAS), Observation Scale of Behavioral Distress (OSBD).
8		o patient or parent satisfaction including preference
9	•	Sedation timing including
0		 length of induction: time from administration of sedation drug to initiation of procedure
2		 recovery: time from completion of procedure to recovery criteria being met or recovery to pre-sedation state
4		o duration of procedure
5 6		 total: time from administration of intervention to when patient has been transferred to the recovery area or has been discharged
7	Adverse ev	ents:
8	•	Aspiration
9	•	Respiratory intervention, including:
20		o oral-pharyngeal airway
21		o endotracheal intubation
22		o assisted ventilation
23	•	Cardiac arrest requiring either/or:
24		o external cardiac massage
25		o defibrillation
26	•	Oxygen desaturation <90%
27	•	Vomiting
28	2.4 Appr	aising the evidence
29	2.4.1 Appro	isal of methodological quality of 'treatment' studies
30		lure adopted
		-

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1	 For each clinical question, the highest level of evidence was sought.
2 3 4	 Randomised control trials (RCTs) were reviewed for drug efficacy and safety outcomes. Only RCTs of N≥20 were included. The largest available cohort studies were also included for drug safety reviews.
5 6	 Meta-analysis of RCT results was performed if the data was sufficiently homogeneous.
7 8 9 10 11	 Studies were appraised for methodological quality using the GRADE[#] scheme. When using GRADE, RCTs start as high quality and observational studies as low quality. Studies were downgraded or upgraded depending upon their risk of bias (see features below and section 2.4.3). Reasoning has been given when studies were downgraded.
12 13	The following features were assessed for the evidence found for each relevant outcome from a systematic review:
14	 study design (as a proxy for bias)
15 16	 limitations in the methodological quality of the study (mainly allocation concealment, blinding and loss to follow-up)
17	 consistency of an effect across studies
18 19	 directness (the degree to which the results directly address the question posed or, for example, are for a somewhat different population).
20	Other considerations:
21	• imprecision *
22	likelihood of reporting bias
23	• strength of association
24	• evidence of a dose–response relationship
25	expected effect of plausible confounders.

GRADE – Grading of Recommendations Assessment, Development and Evaluation

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 $^{^{\}star}$ Precision requires the GDG to decide what are clinically important harms and benefits for that outcome measure. For dichotomous outcomes we used a relative risk reduction of 25% (RR of 1.25 or 0.75) to indicate the clinically important threshold. For positive outcomes, the upper clinically important threshold used depended on the control group rate. When this rate was less than 80% a value of 1.25 was used. When the control group rate was more than 80%, the clinically important threshold was calculated assuming an intervention group rate of 100% and a control group rate based on the median rate where there was more than one study'

2.4.2 Data synthesis for treatment studies

Where possible, meta-analyses were conducted to combine the results of studies for each clinical question using Cochrane Review Manager software. Fixed-effects (Mantel-Haenszel) techniques were used to calculate risk ratios (relative risk) for the binary outcomes: number of adverse events, and the continuous outcome for endpoint or change from baseline IPSS score, QOL question from IPSS score and Qmax was analysed using an inverse variance method for pooling weighted mean differences. Statistical heterogeneity was assessed by considering the chi-squared test for significance at p<0.1 and an I-squared of $\geq 50\%$ to indicate significant heterogeneity.

Where significant heterogeneity was present we explored a number of possible predefined differences including the severity or main symptoms experienced by the participants recruited into the study, study design (open label or masked), and length of follow-up by doing subgroup analyses. Assessments of potential differences in effect between subgroups were based on the chi-squared tests for heterogeneity statistics between subgroups. If no sensitivity analysis was found to completely resolve statistical heterogeneity then a random effects (DerSimonian and Laird) model was employed to provide a more conservative estimate of the effect.

We looked for all outcomes of interest in each paper that was included in the evidence reviews. Where a primary or important decision-making outcome was not reported by a paper, these were not included in the evidence statements or GRADE profiles, in order to highlight an 'absence of evidence'. Where studies reported there were 'no events' for an outcome, this has been denoted in the review evidence statements or GRADE profiles as '0' patients, '0%' or 'no events'.

2.4.3 Grading evidence

The GRADE scheme (GRADE working group 2004) was used to assess the quality of the evidence for each outcome not each study, using the approach described below, and evidence summaries (evidence profiles) across all outcomes were produced.

According to GRADE quality assessments, the evidence is classified as follows:

- High: further research is very unlikely to change our confidence in the estimate of effect
- Moderate: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate
- Low: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate
- Very low: any estimate of effect is very uncertain.

36 The procedure adopted when using GRADE was:

- A quality rating was assigned, based on the study design.
- This rating was up- or down-graded according to specified criteria: study quality, consistency, directness, preciseness and reporting bias. Criteria were

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- 1 given a downgrade mark of -1 or -2 depending on the severity of the limitations.
 - The downgrade/upgrade marks were then summed and the quality rating revised. For example, a decrease of -2 points for an RCT would result in a rating of 'low'. Reasoning was explained for the downgrade marks.

The GRADE scheme was only used to assess the quality of evidence for RCTs. Full evidence profiles for efficacy and safety were produced and are contained on the relevant drug section.

The GDG recognised that research from non RCT observational studies is subject to the usual limitations of observational work, including dependence on the quality of medical record documentation and potential for bias secondary to non randomisation, and unblinded participants. In these studies, there were no interventions or comparisons but merely data collection of adverse events. The datasets were generally large, and were expected to provide more information on a range of adverse events than the small RCTs available for review. Due to these limitations, we only assigned quality rating ('very low' quality) based on the GRADE scheme. It was considered more comprehensive to present separately this supplementary observational data in the form of concise, customised summary tables which also contain the GRADE ratings.

2.5 Consensus

There are generally three main methods reported for developing consensus. These are Delphi, consensus development panels and nominal group processes³³. The nominal group technique (NGT) was originally developed by Delbecq et al⁵² as an organisational planning tool. The methodology varies from the Delphi process, which by design allows individuals to work in the presence of others, but verbal interaction is discouraged and facilitated through sequential questionnaires or summary processes, enabling consensus to be developed without the social pressures normally exerted through open dialogue²³⁶. Individual ideas are shared within the group, with facilitated discussion enabling the group to see how individuals are expressing their ideas. Normal practice is for the facilitator to then ask the group to prioritise, with aggregated rankings recorded. NGT uses this approach but with participant dialogue encouraged. This methodology works extremely well in clinical guideline development for those clinical questions identified and agreed as areas to be explored through consensus process, and towards the end of guideline development and in particular when working with the GDG in prioritising recommendations for targeted implementation.

The GDG in working together for a fourteen month period during development of the guideline is by nature a mature working group; individuals within the group are able to express their views relating to key issues in relation to clinical questions and key recommendations addressed through consensus methods within a social setting (the last GDG meeting). This is important for the group, who are able to use this experience and the content of discussion to then go into a formal agreement of consensus recommendations and formal voting as part of recommendation sign off. Developing consensus through validated instruments is important in ensuring the integrity of final recommendations that reflect the group as a whole, and benefit from the wealth of clinical and patient experience considered. The process itself enables all constituent members of the GDG to have equal weighting of opinion as their opinion moves towards a consensus group position. Typically, NGT works well for small groups, with 12 to 15

people widely acknowledged in the literature as the maximum number of people involved in this process

2.6 Cost-effectiveness review methods

Economic evaluations are useful in guideline development as they assess the costs and benefits of alternative courses of action which could be recommended within the guideline. Relevant published economic information may be used by the GDG to determine whether a particular recommendation would result in the efficient use of NHS resources, but in order to do so it must provide an estimate of both the costs to the NHS and the health benefits to patients. Relevant study designs are cost-effectiveness, cost-utility or cost-benefit analyses. Cost-minimisation analyses are only relevant when supported by evidence demonstrating that there is no difference in health outcome between the alternative health care interventions. Cost studies which focus solely on the cost of alternative health care interventions are not suitable for informing decisions on the efficient use of NHS resources as they do not take into account any differences in the benefits for patients. Studies reporting analyses in non-OECD member countries or prior to 1990 were also excluded as these were felt to be less relevant to current practice in the UK.

We have excluded analyses where the estimates of clinical effectiveness used to inform the economic evaluation are not based on evidence from randomised controlled trials (RCTs) or quasi-randomised controlled trials. This was done to minimise the potential for bias and to ensure consistency with the clinical effectiveness reviews.

The search strategy for existing literature is described in section 2.2.3 (Economic literature search). There were 226 papers identified by the search. After considering titles and abstracts, 24 papers were identified as potential cost or cost-effectiveness studies and all of these were ordered to cross check whether they reported both cost and health outcomes even in a disaggregated way.

Of the 24 full text papers considered, 7 were found to be not relevant to the review question as they were found either to report clinical outcomes only, or they compared interventions that were not relevant to the guideline, or they were in predominantly adult populations (minimum age of 16 and a mean age >45).

Of the 17 remaining studies, 12 were economic evaluations carried out within studies using non-RCT designs in which the estimates of clinical effectiveness were considered to be open to bias due to the trial design. These were excluded from the cost-effectiveness review. A list of the excluded studies and reasons for exclusion are listed in appendix F. Two (Martinez 2002^{159} , lannalfi 2005^{98}) of the remaining 5 studies were economic evaluations carried out within RCTs and three (Lee 2000^{136} , Jameson 2007^{100} , Pershad 2006^{179}) were model based evaluations. A description of the five studies is also given in appendix F. We carried out update searches up to 18^{th} January 2010 but did not identify further useful studies.

None of the identified five studies was of high quality, and they provided little relevant evidence on the cost-effectiveness of sedation techniques considered in the guideline. It was therefore necessary to construct an original economic evaluation model to determine the cost-effectiveness of sedation techniques.

2.7 Cost-effectiveness modelling

The details of the economic model are described in Appendix F.

Cost-effectiveness information helps the GDG to weigh the balance of the cost and health benefit of applying intervention strategies in the different population groups considered in the guideline. At the early stages of the sedation guideline development, the health economist worked with the GDG to identify two high priority areas for cost-effectiveness evidence. The first area of priority was on the cost-effectiveness evidence to enable the GDG determine which sedation technique is most appropriate. The second area was on the cost-effectiveness of using a combination of non-pharmacological techniques and sedation drugs as sedation sparing technique.

These were classified as high priority because appropriate sedation technique should have the potential to prevent the need to abandon and reschedule procedures when sedation is unsuccessful. This will reduce the use of the National Health Services (NHS) or Personal Social Services (PSS) resources. It should minimise distress, discomfort for and risk of harm to patients as well as reduce the potential for QALY loss due to long term morbidity or mortality. There was the need to gather health economic information on different sedation strategies. As we did not identify directly applicable reports, it became necessary to consider carrying out a de novo economic evaluation to determine the cost-effectiveness of different techniques.

We did not construct any cost-effectiveness model for using a combination of pharmacological techniques and sedation drugs as sedation sparing technique. The GDG did not consider it worthwhile to build this model as there was no evidence that a combination of non-pharmacological techniques and sedation drugs has a sedation sparing effect. The health economic work for this guideline was therefore focussed on the first area of priority, the most appropriate sedation technique.

Cost-effectiveness was determined by comparing the cost per patient for the different strategies. The technique with the lowest cost per patient is considered to be the optimal strategy from a cost-effectiveness perspective.

The model was constructed using the best available evidence. Clinical and safety evidence was taken from a systematic review (chapter 6 on clinical effectiveness and safety review) and costing was based on the perspectives of the NHS and PSS. When the evidence was weak or absent the GDG expert opinion was used to determine the input parameters of the model. The assumptions made in the model and the uncertainties in the input parameters are described explicitly. These were considered by the GDG when interpreting the model results. The impact of uncertainties in the model structure and input parameters were explored through deterministic sensitivity analyses. We did not do a probabilistic sensitivity analysis as the estimate for a number of key input parameters were ascertained by expert opinion. The limitations of the model are discussed.

We have not prioritised all the clinical questions for economic evaluation. For those which were not prioritised, the GDG considered the likely cost-effectiveness of available options by making a qualitative judgement on the likely costs, health benefits and potential harms.

2.8 Developing recommendations

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- Over the course of the guideline development process, the GDG was presented with the following:
 - The clinical and economic evidence reviews. All evidence tables are in Appendices D, E and G.
- Forest plots of results from studies, including meta-analyses where appropriate.
- A description of the methods and results of the cost-effectiveness analysis
- 8 Recommendations were drafted on the basis of this evidence whenever it was available.
- When clinical and economic evidence was poor or absent, the GDG proposed recommendations based on their expert opinion.
- 11 The GDG also developed a care pathway algorithm according to the recommendations.

12 2.9 Research recommendations

- When areas were identified for which good evidence was lacking, the GDG considered making recommendations for future research. Decisions about inclusion were based on factors such as:
- the importance to patients or the population
- 17 national priorities
- potential impact on the NHS and future NICE guidance
- ethical and technical feasibility
- The GDG identified four high priority research recommendations, after discussion and voting (appendix G).

22 2.10 Validation of the guideline

The first draft of this guideline was posted on the NICE website for an 8-week consultation period between 7 May – 2 July 2010 and registered stakeholders were invited to comment. The GDG responded to comments and an amended version of the guideline was produced.

2.11 Disclaimer and funding

Healthcare providers need to use clinical judgement, knowledge and expertise when deciding whether it is appropriate to apply guidelines. The recommendations cited here are a guide and may not be appropriate for use in all situations. The decision to adopt any of the recommendations cited here must be made by the practitioner in light of individual patient circumstances, the wishes of the patient, clinical expertise and resources.

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1 2 3	The National Clinical Guideline Centre disclaim any responsibility for damages arising out of the use or non-use of these guidelines and the literature used in support of these guidelines.
4 5 6	The Collaborating Centre for Nursing and Supportive Care (now a part of the National Clinical Guideline Centre) were commissioned by the National Institute for Health and Clinical Excellence to undertake the work on this guideline.
7	2.12 Updating the guideline

2.12 Updating the guideline

This guideline will be updated in concordance with NICE guidelines manual (NICE 2009) 170 . 8 9

3 Summary of Recommendations

2 3 4		Below are the recommendations that the GDG selected as the key priorities for implementation followed by the complete list of recommendations and research recommendations.
5	3.1	Key priorities for implementations
6 7		The GDG identified ten key priorities for implementation. The decision was made after discussion and voting by the GDG. They selected recommendations that would:
8		 Have a high impact on outcomes that are important to patients (A)
9		• Have a high impact on reducing variation in care and outcomes (B)
10		• Lead to a more efficient use of NHS resources (C)
11		• Promote patient choice (D)
12		• Promote equalities.(E)
13 14		In doing this the GDG also considered which recommendations were particularly likely to benefit from implementation support. They considered whether a recommendation:
15		 Relates to an intervention that is not part of routine care (U)
16		• Requires changes in service delivery (V)
17		• Requires retraining staff or the development of new skills and competencies (W)
18		 Highlights the need for practice to change (X)
19 20		 Affects and needs to be implemented across various agencies or settings (complex interactions) (Y)
21 22		 May be viewed as potentially contentious, or difficult to implement for other reasons (Z)

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support points are indicated by the use of the letters shown in brackets above.

For each key recommendation listed below, the selection criteria and implementation

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1	
2	
3 4	Ensure that trained healthcare professionals carry out pre-sedation assessments and document the results in the healthcare record.
5	(Selection criteria: A, B, C, D, E. Implementation support: W, X, Y)
6	
7	Establish suitability for sedation by assessing:
8	o current medical condition and any surgical problems
9	 weight (growth assessment)
10 11	 past medical problems (including any associated with previous sedation or anaesthesia)
12	o current and previous medication (including any known allergies)
13	o physical status (including the airway)
14	o psychological and developmental status.
15	(Selection criteria: A, B, C, D, E. Implementation support: W, X, Y)
16	
17	Seek further specialist advice before delivering sedation if either:
18	o there is concern about a potential airway or breathing problem, or
19	o the child or young person is assessed as ASA* grade 3 or greater.
20	(Selection criteria: A. Implementation support: W)
21	
22	Choose the most suitable sedation technique based on:
23	 what the procedure involves
24	o target level of sedation
25	o contraindications
26	o side effects

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^{*} ASA=American Society of Anesthesiologists

1	o patient preference.
2	(Selection criteria: A, B, C, D, E. Implementation support: W, X, Y)
3	
4	Ensure that both the following are available when considering sedation:
5 6	 a healthcare professional and assistant trained* in delivering and monitoring sedation in children and young people and
7	o immediate access to resuscitation and monitoring equipment [#] .
8	(Selection criteria: A, B. Implementation support: V, W, X, Y)
9	
10 11	Healthcare professionals delivering sedation should have knowledge and understanding of:
12	 Sedation drug pharmacology and applied physiology
13	 Assessment of the child or young person
14	o Monitoring
15	o Recovery care
16 17	 Complications and their immediate management, including paediatric life support.
18	(Selection criteria: A, B. Implementation support: U, W, X, Y, Z)
19	
20	Healthcare professionals delivering sedation should have practical experience of
21 22	 effective delivery of the sedation technique used and management of complications
23 24	 observing clinical signs (for example airway patency, breathing rate and depth, pulse, pallor and cyanosis, depth of sedation)
25	o using monitoring equipment.
26	(Selection criteria: A, B. Implementation support: U, W, X, Y)
27	

^{*} See recommendations on monitoring Sedation in children and young people: full guideline DRAFT (May 2010) Page 48 of 371

1 Sensure that all members of the sedation team have the following competencies:

Minimal	Moderate	Deep
sedation*	sedation	sedation
All members have	All members have	All members have
basic life support	basic life support	basic life support
skills	skills	skills
	At least one team member should have intermediate life support skills in airway management using mask ventilation and use of defibrillator	At least one team member should have advanced life support skills including advanced airway skills and cardiac arrest management

*and sedation with nitrous oxide alone (up to 50% in oxygen) (Selection criteria: A, B. Implementation support: U, W, X, Y, Z)

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- * Healthcare professionals delivering sedation should have documented evidence (for example, a certificate or a comprehensive record) of competency including:
 - satisfactory completion of a theoretical training course covering the principles of sedation practice;
 - o practical experience of sedation techniques, including details of:
 - children and young people managed under supervision
 - successful completion of work-based assessments.
- 12 (Selection criteria: A, B. Implementation support: U, W, X, Y, Z)

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- For deep sedation continuously monitor, interpret and respond to all of the following:
- o Respiration
- 17 Oxygen saturation
- 18 o Heart rate
- 0 End tidal CO₂ (capnography)
- 20 o three-lead electrocardiogram (ECG)
- O Blood pressure (monitor every 5 minutes)
- 22 O Depth of sedation
- o Pain

1	o Coping
2	o Distress.
3 4	(Selection criteria: A, B. Implementation support: U, W, X, Y, Z)
5	3.2 Complete list of recommendations
6	3.2.1 Recommendations on pre-sedation assessment, communication, patient information
7	and consent
8 9	Ensure that trained healthcare professionals carry out pre-sedation assessments and document the results in the healthcare record.
0	
1	Establish suitability for sedation by assessing:
2	o current medical condition and any surgical problems
3	 weight (growth assessment)
4 5	 past medical problems (including any associated with previous sedation or anaesthesia)
6	o current and previous medication (including any known allergies)
7	o physical status (including the airway)
8	o psychological and developmental status.
9	
20	> Seek further specialist advice before delivering sedation if either:
21	o there is concern about a potential airway or breathing problem, or
22	o the child or young person is assessed as ASA* grade 3 or greater.
23	
24	Choose the most suitable sedation technique based on:
25	 what the procedure involves
26	o target level of sedation

^{*} ASA=American Society of Anesthesiologists Sedation in children and young people: full guideline DRAFT (May 2010) Page 50 of 371

1		o contraindications
2		o side effects
3		o patient preference.
4		
5	>	Ensure that both the following are available when considering sedation:
6 7		 a healthcare professional and assistant trained[*] in delivering and monitoring sedation in children and young people and
8		o immediate access to resuscitation and monitoring equipment #
9		
10 11	>	To enable the child or young person and their parents or carers to make an informed decision, offer them verbal and written information on:
12		o the proposed sedation technique
13		o the alternatives
14		o the associated risks and benefits.
15 16	>	Obtain and document informed consent for sedation (see recommendation above).
17		
18	3.2.2 Recom	mendations on fasting
19 20	>	Before starting sedation, confirm and record the time of last food and fluid intake in the healthcare record.
21		
22	>	For elective procedures, apply the 2-4-6 rule ¹ .
23		

* See recommendations on training

- 2 hours for clear fluids
- 4 hours for breast milk
- 6 hours for solids

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[#] See recommendations on monitoring

1 Fasting times should be as for general anaesthesia:

1 2 3	>	For urgent procedures in a child or young person who has not fasted, base the decision to proceed with sedation on clinical emergency and the target depth of sedation.
4 5	>	Fasting is not required for minimal sedation and for sedation with nitrous oxide alone (up to 50% in oxygen).
6		
7	3.2.3 Recon	nmendations on psychological preparation
8 9	>	Ensure that the child or young person is prepared psychologically for sedation by offering advice about:
10		o the procedure itself
11 12		 what the child or young person should do and what the healthcare professional will do
13 14		 the sensations associated with the procedure (for example, a sharp scratch, numbness)
15		o how to cope with the procedure.
16		
17 18	>	Ensure that the information is appropriate for the developmental stage of the child or young person.
19		
20 21 22	>	Offer parents and carers the opportunity to be present during sedation when appropriate. If a parent or carer decides to be present, offer them advice about their role during the procedure.
23		
24 25	>	For an elective procedure, consider referral to a mental health specialist for children who are severely anxious or who have a learning disability.
26		
27	3.2.4 Recon	nmendations on personnel and training
28 29	>	Healthcare professionals delivering sedation should have knowledge and understanding of:
30		 Sedation drug pharmacology and applied physiology
31		Assessment of the child or young person
32		 Monitoring
33		Recovery care

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1 2	 Complications and their immediate management, including paediatric life support 					
3						
4	≻ н	ealthcare professionals	delivering sedation	on should have pro	actical experience of:	
5 6		 effective delivery of the sedation technique used and management of complications 				
7 8		 observing clinical signs (for example airway patency, breathing rate and depth, pulse, pallor and cyanosis, depth of sedation) 				
9		 using monitoring e 	equipment.			
10						
11	≻ Er	nsure that all members o	of the sedation tec	ım have the follow	ing competencies:	
		Minimal sedation*	Moderate sedation	Deep sedation		
		All members have basic life support skills	All members have basic life support skills	All members have basic life support skills		
			At least one team member should have intermediate life support skills in airway management using mask ventilation and use of defibrillator	At least one team member should have advanced life support skills including advanced airway skills and cardiac arrest management		
12 13		*and sedation with ni	trous oxide alone (up to	50% in oxygen)	I	
14						
15 16		Ensure that a healthcare professional trained in delivering anaesthetic agents is available to administer the following sedatives:				
17	 Sevoflurane 					
18	o Propofol					
19		O Opioids combined	d with ketamine.			
20						
21 22		ealthcare professionals or example, a certificat	-			
23 24	 satisfactory completion of a theoretical training course covering the principles of sedation practice 					
25	o practical experience of sedation techniques, including details of:					
26	 children and young people managed under supervision 					

1			successful completion of work-based assessments.
2			
3 4 5	>	they u	ealthcare professional and their team delivering sedation should ensure pdate their knowledge and skills through programmes designed for ing professional development.
6			
7	3.2.5 Recor	nmenda	tions on clinical environment and monitoring
8 9	>		oderate sedation continuously monitor, interpret and respond to changes in the following:
10		0	Coping
11		0	Depth of sedation
12		0	Pain
13		0	Distress
14		0	Respiration
15		0	Oxygen saturation
16		0	Heart rate.
17			
18 19	>	For de	ep sedation continuously monitor, interpret and respond to all of the ing:
20		0	Respiration
21		0	Oxygen saturation
22		0	Heart rate
23		0	End tidal CO ₂ (capnography)
24		0	three-lead electrocardiogram (ECG)
25		0	Blood pressure (monitor every 5 minutes)
26		0	Depth of sedation
27		0	Pain
28		0	Coping
29		0	Distress.
30			

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1 2	>	Ensure that data from continuous monitoring during sedation are clearly documented in the healthcare record.					
3							
4	>	After the procedure, continue monitoring until:					
5		o the airway is patent					
6		o protective airway and breathing reflexes are present					
7		o the child or young person is haemodynamically stable					
8		o the child or young person has returned to baseline level of consciousness.					
9							
10	3.2.6 Recon	nmendations on discharge criteria					
11 12	>	Ensure that all of the following criteria are met before the child or young persor is discharged:					
13		o vital signs* have returned to normal levels					
14 15	 the child or young person has returned to baseline level of responsivenes and orientation 						
16		o nausea, vomiting and pain have been adequately managed					
17		o there is no risk of further reduced level of consciousness.					
18							
19 20	>	Consider referring to an anaesthesia specialist if the child or young person is no able to tolerate the procedure under sedation.					
21							
22	3.2.7 Recon	nmendation on painful procedures					
23 24	>	For painful procedures (for example suture laceration or manipulation of fracture) consider using:					
		For minimal sedation For moderate sedation					
		Oral or intranasal Nitrous oxide Intravenous ketamine (or intramuscular if					

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^{*} Vital signs are measures of various physiological statistics and usually include body temperature, heart rate, blood pressure and respiratory rate.

	in oxygen)	intravenous is difficult)
Nitrous oxide alone (up to 50% in oxygen)	Intravenous midazolam with or without fentanyl	Propofol with or without fentanyl

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3.2.8 Recommendations on painless imaging

Do not routinely use ketamine or opioids for sedating children or young people for painless imaging procedures.

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- For children and young people who are unable to tolerate a painless procedure (for example during diagnostic imaging) consider one of the following:
- 8 o chloral hydrate (oral) for children under 15Kg or
 - propofol or
- 10 o sevoflurane.

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3.2.9 Recommendations on endoscopy

Consider intravenous midazolam to achieve minimal or moderate sedation for upper gastrointestinal endoscopy.

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Consider using fentanyl (or equivalent opioid) and intravenous midazolam to achieve moderate sedation for lower gastrointestinal endoscopy.

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3.2.10 Recommendation on dental procedures

- For a child of young person who cannot tolerate a painful dental procedure with local anaesthesia alone, consider one of the following techniques to achieve moderate sedation:
 - nitrous oxide and oxygen (titrated according to needs and using a maximum of 70% nitrous oxide) or
 - o midazolam.

If these sedation techniques are not suitable or effective, consider referral to a specialist team for other sedation techniques (for example midazolam in combination with nitrous oxide and/or sevofluorane).

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3.3 Algorithms

3 See separate file

3.4 Research recommendations

3.4.1 Research recommendation on assessment

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation, what factors should be assessed to both establish the need for sedation and to reduce the risk of adverse events?

Why it is important

Some children need sedation, some need anaesthesia, and some need behavioural management alone. There is wide variation in how this choice is made and a recommended standard method of assessment may reduce variation and improve both success and safety of sedation when it is chosen. Furthermore an assessment tool may prevent inappropriate choices and improve the overall management of children having procedures. The GDG suggest an observational study to determine the important factors, followed by a consensus study to develop a tool. The assessment of the tool should be tested by a randomised comparison of children and young people who have been assessed routinely with those who have been assessed using the tool. The assessment tool aims to improve sedation success and quality, and reduce any complications.

3.4.2 Research recommendation on fasting

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation how long should they be fasted to prevent adverse events?

Why it is important

Inhalation of gastric contents can be fatal. Loss of consciousness is associated with the loss of vital airway reflexes and inhalation of gastric contents is possible. Consequently fasting (in order to keep the stomach empty) is standard practice before general anaesthesia and has become standard before any sedation technique that may cause loss of consciousness. Prolonged fasting however is distressing and can cause dehydration and hypoglycaemia. It would be helpful to know the minimum length of time necessary to fast a child before sedation in order to ensure that he stomach is empty, and to know that likelihood of regurgitation or vomiting is very small.

3.4.3 Research recommendation on psychological preparation

For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures under sedation what psychological techniques can lead to sedation sparing, improve patient/family satisfaction, and ensure safe completion of the procedure?

3.4.4 Research recommendation on personnel and training

For personnel involved in delivering sedation to children and young people under the age of 19 undergoing diagnostic and therapeutic procedures what training is required to both achieve and maintain essential skills?

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Why it is important

Potent drugs can cause unintended airway obstruction. Anaesthetists are skilled at managing airway obstruction because they practise this regularly. However, anaesthetists are a scarce resource so non-anaesthetists need to learn how to manage airway obstruction. The skills that are needed have been identified but can these skills be attained and maintained by professionals who need them occasionally? The GDG suggests that a standard teaching method and assessment tool are developed. This would involve an observational study of a cohort of trainees, who can be assessed, trained and then reassessed at varying intervals to determine whether the training is successful and how often it is necessary.

3.4.5 Research recommendations clinical environment and monitoring

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation is the introduction of system of capnography monitoring cost-effective, compared to standard monitoring, in reducing adverse events?

Why it is important

Monitoring of airway patency and breathing during sedation is crucial. During anaesthesia this is achieved with capnography. During sedation this can be achieved by using soft nasal catheters. Capnographs are expensive and their ability to recognise airway obstruction and respiratory depression in sedated children is uncertain.

Which depth of anaesthesia monitors can be used to monitor depth of sedation in children and which is best?

Why it is important

Several depth of anaesthesia monitors are in use around the world. Most use processed EEG signals while some use stimulation of the brainstem by auditory stimuli. It is not yet clear whether the available monitors can follow children through different levels of sedation accurately and this study would set out to determine which monitor best tracks the transition from moderate to deep sedation in children of different ages.

3.4.6 Research recommendations on drugs for sedation

For children and young people under the age of 19 undergoing minor painful procedures, what potent analgesic drugs can be combined with midazolam to provide safe moderate sedation?

Why it is important

Midazolam has a strong safety profile in inducing either minimal or moderate sedation. For painful procedures midazolam should be combined with analgesia. Ideally analgesia

is achieved by local anaesthesia. Sometimes local analgesia is insufficient and potent opioid analgesia is necessary. The combination of potent opioid and midazolam can cause deep sedation and airway obstruction. These effects can be managed safely but will involve extra resources. If would be safer if a technique could be developed that was both reliable and had a wide margin of safety. Prospective and retrospective audit data are available to help guide the choice of opioid and the doses. A randomised controlled trial is needed to test the efficacy and safety of these combinations.

For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures under sedation with ketamine, how can the vomiting be reduced?

Why it is important

14 15 16

Ketamine is demonstrated to have a strong efficacy and safety profile in enabling safe sedation and as an analgesic drug useful for painful procedures in children and young people. Its main side effect is vomiting in approximately 10% of patients. No data is available on whether antiemetic drugs prevent vomiting. The GDG suggested an RCT study comparing ketamine + placebo versus ketamine with antiemetic

For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, are the procedures carried out under sedation delivered more cost effective compared to general anaesthesia?

Why it is important

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Anaesthesia or an "Anaesthetist led service" has the advantage over sedation because it usually has faster onset and offset and is more predictable. It may be more expensive and is a scarce resource. Data comparing the efficiency of sedation in comparison with anaesthesia for certain procedures are not available. Models of care need to be developed and studied to whether anaesthesia or sedation gives the best value for money. With such data, efficient services can be planned.

30 31

For children and young people under the age of 19 undergoing endoscopy, is propofol (with or without: analgesia, another drug or psychological techniques) effective, safe and cost effective for sedation (at minimal and moderate levels) in comparison with midazolam (with or without opioids) or with general anaesthesia?

Why it is important

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Propofol is a short acting anaesthetic agent that can be used to achieve any target sedation level. The dose necessary for gastrointestinal endoscopy however usually has a tendency to cause anaesthesia albeit for a short period of time. It would be helpful to know the dose limitation that is unlikely to cause deep sedation because this dose may be effective and safe enough. Moderate sedation with propofol could be compared with another sedation technique such as midazolam with or without opioid. It caoul also be compared with a general anaesthetic dose of propofol.

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For children and young people under the age of 19 undergoing painful procedures, is ketamine effective and safe for sedation in comparison with propofol?

Why it is important

Both ketamine and propofol are safe and effective drugs suitable for painful procedures. Propofol however has a tendency to cause deep sedation and anaesthesia in which the airway and breathing may need an intervention or support. Ketamine has few appreciable effects on the airway and breathing but has a longer recovery time than propofol and causes vomiting.

What are the safety and efficacy profiles of sedation techniques in current practice?

Why it is important

Data on the safety of sedition in the UK are not available. A large prospective database of sedation cases, including data on drugs, procedures, the level of sedation and any complications, would be beneficial in not only providing definitive data on the safety of sedation but also actively promoting safe practice. The GDG suggests that a national registry for paediatric sedation is established for the purpose of creating a database with sufficient data.

Is patient-controlled sedation with propofol feasible in adolescents and children?

Why it is important

Propofol in low dose is an excellent anxiolytic. Patient-controlled sedation has been validated in adults undergoing dental procedures and endoscopy for safety and efficacy. Giving the patient control of their sedation has important psychological benefits. The study would involve developing new pump technology, paediatric software and a child friendly patient-activation system. There would have to be an open pilot evaluation to establish safety and efficacy followed by a randomised-controlled trial versus IV midazolam.

4 Key considerations in supporting the patient's

2 journey

The patient journey is the experience of the patient and their family or carers before, during and after sedation for a procedure. It includes key stages of management by healthcare professionals including patient assessment and preparation. Each stage of the journey has been considered by the GDG for the purpose of maximising the success and safety of sedation. It is the healthcare practitioners themselves who will ensure that sedation is managed well and therefore their training has been discussed at length.

4.1 Pre-sedation assessment, communication, patient information and

10 consent

11 4.1.1 Clinical introduction

Assessment of the patient is crucial to determine their needs for the procedure. Some patients will cooperate or tolerate procedures without alteration of their conscious level. Others will need sedation and the target level will vary according to the patient that the procedure. For example the target sedation level for dental procedures is conscious sedation whereas a small child having an MRI scan needs to be unconscious either by deep sedation or anaesthesia. Many patients will have medical problems that could give rise to difficulties with sedation and anaesthesia. These will need careful assessment so that the risks of any chosen sedation technique can be appreciated. Communication of all these factors to the patient and their family is important to the consenting process. The presentation of clear and relevant information is likely to help patients and their families make reasoned choices.

4.1.2 Clinical methodological introduction

CLINICAL QUESTIONS:

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques:

- 1. What factors should be assessed to justify the use of sedation rather than no sedation or general anaesthesia?
- 2. What validated tools should be used to support assessment?

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1	3. Who should make the assessment and how should the assessment be recorded?
2	4. How should consent be obtained for sedation?
3	Clinical questions 1, 3 and 4
4 5	GDG sought to provide guidance to these questions based on their expert experience and opinion.
6	Clinical question 2
7	The literature was searched but no relevant papers were identified for review.
8	4.1.3 GDG discussion
9	Factors to consider in assessment
10 11 12 13	The GDG agreed that clear guidance should be given about the components of the assessment of a child or young person prior to sedation. These components feature in the recommendation and although others maybe important, the specified components were considered to be essential and have been arranged in order of priority.
14 15 16 17 18 19 20 21	The assessment should begin by understanding the child or young person's medical (or surgical) problem that has led them to require the procedure. Other non-related problems or illnesses, such as diabetes mellitus or an upper airway viral infection, should be identified and assessed. Measurement of the body weight is a simple method of identifying children who are not following normal growth development (or those who are obese). Growth failure may suggest that the disease is severe. Obesity is associated with other medical problems and can impair effective breathing during deep sedation. The doses of all drugs, except vapours and gases, must be calculated or adjusted according to the body weight.
23 24 25 26 27 28 29	Details of previous sedation or anaesthesia, or any medication, may identify problems that can be avoided. An assessment of the airway, breathing and circulation may find dangerous risk factors and problems that require additional equipment and technical expertise. Pulse oximetry is a reliable estimate of oxygen saturation of arterial blood and heart rate. The GDG considered that this tool should be available in the presedation assessment because it is easy to use and will identify some important respiratory and cardiovascular problems.
30 31 32	Some problems are well known to increase the risk of sedation so the benefit of the intended procedure needs to be considered. Physical examination requires training and experience.
33 34	Access to the patient's healthcare record is essential for information about previous problems with sedation or anaesthesia.
35 36 37 38 39 40	Children and young people who are unable to understand or cooperate with the sedation may be identified by assessment of their psychological and developmental status. Pre-sedation assessment should establish what the patient is able to understand and appreciate. This aids communication and gains assent. It should be determined if restraint or clinical holding have been used previously and how this was managed. Guidance on the appropriate use of restraint in children has been published by the Royal College of Nursing ⁷ .

The GDG discussed assessment of sedation in the emergency situation. It was agreed that in an emergency the medical needs should take priority until the patient has been stabilized. Once the child or young person has been stabilized, they can be assessed for sedation.

The GDG considered that it was important to make sure that there were safe facilities available to deliver the chosen sedation technique, and this led to discussion about who should be present and what equipment was necessary. The number of required healthcare professionals was discussed and the type of equipment. The GDG emphasized that these resources are essential and need to be present during sedation. Having them nearby may not prevent a problem soon enough, so they need to be next to the patient. If there is a respiratory complication, the healthcare professional will need to react promptly. If monitoring is used effectively, most problems will be prevented and others will be identified as soon as possible. Resuscitation equipment needs to be ready at hand. This includes airway and breathing devices that may needed to be inserted promptly to avoid or treat hypoxia and cardiac arrest.

The GDG discussed how many healthcare professional were needed according to the type of sedation and the intended procedure. It was noted that for some procedures the professional performing the procedure could control or assist in the sedation. In other situations two professionals were needed to concentrate on the patient during sedation and could not therefore be involved in the procedure. Overall, two professionals have to be available to look after a sedated patient; one of these may be involved with procedure provided they can stop the procedure and help with any complications of sedation.

Use of validated tools in assessment

As no evidence was found to support the use of validated tools in the assessment of children prior to sedation, the recommendations are based on the specialist experience and opinion of the GDG.

There are no validated tools for assessment of children and young people for sedation. There is however a widely used American Society of Anesthiologists (ASA)^{1,6,44} scoring tool to grade risk in patients having anaesthesia. The GDG considered that this was widely understood, simple to use and therefore should be used in describing the physical status of children and young people who need sedation. The sedation management of a patient who is assessed at ASA grade 3 or 4 should be managed after discussion with a specialist in sedation or anaesthesia.

Who should make the assessment?

Whichever professional group is involved with sedation, assessment of children and young people should be sufficient to identify important factors that affect the management of sedation. The importance of assessment is emphasised and should be carried out by a trained healthcare professional experienced in supporting children and young people undergoing sedation.

The assessment (and other details of sedation management) should be recorded in the healthcare record so that important details are available for any subsequent sedation or anaesthesia. Clear healthcare records may prevent mistakes and reduce risks.

Information and consent

The GDG agreed that each child or young person should be assessed concerning their capacity to make decisions, taking into account their previous experiences, level of maturity and cognitive development. Children and young people who have capacity to consent should be encouraged to do so.

Valid consent should be voluntary, fully informed and the person giving consent should have capacity. Besides their parents or guardians, children and young people might like to know about their illnesses, investigations and treatment and what is likely to happen to them. They should be involved in decisions about their care, even if they are not able to make decisions on their own and should be given the opportunity to ask questions. It is important that patients are given choice about which sedation technique, if any, should be used. The choice will depend upon the risks, the side effects and the patient's ability to cope with discomfort or anxiety. In essence, the choice is between sedation techniques, no sedation or angesthesia.

Healthcare professionals have a duty to explain fully to the child or young person about the proposed sedation technique and any alternatives. The explanation should be given in a way that the patient can understand and it should be supported by illustrations, or in other formats, and in the language of the patient and family. High quality patient information provision is the cornerstone of good clinical care and is essential for consent to be valid. Department of Health guidance on obtaining consent and what to expect if you are a young person/parent/carer²⁻⁵ is available online at:

http://www.dh.gov.uk/en/Publichealth/Scientificdevelopmentgeneticsandbioethics/Consent/Consentgeneralinformation/index.htm

Children and young people should be provided with timely, accessible information that is easy to understand and appropriate to their level of understanding and maturity.

Details of consent and relevant discussions should be available in the healthcare record to help any future patient management.

4.1.4 Health economic considerations

An economic analysis was not carried out. The need for assessment is the same for all the sedation techniques considered and it is expected to have a low impact on the NHS resources.

4.1.5 Recommendations

Recommendation 1	Ensure that trained healthcare professionals carry out pre-
	sedation assessments and document the results in the
	healthcare record.

Recommendation 2	Establish suitability for sedation by assessing:		
	- Current medical condition and any surgical problems		
	- Weight (growth assessment)		
	- Past medical problems (including any associated with previous sedation or anaesthesia)		
	- Current and previous medication (including any known allergies)		
	- Physical status (including the airway)		
	- Psychological and developmental status.		
Recommendation 3	Seek further specialist advice before delivering sedation if either:		
	- there is concern about a potential airway or breathing problem, or		
	- the child or young person is assessed as ASA^* grade 3 or greater.		
Recommendation 4	Choose the most suitable sedation technique based on:		
Recommendation 1	- What the procedure involves		
	- Target level of sedation		
	- Contraindications		
	- Side effects		
	- Side effects		
	- Patient preference.		

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^{*} ASA=American Society of Anesthesiologists Sedation in children and young people: full guideline DRAFT (May 2010) Page 66 of 371

Recommendation 5	Ensure that both the following are available when considering sedation:
	- A healthcare professional and an assistant trained in delivering and monitoring sedation in children and young people and
	- Immediate access to resuscitation and monitoring equipment [#] .
Recommendation 6	To enable the child or young person and their parents or carers to make an informed decision, offer them verbal and written information on:
	- the proposed sedation technique
	- the alternatives
	- the associated risks and benefits.
Recommendation 7	Obtain and document informed consent for sedation (See recommendation 6).
416 Posograh	rocommondation
For children therapeutic	recommendation and young people under the age of 19 undergoing diagnostic and procedures under sedation, what factors should be assessed to both e need for sedation and to reduce the risk of adverse event?
For children therapeutic	and young people under the age of 19 undergoing diagnostic and procedures under sedation, what factors should be assessed to both e need for sedation and to reduce the risk of adverse event?

* See recommendations 12 to 17 on personnel and training.

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management alone. There is wide variation in how this choice is made and a

recommended standard method of assessment may reduce variation and improve both

success and safety of sedation when it is chosen. Furthermore, an assessment tool may

children. The GDG suggest an observational study to determine the important factors,

prevent unsuitable choices and improve the overall management of procedures in

[#] See recommendations 18 to 21 on clinical environment and monitoring.

followed by a consensus study to develop a tool. The assessment tool should be tested by a randomised comparison of children and young people who have been assessed routinely with those who have been assessed using the tool. The assessment tool aims to improve sedation success and quality, and reduce any complications.

4.2 Fasting

4.2.1 Clinical Introduction

The importance of safety in any clinical procedure is paramount, and in relation to sedation the question 'should a child or young person be fasted before the procedure?' is important. Currently, local policy in relation to the administration of general anaesthesia is shaped by the joint Royal College of Nursing (RCN) /Royal College of Anaesthetists (RCA) Clinical Guideline 'Perioperative Fasting in Adults and Children' (2005) ⁸. However, there is acknowledged variation in practice to routine fasting (or not) when applied to the management of children and young people receiving sedation. This guideline is timely in providing standard recommendations for practice.

4.2.2 Clinical methodological introduction

CLINICAL QUESTION:

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques:

5. Should fasting versus no fasting be implemented to prevent adverse outcomes?

The review for this question consisted of three evaluation processes:

- 1) The joint RCN/RCA guideline 'Perioperative fasting in adults and children' (2005)⁸ was assessed using the AGREE instrument for appraisal of clinical guidelines.
- 2) An update search was conducted for perioperative fasting in children from 2004 to 2009, using key words 'anaesthesia,' 'fasting,' and 'children.' The purpose of this search to was identify recent publications which might impact recommendations in the joint RCN/RCA guideline 'Perioperative fasting in adults and children' (2005)8.
- 3) A full search of the literature relevant to fasting for paediatric sedation was conducted, using key words 'sedation,' 'fasting,' and 'children'.
- One RCT met inclusion criteria. Six observational studies were also included in this review, due to lack of further RCT data.
- **Population:** Infants, children and young people under the age of 19 who are receiving sedation by any technique for diagnostic or therapeutic procedures including dental surgery and minor operations carried out under local anaesthesia.
- **Intervention:** Fasting before sedation with one of the following drugs: midazolam, ketamine, propofol, chloral hydrate, nitrous oxide, sevoflurane, fentanyl, morphine intravenous or intramuscular, or diamorphine.

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1	Comparison: Fasting versus no fasting.
2	Outcomes for adverse events as evidenced by:
3	 Aspiration
4	 Vomiting
5	Oxygen saturation <90%
6	Respiratory intervention, including:
7	o oral-pharyngeal airway
8	o intubation
9	o assisted ventilation
10	AGREE appraisal
11 12 13	The AGREE instrument was used to appraise the joint RCN/RCA clinical guideline 'Perioperative fasting in adults and children' (2005) ⁸ . The full instrument with reviewer's comments is available in Appendix I. The overall assessment was as follows:
14	This guideline is recommended with following provisos:
15 16	 Update searches for the period from 2005 – 2009 are carried out, as the guideline is scheduled for review in 2009.
17 18	 Description of consensus methodology used for any Grade D recommendations is described.
19	 Conflict of interest records for GDG are summarised.
20 21 22 23	At the request of the GDG, an update search was carried out for review of perioperative fasting in children and young people (2004-2009). A full search was conducted for fasting for sedation in children and young people for diagnostic and therapeutic procedures.
24	Perioperative fasting in children and young people (2004-2009)
25 26 27	No RCTs or observational studies were identified in the update search which met the inclusion criteria for a review of fasting in this population in preparation for general anaesthesia.
28	Fasting for sedation in children and young people (all dates)
29 30	One RCT and six observational studies were identified in the search for fasting prior to sedation in this population.
31 32	Fasting State and Episodes of Vomiting in Children Receiving Nitrous Oxide for Dental Treatment ¹³² .

This controlled crossover study was performed to determine the frequency of vomiting during nitrous oxide/oxygen administration and to assess the relationship between fasting status and vomiting. A convenience sample of children (n=113) was randomly assigned to be fasting from solids for six hours and clear liquids for two hours before the procedure and their first dental treatment and non-fasting for the second treatment or alternatively, non-fasting initially and fasting for the next visit. The treatment time was under 35 minutes in all cases. The average fasting time was six hours before treatment in the fasting group and one hour before treatment in the non-fasting group. Vomiting occurred in only one subject, a child who was not fasting (1/113). This was a non significant result.

The following six studies represent observational data which records incidence of adverse events related to fasting status of children undergoing sedation. The data is summarised in Table 2. Adverse events related to pre-sedation fasting status

Table 2. Adverse events related to pre-sedation fasting status

Author	Total N	Adverse events/children	Adverse events/children not	Results
Study design	Age Range	fasted per guidelines (%)	fasted (%)	
Setting		3		
Drug				
Agrawal 2003 ¹¹	905	32/396 (8.1%) total adverse events	35/509 (6.9%) total adverse events	No association between fasting
Prospective case series	5 days – 18 years Median age: 5.4 years			state and adverse events. All adverse events were minor.
Emergency Dept, USA				Emesis occurred in 15 (1.5%) of patients. There
Mixed:				were no episodes of
47% ketamine				aspiration.
23% fentanyl and midazolam				
24% Chloral				
hydrate and				
pentobarbital				
Babl 2005 ²²	218	4/63 (6.3%)	11/155 (7.1%)	There were no
		vomiting	vomiting	serious adverse
Prospective case	14 months – 17			events and no
series	years			episodes of
F	Median age:			aspiration. The
Emergency Department,	8 years 3 monghs			recorded represent
Australia				emesis which
7.0311 4114				occurred in 15
50 – 70% Nitrous				children in total.
oxide				There was no
				significant
				association between
				preprocedural
				fasting and emesis
Heistein 2006 ⁹¹	1005			in this series.
⊓eistein ∠000′′	1095			Multivariate analysis showed
Retrospective review	1 month -3 years			that fasting times
Kenospeciive review	i monini -o years			(0.6-72 hours) were
Echocardiography,				not significantly

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Author	Total N	Adverse events/children	Adverse events/children not	Results
Study design Setting	Age Range	fasted per guidelines (%)	fasted (%)	
· ·				
Drug USA				associated with
Chloral Hydrate				adverse events (p=0.36) including apnea, airway obstruction, hypoxia, hypercarbia, hypotension, vomiting and prolonged sedation.
Keidan 2004 ¹²⁷ Retrospective review Auditory brainstem	200 infants (mean age 16 months + 10 in Groups A – fasted group and mean age 14	3/100 transient desaturation 25/100 prolonged sedation (>120	1/100 transient desaturation 5/100 prolonged sedation (>120	The fasted group showed significantly higher failure rate to achieve sedation with first dose
chloral hydrate, 50-	months + 13 in Groups B — not fasted group)	minutes) 2/100 agitation	minutes) 0/100 agitation	(p=0.03) and hence needed higher doses (p<0.01) and were sedated for
60 mg/kg		0/100 vomiting	0/100 vomiting	longer periods p<0.001). No
		21/100 failure to achieve adequate sedation with first dose	11/100 failure to achieve adequate sedation with first dose	difference was found in the adverse effect rate.
Roback 2004 ¹⁸⁸ Prospective cohort Emergency Department, USA Ketamine, midazolam and 53/2085 'other' drugs	2085 19 days -18 years Median age: 6.7 years	Fasted 2-4 hours: Respiratory (apnea, laryngospasm, osygen saturation <90%): 30/391 (7.7%) Vomiting: 40/391 (10.2%) Fasted 4-6 hours: Respiratory(apnea, laryngospasm, osygen saturation <90%): 31/430 (7.2%) Vomiting: 10/150 (6.7%) Fasted 6-8 hours: Respiratory(apnea, laryngospasm, osygen saturation <90%): 7/281 (9.6%) Vomiting: 18/281 (6.4%) Fasted >8 hours: (apnea, laryngospasm, osygen saturation <90%): 7/281 (9.6%)	Fasted 0-2 hours: Respiratory apnea, laryngospasm, osygen saturation <90%): 11/150 (7.3%) Vomiting: 30/430 (7.0%)	No significant differences were found in adverse events according to fasting times. No patients experienced clinically apparent aspiration.

Author Study design Setting	Total N Age Range	Adverse events/children fasted per guidelines (%)	Adverse events/children not fasted (%)	Results
Drug				
		osygen saturation <90%): 19/303 (6.3%) Vomiting: 27/303 (8.9%)		
Treston 2004 ²¹⁵	257	Longer than 3 hours: 20/127 (15.7%)	2-3 hours: 14/100 (14%) vomited	There was a non- significant trend to
Prospective cohort	1-12 years	vomited	1 hour: 2/30	increased incidence of vomiting with
Emergency			(6.6%) vomited.	increased fasting
Department,				times (p=0.08)
Australia				
Ketamine				

Other Relevant Publications

The Dental Clinical Guidance for conscious sedation in dentistry was published in 2006 ⁴⁵ and highlighted fasting before conscious sedation as an area requiring further high-quality research.

Another prospective cohort study in which children were sedated for gastroscopy with demerol or diazepam showed that there was no significant correlation between duration of fasting from 0.5 to 24 hours and either gastric volume or pH ⁹⁹.

4.2.3 GDG discussion

When considering what guidance should be provided in relation to fasting, the GDG looked at a range of possible recommendations. This ranged from no fasting is necessary prior to administration of sedation through to the application of standard fasting policy throughout the UK shaped by the joint RCN/RCA clinical guideline 'Perioperative fasting in adults and children' (2005)⁸, known colloquially as the "2-4-6"rule, namely 2 hours for clear fluid, 4 hours for breast milk and 6 hours for solids (including formula milk). This guideline was positively appraised as per NICE Technical Manual (2009)¹⁷⁰ using the AGREE instrument, and the initial GDG position was to apply standard fasting policy.

During GDG discussion, two main concerns emerged, these were that children and young people undergoing sedation should not be unnecessarily fasted and the importance of safety. One pharmacological intervention, nitrous oxide alone (up to 50% in oxygen), was felt to have no safety concerns and on this basis the GDG accepted that recommendations should reflect this. Given the publication date of the RCN/RCA guideline⁸, the original search strategy was re run to end of 2009, with an additional search applied to the target population of this guideline: children and young people receiving sedation and not general anaesthesia. While a number of studies were found, the quality of the evidence was weak, with the GDG choosing to apply the standard fasting recommendation from the Clinical Guideline 'Perioperative Fasting in Adults and

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1 Children' (2005) 8. The wording of the recommendation which focuses on 'elective 2 procedures', reflects an important GDG discussion on administering sedation for 3 emergency procedures. Clinical decision making in this context was recognised to 4 balancing the risks and benefits of sedation. The GDG noted that the fasting status of a 5 child presenting in the emergency context cannot be guaranteed and recognise the 6 importance of local clinical decision making given the clinical circumstances. It was also 7 noted by the GDG that recording pre sedation fasting was important and should be 8 inserted into the healthcare record. 9 4.2.4 Health economic considerations 10 An economic analysis was not carried out. It was anticipated that fasting will not significantly increase the health care resources required to manage a patient undergoing 11 12 a procedure. 13 4.2.5 Recommendations Recommendation 8 Before starting sedation, confirm and record the time of last food and fluid intake in the healthcare record. 14 **Recommendation 9** For elective procedures, apply the 2-4-6 rule.* 15 Recommendation 10 For urgent procedures in a child or young person who has not fasted, base the decision to proceed with sedation on clinical emergency and the target depth of sedation. 16 Recommendation 11 Fasting is not required for minimal sedation and for sedation with nitrous oxide alone (up to 50% in oxygen). 17

- * Fasting times should be as for general anaesthesia:
- 2 hours for clear fluids
- 4 hours for breast milk
- 6 hours for solids .- .

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4.2.6 Research recommendation

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation how long should they be fasted to prevent adverse events?

Why it is important

Inhalation of gastric contents can be fatal. Loss of consciousness is associated with the loss of vital airway reflexes and inhalation of gastric contents is possible. Consequently fasting (in order to keep the stomach empty) is standard practice before general anaesthesia and has become standard before any sedation technique that may cause loss of consciousness. Prolonged fasting however is distressing and can cause dehydration and hypoglycaemia. It would be helpful to know the minimum length of time necessary to fast a child before sedation in order to ensure that he stomach is empty, and to know that likelihood of regurgitation or vomiting is very small.

4.3 Psychological preparation

For a full narrative review on psychological preparation see chapter 5.

4.3.1 Clinical Introduction

A substantial body of research from different paradigms, affirms that children who have been repeatedly exposed to anxiety provoking painful medical events are at increased risk for developing adult dysfunctional cognitions and avoidant attitudes toward health care. In some cases, serious mental health problems, such as post-traumatic stress can occur. The pharmacological management of acute pain and anxiety in children undergoing therapeutic and diagnostic procedures outside the operating room has developed substantially in the past 15 years and procedural sedation is frequently used for the care of children in many medical settings. Pharmacological sedation and analgesia, however, do not adequately address the emotional, cognitive, and behavioural components that are integral to the sedation experience. Consequently, effective patient management requires an interdisciplinary approach and should include psychological techniques, which can be used alone or in combination with pharmacological treatment.

4.3.2 Clinical methodological introduction

CLINICAL QUESTION:

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques

7. what standard psychological preparation, coping skills and strategies should be used?

Population: Infants, children and young people under the age of 19 who are receiving sedation by any technique for diagnostic or therapeutic procedures including dental surgery and minor operations carried out under local anaesthesia.

Intervention: Psychological preparation.

Comparisons:

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ı	No intervention, usual care					
2	Pre-medication					
3	Another non-pharmacological treatment					
4	Outcomes for efficacy of psychological preparation:					
5	Completion of procedure					
6	Behavioural ratings including:					
7	 Pain as assessed using validated pain scales such as FACE, VAS, 					
8 9	 Children's Hospital of Eastern Ontario Pain Scale (CHEOPS), Spielberger State- Train Anxiety Inventory (STAI). 					
10 11	 procedural distress as assessed by validated scales such as Observational Scale of Behavioural Distress (OSBD) 					
12	 Parent/patient satisfaction 					
13	Sedation timing including					
14	 Length of induction (defined as time from administration of sedation 					
15	o drug to initiation of procedure)					
16 17	 Length of recovery (defined as time from completion of procedure to recovery criteria being met) 					
18 19 20 21 22	The search for psychological preparation for paediatric sedation included both quantitative and qualitative literature. Only two RCTs were identified and therefore the review for this intervention was primarily a narrative review of observational studies and randomized controlled clinical trials conducted in other relevant contexts, that is, induction for anaesthesia and medical procedures (See chapter 5).					
23	4.3.3 Clinical Evidence Statements					
24 25	The effects of a psychological preparation program on anxiety in children and adolescents undergoing gastrointestinal endoscopy; Mahajan 1998 ¹⁵⁴ .					
26 27 28 29 30 31 32	This study was carried out at the Cleveland Clinic in the USA in a population of children and young people ages 6-19 years. In a sample of 60 patients, the control group received usual patient education and the intervention group received psychological preparation consisting of demonstration of materials that would be used in the procedure. A doll was used as a model, if age appropriate. A book with photographs of a child undergoing the procedure was also shown. The same child life specialist provided all of psychological preparation.					
33 34 35 36	In this study, the outcomes of anxiety and distress were measured using validated scales. The Speilberger State-Trait Anxiety Inventory (STAI) was administered to patients after the psychological intervention but before the endoscopic procedure. The Observational Scale of Behavioural Distress (OSBD) was administered during the procedure.					

- Compared to usual care the children receiving psychological preparation had significantly less anxiety before the procedure [low quality evidence].
- There was no significant difference between the groups in distress levels as measured by the OSBD instrument although patients in the intervention group had a lower weighted mean score interval (1 versus 1.3).

Author(s): Mahajan 1998 **Date:** 2009-11-14

Question: Should psychological preparation versus usual care be used for paediatric sedation?

	Quality accessment							Summary of findings				
	Quality assessment							No of patients Effect				
No of studie s	Design	Limitatio ns	Inconsiste ncy	Indirectness	Imprecis ion	consider	psychol ogical preparati on	usual care	Relati ve (95% CI)	Absolute	Quality	Import ance
Anxiety	Anxiety (range of scores: -; Better indicated by less)											
1	randomise d trial	serious ¹		no serious indirectness	serious ²	none	30	30	-	MD -10.10 (-13.77 to -6.43)	LOW	
Distres	Distress (range of scores: -; Better indicated by less)											
1	randomise d trial	serious ¹		no serious indirectness	serious ²	none	30	30	-	MD -0.30 (-0.88 to 0.28)	LOW	

1 Method of randomisation and allocation not described. Blinding of assessors not described.

9 Anticipatory anxiety in children visiting the dentist: lack of effect of preparatory information; Olumide 2009 173

This study was carried out at the Kings College Hospital paediatric dental clinic, London, in a population of children ages 8-12 years. In a sample of 50 patients, the intervention group received a preparatory leaflet and the control group received a leaflet about healthy eating. Anxiety levels were measured using the Facial Image Scale before and after children read their leaflets. Intra-group comparisons were made. No inter-group statistics were calculated

In both groups there was no significant difference in anxiety levels before or after reading the leaflets [moderate quality evidence].

² Small study with 30 participants in each group. Outcome measures dependant upon subjective perception of anxiety and distress despite validation.

Author(s): Olumide 2009
Date: 2010-02-01

Question: Should preparatory leaflet be used for anxiety?

Settings: dental treatment

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			Quality assesse	Summary of findings								
	Quality assessment								No of patients Effect			
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	preparatory leaflet	control	Relative (95% CI)	Absolute	Quality	Importance
Anxiety wit	Anxiety with preparatory leaflet (range of scores: -; Better indicated by less)											
1			no serious inconsistency	no serious indirectness	serious ¹	none	25	25	-	MD 0.56 (0.08 to 1.04)	MODERATE	
Anxiety with healthy eating leaflet (range of scores: -; Better indicated by less)												
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	25	25	-	MD 0.24 (-0.16 to 0.64)	MODERATE	

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2 No explanation was provided

¹ Although sample size calculations were acceptable for 80% power, this remains a small study and should be repeated in larger population

1 4.3.4 **GDG** discussion 2 The GDG noted that sedation is only one of the management options available for 3 children and young people undergoing therapeutic or diagnostic procedures. 4 Psychological interventions can be used to reduce anxiety and manage behaviour in 5 combination with sedation. 6 Parental involvement in the preparation of the child and during the procedure may 7 reduce the distress caused by separation anxiety, particularly in young children. 8 The GDG believe psychological techniques (for example information for the 9 patient/carer about before during and after sedation, cognitive behavioural therapy, 10 distraction, guided imagery, hypnosis, demonstration play therapy, music therapy) form 11 part the child/family preparation. An individualised approach to using these techniques 12 will benefit the child and minimise fear, anxiety, pain and distress. 13 In making the recommendations, the GDG agreed that health care professionals involved 14 in sedating should: 15 Have knowledge and understanding of psychological methods of patient 16 preparation and coping skills and strategies e.g. the "tell-show-do" method, simple 17 distraction techniques 18 Consider psychological techniques for the child and family as part of patient 19 preparation and tailor to the age, understanding and needs of the child/parent 20 Involve the parent/carer in the preparation of the child and during the procedure 21 Offer factual information about the clinical setting, the procedure itself, the different 22 steps of the procedure 23 Offer information and discussion about what the child may experience before, 24 during and after the procedure 25 Discuss coping strategies and skills with the child/family 26 Consider using trained psychosocial professionals for patient preparation 27 Modify psychological methods of preparation according to the urgency of the 28 procedure 29 4.3.5 Health economic considerations 30 An economic analysis was not conducted. Preparation for children and young people 31 undergoing diagnostic and therapeutic procedures under sedation techniques was felt to 32 be part of routine care. Providing patients and their families with information on coping 33 strategies was felt to be part of a routine.

1 4.3.6 Recommendations

Recommendation 12 Ensure that the child or young person is prepared psychologically for sedation by offering advice about: the procedure itself what the child or young person should do and what the healthcare professional will do the sensations associated with the procedure (for example, a sharp scratch, numbness) how to cope with the procedure.

Recommendation 13

Ensure that the information is appropriate for the developmental stage of the child or young person.

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Recommendation 14

Offer parents and carers the opportunity to be present during sedation when appropriate. If a parent or carer decides to be present, offer them advice about their role during the procedure.

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Recommendation 15

For an elective procedure, consider referral to a mental health specialist for children who are severely anxious or who have a learning disability.

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4.3.7 Research recommendation

For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures under sedation what psychological techniques can lead to sedation sparing, improve patient/family satisfaction, and ensure safe completion of the procedure?

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4.4 Personnel and training

4.4.1 Clinical Introduction

All healthcare professionals involved in the care of sedated children and young people should be appropriately trained. The training of health care professionals delivering sedation currently varies by speciality. There are a number of reports that provide guidance on the types of training courses available (for example "Conscious Sedation in the provision of dental care" ²⁰⁶) but there remains significant variability between different healthcare providers and specialities.

The aim of this section is to provide clear advice on training requirements to ensure that every healthcare professional is competent in the sedation techniques they use and in the management of complications that might arise when using these techniques. This is important because there is currently no uniform requirement for assessing sedation skills, nor any consistent requirement for revalidation of skills.

Training may be delivered by Trusts, Universities, Royal Colleges or other independent providers but the responsibility for ensuring that health care professionals have undergone appropriate training should lie with the local NHS Trust providing sedation services.

4.4.2 Clinical methodological introduction

CLINICAL QUESTIONS:

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation

- 8. what generic and specific skills are required for different team members and for different levels of sedation?
- 9. what training and competences are required for the personnel involved?
- 10. what assessment and maintenance of skills is required for the personnel involved?

GDG sought to provide guidance to these questions based on their expert experience and opinion.

Skills required for sedation

The GDG agreed that sedation should be administered by a team and someone in the team should have the skills to ensure the sedation is effective and that any complications are managed successfully. Many types of skills were discussed including pre-sedation patient assessment and communication. During sedation until the end of recovery the skills of observation and monitoring were considered to be essential for safety. These include airway patency, breathing rate and depth, pulse, pallor and cyanosis and depth of sedation. The complications of airway obstruction and respiratory arrest can be readily overcome by prompt recognition and management; if they occur serious consequences should be unlikely. These skills need to be practised regularly. The skills for the management of cardiac arrest are also essential.

Training and competencies

The GDG agreed that all healthcare practitioners administering sedation need to be trained in the practice of delivering effective sedation. Since there are a number of sedation techniques, the training and competencies would need to be specific to the sedation technique. Some generic skills were agreed such as the assessment of conscious level and pain. In respect to the complications of sedation, however, the GDG accepted that some sedation techniques were not safe enough to be used unless healthcare practitioners had specific training. They would need to be trained to manage the complications of that technique. If airway or respiratory complications were considered to be extremely unlikely, then some skills may be considered unnecessary. The recommendations took account of the likelihood of airway and respiratory complications of the sedation according of the technique and the target level of sedation.

Techniques with a narrow margin of safety readily cause airway obstruction and apnoea. Consequently the GDG believed that these drugs could only be recommended for use by teams with special expertise. This situation applies to anaesthetic agents and also the use of some combinations of drugs with opioids. The risk of opioids relates to judging the correct dose to overcome the pain. If the pain reduces (for example after the extraction of a tooth) the opioid causes the respiratory depression and this is made more likely if the patient is deeply sedated.

The GDG noted it is essential that healthcare practitioners undergo competency-based assessment upon completion of training to ratify their ability to undertake sedation on children and young people. Current practice varies between providers and specialities and there is currently no uniform requirement for assessing sedation skills, nor any consistent requirement for healthcare practitioners to revalidate their skills.

Assessment and maintenance of skills

GDG pointed out that there are a number of reports which have provided guidance on the nature of training but there remains variability across different health care providers and specialities.

The GDG considered the following in making recommendations by consensus:

- Health care professionals practising sedation should have documented evidence of competency. This should include:
 - Satisfactory completion of knowledge-based learning. (for example certificate confirming completion of a didactic training course covering the theoretical principles of sedation practice)
 - Log/record of satisfactory acquisition of practical and clinical skills relevant to the type of sedation being used including:
 - log-record of patients managed under supervision
 - a record of successful completion of work-based assessments (for example Direct Observation of Procedural Skills - DOPS)
- Healthcare professionals practising sedation should keep their skills up-to-date by regular practice of sedation techniques and reinforcement of theoretical and practical skills, undertaken as an essential component of Continuing Professional Development

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Healthcare professionals should maintain documentary evidence of clinical activity and continuing professional development in sedation

3 4.4.3 Health economic considerations

An economic analysis was not carried out. The cost of training health care professionals is not normally considered within cost-effectiveness analysis but may be included in the budget impact analysis.

4.4.4 Recommendations

Recommendation 16

Healthcare professionals delivering sedation should have knowledge and understanding of:

- Sedation drug pharmacology and applied physiology
- Assessment of the child or young person
- Monitoring
- Recovery care
- Complications and their immediate management, including paediatric life support.

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Recommendation 17

Healthcare professionals delivering sedation should have practical experience of:

- Effective delivery of the sedation technique used and management of complications
- Observing clinical signs (for example airway patency, breathing rate and depth, pulse, pallor and cyanosis, depth of sedation)
- Using monitoring equipment.

Recommendation 18

Ensure that all members of the sedation team have the following competencies:

Minimal sedation*	Moderate sedation	Deep sedation
All members have	All members have	All members have
basic life support	basic life support	basic life support
skills	skills	skills
	At least one team	At least one team
	member should	member should
	have intermediate	have advanced life
	life support skills in	support skills in
	airway management	advanced airway
	using mask	management and
	ventilation and use	cardiac arrest
	of defibrillator	management

^{*} and sedation with nitrous oxide alone (up to 50% in oxygen)

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Recommendation 19

Ensure that a healthcare professional trained in delivering anaesthetic agents is available to administer the following sedatives:

- Sevoflurane
- Propofol
- Opioids combined with ketamine.

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Recommendation 20

Healthcare professionals delivering sedation should have documented evidence (for example, a certificate or a comprehensive record) of competency including:

- evidence (for example a certificate) of satisfactory completion of a theoretical training course covering the principles of sedation practice
- a comprehensive record of practical experience of sedation techniques, including details of:
- children and young people managed under supervision
 - successful completion of work-based assessments.

Recommendation 21

Each healthcare professional and their team delivering sedation should ensure they update their knowledge and skills through programmes designed for continuing professional development.

4.4.5 Research recommendation

For personnel involved in delivering sedation to children and young people under the age of 19 undergoing diagnostic and therapeutic procedures what training is required to both achieve and maintain essential skills?

Why it is important

Potent drugs can cause unintended airway obstruction. Anaesthetists are skilled at managing airway obstruction because they practise them regularly. However, anaesthetists are a scarce resource so non-anaesthetists need to learn how to manage airway obstruction. The skills that are needed have been identified but can these skills be attained and maintained by professionals who need them occasionally? The GDG suggests that a standard teaching method and assessment tool are developed. This would involve an observational study of a cohort of trainees, who can be assessed, trained and then reassessed at varying intervals to determine whether the training is successful and how often it is necessary.

4.5 Clinical environment and monitoring

16 4.5.1 Clinical introduction

Sedation of children and young people happens in a variety of clinical environments, with a range of specialty staff, and a selection of different sedative agents.

Sedation carries a risk of serious adverse events, including hypoxia, reduced consciousness, apnoea, and loss of airway control. In some sedation techniques the sedation level can become deep rapidly, so, in order to ensure safety of the child or young person, it should be possible to monitor a child for a deeper level of sedation than planned.

Assessment of requirements for monitoring should be undertaken prior to any sedation event, and monitoring should start prior to administration of any sedation agent. Monitoring will depend not only on sedation technique but the child's tolerance, and may become less intrusive as the child becomes more awake.

This section makes recommendations for minimum levels of monitoring for all children receiving sedation, to reduce the risk of adverse events, and improve patient safety.

4.5.2 Clinical methodological introduction

CLINICAL QUESTIONS:

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation techniques:

11. during moderate or deep sedation techniques, what monitoring and equipment is required to reduce the risk of complications?

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12. when should monitoring stop?

GDG sought to provide guidance to these questions based on their expert experience and opinion.

4.5.3 GDG discussion

What monitoring is required?

The GDG aimed to provide consistency in monitoring standards, to provide some evidence around the use of capnography, to inform judgement and to reduce the risk of adverse events to patients.

The GDG noted that monitoring varies across specialties. In emergency care, monitoring commences prior to sedation. Vital signs are taken prior to commencement and documented at intervals throughout the procedure. The healthcare team's approach in determining frequency of observation/monitoring interventions is dependant on the procedure itself, level of sedation to be achieved and child's tolerance. The GDG indicated that in some sedation techniques the sedation level can become deep rapidly and monitoring should be increased if patient becomes unrousable or unconscious.

The GDG noted that patient monitoring needs to begin prior to administration of the agent(s) unless this causes unnecessary distress. GDG described sedation monitoring as a continuum from awake to anaesthesia which becomes less intrusive as the child becomes more awake.

The GDG raised concern about the difficulty in dealing with monitoring of children who are uncooperative, distressed or anxious; and on the lack of understanding of the potential effects/side-effects of drugs used, and risks of a changing target state. This concern reflects the range of possible behaviours and compliance observed in practice and the various techniques that healthcare professionals may apply in effectively managing this. Factors for consideration are seen in the recommendation and provide direction for the sedation team.

When should monitoring stop?

GDG noted that practitioners do sometimes take their 'eye off the ball' when the procedure is complete, but the child is still sedated. The GDG agreed, by general consensus, that the point at which monitoring stops is not the same as discharge criteria as sedation state may vary throughout recovery period, with the level of monitoring.

Staff and facilities should be available to manage an unconscious or an acutely sick patient until either they have recovered or they can be transported to another facility who can continue their care.

4.5.4 Health economic considerations

An economic analysis was not carried out. The appropriate monitoring will be largely determined by safety considerations. If the use of particular sedation technique increases the duration and intensity of monitoring, then this should have been captured in the cost estimate of that sedation technique. We have included the cost of staff and consumables associated with different sedation techniques in our economic analysis.

1 4.5.5 Recommendations

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For moderate sedation continuously monitor, interpret and respond to changes in all of the following:
- Coping
- Depth of sedation
- Pain
- Distress
- Respiration
- Oxygen saturation
- Heart rate.

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Recommendation 23	For deep sedation continuously monitor, interpret and respond to all of the following: - Respiration - Oxygen saturation - Heart rate - End tidal CO ₂ (capnography)
	- three-lead electrocardiogram (ECG)- Blood pressure (monitor every 5 minutes)
	- Depth of sedation
	- Pain
	- Coping
	- Distress.

Recommendation 24	Ensure that data from continuous monitoring during sedation
	are clearly documented in the healthcare record.

Recommendation 25

After the procedure, continue monitoring until:

- The airway is patent
- Protective airway and breathing reflexes are present
- The child or young person is haemodynamically stable
- The child or young person has returned to baseline level of consciousness.

2 4.5.6 Research recommendations

For children and young people under the age of 19 undergoing diagnostic and therapeutic procedures under sedation is the introduction of system of capnography monitoring cost-effective, compared to standard monitoring, in reducing adverse events?

Why it is important

Monitoring of airway patency and breathing during sedation is crucial. During anaesthesia this is achieved with capnography. During sedation this can be achieved by using soft nasal catheters. Capnographs are expensive and their ability to recognise airway obstruction and respiratory depression in sedated children is uncertain.

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Which depth of anaesthesia monitors can be used to monitor depth of sedation in children and which is best?

Why it is important

Several depth of anaesthesia monitors are in use around the world. Most use processed EEG signals while some use stimulation of the brainstem by auditory stimuli. It is not yet clear whether the available monitors can follow children through different levels of sedation accurately and this study would set out to determine which monitor best tracks the transition from moderate to deep sedation in children of different ages.

4.6 Discharge criteria

4.6.1 Clinical introduction

The aim of establishing discharge criteria is to ensure children go home from a sedation event only when it is safe for them to do so. Recovery from sedation is a continual process and some children might benefit from a longer period of less-intense observation before discharge home. This is particularly important when using sedation agents that Sedation in children and young people: full guideline DRAFT (May 2010) Page 88 of

have a prolonged effect and may delay a child's complete recovery, or pose the risk of re-sedation.

4.6.2 Clinical methodological introduction

CLINICAL QUESTION:

For children and young people under the age of 19 after diagnostic and therapeutic procedures under sedation techniques

13. what discharge criteria are required?

GDG sought to provide guidance to this question based on their expert experience and opinion.

4.6.3 GDG discussion

The GDG noted that in current practice, discharge criteria varies across specialties and professionals. In emergency care, children will be observed/monitored until they reach a 'pre-sedation' state. They are discharged into the care of a responsible adult, and advice is given on what to expect in the first 24 hours after sedation. Recovery from both the procedure and the sedation takes a variable length of time and depends upon the procedure, its length, the sedation technique and the doses used.

A simple checklist can be used to make sure that children have returned to their presedation states. However this should also take into account the capabilities of the person caring for the child following discharge, the presence of other medical problems and the distance the family has to travel to obtain medical assistance It is more important to individualize the times of discharge rather insist on a minimum length of stays.

Recovery from sedation caused by some drugs and techniques can be prolonged and unpredictable and there is a risk that after discharge the patient may become resedated. In this situation there may be a danger of respiratory depression and hypoxia. Prolonged sedation may also mean that intake of drink and food may be delayed leading to dehydration and hypoglycaemia. These problems may be more common with orally administered drugs because absorption can be delayed and unpredictable.

Sedation may not always succeed; the drugs may not be effective enough at the desired target level of sedation. If a patient becomes too distressed and cannot cope or cooperate with a painful procedure, increasing the doses of sedation drugs may only be effective if they cause deep sedation of anaesthesia. Likewise, if sedation does not cause a child to sleep during painless imaging, increasing the doses may only be effective if the child becomes unconscious. Deep sedation techniques often cause a prolonged recovery and have the associated hazards of suppression of vital airway and breathing reflexes. In these circumstances anaesthesia drugs are more suitable because they can be given in the dose required to cause the sedation level that the patient needs. Moreover they are short acting drugs and can be given to cause sedation or anaesthesia over the period of the procedure; they do not cause prolonged recovery times. If the healthcare professional is suitably trained and has the facilities for anaesthesia, anaesthesia is feasible as soon as patient needs it. Often, the skills and facilities are not available and anaesthesia will need to be arranged at another time and place.

4.6.4 Health economic considerations

An economic analysis was not carried out. The choice of discharge criteria should be based on minimizing the risk that a patient will experience an adverse event after discharge. If the use of a particular sedation technique results in the patient taking longer to meet the discharge criteria, and is associated with increased duration of stay, this should have been accounted for in the costing of the technique. We have included the cost of recovery in our costing of sedation techniques.

4.6.5 Recommendations

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Recommendation 26 Ensure that all of the following criteria are met before the child or young person is discharged:

- Vital signs* have returned to normal levels
- The child or young person has returned to baseline level of responsiveness and orientation
- Nausea, vomiting and pain have been adequately managed
- There is no risk of further reduced level of consciousness.

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Recommendation 27	Consider referring to an anaesthesia specialist if the child or young person is not able to tolerate the procedure under sedation.

^{*} Vital signs are measures of various physiological statistics and usually include body temperature, heart rate, blood pressure and respiratory rate.

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5 Psychological preparation

5.1 Narrative review

5.1.1 Introduction

This narrative review provided material to inform the GDG and to enable consensus decisions leading to recommendations on how children and young people should be prepared prior to their sedation experience. The nature of the evidence base in this area lends itself to this approach.

The benefits of a systematic narrative review of the clinical evidence are highlighted by Oxman¹⁷⁴ and colleagues. Applying the quality assurance principles advocated by Oxman¹⁷⁴, a valid review article can provide the best possible source of information that can lay a foundation for clinical decisions to be made. There is argument that focused narrative reviews for these important areas of preparation and assessment of the child prior to sedation are more likely to provide valid results that are useful for clinicians. Having provided the background and context for this review, we begin by defining psychological preparation and stating its aims and factors that affect its exact nature and content. It continues by summarising the evidence for the efficacy of psychological preparation for anaesthesia induction and other medical procedures. Following this, the literature regarding parental and children's desire for information is reviewed. Next the evidence regarding the effects of parental presence during anaesthesia induction and other medical procedures is discussed along with the role that parents play when present. The review concludes by summarising the existing evidence and good clinical practice and making recommendations for the preparation of children and their parents for sedation.

5.1.2 What is psychological preparation

Psychological preparation includes specific interventions to provide information and reduce anxiety. Providing three types of information is central: (a) information is provided about the procedure itself (i.e. steps that children must perform and steps that healthcare professionals will perform); (b) the sensations the patient can expect to feel (e.g. sharp scratch, numbness); and (c) about how to cope with the procedure¹⁴¹.

The aim of presedation and/or preprocedure psychological preparation in children and young people is to:

- reduce anxiety for patients and their parents
- improve patient cooperation

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- enhance patient postprocedure recovery
- increase self-control for patients and their parents
 - improve long-term emotional and behavioural adjustment in patients and their parents

The factors affecting presedation and/or preprocedure preparation are (Kain & Caldwell-Andrews, 2005)¹⁰²:

- the developmental stage of the child or young person
- previous medical experiences
- timing relative to the procedure
- temperament, current anxiety levels and coping style
- role of parents

There is limited evidence regarding the best way to prepare children and young people for sedation therefore the extensive related literature on preparation for painful medical procedures and anaesthesia were reviewed and the results of this body of knowledge informed the present recommendations. Overall, published evidence supports the view that good preparation results in improved sedation outcomes (e.g. less distress and improved adjustment for the parent and patient ^{138,157}. A number of studies have shown that adequate preprocedural preparation can also reduce anxiety and procedural pain for a range of medical events, including venipuncture¹²⁹, dental procedures ¹⁶⁵ surgery¹⁰² and voiding cystourethrography²⁰⁷.

5.1.3 Psychological preparation for anaesthesia induction

Children have numerous concerns related to anaesthesia and surgery including fear of separation, fear of physical harm, fear of the unknown, fear of death, fear of losing control, and uncertainty of the limits of acceptable behaviour^{79,193}. It has been estimated that 50% to 75% of children undergoing surgery will develop extreme anxiety and distress during the perioperative period¹²¹. Anxiety experienced by children at induction is associated with distress on awakening in the recovering area and with later postoperative behavioural problems²²⁸. Younger age, behavioural problems with previous health care attendances, longer duration of procedure, having more than five previous hospital admissions and anxious parents at induction are associated with high anxiety at induction⁵⁰. Interestingly, mother's prediction of uncooperative behaviour is a good predictor of anxiety during induction¹⁵³. 54% of all children undergoing general anaesthesia and surgery exhibit new onset maladaptive behavioural responses including general anxiety, night-time crying, enuresis, separation anxiety, eating disturbances, sleep related problems, and temper tantrums at 2 weeks postoperatively^{104,112,120}.

Behavioural preoperative preparation has been advocated in the psychological and medical literature as a way to ameliorate children's preoperative anxiety and facilitate post procedure recovery. An estimated 78% of all major hospitals offer such programmes to children and their parents. These preparation programmes may provide narrative information, an orientation tour, role rehearsal using dolls, a puppet show, child life preparation or the teaching of coping and relaxation skills to children and their

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parents. Although there is a general consensus about the desirability of these programmes, recommendations regarding the content of preoperative preparation for children differ widely. O'Byrne and colleagues ¹⁷² asked a panel of psychological experts to rate the effectiveness of behavioural preparation programs used in the United States prior to surgery. Experts rated each program on a 1 (least effective) to 9 (most effective) Likert scale. Coping skills instruction was ranked as the most effective preoperative intervention, followed by modelling, play therapy, operating theatre (OR) tours, and printed materials.

Kain and Caldwell-Andrews¹⁰² suggest that a number of variables are important to consider when designing a preparation program including child age, timing relative to surgery, and the child's previous hospitalization history. For example, participation in a preparation program more than 5 to 7 days prior to surgery has been found to be most beneficial for children 6 years and older and the least beneficial when the program is given 1 day before surgery^{114,166,191}. Previous hospitalization history can be a particular challenge for designing a preparation program as well¹⁰². Information about what to expect on day of surgery does not offer new knowledge to these children and ⁶⁵ have further demonstrated that simple modelling and play programs are not beneficial for children with previous hospitalizations. Individualized coping skills training in combination with actual practice have been identified as strategies that are more helpful for these children ¹¹⁵. Kain and Caldwell-Andrews¹⁰² suggest that the latter types of programs should be designed with the child's specific past experiences in mind.

5.1.4 The benefit of preoperative anxiety reduction programmes – what the evidence says

- Kain and colleagues ¹⁰⁹ in an RCT compared three types of behavioural preoperative preparation programs including a tour of the OR (information based), an information based + modelling based programme (OR tour + commercially available videotape), or an information + modelling + coping based programme (OR tour + videotape + Child Life preparation) with 75 children aged 2 to 12 years. Children and parents who received Child Life coping skills preparation exhibited less anxiety immediately following the preparation in the holding area on the day of surgery and on separation to the OR than children and parents who did not receive this preparation. There were no significant differences in anxiety levels across the groups during anaesthetic induction, in the recovery room, or at 2 weeks following the operation.
- Golan, Tighe, Dobija, Perel and Keidan ⁷⁸ found that the use of preoperative medically trained clowns for children undergoing surgery can significantly alleviate preoperative anxiety. In a randomised, controlled and blinded study conducted with 3-8 year olds undergoing GA for elective outpatient surgery patients were assigned to three groups: Group 1 did not receive midazolam or clown presence (n=22), Group 2 received 0.5mg/kg oral midazolam 30min before surgery up to a maximum of 15mg (N=22), and Group 3 had two specially trained clowns (N=21) present upon arrival to the preoperative holding area and throughout operating theatre entrance and mask application for inhalation induction of anaesthesia. The intervention lasted approximately 20min and the clowns used developmentally appropriate techniques such as magic tricks, gags, music, games, puppets, word games and bubbles. In all groups parents were present. All children in the study were videotaped in the holding area until the induction of anaesthesia and blinded evaluators used the tapes to rate children's anxiety. The clown group had a statistically significant lower

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modified-Yale preoperative Anxiety Scale score (m-YPAS; Kain, Mayes, Cicchetti et al., 1997¹¹⁶) in the preoperative holding area compared to a control and a midazolam group. The clowns' effect on anxiety reduction continued when the children entered the operating theatre but was equal at this point to the midazolam group. Upon application of the anaesthesia mask no statistically significant differences were detected between groups but the clown group had the largest increase in m-YPAS score which surpassed the other two groups m-YPAS scores.

- Kain, Caldwell- Andrews, Krivutza, Weinberg, Gaal, and colleagues¹⁰³ compared the effectiveness of an interactive music intervention and midazolam in alleviating preoperative anxiety in 123 children aged 3 to 7 years old. The results of this study suggested that interactive music therapy may be useful in alleviating preoperative anxiety on separation from parents and entrance to the OR, but that music therapy did not appear to alleviate children's anxiety at anaesthetic induction.
- Kain and colleagues¹⁰⁷ randomly assigned 408 children and their parents to one of four groups: (1) control which received standard of care; (2) parental presence which received standard parental presence during induction of anaesthesia; (3) ADVANCE: received standard-of-care treatment plus multicomponent family-centered behavioural preparation (Anxiety-reduction, Distraction, Video modelling and education, Adding parents, No excessive reassurance, Coaching, and Exposure/shaping); and (4) oral midazolam. Parents and children in the ADVANCE group exhibited significantly lower anxiety in the holding area as compared with all three other groups (34.4+)-16 vs. 39.7+15; P=0.007) and were less anxious during induction of anaesthesia as compared with the control and parental presence groups (44.9+/-22 vs.)51.6+/-25 and 53.6+/-25, respectively; P=0.006). Anxiety and compliance during induction of anaesthesia was similar for children in both the ADVANCE and midazolam groups (44.9+/-22 vs. 42.9+/-24; P=0.904). Children in the ADVANCE group exhibited a lower incidence of emergence delirium after surgery (P=0.038), required significantly less analgesia in the recovery room (P=0.016), and were discharged from the recovery room earlier (P=0.04) as compared with children in the three other groups.
- A recent meta-analysis²³⁵ that assessed the effects of non-pharmacological interventions in assisting induction of anaesthesia in children by reducing their anxiety, distress or increasing their co-operation concluded that nonpharmacological interventions such as parental acupuncture; clown doctors; hypnotherapy; low sensory stimulation; and handheld video games are promising and need to be investigated further. More specifically six trials assessed interventions for children. Preparation with a computer package improved cooperation compared with parental presence³⁶. Children playing hand-held video games before induction were significantly less anxious than controls or premedicated children¹⁷⁷. Compared with controls, clown doctors reduced anxiety in children (modified Yale Preoperative Anxiety Scale (mYPAS): mean difference (MD) 30.75 95% Cl 15.14 to 46.36; Vagnoli 2005²¹⁸). In children undergoing hypnosis, there was a nonsignificant trend towards reduced anxiety during induction (mYPAS < 24: risk ratio (RR) 0.59 95% CI 0.33 to 1.04 - 39% versus 68%: Calipel 2005³⁴) compared with midazolam. A low sensory environment improved children's co-operation at induction (RR 0.66, 95% Cl 0.45 to 0.95; Kain 2001¹²⁰) and no effect on children's anxiety was found for music

therapy¹⁰³. Parental interventions were assessed in three trials. Children of parents having acupuncture compared with parental sham-acupuncture²²⁶ were less anxious during induction (mYPAS MD 17, 95% CI 3.49 to 30.51) and more children were co-operative (RR 0.63, 95% CI 0.4 to 0.99). Parental anxiety was also significantly reduced in this trial. In two trials^{161,238}, a video viewed preoperatively did not show effects on child or parental outcomes.

5.1.5 Psychological preparation/interventions for other medical procedures-what the evidence says

- Megel et al¹⁶⁴ examined how parents prepared their children before preschool immunizations. Five types of preprocedural preparation/discussion were postulated: information sharing (what will happen), sensory information (how it will feel), justifying the procedure (explaining why the procedure is necessary), teaching relaxation strategies, and role playing. The results suggested that parents used a mixture of various types of preparation. Seventy-five percent of children received informational preparation from their parents, typically involving a description of the events that would occur. Of the 25% of children who received no information, 9 children were <3 years of age. Forty-two percent of parents also used some sensory information in their description. Forty percent of parents offered a rationale for receiving the injection. Relatively few parents (10%) offered the children any strategies for how to cope with the procedure (eg, relaxation, breathing, or distraction). Unfortunately, the relationship between the type of preparation and the child's subsequent distress was not reported by the researchers.
- Uman et al²¹⁷ assessed the efficacy of cognitive-behavioural psychological interventions for needle-related procedural pain and distress in children and adolescents. Only randomized controlled trials (RCTs) with at least five participants in each study arm comparing a psychological intervention group with a control or comparison group were eligible for inclusion. Twenty-eight trials with 1951 participants were included. Together, these studies included 1039 participants in treatment conditions and 951 in control conditions. The most commonly studied needle-procedures were immunizations and injections. The largest effect sizes for treatment improvement over control conditions exist for distraction 37,62,184 (self-reported pain: SMD = -0.24, 95% CI = -0.45 to -0.04), hypnosis 142,143,145,146 (self-reported pain: SMD = -1.47, 95% CI = -2.67 to -0.27; self-reported distress: SMD = -2.20, 95% CI = -3.69 to -0.71; and behavioural measures of distress: SMD = -1.07, 95% CI = -1.79 to -0.35), and combined cognitive-behavioural interventions^{29,40,41,142} (other-reported distress: SMD = -0.88, 95% CI = -1.65 to -0.12; and behavioural measures of distress: SMD = -0.67, 95% CI = -0.95 to -0.38). The authors commented that while there may be preliminary evidence to support the efficacy of information/preparation there is not enough evidence at this time to make strong conclusions. More specifically, Harrison⁸⁸;Tak et al.²⁰⁹ reported information/preparation was effective in reducing observer-reported distress (SMD= -0.77, 95%CI = -0.17 to -0.38) and pulse rates (SMD = -0.47, 95% CI = -0.87 to -0.07). Although SMDs for self-reported pain and observer-reported distress both fell in the negative range (-0.22 and -0.15), their Cls passed into the positive range, indicating that while there may be preliminary evidence to support the efficacy of information/ preparation on these outcome, there is not enough evidence at this time to make strong conclusions. Information / Preparation did not appear to be effective in

reducing distress as assessed by behavioural measures (SMD = 0.24, 95% Cl = -0.30 to 0.78), as the SMD fell in the positive range.

- Sinha et al (2006)²⁰⁴ assessed the effectiveness of distraction techniques in reducing the sensory and affective components of pain among paediatric patients undergoing laceration repair in the ED. 240 children between 6 and 18 years of age were randomly assigned to an intervention or control arm. Those assigned to the intervention arm were given a choice of age-appropriate distracters during laceration repair. Quantitative measures of pain intensity, situational anxiety, and pain distress (as perceived by the parent) were assessed by using the 7-point Facial Pain Scale, State Trait Anxiety Inventory for Children, and a visual analogue scale, respectively, before and after laceration repair. The State Trait Anxiety Inventory for Children was performed in children > or =10 years of age. There was no difference in mean change in Facial Pain Scale scores between the control and the intervention groups in children < 10 years of age. Multivariate analysis in this same age group showed that the intervention was independently associated with a reduction in pain distress as perceived by parents based on the mean change in visual analog scale scores. In older children, the intervention was independently associated with reduction in situational anxiety but not in pain intensity or in parental perception of pain distress.
- Haeberli S et al (2008)⁸⁶ examined whether a psychoeducational intervention might reduce the need for anaesthesia during radiotherapy. 223 consecutive paediatric cancer patients receiving 4141 RT fractions during 244 RT courses were studied. Whereas in 154 RT courses corresponding with 2580 RT fractions patients received no psychoeducational intervention (group A), 90 RT courses respectively 1561 RT fractions were accomplished by using psychoeducational intervention (group B). This tailored psychoeducational intervention in group B included a play program and interactive support by a trained nurse according to age to get familiar with staff, equipment and procedure of radiotherapy. Group A did not differ significantly from group B in age, gender, diagnosis, localization of RT and positioning during RT. Whereas 33 (21.4%) patients in group A got anaesthesia, only 8 (8.9%) patients in group B needed anaesthesia. The median age of cooperating patients without anaesthesia decreased from 3.2 to 2.7 years. In both uni- and multivariate analyses the psychoeducational intervention significantly and independently reduced the need for anaesthesia.
- Train et al $(2006)^{214}$ evaluated the effect of a psychological approach on distress and sedation rates in children undergoing dimer captosuccinic acidlabelled with technetium-99 (99mTc) DMSA imaging. Baseline data, on a retrospective consecutive sample of children examined using DMSA over a 6-month period (n = 81), were collected via medical note search and postal questionnaire. A further consecutive sample of 40 children was recruited prospectively to the intervention, which consisted of distraction during medical procedures and environmental manipulation. In addition half of the intervention group were provided with a photo-booklet depicting a coping child model, together with a letter offering advice to parents on how to prepare their child for the procedure. Sedation rates were lower (p = 0.003) and service satisfaction ratings higher (p = 0.002) in the Intervention group as compared with the Baseline group. Within the intervention condition, children who received the photo-booklet displayed less distress before the procedure (p = 0.01) than those

who did not. Also families who received the photo-booklet were more likely to attend the appointment (p = 0.024).

5.1.6 Psychological preparation for dental procedures

In dentistry, the American Academy of Pediatric Dentistry (AAPD) recognises that, in providing oral health care for young patients a continuum of both nonpharmacological and pharmacological behaviour guidance techniques may be used by dental health care providers and recommends behavioural guidance to be used in combination with pharmacological interventions for the management of the young dental patient¹⁴. Techniques recommended include:

- Tell-show-do is a technique of behaviour shaping first described by Addelston¹⁰ that involves verbal explanations of procedures in phrases (what, why and how a procedure will be performed) appropriate to the developmental level of the patient (tell); demonstrations for the patient of the visual, auditory, olfactory, and tactile aspects of the procedure in a carefully defined, nonthreatening setting (show); and then smoothly with no break in time and without deviating from the explanation and demonstration, completion of the procedure (do). The tell-show-do technique is used with communication skills (verbal and nonverbal) and positive reinforcement^{66,94}.
- Voice control is a controlled alteration of voice volume, tone, or pace to influence and direct the patient's behaviour.
- Positive reinforcement involves the reward of desired behaviours with social reinforcers such as positive voice modulation, facial expression, verbal praise, and appropriate physical demonstrations of affection by all members of the dental team and nonsocial reinforcers such as tokens and toys.

5.1.7 Parental desire for information

Parents are frequently dissatisfied with the lack of information they are offered and express a strong desire for perioperative information. Many health care professionals may withhold information because of a belief that details will induce anxiety in parents which in turn will be communicated and increase the anxiety of children. Empirical evidence does not support this view.

- Kain et al.¹¹⁹ explored parents' desire for perioperative and anaesthetic information at a pre-surgical assessment clinic visit or on the day of their children's outpatient surgery. Almost all parents (95%; n = 317) wished to receive comprehensive information concerning their child's anaesthetic including information about all possible complications.
- Waisel and Troug²²⁵ evaluated parents' perceived understanding and anxiety related to the discussion of the general anaesthesia risks for children that occurred during the preoperative interview with the anaesthetist, immediately prior to surgery. Approximately half the sample (N = 55) was most concerned about the anaesthetic aspects of surgery (n = 25), and 39% (n = 21) were equally concerned about anaesthesia and surgery. Over 90% (n = 50) of parents reported that the discussion of anaesthetic risks was desirable and that they understood the information. Half of the sample (n = 25) felt the discussion

did not change their anxiety, whereas 25% (n = 13) felt it decreased anxiety and 24% (n = 12) felt it increased anxiety.

- Litman et al.¹⁴⁷ examined parental knowledge and desire for information regarding risk of death from anaesthesia in 115 parents of healthy children undergoing elective surgery. The majority (87%) wanted to know the chance of death after anaesthesia and over half of parents (68%) had accurate knowledge of risk of death from anaesthesia. Most parents (75%) also wanted to know all possible risks, however, this was greater for mothers than fathers. A separate group of parents (n = 121) were surveyed after participating in a preanaesthetic discussion with the anaesthetist. In 60% of cases, risk of death from anaesthesia was mentioned or implied and the proportion of parents who said they had wanted this information was similar to the previous survey. No demographic factors influenced the responses. However, several parents did not want the risk of death discussed in front of the children, who were sometimes present during the discussions.
- Franck and Spencer⁷⁰ critically analysed the published research literature (6 descriptive and 5 intervention studies) on providing information about children's anaesthesia to parents. The intervention studies tested different methods of providing information, including verbal, video or written modalities and showed some improvements in knowledge, anxiety and satisfaction. The authors concluded that parents want detailed information about the specifics of anaesthetic procedures, risks, and personnel roles.

5.1.8 Children's desire for information

There is widespread agreement that children should be given information prior to anaesthesia, surgery and medical procedures but continuing debate about the most appropriate form and content of that information. There is little research evidence about children's concerns, fears, and misconceptions about hospitals, anaersthesia and medical procedures and paucity of data regarding children's desire for perioperative information²⁰⁵.

Fortier et al 68 studied the perioperative information children want to receive from the medical staff. On the day of surgery, 143 children aged 7-17 yr (ASA I or II) completed a 40-item assessment of desired surgical information and the State-Trait Anxiety Inventory for Children. Parents completed a measure assessing their child's temperament (Emotionality, Activity, Sociability, and Impulsivity Survey) and the State-Trait Anxiety Inventory. The vast majority of children had a desire for comprehensive information about their surgery, including information about pain and anaesthesia, and procedural information and information about potential complications. The most highly endorsed items by children involved information about pain, including whether they would experience pain, how long it would last, and how bad it would be. Children who were more anxious endorsed a stronger desire for pain information and lesser tendency to avoid information. Younger children wanted to know what the perioperative environment would look like more than adolescent children. There were no significant correlations among child age, gender, and temperament on desire for information. Interestingly, children with a history of surgery did not require less perioperative information as compared with children who never had surgery.

5.1.9 Parental presence in anaesthesia induction

Permitting parental presence during anaesthesia induction varies widely between and within hospitals and countries¹¹⁰ and is surrounded in controversy. While parental presence is routine in some hospitals and actively discouraged in others, in many cases it is based on parental advocacy balanced with the preference of individual anaesthetists carrying out the induction. Supporters of parental presence during induction of anaesthesia argue that the trauma of separation is avoided, it increases child cooperation, minimises the need for premedication, decreases the child's anxiety during induction, facilitates the long term behavioural sequelae of surgery and enhances parental satisfaction. Arguments against parental presence include the potentially unpredictable response of the parent to the situation, increased parental anxiety and distress levels, the logistics of moving parents in and out of the induction area, the extra stress on the anaesthetist due to the presence of an emotionally involved observer, and potential legal ramifications of having a parent present^{32,74,87,122,202,238}.

The question of whether parents should stay with their child during a medical procedure has been empirically studied in many contexts apart from induction of anaesthesia including venipuncture and immunization, dental procedures, burn debridement, lumbar puncture, bone marrow aspirations and minor emergency procedures. In all of these contexts empirical evidence is inconclusive.

- Three studies have focused on parental presence during anaesthesia induction in relation to parents' anxiety. In a prospective study, Bevan et al.²⁶ examined parents of children aged 2-10 years (ASA physical status I or II) undergoing ear, nose and throat, plastic, dental, eye, or urologic surgery. Of the 134 children enrolled in the study, 67 had parents present during induction (treatment group) and 67 did not (control group). Group assignment was determined by day of surgery. Parents' in-hospital anxiety was assessed in the reception and induction areas with the VAS, a 100 mm linear scale ranging from 0 to 100 ("no fear" to "great anxiety"). Parents in the treatment group had a mean VAS score of 42.8 \pm 32.2 in the reception area compared to 41.9 \pm 28.9 in the control group. In the induction area, the treatment group had a mean VAS score of 54.1 ± 36.4 compared to 52.3 ± 33.1 in the control group. Neither of these between-group differences were significant. Subgroups of "calm" and "anxious" parents were identified by a median split of their preoperative VAS scores. Children in the "calm treatment" "calm control" and "anxious control" subgroups were similarly upset at induction. Children in the "anxious-treatment" subgroup were the most disturbed at induction and significantly more than those in the "anxious control" subgroup. Preoperative parental anxiety levels also correlated with the child's fears (measured with the Hospital Fears Inventory 197) and behaviours (measured with the Behavioural Questionnaire ²²³) one week after surgery.
- Blesch and Fisher²⁸ carried out a RCT of parents of children aged 10 years or younger undergoing elective myringotomy with tube insertion, tonsillectomy, and/or adenoidectomy. Of the 75 parents in the study, based on the week that their children were scheduled for surgery, 41 were assigned to be present for induction (treatment group) and 34 were not (control group). Parents' blood pressure and pulse rates were obtained as measures of anxiety at the following intervals: after consenting to the study, after separation from their children, and before discharge. The state scale of the State-Trait Anxiety Inventory (STAI) was used to measure parents' subjective anxiety. After consent, the treatment group's mean blood pressure was 115/76 ± 13.7/9.8 mmHg compared to 112/72 ±

13.4/8.8 mmHg in the control group. After consent, the treatment group's mean pulse rate was 77 \pm 10.2/min compared to 73 \pm 10.5/min in the control group. After separation from children, the treatment group's mean blood pressure and pulse rate were $132/78 \pm 19/10.9$ mmHg and 81 ± 12.7 /min, respectively, compared to $125/80 \pm 15.4/11.5$ mmHg and 75 ± 14.9 /min, respectively, in the control group. Before discharge, the treatment group's mean blood pressure was $118/73 \pm 12.8/11$ mmHg compared to $110/71 \pm 9.2/7.9$ mmHg in the control group. Before discharge, the treatment group's mean pulse rate was 73 \pm 7.3/min compared to $74 \pm 12.6/\text{min}$ in the control group. The only significant differences found between the treatment and control groups were between time after consent and time after separation from their children mean diastolic blood pressures (-2.49 ± 10.63 vs. -8.24 ± 11.01 , respectively; P = 0.025) and time after separation from their children and time before discharge mean pulse rates $(7.66 \pm 10.30 \text{ vs. } 2.00 \pm 9.07, \text{ respectively; P} = 0.016)$. Subjective anxiety was not significantly different between the treatment and control group (39.05 \pm 11.53 vs. 44.61 \pm 14.51, respectively; P = 0.077).

- In a RCT Palermo et al. 175 assessed parents of infants aged one to 12 months (ASA class I and II), undergoing outpatient surgery. Of the 73 parents in the study, 37 were present during induction and 36 were not. Parental anxiety was measured with the STAI before and after surgery. There were no significant differences in anxiety between the two groups. Before surgery, parents of accompanied children had a mean STAI score of 57.6 ± 5.4 compared to 56.9 ± 6.4 for parents of unaccompanied children. After surgery, parents of accompanied children had a mean STAI score of 47.2 ± 4.8 compared to 45.2 ± 5.2 for parents of unaccompanied children. Interestingly, parents who were present during induction demonstrated comparable health care attitudes (measured with the Health Care Attitudes Questionnaire 85) before and after surgery, as well as comparable levels of satisfaction with the surgical experience (measured with a modified version of the Perception of Procedures Questionnaire 126) compared to parents who were absent during induction.
- Four studies have examined parental presence during anaesthesia induction in relation to children's anxiety. Hickmott et al.⁹² undertook a RCT of children aged 1-9 years undergoing general anaesthesia for minor elective surgery. Of 49 children in the study, 26 had their mothers present during induction and 23 did not. Allocation to each group was determined by the week in which the children's surgery took place. A recovery room or ward nurse, not involved in the anaesthetic procedure, was responsible for observing and measuring children's anxiety levels in the anaesthesia room. Time in the anaesthesia room was separated into the 'waiting period' (time from the children's arrival until the anaesthetist arrived) and the 'induction period' (time from the anaesthetist's arrival). Children's anxiety was measured using a pre-determined scale ranging from 0 (no anxiety) to 2 (marked anxiety) during the waiting period and 0 (calm) to 4 (screaming and uncontrollable) during the induction period. During the waiting period in the mother-present group, five children scored 0 and two children scored 2; whereas, in the mother-absent group, seven children scored 0 and one each scored 1 and 2. During the induction period in the mother-present group, 13 children scored 0, nine scored 1, and two each scored 2 and 3; whereas, in the mother-absent group, 15 children scored 0, four scored 1, three scored 2, and one scored 3. Children's anxiety levels did not differ significantly

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between the two groups during either the waiting or the induction period (Mann– Whitney U test).

- In a RCT, Amanor-Boadu¹³ assessed 118 children aged 1–12 years undergoing inguinal surgery as day cases. Children undergoing surgery were randomly assigned to be accompanied or unaccompanied. Of the 118 children in the study, 52 were accompanied by a parent and 66 were not. Children were evaluated according to their age group, i.e., aged 5 years or less and more than 5 years. Heart rates using a stethoscope were taken both on the ward and before induction as a measure of anxiety. For children 5 years or less, unaccompanied children had a mean heart rate of 109 ± 13 /min on the ward compared to 111 \pm 12/min for accompanied children. For children more than 5 years, unaccompanied children had a mean heart rate of $101 \pm 11/\text{min}$ on the ward compared to 100 ± 10 /min for accompanied children. These two differences were not significant. Mean heart rates before induction, for children 5 years or less, was 128 ± 20 /min for unaccompanied children compared to 118 ± 16 /min for accompanied children. For children more than 5 years, it was $108 \pm 10/\mathrm{min}$ for unaccompanied children compared to 97 ± 19 /min for accompanied children. Both of these differences were significant at P = 0.001.
- In a retrospective study using a multiple matched concurrent cohort, Kain et al. 105 examined children's anxiety in relation to parents'. The participants were selected from a database of children from a number of previous prospective and randomized studies that the authors conducted comparing parental presence with no parental presence. Of the 568 children included in the study (aged 2–12 years undergoing general anaesthesia for elective outpatient surgery), 284 had their parent present during induction and 284 did not. For children, anxiety was measured with the modified Yale Preoperative Anxiety Scale (mYPAS) and children were categorized as "anxious" if they scored >40 on the mYPAS, and as "calm" if they scored <30 on the mYPAS. For parents, anxiety was measured with the STAI and parents were categorized as "anxious" if they scored in the upper 50% on the STAI, and as "calm" if they scored in the lower 50% on the STAI. Four groups of child-parent pairs were then retrospectively compared for the parent-present and parent-absent groups: calm parent-calm child, anxious parent-calm child, calm parent-anxious child, and anxious parent-anxious child. Anxious children with calm parents present were significantly less anxious during induction than anxious children with no calm parents present (mean mYPAS = 51.9 ± 24 vs. 64.6 ± 26 , respectively; P = 0.03). Calm children with anxious parents present were significantly more anxious during induction than calm children with no anxious parents present (mean mYPAS = 52.4 ± 28 vs. $39.4 \pm$ 21, respectively; P = 0.002). On the other hand, there was no significant difference in anxiety during induction between calm children with calm parents present and calm children with no calm parents present (mean mYPAS = 39.9 \pm 22 vs. 34.7 ± 20 , respectively; P = 0.150), and no significant difference in anxiety during induction between anxious children with anxious parents present and anxious children with no anxious parents present (mean mYPAS = 71.0 ± 23 vs. 66.6 \pm 27, respectively; P = 0.490). The authors concluded that the presence of a calm parent does benefit an anxious child during induction of anaesthesia and the presence of an overly anxious parent has no benefit.
- In a RCT, Patel et al.¹⁷⁷ examined 112 children aged 4–12 years undergoing outpatient surgery. Children's change in anxiety was assessed from baseline to introduction of the anaesthesia mask using the mYPAS. Children were randomly

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assigned to one of three groups using sealed envelopes: parental presence (n = 36), parental presence plus 0.5 mg/kg oral midazolam (n = 38), or parental presence plus a hand-held video game (n = 38). Children who received parental presence plus a hand-held video game experienced a statistically significant decrease in anxiety from baseline to introduction of the anaesthesia mask compared to children who received parental presence alone (median change in mYPAS = -3.3 vs. + 11.8, respectively; P = 0.04). Children who received parental presence plus midazolam did not experience a statistically significant change in anxiety from baseline to introduction of the anaesthesia mask compared to the other two groups (median change in mYPAS = +7.3).

- Seven studies examined both parents' and children's anxiety in relation to parental presence during anaesthesia induction. Johnston et al. 101 carried out a prospective study of parents and their children aged 2-8 years undergoing day surgery. Of the 134 children in the study, 67 had their parent present and 67 did not. Parents and children were assigned to each group based on the day of the week that surgery was scheduled. Anxiety was measured before induction. For parents, the VAS, a 10 cm line ranging from 0 ("no anxiety") to 10 ("most anxiety") was used to measure anxiety. For children, the Global Mood Scale (GMS), an observation scale ranging from 1 (child attentive and happily active) to 7 (child screaming), was used. Overall, there were no differences in parents' or children's anxiety between parent-present and parent-absent groups. To conduct further analysis, the authors separated parents into low-anxiety and high-anxiety groups based on their VAS scores; i.e., those who scored ≤3 on the VAS were considered low-anxiety, and those who scored ≥6 on the VAS were considered high-anxiety. The authors found that high-anxiety parents who were present for induction were more anxious than high-anxiety parents who were not present for induction. Low-anxiety parents who were present for induction were less anxious than low-anxiety parents who were not present for induction. Children with highanxiety parents who were present were more anxious than children with highanxiety parents who were not present. Children with low-anxiety parents experienced the same level of anxiety whether they were in the parent-present or parent-absent group.
- In a non randomised prospective study Cameron et al.³⁵ assessed 74 parents and their children aged 1-8 years undergoing day surgery. Parents were only allowed to be present for induction if the anaesthetist carrying out the induction granted them permission. The treatment group consisted of 38 parents who were granted permission and decided to be present. The control group consisted of 36 parents who were either not permitted or decided not to be present. In the control group, 22 parents chose to separate from their children in the theatre holding bay area and 14 parents chose to separate from their children in the day surgery ward. Parents' anxiety was measured immediately upon separation from their children using a VAS with scores ranging from 1 ("no anxiety at all") to 10 ("most anxiety anyone could have"). A five-point scale with scores ranging from 1 (cheerful and attentive) to 5 (very distressed and uncontrollable) was used by parents to assess their children's anxiety right before separation from them. Parents in the treatment group were significantly less anxious, as measured by the VAS, than parents in the control group (mean = 3.4 ± 1.6 vs. 6.5 ± 2.2 , respectively; P < 0.001). Parents who were present for induction reported their children to be significantly less anxious than parents who were not present for induction (mean = 1.9 \pm 1.1 vs. 2.8 \pm 1.1, respectively; P < 0.001).

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In a RCT, Kain et al.¹¹⁵ examined parents and their children aged 1–6 years undergoing general anaesthesia for elective outpatient surgery. Of the 84 children in the study, using a random numbers table generated by a computer, 43 were randomised to have their parent present during induction (intervention group) and 41 did not (control group). For children, anxiety was measured with the Yale Preoperative Anxiety Scale (YPAS), Clinical Anxiety Rating Scale (CARS), VAS, and cortisol. For parents, anxiety was measured with the STAI, VAS, heart rates, and blood pressure. The VAS, a 100-mm line ranging from 0 ("not anxious") to 100 ("extremely anxious"), was used as an observational measure for children and a self-report measure for parents. Using these measures, no significant differences were found between the two groups for either children's or parents' anxiety. For children, anxiety was reported as medians and 25–75% interquartile ranges for the holding area, induction 1 (entering the induction room), and/or induction 2 (introduction of anaesthesia mask). On the VAS, children in the control group compared to those in the intervention group scored the following: holding area = 11 (0-28) vs. 6 (0-33), respectively; induction 1 =38 (0-89) vs. 37 (0-82), respectively; and induction 2 = 43 (5-78) vs. 45 (8-86), respectively. On the YPAS, children in the control group compared to those in the intervention group scored the following: induction 1 = 34 (24-41) vs. 30 (25-41), respectively, and induction 2 = 38 (24-65) vs. 42 (30-62), respectively. On the CARS, children in the control group compared to those in the intervention group scored the following: induction 1 = 0 (0-1) vs. 0 (0-1), respectively, and induction 2 = 1 (0-4) vs. 1 (0-4), respectively. With respect to cortisol ($\mu g/mL$) for induction 2, the results for children in the control group compared to those in the intervention group were 73 (51–100) vs. 76 (48–91), respectively. For parents, anxiety was reported as means and standard deviations or as medians and 25–75% interquartile ranges for the holding area and/or post-induction (after parents left their children). State-Trait Anxiety Inventory scores for the control and intervention group parents were 46 \pm 12 vs. 43 \pm 12, respectively, post-induction. Visual analogue scale scores for the control group parents compared to the intervention group parents were 43 (20–58) vs. 38 (13–49), respectively, in the holding area and 49 (18-73) vs. 41 (5-66), respectively, post-induction. Systolic blood pressure (mmHg) for the control group parents compared to the intervention group parents was 114 ± 11 vs. 116 ± 17 , respectively, in the holding area and 122 \pm 12 vs. 121 \pm 13, respectively, postinduction. Diastolic blood pressure (mmHg) for the control group parents compared to the intervention group parents was 71 \pm 8 vs. 67 \pm 10, respectively, in the holding area and 77 \pm 9 vs. 75 \pm 7, respectively, postinduction. Heart rates (beats min-1) for the control group parents compared to the intervention group parents were 81 ± 9 vs. 78 ± 8 , respectively, in the holding area and 85 ± 10 vs. 84 ± 8 , respectively, post-induction. The authors concluded that only children who were older than 4 years, had a parent with a low trait anxiety level or a low baseline level of activity as assessed by temperament ratings benefited from parental presence during induction of anaesthesia. In contrast, there was a trend among children younger than 4 years to be more anxious during induction in the presence of their parent.

• Kain et al.¹¹⁷ in a RCT examined 88 parents and their children aged 2–8 years undergoing general anaesthesia for elective outpatient surgery. The children were randomized into one of three groups according to a random numbers table: (a) parental presence (n = 29), (b) premedication with 0.5 mg/kg oral midazolam mixed in 10 mg/kg acetaminophine syrup at least 20 min before surgery (n = 33), (c) no parental presence and no sedative premedication (n =

- 26). Anxiety was measured for parents with the STAI and for children with the Procedural Behavior Rating Scale (PBRS¹²⁵). There were no significant differences between the three groups regarding children's anxiety in the preoperative holding area. Upon separation from their parents, children in the midazolam group were significantly less anxious than children in the other two groups [PBRS = 0 (0–1) vs. 4 (0–5); P = 0.02]. Children in the midazolam group were also significantly less anxious than children in the other two groups at both entrance to the operating room (P = 0.0171) and introduction of the anesthesia mask (P = 0.0176). Parents in the midazolam group were significantly less anxious after separation than parents in the parental presence group and parents in the control group (mean STAI score = 43 ± 12 vs. 50 ± 10 vs. 47 ± 10 , respectively; P = 0.048). The percentage of inductions in which compliance of the child was poor was significantly greater in the control group compared with the parental presence and midazolam groups (25% vs. 17% vs 0%, P= 0.013)
- Kain et al.¹¹⁸ in a RCT assessed 103 parents and their children aged 2–8 years. Parents and their children were randomly assigned to each group using a random numbers table. The intervention group had parental presence and received premedication with oral midazolam syrup (0.5 mg/kg) at least 20 min before surgery. The control group received premedication with oral midazolam syrup (0.5 mg/kg) at least 20 min before surgery only. Anxiety was measured for children with the mYPAS and for parents with the STAL Children's anxiety was not significantly different between the two study groups (P = 0.49). Parents' anxiety, on the other hand, was significantly lower after separation for those who were present compared to those who were not present (mean = 43 ± 11 vs. 48 ± 12, respectively; P = 0.037). Parental satisfaction with the overall care provided and with the separation process was significantly higher among the premedication and parental presence group compared with the premedication only group.
- Kain et al.¹⁰⁶ undertook a RCT of parents and their children undergoing general anaesthesia and elective outpatient surgery. Of the 80 children in the study, 29 had their parent present, 27 had their parent present and received oral midazolam (0.5 mg/kg) about 30 min before induction, and 24 did not have their parent present (control group). They were randomly assigned to the three groups based on a random number table. For children, anxiety was measured with the mYPAS and for parents with the STAI. Heart rates, skin conductance levels (SCL), and blood pressure levels were also used to measure parents' anxiety. Children in parental presence plus midazolam group were less anxious than children in either the control group or the parental presence only group (P =0.023). At different time points, parents in both parental presence groups had higher anxiety, as measured by heart rates, than the control group (P < 0.05). However, there was no significant difference in heart rates between the parental presence and parental presence plus midazolam groups. Skin conductance level was higher in the two parental presence groups than in the control group (P < 0.05). However, there was no significant difference in SCL between the two parental presence groups. The SCLs were not provided by the authors. There were no significant differences between the parental presence, parental presence plus midazolam, and control groups with regards to systolic blood pressure (123 \pm 21 vs. 128 \pm 16 vs. 126 \pm 19, respectively; P = 0.59) and diastolic blood pressure (82 \pm 14 vs. 85 \pm 13 vs. 81 \pm 15, respectively; P = 0.88) after induction. In addition, there were no significant differences in parents'

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48 49 50 self-reported anxiety, as measured by the STAI, between the three groups (STAI scores and P-values were not provided).

- Kain et al.¹⁰⁸ undertook a prospective study of parents and their children (mean age = 4.9 years) who were part of a previous investigation by the authors at their initial surgery and were undergoing a subsequent surgery. At their initial surgery, the children had been assigned to the following preoperative intervention: parental presence (n = 27), oral midazolam (n = 13), parental presence plus oral midazolam (n = 10), and no intervention (n = 33). The authors allowed parents to choose their preoperative intervention group at the subsequent surgery. The parents of the 83 children in the study chose the following preoperative intervention: parental presence (n = 46), oral midazolam (n = 8), parental presence plus oral midazolam (n = 21), and no intervention (n = 21)8). Anxiety was measured for children with the mYPAS and for parents with the STAI. There were no significant differences between the groups regarding children's anxiety upon entering the operating room [median mYPAS score (range): parental presence = 45.8 (22.9–91.7), oral midazolam = 54.2 (22.9– 95.8), parental presence plus oral midazolam = 35.4 (22.9–100.0), and no intervention = 23.2 (22.9-45.8); P = 0.31] or during induction [median mYPAS score (range): parental presence = 45.8 (22.9-100.0), oral midazolam = 65.5(22.9–95.8), parental presence plus oral midazolam = 34.2 (22.9–100.0), and no intervention = 24.5 (22.9-50.0); P = 0.15]. There was also no significant difference in parents' anxiety at separation (mean STAI score: parental presence = 42.8 \pm 11.1, oral midazolam = 49 \pm 6.5, parental presence plus oral midazolam = 43.3 ± 13.0 , and no intervention = 37.8 ± 6.5 ; P = 0.28). Children in the midazolam group experienced significantly higher anxiety in the preoperative holding area than children in the other groups [median mYPAS score (range): parental presence = 23.3 (23.3-70.0), oral midazolam = 37.5(23.3-68.8), parental presence plus oral midazolam = 45.8 (23.3-96.7), and no intervention = 23.3 (23.3-55.0); P = 0.03]. Parents of children in the midazolam group were also significantly more anxious than parents of children in the other groups in the preoperative holding area (mean STAI score: parental presence = 38.6 ± 9.1 , oral midazolam = 47.3 ± 8.4 , parental presence plus oral midazolam = 42.5 ± 12.2 , and no intervention = 36.8 ± 5.1 ; P = 0.09). Interestingly, of parents whose children received parental presence at the initial surgery, 70% chose to be present during induction again. In contrast only 23% of the patients who received midazolam at the initial surgery requested midazolam at the subsequent surgery and only 15% of the patients who received no intervention at the initial surgery requested no intervention at the subsequent surgery. Parents' intervention preferences at the subsequent surgery were influenced by children's anxiety at the initial surgery.
- Arai et al., 17 in 22 pairs of mothers and children (1-3 years old) scheduled for minor plastic surgery under general anaesthesia found that higher parental anxiety pre-surgery, as indicated by higher amounts of maternal salivary amylase activity, was significantly correlated higher children's anxiety during induction (r_s = -0.667, P < 0.0001) and severer children's emergence agitation (r_s = 0.705, P< 0.0001). Both children's anxiety and agitation were rated by a blind observer.
- In another study¹⁶ the same authors randomised, using computer-generated random numbers, 58 children, aged 1-3 years, classified as ASA I, undergoing minor plastic surgery under general anaesthesia to one of three groups: (a) a

sedative group (0.5 mg/kg oral midazolam) (n= 19), (b) parental presence (20), (c) a sedative and parental presence (19). Children in the midazolam group showed a better quality of mask induction compared with those on the parental presence group but the addition of parental presence to oral midazolam did not provide additional improvement of mask induction. In contrast, the children in the midazolam and parental presence group were less agitated than those in the other groups at emergence from anaesthesia.

• A recent meta-analysis²³⁵ that assessed the effects of non-pharmacological interventions in assisting induction of anaesthesia in children by reducing their anxiety, distress or increasing their co-operation concluded that the presence of parents during induction of general anaesthesia does not reduce their child's anxiety. However, the authors commented further that calm parents may be helpful and parental presence should be considered on an individual basis.

Taken in combination the results of the above randomised studies point that current evidence shows that there is no apparent benefit of parental presence during anaesthesia induction in relation to decreasing parents' and children's anxiety ³⁹. In many cases, midazolam or distraction techniques appear to be a suitable substitute. Overall, positive effects for parental presence, including lower levels of child anxiety and distress, have been reported in studies in which parents were not randomly assigned to condition but were permitted to self-select presence or absence. In terms of child characteristics, a prospective cohort study has demonstrated that children who benefit from parental presence are older, have lower levels of activity in their temperament, and have parents who are calmer and who value preparation and coping skills for medical situations¹¹³.

5.1.10 Parental presence during medical procedures

Piira et al.¹⁸¹ conducted a systematic review, of controlled studies investigating parental presence in the paediatric treatment room at the time of their child's medical procedure. A total of 28 studies met inclusion criteria which were as follows: the studies evaluated the effects of parental presence on child, parent or health professional outcomes; concurrent control groups were used; only primary data were used to avoid bias resulting from the use of duplicate results. The age of the children participating in the studies ranged from 2 weeks to 18 years. 1256 children had a parent present and 1025 children did not have a parent present. The medical experiences included routine immunizations, venipunctures, dental procedures, lumbar punctures, burns treatments, intubation, central line placement, chest tube placement and anaesthesia induction, with some studies including a number of different painful contexts. There were mixed findings regarding the effect of parental presence on measures of child distress and affect, however, studies of lower levels of evidence were more likely to report significant results. Parents who were present during their child's medical intervention were either better off or no different from parents who were absent with regard to their levels of distress and satisfaction. There was no evidence of increased technical complications nor elevated staff anxiety for health professionals attending to children with a parent present as compared to attending to children without their parents.

5.1.11 The role of the parents during medical procedures and/or anaesthesia induction

In the paediatric pain literature a number of studies point to the role that parents play in shaping their child's pain perception and distress response. Certain parental behaviours

are associated with child coping and others with child distress when children undergo painful medical procedures. Parenting behaviours such as agitation, provision of reassurance, empathic comments, giving control, excessive explanations and apologies to their children have been shown to be associated with (and indeed precede) elevated distress and increased pain intensity during medical procedures^{30,31,48}. Humor, commands to use coping strategies, and non procedural talk are associated with increases in child's coping. Dahlquist and colleagues⁴⁷ demonstrated the influence of speech function on pain distress. Their results showed that vague commands by caregivers were positively associated with child distress during painful procedures. Liossi and colleagues¹⁴⁴ showed that parental expectancies are highly predictive of experienced pain in children undergoing lumbar punctures.

Parents are often anxious not only about their child's distress but also about their own ability to support and comfort their child through a painful experience. Thus, parents need to be included in interventions and helped to control their own anxiety which in turn will ensure less anxiety being communicated to the child. Simple educational leaflets can give useful information and more extensive training programmes can teach parents what to do¹⁸³.

5.1.12 Summary - Preparation for sedation

In summary, current evidence from the broader preparation literature i.e., preparation for anaesthesia and medical procedures suggests:

- that preparation for sedation is important for young people and their parents
- there is some helpful direction informing what this should and should not include and how it is performed

For children, the extensiveness and style of preparation should be guided by each child's age and developmental level (see Table 1 which provides recommendations about the timing and content of preparation according to children's developmental level).

In general, specific discussion about the sedation and procedure has more relevance for children >2 years of age. The outcome from this narrative review suggests that preparation should have at least 3 components, namely:

- what will happen (where, how long it will last, and what will be done)
- how it will feel (pressure, temperature, and level of discomfort to be expected)
 - strategies to cope with the stressor (which may be related to the sedation technique and/or procedure 57,180,208.

Given this, children can be asked what strategies they think will help them to cope and, if possible, those strategies should be incorporated into the sedation administration. In addition, given the strong data supporting distraction, distraction techniques should be used during the induction of sedation. Evidence supporting the use of behavioural strategies such as teaching children coping techniques to alleviate their preoperative anxiety has emerged throughout the literature²³⁴. Teaching children coping skills allows

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them to learn how to calm themselves in times of stress and thus may be useful not just at the time of the procedure in question but at subsequent procedures as well.

For parents, there is inconclusive evidence indicating whether parents should be encouraged or discouraged to be present at their child's induction. The offer to be present is therefore based on negotiation with the care team. Although parental presence may not have a clear, direct influence on child distress and behavioural outcomes, there are potential advantages for parents and children, offering the option of parental presence is clearly in line with a paradigm shift to family centred care during hospitalization¹¹¹. Parental inclusion in supporting interventions may also help their own anxiety, lessening the potential for this to be communicated to their child.

6 Drugs for sedation in infants, children and

young people

3	6 1	General	clinical	introduction:	drugs for	sedation i	n infants	children and
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- The Guideline Development Group (GDG) considered that many potentially useful sedation drugs could be reviewed. The GDG decided to limit the literature searches and discussions to sedative drugs that were both currently available and in common use in the UK. All commonly used routes of administration of the chosen drugs, for example by injection, by mouth or by inhalation, were considered.
- The GDG was mindful of the fact that some classes of sedative drugs may be used for analgesia, pre-operative or pre-induction medication and in some situations, may cause general anesthesia. Evidence for sedation was considered only if the studies reviewed specifically intended to assess the sedative effects of the drug. The GDG made a judgment on whether the doses used were likely to cause anesthesia.
- 15 The GDG reviewed evidence on the following sedative drugs:
- Midazolam: Oral, IV, rectal, transmucosal
- Ketamine: IV, IM
- Chloral Hydrate: Oral
- Triclofos sodium: Oral
- Nitrous oxide: Inhalation
- Sevoflurane: Inhalation
- 22 Propofol: IV
- Opioids: IV Fentanyl, IV Morphine and intranasal (IN) Diamorphine
- 24 Midazolam is a short acting benzodiazepine with a short half life. It has potent 25 anxiolytic, amnestic, hypnotic, anticonvulsant, skeletal muscle relaxant, and sedative

1 2	properties. It can be administered by several different routes and is often given in combination with other sedative agents.
3 4 5 6 7	Ketamine is an N-methyl d-aspartate (NMDA) receptor antagonist which causes a trance- like sedation with few appreciable effects on the respiratory and cardiovascular systems. Its analgesic effect is a major advantage. Administered intravenously it can be titrated. A single intramuscular dose is predictable and effective whenever venous access is impractical.
8 9 10 11	Chloral hydrate was the first synthetic drug employed for its sedative-hypnotic effect. Unlike opioids, it produces sedation without significant adverse effects on cardiovascular or respiratory function at therapeutic doses. In children it is orally administered for painless imaging.
12 13 14	Nitrous oxide gas, delivered with oxygen, also acts as an NMDA receptor antagonist. It has a rapid anxiolytic/sedative/analgesic effect and is delivered by inhalation. Doses may be titrated to achieve target effect.
15 16 17 18 19 20	Opioid drugs can be used as sedatives for painful procedures however it is important to separate the use of opioids used as sedation from when they are used specifically for analgesia alone. Intravenous morphine and fentanyl are commonly used opioids whose sedative action can be improved by the addition of another sedative such as midazolam. Intranasal diamorphine has been considered in the review because it has the potential to be rapidly effective and easily administered.
21 22 23 24	Propofol is a short acting hypnotic agent that can be given in low doses to achieve short acting and controlled sedation. Propofol is not considered an analgesic, so opioids such as fentanyl may be combined with propofol to alleviate pain. Propofol is administered intravenously.
25 26	Sevoflurane is a fluorinated isopropyl ether which has a rapid induction and quick elimination effect. It is delivered by inhalation and may be titrated for sedative effect.
27 28	Triclofos is a sedative-hypnotic drug, similar to chloral hydrate but with less gastric irritation. It is orally administered for painless imaging.
29 30	In some settings, the use of local anaesthesia was included because the effect of analgesia is likely to be crucial to the success of any sedation for painful procedures.
31	The GDG reviewed evidence on sedative drugs with the following comparisons:
32	Placebo; non-pharmacological treatment
33	Head to head
34	Combination (including analgesia and general anaesthesia)
35	Route of administration
36	• Dose
37 38	In some settings, the use of local anaesthesia was included because the effect of analgesia is likely to be crucial to the success of any sedation for painful procedures.

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1 In general, for the purposes of categorisation of RCTs, a drug combination is defined as 2 two or more drugs that have sedative potential. In some RCTs, single sedation drugs have 3 been combined with interventions that do not cause sedation such as local anaesthesia, 4 mild analgesics (such as paracetamol) or a non-pharmacological intervention. For the 5 purposes of categorisation of the RCTs, these additional interventions are not considered 6 to be part of a sedation drug combination when they have been applied equally to both 7 groups. For example in a RCT in which one group receives sedation drug A and the other 8 has sedation drug B, but both groups receive local anaesthesia, the RCT is categorised as 9 a single drug comparison. However if local anaesthesia had been used only in one group 10 the RCT would be categorised as a comparison of a drug combination. 11 6.2 General methodological introduction: drugs for sedation in infants, 12 children and young people

Efficacy outcome data for this review was taken from RCTs alone. Each outcome was quality assessed using a GRADE evidence profile. The outcome measures for drug efficacy that were considered by the GDG were as follows:

Primary outcome:

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- Successful completion of diagnostic or therapeutic procedure
 - measured as the number of patients for whom the diagnostic or therapeutic procedure was carried out and completed.

Secondary outcomes:

- Behavioural ratings including:
 - pain as assessed by the patient or parent or other observer using validated pain scales for example Visual Analogue Scale (VAS),
 Children's Hospital of Eastern Ontario Pain Scale (CHEOPS), Faces Pain Scale (FPS)...
 - procedural distress and/or anxiety as assessed by the patient or parent or other observer using validated scales e.g. Visual Analogue Scale (VAS), Observation Scale of Behavioral Distress (OSBD).
 - o patient or parent satisfaction including preference
- Sedation timing including
 - length of induction: time from administration of sedation drug to initiation of procedure
 - recovery: time from completion of procedure to recovery criteria being met or recovery to pre-sedation state
 - o duration of procedure
 - o total: time from administration of intervention to when patient has been transferred to the recovery area or has been discharged

Evidence of safety was sought from both RCTs and non RCT observational studies. Each RCT was quality assessed using a GRADE evidence profile. The GDG recognised that research from non RCT observational studies is subject to the usual limitations of observational work, including dependence on the quality of medical record documentation and potential for bias secondary to non randomisation, and un-blinded participants. In these studies, there were no interventions or comparisons but merely data collection of adverse events. The datasets were generally large, and were expected to provide more information on a range of adverse events than the small RCTs available for review. Due to these limitations, we only assigned quality rating ('very low' quality) based on the GRADE scheme. It was considered more comprehensive to present separately this supplementary observational data in the form of concise, customised summary tables which also contain the GRADE ratings.

The outcomes measures for safety were limited to short term effects. Long term effects of sedation drugs were considered to be too rare for inclusion in this review. The outcome measures for drug safety and adverse effects that were considered by the GDG are as follows:

- Aspiration
- Respiratory intervention, including:
 - o oral-pharyngeal airway
 - o endotracheal intubation
 - assisted ventilation
- Cardiac arrest requiring either/or:
 - o external cardiac massage
 - o defibrillation
- Oxygen desaturation <90%
- Vomiting

A decrease in oxygen saturation to below 90% was chosen as a safety outcome because saturations between 90 and 95% are commonplace during recovery from anaesthesia, especially if supplemental oxygen is not administered; saturations above 90% are not necessary evidence of an appreciable adverse event. The GDG agreed however that desaturation less than 90% is concerning.

The GDG agreed that the dose of drugs was an important consideration. Matching the dose to the target sedation level is essential and when robust data has been published, it has been quoted. Yet the dose *question* is not straightforward. When a drug is given by mouth, only a single dose is practical because its absorption, and therefore its maximum effect, can take a variable time. In contrast, intravenous drugs can be titrated to achieve the target level of sedation although it must be appreciated that there is considerable variation and the practitioner will need to continually assess the conscious level and adjust the dose accordingly. Prolonged recovery is a hazard that can be avoided if the lengths of action of the sedation drugs match the length of the procedure. This is a

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example after a dental extraction, because the sedation is no longer opposed by the stimulation of painful procedure.	1	notoriously dangerous problem following painful procedures when pain has subsided, for
3 stimulation of painful procedure.	2	example after a dental extraction, because the sedation is no longer opposed by the
	3	stimulation of painful procedure.

1 6.3 Midazolam

Matrix of midazolam comparators

Key:

Chloral hydrate = CH

Fentanyl = F

Morphine = Mo

Meperidine = Me

lsoflurane = 1

Ketamine = K

 $Local\ anaesthesia = LA$

Topical anaesthesia = TA

Midazolam = M

Nitrous oxide = N_20

Nitrous oxide and oxygen = N_20+02

Opioids = 0

Propofol= P

 ${\sf Sevoflurane} = {\sf S}$

Triclofos sodium = TS

Midazolam vs.

	Reference	Tables	Evidence statements page
Placebo			
	Liacouras, 1998 ¹³⁹ Mortazavi, 2009 ¹⁶⁸	Table 3	158
	Fatovich 1995 ⁶³ , Luhman 2001 ¹⁵¹	Table 4	162 159
	Kapur 2004 ¹²⁴	Table 5	159
	Fishbein 1997 ⁶⁷	Table 6	162
	Ljungman 2000 ¹⁴⁸ Theroux 1993 ²¹⁰	Table 7	159 160
Head to head			
M vs TS	Singh 2002 ²⁰³	Table 8	160
M vs CH	Layangool 2008 ¹³⁵	Table 9	160
M + non-pahrma vs N ₂ 0 + pharma	Zier 2008 ²³⁷	Table 10	161
Combinations			
M vs M + N ₂ 0+02	Al-zahrani 2009 ¹²	Table 11	161
M + N20 vs N20	Luhman 2001 ¹⁵¹	Table 12	162

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Description	Г		T =	1
M + Morphine vs	M + P vs P			
Propofol + Morphine		Disma 2005 ⁵⁶	fable 14	162
Propofol + Morphine	M + Morphine vs	Havel 1999 ⁹⁰	Table 16	163
Meperidine Memoral M				
Meperidine Memoral M	M + Meperidine vs	Fishbein 1997 ⁶⁷	Table 15	162
M + Remifentanil vs Remifentanil vs Remifentanil Table 18 164	-			.02
Remifentanil	M + F vs F	Antmen 2005 ¹⁵	Table 17	163
Page		Antmen 2005 ¹⁵	Table 18	164
Dilli 2008 ⁵⁵ Dilli 2008 ⁵⁵	M + K vs K +	Sherwin 2000 ²⁰¹	Table 19	164
RCTs	placebo			
Luhmann 2001 151 Liungman 2000 148 Layangool 2008 135 Zier 2008 2005 168	Safety			
Ljungman 2000148 Layangool 2008135 Zier 2008237 Disma 200556 Havel 199990 Antmen 200515 Wathen 2000227 Sherwin 2000201 Dilli 200954 Aspiration	RCTs			
Layangool 2008135 Zier 2008237 Disma 200556 Havel 1999% Antmen 200217 Sherwin 2000201 Dilli 200954			Table 25	
Zier 2008 ²³⁷ Disma 2005 ⁵⁶ Havel 1999 ⁹⁰ Antmen 2000 ²²⁷ Sherwin 2000 ²²¹ Dilli 2009 ⁵⁴				
Disma 2005 ⁵⁶ Havel 1999 ⁹⁰ Antmen 2000 ²¹⁵ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Dilli 2009 ⁵⁴				
Havel 1999% Antmen 200515 Wathen 2000227 Sherwin 2000201 Dilli 200954 Aspiration				
Antmen 2005 ¹⁵ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Dilli 2009 ⁵⁴ Aspiration Luhmann 2001 ¹⁵¹ Havel 1999 ⁹⁰ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Desaturation Liacouras 1998 ¹³⁹ Disma 2005 ⁵⁶ Havel 1999 ⁹⁰ Antmen 2005 ¹⁵ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Hartgraves 1994 ⁸⁹ Needleman 1995 ¹⁷¹ Respiratory Intervention Respiratory Intervention Respiratory Intervention Disma 2005 ⁵⁶ Havel 1999 ⁹⁰ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Hartgraves 1994 ⁸⁹ Needleman 1995 ¹⁷¹ Kanegaye 2003 ¹²³ Vomiting Luhmann 2001 ¹⁵¹ Luhmann 2001 ¹⁵¹ Table 27 Table 28 Table 28				
Sherwin 2000 ²⁰¹ Dilli 2009 ⁵⁴				
Dilli 2009 ⁵⁴				
Aspiration				
Havel 1999°0 Wathen 2000²27 Sherwin 2000²21 Table 27 Table 28		Dilli 2009 ⁵⁴		
Havel 1999°0 Wathen 2000²27 Sherwin 2000²21 Table 27 Table 28	Aspiration	Luhmann 2001 ¹⁵¹		168
Sherwin 2000 ²⁰¹	- 1			
Desaturation				
Disma 2005 ⁵⁶ Havel 1999 ⁹⁰ Antmen 2005 ¹⁵ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Hartgraves 1994 ⁸⁹ Needleman 1995 ¹⁷¹ Table 27 Table 28 Table 28		Sherwin 2000 ²⁰¹		
Disma 2005 ⁵⁶ Havel 1999 ⁹⁰ Antmen 2005 ¹⁵ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Hartgraves 1994 ⁸⁹ Needleman 1995 ¹⁷¹ Table 27 Table 28 Table 28	Desaturation	Liacouras 1998 ¹³⁹	Table 27	168
Antmen 2005 ¹⁵ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Hartgraves 1994 ⁸⁹ Needleman 1995 ¹⁷¹ Respiratory intervention Luhmann 2001 ¹⁵¹ Disma 2005 ⁵⁶ Havel 1999 ⁹⁰ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Needleman 1995 ¹⁷¹ Kanegaye 2003 ¹²³ Vomiting Luhmann 2001 ¹⁵¹ Ljungman 2000 ¹⁴⁸ Table 27 Table 28	2 course anon			
Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Hartgraves 1994 ⁸⁹ Needleman 1995 ¹⁷¹ Table 27 Table 28 Table 28		Havel 1999 ⁹⁰		
Sherwin 2000 ²⁰¹				
Hartgraves 1994 ⁸⁹ Needleman 1995 ¹⁷¹				
Needleman 1995 ¹⁷¹				
Respiratory Luhmann 2001 ¹⁵¹ Table 27 Table 28 Table 28				
Disma 2005 ⁵⁶				
Havel 1999 ⁹⁰ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Needleman 1995 ¹⁷¹ Kanegaye 2003 ¹²³ Vomiting				168
Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Needleman 1995 ¹⁷¹ Kanegaye 2003 ¹²³ Vomiting Luhmann 2001 ¹⁵¹ Table 27 Ljungman 2000 ¹⁴⁸ Table 28 168	intervention		Table 28	
Sherwin 2000 ²⁰¹				
Needleman 1995 ¹⁷¹ Kanegaye 2003 ¹²³				
Vomiting Luhmann 2001 ¹⁵¹ Table 27 168 Ljungman 2000 ¹⁴⁸ Table 28				
Ljungman 2000 ¹⁴⁸ Table 28				
Ljungman 2000 ¹⁴⁸ Table 28	Vamiting	Luhmann 2001 151	Table 27	168
	v omining			100
		Layangool 2008 ¹³⁵		

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	Zier 2008 ²³⁷ Antmen 2005 ¹⁵ Wathen 2000 ²²⁷ Sherwin 2000 ²⁰¹ Everitt 2002 ⁶⁰ Shashikiran 2006 ¹⁹⁹ Fuks 1994 ⁷¹ Needleman 1995 ¹⁷¹ Kanegaye 2003 ¹²³		
Observational	Peña 1999 ¹⁷⁸ Hulland 2002 ⁹⁷ Pitetti 2003 ¹⁸² Roback 2005 ¹⁸⁹ Mamula 2007 ¹⁵⁶ Sacchetti 2007 ¹⁹⁵ Lightdale 2009 ¹⁴⁰	Table 27 Table 28	168
Route of administration			
Oral / intranasal	Connors 1994 ⁴² Everitt 2002 ⁶⁰ Hartgraves 1994 ⁸⁹ Lightdale 2009 ¹⁴⁰	Table 21 Table 22	165
Intranasal / IM	Shashikiran 2006 ¹⁹⁹	Table 23	166
Dose			
	Fuks 1994 ⁷¹ Fukuta 1994 ⁷² Kanegaye 2003 ¹²³	Table 24 Table 26	167

1

6.3.1 Clinical methodological introduction

2		CLINICAL QUESTIONS
3 4 5		For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is midazolam (with or without: analgesia, another drug or psychological techniques):
6 7 8		- effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?
9		- safe for sedation (at mild, moderate, and deep levels) in different settings?
10 11 12		The literature was searched for systematic reviews and RCTs for the clinical efficacy of midazolam. The search was expanded to include non-RCT observational studies for the safety of midazolam.
13 14		There were no systematic reviews identified for the use of midazolam in paediatric sedation.
15 16		Twenty seven RCTs comparing midazolam in any route with other sedative drugs were assessed for efficacy and safety.
17 18		Seven non-RCTs observational studies in 5,412 patients assessed the safety of midazolam.
19 20		Crossover trials were treated separately from parallel armed trials unless there was sufficient data to allow their combination.
21 22 23 24		Meta-analyses for RCTs were performed where drug interventions and comparisons and outcomes were sufficiently homogenous and studies were combined regardless of dose, duration of intervention, procedure (within painful and non-painful groups), setting (e.g. dentistry, accidents and emergencies) and age.
25	6.3.2	2 Evidence profiles
26	6.3.2	2.1 RCT evidence profiles for efficacy and safety
27 28		Study characteristics and methodological quality of the study are provided in Appendix D. GRADE tables for quality assessment and summary of findings are provided below.
29		

PLACEBO COMPARISONS OR NON-DRUG TREATMENT

Table 3: Oral midazolam vs. placebo/no drug treatment; Liacouras 1998¹³⁹, Mortazavi 2009^{139,168}

Question: Should oral midazolam vs. placebo be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: gastroenterology and outpatients (dental postgraduate paediatric clinic)

Bibliography: Liacouras 1998 (intravenous placement); Mortazavi 2009 (dental extractions, teeth restorations, pulpotomies)

			Quality assessi	mont				,	Summary of fin	dings		
			Quality assessi	nent	No of pa	atients	E		Imp			
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	oral midazolam	placebo	Relative (95% CI)	Absolute	Quality	orta nce
Completi	ompletion of procedure (Mortazavi 2009)											
1	randomised trial	- /	no serious inconsistency		no serious imprecision	none	9/20 (45%)	20/20 (100%)	RR 2.16 (1.34 to 3.47) ²	0 more per 1,000	LOW	
Completi	on of procedure (L	iacouras 19	98)									
1	randomised trial		no serious inconsistency		no serious imprecision	none	59/62 (95.2%)	47/61 (77%)	RR 1.24 (1.07 to 1.43) ⁴	185 more per 1000 (from 54 more to 331 more) 0 more per 1,000	MODERA TE	
Adverse	events: Oxygen de	esaturation •	<90% (Mortazavi 2009	9)							•	
1	randomised trial	1 1	no serious inconsistency		no serious imprecision ⁵	none	0/20 (0%) ^{2,5}	0/20 (0%)	not pooled	-	LOW	

Mortazavi 2009: double blind study however partial allocation concealment and unclear blinding of outcome assessor and unclear ITT and N=20 (small study)

Note: The Mortazavi (2009) study used the Houpt scale to evaluate overall behaviour. One of the six ratings within this scale is called 'aborted', defined as 'no treatment rendered', so we used those data to calculate the number of patients who completed the procedure in each group.

² p=0.002

³ Liacouras 1998: unclear if ITT analysis was done; also large loss to follow up (>20%) for the outcome of patients satisfaction: for 32/123 (26%) patients, data was not available and this was greater in the control arm (18/61=30%) compared to the intervention arm (14/62=23%)

4 p=0.005

⁵ Mortazavi 2009: study stated that all patients remained close to 100% oxygen desaturation during procedure

⁶ For two RCTs, there was highly significant heterogeneity (I2=83%; p=0.02). Thus, the studies are presented individually.

Table 4: Oral midazolam vs. placebo; Fatovich 1995⁶³, Luhman 2001¹⁵¹

Question: Should oral midazolam vs. placebo (with local anesthesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: A & E

Bibliography: Luhman 2001 (suturing for laceration repair) Fatovich 1995 a) (suturing for laceration repair) Fatovich 1995 b) (suturing for laceration repair)

		0	uality accessm	ant	Summary of findings							
		Q	uality assessm	ent			No of pa	atients	Effe	ect		Importa
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other consideratio ns	oral midazolam	placebo	Relative (95% CI)	Absolute	Quality	nce
Completion	of procedure	e (Luhman 2001)										
1	randomised trial	serious		no serious indirectness	no serious imprecision	none	51/52 (98.1%)	50/50 (100%) ¹	RR 0.98 (0.93 to 1.04)	20 fewer per 1000 (from 70 fewer to 40 more)	MODER ATE	
Anxiety - as	ssessed by o	bservers using a	validated scale	(Herbertt-Mich	naelinees-Venh	am scale) (Fa	tovich 1995)					
1	randomised trial	no serious limitations ²		no serious indirectness	serious ³	none	33/57 (57.9%)	32/50 (64%)4	RR 0.89 (0.66 to 1.21)	70 fewer per 1000 (from 218 fewer to 134 more)	MODER ATE	
Distress - a	ssessed by p	parents using a va	lidated scale (r	neasured with	Visual Analog	gue Scale (VA	S); Better indica	ted by less) (F	atovich 1995)		•	•
1	randomised trial	no serious limitations ²		no serious indirectness	serious ³	none	57	50	-	MD -1.6 (-2.81 to -0.39) ⁵	MODER ATE	
Adverse ev	ents: Aspirat	ion (Luhman 200	1)		·	,	·					
1	randomised trial	serious ⁶		no serious indirectness	no serious imprecision ⁷	none	0/51 (0%) ⁷	0/50 (0%)	not pooled	-	MODER ATE	
Adverse ev	ents: Respira	atory intervention	(Luhman 2001))								
1	randomised trial	serious ⁶		no serious indirectness	no serious imprecision ⁸	none	0/51 (0%)8	0/50 (0%)	not pooled	-	MODER ATE	
Adverse ev	ents: Vomitir	ng (Luhman 2001)										
1	randomised trial	serious ⁶		no serious indirectness	no serious imprecision ⁹	none	0/51 (0%) ⁹	0/52 (0%)	not pooled	-	MODER ATE	

Luhman 2001: p=0.49
² Fatovich 1995: unclear ITT and unclear drop out rate; otherwise adequate allocation concealment and double blind

³ wide confidence intervals

⁴ Fatovich 1995: p=0.47 ⁵ Fatovich 1995: p=0.009

⁶ Luhman 2001: adequate concealment and low loss of follow up (1 patient in the midazolam group); however single blind study (only assessors were blind) and ITT was not performed -per protocol analysis instead

Luhman 2001: stated that not clinically apparent aspiration occurred in any patient

Luhman 2001: stated that no cardio respiratory adverse events occurred in any patient at any time

Luhman 2001: no incidents of vomiting in any patient in either group were observed

Table 5: Oral midazolam + non-pharmacological* vs. placebo + non-pharmacological*; Kapur 2004 124

*Love care, Tell show do techniques, physisical restrain

Question: Should oral Midazolam plus non-pharmacological technique vs. placebo plus non-pharmacological technique be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: dental hospital

Bibliography: Kapur 2004 (dental: restorations)

	. , ,	,	Ovelity and	accoment.				Summary of f	indings			
			Quality ass	essment			No of patients Effect					
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other	oral Midazolam plus non- pharmacological technique	placebo plus non- pharmacological technique	Relative (95% CI)	Absolute	Quality	Importance
Completion of procedure												
	randomised trial	, ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	18/20 (90%)	7/20 (35%)	RR 2.57 (1.39 to 4.76) ²	549 more per 1000 (from 136 more to 1000 more) 0 more per 1,000	LOW	
Duration of procedure (Better indicated by less)												
	randomised trial	. 1		no serious indirectness	serious ³	none	20	20	-	MD -9.83 (-17.22 to -2.44) ⁴	VERY LOW	

¹ Kapur 2004: assessors and patients blinded; however unclear allocation concealment, unclear if ITT was performed and dropouts not stated ² Kapur 2004: p=0.003 ³ Kapur 2004: wide confidence intervals ⁴ Kapur 2004: p=0.009

Table 6: Intranasal midazolam vs. placebo; Fishbein 1997 67

Question: Should intranasal midazolam vs. placebo be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: gastroenterology

Bibliography: Fishbein 1997 (Venipuncture)

			Quality assess	mont					Summary of t	findings		
			Quality assessi	ment		No of patients Effect					Importance	
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intranasal midazolam	placebo	Relative (95% CI)	Absolute	Quality	Importance
Distress - a	bistress - assessed by an observer using a validated scale (OBRS)											
	randomised trial			no serious indirectness	serious ²	none	15/19 (78.9%)	16/19 (84.2%)	RR 0.94 (0.69	51 fewer per 1000 (from 261 fewer to 227 more) 0 fewer per 1,000	LOW	

¹ Fishbein 1997: unclear allocation concealment; not true ITT performed -available case analysis only; otherwise double blind and low dropout (<20%) (venipuncture was not performed in 1 patient in each arm but reasons not stated)
² Fishbein 1997: wide confidence intervals
³ Fishbein 1997: p=0.68

Table 7: Intranasal midazolam vs. placebo; Ljungman 2000 148, Theroux 1993 210

Question: Should intranasal midazolam vs. placebo (with local anaesthesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: A & E and oncology
Bibliography: Theroux 1993 (suturing for laceration repair) Ljungman 2000 (cross over) (needle insertion)

			Quality assessme	ant.				Sı	ımmary of findir	igs		
			Quality assessme	ent	No of	patients	Ef	ffect				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations		placebo plus analgesia	Relative (95% CI)	Absolute	Quality	Import ance
Parent satis	sfaction (Ther	oux 1992)										
	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	15/22 (68.2%)	9/27 (33.3%)	RR 2.05 (1.12 to 3.75) ³	350 more per 1000 (from 40 more to 916 more) 0 more per 1,000	LOW	
Patients' pr	eference (Lju	gman 2000)					ļ			· ·	<u> </u>	
	randomised trial	very serious ⁴	no serious inconsistency	no serious indirectness	very serious ⁵	none	3/15 (20%) ⁶	0/10 (0%)	RR 4.81 (0.28 to 84.2) ⁷	0 more per 1000 (from 0 fewer to 0 more) 0 more per 1,000	VERY LOW	
Parents' pre	eference (Ljug	gman 2000)		•		•				·	ı	
	randomised trial	very serious ⁴	no serious inconsistency	no serious indirectness	no serious imprecision ⁵	none	13/27 (48.1%)	0/22 (0%)	RR 22.18 (1.39 to 353.32) ⁸	0 more per 1000 (from 0 more to 0 more) 0 more per 1,000	LOW	
Pain - asses	ssed by parer	nts using a v	alidated scale (meas	sured with: Vis	sual analogue	e scale; range o	f scores: 1-1	100; Better inc	dicated by less)	(Ljugman 2000)		
	randomised trial	serious ⁴	no serious inconsistency	no serious indirectness	serious ^{9,10}	none	22	27	-	not pooled	LOW	
Pain - asses	ssed by patie	nts using a	validated scale (mea	sured with: Vi	sual analogu	e scale; range c	of scores: 1-	100; Better in	dicated by less)	(Ljugman 2000)	*	
-	randomised trial	serious ⁴	no serious inconsistency	no serious indirectness	serious ^{10,11}	none	22	27	-	not pooled	LOW	
Adverse eve	ents: Vomitin	g after disch	narge (Theroux 1993))								
	randomised trial		no serious inconsistency		no serious imprecision ¹²		0/22 (0%) ¹²	0/27 (0%)	not pooled	-	LOW	

Theroux 1993: ITT appeared to have been performed and no dropouts were reported; however unclear allocation concealment and blinding of patients and assessors was partially possible only: the control group received no treatment while intervention and placebo groups blinded

² Theroux 1992: wide confidence intervals ³ Theroux 1992: p=0.02

⁴ Ljungman 2000: patients and assessors blinded; however, unclear allocation concealment, ITT not performed -available case analysis for the outcomes of pain and patient's preference- and

large amount (>20%) of loss of follow up at interview questionnaires for the outcomes of pain and preference; >35% of parents (25/74) and children (49/74) not contacted for the outcome of preference; for the outcome of pain, 38% (25/74) of children and 3% (2/74) of parents were not contacted

⁵ Ljungman 2000: very wide confidence intervals

⁶ Ljugman 2000: information/data available from only 25 parents/children; 15 in the first visit and 10 in the second visit

⁷ Ljungman 2000: p=0.28

- ⁸ Ljungman 2000: p=0.03
- ⁹ Ljungman 2000: point estimate not possible to calculate based on reported data. Study stated that pain assessed by parents was significantly less in the placebo group (median 81, IQR 46.7 to 92) than the intranasal midazolam (median 90, IQR 76.3 to 98; p=0.39)

 Ljungman 2000: median and IQR indicatives of skewed data

- Ljungman 2000: point estimate not possible to calculate based on reported data. Study stated that pain assessed by patients was no significant between groups; placebo group (median 87, IQR 41to 97), intranasal midazolam (median 87.5, IQR 78.3 to 100; p=0.625)
- 12 Theroux 1992: there was no evidence of children having vomited after discharge; vomiting was included as part of the follow up data collected from parents by telephone interview

2

8 9 10

11

Table 8: Oral midazolam vs. oral triclofos sodium; Singh 2002 203

Question: Should oral midazolam vs. oral triclofos sodium be used in children and young people undergoing diagnostic and therapeutic procedures?

Settings: dental hospital Bibliography: Singh 2002

			Quality asses	cmont				S	ummary	of findings		
			Quality asses	Silielit			No of p	patients		Effect		Imp
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	oral midazolam	oral triclofos sodium	Relative (95% CI)	Absolute	Quality	orta nce
Completic	n of procedu	re										
	randomised very no serious inconsistency y (when the patient was able to sit or sit)			no serious indirectness	no serious imprecision	none	30/30 (100%) ²	30/30 (100%)	not estimabl e	1000 fewer per 1000 (from 1000 fewer to 1000 fewer) 0 fewer per 1,000	LOW	
Recovery	(when the pa	tient was ab	le to sit or stand a	lone with minima	al assistance:	Better indicated	l by less)					
	randomised trial	very serious	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	30	-	MD -38.23 (-44.94 to -31.52)	LOW	
Length of	induction (Be	etter indicate	ed by less)									
		1 7 4	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	30	-	MD -16.10 (-18.11 to -14.09)	LOW	

TSingh 2002: patients and outcome assessors blinded however concealment, ITT and attrition details not stated
2 Singh 2002: all completed - ease of treatment completion rated as 1-excellent, 2-difficult and 3-impossible; study stated that treatment was most convenient for midazolam group than for triclofos group. Difficulty in treatment was significantly more for group of promethazine than for midazolam (p<0.01) and for triclofos (p<0.05)

Table 9: Sublingual midazolam vs. oral chloral hydrate; Layangool 2008 135

Question: Should sublingual midazolam vs. oral chloral hydrate be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: Outpatients' cardiology unit

Bibliography: Layangool 2008 (echocardiogram)

			Quality access	mont				Summary	of findings			
			Quality assess	sinent			No of pa	itients	Ef	fect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	sublingual midazolam	oral chloral hydrate	Relative (95% CI)	Absolute	Quality	ance
Completion	n of procedur	e (number c	f patients)									
1		,	no serious inconsistency	no serious indirectness	serious ²	none	127/132 (96.2%)	131/132 (99.2%)	RR 0.97 (0.93 to 1.01) ³	30 fewer per 1000 (from 69 fewer to 10 more)	VERY LOW	
Induction t	ime (Better ir	ndicated by	ess)									
1		very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	131	131	-	MD -13.80 (- 17.56 to - 10.04) ⁴	LOW	
Duration of	f procedure (Better indica	ated by less)									
1	randomised trial	1	no serious inconsistency	no serious indirectness	no serious imprecision	none	131	131	-	MD -0.40 (- 1.59 to 0.79) ⁵	LOW	
			Total time covered wed full recovery.	from administration	on to recover	y in full, determ	nined by vital sigi	ns, oxygen sat	uration and	conscious le	vel which	n were
1		,	no serious inconsistency	no serious indirectness	no serious imprecision	none	131	131	-	MD 38.80 (33.18 to 44.42) ⁶	LOW	
Adverse ev	ents: Vomiti	ng										
1	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ⁷	none	1/132 (0.8%)	14/132 (10.6%)	RR 0.07 (0.01 to 0.54) ⁸	99 fewer per 1000 (from 49 fewer to 105 fewer)	LOW	
								0.8%	0.04)	7 fewer per 1,000		

Layangool 2008: stated as double blinded study; however, partial allocation concealment and ITT not performed, available case analysis instead for children who both completed procedure in full plus children who completed procedure partially; and <20% lost of follow up

Layangool 2008: crosses left precision limit

³ Layangool 2008: p=0.10

⁴ Layangool 2008: P<0.00001 ⁵ Layangool 2008: p=0.51

Layangool 2008: p<0.00001

⁷ Layangool 2008: precise

⁸ Layangool 2008: p=0.01

Note: For Layangool (2008), the ability to complete the procedure was described in four different levels. Level 0 was defined as 'unable to perform the study'; level 1 was stated as 'important part of the study accomplished, but study shortened'; level 2 defined a 'complete study possible with coaxing'; and level 3 was defined as 'complete study easily accomplished'. Furthermore, the RCT stated that procedure was incompletely performed in four cases in the midazolam group and it was failed in one case in each group. Thus we dichotomised the four levels into procedure completely performed (level 2 + level 3) and procedure not or partly performed (level 0 and level 1).

Table 10: Rectal midazolam + non-pharmacological intervention* versus nitrous oxide (70%) + non-pharmacological intervention*; Zier 2008 ²³⁷

*distraction: storytelling, soothing discourse

Question: Should rectal midazolam vs. nitrous oxide (with topical anaesthesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and

therapeutic procedures? Settings: gastroenterology

Bibliography: Zier 2008 (injections for spasticity)

			Quality asse	eemant				Summary of fin	dings			
			Quality asse	SSIIICIII			No of p	atients	E	ffect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	rectal midazolam plus placebo plus topical anaesthesia plus non- pharmacological intervention		Relative (95% CI)	Absolute	Quality	Import ance
Pain - n	umber of pa	tients - asse	ssed by a train	ed observer ι	ısing a valid	ated scale (Face	, Legs, Activity, Cry, C	Consolability (FLACC))			
	randomised trial			no serious indirectness	serious ²	none	24	25	not pooled ²	-	MODERATE	
Parents	satisfaction	assessed o	n a 1 to 10 sca	le (measured	with: arbitra	ry scale; range o	of scores: 1-10; Better	indicated by less)				
	randomised trial			no serious indirectness	serious ³	none	22	25	-	not pooled4	MODERATE	
Total tin	ne (Better in	dicated by l	ess)							<u> </u>		
	randomised trial	1		no serious indirectness	serious ⁵	none	24	25	-	not pooled5	MODERATE	
Adverse	events: Vo	miting durin	g drug nitrous	oxide adminis	stration							•
	randomised trial			no serious indirectness	serious ^{2,6}	none	0/25 (0%)	4/25 (16%)	RR 0.11 (0.01 to 1.96) ⁷	142 fewer per 1000 (from 158 fewer to 154 more) 0 fewer per 1,000	MODERATE	

Zier 2008: adequate concealment, ITT appeared to be performed and there were no loss of follow up reported, adequate allocation concealment and both patients and outcome assessors were blind

² Zier 2008: reported p-value=0.010; sample size small; median scores were 6 for the midazolam group and 4 for the nitrous oxide group

³ Zier 2008: reported satisfaction was no significant between groups; p=0.10; assessed on a 1 to 10 arbitrary scale where 1=satisfaction and 10=dissatisfaction; median scores were 2 for the midazolam group and 1 for the nitrous oxide group; small study

⁴ Zier 2008: reported p=0.10

⁵ Zier 2008: stated that there was no difference between groups regarding the time each group stayed in the clinic, did not report p-value; small study ⁶ Zier 2008: very wide confidence intervals

⁷ Zier 2008: p=0.13

2

COMBINATION COMPARISONS

Table 11: Oral midazolam vs. oral midazolam + nitrous oxide/oxygen; Al-zahrani 2009 12

Question: Should oral midazolam vs. oral midazolam plus nitrous oxide/oxygen (with topical and local anaesthesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: dental hospital

Bibliography: Alzahrani 2009 (dental restorative procedures)

			Quality asse	coment				Summary of findir	ngs			
			Quality asse	SSIIIEIII			No o	of patients	Ef	fect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	oral midazolam plus topical anaesthesia plus local anaesthesia	oral midazolam plus nitrous oxide/oxygen plus topical anaesthesia plus local anaesthesia	Relative (95% CI)	Absolute	Quality	Importan ce
Complet	tion of procedure (number of patients)											
	randomised trial		no serious inconsistency	no serious indirectness	serious ^{2,3}	none	30/30 (100%)	30/30 (100%)	not estimable	-	LOW	
Induction	n time (meas	sured with:	minutes; Better	indicated by le	ess)							
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	30	30	-	MD -0.70 (- 2.59 to 1.19) ⁴	LOW	
Duration of procedure (time from bringing the patient to the operating room until the planned dental procedures were completed Better indicated by less)												
	randomised trial		no serious inconsistency	no serious indirectness	very serious ⁵	none	30	30	-	MD 0.10 (- 2.79 to 2.99) ⁶	VERY LOW	

Alzahrani 2009: cross-over trial, unclear concealment, unclear blinding of outcome assessors but all patients completed the trial and all patients appeared to be included in analyses

² Alzahrani 2009: imprecise as crosses left precision limit; small sample ³ Al-zahrani 2009: small sample ⁴ Alzahrani 2009: p=0.47

⁵ Alzahrani 2009: imprecise, crosses right precision limit and very wide confidence intervals; small sample

Alzahrani 2009: p=0.95

Note: For Alzahrani (2009), the completion of procedure was based on assessment of overall behaviour using the Houpt scoring system (sleep, crying, movement, behaviour), most of the patients movement did not interrupt dental treatment on both visits and most of the patients showed good or very good behaviour in both groups; with no poor behaviour or treatment aborted.

Table 12: Oral midazolam + nitrous oxide vs. nitrous oxide + placebo; Luhman 2001 151

Question: Should oral midazolam plus nitrous oxide vs. nitrous oxide plus placebo (with local anaesthesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: accidents and emergencies

Bibliography: Luhman 2001 (suturing and laceration repairs)

			Quality asse	neemont				Summ	ary of findin	gs		
			Quality ass	essillelli			No of pa	atients	E	Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	oral midazolam plus nitrous oxide plus analgesia	nitrous oxide plus placebo plus analgesia	Relative (95% CI)	Absolute	Quality	Import ance
Complet	ion of proce	edure			•	•		•				•
1	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision ²	none	52/52 (100%)	51/51 (100%)	not estimable ²	1000 fewer per 1000 (from 1000 fewer to 1000 fewer) 0 fewer per	MODERATE	
										1,000		
Adverse	events: As	piration								<u>, </u>		
1	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	0/52 (0%) ³	0/51 (0%)	not pooled	-	MODERATE	
Adverse	events: Res	spiratory int	ervention									
1	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	0/52 (0%)4	0/51 (0%)	not pooled	-	MODERATE	
Adverse	events: Vo	miting		<u> </u>		!				!		!
	randomised trial		no serious inconsistency	no serious indirectness	serious ⁵	none	1/52 (1.9%)	5/51 (9.8%)	RR 0.20 (0.02 to 1.62) ⁶	78 fewer per 1000 (from 96 fewer to 61 more) 0 fewer per 1,000	LOW	
² Luhmar ³ Luhmar ⁴ Luhmar ⁵ Luhmar	n 2001: not e n 2001: state n 2001: state	estimable, all ad that not clir ad that no car confidence in	patients complete nically apparent a dio respiratory ac	ed the procedures is piration occur	re red in any patie	• , •	assessors were bli	nd) and ITT was	not performe	ed		

Table 13: Oral midazolam + intravenous propofol vs. intravenous propofol; Paspatis 2006 176

Question: Should oral midazolam plus intravenous propofol vs. intravenous propofol (with local anaesthesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: gastroenterology
Bibliography: Paspatis 2006 (endoscopy)

			Quality ass	ocemont				Summary of	of finding	S		
			Quality ass	essillelit			No of patie	ents		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	oral midazolam plus intravenous propofol	intravenous	Relative (95% CI)	Absolute	Quality	ance
Duration	of procedure	(Better indi	cated by less)	•								
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	26	28	-	MD 0.10 (-2.5 to 2.7) ³	LOW	
Recovery	from comple	etion of proc	edure to recover	y/discharge crite	eria met (meası	red with: REACT	score; range of scores	s: 0-10; Better i	ndicated	by more)		
1	randomised trial		no serious inconsistency		no serious imprecision	none	26	28	-	MD 18.20 (16.14 to 20.26) ⁴	MODERATE	

Paspatis 2006: ITT appeared to have been performed and no loss of follow up were reported; however, allocation concealment and blinding of patients were not stated and blinding of outcome assessors was not clear

² Paspatis 2006: wide confidence intervals

³ P=0.94

⁴ P<0.00001

Table 14: Intravenous midazolam + intravenous propofol vs. intravenous propofol; Disma 2005 56

Question: Should intravenous midazolam plus intravenous propofol vs. intravenous propofol be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: gastroenterology

Bibliography: Disma 2005 (Endoscopy)

			Quality ass	occment				Summa	ary of finding	js .		
			Quality ass	essment			No of pati	ents		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous midazolam plus intravenous propofol	intravenous propofol	Relative (95% CI)	Absolute	Quality	Import ance
Completi	ion of proced	dure (numbe	er of patients)	•	•	•						
	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision ²	none	78/78 (100%)	80/80 (100%)	not estimable ³	-	MODERATE	
Duration	of procedur	e (Better inc	dicated by less)									
	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	78	80	-	MD -0.20 (-0.98 to 0.58) ⁴	MODERATE	
Recovery	y from comp	letion of pro	cedure to recov	ery/discharge	riteria met (Be	etter indicated by	less)					
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	78	80	-	MD 2.50 (-0.4 to 5.4) ⁵	LOW	
Adverse	events: Assi	isted ventila	tion (bag-valve i	mask)								
	randomised trial		no serious inconsistency	no serious indirectness	serious ⁶	none	0/78 (0%)	5/80 (6.3%)	RR 0.09 (0.01 to 1.66) ⁷	57 fewer per 1000 (from 62 fewer to 42 more) 0 fewer per 1,000		
Adverse	events: Oxy	gen desatur	ation <90%									
	randomised trial		no serious inconsistency	no serious indirectness	serious ⁶	none	2/78 (2.6%)	3/80 (3.8%)	RR 0.68 (0.12 to 3.98) ³	12 fewer per 1000 (from 33 fewer to 113 more)	LOW	

Disma 2005: ITT appeared to be performed and there were no loss of follow up reported; however, allocation concealment and blinding of patients were not stated and blinding of outcome assessors was not clear

² Disma 2005: wide confidence intervals

³ P=0.67

⁴ P=0.62

⁵ P=0.09

⁶ Disma 2005: wide confidence intervals

⁷ P=0.11

Table 15: Intravenous midazolam + intravenous meperidine vs. intravenous meperidine; Fishbein 1997 67

Question: Should intravenous midazolam plus intravenous meperidine vs. intravenous meperidine be used in children and young people under 19 years of age undergoing diagnostic and

therapeutic procedures?

Settings: gastroenterology Bibliography: Fishbein 1997 (esophagogastroduodenoscopy)

			Quality ass	ocement				Summary	of findings	3		
			Quality ass	essinent			No of pa	tients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous midazolam plus intravenous meperidine	placebo plus intravenous meperidine	Relative (95% CI)	Absolute	Quality	Import ance
Distress	assessed by	y an observe	er using a valida	ted scale (Obs	ervational Beh	aviour Rating Sca	ale (OBRS) - data for	major behaviours	s)			
	randomised trial		no serious inconsistency		no serious imprecision ²	none	18/20 (90%)	19/20 (95%)	RR 0.95 (0.79 to 1.13) ³	48 fewer per 1000 (from 199 fewer to 123 more) 0 fewer per 1,000	MODERATE	
Duration	uration of procedure (Better indicated by less)											
	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ⁴	none	20	20	-	MD 0.40 (-1.22 to 2.02) ⁵	LOW	

Fishbein 1997: unclear allocation concealment; not true ITT performed -available case analysis only; otherwise double blind and low dropout (<20%) (venipuncture was not performed in 1 patient in each arm but reasons not stated)

² Fishbein 1997: precise

³ Fishbein 1997: p=0.55 ⁴ Fishbein 1997: imprecise ⁵ Fishbein 1997: p=0.63

Table 16: Intravenous midazolam + intravenous morphine vs. intravenous propofol + intravenous morphine + local anaesthesia; Havel 1999⁹⁰

Question: Should intravenous midazolam plus intravenous morphine vs. intravenous propofol plus intravenous morphine plus local anaesthesia (with placebo in both arms) be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: accidents and emergencies

Bibliography: Havel 1999 (fractures of the forearm, humerus, femur, lower leg, or hand, hip dislocation)

			Quality ass	accment				Summary of fi	ndings			
			Quality assi	essinent			No of	patients	E	Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous midazolam plus intravenous morphine plus placebo	intravenous propofol plus intravenous morphine plus placebo plus lidocaine	Relative (95% CI)	Absolute	Quality	Import ance
Complet	ion of proce	dure										
	randomised trial	very serious¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	46/46 (100%)	43/43 (100%)	not estimable 0 (0 to 0) ²	0 fewer per	LOW	
Industic	a tima /Datta	w indicated	hy less)							1,000		
	n time (Bette randomised			no serious	no serious	none				MD 0.20 (-1.89		1
		- /		indirectness	imprecision	none	46	53	-	to 2.29) ³	LOW	
Duration	of procedur	re (Better in	dicated by less)	<u> </u>				!		· · · · · ·		
	randomised trial	- /	no serious inconsistency	no serious indirectness	no serious imprecision	none	46	43	-	MD 0.70 (-5.34 to 6.74) ⁴	LOW	
Pain (nu	mber of pati	ents who re	ported pain)		•	•		•				
	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	very serious⁵	none	2/46 (4.3%)	3/43 (0%)	RR 0.61 (0.1 to 3.82) ⁶	0 fewer per 1,000	VERY LOW	
Recover	y time (Bette	r indicated	by less)		•	•		•				
	randomised trial	very serious ¹		no serious indirectness	no serious imprecision	none	46	43	-	MD 46.80 (40.76 to 52.84) ⁷	LOW	
Total tim	e (from adn	nission unti	l having been di	scharged from	the clinic; Be	etter indicated by	less)					
	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	46	43	-	MD 23.80 (0.93 to 46.67)	LOW	
Adverse	events: Asp	iration										
1	randomised	very	no serious	no serious	no serious	none	0/46 (0%)8	0/43 (0%)	not pooled8	-		

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				1								
	trial	serious ¹	inconsistency	indirectness	imprecision ⁸					-	LOW	
Adverse	events: Ass	sisted ventil	ation									
1	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ⁹	none	0/46 (0%) ⁹	0/43 (0%)	not pooled9	-	LOW	1
Adverse	events: End	dotracheal i	ntubation									
1	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	0/46 (0%)	0/43 (0%)	not pooled	0 fewer per 1000 (from 0 fewer to 0 fewer) 0 fewer per 1,000	LOW	
Adverse	events: Oxy	gen desatu	ration < 90%						,		•	
1	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	serious ¹⁰	none	_10	_10	not pooled ¹⁰	-	VERY LOW	

Havel 1999: patients and outcome assessors were blind and low loss of follow-up; however, inadequate allocation concealment, the sedationist knew medications, infusion tubing and intravenous site and ITT was no performed -per protocol analysis instead

Havel 1999: not estimable; all patients completed the procedure

Havel 1999: p=0.85

Havel 1999: p=0.82

Havel 1999: p=0.59

Havel 1999: p<0.00001

Havel 1999: stated that not clinically apparent aspiration occurred in any patient in either sedation group

Havel 1999: no patient in either sedation group required assisted ventilation

Havel 1999: stated that groups had similar frequencies of hypoxemia but explicit data for this was not reported; selective reporting

Table 17: Intravenous midazolam + intravenous fentanyl (analgesic) vs. intravenous fentanyl (analgesic); Antmen 2005 15

Question: Should intravenous midazolam plus intravenous alfentanyl vs. intravenous alfentanyl be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: paediatric haematology outpatients

Bibliography: Antment 2005 (bone marrow aspiration)

			Quality ass	acemont				Summary of f	indings			
			Quality ass	essinent			No of patier	nts	E	ffect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous midazolam plus intravenous alfentanyl	intravenous alfentanyl	Relative (95% CI)	Absolute	Quality	
Completion	on of proced	ure			•	•			•			
1		- / 1	no serious inconsistency	no serious indirectness	no serious imprecision	none	20/20 (100%)	20/20 (100%)	not estimable	-	LOW	
Pain asse by less)	essed by the	anaesthetis	t using a validate	d scale - 2 (mea	sured with: Chi	Idren's Hospital o	f Eastern Ontario Pain So	cale (CHEOPS);	range of so	cores: 0-13; B	etter indi	icated
1		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	20	20	-	MD -0.15 (- 1.05 to 0.75)	VERY LOW	
Pain asse	essed by the	anaesthetis	t using a validate	d scale - 1 (mea	sured with: Vis	ual analogue scal	e (VAS); range of scores:	0-10; Better in	dicated by	ess)		•
1		- 1	no serious inconsistency	no serious indirectness	serious ²	none	20	20	-	MD -0.30 (- 1.8 to 1.2)	VERY LOW	
Adverse (events: Oxyg	en desatura	tion <90%	•	•	•						
1		very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	0/20 (0%)	0/20 (0%)	not pooled	-	LOW	
Adverse (events: Vomi	iting	•	•					•			
1			no serious inconsistency	no serious indirectness	no serious imprecision	none	0/20 (0%)	0/20 (0%)	not pooled	-	LOW	

Antment 2005: ITT appeared to be performed and there were no loss of follow up reported; however, allocation concealment and blinding of patients were not stated and blinding of outcome assessors was not clear; small study N=20

² Antment 2005: confidence intervals cross precision limits

Table 18: Intravenous midazolam + intravenous remifentanil (analgesic) vs. intravenous remifentanil (analgesic); Antmen 2005 15

Question: Should intravenous midazolam plus intravenous remifentanil vs. intravenous remifentanil be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: paediatric haematology outpatients

Bibliography: Antment 2005 (bone marrow aspiration)

			Ovality	acamant.				Summary of fi	ndings			
			Quality ass	essment			No of patier	nts	Е	ffect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous midazolam plus intravenous remifentanil	intravenous remifentanil	Relative (95% CI)	Absolute	Quality	ance
Completi	on of proced	ure							•			
1	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	20/20 (100%)	20/20 (100%)	not estimable	-	LOW	
Pain asso by less)	essed by the	anaesthetist	t using a validate	ed scale - 2 (mea	sured with: Ch	ildren's Hospital o	of Eastern Ontario Pain S	cale (CHEOPS);	range of so	cores: 0-13; B	etter ind	icated
1		,	no serious inconsistency	no serious indirectness	serious ²	none	20	20	-	MD -0.05 (- 0.68 to 0.58) ³	VERY LOW	
Pain asse	essed by the	anaesthetist	using a validate	d scale - 1 (mea	sured with: Vis	ual Analogue Sca	le (VAS); range of scores	: 0-10; Better in	dicated by	less)	•	•
1	randomised trial	1	no serious inconsistency	no serious indirectness	serious ²	none	20	20	-	MD -0.05 (- 0.86 to 0.76)	VERY LOW	
Adverse	events: Oxyg	en desatura	tion <90%			•						
1	randomised trial	, ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	0/20 (0%)	0/20 (0%)	not pooled	-	LOW	
Adverse	events: Vom	iting		1			'		•		,	,
1	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	0/20 (0%)	0/20 (0%)	not pooled	-	LOW	

Antment 2005: ITT appeared to be performed and there were no loss of follow up reported; however, allocation concealment and blinding of patients were not stated and blinding of outcome assessors was not clear; small study N=20

Antment 2005: confidence intervals cross precision limits

Antment 2005: p=0.88

Table 19: Intravenous midazolam + intravenous ketamine vs. intravenous ketamine + placebo; Sherwin 2000 ²⁰¹; Wathen 2000 ²²⁷; Dilli 2008 ⁵⁵

Question: Should intravenous midazolam plus intravenous ketamine vs. intravenous ketamine plus placebobe used in children and young people under 19 years of age undergoing

diagnostic and therapeutic procedures? **Settings:** A & E and hospital outpatients

Bibliography: Wathen 2000 (fractures, lacerations, other including joint aspiration, abscess drainage, vaginal laceration, dog bite, wound care, chest tube placement, nail bed injury, vaginal foreign body removal, inguinal hernia, urologic procedures); Sherwin 2000 (intravenous catheter insertion for orthopaedic, wound or thermal, other procedures) Dilli 2008 (lumbar puncture)

			Ovelity assess	mant.	Summary of findings							
			Quality assess	nent			No of patients Effect					
No of studies	Design	Limitation s	Inconsistency	Indirectness	Imprecision	Other considerations	efficacy of intravenous midazolam plus intravenous ketamine	intravenous ketamine plus placebo	Relative (95% CI)	Absolute		Import ance
Completion of procedure (Sherwin 2000; Wathen 2000)												
2	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	no serious imprecision ^{3,4}	none	190/190 (100%)	180/180 (0%)	not estimable	-	MODERAT E	
Induction t	ime (Better ir	dicated by	less) (Dilli 2008)	•		•						
1	randomised trial	very serious⁵	no serious inconsistency	no serious indirectness	serious ⁶	none	48	51	-	MD -0.80 (- 1.36 to - 0.24)	VERY LOW	
Parents' sa	atisfaction (Di	Ili 2008)										
1	randomised trial	very serious⁵	no serious inconsistency		no serious imprecision ⁷	none	-	-	p=0.001 ⁷	-	LOW	
Recovery t	ime (Better ir	dicated by	less) (Dilli 2008)	•		•						
1	randomised trial	very serious⁵	no serious inconsistency	no serious indirectness	serious ⁸	none	48	51	-	MD 2.20 (- 0.79 to 5.19)	VERY LOW	
Parents' sa	atisfaction (W	athen 2000)										
1	randomised trial	serious ¹	no serious inconsistency		no serious imprecision	none	112/137 (81.8%)	115/129 (89.1%)	RR 0.92 (0.83 to 1.01)	71 fewer per 1000 (from 151 fewer to 9 more) 0 fewer per 1,000		
Duration of	f procedure (Better indic	ated by less) (Wath								_	
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ⁹	none	112	115	-	MD -1(IQR, 95% CI -5 to 1)	LOW	

DRAFT FOR CONSULTATION

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- ¹ Wathen 2000: adequate allocation concealment and patients and assessors were blinded; however, ITT was not performed -per protocol analysis instead; low amount of loss of follow up: 3 randomised patients dropped out, 2 in the intervention and 1 in the control group had protocol violation and received intramuscular vial instead of intravenous
- ² Sherwin 2000: adequate allocation concealment and patients and were blinded, ITT appeared to have been performed and there were no loss of follow up reported
- ³ Wathen 2000: not estimable, all patients completed the procedure
- ⁴ Sherwin 2000: not estimable, all patients completed the procedure
- ⁵ Dilli 2008: adequate allocation concealment, and outcome blinded; however ITT was not performed -per protocol analysis instead: 104 randomised but 99 analysed: midazolam+ketamine=48, ketamine=51; loss of follow up: midazolam+ketamine group: 4%(2/50) one patient did not received allocated intervention and one was lost to follow-up; 6%(3/54) one patient did not received allocated intervention and two were lost to follow-up; patients were not blind
- ⁶ Dilli 2008*: crosses left confidence limit
- ⁷ Dilli 2008: stated that parental satisfaction was significantly higher in patients in the midazolam group, p=0.001
- ⁸ Dilli 2008: crosses right confidence limit
- ⁹ Wathen 2000: based on data reported on the study, it was not possible to calculate a RR; the study stated that the difference between ketamine+midazolam versus ketamine plus placebo was no significant; mean difference -1 minute (IQR, 95% CI-5 to 1)
- *Note: Dilli (2008), also stated that patients were discharged two hours after procedure and after being awake, coherent and able to tolerate oral food

Table 20: safety of intravenous midazolam plus intravenous ketamine vs. intravenous ketamine plus placebo

Question: What is the safety of intravenous midazolam plus intravenous ketamine vs. intravenous ketamine plus placebo in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: A & E and hospital outpatients

Bibliography: Wathen 2000 (fractures, lacerations, other including joint aspiration, abscess drainage, vaginal laceration, dog bite, wound care, chest tube placement, nail bed injury, vaginal foreign body removal, inguinal hernia, urologic procedures); Sherwin 2000 (intravenous catheter insertion for orthopaedic, wound, thermal injury or other procedures) Dilli 2008 (lumbar puncture)

			Quality acc	ncomont		Summary of findings						
			Quality asso	essinent			No of patients			Effect		П
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	adverse events of intravenous midazolam plus intravenous ketamine	intravenous ketamine plus placebo	Relative (95% CI)	Absolute		Impo rtanc e
Adverse events: Vomiting (during visit and at home 12 hrs after discharge and well into recovery) (Wathen 2000; Sherwin 2000)												
2	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	serious ^{3,4}	none	14/190 (7.4%)	31/180 (17.2%)	RR 0.43 (0.24 to 0.77) ⁵	98 fewer per 1000 (from 40 fewer to 131 fewer) 0 fewer per 1,000	LOW	
Adverse	events: As	sisted ventila	tion (bag mask)	(Sherwin 2000	; Wathen 2000)						
2	randomised trial	no serious limitations ²	no serious inconsistency	no serious indirectness	serious imprecision ⁶	none	1/190 (0.5%)	1/180 (0.6%)	RR 0.94 (0.06 to 14.9) ⁹	0 fewer per 1000 (from 0 fewer to 0 fewer)	LOW	
									14.9)	0 fewer per 1,000		
Adverse	events: As	piration (Watl	hen 2000; Sherw	in 2000)	•							
2	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	no serious imprecision ⁷	none	0/190 (100%) ⁷	0/180 (100%) ⁷	not pooled	-	MODERATE	E
Adverse	events: En	dotracheal in	tubation (Wathe	n 2000)								
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ⁸	none	0/137 (100%) ⁸	0/129 (100%)8	not pooled	-	MODERATE	E
Adverse	events: Ox	ygen desatur	ation 90% (Wath			2008)						
3	randomised trial	very serious ^{1,2,10}	no serious inconsistency ¹¹	no serious indirectness	no serious imprecision	none	14/238 (5.9%)	3/231 (1.3%)	RR 4.01 (1.27 to 12.68) ¹²	39 more per 1000 (from 4 more to 152 more) 0 more per 1,000	LOW	

Wathen 2000: adequate allocation concealment and patients and assessors were blinded; however, ITT was not performed -per protocol analysis instead; low amount of loss of follow up: 3 randomised patients dropped out, 2 in the intervention and 1 in the control group had protocol violation and received intramuscular vial instead of intravenous

² Sherwin 2000: adequate allocation concealment and patients and assessors were blinded, ITT appeared to have been performed and there were no loss of follow up reported

³ Wathen 2000: very wide confidence intervals

⁴ imprecise: cross left precision limits

⁵ Wathen 2000 and Sherwin 2000: p=0.005

Wathen 2000 and Sherwin 2000: there was no incidence of aspiration in any patient in either group

⁸ Wathen 2000: stated that endotracheal intubation was not performed in any patient

⁹ Wathen 2000: p=0.97

¹¹ Wathen 2000, Sherwin 2000 and Dilli 2008: not significant heterogeneity=0%, p=0.53

⁶ Sherwin 2000; Wathern 2000: imprecise, wide confidence intervals; no assisted ventilation was required in any patients in either group in the study by Sherwin (2000) while one patient in each arm required assisted ventilation in the study by Wathen (2000)

¹⁰ Dilli 2008: adequate allocation concealment, and outcome assessors blinded; however ITT was not performed -per protocol analysis instead: 104 randomised but 99 analysed: midazolam+ketamine=48, ketamine=51; loss of follow up: midazolam+ketamine group: 4%(2/50) one patient did not received allocated intervention and one was lost to follow-up; 6%(3/54) one patient did not received allocated intervention and two were lost to follow-up; page 150.

¹² Wathen 2000, Sherwin 2000 and Dilli 2008: p=0.02

ROUTE OF ADMINISTRATION COMPARISONS

Table 21: Oral midazolam vs. intranasal midazolam; Connors 1994 42; Everitt 2002 60

Question: Should oral midazolam vs. intranasal midazolam be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: accidents and emergencies

Bibliography: Connors 1994 (suturing for laceration repair) Everitt 2002 (suturing for laceration repair)

Quality assessment								Summary of findings					
			Quality ass	essment	No of patients		Effect			Importance			
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	oral midazolam	Ansoluta		Quality	Importance		
Completion of procedure (Connors 1994)													
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	26/26 (100%)	28/28 (100%)	not estimable	-	MODERATE		
Distress a	Distress assessed by an observer using a validated scale (measured with VAS; range of scores: 1-100; Better indicated by less) (Everitt 2002)												
1		very serious ²	no serious inconsistency	no serious indirectness	serious ³	none	45	42	=	MD -13 (-25.83 to -0.17) ⁴	VERY LOW		
Total time	: administrati	on to recove	ry area/discharge	criteria (measure	d with: minutes	Better indicated b	y less) (Conn	ors 1994; Everit	tt 2002)				
1	randomised trial	very serious ^{1,2}	no serious inconsistency	no serious indirectness	serious ³	none	71	70	-	MD 3 (-1.44 to 7.44) ⁵	VERY LOW		
Adverse e	Adverse events: Vomiting (Everitt 2002)												
1		very serious ¹	no serious inconsistency	no serious indirectness	serious ⁶	none	6	6	not pooled ⁶	-	VERY LOW		

Connors 1994: double blind (patients and outcome assessors), double placebo trial with low loss of follow up: 7% (4/58) of patients were excluded from analyses (2 in each arm had protocol violation and for 2 data collection was not available); however, allocation concealment was not stated and ITT was not performed -per protocol analysis instead; small study

² Everitt 2002: unclear allocation concealment, outcome assessors partially blinded (assessors: staff participating were unaware of sedative being given but parents who also assessed children for anxiety were aware of sedative given), ITT and amount of loss of follow up were unclear or not stated; also, study stated to have obtained data on parents' satisfaction after discharge but results data were not reported (selective outcome reporting)

³ Everitt 2002: imprecise; wide confidence intervals

⁴ Everitt 2002: p=0.05

⁵ Everitt 2002: p=1.00

⁶ Everitt 2002: selective outcome reporting; data was collected for analysis but results were not shown

Question: Should oral midazolam plus nitrous oxide vs. intranasal midazolam plus nitrous oxide (with local anaesthesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: dental hospital

Bibliography: Hartgraves 1994 (dental: extractions, restorations, pulpotomies, brief) Lee-kim 2004 (dental: amalgam, composite restorations, pulpotomy procedures, stainless steel crowns, extractions)

			Quality	rocement		Summary of findings						
			Quality ass	sessment			No of patients		Effect			
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	oral midazolam plus nitrous oxide plus lidocaine	intranasal midazolam plus nitrous oxide plus lidocaine	Relative (95% CI)	Absolute	Quality	Importance
Completi	Completion of procedure (Hartgraves 1994)											
1	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	45/50 (90%)	47/50 (94%)	RR 0.96 (0.85 to 1.08) ³	38 fewer per 1000 (from 141 fewer to 75 more) 0 fewer per 1,000	LOW	
Induction	time (Bette	r indicated	by less) (Lee-Kim	n 2004)								
1	randomised trial	serious ⁴	no serious inconsistency	no serious indirectness	no serious imprecision⁵	none	20	20	-	MD 9.95 (7.56 to 12.34) ⁶	MODERATE	
Total time	e (from adm	inistration t	o recovery area/o	discharge criteri	a -defined as dr	ugs working time	e) (Lee-Kim 200	4)				
1	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	serious ⁷	none	20	20	-	MD 8.80 (2.73 to 14.87) ⁹	LOW	
Adverse (Adverse events: Oxygen desaturation <90% (Hartgraves 1994)											
1	randomised trial	very serious ¹	no serious inconsistency	no serious indirectness	serious ⁸	none	2/50 (4%)	1/50 (2%)	RR 2 (0.19 to 21.36) ¹⁰	20 more per 1000 (from 16 fewer to 407 more) 0 more per 1,000	VERY LOW	

Hartgraves 1994: allocation concealment not stated, blinding of assessors was unclear and patients not blinded; also, it was unclear whether ITT was performed and unclear loss of follow up

² Hartgraves 1994: precise ³ Hartgraves 1994: p=0.46

⁴ Lee-Kim 2004: assessors were blinded, ITT appeared to have been performed and no loss of follow up were reported; however, unclear allocation concealment and patients were not blinded

⁵ Lee-Kim 2004: precise ⁶ Lee-Kim 2004: p<0.00001

⁷ Lee-Kim 2004: imprecise, wide confidence intervals

 $^8\text{Hartgraves}$ 1994: imprecise, very wide confidence intervals 9 Lee-Kim 2004; p=0.005 10 Hartgraves 1994: p=0.57

Table 23: Intranasal midazolam vs. intramuscular midazolam; Shashikiran 2006 199

Question: Should intranasal midazolam vs. intramuscular midazolam (with analgesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and

therapeutic procedures?

Settings: dental hospital Bibliography: Sashikiran 2006 (dental)

			Quality ass	accmont				Summary of	findings			
			Quality ass	essment			No of	patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intranasal midazolam plus analgesia		Relative (95% CI)		Quality	Importance
Induction	time (Better	indicated b	y less)									
	randomised trial		no serious inconsistency		no serious imprecision²	none	20	20	-	MD -4.90 (- 6.14 to -3.66) ³	MODERATE	
Recovery	from compl	etion of pro	cedure to recove	ry/discharge cr	iteria met (Bett	er indicated by le	ss)					
	randomised trial		no serious inconsistency		no serious imprecision ²	none	20	20	-	MD -24.40 (- 26.48 to - 22.32) ^{3,4}	MODERATE	
Adverse	events: Vom	iting										
	randomised trial		no serious inconsistency		no serious imprecision ⁵	none	0/20 (0%) ⁵	0/20 (0%)	not pooled	-	MODERATE	

Sashikran 2006: ITT appeared to have been performed and no loss of follow up were reported; however, unclear allocation concealment, blinding of outcomes assessors was unclear and patients were not blinded

² Sashikran 2006: precise ³ Sashikran 2006: p<00001 ⁴ Sashikran 2006: p<00001

⁵ Sashikran 2006: stated that there was not a single incidence of vomiting

DOSE COMPARISONS

- Table 24: Intranasal midazolam 0.3mg/kg + nitrous oxide vs. intranasal midazolam 0.2 mg/kg + nitrous oxide;
- Fuks 1994 71; Fukuta 1994 72

Question: Should intranasal midazolam 0.3 mg/kg plus nitrous oxide vs. intranasal midazolam 0.2 mg/kg plus nitrous oxide be used in children and young people under 19 years of age undergoing

diagnostic and therapeutic procedures?

Settings: dental hospital

Bibliography: Fuks 1994 (dental restorations); Fukuta 1994 (dental restorations)

			Quality assess	mont				Sumi	mary of finding	gs		
			Quality assess	sment			No of	patients	Eff	ect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	efficacy of intranasal midazolam 0.3 mg/kg plus nitrous oxide	intranasal midazolam 0.2 mg/kg plus nitrous oxide	Relative (95% CI)	Absolute	Quality	Importance
Completion	of Procedure (F	Fuks 1994 - d	cross over)									
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision ²	none	30/30 (100%)	30/30 (100%)	not estimable	1	MODERATE	
Completion	of Procedure (F	Fukuta 1994	- parallel)									
1	randomised trial	serious	no serious inconsistency	no serious indirectness	serious	none	20/21 (95.2%)	16/22 (72.7%)	RR 1.31 (1 to 1.72)	225 more per 1000 (from 0 more to 523 more) 0 more per 1,000	LOW	
Duration of p	procedure (Bett	ter indicated	by more)(Fukut		•	•					•	
1	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ^{4,5}	none	20	22	-	MD 0.60 (- 7.23 to 8.43) ^{6,7}	LOW	
 Fuks 1994: Fukuta 1994 Fukuta 1994 	not estimable, a 4: patients and a 4: imprecise 4: imprecise; ver 4: p=0.05	Il patients con essessors we	mpleted the proce re blind, ITT appe	edure		were reported; how						tated

Table 25: Safety of intranasal midazolam 0.3 mg/kg plus nitrous oxide vs. intranasal midazolam 0.2 mg/kg plus nitrous oxide

Question: What is the safety of intranasal midazolam 0.3 mg/kg plus nitrous oxide vs. intranasal midazolam 0.2 mg/kg plus nitrous oxide in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: dental hospital

Bibliography: Fuks 1994 (dental restorations); Fukuta 1994 (dental restorations)

			Quality	ecomont				Sı	ımmary of findir	ngs		
			Quality ass	essment			No of	patients	E	ffect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	adverse events of intranasal midazolam 0.3 mg/kg plus nitrous oxide	intranasal midazolam 0.2 mg/kg plus nitrous oxide	Relative (95% CI)	Absolute	Quality	Importance
Adverse	events: Vor	niting - (Fuk	s 1994)									
	randomised trial		no serious inconsistency		no serious imprecision ²	none	0/30 (0%) ²	$0/30 (0\%)^2$	not pooled	-	MODERATE	
Adverse	events: Oxy	/gen desatu	ration <90% (Fu	kuta 1994)								
1	randomised trial		no serious inconsistency	no serious indirectness	very serious ⁴	none	1/21 (4.8%)	0/22 (0%)	RR 3.14 (0.13 to 72.96) ⁵	0 more per 1000 (from 0 fewer to 0 more)	VERY LOW	
									10 / 2.00)	0 more per 1,000		
Adverse	events: Ass	sisted respir	ation (during an	d post dental tr	eatment) (Fuku	ta 1994)						
	randomised trial		no serious inconsistency		no serious imprecision ⁶	none	0/21 (0%) ²	0/22 (0%)	not pooled	-	MODERATE	
Adverse	events: Vor	niting durin	g dental procedu	•	,							
1	randomised trial		no serious inconsistency	no serious indirectness	very serious⁴	none	1/21 (4.8%)	0/22 (0%)	RR 3.14 (0.13 to 72.96) ⁵	0 more per 1000 (from 0 fewer to 0 more)	VERY LOW	
Adverse	events: Vor	miting post	dental procedure	e (Fukuta 1994)								
	randomised trial		no serious inconsistency	indirectness	no serious imprecision ⁷	none	0/21 (0%) ⁷	0/22 (0%)	not pooled	- unclear and blinding	MODERATE	

Fuks 1994: assessors were blind, ITT appeared to have been performed and no loss of follow up were reported; however, allocation concealment was unclear and blinding of patients was not stated

² Fuks 1994: no adverse events such as vomiting were observed

³ Fukuta 1994: patients and assessors were blind, ITT appeared to have been performed and no loss of follow up were reported; however, allocation concealment was no stated

⁴ Fukuta 1994: too wide confidence intervals

⁵ Fukuta 1994: p=0.48

⁶ Fukuta 1994: stated that no patients needed assisted respiration during and post dental treatment ⁷ Fukuta 1994: there were no incidents of vomiting post dental procedure in any patient in either group

Table 26: Rectal midazolam 2mg/kg vs. rectal midazolam 1mg/kg; Kanegaye, 2003 123

Question: Should rectal midazolam 2mg/kg vs. rectal midazolam 1mg/kg (with local anaesthesia in both arms) be used in children and young people under 19 years of age undergoing diagnostic and therapeutic procedures?

Settings: accidents and emergencies

Bibliography: Kanegaye 2003 (suturing for laceration repair)

			Ouglity and	accoment.				Summa	ry of finding	gs		
			Quality ass	essment			No of p	oatients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	rectal midazolam 2mg/kg plus lidocaine	rectal midazolam 1mg/kg plus lidocaine	Relative (95% CI)	Absolute	Quality	Importance
Parents'	satisfaction											
	randomised trial			no serious indirectness	serious ²	none	24/28 (85.7%)	18/26 (69.2%)	RR 1.24 (0.92 to 1.67) ³	166 more per 1000 (from 55 fewer to 464 more) 0 more per 1,000	LOW	
Total time	e: from adm	inistration to	recovery/disch	arge criteria (B	etter indicated	by less)						•
	randomised trial		no serious inconsistency	no serious indirectness	serious ⁴	none	33	32	-	MD 6 (-9.35 to 21.35) ⁵	LOW	
Recovery	(total recov	ery time fro	om completion o	f procedure to	recovery/disch	arge criteria met	Better indicated I	by less)				
	randomised trial		no serious inconsistency	no serious indirectness	serious ⁴	none	33	32	-	MD -1 (-15.21 to 13.21) ⁶	LOW	
Adverse	events: Card	dio respirato	ry complication	S								
1	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision ⁷	none	0/28 (0%) ⁷	0/26 (0%)	not pooled	-	MODERATE	
Adverse	events: Vom	niting										•
1	randomised trial			no serious indirectness	serious ⁸	none	_8	_8	not pooled8	-	LOW	

Kanegaye 2003: adequate allocation concealment, assessors were blind and some patients were blind; ITT appeared to have been performed for the outcome of recovery and total time; however for the outcome of parents' satisfaction, case analysis was available although loss of follow up was 17% (11/65: 5 in the intervention and 6 in the control groups) due lack of data collection

- ² Kanegaye 2003: imprecise
- ³ Kanegaye 2003: p=0.16
- ⁴ Kanegaye 2003: imprecise; wide confidence intervals
- ⁵ Kanegaye 2003: p=0.44
- ⁶ Kanegaye 2003: p=0.89
- Kanegaye 2003: stated that no cardio respiratory complications occurred in any patient
 Kanegaye 2003: selective outcome reporting: vomiting was part of the outcome data collected but results were not reported

6.3.2.2 Non RCT evidence profiles for safety

- Seven non RCT observational studies (n=5,412) assessed the safety of midazolam

 97,140,156,178,182,188,195. There were six prospective studies, and one retrospective study

 conducted for the following procedues: dental (1), imaging procedures (1), accidents and
 emergencies procedures (4) as well for GI procedures (1).
- 6 The non RCT study characteristics for midazolam are presented in Table 27.
- The non RCT adverse event table for midazolam is presented in Table 28.

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NON-RCT OBSERVATIONAL STUDIES FOR MIDAZOLAM

2 Table 27: Midazolam Non RCT Study Characteristics Safety Review

	dazolalii Noli Kel 3loa	,	Transfer Durery Review	<u> </u>	<u></u>	I
Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Peña 1999, USA ¹⁷⁸	pediatric emergency department for diagnostic imaging, oral and rectal sedation and analgesia. IM and IV in radiology suite	ASA I-II	described as procedural sedation and analgesia (depressed level of consciousness)	62% (733/1188)	IM ketamine+midazolam: 0.01-0.05 mg/kg IV keatmine+midazolam: 0.025-0.05 mg/kg IM or IV atropine 0.02 mg/kg	Not stated
Hulland 2002, Canada ⁹⁷	Paediatric outpatients	ASA I-III	Conscious sedation	N2O 53% (126/240) Midazolam 54% (310/579)	Oral midazolam: 0.5 mg/kg max 10 mg per appointment. mean 8.6 mg/kg Nitrous oxide/Oxygen: no higher than 70% concentration	Not stated
Pltetti 2003, USA ¹⁸²	Accidents and emergencies	81% were Class I; 17% were class II; 1.3% were class III and 0.1% were class IV.	Procedural sedation	65.1% boys in total sample (791)	IV fentanyl citrate + midazolam & IV morphine sulphate + midazolam and IV midazolam Mean fentanyl dose: 2.7 mcg /kg Midazolam 0.1 mg/kg; Morphine not stated	Mean fasting 5.0 + 2.8 hours before sedation.
Roback 2005, USA ¹⁸⁹ (*update of	Accidents and emergencies	ASA I-II	described as procedural sedation and analgesia	lv/im midazolam: 52.7% (137/260)	iv or im midazolam iv or im midazolam (0.1 mg/kg) +	of 2085 children from previous reports*

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Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Roback 2004 & follow-up from Wathen 2000)				lv/im midazolam + ketamine + glycopyrrolate: 56.9% (170/299) lv/im midazolam + fentanyl 56.8% (191/336) lv or im ketamine 63.1 (941/1492)	ketamine (1 mg/kg) iv or im midazolam + fentanyl lv or im ketamine	up to 8 hrs in 60% more than 8 hrs in 14.5% not documented in 25.4%
Mamula 2007, USA ¹⁵⁶	Operating Room	ASA I-III	Intravenous or general anaesthesia	55% (674/1226)	IV midazolam (2 mg/2mL) & fentanyl (100 mcg/2mL) during 1 minute. Midazolam 0.05 to 0.1 mg/kg max 2 mg; fentanyl 1 mcg/kg max 75 mcg Oral midazolam fo anxious patients; IV diphenhydramine as additional sedative	3 hours
Sacchetti 2007, USA ¹⁹⁵	Accidents and emergencies prospective observational database	94.1% of total cohort Class I, 5.3% class II and 0.6% class III.	Procedural sedation	Not stated	Fentanyl & Morphine	Not stated
Lightdale, 2009 USA ¹⁴⁰	outside operating room retrospective analysis of a database of clinical and adverse events records of all procedures requiring	Not stated	described as procedural sedation	56% (2,825/5,045)	IV midazolam (N=1,059) IV fentanyl (N=762) Chloral hydrate (N=604) Ketamine (N=513) Meperidine (N=21) Pentobarbital (N=2959)	Not stated

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Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
	sedation occurring outside of an operating room at a large tertiary care hospital				20% (1017/5045) had two sedatives in combination	
	82% of patients had underlying medical conditions					
	clinical and adverse events recorded by institutionally credentialed nurses					

Table 28: Midazolam Safety: Non RCTs

							AD	VERSE EVEN	ΓS, rate: %	(n)			GRADE PROFILE
Study type,	Drug /	Procedure	Age	Total N						c arrest either/or		oxygen	
reference, country	Comparison	Troccaore	Age	Total IV	Aspiratio n	oral- pharyn geal airway	endotrac heal intubatio n	assisted ventilation	external cardiac massage	defibrilla tion	vomitin g	desaturat ion <90%	EVIDENCE QUALITY
Peña, 1999 USA ¹⁷⁸	iv midazolam + fentanyl	laceration repairs, fracture	range (of 1,188 patients):	391			0%	0.51% (2/391)bag and mask			1.02% (4/391)	2.56% (10/391)	VERY LOW
	im midazolam + ketamine + atropine	reduction, CT, abscess drainage laceration repairs	median: 48 mo	180			0%	0.55% (1/180) bag and mask			0.55% (1/180)	1.11% (2/180)	VERY LOW
	iv midazolam + ketamine + atropine	fracture reduction, lumbar puncture, bone		40			0%				2.5% (1/40)		VERY LOW
	in midazolam + sufentanyl	marrow aspiration, foreign body		25			0%					4% (1/25)	VERY LOW
	inh nitrous oxide	removal, hernia reduction, arthrocentesis		168			0%	0.60% (1/168) bag and mask				0.60% (1/168)	VERY LOW
	iv fentanyl			21			0%						VERY LOW
	iv midazolam +morphine			1			0%						VERY LOW
	oral midazolam			62			0%						VERY LOW
	in midazolam			3			0%						VERY LOW

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	iv midazolam			67		0%				VERY LOW
Hulland, 2002 Canada ⁹⁷	oral midazolam	dental	range: 0.9-10.5y mean: 5.4 y	579					1.55% (9/579)	VERY LOW
	Inh nitrous oxide	dental	range: 3-14y mean: 10.8y	240 (326 sedation s)					1.54% (5/326)	VERY LOW
Pltetti 2003, USA ¹⁸²	IV fentanyl citrate + midazolam hydrochloride Vs. midazolam alone	A & E	0-21 years (of 1244 patients, mean age:6.9 (SD4.5)	686 vs 65 Total adverse events: 23.5% vs. 1.5%		0%				VERY LOW
	IV morphine sulphate + midazolam Vs. IV midazolam	A & E	0-21 years (of 1244 patients, mean age:6.9 (SD4.5)	48 vs. 65 Total adverse events: 16.7% vs. 1.5%		0%				VERY LOW
Roback et al, 2005 ¹⁸⁹	iv or im midazolam	fracture reduction, laceration repair, lumbar puncture, imaging, other	range: 42d-32y median: 4.91y	260				0.8% (2/260)		VERY LOW
	iv or im midazolam + ketamine + glycopyrrolate	fracture reduction, laceration repair, lumbar puncture,	range: 4.8mo-18y median: 6.21 y	299				5.4% (16/299)		VERY LOW

		l								
	iv or im midazolam + fentanyl	fracture reduction, laceration repair, lumbar puncture, imaging, other	range: 19d-28y mean: 7.84 y	336					1.8% (6/336)	VERY LOW
	lv or im ketamine	fracture reduction, laceration repair, lumbar puncture, imaging, other	range 39 days-22 y median 6.85y	1492					10.1% (151/14 92)	VERY LOW
Mamula, 2007 USA ¹⁵⁶	iv midazolam/fenta/ only when needed: oral Mid for anxious children & diphenhydramine to reach desired effect		0.1-34 y	1226	0% (pulmonar y aspiration)	0%	0.16% (2/1226) (bag/mask ventilation)	0% (0/1226) (cardiac arrest)	5.2% (64/122 6) (during recovery)	VERY LOW
Sacchetti 2007, USA ¹⁹⁵	Fentanyl	A & E	0-20 years	51/977 *episode of apnea with fentanyl and etomidat e which required reversal was only adverse						VERY LOW

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				event reported						
Lightdale, 2009 USA ¹⁴⁰	IV midazolam (N=1,059) IV fentanyl (N=762) Chloral hydrate (N=604) Ketamine (N=513) Meperidine (N=21) Pentobarbital (N=2959) 20% (1017/5045) had two sedatives in combination	Mixed procedures* 81% (4072/5045) underwent sedation for imaging procedures 48% (2408/5045) underwent MRI; 969 non-imaging procedures were painful and 34 nonpainful	≤30 years old Median age: 3.3 years (IQR 1.4, 6.4) with 75% of children ≤6.4 years old	&There were 329 adverse events in total	0% (serious AE)		(serious AE)	0.04% (2/5045) (serious AE) (need for resuscitati on)	5) (minor AE)	VERY LOW

ı	
2	6.3.3 Evidence statements
3	6.3.3.1 RCT efficacy and safety
4	PLACEBO COMPARISONS or NON-DRUG TREATMENT
5	Oral midazolam vs. placebo/no drug treatment
6 7 8 9 10 11 12	For the outcome of completion of procedure, we found evidence of highly significant heterogeneity ($I^2=83\%$; $p=0.02$) between two RCTs 139,168 . Possible sources of heterogeneity could be attributed to the differences between the studies in procedure performed (dental versus venous placement) and length of procedure (dental is likely to be longer), setting (outpatients versus gastroenterology) and dose [0.25 mg/kg (dental) versus 0.5 mg/kg (for intravenous placement)]. We therefore felt it was not appropriate to pool the RCTs together in a meta-analysis and the studies are presented separately for this outcome.
14	
15	Mortazavi 2009 ¹⁶⁸
16	Compared with placebo/no drug treatment, the oral midazolam group had significantly:
17	More completed procedures [low evidence quality]
18	There were no events of:
19 20	 Oxygen desaturation <90% [low evidence quality]
21	Liacouras 1998 ¹³⁹
22	Compared with placebo/no drug treatment, the oral midazolam group had significantly
23	More completed procedures [moderate quality evidence]
24	There was no significant difference in:
25 26	 Duration of procedure [the study stated that and time to discharge were not significant (data was not shown)].
27	
28	Oral midazolam vs. placebo
29	Luhman 2001 ¹⁵¹
30	There were no events of:
31	Aspiration [moderate quality evidence]

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1	 Respiratory intervention [moderate quality evidence]
2	Vomiting during procedure and recovery [moderate quality evidence]
3	There was no significant difference in:
4	Completion of procedure [moderate quality evidence]
5	
6	Fatovich 1995 ⁶³
7 8	Compared with placebo + analgesia, the oral midazolam + analgesia group were significantly:
9	 Less distressed (assessed by parents, VAS) [moderate quality evidence]
10	There was no significant difference in:
11 12	 The level of anxiety (Herbertt-Michaelinnees-Venham scale) [moderate quality evidence]
13	
14	Oral midazolam plus non-pharmacological vs. placebo plus non-pharmacological
15	Kapur 2004 ¹²⁴
16 17	Compared with placebo \pm non-pharmacological intervention, the oral midazolam \pm non pharmacological intervention had significantly:
18	More completed procedures [low quality evidence]
19	Shorter duration of procedure [very low quality evidence]
20	
21	Intranasal midazolam vs. placebo
22	Fishbein 1997 ⁶⁷
23	There was no significant difference in:
24	Distress (Observational Behaviour Rating Scale) (OBRS) [low quality evidence]
25	
26	Intranasal midazolam vs. placebo
27	Ljungman 2000 ¹⁴⁸
28 29	Compared with placebo \pm analgesia, the intranasal midazolam \pm analgesia group had significantly:

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1	• more parents who preferred midazolam + analgesia [low quality evidence]
2	There was no significant difference in:
3	• patients' preference [very low quality evidence]
4	
5	Pain assessment:
6 7 8	It was not possible to calculate the point estimate for this outcome based on the data provided. The study gave the median and interquartile ranges with the corresponding p-values indicating a source of bias (spread of skewed or non-normally-distributed data).
9	 Pain assessed by parents (VAS) [low quality evidence]
10	Pain assessed by patients (VAS) [low quality evidence]
11	
12	Theroux 1993 ²¹⁰
13 14	Compared with placebo + analgesia, the intranasal midazolam + analgesia group had significantly:
15	• more parents who felt satisfied with the treatment [very low quality evidence]
16	There were no events of:
17	Vomiting after discharge [low quality evidence]
18	
19	HEAD to HEAD COMPARISON
20	Oral midazolam vs. oral triclofos sodium
21	Singh 2002 ²⁰³
22	All patients completed the procedure [low quality evidence]
23	Compared with triclofos sodium group, the oral midazolam group had significantly:
24	Shorted induction time [low quality evidence]
25	Faster recovery time [low quality evidence]
26	
27	Sublingual midazolam vs. oral chloral hydrate
28	Layangool 2008 ¹³⁵
29	Compared to oral choral hydrate, the sublingual midazolam group had significantly:

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Longer total time [low quality evidence]
Longer fordi filite [low quality evidence]
Less vomiting [low quality evidence]
There was no significant difference in:
Completion of procedure [very low quality evidence]
Duration of procedure [low quality evidence]
Enteral midazolam vs. nitrous oxide (70%)
Zier 2008 ²³⁷
Based on the data provided, it was not possible to calculate the point estimate for the
outcome of pain assessment, parental satisfaction and total time.
Compared to nitrous oxide + placebo + topical anaesthesia + non-pharma intervention (distraction), the midazolam + placebo + topical anaesthesia + non-pharma intervention (distraction) group had significantly:
 More pain (FLACC); reported p=0.010 [moderate quality evidence]
Parents satisfaction [moderate quality evidence]
Total time [moderate quality evidence]
There was no significant difference in:
Vomiting during drug nitrous oxide administration [moderate quality evidence]
COMBINATION COMPARISONS
Oral midazolam vs. oral midazolam + nitrous oxide/oxygen
Al-zahrani 2009 ¹²
All patients completed the procedure [low quality evidence]
There was no significant difference in:
Induction time [low quality evidence]
Duration of procedure [very low quality evidence]

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1	Luhman 2001 ¹⁵¹
2	All patients completed the procedure [moderate quality evidence]
3	There were no events of:
4	Aspiration [moderate quality evidence]
5	Respiratory intervention [moderate quality evidence]
6	There was no significant difference in:
7 8	 Vomiting during visit (during procedure and after the last suture was placed) [low quality evidence]
9	
0	Oral midazolam + IV propofol vs. IV propofol
1	Paspatis 2006 ¹⁷⁶
2	Compared with intravenous propofol + lidocaine, the oral midazolam + intravenous propofol + lidocaine group had significantly:
4	• slower recovery time [moderate quality evidence]
5	There was no significant difference in:
6	Duration of procedure [low quality evidence]
7	
8	IV midazolam + IV meperidine vs. placebo + IV meperidine
9	Fishbein 1997 ⁶⁷
20	There was no significant difference in:
21 22	 Distress with major negative behaviours as assessed by an observer using the Observational Behaviour Rating Scale (OBRS) [moderate quality evidence]
23	Duration of procedure [low quality evidence]
24	
25	IV midazolam + IV propofol vs. IV propofol
26	Disma 2005 ⁵⁶
27	All patients completed the endoscopy procedure [moderate quality evidence]
28	There was no significant difference in:
29	The duration of procedure [moderate quality evidence]

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1	The recovery time [low quality evidence]
2	Assisted ventilation (bag-mask) [low quality evidence]
3	 Oxygen desaturation < 90% [low quality evidence]
4	
5	IV midazolam + IV morphine vs. IV propofol + IV morphine + local anaeshtesia
6	Havel 1999 ⁹⁰
7	All patients completed the procedure [low quality evidence]
8 9	Compared to children in the intravenous propofol group, children in the intravenous midazolam group had significantly:
10	Slower recovery time [low quality evidence]
11	Longer total time [low quality evidence]
12	There were no events of:
13	Aspiration [low quality evidence]
14	Assisted ventilation [low quality evidence]
15	Endotracheal intubation [low quality evidence]
16	There was no significant difference in:
17	• Induction time [low quality evidence]
18	Duration of procedure [low quality evidence]
19	Pain (number of patients) [very low quality evidence]
20	The groups had similar frequencies of:
21	 Hypoxemia (oxygen desaturation <90%)* [very low quality evidence]
22 23	*As stated in the study. It was not possible to calculate the point estimate for this outcome based on the information reported in the study.
24	
25	IV midazolam + IV fentanyl (analgesic) vs. IV fentanyl (analgesic)
26	Antmen 2005 ¹⁵
27	All patients completed the procedure [low quality evidence]
28	There were no events of:

1	 Oxygen desaturation < 90% [low quality evidence]
2	Vomiting [low quality evidence]
3	There was no significant different in:
4	Pain (CHEOPS) [very low quality evidence]
5	Pain (VAS) [very low quality evidence]
6	
7	IV midazolam + IV remifentanil (analgesic) IV remifentanil (analgesic)
8	Antmen 2005 ¹⁵
9	All patients completed the procedure [low quality evidence]
10	There were no events of:
11	• Oxygen desaturation < 90% [low quality evidence]
12	Vomiting [low quality evidence]
13	There was no significant different in:
14	Pain (CHEOPS) [very low quality evidence]
15	Pain (VAS) [very low quality evidence]
16	
17	IV midazolam + IV ketamine vs. IV ketamine + placebo
18	Sherwin 2000 ²⁰¹ ; Wathen 2000 ²²⁷
19	All patients completed the procedures [moderate quality evidence]
20 21	Compared with intravenous ketamine + placebo, the intravenous midazolam + intravenous ketamine group had significantly:
22	Less vomiting* [low quality evidence]
23	*during visit and at home 12 hrs after discharge ²²⁷ and well into recovery ²⁰¹
24	There was no significant difference in:
25	Assisted ventilation (bag mask) [low evidence quality]
26	There were no events of:
27	Aspiration [moderate auglity evidence]

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1	
2	Dilli 2008 ⁵⁵
3 4	Compared with intravenous ketamine + placebo, the intravenous midazolam + intravenous ketamine group had significantly:
5	Shorter induction time [very low quality evidence]
6	More satisfied parents [low quality evidence]
7	There was no significant difference in:
8	Recovery time [very low quality evidence]
9 10	Wathen 2000 ²²⁷
11	There was no significant difference in:
12	Parents' satisfaction [moderate quality evidence]
13	Duration of procedure* [low evidence quality]
14 15	*As stated in the study. It was not possible to calculate the point estimate for this outcome based on the information reported in the study.
16	There were no events of:
17	Endotracheal intubation [moderate quality evidence]
18	
19	Sherwin 2000 ²⁰¹ ; Wathen 2000 ²²⁷ ; Dilli 2008 ⁵⁵
20 21	Compared with intravenous ketamine + placebo, the intravenous midazolam + intravenous ketamine group had significantly:
22	 More oxygen desaturation < 90% [low quality evidence]
23	
24	
25	ROUTE OF ADMINISTRATION COMPARISONS
26	Oral midazolam vs. intranasal midazolam
27	Connors 1994 ⁴²
28	All patients completed the suturing procedure [moderate quality evidence]
29	

1	Everitt 2002 ⁶⁰
2	Compared with intranasal midazolam, the oral midazolam group had significantly:
3	• Lower distress scores (VAS) [very low quality evidence]
4	It was not possible to calculate the point estimate for:
5	Vomiting data not reported [very low quality evidence]
6	
7	Connors 1994 ⁴² ; Everitt 2002 ⁶⁰
8	There was no significant difference in:
9	 Total time from administration to recovery area/discharge criteria being met [very low quality evidence]
1	
2	Oral midazolam + nitrous oxide (40/45%) vs. intranasal midazolam+ nitrous oxide (40/45%)
4	Hartgraves 1994 ⁸⁹
5	There was no significant difference in:
6	The completion of procedure [low quality evidence]
7	• Oxygen desaturation < 90% [very low quality evidence]
8	
9	Lee-Kim 2004 ¹³⁷
20	Compared with intranasal midazolam, the oral midazolam group had significantly:
21	longer induction time [moderate quality evidence]
22	longer total time [low quality evidence]
23	
24	Intranasal midazolam vs. intramuscular midazolam
25	Shashikiran 2006 ¹⁹⁹
26 27	Compared with intramuscular midazolam, the intranasal midazolam group had significantly:
28	Shorter induction time [moderate quality evidence]

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I	Snorter recovery time [moderate quality evidence]
2	There were no events of:
3	Vomiting in either sedation group [moderate quality evidence]
4	
5	DOSE COMPARISONS
6 7	Intranasal midazolam 0.3mg/kg + nitrous oxide vs. intranasal midazolam 0.2 mg/kg + nitrous oxide
8	Fuks 1994 ⁷¹
9	All patients completed the procedure [moderate quality evidence]
10	There were no events of:
11	Vomiting [moderate quality evidence]
12	
13	Fukuta 1994 ⁷²
14	There were no events of:
15 16	 Assisted respiration during and post dental treatment [moderate quality evidence]
17	Vomiting post dental procedure [moderate quality evidence]
18	There was no significant difference in:
19	The completion of procedure [low quality evidence]
20	The duration of procedure [low quality evidence]
21	 Oxygen desaturation <90% [very low quality evidence]
22	Vomiting during dental procedure [very low quality evidence]
23	
24	Rectal midazolam 2mg/kg vs. rectal midazolam 1mg/kg
25	Kanegaye 2003 ¹²³
26	There were no events of:
27	Cardiorespiratory complications [moderate quality evidence]
28	It was not possible to calculate the point estimate for:

'	volining (data not reported) [low quality evidence]
2	There was no significant difference in:
3	Parents' satisfaction [low quality evidence]
4	Total time [low quality evidence]
5	Recovery time [low quality evidence]
6	6.3.3.2 Non RCT safety (adverse events)
7 8	For the characteristics of studies and outcome data on midazolam refer to Table 27 and Table 28.
9	Two studies reported rates of aspiration: 0% 140,156*
10	• Three studies reported rates of endotracheal intubation: 0% 156,178,182
11	 Two studies reported rates of assisted ventilation: from 0.16% to 0.60% 156,178
12 13	• Three studies reported rates of external cardiac massage: from 0% 156,196 to 0.02% 140*
14	 Two studies reported rates of defibrillation: from 0% ¹⁹⁶ to 0.04% ^{140*}
15	 Five studies reported rates for vomiting: from 0.55% to 5.4% 97,140,156,178,189*).
16 17	• Three studies reported rates for oxygen desaturation <90%: from 0.60% to 4% 97,156,178
18 19	 One study reported two case episodes of apnoea with fentanyl and etomidate which required reversal ¹⁹⁵
20 21	*Lightdale 2009 ¹⁴⁰ : reported adverse events were based on a total sample of 5045 patients who received treatment as follows:
22 23	 IV midazolam (N=1,059); IV fentanyl (N=762); Chloral hydrate (N=604); Ketamine (N=513); Meperidine (N=21); Pentobarbital (N=2959)
24	 20% (1017/5045) had two sedatives in combination
25	6.3.4 GDG discussion of the evidence for midazolam
26 27 28 29 30 31	In clinical practice the GDG felt that midazolam is the most common sedative drug used however there was agreement that midazolam was probably not an effective sedative drug on its own apart from achieving mildly sedative effects. Midazolam can be combined with various drugs including fentanyl, ketamine, propofol or nitrous oxide and evidence was found for these combinations. Overall the GDG felt that midazolam is a useful sedation drug and, based upon the evidence reviewed, that it is best used in combination with other drugs chosen to suite the needs of the clinical situation

2 3 4	reasons. The data derived from the studies were based upon different routes of administration together with differing drug combinations and doses. The sample sizes were small and the quality of the data was judged to be low.
5 6 7	The GDG noted that there were no UK studies and that in the UK there is a different ethos to the administration of midazolam; for example midazolam is not given intramuscularly in dentistry for children aged 1 to 5 years old.
8 9 10 11 12 13 14	Concerning the route of administration the GDG noted that oral and intranasal routes achieved a similar effect. There was no evidence comparing intravenous (IV) administration to other routes. The GDG agreed that IV drug administration acts more quickly than oral administration and once IV access is established further doses require little further cooperation unlike further doses via the intranasal or oral routes. However gaining IV access may cause distress. Overall the GDG agreed that Midazolam administered by any route helped to calm children prior to minor procedures or before the administration of more potent sedative drugs for painful procedures.
16 17 18 19	The GDG acknowledged that the safety data derived from both RCT data and observational studies showed that midazolam used on its own is a remarkably safe drug provided that doses are limited. The GDG were aware of cases of paradoxical excitement.
20 21	When considering midazolam in combination with other drugs the GDG noted that evidence was available for ketamine, opioids, nitrous oxide (N20) and propofol.
22 23	In combination with ketamine the GDG felt that the evidence demonstrated no more of an effect than for ketamine alone.
24 25 26 27	The GDG agreed that the evidence suggested that midazolam in combination with either opioids or N20 was effective for painful procedures. However when midazolam is used in combination with propofol it does not seem to result in any additional improvement in efficacy and the GDG agreed that midazolam is not necessary when using propofol.
28 29 30 31 32 33	The GDG debated vomiting as a side effect result of drug administration. It was noted that vomiting seems to be increased by approximately 10% when midazolam is combined with ketamine. No evidence was available to determine if antiemetics were effective with this drug combination. For the combination of midazolam with opioids observational data suggested that vomiting was increased by approximately 5% however the GDG felt that an antiemetic may be effective with this drug combination.
34 35 36	When midazolam is combined with either ketamine, opioids or N20 deep sedation can result and the harms of using a combined drug approach for achieving sedation in children should be weighed against benefits of relieving the pain of the procedure.
37 38 39 40 41 42 43	Combination sedation with ketamine, opioids or N20 all risk possible oxygen desaturation and the need for airway intervention. The GDG noted the small numbers when looking at the adverse event data for assisted ventilation resulting from midazolam used in combination with other sedative drugs. When combined with ketamine one case (out of 180 children) of assisted ventilation was noted, for opioids 2/391 and 2/1226 (ref 153) and for N20 one case out of 168 children resulted in assisted ventilation. The GDG did noted that there was more desaturation with the of midazolam ketamine combination than with ketamine alone.

 The general principle agreed by the GDG is that only sedation techniques commonly available in the NHS should be included in the economic analysis. Economic analysis was conducted for six broad groups (dental procedure in children, dental procedure in adolescents, short painful procedures, painless imaging, oesophago-gastroscopy and colonoscopy). Midazolam combined with fentanyl was felt to be a strategy commonly used in colonoscopy, and there is some evidence that it is effective and safe. The GDG therefore agreed that this strategy should be included in the economic analysis. Details of the considerations of cost-effectiveness with respect to using this combination strategy in this population group are given in section 6.12.3.2.

The GDG also felt that the use of midazolam alone in dental procedures in adolescents and in oesophago-gastroscopy is common, and there is some evidence on the effectiveness and safety of using midazolam alone. The GDG agreed that an economic analysis should be done on the use of midazolam alone in dental procedures in adolescents, and in children undergoing oesophago-gastroscopy. The details of the considerations of the cost-effectiveness for using this strategy in the two population groups are given in sections 6.12.4.2. and 6.12.3.2 respectively.

1 **6.4 Ketamine**

Matrix of ketamine comparators

Key:

Chloral hydrate = CH

FentanyI = F

Isoflurane = I

Ketamine=K

Local anaesthesia = LA

Midazolam = M

Nitrous oxide = N_20

Nitrous oxide and oxygen = N_20+02

Opioids = O

Propofol= P

 \dot{S}

Triclofos sodium = TS

Ketamine vs

	Reference	Tables	Evidence statements page
Placebo			
Nil			
Head to head			
Nil			
Combinations			
K + M vs M + F	Kennedy 1998 ¹²⁸ Lucas Da Silva 2007 ¹⁵⁰ Tosun 2007 ²¹³	Table 29 Table 30	195
K + M vs M	Acworth 20019	Table 31	196
K + P vs P + F	Tosun 2007 ²¹³	Table 32	196
K + M vs P + F	Godambe 2003 ⁷⁷	Table 33	197
K + M vs regional block	Kriwanek 2006 ¹³¹	Table 34	197
K + M vs N ₂ 0+haematoma block	Luhmann 2006 ¹⁵²	Table 35	197
P + F + K vs P + F	Erden 2009 ⁵⁹	Table 36	198

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Safety			
-			
RCTs			
Desaturation	Kennedy 1998 ¹²⁸ Lucas Da Silva 2007 ¹⁵⁰ Acworth 2001 ⁹ Tosun 2007 ²¹³ Godambe 2003 ⁷⁷ Erden 2009 ⁵⁹ Roback 2006 ¹⁹⁰	Table 38 Table 39	199
Vomiting	Kennedy 1998 ¹²⁸ Acworth 2001 ⁹ Tosun 2007 ²¹³ Godambe 2003 ⁷⁷ Luhmann 2006 ¹⁵² Roback 2006 ¹⁹⁰	Table 38 Table 39	199
Observational studies	McGlone 2004 ¹⁶² Sacchetti 2007 ¹⁹⁵ Roback 2005 ¹⁸⁹ Green 1998 ⁸² Green 1998 ⁸¹ Green 2001 ⁸⁰ Gilger 2004 McQueen 2009 ¹⁶³ Ramaswamy 2009 ¹⁸⁶ Thorp 2009 ²¹² Treston 2009 ²¹⁶	Table 38 Table 39	199
Route of administration			
IV / IM	Roback 2006 ¹⁹⁰	Table 37	198
Dose			
Nil			

1

6.4.1 Clinical methodological introduction

2		CLINICAL QUESTIONS:
3 4 5		For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is ketamine (with or without: analgesia, another drug or psychological techniques):
6 7 8		- effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?
9		- safe for sedation (at mild, moderate, and deep levels) in different settings?
10 11 12		The literature was searched for systematic reviews and RCTs for the clinical efficacy and safety of ketamine. The search was expanded to include non RCT observational studies for the safety of ketamine.
13 14		There were no systematic reviews identified for the use of ketamine in paediatric sedation.
15		There were no placebo controlled studies identified.
16 17		Nine RCTs comparing IV/IM ketamine with other sedative drugs and with regional anesthesia were assessed for efficacy.
18		Seven RCTs met the inclusion criteria for the review of the safety of ketamine.
19 20 21		Meta-analysis was not performed as there were no studies in which comparisons and outcome measures were sufficiently homogenous to calculate a meaningful summary statistic.
22		Eleven non RCT observational studies in a 6892 patients assessed the safety of ketamine
23	6.4.2	2 Evidence profiles
24	6.4.2	2.1 RCT evidence profiles for efficacy and safety
25 26		Study characteristics and methodological quality of the study are provided in Appendix D. GRADE tables for quality assessment and summary of findings are provided below.
27		

DRUG COMBINATION COMPARISONS

Table 29: Ketamine/midazolam vs. midazolam/fentanyl; Kennedy 1998¹²⁸

Author(s): Kennedy 1998

Question: Should ketamine/midazolam IV vs. fentanyl/midazolam be used for pediatric orthopedic emergencies?

Settings: A & E

			Quality acco	coment			Summary of findings					
			Quality asse	essment			No of pa	atients		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Ketamine/midazolam IV	Fentanyl/Midazolam	Relative (95% CI)	Absolute	Quality	ance
Completi	on of proced	dure (follow-	-up mean 121 mi	inutes)								
	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	serious ³	none	129/130 (99.2%)	127/130 (97.7%)	OR 3.05 (0.31 to 29.68)	799 more per 1000 (from 585 fewer to 1000 more)	LOW	
								0%		0 more per 1,000		
			server: VALIDAT	ED scales (follo	•	ated minutes; me	asured with: OSBD-R	score; range of score	es: 0-23.5; E	Better indicated by	less)	
	randomised trial		no serious inconsistency	no serious indirectness	serious ^{3,4}	none	130	130	i	MD -1.62 (-2.04 to -1.2)	LOW	
	Pain score- assessed by parent: VALIDATED scales (follow-up Not stated minutes; measured with: 10 point VAS; higher scores indicate greater pain; range of scores: 0-10; Bette indicated by less)										Better	
	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	serious ^{3,5}	none	130	130	i	MD 1.34 (-2.15 to 0.53)	LOW	
•	score - asses dicated by le		ent: VALIDATED	scales (follow-	up Not state	d minutes; meası	ured with: 10 point VA	S; higher scores indi	cate greate	r anxiety; range of	scores:	0-10;
	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	serious ^{3,5}	none	130	130	-	MD -1.01 (-1.8 to - 0.22)	LOW	
Induction indicated		in minutes l	between first mid	dazolam dose a	and first orth	opedic manipulat	ion (follow-up mean 1	3 minutes; measured	with: minu	tes; range of score	es: -; Be	tter
	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	serious ³	none	130	130	-	MD -0.30 (-3.1 to 2.5)	LOW	
	e: time from ; Better indic			on to when pat	ient has beer	transferred to th	ne recovery area (follo	w-up mean 120 minu	tes; measui	red with: minutes;	range o	f
	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	serious ³	none	130	130	i	MD 13.90 (2.34 to 25.46)	LOW	
Adverse	event: oxyge	en saturatio	n <90% (follow-ບ	p Throughout	sedation min	utes; Pulse oxim	etry)					
	randomised trial	serious ^{1,2}	no serious inconsistency	no serious indirectness	serious ³	none	8/130 (6.2%)	31/130 (23.8%)	OR 0.21 (0.09 to 0.48)	181 fewer per 1000 (from 113 fewer to 213	LOW	

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										fewer) 0 fewer per 1,000			
Adverse	dverse event: vomiting during sedation and recovery (follow-up during sedation and recovery minutes)												
	randomised trial			no serious indirectness	serious ³	none	9/130 (6.9%)	2/130 (1.5%)	OR 4.76 (1.01 to 22.48)	53 more per 1000 (from 0 more to 242 more) 0 more per 1,000	LOW		

The study was quasi randomised. Subjects were stratified according to initial parental choice to remain in the room or not during reduction. Subjects were then randomly assigned in blocks of 20 within strata to receive fentanyl or ketamine. A random number generator used.

The study was not fully blinded. Two trained observers were blinded to study purpose and design reviewed the videotape of each study. Unable to blind sedators. Blinding of patients and parents

was not described.

³ Small sample size

OBSD-R may be biased by subjectivity of observer
 Parental observations may be subjective and therefore biased

Table 30: Midazolam/ketamine vs. midazolam/fentanyl; Lucas Da Silva 2007 150

Author(s): Lucas Da Silva 2007

Question: Should midazaolam/ketamine IV vs.midazolam/fentanyl be used for procedural sedation for insertion of CV catheter?

Settings: In hospital CV catheter insertion

			Quality asse	sement			Summary of findings					
			Quality asse	SSIIIeIII			No of pa			Effect	Quality	Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Midazaolam/ketamine IV	Midazolam/fentanyl	Relative (95% CI)	Absolute		ance
Completi	ion of proced	dure (follow-	up mean 101 mi	nutes)								
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	29/29 (100%)	28/28 (100%)	RR 1 (0 to 0)	0 fewer per 1000 (from 1000 fewer to 1000 fewer) 0 fewer per 1,000	LOW	
Recovery	y time: Time	elapsed froi	m end of proced	ure to awakenii	ng (follow-up	median 20 minu	tes; measured with: mi	nutes; range of sco	res: -; Better	r indicated by less	5)	
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	29	28	-	-5.0 (-15 to 7.9)	LOW	
	e: Time elap l by less)	sed from ini	tial sedative adn	ninistration to s	spontaneous	eye opening (fol	low-up median 101 min	nutes; measured with	n: minutes;	range of scores: -	; Better	
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	29	28	=	6.5 (-19 to 33)	LOW	
	n time: Time I by less)	elapsed froi	n initial sedative	administration	to onset of	the procedure (fo	ollow-up median 7.5 mi	nutes; measured wit	h: minutes;	range of scores:	-; Better	٢
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	29	28	-	2 (-0.002 to 5.998)	LOW	
Adverse	event: oxyge	en saturation	n <90% (follow-u	p median 101 r	ninutes; Pul	se oximeter)						
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	2/29 (6.9%)	0/28 (0%)	OR 5.18 (0.24 to 112.89)	0 more per 1000 (from 0 fewer to 0 more) 0 more per 1,000	LOW	

Double blinding was deemed impractical because of different dosing algorithms of the drugs used and because medications used present clinically distinguishable effects. ² Small sample size

³ Recovery time, induction time and total times were reported as median differences

Author(s): Acworth 2001

Question: Should intravenous ketamine plus midazolam vs. intranasal midazolam be used for emergency paediatric procedural sedation?

Table 31: Ketamine + midazolam vs. intranasal midazolam; Acworth 20019

Settings: A & E

			Quality asse	eemont			Summary of findings					
			Quality asse	SSIIICIIL			No of pa	tients		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	IV ketamine plus midazolam	intranasal midazolam	Relative (95% CI)	Absolute	Quality	ance
Completion	on of proced	ure (follow-ເ	ıp mean 88 minut	es)								
	randomised trial		no serious inconsistency	no serious indirectness	serious	none	26/26 (100%)	24/26 (92.3%)	OR 5.4 (0.25 to 118.34)	1000 more per 1000 (from 618 fewer to 1000 more) 0 more per 1,000	LOW	
Induction indicated		om adminis	tration of sedatio	n until sedation	score reache	ed 3 or less) (follo	w-up mean 5 minu	ıtes; measured	d with: minute	s; range of scores: -; E	Better	
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	26	26	-	MD 5.32 (3.2 to 7.4)	LOW	
	e: timing - tot Better indica			of intervention to	when patier	nt met all the crite	ria for discharge	follow-up mea	n 88 minutes;	measured with: minut	es; ranç	ge of
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	26	26	-	MD -18.9 (-33.4 to -4.4)	LOW	
Adverse e	events: oxyge	en saturatio	n <90% (follow-uj	mean 88 month	ns)							
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	1/26 (3.8%)	0/26 (0%)	OR 3.12 (0.12 to 80.12)	0 more per 1000 (from 0 fewer to 0 more) 0 more per 1,000	LOW	
Adverse e	event: vomiti	ng during pr	rocedure		•						•	
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	0/26 (0%)	1/26 (3.8%)	OR .32 (0.01 to 8.24)	26 fewer per 1000 (from 38 fewer to 211 more) 0 fewer per 1,000	LOW	

Drug route precluded double blinding and allocation concealment but the doctor and nurse responsible for scoring sedation level were not present during drug administration and were blinded to allocation by use of dummy armboard applied to children receiving the intranasal medication

The sample size was only 26 in each arm

Table 32: Ketamine + propofol vs. propofol + fentanyl; Tosun 2007 213

Author(s): Tosun 2007

Question: Should intravenous ketamine plus propofol vs. propofol plus intravenous fentanyl be used in children undergoing upper GI endoscopy?

Settings: Gastroenterology

			Quality acc	acemont			Summary of findings					
			Quality ass	essment			No of p	atients		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	ketamine IV plus propofol	propofol plus fentanyl IV	Relative (95% CI)	Absolute	Quality	ance
Completion	on of proced	ure (follow-	up mean 116 min	utes(time to dis	charge))							
	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	46/46 (100%)	44/44 (100%)	RR 1 (0 to 0)	0 fewer per 1000 (from 1000 fewer to 1000 fewer)	MODERATE	
								0%		0 fewer per 1,000		
Pain (Nun	nber of patie	nts who nee	eded additional p	ropofol during i	nduction as evi	denced by discor	nfort/moving d	uring procedu	re (follow-up	0-1 minute after inde	uction)	
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	8/46 (17.4%)	22/44 (50%)	RR 0.35 (0.17 to 0.7)	325 fewer per 1000 (from 150 fewer to 415 fewer)	LOW	
								0%		0 fewer per 1,000		
Pain (Nur	nber of patie	nts who nee	eded additional p	ropofol during a	as evidenced by	discomfort/mov	ing during pro	cedure)				
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	32/46 (69.6%)	41/44 (93.2%)	RR 0.75 (0.61 to 0.92)	233 fewer per 1000 (from 75 fewer to 363 fewer)	LOW	
								0%	Ì	0 fewer per 1,000		
Recovery	time (time fi	rom comple	tion of procedure	to recovery/dis	charge criteria	being met) (follow	w-up mean 4.5	minutes; rang	e of scores:	; Better indicated by	/ less)	
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	46	44	-	MD 1.60 (-0.42 to 3.62)	LOW	
Adverse 6	events: oxyg	en saturatio	n <90% (follow-u	p mean 116 mir	nutes; Pulse ox	imetry)						
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	3/46 (6.5%)	4/44 (9.1%)	OR 0.73 (0.15 to 3.47)	23 fewer per 1000 (from 76 fewer to 174 more)	LOW	
								25%	ĺ	58 fewer per 1,000		
Adverse 6	events: vomi	ting (follow-	up mean 116 mir	nutes, observati	on)	,	•		•		•	
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	7/46 (15.2%)	0/44 (0%)	OR 16.9 (0.93 to 305.47)	0 more per 1000 (from 0 fewer to 0 more)	LOW	
								0%	303.47)	0 more per 1,000		

Unclear allocation concealment;small trial, total n=90;no loss to follow up; double blind Wide confidence interval; few events

Table 33: Ketamine/midazolam vs. propofol/fentanyl; Godambe 2003 77

Author(s): Godambe 2003
Question: Should ketamine/midazolam vs. Propofol/Fentanyl be used for Procedural Sedation?
Settings: Pediatric Emergency Department

			Ouglity sass	oom on t			Summary of findings					
			Quality asse	ssment			No of pa	tients		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Ketamine/Midazolam	Propofol/Fentanyl	Relative (95% CI)	Absolute	Quality	ance
Complet	ion of proced	lure										
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	50/54 (92.6%)	53/59 (89.8%)	OR 1.42 (0.38 to 5.31)	226 more per 1000 (from -472 fewer to 1000 more)		
Dagayar	time leet d	ooo of modi	aatian ta vatuun t	e bassline /fell	our up time t	a return to bood!	no minutos, mossuro	0%		0 more per 1,000	d by los	201
Recover	1			,			ne minutes; measure	a with: minutes; rai	ige of score	,	a by les	<i>(S)</i>
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	54	59	-	MD 33.4 (26.1 to 40.8)	LOW	
	e; from first dicated by le		lication to return	to baseline (fo	llow-up Tota	I time from begin	ning of sedation to re	covery minutes; me	easured with	n: minutes; range o	f scores	: -;
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	54	59	-	MD 23.2 (15.4 to 30.4)	LOW	
Adverse	events: vom	iting (follow	-up Immediate ad	dverse effects r	ninutes)							
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	2/54 (3.7%)	0/59 (0%)	OR 5.67 (0.27 to	0 more per 1000 (from 0 fewer to 0 more)	LOW	
								0%	120.73)	0 more per 1,000		
Adverse	outcome: ox	ygen satura	tion <90% (follow	v-up Any amou	nt of time du	ring procedure a	nd recovery minutes)				•	
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	4/54 (7.4%)	18/59 (30.5%)	OR 0.18 (0.06 to 0.58)	240 fewer per 1000 (from 113 fewer to 283 fewer)		
								0%	0.36)	0 fewer per 1,000		
Pain sco	re - assessed	d by parent:	VALIDATED sca	les (measured	with: VAS sc	ore, range of sco	res: 0mm-100mm; Be	tter indicated by le	ss)			
1	randomised trial		no serious inconsistency	no serious indirectness	serious ^{2,3}	none	30	38	-	MD 4.30 (-5.28 to 13.88)	LOW	
Distress	score- asses	sed by obse	erver - VALIDATE	D scales (follo	w-up Vidoe t	apes assessed at	fter procedure minute	s; range of scores:	0-23.5; Bett	er indicated by les	s)	
1	randomised trial		no serious inconsistency	no serious indirectness	serious ^{2,4}	none	54	59	-	MD -0.19 (-0.39 to 0)	LOW	

Quasi randomised - Odd or even day assignment Small sample size

Assessment by parents may be subjective and therefore biased
 There is potential for the OSBD to be subjective and therefore biased

Table 34: Ketamine/midazolam vs. axillary block regional anesthesia (intra arterial block); Kriwanek 2006131

Author(s): Kriwanek 2006

Question: Should ketamine plus midazolam vs. axillary (brachial plexus) block regional anesthesia be used for forearm fracture in children? Settings: Pediatric Emergency Department

			Quality asse	eemont				Summary of find	ings			
			Quality asse	SSIIIEIII				No of patients		Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	IV Ketamine +IV midazolam Axillary (brachial plexus) blo regional anesthesia(ABRA (intra arterial lidocaine + epinephrine block)		Relative (95% CI)	Absolute	Quality	Import ance
Complet	ion of proce	dure		•							-	
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	21/21 (100%)	18/20 (90%)	OR 5.81 (0.26 to 128.9)	1000 more per 1000 (from 593 fewer to 1000 more)	LOW	
								0%		0 more per 1,000		
Pain -sc	ore - assesse	ed by patien	t: VALIDATED s	cales (measur	ed with: FPS	-R ; range not pr	ovided; range	of scores: -; Better indicated by I	ess)			
1	randomised trial			no serious indirectness	serious ^{2,3}	none	21	20	-	MD 0.90 (-0.27 to 2.07)		
Pain -sc	ore - assesse	ed by obser	ver: VALIDATED	scales (meas	ured with: Cl	HEOPS during fr	acture reduction	on; range of scores: 4-13; Better i	indicated b	y less)		
1	randomised trial			no serious indirectness	serious ^{2,3}	none	21	20	-	MD 1.10 (-0.31 to 2.51)		

¹ Blinding not possible and allocation concealment not described.
² Small sample size
³ Pain scales have potential for subjective interpretation and therefore bias

Table 35: Ketamine + midazolam vs. nitrous oxide + hematoma block; Luhmann 2006¹⁵²

Author(s): Luhmann 2006

Question: Should ketamine plus midazolam vs. nitrous oxide plus hematoma block be used for forearm fracture reduction in children?

Settings: Emergency Department

			Quality assa	comont				Summ	ary of finding	js		
			Quality asse	ssment			No of	patients		Effect		Importan
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Ketamine +Midazolam	Nitrous Oxide + Hematoma Block	Relative (95% CI)	Absolute	Quality	ce
Completi	ion of proced	ure (follow-ເ	ıp mean 50 minut	tes)								
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	55/55 (100%)	47/47 (100%)	RR 1 (0 to 0)	0 fewer per 1000 (from 1000 fewer to 1000 fewer)	LOW	
								0%		0 fewer per 1,000		
Adverse	event: vomiti	ng (follow-u	p mean 50 month	ıs)								
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	13/55 (23.6%)	12/47 (25.5%)	OR 0.90 (0.37 to 2.23)	22 fewer per 1000 (from 149 fewer to 216 more) 0 fewer per 1,000	LOW	
	time (follow; Better indicate			sured with: P val	lue reported f	for mean difference	e of 83 minutes	for KM group and	16 minutes f	or NO/HB group: p<0	.0001; ra	ange of
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	55	47	-	not pooled ⁶	LOW	
Distress	during proce	dure (measu	red with: OR of N	ID reported: OR	0.9 (95% CI	0.5-2.1); range of	scores: 5-25; B	etter indicated by le	ess)		•	•
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ^{2,3}	none	55	47	-	not pooled ⁷	LOW	
	ported by pati	ient (follow-	up mean 49.5 mir	utes; measured	with: OR of	mean difference i	n VAS scores re	eported: OR 1.1 (95	% CI 0.0-2.1);	range of scores: 1-1	0; Bette	r
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ^{2,4}	none	55	47	-	not pooled ⁷	LOW	
	ported by par- dicated by les		rocedure (follow-	up mean 49.5 m		sured with: OR of	mean difference	e in VAS scores rep	oorted: OR 1.	6 (95% CI 0.6-2.6); rar	nge of so	cores:
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ^{2,5}	none	55	47	_	not pooled ⁷	LOW	

¹ Single blinding only

Single billing Siny

2 Small sample size

3 Distress scale has potential for subjectivity and therefore bias

4 Pain as assessed by patient is inherently subjective and therefore subject to bias

⁵ Pain as assessed by parent is inherently subjective and therefore subject to bias ⁶ Mean recovery times reported in minutes with no SDs reported. Recovery time was significantly shorter for children who received NO/HB vs. KM,p<0.0001

⁷ Distress and pain scores were reported as OR for the mean difference between K/M and NO/HB. SDs for means were not provided. OR for distress score during procedure as measured by PBCL was OR 0.9 (-0.4-2.2); OR for pain as reported by patient was OR1.1 (0.00-2.1); OR for pain score during procedure as reported by parent was 1.6 (0.6-2.6)

Table 36: Propofol-fentanyl + ketamine vs. propofol-fentanyl; Erden 2009 59

Author(s): Erden **Date:** 2010-01-05

Question: Should ketamine plus propofol-fentanyl vs. propofol-fentanyl be used in paediatric sedation? **Settings:** Interventional radiology

			Quality assess	emont								
			Quality assess	Silielit			No of patients Effect					Importan
No of studies	Design	Limitations	mitations Inconsistency Indirectness Imprecision Other considerations Relative fentanyl Fentanyl Relative (95% CI)		Absolute	Quality	ce					
Oxygen s	aturation <90)%	•									
1	randomised trial			no serious indirectness	serious ¹	none	3/30 (10%)	9/30 (30%)	RR 0.33 (0.10 to 1.11)	201 fewer per 1000 (from 270 fewer to 33 more) 0 fewer per 1,000	MODERATE	

¹ Sample size small and characterised as 'about' 30 patients for each group would be sufficient to detect a fall from 30% to 5%

ROUTE OF ADMINISTRATION COMPARISONS

Table 37: Intravenous ketamine vs. intramuscular ketamine; Roback 2006¹⁹⁰

Author(s): Roback 2006

Question: Should intravenous ketamine vs. intramuscular ketamine be used for sedation of pediatric patients?

Settings: Emergency Department Orthopedic Procedures

			Quality asse	coment					Summary of f	indings		
			Quality asse	ssment			No of p	atients		Effect		Importa
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	IV ketamine	IM ketamine	Relative (95% CI)	Absolute	Quality	nce
Completion	on of procedu	re (follow-up	median 13.0 minu	tes)								
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	101/109 (92.7%)	95/99 (96%)	OR 0.53 (0.15 to 1.82)	358 fewer per 1000 (from 760 fewer to 416 more) 0 fewer per 1,000	LOW	
Adverse e	vents: oxyger	n saturation	<90% (Pulse oxim	etry)								
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	9/109 (8.3%)	4/99 (4%)	OR 2.14 (0.64 to 7.17)	42 more per 1000 (from 14 fewer to 193 more) 0 more per 1,000	LOW	
Adverse e	vent: vomitin	g	'		•	'	*	!				
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	13/109 (11.9%)	26/99 (26.3%)	OR 0.32 (0.18 to 0.79)	168 fewer per 1000 (from 47 fewer to 208 fewer) 0 fewer per 1,000	LOW	
Pain scor	e - number of	patients - as	sessed by patient:	VALIDATED sca	les	<u> </u>	<u> </u>		<u> </u>	0 10 mer per 1,000	<u> </u>	
1		serious ¹	no serious inconsistency	no serious indirectness	serious ^{2,3}	none	49/84 (58.3%)	57/70 (81.4%)	OR 0.32 (0.15 to 0.67)	fewer)	LOW	
					L . ,			. ,		0 fewer per 1,000	<u> </u>	
Distress -					serious ^{2,4}	e of scores: -; Bett	ter indicated	by less)	I			l
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious	none	97	93	-	MD 0.47 (0.13 to 0.82)	LOW	
						nsferred to the rec r indicated by less		ollow-up me	edian 104.5 mi	nutes; measured with:	Reported	d as
1			no serious inconsistency	no serious indirectness	serious ²	none	109	99	-	not pooled ⁵	LOW	

¹ Single blinding only
² Small sample size
³ The FACES scale is a subjective measurement and is subject to bias
⁴ OBSD scale has potential to be subjective and therefore biased
⁵ Total time reported as median minutes in IM vs. IV; p<0.001

6.4.2.2 Non RCT evidence profiles for safety

2	Eleven non RCT observational studies in 6892 patients assessed the safety of ketamine
3	76,80-82,162,163,186,189,195,212,216. There were six prospective reviews and five retrospective
4	studies conducted primarily for emergency procedures (9) as well as studies of ketamine
5	for gastrointestinal (GI) procedures.

- 6 The non RCT study characteristics for ketamineare presented in Table 38.
- 7 The non RCT adverse event table for ketamine is presented in Table 39.

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1 Table 38: Ketamine Non RCT study characteristics safety review

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Prospective Cohort						
McGlone et al, 2004 ¹⁶² UK	Lancaster Royal Infirmary, Lancaster, UK Accident and emergency department		IM ketamine sedation for minor painful procedures		IM ketamine: 301 children received 2.0 mg/kg and 191 received 2.5 mg/kg; 26 children received a second dose.	
Sacchetti et al, 2007 ¹⁹⁵ USA Results from ProSCED Registry for Ketamine	14 community emergency departments	321 (94.1%) were ASA I, 18 were ASA class II (5.3%) and 2 were ASA class III (0.6%)	41.3% received ketamine – route of delivery not described			
McQueen et al, 2009 ¹⁶³	A children's hospital emergency department, USA		66% (363) received ketamine alone; 19% (106) received ketamine/midazolam; 15% (85) received non- ketamine drugs	62% (341) were male; 38% (213) were female		

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Ramaswamy et al, 2008 ¹⁸⁶	Royal Children's Hospital Melbourne, Australia ED		Ketamine IM or IV	138 male (60.3%)	Ketamine 3-4 mg/kg IM or 1-1.5 mg/kg IV	
Thorp et al, 2009 ²¹²	Pediatric Emergency Department, Loma Linda University Medical Center andChildren's Hospital, Loma Linda, California USA	ASA I 93% (959); ASA II 6% (66); ASA III 1% (14)	Ketamine	62% (649) male	Ketamine initial dose (0.2-2.4 mg/kg) and total dose (0.3 to 23.8 mg/kg)	
Treston et al, 2009 ²¹⁶	Redcliffe Hospital Brisbane, Australia		Ketamine for minor procedures or		Ketamine from 0.23 to 3.8 mg/kg (mean 1.15	

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
	ED		examination		mg/kg). Titrated IV ketamine used in 691 cases and IM in 54 cases	
Roback et al, 2005 ¹⁸⁹ USA	Paediatric Emergency Department		IV or IM procedural sedation		Ketamine alone; ketamine/midazolam	
Green et al, 1998 ⁸² USA	Emergency Department		IM Ketamine		Ketamine 4 mg/kg combined with atropine .01mg/kg IM; repeat ketamine dose (2-4 mg/kg) without atropine if required	Children who had eaten a full meal within 3 hours were excluded but not those with lesser degrees of oral intake
Green et al, 1998 ⁸¹ USA	Emergency Department		IV Ketamine		The mean loading dose of ketamine was 1.5 + 0.5 mg/kg and was then titrated as necessary. The total mean dose used was 2.5 + 1.6 mg/kg.	Children who had eaten a full meal within 3 hours were excluded but not those with lesser degrees of oral intake
Green et al, 2001 ⁸⁰ USA	University medical centre - Department of Gastroenterology	Ketamine administered at all levels of ASA stratification	IV Ketamine: 98.3% of patients and IM Ketamine: 1.7% of patients Concurrent midazolam was administered in 97% (614) of patients	54.4%	The median IV loading dose of ketamine was 1.00 mg/kg and titrated if necessary. The median total IV dose was 1.34 mg/kg.	

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Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Gilger et al, 2004 ⁷⁶ USA	Children's Hospital: endoscopy		Ketamine + midazolam; Ketamine + midazolam + meperidine	48% male in ketamine/midazolam group; 63% male in the Ketamine + midazolam + meperidine group	Ketamine 0.75-2.0 mg/kg dose	

-

1 Table 39: Ketamine Safety: Non RCTs

		ety: Non RCTs						ADVE	RSE EVENT	S, rate: %	o (n)			GRADE PROFILE
Study type, reference,	Drug / Comparison	Procedure	Age	Total N	N A	Respir	atory inte	ervention	Cardiac arrest requiring either/or			oxygen	D	F. dana
country	Companison				Aspirati on	oral- pharyn geal airway	endotr acheal intubat ion	assisted ventilatio n	external cardiac massage	defibril lation	vomiting	ion <90%	Recovery agitation	Evidence quality
Prospective C	Cohort studies													
McGlone et al, 2004 ¹⁶² UK	IM Ketamine	Injuries in A&E requiring wound toilet and suturing, minor surgery such as nail bed repair, and removal of foreign bodies	Not stated	501							17% (in recovery or at home) (85/501)	.5% (3/501))	Mild: 15% (71/501) Moderat e: 3% 16/501 Pronounc ed: 0.8% (4/501)	VERY LOW
Sacchetti et al, 2007 ¹⁹⁵ USA Results from ProSCED Registry for Ketamine	Ketamine	Minor trauma including laceration repairs, foreign body removal, fracture care, join relocation and also lumbar puncture, radiology, tube thoracostomy and cardioversion	Ages 0- 20 years	This registry reports a total of 1028 procedur al sedations. 141 children received ketamine										VERY LOW
McQueen et	Ketamine	Emergency	3 months	422							25/422			VERY

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Study type, reference,	Drug / Comparison	Procedure	Age	Total N		ADVER	SE EVENTS,	rate: % (n)			GRADE PROFILE
al, 2009 ¹⁶³		Department procedures	-18 years					(5.9%) Before discharge			LOW
	Ketamine/mi dazolam			123				13/123 (10.5%) before discharge			VERY LOW
Ramaswamy et al, 2008 ¹⁸⁶	IM Ketamine vs. IV ketamine	Emergency Department procedures	1.8-4.3 years	229 total; IM, n= 110; IV, n= 119.				IM: 17.3% (95% CI = 10.7% to 25.7%) vs. IV: 11.8% (95% CI = 6.6% to 18.9%); P=0.24			VERY LOW
Thorp et al, 2009 ²¹²	Ketamine	Emergency Department procedures	No emesis: 6.1 years median age; With emesis 9.8 years median age	1039				Rate of emesis was 7.0% when the total dose was 7 mg/kg or less and 11.1% when greater than 7 mg/kg			VERY LOW
Treston et al, 2009 ²¹⁶	Ketamine	Emergency Department procedures	12 months – 13 years	745						16/745 (2,1%)	VERY LOW
Retrospective	,										
Roback et al, 2005 ¹⁸⁹	Ketamine	Fracture reduction;	39 days to 22	1,492					6.1% 91/1492		VERY LOW

Study type, reference,	Drug / Comparison	Procedure	Age	Total N	ADVER	SE EVENTS, rate: %) (n)			GRADE PROFILE
USA		laceration repair; lumbar puncture; imaging; other dental	years; median age 6.58 years					Includes oxygen saturation >90% and laryngos pasm		
	Ketamine/mi dazolam		4.8 mo to 18 y; median age 6.21 years	299				10%30/ 299 Includes oxygen saturation >90% and laryngos pasm		VERY LOW
Green et al, 1998 ⁸² USA	IM Ketamine	Emergency procedures including wound and dermal repair, orthopaedic, GU, GI eye procedures and line placement, lumbar puncture, CT scan chest tube and ET tube placement	0-15 years	1,022	.4% (5/1022) Bag mask ventilatio n		6.7% (68/1022)	.9% (9/1022)	Total events by chart document ation and as assed by physician: 19.3% (197/10 22) Moderat e to severe: 1.6% (16/102 2)	VERY LOW
Green et al, 1998 ⁸¹ USA	IV Ketamine 31% received	Emergency procedures including	0-15 years	156	.6% (1/156) Bag mask		3.8% (6/156) 1 while	.6% (1/156)	Total events by chart	VERY LOW

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Study type, reference,	Drug / Comparison	Procedure	Age	Total N	ADVERS	SE EVENTS, rate: %	(n)		GRADE PROFILE
	concurrent midazolam	wound and dermal repair, orthopaedic, GU, GI eye procedures and line placement, lumbar puncture, CT scan chest tube and ET tube placement			ventilatio n		sedated and 5 in recovery	document ation: Mild:1.3 % (2/156) Moderte to severe: 0	
Green et al, 2001 ⁸⁰ USA	IV Ketamine: 98.3% of patients and IM Ketamine: 1.7% of patients Concurrent midazolam was administered in 97% (614) of patients 15% of patients received other sedatives: meperidine (n=90), diazepam (n=4) and morphine	GI procedures	Median age 5.2 years	636 46% of patients had severe underlyin g illness (ASA >3)	3% (19/636) Bag mask ventilatio n		4.1% (26/636)	1.4% (9/636) mild .9% Moderat e to severe	VERY LOW

Study type, reference,	Drug / Comparison	Procedure	Age	Total N	ADVERSE EVENTS, rate: % (n)								GRADE PROFILE	
	(n=3)													
Gilger et al, 2004 ⁷⁶ USA	Ketamine + midazolam	GI endoscopy	5.9 years mean age (SD 4.77)	128	0						0	*data recorded was oxygen saturation <95%	0	VERY LOW
	Ketamine + midazolam + meperidine		7.68 years mean age (SD 4.22)	82				1.2% (1/82)			0	*data recorded was oxygen saturation <95%	0	VERY LOW

iv= intravenous; in= intranasal; im= intramuscular; inh= inhaled

1	6.4.3 Evidence statements
2	6.4.3.1 RCT efficacy and safety
3	DRUG COMBINATION COMPARISONS
4	Ketamine/midazolam/ vs. midazolam/fentanyl
5	Kennedy 1998 ¹²⁸
6 7	Compared with midazolam + fentanyl, the midazolam + ketamine group had significantly:
8	 Less distress on OSBD [Low quality evidence]
9 10	 Less anxiety as reported by parent on VAS after procedure [Low quality evidence]
11	• Less pain as reported by parent on VAS after procedure [Low quality evidence]
12	Longer total time [Low quality evidence]
13	 Less oxygen desaturation (O₂ saturation <90%) [Low quality evidence]
14	 More vomiting during recovery; p=.03 [Low quality evidence]
15	There was no significant difference in:
16	Completion of procedure [Low quality evidence]
17	Length of induction [Low quality evidence]
18	Vomiting during procedure [Low quality evidence]
19	Valve-mask ventilation [Low quality evidence]
20	
21	Midazolam/ketamine vs. midazolam/fentanyl
22	Lucas Da Silva 2007 ¹⁵⁰
23 24	Median results were reported on this RCT. It was not possible to combine these results with other studies for meta-analysis.
25 26	Compared with midazolam + fentanyl, the midazolam + ketamine group had significantly:
27	Shorter induction time [Low quality evidence]
28	There was no significant difference in:

1 2	 Completion of procedure – all procedures were completed [Low quality evidence]
3	Recovery time [Low quality evidence]
4	Total sedation time [Low quality evidence]
5	 Oxygen saturation <90% [Low quality evidence]
6 7	It was stated that neither cardiac rhythm abnormalities nor increase in cardiac rate were detected.
8	
9	Midazolam + ketamine vs. intranasal midazolam
10	Acworth 20019
11 12	Compared with midazolam + ketamine, the intranasal midazolam group had significantly:
13	Shorter induction time [Low quality evidence]
14	Longer total time [Low quality evidence]
15	There was no significant difference in:
16 17	 Completion of procedure – all procedures were completed [Low quality evidence]
18	 Oxygen saturation <90% [Low quality evidence]
19	Vomiting [Low quality evidence]
20	
21	Propofol-ketamine vs. propofol-fentanyl
22	Tosun 2007 ²¹³
23	All patients completed the procedure [Moderate quality evidence]
24	Compared with Propofol-fentanyl, the propofol-ketamine group had significantly:
25 26	 Less pain as measured by the number of patients requiring additional propofol in the first minute after induction [Low quality evidence]
27	More vomiting [Low quality]
28	There was no significant difference in:
29	 Length/duration of procedure [Low quality evidence]

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1	Recovery time [Low quality evidence]
2	 Oxygen saturation <90% [Low quality evidence]
3	
4	Ketamine + midazolam vs. propofol + fentanyl
5	Godambe 2003 77
6	Compared with propofol + fentanyl, the midazolam + ketamine group had significantly
7	 Less distress on OSBD [Low quality evidence]
8	• Less pain as reported by parent on VAS after procedure [Low quality evidence]
9 10	 Less anxiety as reported by parent on VAS after procedure [Low quality evidence]
11 12	 Less oxygen desaturation (O₂ saturation <90%) [Low quality evidence] More vomiting during recovery [Low quality evidence]
13	Longer total time [Low quality evidence]
14	There was no significant difference in:
15 16	 Completion of procedure – all procedures were completed [Low quality evidence]
17	Length of induction [Low quality evidence]
18	
19	Ketamine + midazolam vs. axillary block regional anaesthesia
20	Kriwanek 2006 ¹³¹
21	There was no significant difference in:
22 23	 Completion of procedure – all procedures were completed [Low quality evidence]
24	Pain assessed by patient using FPS-R [Low quality evidence]
25 26	 Distress during the procedure as measured by CHEOPS scale, [Low quality evidence]
27	
28	Ketamine + midazolam vs. nitrous oxide + haematoma block
29	Luhmann 2006 ¹⁵²

1 2	Compared with nitrous oxide + haematoma block, the ketamine + midazolam group had significantly:
3 4	 Longer recovery time (from cast molding to Aldrete score of 10) [Low quality evidence]
5	There was no significant difference in:
6 7	 Completion of procedure – all procedures were completed [Low quality evidence]
8	Distress as assessed by PBCL score [Low quality evidence]
9	 Pain assessed by patient using VAS [Low quality evidence]
10	Pain assessed by parent using VAS [Low quality evidence]
11	Vomiting [Low quality evidence]
12	
13	Propofol-fentanyl vs. propofol-fentanyl-ketamine
14	Erden 2009 ⁵⁹
15 16	Compared with Propofol-fentanyl, the propofol-fentanyl-ketamine group required significantly:
17	Less supplemental propofol [Moderate quality evidence]
18	There was no significant difference in:
19	 Oxygen saturation <90% [Moderate quality evidence]
20	
21	ROUTE OF ADMINISTRATION COMPARISONS
22	Intravenous ketamine vs. intramuscular ketamine
23	Roback 2006 ¹⁹⁰
24	Compared with ketamine IM, the ketamine IV group had significantly:
25 26	 Less distress during the procedure as measured by CHEOPS scale [low quality evidence]
27	Less total time [low quality evidence]
28	Less vomiting [low quality evidence]
29	There was no significant difference in:

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 Oxygen saturation <90% [low quality evidence]
 Pain assessed by patient using FPS-R [low quality evidence]
• Parental satisfaction assessed on 7 point Likert scale [low quality evidence]
6.4.3.2 NON-RCT safety (adverse events)
For the characteristics of studies and outcome data on ketamine refer to Table 38 and Table 39.
• Four studies reported rates of assisted ventilation: 1.2% 76 , ; 0.6% 82 ; 0.4% 81 ; 3% 80
There were no cardiac events reported in 11 studies.
• Vomiting was reported in nine studies ^{76,80-82,162,163,186,212,216} . and rates ranged from 0% ⁷⁶ , to17% (¹⁶² . The mean vomiting rate for the nine studies was 7.9% A dose response effect was noted in one study ²¹² where the rate of emesis was 7.0% when the total dose was 7 mg/kg or less and 11.1% when greater than 7 mg/kg. A non significant difference was noted between IV and IM routes ¹⁸⁶ .
• Oxygen saturation <90% was reported in five studies 81,82,130,162,186,189 . and rates ranged from 0.5% 162 to 10% 189 . The mean desaturation rate for five studies was 3.8% .
 Recovery agitation was reported in seven studies and was classified as mild, moderate and severe. Definitions of these classifications were not standardised. Mild recovery agitation ranged from 1.3⁸¹, -15% ¹⁶²,; moderate to severe recovery agitation ranged from 0% ⁸¹, to 1.6% ⁸¹.
6.4.4 GDG discussion of the evidence for Ketamine
The GDG noted that out of 16 studies considered 11 were in patients undergoing painful procedures in the Emergency Department (ED) setting. One study was in children undergoing painful insertion of central intravenous catheters and the remainder were in children undergoing gastrointestinal endoscopy.
The GDG discussed four studies ¹⁹⁰ , ⁸² , ¹⁶² , ¹⁹⁵ in which ketamine was used alone in the ED setting. Only one of these studies ¹⁹⁰ was an RCT and it compared IV with IM ketamine. The quality of the evidence was low yet, together with the three large non-RCTs ⁸² , ¹⁶² , ¹⁹⁵ , the GDG agreed that there was much evidence to show that ketamine was effective over a wide range of painful procedures.
Discussions highlighted the difficulty of research in this area. The main problem was that any sedation technique being compared with ketamine would need to be of a similar efficacy. That there were so few studies may indicate that few sedation techniques are as effective as ketamine. The GDG thought that combinations of drugs such as midazolam and fentanyl were potentially as effective.

• Completion of procedure [low quality evidence]

 In the RCT of ketamine alone¹⁹⁰ the GDG noted that the evidence of efficacy was limited to the successful outcome of the procedure. There were no data about the level of sedation achieved. The GDG agreed that the level of sedation achieved by ketamine alone was dependent on dose but that the sedation level was often uncertain because ketamine induces a sedated state in which the patient is not responsive and yet maintains their eyes open. In this state, known as dissociative sedation, vital reflexes remain intact to maintain breathing and prevent aspiration. The GDG discussed whether or not some of the patients were anaesthetised rather than sedated and it was appreciated that high doses could cause anaesthesia in which vital reflexes may be obtunded. It was agreed that it was not possible to be certain about what dose was compatible with sedation rather than anaesthesia.

Evidence showed that intravenous and intramuscular administrations were equally effective for painful procedures in the Emergency department setting and the GDG discussed the advantages and disadvantage of both methods. Intravenous administration facilitates the titration of smaller doses of ketamine and therefore reduces the chance of sedation outlasting the intended procedure. The GDG agreed that that intramuscular is a painful route of administration and should be reserved for situations when intravenous administration is impractical. However it was noted that despite local anaesthesia skin preparation intravenous injections were themselves painful and may need to be repeated if attempts were unsuccessful. Consequently it may be reasonable to offer a single intramuscular injection rather than wait for local anaesthesia to be applied to the skin and become effective in a child in whom venous access may prove to be difficult.

The GDG considered the evidence for ketamine combined other drugs. There were five RCTs¹²⁸, ⁷⁷, ¹⁵⁰, ²¹³, ⁹ in which a combination of ketamine and midazolam had been compared with other drugs. All were low quality evidence. In four studies¹²⁸, ⁷⁷, ¹³¹, ¹⁵² the authors stated that the target level of sedation was deep. The main efficacy outcome was completion of procedure and all procedures were completed in these RCTs. In comparison with a midazolam fentanyl combination the ketamine midazolam combination was associated with lower pain and distress scores. In comparison with propofol and fentanyl combination the ketamine midazolam combination was also associated with lower pain and distress scores although the recovery time was longer. In both comparisons ketamine midazolam combinations were associated with less oxygen desaturations. The GDG agreed that this was likely to be for two reasons. First it may be more difficult to titrate a combination of midazolam and fentanyl than ketamine and second fentanyl causes depresses breathing more than ketamine.

Two studies¹³¹, ¹⁵² compared the ketamine midazolam combination with techniques involving local anaesthesia for reduction of forearm fractures; the local anaesthesia was supplemented by midazolam alone in one and nitrous oxide in the other, and all techniques seemed equally effective.

The GDG discussed the problems of designing a RCT to determine the effect of combining ketamine with other drugs. For example in order to determine the effect of combining ketamine with midazolam it would be necessary to have a comparator group receiving midazolam alone. This however would not be possible because midazolam alone would not be effective for painful procedures. If a ketamine was compared with a ketamine midazolam combination the results would indicate the effect of midazolam. Nevertheless, if it was assumed that ketamine was effective it would be reasonable to consider such a study as evidence of how ketamine alone compared with the combination. The GDG reconsidered two RCTs²²⁷, ²⁰¹ that compared ketamine alone with ketamine combined with midazolam that had already been reviewed in the midazolam

evidence to recommendation discussions. It was agreed that the addition of midazolam conferred no advantage and was associated with more oxygen desaturation.

The GDG discussion focused on airway and breathing effects of ketamine. In some studies 10-15% of children had oxygen desaturation after ketamine but the GDG recognised that these events were usually brief and easily managed with oxygen and simple airway support. The level of desaturation may have been related to the skills of the healthcare practitioner. Nevertheless, evidence showed that potentially dangerous airway effects could occur after ketamine by either route. The need for the use of "bag and mask ventilation" was estimated to be approximately 1-2% but was less than this in some large cohort studies. Laryngospasm was the usual cause or airway obstruction although apnoea is known to be a potential hazard also. The GDG agreed that airway management skills and equipment are essential for this drug.

The GDG discussed the three studies 213 , 80 , 76 of ketamine combined with various drugs for endoscopy procedures. A study comparing the ketamine midazolam combination with a propofol fentanyl combination showed that ketamine was associated with more laryngospasm during gastroscopy. The GDG considered that ketamine causes more salivation than propofol and that the combination of pharyngeal secretions during gastroscopy 213 is likely to lead to laryngospasm.

The problem of fasting before ketamine was also discussed in the emergency department setting. It was agreed that the fasting status of a child in the emergency setting is often uncertain and that the stomach emptying is often delayed after trauma. The GDG felt that in many situations in the emergency setting there was a good trade-off between the benefit of prompt sedation with ketamine and the hazard of vomiting and aspiration during sedation. It was often reasonable therefore to consider using ketamine sedation in situations where a procedure was considered to be sufficiently urgent and that the risk of a full stomach was low. The GDG agreed that ketamine has a safe reputation for use in children who may not be fasted although the quality of evidence for risk of aspiration was very low. In order to prove that ketamine was safe in unfasted children, it was recognised that large numbers of children would need to be studied, some of whom were fasted and others not fasted, before this safety question could be answered with confidence.

Other side effects were also discussed. Vomiting was a common minor side effect but there was no evidence to show that any intervention prevented it. The GDG agreed that there should be research into methods of reducing vomiting with ketamine. Paradoxical excitement and hallucinations are known to cause agitation in the recovery period and the GDG noted that this was uncommon and not reduced by routine administration of midazolam.

Discussions led to how ketamine sedation compared with anaesthesia in the setting of a painful procedure in an emergency department. The GDG could find no evidence to confirm which approach is best but GDG members knew that the issue has been debated recently in the Emergency Medicine professional journals. It was agreed that there were potential economic advantages to providing sedation within a few hours of admission rather than waiting for the services of an anaesthesia team that may involve overnight admission. The GDG recognised that this was a common dilemma. However in many hospitals Emergency department staff were not trained to administer ketamine. Training of a team to deliver ketamine sedation was considered to be essential if ketamine was to be used safely.

The agreement by the GDG is that economic analysis should be conducted only for sedation techniques commonly available in the NHS. Economic analysis was conducted for six broad groups (dental procedure in children, dental procedure in adolescents, short painful procedures, painless imaging, oesophago-gastroscopy and colonoscopy). The GDG felt there is some evidence that ketamine alone is effective and safe. It is commonly used in short painful procedure in the NHS, and it was therefore agreed that this strategy should be compared to other relevant strategies in the economic analysis conducted for this population group. Details of the considerations of cost-effectiveness with respect to using ketamine alone in short painful procedures are given in section 6.12.1.2.

2 6.5 Chloral hydrate

Matrix of chloral hydrate comparators

Key:

Chloral hydrate = CH

Fentanyl = F

Isoflurane = I

Ketamine=K

Local anaesthesia = LA

Midazolam = M

Nitrous oxide = N_20

Nitrous oxide and oxygen = N_20+02

Opioids = O

Propofol= P

Sevoflurane = S

Triclofos sodium = TS

Chloral hydrate vs

	Reference	Tables	Evidence statements page
Placebo			
	Houpt 1989 ⁹⁵	Table 40	220
Head to head			
М	Dallman 2001 ⁴⁹	Table 41	220
General Anaesthetic (GA)	Thompson 1982 ²¹¹	Table 42	220
Music	Loewy 2005 ¹⁴⁹	Table 43	221
Combinations			
CH + hydroxyzine vs M + acetaminophen	Dallman 2001 ⁴⁹ Reeves, 1996 ¹⁸⁷	Table 44	221
Safety			
RCTs	Marti-Bonmati 1995 ¹⁵⁸	Table 46	221
Vomiting	Houpt 1989 ⁹⁵	Table 47 Table 48	222

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Observational studies	Ronchera-Oms 1994 ¹⁹² Napoli 1996 ¹⁶⁹ Greenberg 1991 ⁸³ Greenberg 1993 ⁸⁴ Malviya 2000 ¹⁵⁵ Fox 1990 ⁶⁹ Heistein 2006 ⁹¹ Cortellazzi 2007 ⁴³ Needleman 1995 ¹⁷¹	Table 47 Table 48	222
Route of administration			
Nil			
Dose			
CH high dose vs CH low dose	Houpt 1985%	Table 45	220
CH intermediate dose vs CH high dose	Marti-Bonmati 1995 ¹⁵⁸	Table 46	221

6.5.1 Clinical methodological introduction

2	CLINICAL QUESTIONS:
3 4 5	For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is chloral hydrate (with or without: analgesia, another drug or psychological techniques):
6 7 8	- effective for sedation (at minimal, moderate, and deep levels) in comparison with usu care, with analgesia alone, with another sedation drug, with psychological techniques of with general anaesthesia?
9 10 11	The literature was searched for systematic reviews and RCTs for the clinical efficacy an safety of chloral hydrate. The search was expanded to include observational studies f the safety of chloral hydrate.
12 13	No systematic reviews were identified for the use of chloral hydrate in paediatric sedation.
14	Seven RCTs met the inclusion criteria for the review of the efficacy of chloral hydrate.
15	Two RCTs met the inclusion criteria for the review of the safety of chloral hydrate.
16 17 18	Meta-analysis was not performed as there were no studies in which comparisons and outcome measures were sufficiently homogenous to calculate a meaningful summary statistic.
19	Nine non RCTs in 5,188 patients assessed the safety of chloral hydrate.
20	6.5.2 Evidence profiles
21	6.5.2.1 RCT evidence profiles for efficacy and safety
22 23	Study characteristics and methodological quality of the study are provided in Appendi D. GRADE tables for quality assessment and summary of findings are provided below.
24	

PLACEBO COMPARISONS

Table 40: Chloral hydrate vs. placebo; Houpt 1989 95

Author(s): Houpt 1989

Question: Should chloral hydrate vs. placebo be used in children also receiving nitrous oxide?

Settings: Dental

			Quality asso	remont	Summary of findings							
	Quality assessment								No of patients Effect			Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Chloral Hydrate	Placebo	Relative (95% CI)	Absolute	Quality	ance
			•		-	considerations	пуштаце		(95% CI)			
Vomiting												
1	randomised	serious1	no serious	no serious	serious ²	none	0/40	1/19	DD 0 00 (0 0	53 more per 1000 (from 42		
	trial		inconsistency	indirectness				2/19 (5.3%)	RR 2.00 (0.2 to 20.24)	fewer to 1000 more)	LOW	
							(10.576)	0%	10 20.24)	0 more per 1,000	LOW	

¹ Small sample size and wide confidence levels for relative effect ² Generation code and allocation concealment not described

HEAD TO HEAD COMPARISONS

Table 41: Chloral hydrate vs. intranasal midazolam; Dallman 2001 49

Author(s): Dallman 2001 Question: Should chloral hydrate vs. intranasal midazolam be used for paediatric sedation?

			Quality assoc	semont	Summary of findings							
	Quality assessment							No of patients		Effect		Importan
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Chloral hydrate	intranasal midazolam	Relative (95% CI)	Absolute	Quality	ce
Recovery							,		(22227)			
	randomised trial			no serious indirectness	serious ²	none	24/31 (77.4%)	30/31 (96.8%)	RR 49.00 (3.11 to 771.67)	1000 more per 1000 (from 1000 more to 1000 more)	LOW	

Table 42: Chloral hydrate vs. general anaesthesia; Thompson 1982 211

Author(s): Thompson 1982

Question: Should chloral hydrate vs. GA be used in paediatric sedation?

			Quality ass	rossmont					Summary of	findings		
			Quality as:	bessillerit			No of p	atients		Effect		Importan
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Chloral hydrate	GA	Relative (95% CI)	Absolute	Quality	ce
Complete procedure												•
1	randomised trial	- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	85/101 (84.2%)	101/101 (100%)	RR 0.84 (0.77 to 0.92)	160 fewer per 1000 (from 80 fewer to 230 fewer)	LOW	
Induction	time (range o	of scores: 25	-55; Better indicat	ed by less)				0%		0 fewer per 1,000		
1	randomised trial	- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	101	101	3	MD 30 ³	LOW	
Duration (of procedure	(range of sc	ores: 48-80; Bette	r indicated by les	s)							
1	randomised trial	- ,	no serious inconsistency	no serious indirectness	no serious imprecision	none	101	101	4	MD 32 ⁴	LOW	

¹ Inadequate randomisation, allocation concealment. No blinding. Distribution of ages not equal: 203 infants 0-1month, 82 children ages 1-2 years and remaining equally divided between years 2-0

years.

² No explanation was provided

³Unable to calculate RR as SD not given; mean per group: 55 minutes vs. 25 minutes

⁴Unable to calculate RR as SD not given; mean per group: 48 minutes vs. 80 minutes

Table 43: Chloral hydrate vs. music therapy; Lowey 2005 149

Author(s): Lowey 2005

Question: Should chloral hydrate vs. music therapy be used in paediatric sedation?

Settings: EEG

			Quality asses	semont					Summary o	f findings		
			Quality asses	Sament			No of p	atients		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Chloral hydrate	music therapy	Relative (95% CI)	Absolute		ance
Complete	Complete procedure											
1	randomised trial			no serious indirectness	serious ²	none	12/24 (50%)	33/34 (97.1%)	RR 0.52 (34 to 0.77)	466 fewer per 1000 (from 223 more to 1000 fewer) 0 fewer per 1,000	LOW	
Induction	time (range o	f scores: 23	-32; Better indicate	ed by less)	l				L	•		
1	randomised trial			no serious indirectness	serious ²	none	12	33	3	MD 9.0 ³	LOW	
Total time	Total time (measured with: minutes; range of scores: 66-226; Better indicated by less)											
1	randomised trial			no serious indirectness	serious ²	none	12	33	4	MD 160 ⁴	LOW	

Generation code and allocation concealment not described. Study was unblinded.

² Small sample size
³ Reported mean per group: 23 minutes vs. 32 minutes
⁴ Reported p<0.001; mean per group: 66 minutes vs. 226 minutes

COMBINATION COMPARISONS

Table 44: Chloral hydrate/hydroxyzine vs. midazolam/acetaminophen; Reeves 1996 49,187

Author(s): Reeves, 1996

Question: Should chloral hydrate/hydroxyzine vs. midazolam/acetaminophen be used in paediatric sedation?

Settings: Dental

			Quality assessmen				Summary of fire	ndings				
	Quality assessment							No of patients Effect				Importa
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Chloral Midazolam hydrate/hydroxyzine /acetaminopho		Relative (95% CI)	Absolute	Quality	•
Distress by H	oupt score (ran	ge of scores	s: -; Better indicate	d by less)								
	randomised trial			no serious indirectness	serious ²	none	20	20	-	MD -0.10 (- 0.83 to 0.63)		

Generation code and allocation concealment not described Small sample size. Assessment has elements of subjectivity.

DOSE COMPARISONS

Table 45: High dose vs. low dose chloral hydrate; Houpt 1985 96

3

Author(s): Houpt 1985

Question: Should High dose chloral hydrate vs. Low dose chloral hydrate be used for sedation in children?

Settings: Dental

			Quality acce	coment				Sur	nmary of find	ings		
	Quality assessment							No of patients Effect				Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	High dose chloral hydrate	Low dose chloral hydrate	Relative (95% CI)	Absolute		-
Completion	Completion of procedure											
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	13/17 (76.5%)	16/17 (94.1%)	RR 0.81 (0.61 to 1.09)	179 fewer per 1000 (from 367 fewer to 85 more)	LOW	
								0%	` ´	0 fewer per 1,000		
Induction	Induction time (measured with: minutes; range of scores: 9-24; Better indicated by less)											
1	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	17	17	-	MD 15 (0 to 0)	LOW	

¹ Randomisation and allocation concealment not described. ² Small sample size (<20 patients per arm)

Table 46: Intermediate dose chloral hydrate vs. high dose chloral hydrate; Marti-Bonmati 1995 158

Author(s): Marti-Bonmati et al, 1995

Question: Should intermediate dose chloral hydrate vs. high dose chloral hydrate be used for sedation in children?

Settings: MRI

			Quality acc	occment				Sumi	mary of find	lings		
			Quality ass	essinent			No of pati	ents		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Intermediate dose chloral hydrate	High dose chloral hydrate	Relative (95% CI)	Absolute	Quality	ance
Completi	ompletion of procedure											
	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	46/50 (92%)	47/47 (100%)	RR 0.92 (0.84 to 1.01)	80 fewer per 1000 (from 160 fewer to 10 more) 0 fewer per 1,000	MODERATE	
Length of	ength of induction time (measured with: minutes; range of scores: -; Better indicated by less)											
	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	50	47	-	MD 7 (6.38 to 7.62)	MODERATE	
Recovery	time (measu	red with: m	inutes; range of	scores: -; Bette	r indicated by l	ess)				•		
	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	50	47	=	MD -8.00 (-10.2 to - 5.8)	MODERATE	
All advers	se events											
	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	10/50 (20%)	10/47 (21.3%)	RR 0.94 (0.43 to 2.05)	13 fewer per 1000 (from 121 fewer to 224 more) 0 fewer per 1,000	MODERATE	÷

Method of randomisation and allocation concealment not adequately described.

6.5.2.2 Non RCT safety(adverse events)

- Nine non RCT observational studies studies with greater than 300 subjects (total n= 5,188) assessed the safety of chloral hydrate ¹⁹². There were six prospective reviews and three retrospective studies conducted primarily for imaging procedures (7) as well as one dental and one ophthalmic study.
- The non RCT study characteristics for chloral hydrate are presented in Table 47.
- The non RCT adverse event data for chloral hydrate is presented in Table 48.

8

1

1 Table 47: Chloral hydrate Non RCT study characteristics safety review

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Prospective Cohort						
192 Spain	MRI	Not stated	Chloral hydrate for imaging	55% male	Chloral hydrate syrup 70 mg/ml	Permitted oral fluids before examination
169 USA	Echocardiography	Not stated	Chloral hydrate for imaging	Not stated	Median dose of chloral hydrate was 77 mg/kg	Not stated
83 USA	СТ	Not stated	Chloral hydrate for imaging	63% male	100 mg/kg in a single dose with maximum of 2 grams	Not stated
84 USA	MRI	Not stated	Chloral hydrate for imaging	Not stated	100 mg/kg	Not stated

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Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
155 USA	MRI	72% ASA I; 27% ASA II and 1% as ASA III.	Sedation to facilitate outpatient diagnostic imaging procedures	53% male	64 <u>+</u> 13 mg/kg chloral hydrate	Not stated
usa	Ophthalmic examination	Not stated	Chloral hydrate for ophthalmic procedures in infants and young children	Not stated	80-100 mg/kg chloral hydrate not to exceed 3 g.	NPO for 4 hours prior to administration of chloral hydrate
Retrospective Studies					100 mg/kg chloral hydrate	
91 USA*	Echocardiography	7.3% ASA 1; 54.4% ASA II; 37.4% ASA III and 0.8% ASA IV	Chloral hydrae sedation for echocardiography	Not stated	Oral chloral hydrate (80 mg/kg, maximum 1 g)	Infants less than 6 months could receive formula and solids for up to 6 hours, breast milk for up to 4 hours and clear liquids for up to hours before sedation. Children 6 months or older could receive solids and liquids for up to 6 hours and clear liquids for up to 2 hours before sedation
43 Italy	MRI	Not stated	Level 3 on Skeie Scale – asleep but easily aroused	61% male	50 – 100 mg/kg to a maximum dose of 1.5 g/kg	Determined according to the ASA recommendations

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
171 USA	Dental	ASA I or I	Conscious sedation	56% male	Average dose of chloral hydrate 776 mg (55 mg/kg)	'Pre-operative dietary restrictions'

^{**}In this study potential risk factors were assessed for their association with adverse events. Univariate analysis identified age younger than 6 months, cyanotic heart disease and hospitalization at the time of the study as significant risk factors. Multivariate analysis identified only age younger than 6 months as a significant independent risk factor for the occurrence of an adverse event.

Table 48: Chloral hydrate safety: RCTs and Non RCT's (n = >300 patients)

Church								ADVERSE EV	ENTS, rate: % (n)				GRADE PROFILE
Study type, reference,	Drug / Comparis	Procedur e	Age	Total N		Res	piratory interver	ntion		Cardiac arrest requiring either/or		oxygen	EVIDENCE
country	on				Aspiration	oral- pharyngeal airway	endotracheal intubation	assisted ventilation	external cardiac massage	defibrillation	vomiting	saturation <90%	QUALITY
•	e Cohort stu	dies											
Spain	Chloral hydrate	MRI	Mean age 41 ± 30 months	596							6.9% (41)	0	VERY LOW
USA	Chloral hydrate	Echocardi ography	3 weeks to 14 years; median age 13 months	405							6% (23)	6% (24) defined as greater than 5% drop from baseline in these children with heart disease	VERY LOW
USA	Chloral Hydrate: high dose of 80-100 mg/kg	СТ	Mean age 2.18 years	326	1 aspiration of secretions by child with severe mental retardation		2 due to obstruction of the airway by the tongue. One child was profoundly retarded.				4.3% (14)		VERY LOW
USA	Chloral hydrate	MRI	1 month - 11 years	300							4% (12)		VERY LOW
USA	Chloral hydrate	MRI/CT	3.8 <u>+</u> 3.4 years	336							3% (8) has 'GI effects' in hospital; 26% (78) had 'GI' effects at home		VERY LOW
69	Chloral hydrate	Opthalmic examinatio n	1 month - 5 years	302							0	0	VERY LOW

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Retrospe	Retrospective Studies 1												
91 USA*	Chloral hydrate	Echocardi ography	Birth to 64 months	1095 38% were ASA 3 or 4; 88% had detectable heart disease; 78% received a single agent and 22% received >1 medication	0	0.3% (3) required oral or nasal suctioning	0.1% (1) required intubation	0.1% (1) required bag- mask ventilation			0.4% (4)	5.9% (65) defined as greater than 10% drop from baseline in these children with heart disease	VERY LOW 2 3 4 5 6
taly	Chloral hydrate	MRI	Mean age 28.2 <u>+</u> 18.1 months	888 procedure s using chloral hydrate alone for MRI in neurologic ally impaired children			0	0			0.2% (n=2)	0.5% (n=4)	VERY LOW 8 9 10 11
USA	Chloral hydrate	Dental	Mean age 2.6 years	336							8.1%(27)		VERY LOW 13

In this study potential risk factors were assessed for their association with adverse events. Univariate analysis identified age younger than 6 months, cyanotic heart disease and hospitalization at the time of the study as significant risk factors. Multivariate analysis identified only age younger than 6 months as a significant independent risk factor for the occurrence of an adverse event.

1	6.5.3 Evidence statements
2	6.5.3.1 RCT efficacy and safety
3	PLACEBO COMPARISONS
4	Chloral hydrate vs. placebo
5	Houpt 1989 95
6 7	No efficacy outcomes of interest were reported in this study. One adverse event outcome of interest was reported in this study.
8	There was no significant difference in:
9	number of children who vomited [Low quality evidence]
10	
11	HEAD TO HEAD COMPARISONS
12	Chloral hydrate vs. intranasal midazolam
13	Dallman 2001 ⁴⁹
14	Compared to intranasal midazolam, the chloral hydrate group had significantly:
15	Longer recovery time [Low quality evidence]
16	There was no significant difference in:
17	number of distressed children [Low quality evidence]
18	
19	Chloral hydrate vs. GA
20	Thompson 1982 ²¹¹
21	Compared to GA, the chloral hydrate group had significantly:
22	Fewer completed procedures [Low quality evidence]
23 24	 Longer induction time time (55 mean minutes vs. 25 mean minutes) Unable to calculate RR as SD not given [Low quality evidence]
25 26	 Less procedure time (48 mean minutes vs. 80 mean minutes); Unable to calculate RR as SD not given [Low quality evidence]
27	
28	Chloral hydrate vs. music therapy

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1	Lowey 2005 ¹⁴⁹
2	Compared to music therapy, the chloral hydrate group had significantly:
3	Fewer completed procedures [Low quality evidence]
4	 Longer total time asleep; report ed p<0.001 [Low quality evidence]
5	There was no significant difference in:
6	Induction time [Low quality evidence]
7	
8	COMBINED COMPARISONS
9	Chloral hydrate + hydroxyzine vs. midazolam + acetaminophen
10	Reeves 1996 49,187
11	There was no significant difference in:
12	Distress scores [Low quality evidence]
13	
14	DOSE COMPARISONS
15	High dose vs. low dose chloral hydrate
16	Houpt 1985 %
17 18	Compared to low dose chloral hydrate (mean 708 mg), the high dose chloral hydrate (mean 1062 mg) group had significantly:
19	• fewer procedure failures [Low quality evidence]
20	less induction time [Low quality evidence]
21	
22	Intermediate dose chloral hydrate vs. high dose chloral hydrate
23	Marti-Bonmati 1995 ¹⁵⁸
24 25 26	In this study, sedation was judged a failure if the MRI imaging study could not be completed, if additional sedation other than chloral hydrate was required for completion of the study or if fewer than 95% of the images were acceptable.
27 28	Compared to intermediate dose chloral hydrate (70 mg/kg), the high dose chloral hydrate (96 mg/kg) group had significantly:
29	Less induction time [Moderate quality evidence]

1 2	There was no significant difference in:
3	Completion of MRI [Moderate quality evidence]
4	Recovery time [Moderate quality evidence]
5 6	 Total adverse events (events were not reported individually) [Moderate quality evidence]
7	6.5.3.2 NON-RCT safety (adverse events)
8 9	For the characteristics of studies and outcome data on chloral hydrate refer to Table 47 and Table 48.
10 11 12 13	 One prospective study ⁸³ of high dose chloral hydrate reported 1 aspiration of secretions by child with severe mental retardation and 2 endotracheal intubations due to obstruction of the airway by the tongue. One child was profoundly retarded.
14 15 16	 One retrospective study ⁹¹ reported that 0.3% of children receiving chloral hydrate required oral or nasal suctioning, 0.1% required intubation and 0.1% required bag-mask ventilation.
17	 No respiratory events were reported in seven studies.
18	 No cardiac events were reported in nine studies.
19 20 21	• The mean vomiting rate for 9 non RCT observational studies of chloral hydrate was 4.1% 43,69,83,84,91,155,169,171,192. One study 155 reported that 26% (78) of patients had 'GI' effects at home.
22 23	 One study¹⁶⁹ reported oxygen saturation drop greater than 5% from baseline in 6% of patients.
24 25	 One study⁹¹ reported oxygen saturation drip greater than 10% from baseline in 5.9% of patients.
26	 One study⁴³ reported 0.5% rate of oxygen saturation <90%.
27	6.5.4 GDG discussion of the evidence for chloral hydrate
28 29 30 31 32	Chloral hydrate is an oral drug and unfortunately causes nausea and vomiting when large volumes of the drug are used. The GDG agreed that chloral hydrate is therefore likely to be less successful in larger children. Some GDG members thought more than 1g of chloral hydrate may be vomited and hence be unsuccessful. This may explain why choral is thought to be more effective in smaller children.
33 34 35 36	The GDG considered 14 studies ²¹¹ , ⁸³ , ⁸⁴ , ¹⁵⁵ , ¹⁵⁸ , ¹⁹² , ⁴³ , ¹⁴⁹ , ¹⁶⁹ , ⁹¹ , ⁶⁹ , ⁹⁶ , ⁴⁹ , ¹⁸⁷ of chloral hydrate used alone; two others ⁹⁵ , ¹⁷¹ were of chloral hydrate combined with other drugs. Ten of these studies were in children undergoing painless procedures; five for dental treatment ⁹⁵ , ¹⁷¹ , ⁹⁶ , ⁴⁹ , ¹⁸⁷ and one for ophthalmic examination ⁶⁹ . Of the

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1 2	painless procedure studies five were for MRI ⁸⁴ , ¹⁵⁵ , ¹⁵⁸ , ¹⁹² , ⁴³ and two for CT imaging ²¹¹ , ⁸³ .
3 4 5 6 7	Two RCTs ²¹¹ , ¹⁵⁸ were found for painless imaging. One study ¹⁵⁸ showed that high dose chloral hydrate was not more effective than low dose for MRI but that high dose chloral hydrate caused shorter onset of sedation (the evidence level was moderate). The other study ²¹¹ showed that anaesthesia was more effective than chloral hydrate for CT imaging (the evidence level was low). The other studies were non-RCT.
8 9 10	The GDG concluded that uncooperative children needed to be asleep for imaging and that high doses of chloral hydrate were successful in approximately 90% of children under 15kg. High doses were likely to be more reliable than low doses.
11 12 13 14 15 16 17 18 19 20	The GDG debated as to what sedation level was achieved by chloral hydrate in the painless imaging setting. The GDG noted that the doses of chloral hydrate used caused the children to sleep and, because the success of the scanning required them to be immobile and undisturbed, the true sedation level achieved was uncertain. The GDG members appreciated that all children in the evidence studies were likely to be either moderately or deeply sedated. Nevertheless the GDG agreed that unconsciousness was possible and that appreciable airway and breathing effects could be caused in a small percentage of children. These problems were uncommon but were reported. In one cohort study ⁸³ a child with severe mental retardation suffered pulmonary aspiration during sedation.
21 22 23 24 25	The disadvantages of chloral hydrate are that it is administered as a single oral dose, that it cannot therefore be titrated, and that its effect is variable in terms of depth of sedation, and its onset and recovery times. However there are potential economic advantages of chloral hydrate if its success rate is high enough because anaesthesia resources may be saved (both techniques are equally safe).
26 27 28 29	There was evidence of chloral hydrate being used in other settings. Chloral hydrate combined with nitrous oxide was shown in one study ⁹⁵ to be more effective than nitrous oxide alone in young children having dental treatment. This combination however was associated with vomiting in 10% of cases.
30 31 32	Chloral hydrate was also useful for calming small irritable children for echocardiography and in this setting the GDG appreciated that anaesthesia would not usually be appropriate. ¹⁶⁹ , ⁹¹
33 34 35 36 37	The GDG noted that small children could be sedated successfully chloral hydrate for eye examination. In another study ¹⁴⁹ the GDG noted that children could be calmed for EEG studies more effectively by music rather than chloral hydrate however the GDG thought that this was an unusual setting and that children having EEG are not required to be immobile
38 39 40 41 42 43 44 45 46	The GDG agreed that economic analysis should be conducted only for sedation techniques commonly available in the NHS. Economic analysis was conducted for six broad groups (dental procedure in children, dental procedure in adolescents, short painful procedures, painless imaging, oesophago-gastroscopy and colonoscopy). Chloral hydrate was felt to be effective and safe. It is commonly used in painless imaging in the NHS. The GDG therefore agreed that this strategy should be included in the economic analysis conducted for patients undergoing painless imaging. Details of the considerations of cost-effectiveness with respect to using chloral hydrate in painless imaging are given in section 6.12.2.2.

2 6.6 Triclofos sodium

Matrix of triclofos sodium comparators

Key:

 ${\sf Chloral\ hydrate} = {\sf CH}$

Fentanyl = F

Isoflurane = I

Ketamine=K

Local anaesthesia = LA

Midazolam = M

Nitrous oxide = N_20

Nitrous oxide and oxygen = N_20+02

Opioids = O

Propofol= P

Sevoflurane = S

Triclofos sodium = TS

Triclofos sodium vs

	Reference	Tables	Evidence statements page
	Kolololide	Idoles	Evidence sidiements page
Placebo			
Nil			
Head to head			
TS vs M	Singh 2002 ²⁰³	Table 49	227
Combinations			
Nil			
Safety			
RCTs	-		
Observational studies	Nil		
Route of administration			
Nil			
Dose			
Nil			

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1

6.6.1 Clinical methodological introduction

2	CLINICAL QUESTIONS:
3 4 5	For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is triclofos sodium (with or without: analgesia, another drug or psychological techniques):
6 7 8	- Effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?
9	- Safe for sedation (at mild, moderate, and deep levels) in different settings?
10 11 12	The literature was searched for systematic reviews RCTs for the clinical efficacy of triclofos sodium. The search was expanded to include non RCT observational studies for the safety of triclofos sodium.
13	There were no systematic reviews identified for the use of opioids in paediatric sedation.
14 15 16	One RCT was found that compared triclofos sodium with midazolam. Whilst efficacy data was reported safety data was not. There were no non-RCT observational studies assessing the safety of triclofos sodium.
17	Meta-analyses were not performed as there was only one RCT.
18	6.6.2 Evidence profiles
19	6.6.2.1 RCT evidence profiles for efficacy and safety
20 21	Study characteristics and methodological quality of the study are provided in Appendix D. GRADE tables for quality assessment and summary of findings are provided below.
22	

HEAD to HEAD COMPARISONS

Table 49: Oral triclofos sodium vs. oral midazolam; Singh 2002 203

Question: Should oral triclofos sodium vs. oral midazolam be used in children and young people undergoing diagnostic and therapeutic procedures?

Settings: Dental

Bibliography: Singh 2002

	Quality assessment							Summary of findings				
	Quality assessment							No of patients Effect				
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	oral triclofos sodium	oral midazolam	Relative (95% CI)	Absolute	Quality	ance
Completio	n of procedu	re										
1	randomised trial	- /	no serious inconsistency	no serious indirectness	no serious imprecision	none	30/30 (100%) ²	30/30 (100%)	not estimable	-	LOW	
Induction	time (Better in	ndicated by I	ess)									
1	randomised trial	- /	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	30	-	MD 16.10 (14.09 to 18.11)3	LOW	
Recovery time: when the patient was able to sit or stand alone with minimal assistance (Better indicated by less)											•	
1	randomised trial	1	no serious inconsistency	no serious indirectness	no serious imprecision	none	30	30	-	MD 38.23 (31.52 to 44.94)3	LOW	

³ Singh 2002: p<0.00001

Singh 2002: patients and outcome assessors blinded however concealment, ITT and attrition details not stated; small study

Singh 2002: ease of treatment completion rated as 1-excellent, 2-difficult and 3-impossible; study stated that treatment was most convenient for midazolam group than for triclofos group. Difficulty in treatment was significantly more for the group of promethazine than for midazolam (p<0.01) and for triclofos (p<0.05)

1	6.6.2.2 Non RCT evidence profiles for safety
2	There were no non RCT observational studies of triclofos sodium.
3	6.6.3 Evidence statements
4	6.6.3.1 RCT efficacy and safety
5	HEAD to HEAD COMPARISONS
6	Oral triclofos sodium vs. Oral Midazolam
7	Singh 2002 ²⁰³
8	All patients completed the procedure [low quality evidence]
9 10	Compared with the oral midazolam group, the oral triclofos sodium group had significantly:
11	Longer induction time [low quality evidence]
12	Slower recovery time [low quality evidence]
13	
14	Non RCT safety (adverse events)
15	There were no non RCT observational studies of triclofos sodium.
16	6.6.4 GDG discussion of the evidence for triclofos sodium
17 18 19	Only one study ²⁰³ of triclofos was found and it compared triclofos with midazolam for dental procedures. The GDG noted that triclofos was not effective in this setting and also that the quality of evidence was very low.
20 21 22 23	The GDG noted that the properties of triclofos and chloral hydrate were similar and that triclofos may cause less gastric irritation. The GDG discussed the potential advantages of triclofos but without evidence this drug could not be recommended as more effective that chloral hydrate.
24 25	The GDG felt that triclofos sodium is not among the sedation drugs commonly used in the NHS, and decided that it should not be included in the economic analysis.
26	

1 6.7 Nitrous Oxide

Matrix of nitrous oxide comparators

Key:

Chloral hydrate = CH

Fentanyl = F

Isoflurane = I

Ketamine=K

Local anaesthesia = LA

Midazolam = M

Nitrous oxide = N_20

Nitrous oxide and oxygen = N_20+02

Opioids = 0

Propofol= P

Sevoflurane = S

Triclofos sodium = TS

Nitrous oxide vs

		T =	1
	Reference	Tables	Evidence statements page
Placebo			
N ₂ 0 vs Oxygen	McCann 1996 ¹⁶⁰	Table 50	245
	Primosch 1999 ¹⁸⁵	Table 51	245
N ₂ 0 vs nitrogen and oxygen	Fauroux 2004 ⁶⁴	Table 52	245
Head to head			
N20 vs Behavioural	Veerkamp 1993 ²²²	Table 53	246
management	Veerkamp 1995 ²²⁰	Tuble 50	240
N ₂ 0 vs Midazolam	Wilson 2007 ²³³	Table 54	246
	Wilson 2003 ²²⁹	Table 55	246
	Wilson 2006 ²³⁰ Wilson 2002 ²³¹ Wilson 2002 ²³²	Table 56	247
N ₂ 0 + EMLA vs EMLA	Ekbom 2005 ⁵⁸	Table 57	248
Combinations			
N ₂ 0 + M vs air + M	Averley 2004 ¹⁹	Table 58	248
N ₂ 0 + M vs S + N ₂ 0 + M	Averley 2004 ¹⁹	Table 59	248

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N ₂ 0 + M + S vs air + M	Averley 2004 ¹⁹	Table 60	249	
Safety				
RCTs				
Desaturation	Primosch 1999 ¹⁸⁵	Table 61 Table 62	249	
Observational studies	Babl 2008 ²⁰ Gall 2001 ⁷³ Faddy 2005 ⁶¹	Table 61 Table 62	249	
Route of administration				
Nil				
Dose				
Nil				

6.7.1 Clinical methodological introduction

1

24

2	CLINICAL QUESTIONS:
3 4 5	For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is nitrous oxide (with or without: analgesia, another drug or psychological techniques):
6 7 8	- effective for sedation (at minimal, moderate, and deep levels) in comparison with usua care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?
9	- safe for sedation (at mild, moderate, and deep levels) in different settings?
10 11 12	The literature was searched for systematic reviews and RCTs for the clinical efficacy and safety of nitrous oxide. The search was expanded to include observational studies for the safety of nitrous oxide.
13	No systematic reviews were identified for the use of nitrous oxide in paediatric sedation
14	There were no placebo controlled trials identified.
15	Twelve RCTs met the inclusion criteria for the review of the efficacy of nitrous oxide.
16	Four RCTs met the inclusion criteria for the review of the safety of nitrous oxide.
17	Three non RCTs in 8,220 patients assessed the safety of nitrous oxide.
18 19	Meta-analysis were performed if comparisons and outcome measures were sufficiently homogenous to calculate a meaningful summary statistic 220,222 ; $^{230-232}$.
20	6.7.2 Evidence profiles
21	6.7.2.1 RCT evidence profiles for efficacy and safety
22 23	Study characteristics and methodological quality of the study are provided in Appendix D. GRADE tables for quality assessment and summary of findings are provided below.

PLACEBO COMPARISONS

Table 50: Nitrous oxide vs. oxygen; McCann 1996 160

Author(s): McCann 1996

Question: 50% nitrous oxide vs. 100% oxygen for sedation in children

Settings: Dental

			Quality acces	ement				Sun	nmary of findings	S		
			Quality asses	Silient			No of par	tients	Effe	ect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	50% nitrous oxide	100% oxygen	Relative (95% CI)	Absolute	Quality	ance
Quiet behaviour on OSUBRS												
	randomised trial			no serious indirectness	serious ²	none	19/20 (95%)	15/20 (75%)	RR 1.27 (0.96 to 1.66)	202 more per 1,000	LOW	

¹ Randomisation and allocation concealment not described

Table 51: Nitrous oxide vs. oxygen; Primosch 1999 185

Author(s): Primosch 1999

Question: 40% nitrous oxide vs. 100% oxygen for sedation in children

Settings: Dental

			Quality asses	cmont			Sumn	nary of fin	dings					
			Quality asses	Silicili			No of par	tients		Effect		Import		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	40% nitrous oxide	100% oxygen	Relative (95% CI)	Absolute	Quality	_		
Quiet behav	viour on OSUB	RS (measur	ed with: OSBU ordin	al scale; range of s	er indicated by less)								
	randomised trial			no serious indirectness	serious	none	22	22	3	not pooled3	LOW			
Oxygen sat	cygen saturation (range of scores: -; Better indicated by less)													
	randomised trial			no serious indirectness	serious ²	none	22 ⁴	22	4	MD 0.00 (-0.01 to 0.01) ⁴	LOW			

Randomisation and allocation concealment not described.

8

³ RR behaviour scores not estimable due to use of an ordinal scale and incomplete statistical information; reported scores: 713 for N2O group and 630 for O2 group; reproted p<0.001.
⁴ Values in the two groups were exactly the same, 99+ 0.01.

Table 52: Nitrous oxide vs. nitrogen and oxygen; Fauroux 2004 64

Author(s): Fauroux 2004

Question: 50% nitrous oxide vs. 50% nitrogen & oxygen for sedation in children

Settings: Broncoscopy

			Quality asse	semont				S	ummary of fi	ndings		
			Quality asse	SSIIICIIL			No of	f patients		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	50% nitrous oxide	50% nitrogen & oxygen	Relative (95% CI)	Absolute	Quality	
Completio	mpletion of procedure											
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	11/53 (20.8%)	32/52 (61.5%)	RR 0.34 (0.19 to 0.6)	406 fewer per 1000 (from 246 fewer to 498 fewer)	LOW	
							(20.6%)	61.5%	(0.19 (0 0.6)	405 fewer per 1,000	LOW	
Pain score	e: CHEOPS (r	ange of sco	res: -; Better indic	ated by less)								
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	53	53	-	MD -1.3 (-2.09 to -0.51)	LOW	
Pain: VAS	for children	>6 years (ra	nge of scores: -; E	Better indicated b	y less)							
	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	24	13	-	MD -28 (-34.44 to -21.56)	LOW	

¹ Randomisation and allocation concealment not described.
² Small sample size not adequate to achieve power calculation of 90%

² Small sample size

HEAD TO HEAD COMPARISONS

Table 53: Nitrous oxide vs. behavioural management; Veerkamp 1993 ^{220,222}; Veerkamp 1995 ²²²

Author(s): Veerkamp 1993; Veerkamp 1995

Question: Nitrous oxide vs. behavioural management for sedation in children

Setting: Dental

			Quality acces	cmont				Summar	y of findi	ngs			
			Quality asses	Silielit			No	of patients		Effect		Import	
No of studies	studies Design Limitations Inconsistency Indirectness Imprecision considerati							behavioural management	Relative (95% CI)	Absolute	Quality	ance	
Anxiety (ra	Anxiety (range of scores: -; Better indicated by less)												
	randomised trial	- /		no serious indirectness	serious ²	none	50	51		MD -0.54 (-0.88 to -0.2)	VERY LOW		

¹ Randomisation method and allocation concealment not described. There was only partial blinding ² Two studies by same investigator with small sample sizes

Table 54: Nitrous oxide vs. transmucosal midazolam; Wilson 2007 233

Author(s): Wilson 2007

Question: Nitrous oxide vs. transmucosal (buccal) midazolam for sedation in children

Settings: Dental

			Quality asses	emont				Summary of	of findings			
			Quality asses	Sillelli				No of patients	Eff	ect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrous oxide	transmucosal (buccal) midazolam	Relative (95% CI)	Absolute	Quality	ance
Length of	induction (rar	ge of score	s: -; Better indicate	d by less)								
1	randomised trial			no serious indirectness	serious ²	none	36	36	-	not pooled ³	LOW	
Duration o	f procedure (range of sco	res: -; Better indica	ated by less)								
1	randomised trial			no serious indirectness	serious ²	none	36	36	-	not pooled4	LOW	
Total time	(range of sco	res: -; Bette	r indicated by less)								•	,
1	randomised trial			no serious indirectness	serious ²	none	36	36	-	not pooled ⁵	LOW	
Patient pre	eference				•						•	•
1 Cinala bli	trial			no serious indirectness	serious ²	none	20/36 (55.6%)	10/36 (27.8%)	RR 2 (1.1 to 3.65)	277 more per 1,000	LOW	

¹ Single blind trial
² Small sample size. 80% power calculation required 40 subjects.Only 36 patients completed the study and were analysed.
³ Unable to calculate as SD not given: 7.1 mean minutes vs. 14.4 mean minutes; reported p <0.001
⁴ Unable to calculate as SD not given: 8.0 mean minutes vs. 10.1 mean minutes; reported p <0.001.
⁵ Unable to calculate as SD not given: 34.1 mean minutes vs. 64.7 mean minutes; reported p <0.001.

Table 55: Nitrous oxide vs. intravenous midazolam; Wilson 2003 229

Author(s): Wilson 2003 **Date:** 2009-07-12

Question: Should IV midazolam vs. nitrous oxide be used for paediatric sedation?

Settings: Bibliography:

			Quality asses	remont				S	ummary of findir	ngs		
			Quality asses	Sillelit			No of pa	atients	Ef	fect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	IV midazolam	nitrous oxide	Relative (95% CI)	Absolute	Quality	ance
Duration of	f procedure (m	neasured wit	th: measured with m	edian minutes; rar	nge of scores	: -; Better indicated	l by less)					
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	40	-	not pooled ³	LOW	
Length of i	nduction (mea	sured with:	measured with med	lian minutes; range	of scores: -	; Better indicated by	y less)					
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	40	-	not pooled4	LOW	
Total time	(measured wit	h: median m	ninutes; range of sco	ores: -; Better indic	ated by less							
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	40	-	MD 34.4 (36.42 to 32.38)	LOW	
Patient pre	ference - num	ber of patier	nts									
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	19/37 (51.4%)	14/37 (0%)	RR 1.36 (0.81 to 2.28)	0 more per 1,000	LOW	
Recovery t	ime (range of	scores: -; Be	etter indicated by les	ss)								
1	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	40	40	-	MD 28.3 (26.10 to 30.80)	LOW	

¹ Unable to blind

Small sample size
 Results given as median times thus absolute effect could not be estimated; reported p<0.01
 Results given as median times thus absolute effect could not be estimated; reported p<0.001

Table 56: Nitrous oxide vs. oral midazolam; Wilson 2006; Wilson 2002; Wilson 2002 230-232

Author(s): Wilson 2002, BJD Wilson 2002 Anaesthesia Wilson 2006 Anaesthesia Question: 30% nitrous oxide/70% oxygen vs. oral midazolam for sedation in children

Settings: Dental

			Quality access	mont				Summa	ary of findings	3		
			Quality assess	sinent			No of pat	ients	Effe	ect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	30% nitrous oxide/70% oxygen	oral midazolam	Relative (95% CI)	Absolute	Quality	ance
Induction	time (range o	of scores: -; Be	tter indicated by le	ess)								
2		no serious limitations ¹	no serious inconsistency	no serious indirectness	serious ²	none	61	61	-	not pooled ³	MODERATE	
Recovery	time (range o	of scores: -; Bet	tter indicated by le	ess)								
1		no serious limitations ¹	no serious inconsistency	no serious indirectness	serious ²	none	26	26	-	not pooled4	MODERATE	<u> </u>
Duration (of procedure	(range of score	s: -; Better indicat	ted by less)	•			•			•	
1		no serious limitations ¹	no serious inconsistency	no serious indirectness	serious ²	none	61	61	-	not pooled ⁵	MODERATE	<u> </u>
Total time	(range of sc	ores: -; Better i	ndicated by less)									
2		no serious limitations ¹	no serious inconsistency	no serious indirectness	serious ²	none	61	61	-	not pooled ⁶	MODERATE	
Patient pr	eference (Qu	estionnaire)			•							
2		no serious limitations ¹	no serious inconsistency	no serious indirectness	serious ²	none	39/72 (54.2%)	41/73 (56.5%)	RR 0.97 (0.72 to 1.29)	16 fewer per 1,000	MODERATE	=

These were all randomised crossover trials. Trial data is combined where possible for 1-3 studies.

Small sample size

Two studies ^{230,232} reported mean (range) times and mean differences were not estimable; reported p<0.001 and p<0.0001 respectively. Another study ²³¹ reported induction times as median values

, 5 [5-10] minutes for N2O compared to 20 [5-65] minutes for oral midazolam; reported p<0.001

Two studies ^{230,232}: reported mean (range) times in Wilson 2002 and thus mean differences were not estimable; reported 20 minutes for N2O and 39.7 minutes for midazolam; p<0.0005. Wilson 2002 ²³¹ reported median times: 5 [5-10] minutes for N2O compared to 20 [5-65] minutes for oral midazolam; p<0.001

Studies were not able to be combined to provide a summary statistic due to differences in data reporting and missing data ^{230,232}

Two studies 230,232 reported mean (range) times and mean differences were not estimable; reported p<0.001 and p<0.0005 respectively. Another study 231 reported total time as median values, 35 [30-50] minutes for N2O compared to 100 [70-140] minutes for oral midazolam, p<0.001

Table 57: Nitrous oxide + EMLA vs. EMLA; Ekbom 2005 58

Author(s): Ekbom 2005
Question: Should Nitrous oxide + EMLA vs. EMLA be used for intravenous cannulation?

Settings: Hospital

			Quality asses	cmont				Sui	mmary of finding	s		
			Quality asses	Silielit			No of patie	nts	Effe	ct		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Other considerations	Nitrous oxide + EMLA	EMLA	Relative (95% CI)	Absolute	Quality	ance	
Completion	npletion of procedure											
1		· , 1		no serious indirectness	serious ²	none	25/25 (100%)	21/21 (0%)	RR 1.19 (0.99 to 1.43)	0 more per 1,000	VERY LOW	

¹ Ramdomisation and allocation concealment not well explained. Blinding not possible. ² Small study with no power calculations.

COMBINATION COMPARISONS

Table 58: Nitrous oxide + midazolam vs. medical air + intravenous midazolam; Averley 2004 19

Author(s): Averley 2004 Date: 2009-07-10

Question: 40% Nitrous oxide plus intravenous midazolam vs. medical air for sedation in children

Settings: Dental

			Quality assess	mont				Sumn	nary of finding	ıs		
			Quality assess	illelit			No of patien	ts	Eff	ect		Importan
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	IV midazolam & 40% Nitrous oxide	medical air	Relative (95% CI)	Absolute	Quality	ce
Completio	on of procedu	re: nitrous oxid	de vs. medical air									
1	randomised trial		no serious inconsistency	no serious indirectness	serious ¹	none	204/256 (79.7%)	94/176 (53.4%)	RR 1.49 (1.28 to 1.74)	261 more per 1,000	MODERATE	
Pain by V	by VAS score: nitrous oxide vs. medical air (range of scores: -; Better indicated by less)											
1	randomised trial		no serious inconsistency	no serious indirectness	serious	none	204	94	-	MD 0 (-0.28 to 0.28)	MODERATE	
Recovery	time: nitrous	oxide vs. med	ical air (range of s	cores: -; Better i	ndicated by	less)						
1	randomised trial		no serious inconsistency	no serious indirectness	serious ¹	none	204	94	-	MD -0.8 (-2.03 to 0.43)	MODERATE	
Anxiety: r	nitrous oxide	vs. medical air	(range of scores:	-; Better indicate	ed by less)							
1	randomised trial		no serious inconsistency	no serious indirectness	serious ¹	none	204	94	-	MD 0 (-0.32 to 0.32)	MODERATE	

¹ Greater than 20% did not complete intervention; greater in 1 group and this arm of the study was discontinued

10

2

Table 59: Nitrous oxide + midazolam vs. sevoflurane and nitrous oxide + midazolam; Averley 2004 19

Author(s): Averley 2004

Question 40% nitrous oxide plus intravenous midazolam vs. 3% sevoflurane and 40% nitrous oxide for sedation in children

Settings: Dental

			Ouglity assess					Summary of	of findings			
			Quality assess	Silielit			No of	patients	Ef	fect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	IV midazolam & 40% nitrous oxide	3% sevoflurane and 40% nitrous oxide	Relative (95% CI)	Absolute	Quality	ance
Completi	on of proced	ure	•		•							•
1	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	serious ¹	none	204/256 (79.7%)	249/267 (93.3%)	RR 0.85 (0.8 to 0.92)	139 fewer per 1,000	MODERATE	
Pain: VA	: VAS scale (range of scores: -; Better indicated by less)											
1		no serious limitations	no serious inconsistency	no serious indirectness	serious ¹	none	204	249	-	MD 0 (-0.24 to 0.24)	MODERATE	
Recovery	time (range	of scores: -; E	Better indicated by	y less)								•
1	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	serious ¹	none	204	249	-	MD -0.5 (- 1.21 to 0.21)	MODERATE	
Anxiety (range of sco	res: -; Better iı	ndicated by less)									
1	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	serious ¹	none	204	249	-	MD 0 (-0.24 to 0.24)	MODERATE	

¹ 20% of nitrous oxide group failed to complete procedure and are not included in further analysis.

Table 60: Nitrous oxide + midazolam and sevoflurane vs. medical air + midazolam; Averley 2004 19

Author(s): Averley 2004

Question 40% nitrous oxide and 3% sevoflurane & plus intravenous midazolam vs. medical air for sedation in children

Settings: Dental

			Quality assess	am am t				Summary	of findings			
			Quality assess	sment			No of patients		Ef	fect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	IV midazolam & 3%sevoflurane and 40% nitrous oxide	medical air	Relative (95% CI)	Absolute	Quality	ance
Completi	on of proced	ure	•		•							
1	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	serious ¹	none	249/267 (93.3%)	94/174 (54%)	RR 1.73 (1.5 to 1.99)	394 more per 1,000	MODERATE	:
Pain: VAS	n: VAS scale (range of scores: -; Better indicated by less)											
1			no serious inconsistency	no serious indirectness	serious ¹	none	249	94	-	MD 0.4 (- 0.31 to 0.31)	MODERATE	
Recovery	time (range	of scores: -; B	etter indicated by	/ less)								
1	randomised trial	no serious limitations	no serious inconsistency	no serious indirectness	serious ¹	none	249	94	-	MD -0.3 (- 1.82 to 1.22)	MODERATE	
Anxiety (Anxiety (range of scores: -; Better indicated by less)										•	
1			no serious inconsistency	no serious indirectness	serious ¹	none	249	94	=	MD 0.8 (- 0.31 to 0.31)	MODERATE	:

Greater than 20% of children did not complete procedure and group 1 (medical air) was terminated. Secondary analyses done only for those completing procedure.

1

8

6.7.2.2 Non RCT safety (adverse events)

2 3 4 5	Three non RCT observational studies in 8,220 patients assessed the safety of nitrous oxide. Two prospective cohort studies with greater than 100 subjects specifically assessed the safety of nitrous oxide ^{20,73} . One systematic review which contained information from two relevant paediatric RCTs was also included ⁶¹
6	The non RCT study characteristics for nitrous oxide are presented in Table 61.
7	The non RCT adverse event table for nitrous oxide are presented in Table 62.

1 Table 61: Nitrous oxide Non RCT study characteristics. Safety review

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting			
Prospective Cohort									
Babl et al, 2008 ²⁰ Australia	Tertiary children's hospital emergency department		Procedural sedation	60%	70% nitrous oxide – 72% patients 50% nitrous oxide – 28%	2 hours			
Gall et al, 2001 ⁷³ France	French hospitals; records of paediatric procedures		Procedural sedation with 50% nitrous oxide		50% nitrous oxide				
Retrospective Sy	Retrospective Systematic Review- 12 RCTs (2 paediatric studies with outcomes of interest)								
Faddy & Garlick, 2005 ⁶¹ Australia	Paediatric Emergency Department Laceration repair; fracture reduction		Procedural sedation		50% nitrous oxide				

Table 62: Nitrous oxide safety: Non RCT

		ADVERSE EVENTS, rate: % (n)							GRADE PROFILE				
Study type, reference,	Drug / Comparison	Procedure Aç	Age	Total N	Total N Aspiration	Respiratory in		atory intervention		ac arrest g either/or		oxygen saturation <90%	EVIDENC
country						oral- pharynge al airway	endotrac heal intubation	assisted ventilatio n	external cardiac massage	defibrillatio n	vomiting		E QUALITY
Prospective	Cohort studies			•		•	•		•				
Babel et al, 2008 ²⁰ Australia	70% nitrous oxide	Emergency procedures	0-18 years	72% (548)							4.7% (26/548)	0.18% (1/548)	VERY LOW
	50% nitrous oxide			13% (101)							3.9% (4/101)	0	VERY LOW
Gall et al, 2001 ⁷³ France	50% nitrous oxide	Emergency procedures including laceration repair, fracture reduction, cast remodelling, abscess drainage, lumbar puncture, dressing changes, bone-marrow aspiration, flexible bronchoscopy, gastroscopy, venous puncture and	<19 years	Adverse events reported as 'major' or 'minor' (terms not defined). 375 minor events (5%) and 25 major events (0.3%) All major events resolved within minutes after discontinuation of nitrous oxide. No patient needed		0	0	0					VERY LOW

		other miscellaneous procedures		intervention to maintain their airway.						
Retrospecti	ve Systematic Review-	12 RCTs (2 pae	diatric studi	es with outcome	s of interest)					
Faddy & Garlick, 2005 ⁶¹ Australia	50% nitrous oxide	Laceration repair; fracture reduction;	Mean age Study 1 (Burton et al, 1998): 3.7 (SD 1.6) years. Study 2 (Evans et al 1995) 10 (4- 15) years	60				0	0	VERY LOW

1	6.7.3 Evidence statements
2	6.7.3.1 RCT efficacy and safety
3	PLACEBO COMPARISONS
4	Nitrous oxide vs. oxygen
5	McCann 1996 160
6	There was no significant difference in:
7	Quiet behaviours [Low quality evidence]
8	
9	Nitrous oxide vs. oxygen
10	Primosch 1999 185
11	Compared to 100% oxygen, the nitrous oxide group had significantly:
12 13 14 15 16	 More quiet behaviours. The relative risk behaviour scores were not estimable due to use of an ordinal scale and incomplete statistical information. However, the authors report scores of 713 for the nitrous oxide group and 630 for the oxygen group with p<0.001. [Low quality evidence]
17 18	There was no significant difference between nitrous oxide/oxygen vs. 100% oxygen groups for the following variable:
19 20	 Oxygen saturation. Values in the two groups were exactly the same, 99± 0.01. [Low quality evidence]
21	
22	Nitrous oxide vs. nitrogen and oxygen
23	Fauroux 2004 ⁶⁴
24	Compared to 50% nitrogen and oxygen, the nitrous oxide group had significantly:
25	• fewer procedure failures [Low quality evidence]
26 27	 less pain immediately after the procedure as measured on the CHEOPS scale [low quality evidence]
28 29	 less pain (children >6 years old) immediately after the procedure as measured on a VAS scale [Low quality evidence]
30	
31	HEAD TO HEAD COMPARISONS

1	Nitrous oxide vs. benavioural management
2	Veerkamp 1993 ²²⁰ ; Veerkamp 1995 ^{220,222}
3 4 5	Two studies by the same authors with similar research methods and outcomes were meta- analysed. Anxiety was the only outcome of interest measured in this study. Behavioural observations were made using the Venham clinical rating scale.
6	Compared with behavioural management, the nitrous oxide group had significantly
7 8	 Less anxiety than the behavioural management group [Very low quality evidence]
9	
10	Nitrous oxide vs. transmucosal midazolam
11	Wilson 2007 ²³³
12	Compared to transmucosal midazolam, the nitrous oxide group had significantly:
13	Less induction time. [Low quality evidence]
14 15	Unable to calculate relative risk as SD not given: 7.1 mean minutes vs. 14.4 mean minutes; reported p < 0.001
16	Less procedure time. [Low quality evidence]
17 18	Unable to calculate relative risk as SD not given: 8.0 mean minutes vs. 10.1 mean minutes; reported p $<$ 0.001.
19	Less total time [Low quality evidence]
20 21	Unable to calculate relative risk as SD not given: 34.1 mean minutes vs. 64.7 mean minutes; reported p $<$ 0.001.
22 23	 More patients who preferred this method of sedation [Low quality evidence]
24	
25	Nitrous oxide vs. IV midazolam
26	Wilson 2003 ²²⁹
27	Compared to IV midazolam, the nitrous oxide group had significantly:
28	Shorter induction time [Low quality evidence]
29 30	Results given as median times and therefore absolute effect could not be estimated; reported p $<$ 0.001
31	Shorter procedure time [Low quality evidence]

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2	reported p<0.01
3	Shorter total time [Low quality evidence]
4	Shorter recovery time [Low quality evidence]
5	There was no significant difference in:
6	Patient preference. [Low quality evidence]
7	
8	Nitrous oxide vs. oral midazolam
9	Wilson 2006; Wilson 2002; Wilson 2002 ²³⁰⁻²³²
10	Compared to oral midazolam, the nitrous oxide group had significantly:
11	Shorter induction time [Moderate quality evidence]
12 13 14 15	Results given for two studies 230,232 as mean (range) times and mean differences were not estimable. The authors report p<0.001 and p<0.0001 respectively. The third study 231 reported induction times as median values , 5 [5-10] minutes for nitrous oxide compared to 20 [5-65] minutes for oral midazolam, p<0.001
16	Shorter procedure time in one study [Moderate quality evidence]
17 18	Studies were not able to be combined to provide a summary statistic due to differences in data reporting and missing data 230,232
19	Shorter recovery time [Moderate quality evidence]
20 21 22 23 24	Results given for two studies 230,232 . Mean (range) times were reported in Wilson 2002 and thus mean differences were not estimable. The authors reported 20 minutes for nitrous oxide and 39.7 minutes for midazlam p<0.0005. Median times were reported in Wilson 2002 231 , 5 [5-10] minutes for nitrous oxide compared to 20 [5-65] minutes for oral midazolam, p<0.001
25	Shorter total time [Moderate quality evidence]
26 27 28 29	Results given for two studies 230,232 as mean (range) times and mean differences were not estimable. The authors report p<0.001 and p<0.0005 respectively. The third study 231 reported total time as median values , 35 [30-50] minutes for nitrous oxide compared to 100 [70-140] minutes for oral midazolam, p<0.001
30	There was no significant difference in:
31	 Procedure time in two studies [Moderate quality evidence]^{231,232}
32 33 34	 Patient preferences [Moderate quality evidence] when the results of two studies were meta analysed ^{231,232}. The results of Wilson 2006²³⁰ were non significant but data was not available for meta-analysis.

1	
2	Nitrous oxide + EMLA vs. EMLA
3	Ekbom 2005 ⁵⁸
4 5 6	Compared to conventional treatment for intravenous cannulation with EMLA anaesthetic cream, children who received nitrous oxide + EMLA were reported by the authors to have a statistically significant difference in the following parameter:
7	 Pain as assessed by VAS [Very low quality].
8	There was no significant difference in:
9	Completion of procedure.
0	
1	COMBINATION COMPARISONS
2	Averley 2004 19
3	Nitrous oxide + midazolam vs. medical air + IV midazolam;
4	Nitrous oxide + midazolam vs. sevoflurane and nitrous oxide + midazolam
5	Nitrous oxide + midazolam and sevoflurane vs. medical air + midazolam
6 7	a) 40% nitrous oxide + IV Midazolam vs. medical air + IV midazolam
8	Compared to the medical air group, the nitrous oxide group had significantly:
9	More completed procedures [Moderate quality evidence]
20	There were no significant differences in:
21	Recovery time [Moderate quality evidence]
22	Pain by Vas score [Moderate quality evidence]
23	Anxiety by VAS score [Moderate quality evidence]
24 25 26	b) 40% nitrous oxide + IV Midazolam vs. 0.3% sevoflurane and 40% nitrous oxide + IV midazolam
27	Compared to the sevoflurane group, the nitrous oxide group had significantly:
28	Fewer completed procedures [Moderate quality evidence]
29	There were no significant differences in:

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1	• Recovery time
2	Pain by Vas score
3	 Anxiety by VAS score
4	
5 6	c) 0.3% sevoflurane and 40% nitrous oxide + IV Midazolam vs. medical air + IV midazolam
7 8	Compared to the medical air group, the sevoflurane + nitrous oxide group had significantly:
9	 More completed procedures [Moderate quality evidence]
10	There were no significant differences in:
11	Recovery time [Moderate quality evidence]
12	Pain by Vas score [Moderate quality evidence]
13	 Anxiety by VAS score [Moderate quality evidence]
14	Adverse events were reported for all three arms of this study as follows:
15	Six children in the seveoflurane group vomited clear fluids after treatment
16 17 18	 98% of all children had an oxygen saturation of 98% or above. The lowest saturation of 94% was recorded in one child in the medical air group.
19	6.7.3.2 NON-RCT safety (adverse events)
20 21	 There were no reported incidents requiring respiratory intervention including an oral pharangeal airway, endotracheal intubation or assisted ventilation ^{20,61,73}.
22 23	 There were no reported incidents of cardiac arrest requiring either/or external cardiac massage or defibrillation ^{20,61,73}.
24 25	 One study reported a 4.7% rate of vomiting with 70% nitrous oxide and a 3.9% rate of vomiting with 50% nitrous oxide ²⁰
26 27 28	 One study reported oxygen saturation <90% in 3.9% of patients using 70% nitrous oxide ²⁰. Two studies using 50% nitrous oxide reported that there were no patients with oxygen saturation <90% ^{20,61}.
29	6.7.4 GDG discussion of the evidence for nitrous oxide
30 31 32	The GDG noted that most of the evidence for nitrous oxide came from studies of painful procedures in the Emergency Department or the Dental clinic settings. The evidence level was low except in one RCT where the level was moderate.

The GDG agreed that both the efficacy and safety may be dependent on the concentration of nitrous oxide used. In almost all studies the dose was 50% or less in oxygen. Seventy percent oxygen was reported in a non-RCT in the ED setting.

The GDG noted that the evidence of efficacy in the RCTs was limited to the successful outcome of the procedure and that there were no data to allow the quality of the sedation to be assessed.

The GDG recognised that nitrous oxide is very widely used in UK dental clinics and it was appreciated that the success of administration of nitrous oxide relies on ability of the patient to breathe the gas continuously via a mask placed over the mouth and nose, or over the nose for dental procedures. Gaining and maintaining cooperation of a patient also relies on the skill of the healthcare practitioners.

In small uncooperative children nitrous oxide was not found to be any more effective than oxygen alone 160 but in cooperative children nitrous oxide could be used for a wide range of painful procedures provided the analgesia of the nitrous oxide was sufficient. In the dental setting the injection of local anaesthesia can be uncomfortable and the analgesia from nitrous oxide is effective for the local anaesthesia injection; thereafter, the value of nitrous oxide may relate to its euphoric and anxiolytic effect. The success rate of nitrous oxide in the dental setting was reported as approximately 50% and it was appreciated that this success rate was poor. Nevertheless it was argued by the dentists on the GDG that these studies were in children who had been referred to a dental clinical that specialised in the management of anxious children. In other dental clinics, where children may be less anxious, the success rate was considered to be much higher although no direct evidence was available to support this. Moreover the GDG dentists confirmed that children could be selected into those in whom nitrous oxide would and would not be sufficient for dental treatment; in their experience the success rate of nitrous oxide in selected children was at least 90%.

The advantages of nitrous oxide were considered to be that it was safe and short acting and highly effective in selected patient groups and settings. Occasionally it causes dysphoria and vomiting but this may be related to higher concentrations of nitrous oxide. The GDG appreciated the potential economic advantages of nitrous oxide successfully delivered in the dental clinic setting rather than anaesthesia in the dental hospital setting.

The GDG considered the safety of nitrous oxide. It was agreed that it was extremely unlikely that nitrous oxide concentration of 50% or less would cause unconsciousness provided the patient was fully conscious beforehand and that no other sedation drugs were used.. Equipment failure and medical contraindications to the use of nitrous oxide are rare but the GDG agreed that patients must be assessed and that practitioners must be trained to use nitrous oxide safely. The GDG agreed that nitrous oxide (used alone) was so safe that fasting was not required (although nitrous oxide may induce vomiting if the stomach was full) and that it could be safely administered by the dentist who was treating the patient.

The GDG debated the merits of combining nitrous oxide with other drugs to increase its efficacy. One RCT¹⁹ showed 80% of anxious children undergoing dental procedures were treated successfully by a combination of nitrous oxide with midazolam compared with only 54% of children with midazolam alone. In that study the combination of drugs did not cause unconsciousness but the GDG discussed the risk of unconsciousness caused by combining drugs. It was appreciated that intravenous and inhalational drugs could be titrated to achieve conscious sedation and that unconsciousness was extremely unlikely

provided the dental sedation team were skilled. Nevertheless it was agreed that there was a risk of unintended unconsciousness and that only specially trained dental sedation teams should use combinations of sedation drugs to achieve sedation. The GDG agreed that airway management skills and equipment are essential for combining nitrous oxide with other sedation drugs.

The general principle agreed by the GDG is that only sedation techniques commonly available in the NHS should be included in the economic analysis. Economic analysis was conducted for six broad groups (dental procedure in children, dental procedure in adolescents, short painful procedures, painless imaging, oesophago-gastroscopy and colonoscopy). The GDG agreed that nitrous oxide alone, and nitrous oxide combined with other drugs (nitrous oxide plus sevoflurane, nitrous oxide plus sevoflurane plus midazolam, and nitrous oxide plus midazolam) are commonly used in dental procedures in children, and that there is some evidence that they are effective and safe. It was therefore agreed that these strategies should be included in the economic analysis. Details of the considerations of cost-effectiveness with respect to using these strategies in dental procedure in children are given in section 6.12.4.2.

1 6.8 Sevoflurane and isoflurane

Matrix of sevoflurane / isoflurane comparators										
	/									
Key:										
Chloral hydrate = CH Fentanyl = F Isoflurane = I Ketamine=K Midazolam = M Propofol= P Nitrous oxide = N ₂ 0 Nitrous oxide and oxygen = N ₂ 0+02 Opioids = O Propofol= P Sevoflurane = S Triclofos sodium = TS										
Sevoflurane / isoflurane v	S									
	Reference	Tables and page	Evidence statements page							
Placebo										
Nil										
Head to head										
Nil										
Combinations										
S + NO + M vs. air + M	Averley 2004 ¹⁹	Table 63	261							
S + NO + M vs. NO + M	Averley 2004 ¹⁹	Table 64	261							
S + NO + vs. NO	Lahoud 2002 ¹³³	Table 65	262							
Safety										
RCTs										

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Table 66

Table 67

Table 66

Table 67

Table 66

Table 67

262

262

262

Lahoud 2002¹³³

Averley 2004¹⁹

De Sanctis Briggs

200551

Desaturation

Observational studies

Vomiting

Route of administration		
Nil		
Dose		
Nil		

6.8.1 Clinical methodological introduction

CLINICAL (QUESTIONS
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- For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is sevoflurane or isoflurane (with or without: analgesia, another drug or psychological techniques):
- Effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?
- Safe for sedation (at mild, moderate, and deep levels) in different settings?
- The literature was searched for systematic reviews and RCTs for the clinical efficacy of sevoflurane or isoflurane. The search was expanded to include non RCT observational studies for the safety of sevoflurane or isoflurane.
- 13 There were no systematic reviews identified for the use of opioids in paediatric sedation.
- Two parallel armed RCTs comparing sevoflurane in any route with other sedative drugs were assessed for efficacy and safety.
- One non RCT observational study in 640 patients assessed the safety of sevoflurane.
- There were no relevant studies conducted in children that assessed the safety and efficacy of sedation with isoflurane.
- Meta-analyses for RCTs were performed where drug interventions and comparisons and outcomes were sufficiently homogenous and studies were combined regardless of dose, duration of intervention, procedure (within painful and non-painful groups), setting (e.g. dentistry, accidents and emergencies) and age.

6.8.2 Evidence profiles

24 6.8.2.1 RCT evidence profiles for efficacy and safety

Study characteristics and methodological quality of the study are provided in Appendix D. GRADE tables for quality assessment and summary of findings are provided below.

27

COMBINATION COMPARISONS

Table 63: Sevoflurane + nitrous oxide + intravenous midazolam vs. medical air + intravenous midazolam; Averley 2004¹⁹

3 4 5 Date: 2009-08-27

Question: Should sevoflurane + nitrous oxide + iv midazolam titrated vs. medical air + iv midazolam titrated be used for sedation in children?

Settings: dental hospital

			Quality acco	ssmont			Summary of findings					
			Quality asse	SSIIIGIIL			No of par	tients		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	sevoflurane + nitrous oxide + iv midazolam titrated	medical air + iv midazolam titrated	Relative (95% CI)	Absolute	Quality	ance
number	of people wh	o complete p	rocedure									
	randomised trial	no serious limitations ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	249/267 (93.3%)	94/174 (54%)	RR 1.73 (1.5 to 1.99)	394 more per 1000 (from 270 more to 535 more)	HIGH	
Recover	Recovery time (Better indicated by less)											
	randomised trial	serious ²	no serious inconsistency	no serious indirectness	no serious imprecision	none	249	94	=	MD -0.3 (-1.55 to 0.95)	MODERATE	
child's p	erception of	pain (VAS sc	ore) (measured v	with: VAS; Bett	er indicated by	less)		-				•
	randomised trial	serious ²	no serious inconsistency	no serious indirectness	no serious imprecision	none	249	94	-	MD 0 (-0.28 to 0.28)	MODERATE	
Anxiety I	reported by	hild (VAS sc	ore) (measured v	vith: VAS; Bette	er indicated by	less)						
	randomised trial	serious ²	no serious inconsistency	no serious indirectness	no serious imprecision	none	249	94	-	MD 0 (-0.31 to 0.31)	MODERATE	
Parent's	satisfaction	score (range	of scores: 1-5; E	Better indicated	by more)							
	randomised trial	serious ²	no serious inconsistency	no serious indirectness	no serious imprecision	none	249	94	-	MD 0.1 (-0.05 to 0.25)	MODERATE	
vomiting												
	randomised trial		no serious inconsistency	no serious indirectness		none	6/249 (2.4%)	0/94 (0%)	RR 4.94 (0.28 to 86.84)	0 more per 1000 (from 0 fewer to 0 more)	VERY LOW	

¹ double blind with adequate allocation concealment and randomisation; ITT was performed for this outcome.

² double blind with adequate allocation concealment and randomisation; ITT was not performed for this outcome. ³ very wide 95% CI

Date: 2009-08-27 Question: Should sevoflurane + nitrous oxide + iv midazolam titrated vs. nitrous oxide + iv midazolam titrated be used for sedation in children?

Settings: dental hospital

			Ouglity sees				Summary of findings					
			Quality asse	ssment			No of pa	tients		Effect		Import
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	sevoflurane + nitrous oxide + iv midazolam titrated	nitrous oxide + iv midazolam titrated	Relative (95% CI)	Absolute	Quality	ance
number	of people wh	o complete p	rocedure									
1	randomised trial	no serious limitations ¹		no serious indirectness	no serious imprecision	none	249/267 (93.3%)	204/256 (79.7%)	RR 1.17 (1.09 to 1.25)	135 more per 1000 (from 72 more to 199 more)	HIGH	
Recovery	Recovery time (Better indicated by less)											
1	randomised trial	serious ²		no serious indirectness	no serious imprecision	none	249	204	=	MD 0.5 (-0.21 to 1.21)	MODERATE	
child's p	erception of	pain (VAS sc	ore) (measured v	vith: VAS; Bett	er indicated by	/ less)						
	randomised trial	serious ²		no serious indirectness	no serious imprecision	none	249	204	=	MD 0 (-0.24 to 0.24)	MODERATE	
Anxiety i	reported by o	child (VAS sc	ore) (measured v	vith: VAS; Bette	er indicated by	/ less)						
1	randomised trial	serious ²		no serious indirectness	no serious imprecision	none	249	204	-	MD 0 (-0.24 to 0.24)	MODERATE	
Parent's	satisfaction	score (range	of scores: 1-5; B	etter indicated	by more)							
1	randomised trial	serious ²		no serious indirectness	no serious imprecision	none	249	204	-	MD 0 (-0.1 to 0.1)	MODERATE	
vomiting												
	trial	serious ²		no serious indirectness		none	6/249 (2.4%)	0/204 (0%)	RR 10.66 (0.6 to 188.11)	0 more per 1000 (from 0 fewer to 0 more)	VERY LOW	

Table 64: Sevoflurane + nitrous oxide + intravenous midazolam vs. nitrous oxide + intravenous midazolam; Averley 2004 19

double blind with adequate allocation concealment and randomisation; ITT for this outcome double blind with adequate allocation concealment and randomisation; ITT was not performed for this outcome. very wide 95% CI

8

Table 65: Sevoflurane + nitrous oxide vs. nitrous oxide; Lahoud 2002¹³³

Date: 2009-08-27

Question: Should sevoflurane + nitrous oxide vs. nitrous oxide be used for sedation in children?

Settings: dental hospital

			Quality ass	ossmont			Summary of findings					
			Quality ass	essinent			No of patients Effect				Import	
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	sevoflurane + nitrous oxide	nitrous oxide	Relative (95% CI)	Absolute	Quality	ance
number of children who complete procedure												
1	randomised trial		no serious inconsistency		no serious imprecision	none	215/241 (89.2%)	89/170 (52.4%)	RR 1.7 (1.47 to 1.98)	367 more per 1000 (from 246 more to 514 more)	MODERATE	
number of children who had a score of anxiety (Venham score = 5) (Venham score)												
1	randomised trial	- ,	no serious inconsistency	no serious indirectness	very serious ³	none	0/215 (0%)	2/89 (2.2%)	RR 0.08 (0 to 1.72)	20 fewer per 1000 (from 22 fewer to 16 more)	VERY LOW	
Number of	of children w	ho were sati	isfied with the tre	eatment (rated t	reatment as ex	ccellent)						
1	randomised trial	, ,	no serious inconsistency		no serious imprecision	none	188/215 (87.4%)	74/89 (83.1%)	RR 1.05 (0.95 to 1.17)	42 more per 1000 (from 42 fewer to 141 more)	LOW	
Adverse	events: Oxy	gen desatura	ation <90%									
1	randomised trial		no serious inconsistency		no serious imprecision	none	0/215 (100%)	0/89 (100%)	not pooled	-	LOW	

¹ unclear if assessor was blind and no detail on randomisation generation; adequate allocation concealment; ITT analysis performed for this outcome

² unclear if assessor was blind and no detail on randomisation generation; adequate allocation concealment; ITT analysis was not performed for this outcome

³ very wide 95% CI

1 6.8.2.2 Non RCT evidence profiles for safety

- One non-RCT observational study (n=640) assessed the safety of sevoflurane 51.
- 3 The non RCT study characteristics for midazolam are presented in Table 66.
- The non RCT adverse event table for midazolam is presented in Table 67.

Table 66: Sevoflurane Non RCT Study Characteristics Safety Review

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
De Sanctis Briggs 2005 ⁵¹ , Spain	Centre for MRI	Not stated	Deep Sedation for MRI examinations N= 640 infants age 1 day – 12 months	46.5%	Inhaled sevoflurane 7% in 50% nitrous oxide for induction; followed by sevoflurane 1.8-2% in 50% nitrous oxide for manintenance	Sedation fasting protocol

3

1 Table 67: Sevoflurane Safety: Non RCTs

							ADV	ERSE EVE	NTS, rate:	% (n)			GRADE PROFILE
Study type, reference,	Drug / Comparison	Procedure	Age	Total N	Aspirat -				Cardiac arrest requiring either/or			oxygen desaturat	EVIDENCE
country						oral- pharyn geal airway	endotrac heal intubatio n	assisted ventilati on		defibril lation	vomiting	ion <90%	QUALITY
De Sanctis Briggs 2005 ⁵¹ ,	sevoflurane 1.8-2% in 50% nitrous oxide	MRI	1 day – 12 months old	640							1/640 = 0.16%	0/640 = 0%	VERY LOW
Spain			15% < 1 month old	They state that 627/640 (97.9%) of patients experienced									
			39% 1-6 months old	no complications (defined as vomiting, mild									
			45% 7-12 months old	or severe hypoxia, prolonged sedation, or agitation									

1	6.8.3 Evidence statements
2	6.8.3.1 RCT efficacy and safety
3	COMBINATION COMPARISONS
4	Sevoflurane + nitrous oxide + IV midazolam vs. medical air + IV midazolam
5	Averley, 2004 19
6 7	Compared with medical air and intravenous midazolam group, the sevoflurane + nitrous oxide + intravenous midazolam group had significantly:
8	 More completed procedures [high quality evidence]
9	There was no significant difference in:
10	Recovery time [moderate quality evidence]
11	Child's perception of pain score (VAS) [moderate quality evidence]
12	 Anxiety reported by child (VAS) [moderate quality evidence]
13	Parent's satisfaction score (scale 1-5) [moderate quality evidence]
14	Vomiting [very low quality evidence]
15	
16	Sevoflurane + nitrous oxide + IV midazolam vs. nitrous oxide + IV midazolam
17	Averley, 2004 19
18 19	Compared with nitrous oxide + intravenous midazolam group, the sevoflurane + nitrous oxide + intravenous midazolam group had significantly:
20	More completed procedures [high quality evidence]
21	There was no significant difference in:
22	Recovery time [moderate quality evidence]
23	Child's perception of pain (VAS) [moderate quality evidence]
24	Anxiety reported by child (VAS) [moderate quality evidence]
25	 Parent's satisfaction score (scale 1-5) [moderate quality evidence]
26	Vomiting [very low quality evidence]
27	
28	Sevoflurane + nitrous oxide vs. nitrous oxide

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1	Lahoud 2002 ¹³³
2	Compared with the nitrous oxide group, the sevoflurane + nitrous oxide group had significantly:
4	More completed procedures [moderate quality evidence]
5	There was no significant difference in:
6 7	 Anxiety (proportion of patients) (Venham score = 5) [very low quality evidence]
8	 Parents' satisfaction (number of parents) [low quality evidence]
9	There were no events of:
10	 Oxygen saturation < 90% [low quality evidence]
11	6.8.3.2 Non RCT safety (adverse events)
12	For the characteristics of studies and outcome data refer to Table 66 and Table 67.
13	One study ⁵¹ reported rates of:
14	• Vomiting: 0.16%
15	Oxygen desaturation <90%: 0%
16	6.8.4 GDG discussion of the evidence for sevoflurane and isoflurane
17 18 19 20 21 22 23 24 25 26 27 28	Three studies ¹⁹ , ¹³³ , ⁵¹ informed the GDG discussion on sevoflurane. Sevoflurane is an anaesthetic agent and the GDG discussed whether there was an appreciable risk of accidental anaesthesia. Two ¹⁹ , ¹³³ of the three studies were RCTs in which sevoflurane had been used to sedate anxious children for dental procedures in a specialist dental clinic. The GDG appreciated that sevoflurane was being used in a similar fashion to nitrous oxide in that it required the patient to tolerate breathing the vapour via a nasal mask. In low doses sevoflurane was reported to not cause anaesthesia and its success therefore relied on a degree of cooperation of the patient. The dental studies were in anxious children up to the age of 14. Concentrations of up to 0.3% were used with (or without) 40% nitrous oxide and also with intravenous midazolam titrated to achieve satisfactory compliance for the dental procedure. The addition of sevoflurane was found to increase the completion rate of dental treatment.
29 30 31	The GDG agreed that this is a successful technique but that it required special expertise of a trained sedation team, and that airway management skills and equipment are essential for this drug in this setting.
32 33 34 35	The other study ⁵¹ considered was a descriptive account of 640 infants who were sedated by a combination of sevoflurane and nitrous oxide for painless imaging. The dose of sevoflurane used was 1.8-2% and even though the GDG understood that the conscious level had not been tested, the GDG decided that it was very likely that the

infants had been anaesthetised by this dose.

35

The GDG discussed the advantages of sevoflurane sedation over sevoflurane anaesthesia. In certain settings, in which the patient needs to cooperate with a procedure, such as a dental procedure, sedation may be appropriate. In other situations, such as painless imaging where an uncooperative child needs to be immobile and asleep, the dose of sevoflurane required to cause sleep is likely to cause anaesthesia. The GDG agreed that it was safer to assume that that patients were anaesthetised in this setting and that they would therefore need to managed as though they had a short acting anaesthetic rather than sedation. Overall the GDG agreed that sevoflurane should only be used by specially trained sedation teams.

The GDG agreed that only sedation techniques commonly available in the NHS should be included in the economic analysis. Economic analysis was conducted for six broad groups (dental procedure in children, dental procedure in adolescents, short painful procedures, painless imaging, oesophago-gastroscopy and colonoscopy). Sevoflurane combined with other drugs (sevoflurane plus nitrous oxide, sevoflurane plus nitrous oxide plus midazolam) were felt to be strategies commonly used in dental procedures in children. There is evidence that these drug combinations are effective and safe. The GDG therefore agreed that they should be included in the economic analysis. Details of the considerations of cost-effectiveness with respect to using these combination strategies in dental procedures in children are given in section 6.12.4.2.

1 6.9 Propofol

Matrix of propof	ol comparators		
Key: Chloral hydrate = Cl Fentanyl = F Isoflurane = I Ketamine=K Local anaesthesia = Midazolam = M Nitrous oxide = N ₂ 0 Nitrous oxide and ox Opioids = O Propofol= P Sevoflurane = S Triclofos sodium = TS	LA xygen = N₂0+02		
	Reference	Tables	Evidence statements page
Placebo			
Nil			
Head to head			
Nil			
Combinations			
P vs. M + K + F	Vardi 2002 ²¹⁹	Table 62	278
Safety			
RCTs			
Assisted ventilation	Vardi 2002 ²¹⁹	Table 69 Table 70	278
ET intubation	Vardi 2002 ²¹⁹	Table 69 Table 70	278
Observational studies	Melamed 1976 ¹⁶⁶ Bassett 2003 ²⁵ Barbi 2006 ²⁴ Vespasiano 2007 ²²⁴ Larsen 2009 ¹³⁴ Cravero 2009 ⁴⁶	Table 69 Table 70	278

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Barbi 2003²³

Route of administration		
Nil		
Dose		
Nil		

6.9.1 Clinical methodological introduction

2	CLINICAL QUESTIONS:	
3 4 5	For children and young people under the age of 19 undergoing diagnost therapeutic procedures, is propofol (with or without: analgesia, another dipsychological techniques):	
6 7 8	- Effective for sedation (at minimal, moderate, and deep levels) in compo care, with analgesia alone, with another sedation drug, with psychologica with general anaesthesia?	
9	- Safe for sedation (at mild, moderate, and deep levels) in different setti	ngs?
10 11 12	The literature was searched for systematic reviews and RCTs for the clinic propofol. The search was expanded to include non-RCT observational strafety of propofol.	•
13 14	There were no systematic reviews identified for the use of propofol in pa sedation.	ediatric
15 16	One RCT comparing intravenous propofol with other sedative drug was a efficacy and safety.	ssessed for
17 18	Seven non-RCTs observational studies in 64,115 patients assessed the safintravenous propofol.	ety of
19	Meta-analyses were not performed as there was only one RCT.	
20	6.9.2 Evidence profiles	
21	6.9.2.1 RCT evidence profiles for efficacy and safety	
22 23	Study characteristics and methodological quality of the study are provide D. GRADE tables for quality assessment and summary of findings are pro	• •

2

24

COMBINATION COMPARISONS

Table 68: Intravenous propofol + propofol maintenance + local anaesthesia vs. intravenous midazolam + intravenous ketamine + intravenous fentanyl; Vardi 2002 ²¹⁹

Date: 2009-09-08

2

3

4567890 10

Question: Should intravenous propofol plus propofol maintenance plus local anaesthesia vs. intravenous midazolam plus intravenous ketamine plus intravenous fentanyl be used in children and young people undergoing diagnostic and therapeutic procedures?

Settings: paediatric critical care unit (convenient facility for procedures)

Bibliography: Vardi 2002 (mixed procedures: Intraarticular steroid injection, bronchoscopy, bone marrow aspiration/biopsy, transesophageal echocardiography, PEG/Gastroscopy, Other: central line placement, intrathechal injections, removal of tunnelled central venous catheter, wound care, and chest tube placement)

			Ouglity and					Summary of find	dings			
			Quality ass	essment			No of p	atients	E	ffect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous propofol plus propofol maintenance plus local anaesthesia	intravenous midazolam plus intravenous ketamine plus intravenous fentanyl	Relative (95% CI)	Absolute	Quality	Importance
Complet	ion of proce	dure										
	randomised trial	very serious¹		no serious indirectness	no serious imprecision	none	58/58 (100%)	47/47 (100%)	not estimable	-	LOW	
Induction	nduction time: period between administration of the first dose of sedation and the time when patients were unresponsive to verbal or tactile stimulation (Better indicated by less)											
1	randomised trial	very serious¹		no serious indirectness	serious ²	none	58	47	-	not pooled	VERY LOW	
Duration	of procedu	re (Better in	dicated by less)									
1	randomised trial	1		no serious indirectness	no serious imprecision	none	58	47	-	MD -2 (-9.28 to 5.28) ³	LOW	
Recover	y time: from	administrat	ion of last seda	tion dose to w	hen patients o	pened their eyes	or gave appropriate re	esponse (Better indicate	ed by less)			
	randomised trial	1		no serious indirectness	no serious imprecision	none	58	47	=	MD -27 (-35.22 to -18.78) ⁴	LOW	
							ician, paediatric intens ge of scores: 1-6; Bette	ivist delivering sedation rindicated by more)	n, physiciar	n performing pr	ocedure) using a
1	randomised trial	very serious ¹		no serious indirectness	serious ⁵	none	58	47	_	MD 0.26 (-0.08 to 0.59) ⁶ MD 0.26 (-0.08 to 0.59)	VERY LOW	

Satisfaction at sedation period assessed by four observers (paediatric nurse, resident physician, paediatric intensivist delivering sedation, physician performing procedure) using a validated scale (measured with: Ramsay scale (maximum score = 6) at procedure period; range of scores: 1-6; Better indicated by more)

1	randomised trial			no serious indirectness	serious ⁵	none	58	47	-	MD 0.25 (0.03 to 0.47) ⁷	VERY LOW	
Adverse	events: Ass	isted ventil	ation: bag/mask									
1	randomised trial	, ,		no serious indirectness	very serious ⁸	none	10/58 (17.2%)	3/47 (6.4%)	RR 2.70 (0.79 to 9.26) ⁹	109 more per 1000 (from 13 fewer to 529 more) 0 more per 1,000	VERY LOW	
Adverse	events: End	lotracheal ir	ntubation									
1	randomised trial	, ,		no serious indirectness	very serious ⁸	none	0/58 (0%)	1/47 (2.1%)	RR 0.27 (0.01 to 6.51) ¹⁰	15 fewer per 1000 (from 21 fewer to 116 more) 0 fewer per 1,000	VERY LOW	

Vardi 2002: concealment and ITT not stated and blinding of patients and assessors not stated or unclear; small study
Vardi 2002: it was not possible to calculate RR based on data reported. The study stated that induction time was 40-60 seconds for both groups
Vardi 2002: p=0.59
Vardi 2002: p<0.00001
Vardi 2002: imprecise
Vardi 2002: p=0.13
Vardi 2002: p=0.03
Vardi 2002: p=0.03
Vardi 2002: p=0.01
Vardi 2002: p=0.11
Vardi 2002: p=0.11

6.9.2.2 Non RCT evidence profiles for safety

Seven non RCT observational studies (n=64,115) assessed the safety of propofol ²³^{25,46,134,167,224} There were six prospective studies, and one retrospective study conducted for the following procedues: imaging procedures (2), accidents and emergencies procedures (1) as well for GI and oncology procedures (2) and inpatients and outpatiens (2).

- 7 The non RCT study characteristics for midazolam are presented in Table 69.
- The non RCT adverse event table for midazolam is presented in Table 70.

9

1

Table 69: Propofol Non RCT Study Characteristics Safety Review

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Merola 1995 ¹⁶⁷ , USA	Imaging (MRI and CT suites) (99% ambulatory)	I-II Other ASA I-II: 99.34% (452/455) Other: 0.66% (3/455)	Not stated	Not stated	PRO or CH: PRO: • 2 mg/kg bolus after iv access + dilute PRO by gravity titrated infusion at a rate of 80-140 mcg/kg/min • Children ≥1 y.o. generally received PRO unless they had poor venous access or unless there was a strong parental preference for not inserting an i,v, catheter CH: • Children <1 y.o. generally received CH 75 mg/kg to a maximum of 2g due to difficulty in establishing i.v. access • Younger children were often swaddled and provided with a pacifier • Parents accompanied the children Concurrent: • All patients received O2 at 2 L/min by nasal cannule during procedures (scans)	Not stated
Barbi 2003 ²³ , Italy	Paediatric sedation unit (admitted to paediatric gastroenterology and oncology wards)	I-II	Deep (91% (963/1059) of children experienced transient general angesthesia at any	50% (411/827)	LA/TA/Atropine/PRO/GlucoSol LA: Lidocaine/prilocaine: 1 to 10 mg Lido/PRO for 1st syringe in children without a central line TA: EMLA cream	Clear fluids not allowed for 3 hrs, infant formula and nonhuman milk for 6 hours and solids for hours

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Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
			time during the procedure)		Atropine: 0.01 mg/kg as premedication PRO:	
					• 2mg/kg in children ≤8 y.o.	
					• 1 to 2 mg/kg in >8 y.o.	
					 repeated dose 0.5-1 mg/kg or continuous 6-9 mg/kg per hour for long procedures 	
					GlucoSol: continuous infusion maintenance	
					Concurrent:	
					O2 administered after the 2nd year of study at 6L/min by mask close to face to anticipate hypoxemia; O2 was administered during procedure from beginning of study for children undergoing painful procedures mostly those with cancer	
Bassett 2003 ²⁵ , USA	Emergency department	I-II ASA I: 96% (379/393 procedures) ASA II: 4% (14/393 procedures)	Procedural sedation	67% (263/392)	PRO/Opioid analgesics: PRO: IV initial dose of 1 mg/kg (max 40 mg); IV supplemental doses of 0.5 mg/kg (max 20 mg) at discretion of physician Bolus over 1 to 2 min, 20 secs between each dose; titrated to	Minimum of 3 hrs for solids and liquids
					tolerance of noxious stimuli without patient complaint Morphine:	
					• 0.1 mg/kg (max 5 mg) for	

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Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
					significant pain on presentation to emergency department Fentanyl:	
					1 to 2 mcg/kg (max 50 mcg) for children who had not received narcotics or were still with significant pain	
					Concurrent: supplemental O2 at 10 L/min with a bag-valve mask to face before initiation and during procedure; not used for assistance with respirations unless requested by physician and suction available at bedside	
Barbi 2006 ²⁴ , Italy	Department of gastroenterology (Endoscopic room)	1-11	Procedural sedation	47% (337/716)	TA/Atropine/IV PRO/LA/GlucoSol or Ringer'sSol TA: EMLA cream Atropine: 0.010-0.015 mg/kg PRO infusion: in children up to 8 y.o.:2mg/kg in children >8 y.o.:1-2 mg/kg repeated dose 0.5-1 mg/kg or continuous 6-9 mg/kg per hour for long procedures LA: lidocaine 1 mg for every 10 mg of PRO for the first syringe in all children	Clear fluids not allowed for 3 hrs, infant formula and nonhuman milk for 6 hrs and solids for 8 hrs
					GlucoSol: continuous infusion maintenance (for age and weight) for children >5 y.o. Concurrent:	

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	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
				O2 administered after the 2nd year of study at 6L/min by mask close to face to anticipate hypoxemia; O2 was administered during procedure from beginning of study for children undergoing painful procedures mostly those with cancer	
MRI 42.8% (3126/7304), radiology 22.5% 43/7304), short stay unit 26.2% 914/7304), special iagnostics unit 4.3% 14/7304), PICU 2% 146/7304), Other 2.2% (161/7304)	ASA I-II: 99.7% (7285/7304) ASA > II: 2.5% (18/7304) ASA unassigned: 0.014% (1/7304)	Deep	Not stated	PRO/PRO maintenance/LA PRO: • rarely <2 mg/kg • intermittent bolus doses for shorter interventions and continuous infusion after initial bolus for longer interventions • continuous infusion initiated at 150 mcg/kg/min titrated as required PRO maintenance: • supplemental boluses 1-2 mg/kg LA: • lidocaine doses at discretion of intensivist Concurrent:	Not stated
9 ic	(3126/7304), adiology 22.5% 43/7304), short stay unit 26.2% 214/7304), special agnostics unit 4.3% 4/7304), PICU 2% 46/7304), Other	(3126/7304), adiology 22.5% 43/7304), short stay unit 26.2% 214/7304), special agnostics unit 4.3% 4/7304), PICU 2% 46/7304), Other (7285/7304) (7285/7304) ASA > II: 2.5% (18/7304) ASA unassigned: 0.014% (1/7304)	(3126/7304), adiology 22.5% (7285/7304) ASA > II: 2.5% (18/7304), short stay unit 26.2% (18/7304), special agnostics unit 4.3% (4/7304), PICU 2% (46/7304), Other (7285/7304)	(3126/7304), adiology 22.5% (7285/7304) ASA > II: 2.5% (18/7304), short stay unit 26.2% (18/7304), special agnostics unit 4.3% (4/7304), PICU 2% (46/7304), Other (7285/7304)	MRI 42.8% (3126/7304), adiology 22.5% (13/7304), short starty unit 26.2% (14/7304), PICU 2% (46/7304), PICU 2% (46/7304), Other .2% (161/7304) Beginning of study for children undergoing painful procedures mostly those with cancer Not stated PRO/PRO maintenance/LA PRO: • rarely <2 mg/kg • intermittent bolus doses for shorter interventions and continuous infusion after initial bolus for longer interventions • continuous infusion initiated at 150 mcg/kg/min titrated as required PRO maintenance: • supplemental boluses 1-2 mg/kg LA: • lidocaine doses at discretion of intensivist

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Larsen 2009 ¹³⁴ , USA	Database established by paediatric intensive care to track outpatients requiring propofol sedation for diagnostic therapeutic proceudures Retrospective analysis of database to track each outpatient paediatric procedure requiring propofol	Not stated	Not stated	52% (2463/4716)	Intravenous propofol sedation sufficient to reach a level of sedation not requiring endotracheal intubation	Not stated
Cravero 2009 ⁴⁶ , USA	Outside the operating room Collaborative database of adverse events from 37 locations with data on paediatric sedation/anaesthesia. Prospectively enrolled consecutive patients receiving sedation or sedation/anaesthesia for procedures. Primary inclusion was the need for some form of sedation/anaesthesia to perform a diagnostic or therapeutic procedure outside the operating	ASA ≤ II (41191/49836) ASA > II 18% (8915/49836)	Not clear whether propofol was used for sedation or anaesthesia	55% (27420/48836)	Not clear	Not stated

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1 Table 70: Propofol Safety: Non RCTs

							AD	VERSE EVENT	S, rate: % (r	1)			GRADE PROFILE
Study type, reference,	Drug /	Procedure	Age	Total N						c arrest either/or		oxygen	EVIDENC
country	Comparison		J		Aspiration	oral- pharyngea I airway	endotrach eal intubation	assisted ventilation	external cardiac massage	defibrillatio n	vomiting	desaturati on <90%	E QUALITY
Merola 1995 ¹⁶⁷ , USA	PRO/O2 and CH/O2	Scans of the head, thorax, abdomen, pelvis and spine	Overall age range: <1 mo to 17y PRO range: <1 mo- to 17y ≥1 y: 98% (318/324) <15y: 4% (13/324) CH range: <1 mo to 7 y <1 y: 51% (57/131)	Total: 455 324 PRO 131 CH		0% (airway compro- mise)	0%	0% (controlled ventilation)			0%		VERY LOW
Bassett 2003 ²⁵ , USA	PRO/Mo or Fenta (analgesics)	378/393 procedures)	Overall age range: 1 to 18y Median age: 8 y	393 procedures in 392 children (1 child sedated twice)	0%	3% (11/392) (partial airway obstruct- tion)	0%	0.8% (3/392) (bag- valve-mask)	0% (cardiopulm onary arrest)			5% (20/392)	VERY LOW
Barbi 2003 ²³ , Italy	LA/ TA/Atropine/PRO	Upper endoscopies, colonoscopies, painful procedures	<1y to <10y: 61% (503/827) 10y to <21y:	Total: 1059 procedures in 827 children				Total: 0.5% (5/1059 procedures)			1.05% (3/827) (repeate d vomiting	6.04% (64/1059 procedures)	VERY LOW

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			39% (324/827)	upper endoscopy: 483 procedures in 405 children colonoscopy: 289 procedures in 249 children painful: 287 procedures in 173 children				endoscopies 0.83% (4/483 procedures) colonoscopies 0% (0/289 procedures) painful 0.35% (1/287 procedures)		during procedur e) 0.35% (1/827) (3hr after discharg e)		
Barbi 2006 ²⁴	LA/TA/Atropine/P RO	Upper gastrointestinal endoscopy procedures	<1 to <10y: 65% (463/716) 10y to <21y: 35% (253/716)	811 procedures in 716 children				Total: 0.7% (6/811 procedures) [3 of these required bagvalve-mask: 0.4% (3/811)]			7% (58/811 procedures)	VERY LOW
Vespasiano 2007 ²²⁴ , USA	PRO/LA	MRI, CT, nuclear medicine, lumbar puncture, intratechal chemotherapy, bone marrow aspirates, electroencephal ogram, evoked potentials, hearing tests	Overall age range: 0 mo to 21y 0 to 1mo: 0.4% (29/7304) 1mo to 1y: 1.9% (139/7304) 1 tp 5y: 56% (4076/7304) >5y: 42% (3060/7304)	7, 304	0.01% (1/7304)	0.96% (70/304) (oral airway) 1.57% (115/7304) (nasal trumpet)	0.03% (2/7304)	0.37% (27/7304) (bag and mask)	0% (cardiac arrest)		4.6% (338/7304)	VERY LOW
Larsen 2009 ¹³⁴	IV propofol					0.04% (2/4716) (mask)	0.02% (1/4716)	0.02% (1/4716)((bag- valve)				VERY LOW

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Cravero 2009 ⁴⁶ , USA	Propofol used as the sole or primary sedative in 49,836 sedations/anaesth esia encounters: -20.4% (10149/49836) used in addition to:	diagnoses including: neurological	0 months to 8 years: 71% (35396/49836) > 8 years: 29% (14440/49836)	49, 836 sedation encounters	N=4 rate: 0.9	Airway obstruction: N=432, rate: 93.2 Emergency airway consultation (does not applied to cases delivered by anaesthesiologists): N=7, rate: 1.5	(cardiac arrest) N=2 rate: 0.4		(during sedation) N=49 rate: 10.6	N=716 rate: 154.4	VERY LOW Indirect population Difficult to draw conclusions as unclear whether
	Midazolam [7.5% (3766/49 836)], Ketamine [1.76% (879/49, 836)], Chloral hydrate [0.3% (139/49 836)], Opioids ALL TYPES [10% (5061/49836)], OTHER [0.61% (304/49, 836)] -79.6% (39687/49836) for the remaining encounters	orthopaedic (3.9%) congenital heart disease (2.4%), other defined diagnoses (10.6%)				Reported rates Inadequate anaesthesi		te: 85			sedation used for sedation or anaesthesi a and unclear dose

1 6.9.3 Evidence statements 2 6.9.3.1 RCT efficacy and safety 3 COMBINATION COMPARISONS 4 IV propofol + propofol maintenance + local anaesthesia vs. IV midazolam + IV 5 ketamine + IV fentanyl Vardi 2002 219 6 7 All patients completed the procedure [low quality evidence] 8 Induction time was similar for both groups (as stated in the study) [very low 9 quality] 10 Compared with children receiving intravenous midazolam + intravenous ketamine + 11 intravenous fentanyl, children receiving intravenous propofol + propofol maintenance + 12 local anaesthesia had significantly: 13 Faster recovery time (minutes) [low quality evidence] 14 Better satisfaction at sedation period (Ramsay scale) [very low quality 15 evidencel 16 There was no significant difference in: 17 Duration of procedure (minutes) [low quality evidence] 18 Satisfaction at induction period (Ramsay scale) [very low quality evidence] 19 Assisted ventilation (bag-mask) [very low quality evidence] 20 Endotracheal intubation [very low quality evidence] 21 6.9.3.2 Non RCT safety (adverse events) 22 For the characteristics of studies and outcome data on propofol refer to Table 69 and Table 70 23 24 Two studies reported rates of aspiration: from 0% to 0.01% ^{25,224} 25 Four studies reported rates of oral-pharyngeal airway intervention: from 0% to 3% 25,134,167,224 26 27 Four studies reported rates of endotracheal intubation: from 0% to 0.03% 28 29 Six studies reported rates of assisted ventilation - either bag-valve mask or controlled: from 0% to 0.8% ^{23-25,134,167,224} 30

1 2	 Two studies reported rates of External cardiac massage: there were no events of cardiac ²²⁴ or cardiopulmonary ²⁵ arrest
3	• Three studies reported rates of Vomiting: from 0% to 1.05% ^{23,24,167} .
4 5	 Four studies reported rates of oxygen desaturation <90%: from 0% to 7% ²⁵, ^{23,24,224}
6 7 8 9	In one study 46 it was unclear whether sedation was used for sedation or anaesthesia and how much dose of the propofol was administered. Based on a total of 49, 836 sedations/anaesthesia encounters, the study reported a range of complications with rates (per 10,000) including:
10	 Aspiration: rate 0.9 (n=4)
11	 Airway obstruction: rate 93.2 (n=432)
12 13	 Emergency airway consultation (does not apply to cases delivered by anaesthesiologists): rate 1.5 (n=7)
14	• Cardiac arrest: rate 0.4 (n=2)
15	 Vomiting during sedation: rate 10.6 (n=49)
16	 Oxygen desaturation <90%: rate 154.4 (n=716)
17	 Inadequate anaesthesia: rate: 85, (n=392)
18	6.9.4 GDG discussion of the evidence for propofol
19 20 21 22 23 24 25 26 27 28 29 30	Propofol, being a short acting intravenous anaesthetic agent, can be titrated to achieve any target level of sedation and anaesthesia. In the evidence examined the success rate of propofol was not always specifically stated but was assumed by the GDG to be 100%. The true level of sedation was often not stated. The GDG appreciated that the difference between sedative and anaesthesia doses was small and that unintentional anaesthesia was a risk with this drug. The GDG agreed that doses above 3mg/kg are likely to cause unconsciousness indistinguishable from anaesthesia. It was noted that doses necessary to cause sedation may depend upon the procedure. For example the dose required for a painless procedure would be less than for a painful procedure. The GDG noted that the dose of propofol required for a painful procedure maybe reduced by the use of analgesia and in this respect the combination of an opioid with propofol may reduce the doses of both drugs.
31 32 33 34 35	Seven studies ²²⁴ , ²⁴ , ²³ , ²⁵ , ¹⁶⁷ , ²¹⁹ , were considered by the GDG (very low level evidence). The studies involved procedures ranging from painless imaging, painful ED procedures and endoscopy. The target sedation level was deep or not stated. The GDG considered the doses used and agreed that many of the children would have been anesthetised at some stage.
36 37 38 39	The safety of propofol was discussed. In one large case series ²⁵ the incidence of oxygen desaturation was 7% and the need for an airway device was approximately 3%. The GDG agreed that tracheal intubation would occasionally be required and that propofol should only be used by teams who had adequate training to manage anaesthesia. Sedation in children and young people: full guideline DRAFT (May 2010) Page 279 of 371

The GDG noted that propofol was used in two studies²⁴, ²³ for children undergoing endoscopy. Propofol was being used without any airway device and the GDG agreed that practitioners would need special training to ensure that the airway was not obstructed by the insertion of the endoscope. The GDG believed that laryngospasm was an appreciable risk during this procedure and that sedation teams would need the skills and judgement to manage it.

The GDG discussed the use of a technique combining propofol with other sedation drugs such as midazolam, ketamine and opioids. The GDG understood that combinations of these drugs are being used to provide sedation for dental procedures in the UK. No RCTs were found testing the combinations of these drugs and therefore the efficacy could not be assessed.

The GDG thought that such a technique could cause unintentional deep and prolonged sedation. While it is true that the effects of opioid and midazolam can be reversed by naloxone and flumazenil, the reversal requires prompt administration and sedation may outlast the effects of sedation.

In contrast to drug combinations, the GDG agreed that unconsciousness and airway effects caused by propofol are more likely but are brief. Recovery of full consciousness after propofol is much more rapid and airway obstruction or apnoea can be managed with appropriate skills and equipment.

The GDG discussed the potential economic advantages of using propofol to either sedate or anaesthetise children for a wide variety of procedures. In comparison with almost any other method of sedation, propofol was the most effective apart from ketamine and sevoflurane. Provided intravenous access could be achieved propofol had the advantages of speedy onset and recovery. Propofol could enable a faster turnover of patients than many techniques. The disadvantage however is that propofol would need the same staff and facilities as an anaesthetic. This clearly has resource implications but the GDG agreed that if the demand of procedure was high the rapid nature of propofol sedation/anaesthesia could prove to be economically advantageous

The agreement by the GDG is that economic analysis should be conducted only for sedation techniques commonly available in the NHS. Economic analysis was conducted for six broad groups (dental procedure in children, dental procedure in adolescents, short painful procedures, painless imaging, oesophago-gastroscopy and colonoscopy). Propofol combined with fentanyl was felt to be a strategy commonly used in short painful procedures, and there is some evidence from the systematic review of opioids that propofol plus fentanyl is an effective and safe strategy. The GDG therefore agreed that the combination strategy should be compared to other relevant strategies in the economic analysis conducted for this population group. Details of the considerations of cost-effectiveness with respect to using propofol plus fentanyl in short painful procedures are given in section 6.12.1.2.

1 **6.10 Opioids**

Matrix of opioids comparators

Key:

Chloral hydrate = CH

Fentanyl = F

Isoflurane = I

Ketamine = K

Local anaesthesia = LA

 $\mathsf{Midazolam} = \mathsf{M}$

Nitrous oxide = N_20

Nitrous oxide and oxygen = N_20+02

Opioids = O

Propofol= P

 ${\sf Sevoflurane} = {\sf S}$

Triclofos sodium = TS

Opioids vs

	Reference	Tables	Evidence statements page
Placebo			
Nil			
Head to head			
Nil			
Combinations			
F + P vs. P + placebo	Cechvala 2008 ³⁸ Hollman 2008 ⁹³	Table 71	299
F + P vs. P	Disma 2005 ⁵⁶	Table 72	299
F + P vs. M + P	Disma 2005 ⁵⁶	Table 73	300
F + M vs. M + K	Lucas da Silva 2007 ¹⁵⁰ Kennedy 1998 ¹²⁸	Table 74	300
F + P vs. P + K	Tosun 2007 ²¹³	Table 75	301
Safety			
RCTs			
Assisted ventilation	Cechvala 2008 ³⁸ Hollman 2008 ⁹³	Table 76	302

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	Disma 2005 ⁵⁶ Kennedy 1998 ¹²⁸	Table 77	
ET intubation	Cechvala 2008 ³⁸ Hollman 2008 ⁹³	Table 76 Table 77	302
CPR / defibrillation	Lucas da Silva 2007 ¹⁵⁰	Table 76 Table 77	302
Desaturation	Cechvala 2008 ³⁸ Hollman 2008 ⁹³ Disma 2005 ⁵⁶ Lucas da Silva 2007 ¹⁵⁰ Kennedy 1998 ¹²⁸ Tosun 2007 ²¹³	Table 76 Table 77	302
Vomiting	Cechvala 2008 ³⁸ Hollman 2008 ⁹³ Kennedy 1998 ¹²⁸ Tosun 2007 ²¹³	Table 76 Table 77	302
Aspiration	Kennedy 1998 ¹²⁸	Table 76 Table 77	302
Observational studies	Pitetti 2003 ¹⁸² Sanborn 2005 ¹⁹⁶ Roback 2005 ¹⁸⁹ Mamula 2007 ¹⁵⁶ Sacchetti 2007 ¹⁹⁵	Table 76 Table 77	302
Route of administration			
Nil			
Dose			
Nil			

1

6.10.1 Clinical methodological introduction

2	CLINICAL QUESTIONS:
3 4 5	For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, is intravenous morphine, intravenous fentanyl or intranasal diamorphine (with or without: analgesia, another drug or psychological techniques):
6 7 8	- Effective for sedation (at minimal, moderate, and deep levels) in comparison with usual care, with analgesia alone, with another sedation drug, with psychological techniques or with general anaesthesia?
9	- Safe for sedation (at mild, moderate, and deep levels) in different settings?
10 11 12	The literature was searched for systematic reviews and RCTs for the clinical efficacy of opioids (intravenous morphine, intravenous fentanyl or intranasal diamorphine). The search was expanded to includenon RCT observational studies for the safety of opioids.
13	There were no systematic reviews identified for the use of opioids in paediatric sedation
14 15	Five RCTs comparing intravenous morphine, intravenous fentanyl, and intranasal diamorphine with other sedative drugs were assessed for efficacy and safety.
16 17	Five non RCT observational studies with total n=2439 were assessed for safety of opioids.
18 19	Crossover trials were treated separately from parallel armed trials unless there was sufficient data to allow their combination.
20 21 22 23	Meta-analyses for RCTs were performed where drug interventions and comparisons and outcomes were sufficiently homogenous and studies were combined regardless of dose, duration of intervention, procedure (within painful and non-painful groups), setting (e.g. dentistry, accidents and emergencies) and age.
24 6 .	10.2 Evidence profiles
25 6 .	10.2.1 RCT evidence profiles for efficacy and safety
26 27	Study characteristics and methodological quality of the study are provided in Appendix D. GRADE tables for quality assessment and summary of findings are provided below.

1 COMBINATION COMPARISONS

Table 71: Intravenous fentanyl + intravenous propofol vs. intravenous propofol + placebo; Cechvala, 2008; Hollman 2008 38,93

Date: 2009-09-23

Question: Should intravenous fentanyl plus intravenous propofol vs. intravenous propofol plus placebo be used in children and young people undergoing diagnostic and therapeutic procedures?

Settings: hospital outpatients

Bibliography: Cechvala 2008; Hollman2008 (Lumbar puncture)

			Ouglity sees	noment.			Summary of findings					
			Quality asses	ssment			No of patients Effect					
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous fentanyl plus intravenous propofol	intravenous propofol plus placebo	Relative (95% CI)	Absolute	Quality	Importance
Completi	on of proced	dure										
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	22/22 (100%)	22/22 (0%)	not estimable	-	MODERATE	
Anxiety r	ecorded by	the study inve	estigator using a	validated scale	(modified Ya	ale Preoperative A	Anxiety Scale (mYP	AS))				
	randomised trial		no serious inconsistency	no serious indirectness	serious ^{2,3}	none	0/22 (0%) ³	0/22 (0%)	-	not pooled	MODERATE	
procedur less)	e and when	patient ready	for discharge ar	nd 2) discharge	of patients fr	om sedation prog	edical Centre Scor gram after satisfact			onitoring criteria		
	randomised trial	4	no serious inconsistency	no serious indirectness	serious ⁴	none	22	22	-	MD -12.50 (- 22.4 to -2.6) ⁵	MODERATE	
Parents p	reference											
	trial	limitations ¹	,	indirectness		none	16/21 (76.2%)	5/21 (23.8%)	RR 3.20 (1.44 to 7.13) ⁷	524 more per 1000 (from 105 more to 1000 more) 0 more per 1,000	MODERATE	
Adverse	events: Assi	sted ventilation	on (flow inflating	anaesthesia b	ag)							
			no serious inconsistency	no serious indirectness	very serious	none	1/22 (4.5%)	1/22 (4.5%)	RR 1 (0.07 to 15) ⁸	0 fewer per 1000 (from 42 fewer to 630 more) 0 fewer per 1,000	LOW	
Adverse (events: End	otracheal intu	bation									

1	randomised trial	4		no serious indirectness	serious ^{2,9}	none	0/22 (0%)9	0/22 (0%)	not pooled	-	MODERATE	
Adverse	events: Oxy	gen desaturat	tion <90%									
1	randomised trial			no serious indirectness	very serious ^{2,10,11}	none	0/22 (0%)	1/22 (0%)	RR 0.33 (0.01 to 7.76) ¹⁰	0 fewer per 1,000	LOW	
Adverse	events: Von	niting										
	randomised trial	4		no serious indirectness	very serious ^{2,10,11}	none	0/22 (0%)	1/22 (4.5%)	RR 0.33 (0.01 to 7.76)	30 fewer per 1000 (from 45 fewer to 304 more) 0 fewer per 1,000	LOW	

Cechvala 2008 (Hollman 2008): double blind study - patients and outcome assessors blinded, ITT - yes, all patients followed and adequate allocation concealment; small study

² Cechvala 2008: small study

³ Cechvala 2008: stated that patients were not statistically different between groups in the level of anxiety as assessed by the mYPAS scale either before or after the administration of fentanyl and placebo

Cechvala 2008: imprecise, confidence intervals cross left confidence limit

⁵ Cechvala 2008: p=0.01

⁶ Cechvala 2008: imprecise, outside (right) confidence limits; small study ⁷ Cechvala 2008: p=0.004 ⁸ Cechvala 2008: p=1.00 ⁹ Cechvala 2008: stated there were no events of endotracheal intubation

¹⁰ Cechvala 2008: p=0.49

¹¹ Cechvala 2008: very wide confidence intervals crossing both confidence limits

Table 72: Intravenous fentanyl + intravenous propofol vs. intravenous propofol; Disma 2005 56

Date: 2009-09-23

Question: Should intravenous fentanyl plus intravenous propofol vs. intravenous propofol (with topical and local anaesthesia in both arms) be used in children and young people undergoing

diagnostic and therapeutic procedures?

Settings: gastroenterology

Bibliography: Disma 2005 (Endoscopy)

			Ouglity age				Summary of findings					
			Quality asse	essment			No of par	tients	Е	ffect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous fentanyl plus intravenous propofol plus topical anaesthesia plus local anaesthesia	intravenous propofol plus topical anaesthesia plus local anaesthesia	Relative (95% CI)	Absolute	Quality	Importance
Complet	Completion of procedure											
1	randomised trial		no serious inconsistency		no serious imprecision	none	82/82 (100%)	80/80 (100%)	not estimable	-	MODERATE	
Duration	Duration of procedure (Better indicated by less)											
1	randomised trial			no serious indirectness	serious ²	none	82	80	-	MD -0.60 (- 1.37 to 0.17) ³	LOW	
Recover more)	Recovery assessed using a validated scale: from completion of scan to achievement of Aldrete score of >=8 (measured with: Aldrete score; range of scores: 1-10; Better indicated by											
1	randomised trial			no serious indirectness	serious ⁴	none	82	80	-	MD 2.40 (- 0.09 to 4.89) ⁵	LOW	
Adverse	events: Ass	isted ventil	ation (bag mask	x)								
1	randomised trial			no serious indirectness	very serious ^{6,7}	none	2/82 (2.4%)	3/80 (3.8%)	RR 0.09 (0 to 1.58) ⁸	35 fewer per 1000 (from 38 fewer to 22 more) 0 fewer per 1,000	VERY LOW	
Adverse	events: Oxy	/gen desatu	ration <90%									
1	randomised trial			no serious indirectness	serious ⁶	none	2/82 (2.4%)	3/80 (3.8%)	RR 0.65 (0.11 to 3.79) ⁹	13 fewer per 1000 (from 34 fewer to 106 more) 0 fewer per 1,000	LOW	

Disma 2005: ITT appeared to be performed and there were no loss of follow up reported; however, allocation concealment and blinding of patients were not stated and blinding of outcome assessors was not clear; small study
² Disma 2005: imprecise, crosses left confidence limit

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³ Disma 2005: p=0.13
⁴ Disma 2005: imprecise, crosses right confidence limit
⁵ Disma 2005: p=0.06
⁶ Disma 2005: very imprecise, crosses both confidence limits and very wide confidence interval
⁷ Disma 2005: small study
⁸ Disma 2005: p=0.10
⁹ Disma 2005: p=0.63

Table 73: Intravenous fentanyl + intravenous propofol vs. intravenous midazolam + intravenous propofol; Disma 2005 56

Date: 2009-09-23

Question: Should intravenous fentanyl plus intravenous propofol vs. intravenous midazolam plus intravenous propofol (with topical anaesthesia in both arms) be used in children and young people

undergoing diagnostic and therapeutic procedures?

Settings: gastroenterology

Bibliography: Disma 2005 (Endoscopy)

			Quality acc	ncomont			Summary of findings					
			Quality asse	essinent			No of patients Effect					
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	a	intravenous fentanyl plus intravenous propofol plus topical anaesthesia	intravenous midazolam plus intravenous propofol plus topical anaesthesia	Relative (95% CI)	Absolute	Quality	Importance
Duration	Duration of procedure (Better indicated by less)											
1	randomised trial			no serious indirectness	no serious imprecision ²	none	82	78	ı	MD -0.40 (- 1.17 to 0.37) ³	MODERATE	
Recover more)	Recovery assessed using a validated scale: from completion of scan to achievement of Aldrete score of >=8 (measured with: Aldrete score; range of scores: 1-10; Better indicated by											icated by
1	randomised trial				no serious imprecision⁴	none	82	78	-	MD -0.10 (- 2.46 to 2.26) ⁵	MODERATE	
Complet	ion of proce	dure										
	randomised trial		no serious inconsistency		no serious imprecision	none	82/82 (100%)	78/78 (100%)	not estimable	-	MODERATE	
Adverse	Events: Ass	sisted ventil	ation (bag mask	<u>()</u>								
1	randomised trial				no serious imprecision	none	0/82 (0%) ⁸	0/78 (0%)	not pooled	-	MODERATE	
Adverse	Adverse Events: Oxygen desaturation <90%											
1	randomised trial		no serious inconsistency	no serious indirectness	very serious ⁶	none	2/82 (2.4%)	2/78 (2.6%)	RR 0.95 (0.14 to 6.59) ⁷	1 fewer per 1000 (from 22 fewer to 145 more) 0 fewer per 1,000	VERY LOW	

Disma 2005: ITT appeared to be performed and there were no loss of follow up reported; however, allocation concealment and blinding of patients were not stated and blinding of outcome assessors was not clear; small study

² Disma 2005: precise, within confidence limits ³ Disma 2005: p=0.31 ⁴ Disma 2005: precise within confidence limits ⁵ Disma 2005: p=0.93

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Disma 2005: very imprecise, crosses both confidence limits and too wide confidence intervals
 Disma 2005: p=0.96
 Disma 2005: The study reported that no patients needed bag-mask ventilation for assisted ventilation

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Table 74: Intravenous fentanyl + intravenous midazolam vs. intravenous midazolam + intravenous ketamine Lucas da Silva 2007 150 and Kennedy 1998 128

Date: 2009-09-23

Question: Should intravenous alfentanyl plus intravenous midazolam vs. intravenous midazolam plus intravenous ketamine be used in children and young people undergoing diagnostic and

therapeutic procedures?

Settings: hospital inpatients and accidents and emergencies

Bibliography: Lucas Da Silva 2007 (central venous catheter insertion) - hospital inpatients; Kennedy 1998 (orthopaedic: fracture or joint reduction) - accidents and emergencies

			Quality asse	accmont .				Summary of	findings				
			Quality asse	essinent			No of p	atients		Effect			
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous alfentanyl plus intravenous midazolam	intravenous midazolam plus intravenous ketamine	Relative (95% CI)	Absolute	Quality	Importance	
Completi	on of proced	dure (Lucas	Da Silva 2007; K	ennedy 1998) (follow-up mea	w-up mean 101-121 minutes¹)							
	randomised trial	, ,,	no serious inconsistency		no serious imprecision	none	155/158 (98.1%)	158/159 (99.4%)	RR 0.98 (0.95 to 1.01) ⁴	20 fewer per 1000 (from 50 fewer to 10 more) 0 fewer per 1,000	LOW		
Induction	Induction time (Lucas Da Silva 2007): time from initial sedative administration to onset of the procedure (follow-up mean 7.5 minutes; Better indicated by less)												
	randomised trial			no serious indirectness	serious ⁵	none	28	29	ı	MD 2 (-0.002 to 5.998) ⁶	LOW		
Recovery	time (Luca	s Da Silva 20	007): time from e	nd of procedur	e to awakenin	g (follow-up mear	n 20 minutes; Better	indicated by less)				-	
	randomised trial		no serious inconsistency	no serious indirectness	serious ⁵	none	28	29	-	MD -5 (-15 to 7.9) ⁷	LOW		
Total tim	e (Lucas Da	Silva 2007):	time from initial	sedative admir	nistration to sp	oontaneous eye o	pening (follow-up m	ean 101 minutes; Be	ter indicate	ed by less)		-	
	randomised trial			no serious indirectness	serious ⁵	none	28	29	-	MD 6.5 (-19 to 33) ⁸	LOW		
Adverse	events: Exte	ernal cardiac	massage/defibr	illation (Lucas	Da Silva 2007)					-	•		
1	randomised trial			no serious indirectness	serious ⁹	none	0/28 (0%)	0/29 (0%)	not pooled	0 fewer per 1000 (from 0 fewer to 0 fewer) 0 fewer per 1,000	LOW		
Adverse	events: Oxy	gen desatur	ation <90% (Luca	as Da Silva) (fo	llow-up 101 m	inutes)					•	•	
1	randomised trial		no serious inconsistency	no serious indirectness	serious ¹⁰	none	0/28 (0%)	2/29 (6.9%)	RR 0.21 (0.01 to 4.13)	55 fewer per 1000 (from 68 fewer to 216 more) 0 fewer per 1,000	LOW		

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			petween first mid oulation; Better in			aedic manipulatio	on (follow-up mean 1	3 minutes; measured	d with: time	in minutes betwe	en first r	nidazolam
1	randomised trial		no serious inconsistency	no serious indirectness	serious ¹¹	none	130	130	-	MD 0.30 (-2.5 to 3.1) ¹²	LOW	
	during proc 0-23.5; Bette			using a valida	ted scale (Ken	nedy 1998) (meas	sured with: Observati	ional Scale of Behav	ioural Distre	ess-Revised (OSB	D-R); raı	nge of
1	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ¹³	none	130	130	=	MD 1.62 (1.2 to 2.04) ¹⁴	LOW	
Anxiety (during proce	dure asses	sed by parent us	ing a validated	scale (Kenned	ly 1998) (measure	ed with: Visual Analo	gue Scale, range of	scores: 0-10	; Better indicated	by less)	
	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ¹⁵	none	130	130	=	MD 1.01 (0.22 to 1.8) ¹⁶	LOW	
ain dur	ing procedu	re assessed	by parent using	a validated sca	ale (Kennedy 1	998) (measured v	vith: Visual Analogue	Scale; range of sco	res: 0-10; B	etter indicated by	less)	
1	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ¹⁵	none	130	130	-	MD 1.34 (0.53 to 2.15) ¹⁷	LOW	
Total tim	e (Kennedy	1998): from	administration o	f intervention t	o when patien	t has been transfe	erred to recovery are	a (follow-up mean 12	27.6 minutes	18; Better indicate	ed by les	s)
1	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ¹⁹	none	130	130	-	MD -13.90 (- 25.46 to -2.34) ²⁰	LOW	
Adverse	events: Asp	iration (Ken	nedy 1998) (follo	w-up mean 121	l minutes; thro	ughout procedur	e; number of patient	s)	<u> </u>		-	
1	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ²¹	none	0/130 (0%) ²²	0/130 (0%)	not pooled	-	LOW	
Adverse	events: Ass	isted ventila	tion - bag-valve	mask (Kennedy	y 1998) (follow	-up mean 121 mir	nutes; throughout pro	ocedure)				
Ī	randomised trial	serious ³	no serious inconsistency	no serious indirectness	very serious ²³	none	0/130 (0%)	2/130 (1.5%)	RR 0.20 (0.01 to 4.13) ²⁴	12 fewer per 1000 (from 15 fewer to 47 more) 0 fewer per 1,000	VERY LOW	
Adverse	events: Von	niting during	procedure (Ken	nedy 1998) (fol	llow-up mean 1	121 minutes; thro	ughout procedure)					
1	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ^{11,25}	none	0/130 (0%)	1/130 (0.8%)	RR 0.33 (0.01 to 8.11) ²⁶	5 fewer per 1000 (from 8 fewer to 57 more) 0 fewer per 1,000	LOW	
Adverse	events: Von	niting during	recovery (Kenn	edy 1998)								
1	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ¹¹	none	0/130 (0%)	1/130 (0.8%)	RR 0.27 (0.08 to 0.96) ²⁷	6 fewer per 1000 (from 0 fewer to 7 fewer) 0 fewer per 1,000	LOW	
Adverse	events: Oxy	gen desatui	ration <90% Kenr	nedy (follow-up	mean 121 mir	nutes; throughout	procedure)					
1	randomised trial	serious ³	no serious inconsistency	no serious indirectness	serious ^{11,25}	none	31/130 (23.8%)	8/130 (6.2%)	RR 3.88 (1.85 to 8.11) ²⁸	179 more per 1000 (from 53 more to 441 more) 0 more per 1,000	LOW	

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- ¹ Lucas Da Silva 2007: mean follow up 101 minutes; Kennedy 1998: mean follow up 121 minutes
- ² Lucas Da Silva 2007: double blinding was deemed impractical because of different dosing algorithms of the drugs used and because medications used present clinically distinguishable effects; small study
- ³ Kennedy 1998; quasi randomised; subjects stratified according to initial parental choice to remain in the room or not during reduction and were then randomly assigned in blocks of 20 within strata to receive fentanyl or ketamine; not fully blinded: blinding of patients and parents not stated, two trained observers blinded to study purpose and design reviewed the videotape of each study but unable to blind sedators; ITT performed and all patients followed
- ⁴ Kennedy 1998 and Lucas Da Silva: p=0.31
- ⁵ Lucas Da Silva 2007: median results were reported for the outcomes of induction time, total time and recovery time thus not possible to combine with Kennedy 1998; small sample size
- ⁶ Lucas Da Silva 2007: p=0.03; stated median results with p-values on the study
- ⁷ Lucas Da Silva 2007: p=0.40; stated median results with p-values on the study
- ⁸ Lucas Da Silva 2007: p=0.67; stated median results with p-values on the study
- 9 Lucas Da Silva 2007: study stated there was an increase in cardiac arrest but 'no intervention was required' and 'no cardiac abnormalities were detected'
- ¹⁰ Lucas Da Silva 2007: wide confidence intervals crossing both precision limits
- ¹¹ Kennedy 1998: small sample size
- ¹² Kennedy 1998: p=0.83
- ¹³ Kennedy 1998: precise but OBSD-R may be biased by subjectivity of observer
- ¹⁴ Kennedy 1998: p < 0.00001
- 15 Kennedy 1998: crosses right precision limit
- ¹⁶ Kennedy 1998: p=0.01
- ¹⁷ Kennedy 1998: p=0.001
- ¹⁸ Kennedy 1998: control group had the longest total time 127.6 minutes (SD56.2) compared to 113.7 minutes (SD36.9) in the intervention group
- ¹⁹ Kennedy 1998: crosses left precision limit
- ²⁰ Kennedy 1998: p=0.02
- ²¹ Kennedy 1998: small sample
- ²² Kennedy 1998: study stated there were no events of aspiration
- ²³ Kennedy 1998: crosses both precision limits; too wide confidence intervals
- ²⁴ Kennedy 1998: p=0.30
- ²⁵ Kennedy 1998: precise; wide confidence intervals; no possible to combine with Lucas Da Silva 2007 due to significant heterogeneity (I2=72%; p=0.06) between studies for this outcome
- ²⁶ Kennedy 1998: p=0.50
- ²⁷ Kennedy 1998: p=0.04
- ²⁸ Kennedy 1998: p=0.0003

Table 75: Intravenous fentanyl + intravenous propofol vs. intravenous propofol + intravenous ketamine; Tosun 2007 213

Date: 2009-09-23

Question: Should intravenous fentanyl plus intravenous propofol vs. intravenous propofol plus intravenous ketamine (with topical anaesthesia in both arms) be used in children and young people

undergoing diagnostic and therapeutic procedures?

Settings: gastroenterology Bibliography: Tosun 2007 (upper and lower endoscopy)

			Quality ass	accment			Summary of findings					
			Quality asse	essment			No of p	patients	E	Effect		
No of studies	Design	Limitations	Inconsistency	Indirectness	Imprecision	Other considerations	intravenous fentanyl plus intravenous propofol plus topical anaesthesia	intravenous propofol plus intravenous ketamine plus topical anaesthesia	Relative (95% CI)	Absolute	Quality	Importance
Complet	ion of proce	dure (follow	-up mean 116 m	ninutes)					-			•
	randomised trial		no serious inconsistency	no serious indirectness	no serious imprecision	none	44/44 (100%)	46/46 (100%)	RR 1 (0 to 0)		MODERATE	
Duration	Duration of procedure (follow-up mean 116 minutes; Better indicated by less)											
	randomised trial			no serious indirectness	no serious imprecision ²	none	44	46	-	MD -0.20 (- 1.27 to 0.87)	MODERATE	
Pain: nu	mber of pati	ients who ne	eded additional	l propofol duri	ng induction a	as evidenced by	disconfort/moving du	ring procedure (follow	v-up 0-1 mi	nute after indu	ction)	
	randomised trial	serious ¹	no serious inconsistency	no serious indirectness	serious ³	none	22/44 (50%)	8/46 (17.4%)	RR 2.88 (1.43 to 5.76) ⁴	0 more per 1,000	LOW	
Pain: nu	mber of pati	ients who ne	eded additional	propofol as e	videnced by c	liscomfort/movir	g during procedure				•	
	randomised trial			no serious indirectness	serious ³	none	41/44 (93.2%)	32/46 (69.6%)	RR 1.34 (1.09 to 1.65) ⁵	0 more per 1,000	LOW	
Recover	y: time from	completion	of procedure to	recovery/disc	harge criteria	being met (follo	w-up mean 4.5 minute	es; Better indicated by	y less)			-
	randomised trial		no serious inconsistency	no serious indirectness	serious ²	none	44	46	-	MD 0.80 (- 11.16 to 12.76) ⁶	LOW	
Adverse	events: Oxy	ygen desatu	ration <90% (fol	low-up mean 1	16 minutes; t	hroughout proce	dure)					
	randomised trial			no serious indirectness	serious ⁷	none	4/44 (9.1%)	3/46 (6.5%)	RR 1.39 (0.33 to 5.88) ⁸	25 more per 1000 (from 44 fewer to 317 more)	LOW	

1	6.10.2.2 Non RCT evidence profiles for safety
2 3 4 5	Five non RCT observational studies in 2,439 patients assessed the safety of opioids ^{156,182,189,195,196} . There were four prospective studies, and one retrospective study conducted for the following procedues: imaging procedures (1), accidents and emergencies procedures (3) as well for GI procedures (1).
6	The non RCT study characteristics for opioids are presented in Table 76.
7	The non RCT adverse event table for opioids is presented in Table 77.
8	
9	

Table 76: Opioids Non RCT Study Characteristics Safety Review

Study	Setting	ASA	Sedation type	Gender, % male	Drug (doses)	Fasting
Pitetti 2003 ¹⁸² , USA	Accidents and emergencies Prospective descriptive study	81% were Class I; 17% were class II; 1.3% were class III and 0.1% were class IV.	Procedural sedation	65.1% boys in total sample (791)	IV fentanyl citrate + midazolam & IV morphine sulphate + midazolam and IV midazolam Mean fentanyl dose: 2.7 mcg /kg Midazolam 0.1 mg/kg; Morphine not stated	Mean fasting 5.0 + 2.8 hours before sedation.
Sanborn 2005 ¹⁹⁶ , USA	lmaging Retrospective chart review	44%of total subjects were ASA I; 51% ASA II; 4% ASA III; 0.1% ASA IV; 0.1% ASA V.	IV fentanyl + midazolam and IV fentanyl	56% of total were male	Doses not stated	Not stated
Roback 2005 ¹⁸⁹ , USA	Accidents and emergencies Prospective observational database	Not stated	Procedural sedation	60.4% of total were male	Midazolam + fentanyl vs. midazolam alone	Not stated
Mamula 2007 ¹⁵⁶ , USA	Operating Room	ASA I-III	Intravenous or general anaesthesia	55% (674/1226)	IV midazolam (2 mg/2mL) & fentanyl (100 mcg/2mL) during 1 minute. Midazolam 0.05 to 0.1 mg/kg max 2 mg; fentanyl 1 mcg/kg max 75 mcg Oral midazolam fo anxious patients; IV diphenhydramine as additional sedative	3 hours
Sacchetti 2007 ¹⁹⁵ , USA	Accidents and emergencies Prospective observational database	94.1%of total cohort Class I, 5.3% class II and 0.6% class III.	Procedural sedation	Not stated	Fentanyl & Morphine	Not stated

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1 Table 77: Opioids Safety: Non RCTs

			ire Age	Total N			A	DVERSE EVEN	NTS, rate: % ((n)			GRADE PROFILE
Study type,	Drug /	Procedure						Cardiac arrest requiring either/or			oxygen		
reference, country	Comparison	Trocadic			Aspirat ion	oral- pharyng eal airway	endotrac heal intubatio n	assisted ventilation	external cardiac massage	defibril lation	vomiting	desaturat ion <90%	EVIDENCE QUALITY
Pitetti 2003 ¹⁸² , USA	IV fentanyl citrate + midazolam hydrochloride Vs. midazolam alone	A & E	0-21 years	686 vs 65 Complications reported as total adverse events: 23.5% vs. 1.5%									VERY LOW
	IV morphine sulphate + midazolam Vs. IV midazolam	A & E	0-21 years	48 vs. 65 Complications reported as total adverse events: 16.7% vs. 1.5%									VERY LOW
Sanborn 2005 ¹⁹⁶ , USA	Fentanyl	MR and Ct imaging	Mean age of total sample was 4.8 years + 4.6	42/16467	0	0	0	0	0	0		0	VERY LOW
Roback 2005 ¹⁸⁹ , USA	Midazolam +Fentanyl Vs. Midazolam	A & E	19 days to 32 years; median 6.7 years	336 vs. 260				All patients experiencin g apnea or laryngospas m were managed with administrati			Midazola m/ fentanyl 1.8% (6/336) Midazola m 0.8% (2/260)	Respirator y adverse events reported and included oxygen saturation	VERY LOW

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							on of oxygen, breathing cues, airway positioning or bag-mask ventilation. Numbers of each intervention were not provided.			<90%, apnea or laryngosp asm Midazola m/ fentanyl 19.3% (65/336) Midazola m 5.8% (15/260)	
Mamula, 2007 ¹⁵⁶ , USA	iv midazolam/f enta/ only when needed: oral Mid for anxious children & diphenhydra mine to reach desired effect		0.1-34 y 4%(55/1	1226	0% (pulmon ary aspirati on)	0%	0.16% (2/1226) (bag/mask ventilation)	0% (0/1226) (cardiac arrest)	5.2% (64/1226) (during recovery)		VERY LOW
Sacchetti 2007 ¹⁹⁵ , USA	Fentanyl	A & E	0-20 years	*episode of apnea with fentanyl and etomidate which required reversal was only adverse event reported.							VERY LOW

1	6.10.3 Evidence statements
2	6.10.3.1 RCT efficacy and safety
3	COMBINATIONS COMPARISONS
4	IV fentanyl + IV propofol vs. IV propofol + placebo
5	Cechvala 2008; Hollman 2008 ^{38,93}
6	All patients completed the procedure [moderate quality evidence]
7 8	Compared to intravenous propofol + placebo, the intravenous fentanyl + intravenous propofol group was significantly:
9	Preferred among parents [moderate quality evidence]
10 11	 Faster in recovery time (Connecticut Children's Medical Centre Scoring System [moderate quality evidence]
12	There were no events of:
13	Endotracheal intubation [moderate quality evidence]
14	There was no significant difference in:
15	 Anxiety (mYPAS) [moderate quality evidence]
16	Assisted ventilation (flow inflating anaesthesia bag) [low quality evidence]
17	 Oxygen desaturation < 90% [low quality evidence]
18	Vomiting [low quality evidence]
19	
20	IV fentanyl + IV propofol vs. IV propofol
21	Disma 2005 ⁵⁶
22	All patients completed the procedure [moderate quality evidence]
23	There was no significant difference in:
24	Slower recovery time (Aldrete score) [low quality evidence]
25	Duration of procedure [low quality evidence]
26	 Oxygen desaturation <90% [low quality evidence]
27	Assisted ventilation (bag-mask ventilation) [very low quality evidence]

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1	
2	IV fentanyl + IV propofol vs. IV midazolam + IV propofol
3	Disma 2005 ⁵⁶
4	All patients completed the procedure [moderate quality evidence]
5	There was no significant difference in:
6	Duration of procedure [moderate quality evidence]
7	Recovery time [moderate quality evidence]
8	 Oxygen desaturation <90% [very low quality evidence]
9	There were no events of:
10	Assisted ventilation (bag mask) [moderate quality evidence]
11	
12	IV fentanyl + IV midazolam vs. IV midazolam + IV ketamine
13 14 15 16 17 18 19 20	For the outcome of oxygen desaturation (<90%), we found evidence of highly significant heterogeneity (I ² =72%; p=0.06) between two RCTs ¹⁵⁰ , ¹²⁸ . Possible sources of heterogeneity could be attributed to the differences between the studies in procedure performed (catheter insertion versus orthopaedic fracture or joint reduction) and length of procedure (orthopaedic fracture or joint reduction takes longer), setting (inpatients versus accidents and emergencies) and varying dose of combination agents. We therefore felt it was not appropriate to pool the RCTs together in a meta-analysis and the studies are presented separately for this outcome.
21	Lucas da Silva 2007 ¹⁵⁰ , Kennedy 1998 ¹²⁸
22	There was no significant difference in:
23	Completion of the procedure [low quality evidence]
24	Lucas da Silva 2007 ¹⁵⁰
25	There was no significant difference in:
26	Induction time [low quality evidence]
27	Recovery time (minutes) [low quality evidence]
28	Total sedation time (minutes) [low quality evidence]
29	 Oxygen desaturation <90% [low quality evidence]
30	There were no events of:

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ı	External cardiac massage or defibriliation [low quality evidence]
2	Kennedy 1998 ¹²⁸
3 4	Compared to the intravenous midazolam + intravenous ketamine group, the intravenous midazolam + intravenous fentanyl group had significantly:
5	• Higher distress scores during procedure (OSBD-R) [low quality evidence]
6	Higher anxiety scores (VAS) [low quality evidence]
7	Higher pain scores during procedure (VAS) [low quality evidence]
8	Shorter total time [low quality evidence]
9	 More oxygen desaturation <90% [low quality evidence]
10	Less vomiting during recovery [low quality evidence]
11	There were no events of:
12	Aspiration [low quality evidence]
13	There was no significant difference in:
14	Completion of procedure [Low quality evidence]
15	• Induction time [low quality evidence]
16	Vomiting during procedure [low quality evidence]
17	 Assisted ventilation (valve-mask) [very low quality evidence]
18	
19	IV fentanyl + IV propofol vs. IV propofol + IV ketamine
20	Tosun 2007 ²¹³
21	All patients completed the procedure [moderate quality evidence]
22 23 24	Compared with intravenous propofol + intravenous ketamine + topical anaesthesia, children who received intravenous fentanyl + intravenous propofol + topical anaesthesia had significantly:
25 26	 More pain (number of patients) in the first minute after induction [Low quality evidence]
27	More pain (number of patients) during procedure [Low quality evidence]
28	There was no significant difference in:

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- Length/duration of procedure [moderate quality evidence]
- Recovery time [low quality evidence]
- Oxygen desaturation <90% [low quality evidence]
 - Vomiting [low quality evidence]

6.10.3.2 Non RCT safety (adverse events)

For the characteristics of studies and outcome data refer to Table 76 and Table 77.

- One study reported a 1.6% rate of assisted ventilation ¹⁵⁶. One study reported no events ¹⁹⁶. No other reports of respiratory intervention were elicited from the studies.
- There were no cardiac events reported in 5 studies.
- Vomiting rates were reported in two studies of midazolam + fentanyl: 1.8% ¹⁸⁹ and 5.2% ¹⁵⁶.
 - Adverse respiratory events including oxygen saturation <90%, apnea and laryngospasm were reported with the use of midazolam + fentanyl at 19.3% vs midazolam alone at 5.8% ¹⁸⁹. One study reported no events ¹⁹⁶. No other reports of desaturation were elicited from the studies.

18 6.10.4 GDG discussion of the evidence for opioids

The GDG found no studies that opioids (morphine, fentanyl and diamorphine) were effective for any diagnostic or therapeutic procedure when used alone to cause sedation rather than simply analgesia. In the studies found, opioids were always combined with other sedatives and the GDG agreed that they had been used for their analgesic properties within a sedation technique. The sedative potential of these selected opioids could not be determined from the evidence.

There were no studies on diamorphine.

There was one RCT¹⁸² of morphine in which it was combined with midazolam in the Emergency department setting. The efficacy of this combination could not be determined from the data because the evidence level was very low. The GDG agreed that morphine was a drug that had an analgesic action that was much longer than most painful procedures and for this reason shorter acting opioids such as fentanyl were likely to be more suitable.

All other evidence on opioids was provided from studies of combinations of fentanyl with either midazolam or propofol. Most studies were in the emergency department setting but one was in a hospital in children undergoing lumbar puncture. The GDG agreed that the principles of sedation for painful procedures in the ED were applicable to sedation for similar painful procedures in other settings.

The choice of opioid to be used in combination with midazolam was debated. In the early discussions of the GDG it was agreed that evidence for pethidine would not be sought because it had a longer action than fentanyl and because it was not widely used.

The combination of fentanyl with midazolam was used with the intention of maintaining moderate sedation but the GDG appreciated that it was sometimes difficult to titrate the drugs to provide sedation and analgesia to overcome the pain of the procedure without causing deep sedation or appreciable suppression of airway reflexes or breathing. The hazard of opioid induced respiratory depression occurring after the procedure had been completed was noted by the GDG. In one study¹⁸⁹ of children undergoing procedure in the ED setting, desaturation, apnoea or laryngospasm was reported as occurring in up to 19% of children. In comparison, ketamine has a safer record and has a similar induction and recovery time. The GDG agreed that even with careful titration of fentanyl and midazolam deep sedation and airway obstruction or apnoea are possible and that this combination should only be used by a trained sedation team. Airway management skills and equipment are essential for this drug combination.

Fentanyl combined with propofol was considered by the GDG to be a useful deep sedation or anaesthesia technique. Two RCTs³⁸, ⁵⁶ were considered. One showed that the addition of fentanyl to propofol reduced recovery time and the other found that propofol doses could be reduced. Fentanyl was associated with fewer adverse events.

The general principle agreed by the GDG is that only sedation techniques commonly available in the NHS should be included in the economic analysis. Economic analysis was conducted for six broad groups (dental procedure in children, dental procedure in adolescents, short painful procedures, painless imaging, oesophago-gastroscopy and colonoscopy). A combination of fentanyl and midazolam was felt to be commonly used in colonoscopy, whereas fentanyl plus propofol was felt to be commonly used in short painful procedures. There is some evidence that these combination strategies are effective and safe. The GDG therefore agreed that they should be included in the economic analysis. Details of the considerations of cost-effectiveness with respect to using fentanyl plus propofol in short painful procedures, and using fentanyl plus midazolam in colonoscopy are given in section 6.12.1.2 and 6.12.3.2 respectively.

6.11 SEDATION SPARING

6.11.1 Clinical methodological introduction

CLINICAL QUESTION:

For children and young people under the age of 19 undergoing diagnostic or therapeutic procedures, does a combination of psychological techniques and sedation drugs lead to sedation sparing?

The literature was searched for systematic reviews and RCTs for sedation sparing i.e. how much of the sedation drug is used in each arm alone or in combination with pharmacological intervention.

There were no systematic reviews, RCTs or observational studies that reported relevant outcome measures for analyses of our efficacy and safety outcomes.

6.11.2 Evidence statements

There were no RCTs or observational studies relevant for analyses of our efficacy and safety outcomes.

The GDG felt that sedation sparing techniques are not among the sedation techniques commonly used in the NHS, and decided that an economic analysis should not be done for these techniques.

6.12 Clinical settings: introduction

There are different types of diagnostic and therapeutic procedures. For example, some procedures are painful yet others are painless but require prolonged immobility. The efficacy and safety of sedation depends therefore not only on the drug or technique but also on the procedure itself. After reviewing the drugs, the GDG sought to group the evidence according to the type of procedure to enable the development of guidance on effective and safe sedation for specific procedures. There are many types of procedures and the GDG accepted that guidance on each and every procedure was not practicable. For the purposes of this guidance, the GDG used the classifications of

- painful procedures,
- painless imaging,
- endoscopy,
- dental procedures
- which they believe cover the majority (more than 90%) of common procedures.

Guidance for uncommon procedures can be obtained by applying relevant principles from the guidance below. Before considering sedation for a procedure the practitioner will need to understand what the procedure entails, what is expected of the patient, and what the sedation technique needs to achieve (see chapter 4).

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6.12.1 Painful Procedures

Many children undergo brief painful procedures following injury (such as suture of lacerations and orthopaedic manipulations in emergency departments). In a recent review¹⁸ the prompt administration of analgesia has been promoted not only because it is important and compassionate, but because it can reduce anxiety and increase cooperation of the child or young person to enable the procedure to be carried out with sedation rather than anaesthesia. Recently the term "procedural sedation and analgesia" has been used because it emphasizes that the analgesia component of sedation is crucial.¹³⁰ Many painful procedures can be carried out under local anaesthesia, provided the child or young person is cooperative. If the patient is unable to cooperate local anaesthesia is still important because the dose of sedative drug can be minimized if the patient has no pain. The following recommendations in this section are applicable to any painful procedure not only in the emergency setting but elsewhere such as a hospital ward.

There are several potentially useful sedation techniques for painful procedures. The decision to undertake a particular technique should be influenced by factors such as the type and duration of a painful procedure, the age and developmental stage of the child, and the urgency of a painful procedure. In particular, clinicians should consider the target depth of sedation required, and the relative requirement for analgesia, sedation, immobility and amnesia. Prolonged or complex should be carried out under general anaesthesia.

The sedation techniques recommended for painful procedures are considered in relation to the three target levels of sedation although it should be appreciated that there is variation in the sedation level achieved. Ketamine induces sedation which has different characteristics to any other sedation drug. Ketamine causes 'dissociative sedation' which is a trance-like cataleptic state, with profound analgesia, sedation, amnesia, and immobility. Ketamine tends to preserve airway reflexes, spontaneous respiration, and cardiovascular stability. Nevertheless occasionally ketamine can cause airway complications including laryngospasm. Dissociative sedation has been included in the category of deep sedation because the training and facilities needed for safe practice are similar for both (see sections 4.4 Personnel and training and 4.5 Clinical environment and monitoring).

Wound suture and foreign body removal are common examples of painful procedure usually carried out under minimal sedation. Moderate sedation is required for brief emergency orthopaedic procedures such as transferring a child with a fractured limb or placing the limb into a splint and reduction of a dislocated joint. Titration of the drugs used to achieve moderate sedation is important to avoid excessive respiratory depression. Examples of procedures usually carried out under deep or dissociative sedation are suture of lacerations to the face and nail bed in young children, and orthopaedic manipulations.

In an urgent or emergency situation the time of the last food and drink intake in children and young people is often uncertain. Moreover, trauma may delay gastric emptying. The problem of whether to use sedation (or anaesthesia) within a few hours after admission to hospital in a patient who may not be fasted is common. In most situations the procedure can be delayed although there will be practical problems of arranging for the procedure later. The risk of pulmonary aspiration during deep sedation and anaesthesia will need to be balanced with the risk of delaying the procedure. In many

situations it may be reasonable to use a sedation technique with a wide margin of safety in a patient who is not fasted (see section 4.2 Fasting).

Some of the sedation drugs are anaesthetic agents such as ketamine and propofol, and their use by 'non-anaesthetists' has been controversial. This has arisen because anaesthesia services are not always available. Skills necessary for safe sedation can be achieved by practitioners who are not fully trained anaesthetists (see section 4.4 Personnel and training).

6.12.1.1 Summary of evidence in painful procedures

The GDG extracted essential evidence from each drug review and incorporated this evidence into Table 78 and Table 79 below. The tabular presentation was developed as a way to summarise disparate data, ranging across various drug types, drug combinations, specialty areas and procedural techniques. The tables have thus been organised by setting and include the following: painless procedures (imaging), dentistry, painful procedures and GI procedures. The primary efficacy outcome was completion of procedure.

On the basis of the evidence, the GDG considered the efficacy and safety (benefits and harms) of each drug and drug combination reviewed. They indicated their decision for each drug within the tables by agreeing 'green' (yes) or 'red' (no) for efficacy and for safety.

Table 78: Drugs alone in painful procedures

Key: 'green' (yes) or 'red' (no), in the efficacy and safety columns, indicate the GDG judgment whether the drug was effective and safe.

DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
70% N20	Inhaled	70%	0-18 years	Emergency procedures	N/A	Vomiting 4.7% (26/548); 0.18% (1/548) desaturation	Non RCT	Babl 2008 ²¹
50% N20	Inhaled	50%	0-18 years	Emergency procedures	N/A	Vomiting 3.9% (4/101); 0% (1/548) desaturation	Non RCT	Babl 2008 ²¹
50% N20	Inhaled	50%	<19 years	Mixed procedures including emergency, GI, radiology, lumbar puncture, etc.	N/A	'Minor events' 5% (375/7511) and 0.3% 'major' events (25/7511)	Non RCT	Gall 2001 ⁷³
50% N20	Inhaled	50%	2 studies in systematic review with mean ages of 3.7 years and 10 years.	Laceration repair and fracture reduction	N/A	Yes (green) No reported vomiting or desaturation.	Non RCT	Faddy 2005 ⁶¹
K IV vs IM	IV vs. IM		4mo-18y	Orthopaedic reduction	Yes (Green) Favours IV for distress score but longer total time for IM	Desat: IV 9/109 vs IM 4/99; vomiting: IV 13/109 vs IM 26/99; ventilation: IV 2/109	Low quality	Roback 2006 ¹⁹⁰
Ketamine	IM		0-15y	Emergency procedures miscellaneous	N/A	0.4% (5/1022) bag mask ventilation; desaturation 0.9% (9/1022); vomiting 6.7% (68/1022)	Non RCT	Green 1998 ^{81,82} , Green 1998 ⁸²
Ketamine	IM		Not stated	Suturing, minor surgery in A & E	N/A	17% vomiting in recovery or at home (85/501); 0.5% desaturation; 15% mild recovery agitation (71/501); 3% moderate agitation (16/501); 0.8% pronounced	Non RCT	McGlone 2004 ¹⁶²

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DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
Ketamine	Not stated		0-20 years	Minor trauma including laceration repair, fracture care, lumbar puncture, radiology etc.	N/A	agitation (4/501) No reported adverse events	Non RCT	Sachetti 2007 ¹⁹⁵
IV F/IV P vs IV P/Placebo	IV	Fenta: 1 mcg/kg PRO: 1-2 mg/kg/min infusion	2-17 y	Lumbar puncture	Yes (Green)	Yes (Green)	Low quality	Cechvala 2008 ³⁸
IV F IV M vs IV M/IV K	IV	Mid: 0.15 mg/kg (max:0.5 mg/kg) Fenta: 1 mcg/kg (max 100 mg) Keta: 0.5 mg/kg	3 mo-14 y	Intravenous line placement	Yes (Green)	Yes (Green)	Low quality	Lucas Da Silva 2007 ¹⁵⁰
P/O analgesics (either Mo or F)	IV	PRO: initial dose 1 mg/kg (max 40 mg); supplemental doses 0.5 mg/kg (max 20 mg) Morphine: 0.1 mg/kg (max 5 mg) Fentanyl: 1-2 mcg/kg (max 50 mg)	Overall: 1-18y Median : 8 y	Fractures, dislocations, examination of ocular burn	Not reported	Yes (Green)	Very low quality	Bassett 2003 ²⁵
Oral M vs. Placebo	Oral	0.5 mg/kg	oral M 7.7 y (SD4.4) placebo 7.9 y (SD4.4)	IV insertion	Yes, all completed procedure (Green) and insufficient data (NSD) for procedure duration and	Not reported	Moderate	Liacouras 1998 ¹³⁹

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DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
					recovery (Red)			
IN M vs. Placebo	IN	0.4 mg/kg	median 2.5 y (range: 0.75- 4.9)	Suture/laceration repair	Yes, favours M in patient satisfaction (Green)	Yes (Green) vomiting: no events	Very low - low	Theroux 1993 ²¹⁰
IN M vs. Placebo	IN	0.2 mg/kg	mean age: 5 y (range: 0.8-18 y)	Needle insertion	Insufficient data for pain scores assessed by either patient (NSD) or parents (favours M) (Red) Favours M for parents satisfaction (Green) and NSD for patient satisfaction (Green)	Not reported	Very low - low	Ljungman 2000 ¹⁴⁸
Oral M vs. Placebo	Oral	0.5 mg/kg	mean age 4.1 y (range: 2-6)	Suture/laceration repair	Yes, NSD in completion of procedure (Green)	Yes (Green) no events of aspiration, cardio-repiratory or or cardiac massage	Moderate	Luhman 2001 ¹⁵¹
Oral M vs. Placebo	Oral	0.3 mg/kg	mean age 4.8 y (SD3) (range 0.8-10)	Suture/laceration repair	Yes, favours M for distress score (Green) and NSD for anxiety (Green)	Not reported	Moderate	Fatovich 1995 ⁶³
Rectal M/Non- pharma* vs. N2O/Non- pharma	R	0.35-0.5 mg/kg	mean age: RM 8:7 y (SD4:9), N2O 8:6 (SD3:8)	spasticity injections -cerebral palsy-	No, it is reported to favour N2O group for anxiety (p=0.010) (Red)	Yes (Green) NSD in vomiting	Moderate	Zier 2008 ²³⁷

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DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
					Yes, for parents satisfaction, it is reported NSD between groups (p=0.10) (Green) No, insufficient data for total time, it is reported NSD between groups (Red)			
IV M/IV Mo vs. IVPRO/IV Mo / LA	IV	0.1 mg/kg	(range 2-18 y) mean age: IVM 8.6 y (SD4.2) PRO 9 y (SD3.8)	Reduction of fractures	Yes, all completed, and NSD for induction and procedure time and for pain (Green) No, favours PRO group for recovery and total time (Red)	Yes (Green) no events for aspiration, external cardiac massage or assisted ventilation No (Red) Selective reporting for O2 desat	Very low- low	Havel 1999 ⁹⁰
Oral M vs. IN M	Oral vs. IN	0.5 mg/kg vs. 0.25 mg/kg	(range: 2 -10 y) mean age: Oral M 4.4 y (SD2.5) IN M 3.5 y (SD2)	Suture/laceration repair	Yes, all completed the procedure and NSD for total time (Green)	Yes (Green) NSD for total time	Very low - moderate	Connors 1994 ⁴²
Oral M vs. IN M	Oral vs. IN	1 mg/kg vs. 0.4 mg/kg	(range 1 to 5 y)	Suture/laceration repair	Yes, favours oral M for distress score and NSD for total time	No (red) selective reporting for vomiting	Very low	Everitt 2002 ⁶⁰

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DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
					(Green)			
Rectal M: 2mg vs. 1mg	R 2mg/kg vs. 1 mg/kg	2mg/kg vs. 1 mg/kg	(range 0.5-4) higher dose: 2.5(SD1), lower dose: 2.13(SD0.9).	Suture/laceration repair	Yes, NSD in satisfaction, recovery time and total time (Green)	Yes (Green) Yes, no cardio- respiratory events No (red) selective reporting in vomiting	Low - moderate	Kanegaye 2003 ¹²³

Table 79: Drugs combination in painful procedures

Key: 'green' (yes) or 'red' (no), in the efficacy and safety columns, indicate the GDG judgment whether the drugs combinations was effective and safe.

DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
M/K vs M/F	IV		5-15	All completed* go back to paper and check	Yes (Green) Favours M/K pain, distress, anxiety)	More desat in M/F; more vomiting in M/K; assisted ventilation in 2 M/K pts	Low quality	Kennedy 1998 ¹²⁸
IV P/F vs IV M/K	IV		3-18	All completed	Yes (Green) Favours M/K (pain, distress, anxiety) but longer recovery time	KM: 4/54 desat; P/F 18/59 desat; vomiting KM 2/54; recovery/agitation 3/54	Low quality	Godambe 2003 ⁷⁷
M/K vs IV M/Axillary block	IV		=>8y	All completed	Yes, NSD for pain and distress scores	N/A	Low quality	Kriwanek 2006 ¹³¹
M/K vs haematoma block/entonox	IV		5-17y	All completed	Yes (Green) NSD for pain distress but longer recovery time for K/M	Vomiting: K/M 24/55 vs N2O 26/47	Low quality	Luhmann 2006 ¹⁵²

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DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
M/K vs IN M	IV vs. IN		6 mo-12y	Suturing or painless: all completed	Yes (Green) shorter induction time for Ketamine and longer total time for Mid/K	K/M: 1/27 desat; 2/27 vomiting IN M: 1/26 vomiting	Low quality	Acworth 20019
K (with M in 31% of cases)	IV		0-15y	Emergency procedures miscellaneous	N/A	Desat: 1/156; bag/mask: 1/156; vomiting: 6/156 (1 during sedation; 5 after); recovery agitation: mild 2/156	Non RCT	Green 1998 ^{81,82} Green 1998 ^{8281,82}
K vs. K / M	IV ket	Not stated	39 days to 22 years	Fracture reduction; laceration repair; lumbar puncture; imaging; dental	N/A	Desat and laryngospasm: 6.1% (91/1492)	Non RCT	Roback 2005 ¹⁸⁹
	IV Ket/ Midaz.	Not stated	4.8 months to 18 years		N/A	Desat and laryngospasm: 10% (30/299)	Non RCT	
K/M vs. F/M	IV		3/12 – 15 years-	Insertion CVC	Yes (Green)	Yes (Green)	Non RCT	Lucas Da Silva 2007 ¹⁵⁰
M/F vs. M Procedural sedation	IV	Not stated	19 days to 32 years; median 6.7 years	A & E	N/A	?? Respiratory adverse events reported and included oxygen saturation <90%, apnea or laryngospasm Midazolam/ fentanyl 19.3% (65/336) Midazolam 5.8% (15/260) ;vomiting - Midazolam/ fentanyl 1.8% (6/336) Midazolam 0.8% (2/260)	Non RCT	Roback 2005 ¹⁸⁹
IV F/M vs. M Procedural sedation	IV	Mean fentanyl dose: 2.7 mcg /kg Midazolam 0.1 mg/kg;	0-21 years	A & E	N/A	686 vs 65 Complications reported as total adverse events: 23.5% vs. 1.5%	Non RCT	Pitetti 2003 ¹⁸²
IV Mo/M vs. IV	IV	Midazolam 0.1	0-21 years	A & E	N/A	48 vs. 65 Complications reported as	Non RCT	Pitetti 2003 ¹⁸²

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DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
M Procedural sedation		mg/kg Morphine dose not stated				total adverse events: 16.7% vs. 1.5%		
IVP/LA vs. IV M/IV K/IV F	IV	PRO: initial: 2.5 mg/kg in children, 3 mg/kg in infants; maint: 200 mcg/kg/min Lidocaine: 0.1 mL	overall: 7.3 y (SD5.7) PRO 7.5 y (SD5.7) Mid/Keta/Fenta 6.93 y (SD5.8)	Intraarticular steroid injection, bronchoscopy, bone marrow aspiration/biopsy, trans- oesophageal echocardiography, PEG/Gastroscopy, Other	Yes (Green)	Yes (Green) with appropriate training	Very low	Vardi 2002 ²¹⁹
		M: 0.1 mg/kg Keta: 2mg/kg Fenta: 2mcg/kg						
IV P/O analgesics (either Mo or F)		PRO: initial dose 1 mg/kg (max 40 mg); supplemental doses 0.5 mg/kg (max 20 mg) Morphine: 0.1 mg/kg (max 5 mg) Fentanyl: 1-2 mcg/kg (max 50 mg)	Overall: 1-18y Median : 8 y	Fractures, dislocations, examination of ocular burn	Not reported	Yes (Green)	Very low	Bassett 2003 ²⁵
Oral M/N ₂ O vs. Placebo/N ₂ O	Oral	0.5 mg/kg	mean age 4.1 y (range: 2-6)	Suture/laceration repair	Yes (Green) all completed the procure	Yes (Green) no events of aspiration, cardio-repiratory and NSD in vomiting	Low- moderate	Luhmann 2001 ¹⁵¹
IV M/IV K vs. IV	IV	0.1 mg/kg	(range 2-14 y)	Lumbar puncture	Yes (Green)	No(red)	Low	Dilli 2008 ⁵⁵

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DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
K/Placebo			mean age: IV M/IVKeta 7.1 y (SD3.9) IV Keta 6.0 y (SD3.5)		favours Mid group for induction time and for parents satisfaction it is also reported as significant (p=0.001), NSD for recovery time	favours Keta/Placebo for O2 desat (Red)		
IV M/IV K vs. IV K	IV	0.05 mg/kg	(range 1-15 y) mean age (IQR range) IV M/Keta 7 y (4-11) IV Keta 6 y (2-	IV Catheter insertion	Yes (Green) all completed the procedure	Quite safe, no events for and NSD for assisted ventilation Yes (Green) Favours Mid/Keta for vomiting No (red) Favours Keta/Placebo for O2 desat	Low - moderate	Sherwin 2000 ²⁰¹
IV M/IV K vs. IV K	IV	0.1 mg/kg	(range 0.3-18 y) median age (IQ range): IV M/Keta 5.6 y (3.4-9.6) IV Keta 6.8 y (4.4-10.3)	mixed	Yes (Green) all completed the procedure and NSD for patient satisfaction (Green) and insufficient data (NSD) for duration of procedure (Red)	Quite safe, no events for aspiration or external cardiac massage Yes (Green) NSD for assisted ventilation Yes (Green) Favours Mid/Keta for vomiting Favours Keta/Placebo for O2 desat (Red)	Low - moderate	Wathen 2000 ²²⁷
IV M/F	IV, IM,	0.01-0.05	(age range: of	pediatric emergency	N/A	Yes (Green)	Non RCT	Peña 1999 ¹⁷⁸

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DRUG	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence	Ref
COMBINATION							level	
IM M/K	IN,	mg/kg	1,188 patients:	department for		Yes, no events		
IV M/K	Oral		1 mo-21 y)	diagnostic imaging, oral		endotracheal		
IN M/FI			median: 48 mo	and rectal sedation and		intubation; low rates for		
·				analgesia.		assisted ventilation		
						(0.5% to 0.60%) or		
				IM and IV in radiology		vomiting (0.55% to		
				suite		2.5%)		
						(Green)		
						Quite safe for for O2		
						desat (ranged 0.60%		
						to 4%)		
IV AA /V			(age range: 19	fracture reduction,				
IV M/K	IV or	0.1 /	d-32 y)	laceration repair,	N1/A	Vomiting rate from	NI. DCT	D. L. J. 2005189
or	IM	0.1 mg/kg	mean age:	lumbar puncture,	N/A	0.8% to 10.1%	Non-RCT	Roback 2005 ¹⁸⁹
IM M/K			4.9-10.8 y	imaging, other				

6.12.1.2 Evidence to reccomendations for painful procedures

Management of minor trauma in the ED is the most common scenario for brief painless procedures but the principles of effective and safe sedation in the ED can be applied to other areas such as hospital wards.

The GDG agreed that analgesia was a crucial component in any sedation technique for a painful procedure. If local anaesthesia was not practical analgesia by another method would be necessary. Nitrous oxide is potentially effective for cooperative patients but for many children either an opioid or ketamine would be necessary. Opioids are not effective alone and need to be combined with midazolam or propofol. They should be used with caution because they cause respiratory depression especially after the pain of a procedure has abated. The GDG recognised that it was sometimes difficult to titrate the dose of opioid and sedative without "overshooting" and causing unintended deep sedation or anaesthesia. Airway obstruction is a notorious complication in this situation. Ketamine, in contrast, is effective without any other drug and tends to maintain vital reflexes. Moreover it can be given intramuscularly if venous access is difficult and it is applicable to infants and children. The GDG agreed that ketamine sedation had many advantages and that it was a safe technique provided teams were trained to use it safely.

The main debate was whether ketamine sedation, delivered by an ED team, would have economic advantages over anaesthesia, delivered by an anaesthesia team the day after the trauma. The GDG considered that this was a common and realistic scenario and that guidance on this issue would help healthcare provider manage resources efficiently.

6.12.1.3 Cost-effectiveness

The economic evidence for this group was obtained by modelling the treatment pathway for two sedative drugs (ketamine and a combination of fentanyl plus propofol) and comparing these with general anaesthesia (see Appendix F on cost-effectiveness analysis). This was informed by evidence from clinical and safety review as well as GDG expert opinion. Sedation drugs were shown to be cost-saving compared to general anaesthesia, and ketamine was less costly than fentanyl plus propofol.

In one of the studies included in the economic review¹⁷⁹ (see appendix F), fentanyl plus propofol was compared to other combination strategies in children requiring manipulation of forearm fracture in emergency department. It was suggested that fentanyl plus propofol was a dominant strategy because it had the lowest cost and shortest emergency department duration. We would be cautious about concluding that any one sedation technique is the lowest cost (see Appendix F on cost-effectiveness analysis). The evidence from the economic analysis may have some limitations but it is directly applicable for this population group.

We would be cautious about concluding that any one sedation technique is the lowest cost because of the lack of good quality randomised evidence.

In general, the cost of the drugs is less important than the cost of the staff involved. We found that sedation is clearly cost-saving compared to general anaesthesia in cases where the operating physician and / or a nurse is able to administer sedation without the addition of a sedationist physician. In this case, quite a low success rate is required for sedation to be cost-saving.

In cases where the addition of a sedationist physician is required, sedation could still be cost saving but this will depend primarily on:

- The facility cost: we have not captured this in our analysis. It is particularly important when evaluating sedation techniques being carried out in primary care. However facility costs may also be cheaper in A&E, for example, compared to a surgical theatre.
- The success rate: as the success rate gets lower, the cost of a sedation strategy increases. The GDG reported that very high rates of success (above 95%) are achievable with all techniques if patients are selected carefully.
- The speed at which the operation can be conducted under each technique: It seems unclear whether procedures can be delivered more or less quickly with sedation techniques.

6.12.1.4 Recommendation

Recommendation 28

For painful procedures (for example suture laceration or manipulation of fracture) consider using:

For minimal sedation	For moderate sedation	For deep sedation
Oral or intranasal midazolam	Nitrous oxide alone (up to 50% in oxygen)	Ketamine intravenous (or intramuscular if intravenous is difficult)
Nitrous oxide alone (up to 50% in oxygen)	Intravenous midazolam with or without fentanyl	Propofol with or without fentanyl

6.12.2 Painless imaging

Many children will be able to tolerate painless diagnostic imaging tests without sedation drugs. Adequate patient preparation, parental involvement, and a child-friendly environment are important for success (see section 4.3 Psychological preparation). Non-pharmacological methods such as play therapy and distraction techniques may be also helpful for children who are able to co-operate. The majority of children of school age will manage well with these techniques as an alternative to sedation. Highly anxious children may be helped by having anxiolytic drugs. However there are a large number of children who are too ill, in pain or have behavioural problems that prevent them lying still for prolonged imaging.

The target level of sedation will vary according to the imaging procedure. CT scans and echocardiography can be done under moderate sedation. Some children may need to be asleep in order to tolerate complex or prolonged investigations. Examples include MRI and nuclear medicine imaging that may involve the child keeping still for up to an

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hour. MRI can be particularly frightening because it is noisy and involves lying still in an enclosed space. The level of sedation achieved while the patient is asleep is uncertain; they may be moderately sedated and sleeping naturally, be deeply sedated or be anaesthetised. Determining the level of sedation relies on stimulating the patient which may spoil the image.

Ideally "wide margin of safety" drugs cause the patient to sleep and be either moderately or deeply sedated. Not all children will sleep with these drugs. Anaesthesia, by comparison is always effective and short acting. Low doses of anaesthetic agents also cause sedation of uncertain depth however the true depth may be estimated from the drug dose.

6.12.2.1 Summary of evidence in painless imaging

The GDG extracted essential evidence from each drug review and incorporated this evidence into Table 80 and Table 81 below. The tabular presentation was developed as a way to summarise disparate data, ranging across various drug types, drug combinations, specialty areas and procedural techniques. The tables have thus been organised by setting and include the following: painless procedures (imaging), dentistry, painful procedures and GI procedures. The primary efficacy outcome was completion of procedure.

On the basis of the evidence, the GDG made a decision regarding the efficacy and safety (benefits and harms) of each drug and drug combination reviewed. They indicated their decision in the tables by agreeing 'green' (yes) or 'red' (no) for each of these criteria.

Table 80: Drugs alone in painless procedures
Key: 'green' (yes) or 'red' (no), in the efficacy and safety columns, indicate the GDG judgment whether the drug was effective and safe.

DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
CH vs. GA	oral	80 mg/kg to max of 2 g	Not stated	СТ	Favours GA — Red	NR	Low	Thompson 1982*211
High dose CH for CT	oral	100 mg/kg in a single dose with maximum of 2 grams	Mean age 2.18 years	СТ	(Non-RCT)	1 aspiration (severe mental retardation) 2 ETT due to obstruction by tongue (1 profound retardation) 4.3% vomiting Yes (Green) for ASA 1-2	Prospective cohort, N=326 ? Low	Greenberg Faerber, & Aspinall 1991
High dose CH for MRI	oral	100 mg/kg	Not stated	MRI	(Non-RCT)	Vomiting 4% Yes (Green)	Prospective cohort, N=300 ? Low	Greenberg, 1993 ⁸⁴
CH sedation for diagnostic imaging	oral	64 + 13 mg/kg chloral hydrate	Not stated	MRI	(Non-RCT)	GI side-effects Yes (Green)	Prospective cohort N=336 Low	Malviya 2000 ¹⁵⁵
CH: Intermediate vs. high dose	oral	70 mg/kg vs. 100 mg/kg	Mean: 38 + 31 months	MRI	Ns for complet; induc favour high Yes (Green)	Yes (Green)	Moderate	Marti-Bonmati 1995* ¹⁵⁸
CH for effective and safe sedation	oral	Chloral hydrate syrup 68 +/- 1 mg/kg	Mean age 41 + 30 months	MRI	(Non-RCT)	Vomiting 6.9% Yes (Green)	Prospective cohort N=596 Low	Ronchera-Oms 1994 ¹⁹²
CH Sedation of neurologically impaired children for	oral	50 – 100 mg/kg to a maximum dose	Mean age 28.2 + 18.1	MRI	(Non-RCT)	0.2% vomiting, 0.5% SpO2<90%	Retrospective cohort N=888	Cortellazzi 2007 ⁴³

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DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
MRI		of 1.5 g/kg	months			Yes (Green)	(Neuro impaired)	
CH vs. music therapy*	oral	60 mg/kg with max of 1.5 g	1 month -5 years	EEG	Favours music therapy Green	NR	Low	Loewy 2005 ¹⁴⁹
CH for effective and safe sedation	oral	Median dose of chloral hydrate was 77 mg/kg	3 weeks to 14 years; median age 13 months	Echocardiography	(Non-RCT)	Vomiting 6% Drop in SpO2>5% baseline in 6% (children with heart disease) No (red) for children with heart disease	Prospective cohort N>400	Napoli, Ingall, & Martin 1996 ¹⁶⁹
CH for sedation for echocardiography	oral	Oral chloral hydrate (80 mg/kg, maximum 1 g)	Birth to 64 months	Echocardiography	(Non-RCT)			Heistein 2006 ⁹¹
High dose CH for opththalmic examination	oral	80-100 mg/kg chloral hydrate not to exceed 3 g.	1 month - 5 years	Ophthalmic examination	(Non-RCT)	No vomiting or desaturation Yes (Green)	Prospective cohort N=302	Fox 1990 ⁶⁹
P/LA	IV	PRO rarely: 2 mg/kg PRO maint 1-2 mg/kg PRO cont infusion initiated at 150 mcg/kg/min LA: at discretion of intensivist	Overall range: 0 mo to 21y	MRI, CT, nuclear medicine, lumbar puncture, intratechal chemotherapy, bone marrow aspirates, electroencephalogram, evoked potentials, hearing tests	Not reported	Yes (Green)	Very low	Vespasiano 2007 ²²⁴

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Table 81: Drugs combination in painless procedures

Key: 'green' (yes) or 'red' (no), in the efficacy and safety columns, indicate the GDG judgment whether the drugs combination was effective and safe.

DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
IV F + M and IV F	IV	Not stated	Mean age of total sample was 4.8 years + 4.6	lmaging		No adverse outcomes observed	Very low	Sanborn 2005 ¹⁹⁶
P/O ₂ CH/O ₂	IV	PRO: 2 mg/kg after iv access + dilute PRO infusion at a rate of 80-140 mcg/kg/min CH: children <1 y 75 mg/kg (max 2g)	Overall: <1 mo to 17y PRO range: <1 moto 17y CH range: <1 mo to 7 y	Scans of the head, thorax, abdomen, pelvis and spine	Not reported	Yes (Green)	Very low	Merola 1995 ¹⁶⁷
S+ N ₂ 0	inhal	1.8-2% sevoflurane; 50% N20	1 day-12 months	MRI	Yes (Green)	Yes (Green)	Low	De Sanctis Briggs 2005 ⁵¹

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6.12.2.2 Evidence to recommendations for painless imaging

Of all the imaging techniques MRI is the most common scenario in which sedation may be needed. MRI usually lasts between 30 and 60 minutes and CT imaging is much shorter. To be still enough, the patient usually needs to be sleeping, and the true target level of sedation is uncertain; it may be moderate, deep or anaesthesia. The GDG agreed that the ideal sedation method should not cause sedation much longer than the scan itself. For this reason, techniques such as propofol or sevoflurane have advantages of fast induction time, certainty of completion, and rapid recovery. The GDG agreed however that these techniques should be classified as anaesthesia rather than deep sedation. Many children presenting for imaging are uncooperative because they are young, they have a behavioural problem or because they are distressed or in pain. A further advantage of these techniques is that they can be used in all age groups and all types of patients.

Infants who sleep after a feed may lie still enough without any sedation. Also, many children can be calmed sufficiently and persuaded to lie still without the use of sedation. Occasionally an anxiolytic drug may help them but only if they are cooperative. Otherwise however, children who are uncooperative need sedation or anaesthesia. The GDG considered that sedation with Choral hydrate was an effective and safe alternative to anaesthesia but only in children less than 15kg. The success rate of chloral hydrate may be maximised by careful patient assessment and selection.

Choral hydrate causes sleep lasting approximately one hour and is therefore less appropriate for scans lasting a few minutes. An economic advantage of chloral is that it does not require the services of an anaesthesia team.

Other types of painless imaging such as trans-thoracic echocardiography or EEG do not require the child to be completely immobile and they may therefore be managed with minimal or moderate sedation. Anaesthesia would not be appropriate for these investigations either because the risks outweigh the benefits or, in the case of EEG, anaesthesia may suppress the EEG signal under investigation.

6.12.2.3 Cost-effectiveness

The economic evidence for this group was obtained by modelling the treatment pathway for high dose chloral hydrate and comparing this with general anaesthesia (see Appendix F on cost-effectiveness analysis). This was informed by evidence from clinical and safety review as well as GDG expert opinion. High dose chloral hydrate was more costly than general anaesthesia because this type of sedation was assumed to be less successful but also to require the same staff levels as general anaesthesia.

However, we would be cautious about concluding that any one sedation technique is the lowest cost because of the lack of good quality randomised evidence.

In general, the cost of the drugs is less important than the cost of the staff involved. We found that sedation is clearly cost-saving compared to general anaesthesia in cases where the operating physician and / or a nurse is able to administer sedation without the addition of a sedationist physician. In this case, quite a low success rate is required for sedation to be cost-saving.

In cases where the addition of a sedationist physician is required, sedation could still be cost saving but this will depend primarily on:

- 1 2 3 4
- The facility cost: we have not captured this in our analysis. It is particularly important when evaluating sedation techniques being carried out in primary care. However, facility costs (CT / MRI rooms) may also be cheaper when compared to a surgical theatre.
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- The success rate: as the success rate gets lower, the cost of a sedation strategy increases. The GDG reported that very high rates of success (above 95%) are achievable with all techniques if patients are selected carefully.
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- The speed at which the operation can be conducted under each technique: It seems unclear whether procedures can be delivered more or less quickly with sedation techniques.

11 6.12.2.4 Recommendations

Recommendation 29

Do not routinely use ketamine or opioids for sedating children or young people for painless imaging procedures.

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Recommendation 30

For children and young people who are unable to tolerate a painless procedure (for example during diagnostic imaging) consider one of the following:

- Chloral hydrate (oral) for children under 15Kg
- Propofol
- Sevoflurane.

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6.12.3 Endoscopy

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Gastrointestinal (GI) endoscopy procedures are commonly required in children and young people. The procedures consist of upper GI endoscopy (often called oesophagogastro duodenoscopy [OGD] or gastroscopy) and lower GI endoscopy (colonoscopy). In children and young people the majority of procedures are diagnostic; however, there are some therapeutic techniques performed (for example oesophageal dilatation and polypectomy) that make the procedure more technically difficult and time consuming. Upper endoscopy is uncomfortable but not usually painful. The target level of sedation during upper endoscopy is considered to be no deeper than moderate sedation. The child or young person will need to maintain their airway reflexes for an OGD because vomiting and regurgitation are common. Moreover the endoscope itself may obstruct the airway in an unconscious patient. Colonoscopy may be uncomfortable but can be tolerated by many children and young people under moderate sedation. The use of an analgesic drug is often necessary. If sedation is not successful, anaesthesia should be used and in many centres anaesthesia is the only method used. Nevertheless in a recent survey of members of the British Society of Paediatric Gastroenterology, Hepatology and Nutrition sedation was found to be used in about 30% of units and especially for children of secondary school age and older.

Recently, anaesthesia agents have been used to sedate to the target level that the patient needs in order to tolerate the procedure. This is usually deep sedation but in most cases the patient is anaesthetized albeit for a brief period. Such a method does not necessarily require tracheal intubation and allows effective short acting sedation. Whoever administers anaesthetic agents must be trained to manage the complications of airway obstruction and respiratory depression (see section 4.4 Personnel and training).

6.12.3.1 Summary of evidence in endoscopy

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The GDG extracted essential evidence from each drug review and incorporated this evidence into Table 82 and Table 83 below. The tabular presentation was developed as a way to summarise disparate data, ranging across various drug types, drug combinations, specialty areas and procedural techniques. The tables have thus been organised by setting and include the following: painless procedures (imaging), dentistry, painful procedures and GI procedures. The primary efficacy outcome was completion of procedure.

15 On the basis of the evidence, the GDG made a decision regarding the efficacy and 16 safety (benefits and harms) of each drug and drug combination reviewed. They indicated their decision in the tables by agreeing 'green' (yes) or 'red' (no) for each of 18 these criteria.

Table 82: Drugs alone in endoscopy

Key: 'green' (yes) or 'red' (no), in the efficacy and safety columns, indicate the GDG judgment whether the drug was effective and safe.

DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
IN M vs. Placebo	IX	0.2 mg/kg	age range: 2-12 y	Endoscopy	Yes (Green) NSD in distress score	Not reported	RCT Mod- low	Fishbein 1997 ⁶⁷
IV P vs. IV F/IV P	IV	Fentanyl: 1mcg/kg PRO: 3mg/kg TA: Lidocaine larynx and EMLA cream	PRO/Fenta 6.8 y (SD2.8) PRO/TA 6.7 y (2.9)	Endoscopy	Yes (Green)	Yes (Green)	low	Disma 2005 ⁵⁶
P/LA/TA	IV	PRO: mixed according to age/weight LA: 1 to 10 mg TA: EMLA cream	overall <1 to <21y	Upper gastrointestinal endoscopy procedures	Not reported	Yes (Green)	Very low	Barbi 2006 ²⁴
P/LA/TA	IV	PRO: mixed according to age/weight LA: 1 to 10 mg TA: EMLA cream	overall <1 to <21 y	Upper endoscopies, colonoscopies, painful procedures	Not reported	Yes (Green)	Very low	Barbi 2003 ²³

Table 83: Drugs combination in endoscopy

Key: 'green' (yes) or 'red' (no), in the efficacy and safety columns, indicate the GDG judgment whether the drugs combination was effective and safe.

DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
P/K vs P/F	IV	Ket 1mg/kg Prop 1.2mg/kg Fent 1mcg/kg	1-16years	Endoscopy	Yes (Green)	No (Red) Vom 15% ket (0 fent group) p=0.012 Desat no difference (6-9%)	Low- moderate	Tosun 2007 ²¹³
K/M/Me	IV	0.75-2 mg/kg		Endoscopy	N/A	Yes (Green) 1.2% assisted vent	Non RCT	Gilger 2004 ⁷⁶
Oral M/IV P vs. IV P	Oral	0.5mg/kg	mean age: Oral M/IV PRO 8 y (SD3) PRO 9 y (SD3)	Endoscopy	No, favours PRO in recovery time No (red) and NSD for	Not reported	low - Moderate	Paspatis 2006 ¹⁷⁶

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DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
					duration of procedure Yes (Green)			
IV M/IV P vs. IV P	IV	0.1 mg/kg	(range 1-12 y) mean age: IV M 7.1 y (SD3.1) PRO/Lido 6.7 y (2.9)	Endoscopy	Yes, all completed the procedure and NSD in duration of procedure and recovery time Yes (Green)	Yes (Green) NSD in oral- phryngeal airway and O2 desat	Low - moderate	Disma 2005 ⁵⁶
IV M/IV Me vs. Placebo/IV Me	IV	0.051 mg/kg (max 2 mg)	(range 2-12 y)	Endsoscopy (esophagogastroduodenoscopy)	Yes, NSD in distress score and duration of procedure Yes (Green)	Not reported	Low - moderate	Fishbein 1997 ⁶⁷
P/K vs P/F	IV	Ket 1mg/kg Prop 1.2mg/kg Fent 1mcg/kg	1-16years	Endoscopy	Yes (Green)	No (Red) Vom 15% ket (0 fent group) p=0.012 Desat no difference (6-9%)	Low- moderate	Tosun 2007 ²¹³
M/K (IV 98% or IM)/ 15% Me	IV	Ket 1mg/kg and titrated, median 1.34 mg/kg Dose midaz not known		Endoscopy	N/A	'Amber' 3% bag and mask	Non RCT	Green 2001 ⁸⁰
K/M/Me	IV	0.75-2 mg/kg		Endoscopy	N/A	Yes (Green) 1.2% assisted vent	Non RCT	Gilger 2004 ⁷⁶
IV M/F	IV	0.05-0.1 mg/kg;max 2 mg n.b. oral M to anxious children	(range: 0.1-34 y) median: 10 y mean: 9.05 y (SD 5.8)	Endoscopy (esophagogastro duodenoscopies colonoscopies and combined)	N/A	Quite safe no events: aspiration, cardiac arrest, endotracheal intubation 0.16%(2/1226)	Non RCT	Mamula 2007 ¹⁵⁶

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DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
						needed bag/mask ventilation Yes (Green) 1.2% assisted vent 5.2%(64/1226) vomited during recovery Yes (Green) 1.2% assisted vent		
IV F/IV P vs IV M/IV P	IV	Fenta: 1 mcg/kg PRO: 3 mg/kg Mid: 0.1 mg/kg TA: Lidocaine- larynx and EMLA cream	Fenta/PRO 6.8 y (SD2.8) Mid/PRO 7.1 y (SD3.1)	Endoscopy	Yes (Green)	Yes (Green)	Moderate	Disma 2005 ⁵⁶

^{*} Indicates RCT extracted for efficacy review

6.12.3.2 Evidence to recommendations for endoscopy

Gastroenterological endoscopy is uncomfortable. Gastroscopy requires control of pharyngeal and oesophageal reflexes to overcome retching. Colonoscopy may need opioid analgesia. The GDG felt that a large proportion of children and young people requiring these procedures were old enough to be cooperative and that moderate sedation was effective. It was agreed that deep sedation was potentially hazardous if it was administered by untrained practitioners and without safe resources. The choice of opioid to be used in combination with midazolam combination of midazolam was debated. In the early discussions of the GDG it was agreed that evidence for pethidine would not be sought because it had a longer action than fentanyl and because it was not widely used. In respect of endoscopy however the GDG was advised by one of its members that pethidine may be in common use for colonoscopy. Pethidine may be safer than fentanyl if practitioners were more familiar with its use because they would be less likely to "overshoot" and cause unconsciousness or respiratory depression. Training in the use of any new technique was considered to be crucial.

It was agreed that moderate sedation may not always be effective enough and that sometimes sedation may have to be abandoned. Patient assessment and selection will be important to minimise sedation failure. Occasionally sedation can become too deep and this results in prolonged recovery.

The GDG agreed that whenever moderate sedation is ineffective a short acting titrateable drug such as propofol was ideal. Propofol however readily causes unconsciousness and the hazard of pulmonary aspiration is a special concern with this technique. Staff training and facilities for anaesthesia will be necessary for propofol based techniques. If an anaesthesia team is available either sevoflurane or propofol can be used to induce deep sedation or anaesthesia and this can be applied to children of all ages undergoing procedures of variable length. Tracheal intubation may be needed for gastroscopy and this can be readily achieved by an anaesthesia team.

The GDG agreed that there were potentially important economic advantages of using propofol rather than moderate sedation and that this should be considered by healthcare providers.

6.12.3.3 Cost-effectiveness

The economic evidence for oesophago-gastroscopy was obtained by modelling the treatment pathway for midazolam and comparing it with general anaesthesia (see Appendix F on cost-effectiveness analysis). The economic evidence for colonoscopy was obtained by also modelling the treatment pathway for midazolam plus fentanyl and comparing this combination with general anaesthesia. This was informed by evidence from clinical and safety review as well as GDG expert opinion.

Midazolam was shown to be less expensive than general anaesthesia in oesophago-gastroscopy, and in colonoscopy, the combination sedation strategy, midazolam plus fentanyl, was less expensive than general anaesthesia. However, we would be cautious about concluding that any one sedation technique is the lowest cost because of the lack of good quality randomised evidence.

In general, the cost of the drugs is less important than the cost of the staff involved. We found that sedation is clearly cost-saving compared to general anaesthesia in cases where the operating physician and / or a nurse is able to administer sedation without the

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addition of a sedationist physician. In this case, quite a low success rate is required for sedation to be cost-saving.

In cases where the addition of a sedationist physician is required, sedation could still be cost saving but this will depend primarily on

- The facility cost: we have not captured this in our analysis. It is particularly important when evaluating sedation techniques being carried out in primary care. However facility costs (endoscopy room) may also be cheaper when compared to a surgical theatre.
- The success rate: as the success rate gets lower, the cost of a sedation strategy increases. The GDG reported that very high rates of success (above 95%) are achievable with all techniques if patients are selected carefully.
- The speed at which the operation can be conducted under each technique: It seems unclear whether procedures can be delivered more or less quickly with sedation techniques.

15 **6.12.3.4 Recommendations**

Recommendation 31	Consider intravenous midazolam to achieve minimal or
	moderate sedation for upper gastrointestinal endoscopy.

Recommendation 32

Consider fentanyl (or equivalent opioid) and intravenous midazolam to achieve moderate sedation for lower gastrointestinal endoscopy.

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6.12.4 Dentistry

Dental treatment can be made virtually painless by local anaesthesia. Nevertheless, local anaesthesia insertion itself is painful and anxiety about dental procedures is widespread in children and young people ²²¹. The provision of adequate anxiety control is an integral part of the practice of dentistry. Many anxious children can be satisfactorily treated using minimal or moderate sedation using a technique called relative analgesia (RA) which combines behaviour management techniques with inhaled nitrous oxide and oxygen. In current practice RA is the basis of paediatric dental sedation but this approach is unsuccessful in some children. In such cases, control of pain and anxiety poses a significant barrier to dental care and deep sedation of anaesthesia are often seen as the only alternative options. Whereas these may be appropriate for extensive painful dental procedures, the risks of deep sedation and anaesthesia may be unnecessary or inappropriate for minor treatments. Many anxious patients can be calmed by behavioural therapy over one or more attendances. Moreover special facilities are needed for deep sedation or anaesthesia. The General Dental Council (GDC) has stated that dental treatment under general anaesthesia should "only be carried out when it is judged to be the most clinically appropriate method of anaesthesia; and only take place in a hospital setting with critical care facilities" 75.

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These considerations have resulted in an increased emphasis on the safe provision of conscious sedation instead of a reliance on general anaesthesia. Since the publication of 'A Conscious Decision' in 2000 general anaesthesia for dentistry has ceased in the primary care setting ⁵³. Since 2002, anaesthesia has been prohibited in the non-hospital setting. The vast majority of dental treatment however, is carried out in a primary care setting.

For successful dental treatment under sedation the patient needs to be able to open their mouth and therefore sedation should be limited to moderate sedation. Deep sedation can easily become anaesthesia by accident. In the primary care setting it is particularly important to ensure that sedation technique has a wide margin of safety ^{198,206}. Recent developments in dental sedation have seen the introduction of new techniques such as the combination of sedation drugs including low doses of anaesthetic agents. The evidence pertaining to some of these newer techniques was reviewed by the GDG.

6.12.4.1 Summary of evidence in dentistry

The GDG extracted essential evidence from each drug review and incorporated this evidence into Table 84 and

Table 85 below. The tabular presentation was developed as a way to summarise disparate data, ranging across various drug types, drug combinations, specialty areas and procedural techniques. The tables have thus been organised by setting and include the following: painless procedures (imaging), dentistry, painful procedures and GI procedures. The primary efficacy outcome was completion of procedure.

On the basis of the evidencev the GDG made a decision regarding the efficacy and safety (benefits and harms) of each drug and drug combination reviewed. They indicated their decision in the tables by agreeing 'green' (yes) or 'red' (no) for each of these criteria.

Table 84: Drugs alone in dental procedures
Key: 'green' (yes) or 'red' (no), in the efficacy and safety columns, indicate the GDG judgment whether the drug was effective and safe.

DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
50% N ₂ O vs.100% O ₂	Inhaled	50%	36-55 months	Dental	NSD in quiet behaviours	Not reported	Low	McCann 1996 ¹⁶⁰
40% N ₂ O/O ₂ vs.100% O ₂	Inhaled	40%	5-9 years	Dental	Yes (Green) Distress score favours nitrous oxide	Yes	Low	Primosch 1999 ¹⁸⁵
Oral M/Non-pharma vs. Placebo/Non- pharma	Oral	0.5 mg/kg	< 4 y	Dental	Yes (Green) favours M in procedure completion and duration	Not reported	Very low - low	Kapur 2004 ¹²⁴
Oral M	Oral	0.5 mg/kg; max 10 mg per appt.; mean 8.6 mg/kg	range: 0.9-10.5 y mean: 5.4 y	Dental	N/A	O2 desat: 1.55% (9/579) (Green)	Non RCT	Hulland 2002 ⁹⁷
IN M vs. IM M	IN vs. IM	IN& IM M: 0.2 mg/kg	(range: 1-5 y) mean age: IN M: 3.5 y (SD0.7) (range 2.5-5) IM M: 3.4 y (SD 0.6) (range 2- 4.5)	Dental	Yes (Green) favours IN M in induction time and recovery	Yes, no events for vomiting	Moderate	Shashikiran 2006 ¹⁹⁹
IN M vs. IM M	IN vs. IM	IN& IM M: 0.2 mg/kg	(range: 1-5 y) mean age: IN M: 3.5 y (SD0.7) (range 2.5-5) IM M: 3.4 y (SD 0.6) (range 2- 4.5)	Dental	Yes (Green) favours IN M in induction time and recovery	Yes (Green) no events for vomiting	Moderate	Lee-Kim 2004 ¹³⁷
CH: High dose vs. low dose*	oral	50 mg/kg vs. 75 mg/kg	Mean: 31 months	Dental	Favours high dose Yes, (green)	Not reported	Low	Houpt 1985 ⁹⁶

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DRUG (alone, or with local anaesthesia)	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
					wrong technique for procedure????			
CH vs. IN M	oral/ IN	62.5 mg/kg CH; 0.2 mg/kg midazolam	Mean: 41.8 months + 11.4 months	Dental	NS but recovery favours midazolam Yes (Green)	Not reported	Low	Dallman 2001* ⁴⁹
CH/hydroxyzine vs M/acetaminophen	oral	50 mg/kg not to exceed 1 g and 25 mg hydroxyzine vs 0.5 mg/kg midazolam with acetaminophen 10 mg/kg	Average 48 months in CH group vs. 42 months in Midazo-lam group	Dental	NSD Yes (Green)	NR	Moderate	Reeves 1996*187
Oral TS vs Oral M	Oral	TRI 70 mg/kg M: 0.5 mg/kg	overall: 3-9 y	dental - mixed: extractions, restorations, pulpotomies, brief	No (Red)	Not reported	Very low	Singh 2002 ²⁰³

Table 85: Drugs combination in dental procedures
Key: 'green' (yes) or 'red' (no), in the efficacy and safety columns, indicate the GDG judgment whether the drug was effective and safe.

DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
CH + N ₂ O vs. Placebo + N ₂ O*	Oral/inhaled	50 mg/kg CH + 50% nitrous oxide vs. placebo + 50% nitrous oxide	19-41 months	Dental	Outcomes not reported (Crying & movement scores suggested chloral more effective, but not uniformly so) Yes (Green)	Vomiting in 10.5% chloral group, 5% placebo Green	Moderate	Houpt 1989 ⁹⁵
CH/hydroxyzine and N ₂ O	Oral/inhaled	Average dose of chloral hydrate 776 mg (55 mg/kg)	Mean age 2.6 years	Dental	(Non-RCT)	Vomiting 8.1% Green	Retrospective, non-RCT Low	Needleman, Joshi, & Griffith, 1995 ¹⁷¹
N ₂ O vs. Behavioural management	Inhaled	Not stated	Not stated	Dental	Yes (Green) Anxiety score favours nitrous Oxide	Not reported	Very low	Veerkamp 1993 ²²² & Veerkamp 1995 ²²⁰

DRUG COMBINATION	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
30% N ₂ O vs. Transmucousal M	Inhaled	30%	10-16 years	Dental	Yes (Green)	Yes (Green)	Low	Wilson 2007 ²³³
40% N ₂ O + IV M vs. Medical air + IV M	Inhaled	40%	Mean age 9.5 years	Dental	Yes (Green) Favours nitrous oxide	Yes (Green)	Moderate	Averley 2004 ¹⁹
40% N ₂ O + IV M vs. 0.3% S and 40% N ₂ O + IV M	Inhaled	40%	Mean age 9.6 years	Dental	Favours sevoflurane + nitrous oxide group	Yes (Green)	Moderate	Averley 2004 ¹⁹
0.3% S and 40% N ₂ O + IV M vs. Medical air +IV M	Inhaled	40%	Mean age 9.1 years	Dental	Yes (Green) Favours sevoflurane and nitrous oxide	Yes (Green)	Moderate	Averley 2004 ¹⁹
30% N ₂ O vs. IV M	Inhaled	30%	12-16 years	Dental	Yes (Green) Favours nitrous oxide	Yes (Green)	Low	Wilson 2003 ²²⁹
30% N ₂ O/70% O ₂ vs. Oral M	Inhaled	30%	10-16 years	Dental	Yes (Green) Favours nitrous oxide	Yes (Green)	Low	Wilson 2002 a & b ^{231,232}
50% N ₂ O vs. 50% nitrogen + O ₂	Inhaled	50%	1 month – 18 years	Dental	Yes (Green)	Yes (Green)	Low	Fauroux 2004 ⁶⁴
\$ + N ₂ O vs. N ₂ O	inhal	0.1-0.3% sevoflurane; 40% N20	3-10 years mean age: 6.0 y (sevoflurane + N20); 6.2 y (N20)	dental	Yes (Green)	Yes (Green)	Moderate	Lahoud 2002 ¹³³
IN M/N ₂ O (50%) 0.3 mg/kg vs. 0.2 mg/kg	IN 0.3 mg/kg vs. 0.2 mg/kg	0.3 mg/kg	mean age 2.7 y (range: 1.7- 3.5)	Dental	Yes (Gree) all completed procedure	Yes (Green) no events for vomiting	Moderate	Fuks 1994 ⁷¹
IN M/ N ₂ O (30- 50%) 0.3 mg/kg vs. 0.2 mg/kg	IN 0.3 mg/kg vs. 0.2 mg/kg	0.3 mg/kg vs. 0.2 mg/kg	(range 5-20) average: higher dose: 0.3-11.6 y lower dose: 0.2-13.6 y	Dental	Yes, NSD in completion of procedure and duration of procedure (Green)	Yes (Green), no events for assisted respiration or vomiting after procedure and NSD in O2 desat or vomiting during procedure	Very low – low - moderate	Fukuta 1994 ⁷²

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DRUG	Route	Dose	Age	Procedure	Efficacy	Safety	Evidence level	Ref
COMBINATION								
Oral M/N ₂ O (40%) vs. IN M/N ₂ O (40%)	Oral vs. IN	Oral M 0.5 mg/kg IN M 0.2 mg/kg	(range 1.5-5.9) mean age: Oral M: 3.3 y IN M: 3.1 y	Dental	Yes, NSD in completion of procedure (Green)	Yes (Green), NSD for O2 desat	Very low - low	Hartgraves 1994 ⁸⁹
Oral M/N ₂ O 45% vs. IN M/N2O 45%	Oral vs. IN	0.7 mg/kg vs. 0.3 mg/kg	mean age: Oral M 3.4 y (SD11) IN M 3.2 y (SD10)	Dental	Ye (Green)s, favours IN M for induction and total time	Not reported	Low - moderate	Lee-Kim 2004 ¹³⁷

6.12.4.2 Evidence to recommendations for dental procedures

The GDG acknowledged the considerable sedation experience of UK dentists. Many children currently require both dental extractions and conservative treatment and many are too anxious to allow the insertion of local anaesthesia. Sedation for dentistry requires that the patient opens their mouth and therefore they need to remain conscious. Moderate sedation with intravenous midazolam, is considered to be effective for selected children and young people who are cooperative, and younger children who can tolerate a nasal mask can be managed with nitrous oxide.

In the past, if these were not effective, anaesthesia has often been the only alternative. The GDG agreed that additional sedation techniques could be effective for patients who cannot be managed by midazolam or nitrous oxide. If demand is high, alternative sedation techniques would be necessary. The common concern is that additional sedation drugs, especially in combination, may not be predictable enough for widespread use. Sevoflurane and propofol for example may only be safe enough for use by specialist sedation teams.

The GDG agreed that there were potential important economic advantages of avoiding hospital based anaesthesia services. The training of dental sedation teams was regarded as crucial.

6.12.4.3 Cost-effectiveness

The economic evidence for dental procedures in *children* was obtained by modelling the treatment pathway for four sedation drugs (nitrous oxide plus oxygen, nitrous oxide plus midazolam, sevoflurane plus nitrous oxide, sevoflurane plus nitrous oxide plus midazolam) and comparing these with general anaesthesia (see Appendix F on costeffectiveness analysis). The economic evidence for dental procedures in *adolescents* was obtained by modelling the treatment pathway for midazolam and comparing this with general anaesthesia. This was informed by evidence from clinical and safety review as well as GDG expert opinion.

Nitrous oxide plus oxygen was the least expensive drug for dental procedures in children. Midazolam was less expensive than general anaesthesia for dental procedures in adolescents. However, we would be cautious about concluding that any one sedation technique is the lowest cost because of the lack of good quality randomised evidence.

In one of the studies reviewed for dental procedure in children²⁰⁰ it was suggested that sedation would cost less than general anaesthesia (see appendix F). Nitrous oxide²⁷ and advanced conscious sedation¹⁰⁰ were suggested to be less expensive than general anaesthesia for dental procedure in children.

In general, the cost of the drugs is less important than the cost of the staff involved. We found that sedation is clearly cost-saving compared to general anaesthesia in cases where the operating dentist and / or a nurse is able to administer sedation without the addition of a sedationist dentist. In this case, quite a low success rate is required for sedation to be cost-saving.

In cases where the addition of a sedationist dentist is required, sedation could still be cost saving but this will depend primarily on

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11 6.12.4.4 Recommendation

Recommendation 33

For a child or young person who cannot tolerate a painful dental procedure with local anaesthesia alone, consider one of the following techniques to achieve moderate sedation:

The facility cost: we have not captured this in our analysis. It is particularly

important when evaluating sedation techniques being carried out in primary care

The success rate: As the success rate gets lower, the cost of a sedation strategy increases. The GDG reported that very high rates of success (above 95%) are

The speed at which the operation can be conducted under each technique: It

seems unclear whether procedures can be delivered more or less quickly with

achievable with all techniques if patients are selected carefully.

- Nitrous oxide and oxygen (titrated to the child's needs and using a maximum of 70% nitrous oxide)
- Midazolam.

(e.g. dental procedures).

sedation techniques.

If these sedation techniques are not suitable or effective, consider referral to a specialist team for other sedation techniques (for example midazolam in combination with nitrous oxide and/or sevoflurane).

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6.13 Research recommendations

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For children and young people under the age of 19 undergoing minor painful procedures, what potent analgesic drugs can be combined with midazolam to provide safe moderate sedation?

Why it is important

Midazolam has a strong safety profile in inducing either minimal or moderate sedation. For painful procedures midazolam should be combined with analgesia. Ideally analgesia is achieved by local anaesthesia. Sometimes local analgesia is insufficient and potent opioid analgesia is necessary. The combination of potent opioid and midazolam can cause deep sedation and airway obstruction. These effects can be managed safely but will involve extra resources. If would be safer if a technique could be developed that was both reliable and had a wide margin of safety. Prospective and retrospective audit data are available to help guide the choice of opioid and the doses. A randomised controlled trial is needed to test the efficacy and safety of these combinations.

1 For children and young people under the age of 19 undergoing diagnostic or 2 therapeutic procedures under sedation with ketamine, how can the vomiting be 3 reduced? 4 Why it is important 5 Ketamine is demonstrated to have a strong efficacy and safety profile in enabling safe 6 sedation and as an analgesic drug useful for painful procedures in children and young 7 people. Its main side effect is vomiting in approximately 10% of patients. No data is 8 available on whether antiemetic drugs prevent vomiting. The GDG suggested an RCT 9 study comparing ketamine + placebo versus ketamine with antiemetic 10 11 > For children and young people under the age of 19 undergoing diagnostic or 12 therapeutic procedures, are the procedures carried out under sedation 13 delivered more cost effective compared to general anaesthesia? 14 Why it is important 15 Anaesthesia or an "Anaesthetist led service" has the advantage over sedation because it 16 usually has faster onset and offset and is more predictable. It may be more expensive 17 and is a scarce resource. Data comparing the efficiency of sedation in comparison with 18 anaesthesia for certain procedures are not available. Models of care need to be 19 developed and studied to whether anaesthesia or sedation gives the best value for 20 money. With such data, efficient services can be planned. 21 22 For children and young people under the age of 19 undergoing endoscopy, is 23 propofol (with or without: analgesia, another drug or psychological techniques) 24 effective, safe and cost effective for sedation (at minimal and moderate levels) 25 in comparison with midazolam (with or without opioids) or with general 26 anaesthesia? 27 Why it is important 28 Propofol is a short acting anaesthetic agent that can be used to achieve any target 29 sedation level. The dose necessary for gastrointestinal endoscopy however usually has a 30 tendency to cause anaesthesia albeit for a short period of time. It would be helpful to 31 know the dose limitation that is unlikely to cause deep sedation because this dose may 32 be effective and safe enough. Moderate sedation with propofol could be compared with 33 another sedation technique such as midazolam with or without opioid. It could also be 34 compared with a general anaesthetic dose of propofol. 35 36 For children and young people under the age of 19 undergoing painful 37 procedures, is ketamine effective and safe for sedation in comparison with 38 propofol? 39 Why it is important

Both ketamine and propofol are safe and effective drugs suitable for painful procedures. Propofol however has a tendency to cause deep sedation and anaesthesia in which the airway and breathing may need an intervention or support. Ketamine has few appreciable effects on the airway and breathing but has a longer recovery time than propofol and causes vomiting.

What are the safety and efficacy profiles of sedation techniques in current practice?

Why it is important

Data on the safety of sedition in the UK are not available. A large prospective database of sedation cases, including data on drugs, procedures, the level of sedation and any complications, would be beneficial in not only providing definitive data on the safety of sedation but also actively promoting safe practice. The GDG suggests that a national registry for paediatric sedation is established for the purpose of creating a database with sufficient data.

Is patient-controlled sedation with propofol feasible in adolescents and children?

Why it is important

Propofol in low dose is an excellent anxiolytic. Patient-controlled sedation has been validated in adults undergoing dental procedures and endoscopy for safety and efficacy. Giving the patient control of their sedation has important psychological benefits. The study would involve developing new pump technology, paediatric software and a child friendly patient-activation system. There would have to be an open pilot evaluation to establish safety and efficacy followed by a randomised-controlled trial versus IV midazolam.

7 Swimming in the sea of uncertainty in relation to sedation experience for children and young people undergoing diagnostic and therapeutic procedures

5 "To study the phenomenon of disease without books is to sail an uncharted sea, whilst to study books without patients is not to go to sea at all"

7 Osler (circa 1900)

7.1 Introduction

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The importance of patient input to healthcare is not underestimated, but rarely is it properly achieved in providing real time comment on how the experience has been shaped and the resultant impact of this experience on the patient's approach to future health care interventions. Whilst this has been achieved in adult populations to varying degrees of success, in the children's and young people population, this is extremely rare, and little is reported in the literature. Having children and young people represented on the GDG is of course standard practice in NICE guideline development, but this has almost uniquely been through advocacy of carer's. In trying to understand the challenges of providing a safe and effective sedation service, this feedback is crucial in determining how experts interpret evidence and remain sensitive to key clinical issues that impact on the child or young person receiving sedation. Early in development, NCGC in supporting this guideline and with the agreement from NICE made an ambitious decision to try and establish a snapshot of what it is like to be a child receiving sedation across a range of clinical contexts. The benefit of collecting real-time feedback in informing and shaping recommendations for practice is self evident, and through engagement with a developing methodology (National Paediatric Toolkit) NCGC commissioned some primary data collection at Alder Hey Children's NHS Foundation Trust. The Trust is well positioned as England's first paediatric health promoting hospital accredited by the World Health Organisation and is one of Europe's biggest and busiest children's hospitals, providing care for over 200,000 children each year.

7.2 Development and conduct of the survey

The survey was carried out as part of a pilot project, with this particular survey focus added to a menu of survey's administered within the Trust. The content of the survey was

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shaped by a sub group of the GDG, with clinicians, technical team members and both
patient carer representatives involved in the shaping of the questions asked. These were
reviewed and signed off in consultation with the rest of the GDG and NICE, and were
targeted at children undergoing painful and non-painful procedures requiring sedation.
The questionnaire was administered using the National Paediatric Toolkit (NPT) software
via hand-held, touch screen computers, a developing technology which is easy to use by
even young children (over the age of four).

The NPT concept has been developed by Alder Hey Children's NHS Foundation Trust in partnership with Priority Research Ltd; throughout its development, children and young people were closely involved and contributed many ideas which have been incorporated into the current data collecting system.

The NPT was considered the system of choice for administering this survey because of a variety of unique advantages which it offers. These include:

- an engaging, cartoon format to maintain children's interest
- a large array of over 900 pre-defined questions, each worded differently for four developmental levels
- all questions available in eleven languages
- full voice-over for all text in all languages
- Disability Discrimination Act (1995) compliant for sensory, visual and hearing impairment
- real time data collection and reporting

The pilot ran from early November 09 for 4 months and was conducted by experienced Alder Hey staff members previously engaged in similar types of data collection using the NPT.

7.3 Survey conduct approval

Patient opinion surveys are growing increasingly in both their conduct and importance, and this helps shape and reshape service delivery in different care settings.

Contextually, until recently, this type of opinion seeking would have been viewed as primary research activity and therefore requiring ethics approval via a local committee or through a national committee, particularly relevant if this multi centre. Following changes in approach, seeking patient opinion is more latterly viewed as part of a quality improvement cycle, and is becoming more and more embedded into routine NHS Trust processes.

For this survey, approval and advice was sought and gained from Alder Hey NHS Foundation Trust's Head of Research and Ethics, Dr Matthew Peak.

7.4 Recruitment

A total of 70 patients undergoing a wide range of procedures were invited to take part, and 63 consented to do so. All departments and clinical areas within the hospital where patients receive sedation participated in the pilot.

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7.5 Limitations of the survey

The limitations of the survey are important to note as this methodology will only describe the experience of the target population in one place at one time event. The 'snapshot' nature if surveys are extremely useful in determining the nature of patient experience and care interventions on a particular day. These cannot be generalised to other settings but findings are extremely helpful if repeat measurement is established so that a time series of events are recorded. Data is also useful as in this case when supporting other data (clinical and cost effectiveness reviews, consensus development), as when triangulated with this 'other' data inevitably enables the GDG in this case to build a clearer picture of what is happening and how to plan improvements in care and experience outcome.

7.6 Summary of main findings

7.6.1 Demographics

- The sample had a even spread of male and female patients (44% male, 46% female, 10% not recorded) and covered a broad age range from under four to over 16 years of age.
- All except one were accompanied by a parent or carer, and for those children
 who could not complete the questionnaire themselves, they were in a good
 position to do so (as expected this was mostly younger children). Acceptability
 and usability of the system was such that nearly 1 in 4 (22%) of the under four
 age group were able to complete the survey themselves.
- Of the 23 children aged nine and over, only one child aged eleven did not complete the survey themselves.
- Only four children (6%) were of black or minority ethnic origin.

25 7.6.2 Clinically relevant data

- The most frequent clinical areas, accounting for almost two thirds of the sample, were:
- Burns (21%)
- Medical and Renal Day Cases (17%)
- Radiology (16%)
- Accident & Emergency (10%)
- 31 The most common agents used for sedation were:
- Entonox (48%)
- 33 Midazolam (30%)
- Oral morphine (14%)

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1	Five procedures accounted for over half of the sample:
2	 Change of wound dressings (22%)
3	• Urodynamics (11%)
4	 Intra-articular steroid injections (10%)
5	Cannulation (6%)
6	 Removal of chest drains (6%)
7	7.6.3 Experience of children and young people receiving sedation
8	Ratings of satisfaction with information and consent issues were high:
9	 The people looking after me were nice to me and helped me feel OK (98%)
10	 I was told everything I wanted to know about what would happen (97%)
11 12	 I was told enough about the sedation (medicine that would make me feel OK and sleepy) (95%)
13	 I had time to ask any questions I wanted (91%)
14	 I was told enough about how I might feel (89%)
15	 I was taught things I could do to help me feel OK with what would happen (78%)
16 17 18	Patients were asked to rank their experience of pain, fear and upset on a six-point scale from 'Not at all' to 'As much as I can imagine'. The criterion for a positive result was a rating in the two lowest categories, that is 'Not at all' or 'Just a little bit'.
19 20	 Before the procedure, 56% were either not scared or just a little bit scared, and 11% said 'As much as I can imagine'.
21	 After receiving sedation, these figures were 80% and zero respectively.
22 23	 70% reported no or little pain after sedation, and 86% no or little upset afterwards.
24	7.6.4 Other outcomes of interest
25 26 27 28	As would be expected, the degree of amnesia was dependent on the agent used; 13 of the 16 respondents who said they remembered "everything" had received Entonox, whilst of the 13 who received the benzodiazepine, five remembered "nothing" and five "just a little bit".
29 30 31	Post procedural nausea was related only to the degree of upset felt afterwards; those who felt more upset were more likely to report nausea ($p=0.019$) but the direction of causality is not clear.

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1 Only four patients said that they would not want to receive sedation again if undergoing 2 the same procedure; this was significantly related to only two variables, both ratings of 3 distress during the procedure after sedation. All four reported more than "just a little" 4 pain during the procedure (p = 0.006) and being more than "just a little bit" scared (p 5 = 0.001). 6 **Demographics: Gender** 7 Base: N = 638 9 46% 44% 10 11 10% 12 Male **Female** Not recorded 13 14 The sample had an even spread of male and female patients and covered a broad age 15 range. All except one were accompanied by a parent or carer. 16 17 Demographics: Age range of participants 18 Base: N = 6319 29% 20 13% 8% 5% 5% 3% 2% 3% 3% 3% 2% 2% 2% 21 4 5 N/R Under 6 8 9 10 11 12 13 14 15 16 & 22 over 23

1 Demographics: Ethnic origin

2 Base: N = 63

	White Mixed				Asian or Asian British				Black or Black British			Other							
	British	lrish	Other White	White & Black Caribbean	White & Black African	White & Asian	Other mixed	Indian	Pakistani	Bangladeshi	Chinese	Other Asian	Caribbean	African	Other black	Other ethnic group	Gypsy or traveller	N/R	Base
%	79	1.6	1.6	0	1.6	0	0	1.6	0	0	0	0	0	0	0	0	0	14	100
N	50	1	1	0	1	0	0	1	0	0	0	0	0	0	0	0	0	9	63

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Demographics: Percentage of children completing the survey themselves

5 Base: N = 63 96%

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22%

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Parents or carers assisted children who could not complete the questionnaire themselves, and as would be expected this was mostly the younger children. Nevertheless, the acceptability of the system was such that 22% of the under fours were able to complete the survey themselves.

15 16

Of the 23 children aged nine and over, only one child aged eleven did not complete the survey themselves.

Demographics: Range of clinical areas relating to the child or young person's procedural sedation

3 Base: N = 63

Clinical area	N	%
Medical & renal day cases	11	1 <i>7</i>
Burns 1	11	1 <i>7</i>
Radiology	10	16
Accident & Emergency	6	9.5
Cardiac inpatients	4	6.3
Burns 2	4	6.3
Oncology	2	3.2
General surgery	2	3.2
Orthopaedics	1	1.6
High Dependence Unit	1	1.6
Neuro-medical	1	1.6
General medical	1	1.6
Cardiac outpatients	1	1.6
Not recorded	8	13

Within the survey, a large number of differing clinical contexts and therefore clinical teams are represented, which is very encouraging given the participants positive experience.

Demographics: Range of medication used relating to the child or young person's procedural sedation

9 Base: N = 63

Medication used	N	%
Entonox	30	48%
Midazolam	19	30%
Oral morphine	9	14%
Chloral hydrate	1	1.6%
IV morphine	1	1.6%
Oral ketamine	1	1.6%
IV ketamine	1	1.6%
Not recorded	11	18

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Decisions made by the GDG when reviewing the initial scope and resulting clinical questions helped focus the pharmacological interventions review to what agents were in common use. The survey results reflect those discussions in that all of the above agents were systemically reviewed, with oral morphine being reviewed when used in combination. The single use of oral morphine is not advised. The absence of propofol as a single agent is noted that it was used at all in this large NHS Foundation Trust.

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Demographics: Range of clinical procedure chosen in relation to the child or young person's procedural sedation

Base: N = 63

Procedure	N	%
Change of wound dressings	14	22
Urodynamics	7	11
Intra-articular steroid injections	6	10
Other	6	10
Cannulation	4	6.3
Removal of chest drains	4	6.3
Gamma camera	3	4.8
Botox injections	2	3.2
Removal of sutures	2	3.2
Removal of wound drains	2	3.2
MRI	1	1.6
Lumbar Puncture	1	1.6
Removal of wires	1	1.6
Catheter insertion	1	1.6
Changing of line position	1	1.6
Not recorded	8	13

Survey results again helpful in relation to the types of procedure anticipated in relation to the target guideline population. The one clear obvious omission is dental treatment which the survey was not able to include.

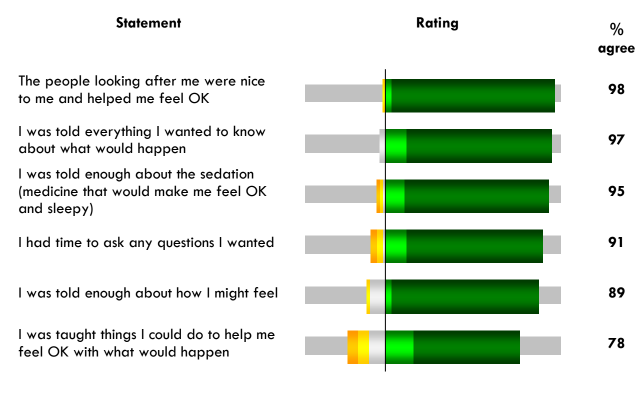
The survey results within the context of the clinically important issues are extremely useful as they by large, affirm the clinical interpretation of the evidence by the GDG in relation to targeting key clinical contexts, key clinical procedures, key clinical interventions. That said, the way children and young people are supported through the sedation experience is of perhaps the greatest interest.

1 The experience of children and young people undergoing procedural sedation:

2 Part 1: Information and support

3 Base: N = 63

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Key: Disagree a lot Disagree a bit In the middle Agree a bit Agree a lot

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The GDG sub group had carefully considered the type of questions we wanted to ask, these covered the pre procedural phase when the child or young person is being prepared (information; consent; visualisation), during procedure (amnesic effect, pain free) and the post procedure phase (amnesic effect, nausea, emotional response, preparedness for repeat intervention under sedation).

14 15 16

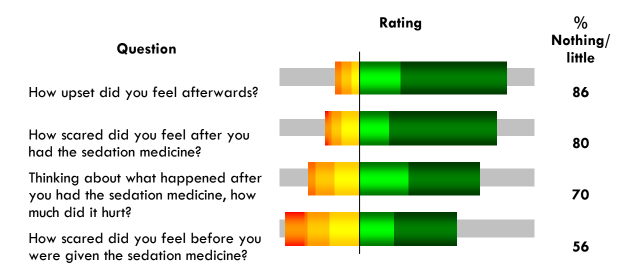
The questions were then in discussion with Priority Research who have experience in conducting this type of survey finalized to ensure they would be understood by all age ranges and that they would readily translate into the range of languages used.

1 The experience of children and young people undergoing procedural sedation:

Part 2: Emotional engagement and memory recall

3 Base: N = 63

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Key: As much as I Loads Qui	te a lot Some	Just a little bit	Not at all
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The survey results are particularly interesting in this area as they indicate that children and young people have an extremely positive experience of sedation in relation to a wide range and variety of clinical procedures and clinical settings. The responses indicate little variation in practice in this one NHS Foundation Trust, and are indicative of the benefit that clinical guidance can bring when clinical and patient pathways are followed to plan and prepare the patient and ensure their experience is a positive.

The results as seen indicate that much of this is bearing this positive outcome

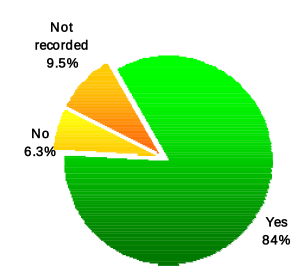
- 1 The experience of children and young people undergoing procedural sedation:
- 2 Part 1 and 2 survey detail in relation to responses and percentage breakdown
- 3 Base: N = 63

	Percentages						Further details				
	Disagree a lot	Disagree a bit	In the middle	Agree a bit	Agree a lot	Base	N/R%	Response	Total base		
I was told everything I wanted to know about what would happen	0	0	3.5	12	84	57	6.6	93	63		
I was told enough about the sedation	1.8	1.8	1.8	11	84	55	9.8	90	63		
I was told enough about how I might feel	0	1.8	9.1	3.6	85	55	9.8	90	63		
I was taught things I could do to help me feel OK with what would happen	6	6	10	16	62	50	18	82	63		
I had time to ask any questions I wanted	3.6	3.6	1.8	13	79	56	8.2	92	63		
The people looking after me were nice to me and helped me feel OK	0	1.8	0	3.6	95	56	8.2	92	63		

	Percentages							Furth	er details	
	As much as I can imagine	Loads	Quite a lot	Some	Just a little bit	Not at all	Base	N/R%	Response	Total base
How upset did you feel afterwards?	0	3.6	5.5	5.5	24	62	55	13	87	63
How scared did you feel after you had the sedation medicine?	1.9	1.9	5.6	11	17	63	54	14	86	63
Thinking about what happened after you had the sedation medicine, how much did it hurt?	0	3.8	11	15	28	42	53	16	84	63
How scared did you feel before you were given the sedation medicine?	11	1.8	13	18	20	36	55	13	87	63

1 2 The experience of children and young people undergoing procedural sedation: 3 **Part 3 Outcomes** 4 Amnesic effect of sedation in relation to memory of procedure 5 6 7 31% 8 24% 20% 9 16% 10 7.8% 11 2.0% 12 **Nothing** Just a little bit Some Quite a lot Loads **Everything** 13 Did you feel sick after the procedure? 14 15 Νo 16 **75%** 17 18 Yes 14% 19 20 21 Not recorded 22 11% 23 24

Would you want sedation again if you had to have more treatment?



The above outcomes were related to other variables. As would be expected, the degree of amnesia was dependent on the agent used; 13 of the 16 respondents who said they remembered "everything" had received Entonox, whilst of the 13 who received the benzodiazepine, five remembered "nothing" and five "just a little bit".

Post procedural nausea was related only to the degree of upset felt afterwards; those who felt more upset were more likely to report nausea (p = 0.019) but the direction of causality is not clear.

Only four patients said that they would not want to receive sedation again if undergoing the same procedure; this was significantly related to only two variables, both ratings of distress during the procedure after sedation. All four reported more than "just a little" pain during the procedure (p = 0.006) and being more than "just a little bit" scared (p = 0.001).

The Questionnaire Content (respondent) and Data Summary can be found in appendixes J and K respectively.

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