1 Appendices: Evidence Tables – Diagnosis of active TB RQ B

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1.1 RQB: What is the most effective method of collecting respiratory samples from children unable to expectorate spontaneously?

1.1.1 Abadco and Steiner, 1992

Bibliographic reference	Abadco DL and Steiner P (1992) Gastric lavage is better than bronchoalveolar lavage for isolation of Mycobacterium tuberculosis in childhood pulmonary tuberculosis. <i>Pediatric Infectious Disease Journal</i> 11(9): 735-8
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? unclear
	Participants blinded? unclear
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes

Bibliographic reference	Abadco DL and Steiner P (1992) Gastric lavage is better than bronchoalveolar lavage for isolation of Mycobacterium tuberculosis in childhood pulmonary tuberculosis. <i>Pediatric Infectious Disease Journal</i> 11(9): 735-8
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 20
Patient characteristics	Inclusion Admitted to the Children's Medical Centre of Brooklyn for gastric lavage and bronchoalveolar lavage Characteristics of included participants 19 children were diagnosed with pulmonary tuberculosis based on a positive tuberculin skin test (induration ≥10 mm) and an abnormal chest roentogram 1 patient with AIDS had an abnormal chest roentogram but a negative tuberculin skin test, but was included in the study because of a history of exposure to active tuberculosis 10 males, 10 females Ages ranged from 4 months to 7.5 years (median = 24 months) 16 participants were <5 years of age, 4 participants were >5 years of age 4 participants were asymptomatic, 16 had cough and/or fever
Intervention	Nasogastric lavage 3 specimens collected on 3 consecutive mornings after an overnight fast Depending on the age of the participant, the child was either allowed to drink 30 to 60 ml of sterile water or it was administered through a nasogastric tube The gastric contents were aspirated through a nasogastric tube and immediately sent for fluorescence microscopy and culture on Löwenstein-Jensen and Middlebrook 7H11 media
Comparator	Bronchoalveolar lavage Single specimen collected on the same day as the gastric lavage Patients received meperidine (2 mg/kg), promethazine (1 mg/kg) and chlorpromazine (1 mg/kg) intramuscularly 30 to 60 minutes before the procedure All patients received supplemental oxygen

Bibliographic reference	Abadco DL and Steiner P (1992) Gastric lavage is better than bronchoalveolar lavage for isolation of Mycobacterium tuberculosis in childhood pulmonary tuberculosis. <i>Pediatric Infectious Disease Journal</i> 11(9): 735-8
	Topical 2% lidocaine was applied to the nose and larynx as needed The flexible bronchoscope was inserted transnasally, advanced into the trachea and wedged into the most involved area as seen on the chest roentogram or into a subsegment of the right middle lobe if the infiltrate was diffuse After wedging, 5 to 10 ml of sterile nonbacteriostatic 0.9% sodium chloride solution was instilled through the suction channel and subsequently aspirated The specimen was immediately sent for fluorescence microscopy and culture on Löwenstein-Jensen and Middlebrook 7H11 media
Location	New York, US
Outcomes measures and effect size	Smear positivity (number positive participants; note: all participants were considered to have tuberculosis) • nasogastric lavage = 0/20 • bronchoalveolar lavage = 0/20 • nasogastric lavage plus bronchoalveolar lavage = 0/20
	Volume of single specimen (mean (range), ml) • nasogastric lavage = 35 (20–55) • bronchoalveolar lavage = 56.5 (45 to 80)
	Need for topical anaesthesia • nasogastric lavage = none documented • bronchoalveolar lavage = 2 participants, each requiring no more than 60 mg
Source of funding	No details provided
Comments	Same population as Chan (1994); duplicate outcomes (culture positivity) extracted from Chan (2004)

1.1.2 Al-Aghbari, 2009

Bibliographic reference	Al-Aghbari N, Al-Sonboli N, Yassin MA, Coulter JB, Atef Z, Al-Eryani A and Cuevas LE (2009) Multiple sampling in one day to optimize smear microscopy in children with tuberculosis in Yemen. <i>PLoS One</i> 4(4): e5140
Study type	Cross-sectional
Study quality	Study limitations Was a consecutive or random sample of patients enrolled? unclear

Bibliographic reference	Al-Aghbari N, Al-Sonboli N, Yassin MA, Coulter JB, Atef Z, Al-Eryani A and Cuevas LE (2009) Multiple sampling in one day to optimize smear microscopy in children with tuberculosis in Yemen. <i>PLoS One</i> 4(4): e5140
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? no, not all collection techniques applied to all participants
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? microscopists blinded; other investigators unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? unclear
	Inconsistency
	Groups comparable at baseline? unclear
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? unclear
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes

Bibliographic reference	Al-Aghbari N, Al-Sonboli N, Yassin MA, Coulter JB, Atef Z, Al-Eryani A and Cuevas LE (2009) Multiple sampling in one day to optimize smear microscopy in children with tuberculosis in Yemen. <i>PLoS One</i> 4(4): e5140
Number of patients	Included = 213 participants • nasopharyngeal aspirate = 197 participants • gastric aspirate = 196 participants • induced sputum = 88 participants
Patient characteristics	Inclusion Children with suspected tuberculosis, as defined using the following criteria: • a history of contact with cases of pulmonary tuberculosis • children not regaining normal health after measles or whooping cough • unexplained weight loss • the presence of cough and wheeze not responding to antibiotic therapy • X-ray findings suggestive of pulmonary tuberculosis Characteristics of included participants Age ranged from 2 months to 15 years, with a median of 5 years and 42 (20%) were <2 years old The most frequent clinical symptoms at presentation were cough (195, 92%), unexplained fever (179, 84%), anorexia (142, 67%), weight loss (125, 59%) and difficult breathing (82, 38%)
Intervention	Nasopharyngeal aspirate 1 specimen collected by direct aspiration via a mucus trap connected to a suction device on the first day, without induction of cough or instillation of saline solutions After the preparation of direct smears, specimens were stained using the hot Ziehl Neelsen method; all smears were read and graded by trained microscopists who were unaware of the grading of the previous specimens All specimens were cultured using Ogawa culture media Nasogastric aspiration 3 specimens collected Specimens were obtained at 6:00 am on three consecutive days by introducing a nasogastric tube and aspiration of the gastric content with a syringe After the preparation of direct smears, specimens were stained using the hot Ziehl Neelsen method; all smears were read and graded by trained microscopists who were unaware of the grading of the previous specimens All specimens were cultured using Ogawa culture media
Comparator	Induced sputum 3 specimens collected using inhaled salbutamol via a metered dose inhaler attached to oxygen at a flow rate of 5 liters per

Bibliographic reference	Al-Aghbari N, Al-Sonboli N, Yassin MA, Coulter JB, Atef Z, Al-Eryani A and Cuevas LE (2009) Multiple sampling in one day to optimize smear microscopy in children with tuberculosis in Yemen. <i>PLoS One</i> 4(4): e5140
	minute on 5 ml of 5% sterile saline for 15 minutes, followed by physiotherapy (chest percussion, vibration and active cycle breathing) and sputum collection by expectoration or from the naso/orophrarynx using a mucus extractor in those unable to expectorate After the preparation of direct smears, specimens were stained using the hot Ziehl Neelsen method; all smears were read and graded by trained microscopists who were unaware of the grading of the previous specimens All specimens were cultured using Ogawa culture media
Location	Sana'a, Yemen
Outcomes measures and effect size	Culture positivity (number positive/total number of specimens) • nasopharyngeal aspirate = 14/200 • nasogastric aspirate = 49/564 • induced sputum = 31/216
	Smear positivity (number positive/total number of specimens) • nasopharyngeal aspirate = 10/200 • nasogastric aspirate = 17/564 • induced sputum = 9/216
Source of funding	Dr Al-Aghbari received a study scholarship from the Special Programme for Research in Tropical Diseases of the World Health Organization The funder had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript
Comments	

1.1.3 **Bhandari**, 1971

Bibliographic reference	Bhandari B, Singh SV and Sharma VK (1971) Bacteriological diagnosis of pulmonary tuberculosis. A comparative study of gastric wash, laryngeal swab and lung puncture. <i>Indian Journal of Pediatrics</i> 38(284): 349-53
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? <i>unclear</i>

Bibliographic reference	Bhandari B, Singh SV and Sharma VK (1971) Bacteriological diagnosis of pulmonary tuberculosis. A comparative study of gastric wash, laryngeal swab and lung puncture. <i>Indian Journal of Pediatrics</i> 38(284): 349-53
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? unclear
	Participants blinded? unclear
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 30

	Bhandari B, Singh SV and Sharma VK (1971) Bacteriological diagnosis of pulmonary tuberculosis. A
Bibliographic reference	comparative study of gastric wash, laryngeal swab and lung puncture. <i>Indian Journal of Pediatrics</i> 38(284): 349-53
Patient characteristics	Inclusion
	Children with suspected pulmonary tuberculosis who had not received previous antituberculosis treatment
	Characteristics of included participants
	Age range: 4 months to 13 years
	≤5 years = 17
	12 females
	Positive tuberculin skin test $(n = 19) = 9$
	Erythrocyte sedimentation rate (mean±SD (range), mm at 1 st hour) = 32±19 (5–61)
Intervention	Laryngeal swab
	3 specimens collected on 3 consecutive days
	A sterilised cotton swab was moistened with distilled water and the swab collected with the aid of a laryngoscope
	Specimens were examined by Gram and Ziehl-Neelsen microscopy, and cultured on Löwenstein-Jensen medium for 6 to 10 weeks
	Lung puncture aspiration
	1 specimen collected from every patient
	The site of the pulmonary lesion was identified clinically and radiologically
	Skin preparation was achieved with 2% iodine, applied for 2 minutes, and removed with 95% ethanol
	The thorax was entered with a 4 cm needle and continuous suction applied with a 10 ml syringe
	Specimens were examined by Gram and Ziehl-Neelsen microscopy, and cultured on Löwenstein-Jensen medium for 6 to 10 weeks

Bibliographic reference	Bhandari B, Singh SV and Sharma VK (1971) Bacteriological diagnosis of pulmonary tuberculosis. A comparative study of gastric wash, laryngeal swab and lung puncture. <i>Indian Journal of Pediatrics</i> 38(284): 349-53
Comparator	Gastric lavage
	3 specimens collected early in the morning on 3 consecutive days after an overnight fast
	A stomach tube was passed and the contents aspirated
	30 to 50 ml of distilled water was pushed through the tube and the stomach was, again, aspirated
	Specimens were examined by Gram and Ziehl-Neelsen microscopy, and cultured on Löwenstein-Jensen medium for 6 to 10 weeks
Location	Udaipur, India
Outcomes measures and	Culture positivity (number of participants with a positive result)
effect size	• laryngeal swab (3 specimens) = 4/30
	• lung puncture aspiration (1 specimen) = 16/30
	• gastric lavage (3 specimens) = 3/30
	≤5 years
	• laryngeal swab (3 specimens) = 3/17
	• lung puncture aspiration (1 specimen) = 10/17
	• gastric lavage (3 specimens) = 3/17
	>5 years
	• laryngeal swab (3 specimens) = 1/13
	• lung puncture aspiration (1 specimen) = 6/13
	• gastric lavage (3 specimens) = 0/13

Bibliographic reference	Bhandari B, Singh SV and Sharma VK (1971) Bacteriological diagnosis of pulmonary tuberculosis. A comparative study of gastric wash, laryngeal swab and lung puncture. <i>Indian Journal of Pediatrics</i> 38(284): 349-53
	Smear positivity (number of participants with a positive result)
	• laryngeal swab (3 specimens) = 6/30
	• lung puncture aspiration (1 specimen) = 5/30
	• gastric lavage (3 specimens) = 4/30
	≤5 years
	• laryngeal swab (3 specimens) = 4/17
	• lung puncture aspiration (1 specimen) = 4/17
	• gastric lavage (3 specimens) = 3/17
	>5 years
	• laryngeal swab (3 specimens) = 2/13
	• lung puncture aspiration (1 specimen) = 1/13
	• gastric lavage (3 specimens) = 1/13
Source of funding	No details provided
Comments	
Abbreviations: SD, standard	d deviation

1.1.4 Buonsenso, 2014

Bibliographic reference	Buonsenso D, Barone G, Valentini P, Pierri F, Riccardi R and Chiaretti A (2014) Utility of intranasal Ketamine and Midazolam to perform gastric aspirates in children: a double-blind, placebo controlled, randomized study. <i>BMC</i>
	Pediatrics 14: 67

Bibliographic reference	Buonsenso D, Barone G, Valentini P, Pierri F, Riccardi R and Chiaretti A (2014) Utility of intranasal Ketamine and Midazolam to perform gastric aspirates in children: a double-blind, placebo controlled, randomized study. <i>BMC Pediatrics</i> 14: 67
Study type	Randomised controlled trial
Study quality	Study limitations
	Appropriate method of randomisation? yes
	Adequate allocation concealment? yes
	Participants blinded? unclear
	Individuals administering care blinded? yes
	Investigators blinded to intervention? yes
	Investigators blinded to confounding and prognostic factors? yes
	Appropriate length of follow-up? yes
	Precise definition of outcome? yes
	Valid and reliable method of outcome measurement? yes
	Intent-to-treat principle adhered to? yes
	Inconsistency
	Groups comparable at baseline? yes, although information provided relates only to age and disease status
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups equivalent for intervention completion? yes
	Groups comparable for availability of data? yes
	Indirectness

Bibliographic reference	Buonsenso D, Barone G, Valentini P, Pierri F, Riccardi R and Chiaretti A (2014) Utility of intranasal Ketamine and Midazolam to perform gastric aspirates in children: a double-blind, placebo controlled, randomized study. <i>BMC Pediatrics</i> 14: 67
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Randomised/analysed = 36
·	• nasogastric aspiration with sedation = 19
	• nasogastric aspiration with placebo = 17
	Number of procedures performed = 108
	• nasogastric aspiration with sedation = 57
	• nasogastric aspiration with placebo = 51
Patient characteristics	Inclusion
	Children <14 years old
	'Uncooperative' children undergoing gastric aspirates for suspected tuberculosis Exclusion
	ASA physical status classification of III (patient with severe systemic disease) or higher
	Known allergy to benzodiazepines
	Known allergy to ketamine
	Upper respiratory tract infection with nasal discharge
	Known liver disease or respiratory distress
	Characteristics of included participants
	Age (mean±SD, months)
	• nasogastric aspiration with sedation = 41.5±36.0
	• nasogastric aspiration with placebo = 40.6±34.8
	Final diagnosis
	 nasogastric aspiration with sedation: pulmonary tuberculosis = 15; latent tuberculosis = 1; non-tuberculous pneumonia = 3 nasogastric aspiration with placebo: pulmonary tuberculosis = 17
	The two groups did not differ significantly with respect to weight and ethnicity
Intervention	Nasogastric aspiration with intranasal sedation
	Intranasal sedation using 2 mg/kg of ketamine hydrochloride in both nostrils, followed by 0.5 mg/kg (maximum dose 10 mg)
	of midazolam using a mucosal atomizer device

Bibliographic reference		ini P, Pierri F, Riccardi R and Chiaretti A (20 spirates in children: a double-blind, placebo	
	A nasogastric tube was inserted in topical anaesthetics on nasogastic volume of gastric aspirates was au aspirating back	on three consecutive days early in the morning to the child's stomach after intranasal administration and then aspirated its contents; in case agmented as needed by injecting in the stomaches within 60 minutes from intranasal administration.	ration (without any additional drugs nor of unsuccessful or poor aspiration, the h 5 mL of saline solution (sterile water) and
Comparator	Nasogastric aspiration with placebo Intranasal administration of normal saline solution (the same volume the child would have received if in the sedation-group) in each nostril (twice, in order to pretend the two different drugs of the sedation-group) using a mucosal atomizer device Gastric washings were performed on three consecutive days early in the morning and after an overnight fasting A nasogastric tube was inserted into the child's stomach after intranasal administration (without any additional drugs nor topical anaesthetics on nasogastic tube) and then aspirated its contents; in case of unsuccessful or poor aspiration, the volume of gastric aspirates was augmented as needed by injecting in the stomach 5 mL of saline solution (sterile water) and aspirating back The procedure began in every case within 60 minutes from intranasal administration		
Location	Rome, Italy		
Outcomes measures and	Acceptability of the procedure to parents ^c		
effect size		Median score (range) provided by parents of those receiving sedation ($n = 19$)	Median score (range) provided by parents of those receiving placebo ($n = 19$)
	Did the sedation help?	10 (10–10)	5 (3–7)
	Level of child's outlook	8.9 (7–10)	5.8 (5–7)
	Level of parents' outlook	9.1 (8–10)	4.9 (3–7)
	Level of child's tolerance of procedures	8.7 (7–10)	8.5 (7–10)
	Would recommend to other parents	9.3 (9–10)	4 (3–6)
	Would like to see the mucosal atomizer device used routinely	9.8 (9–10)	4 (3–6)
	Acceptability of the procedure to cl	linicians ^c	
		Median score (range) provided by parents of those receiving sedation	Median score (range) provided by parents of those receiving placebo

Bibliographic reference		Buonsenso D, Barone G, Valentini P, Pierri F, Riccardi R and Chiaretti A (2014) Utility of intranasal Ketamine and Midazolam to perform gastric aspirates in children: a double-blind, placebo controlled, randomized study. <i>BMC Pediatrics</i> 14: 67	
	Did the sedation help?	10 (10–10)	3 (2–4)
	Level of child's outlook	8 (7–9)	3 (2–4)
	Level of clinicians' outlook	9.5 (9–10)	4 (3–5)
	Level of child's tolerance of procedures	8.2 (7–9)	8 (7–9)
	Would recommend to other clinicians	9.4 (9–10)	3 (1–5)
	Would like to see the mucosal atomizer device used routinely	10 (10–10)	3 (1–5)
	Made the procedure more acceptable	10 (10–10)	3 (1–5)
	Adverse events - transitory postsedation a Number of procedures after which transito • nasogastric aspiration with sedation (n = • nasogastric aspiration with placebo (n =	ory postsedation agitation 57 procedures) = 6	was experienced
Source of funding	No funding have been received by the aut	hors	
Comments			
· ·	• nasogastric aspiration with placebo (n =	51 procedures) = 0	

^a Pre-sedation behaviour was assessed on a 4-point scale (1 = calm, cooperative; 2 = anxious but reassurable; 3 = anxious and not reassurable; 4 = crying or resisting) by an anesthesiologist blinded to the group of the child; children were included if they were <14 years old and had a pre-sedation behaviour ≥3

Abbreviations: SD, standard deviation

1.1.5 Cakir, 2008

Bibliographic reference	Cakir E, Uyan ZS, Oktem S, Karakoc F, Ersu R, Karadag B and Dagli E (2008) Flexible bronchoscopy for diagnosis and follow up of childhood endobronchial tuberculosis. <i>Pediatric Infectious Disease Journal</i> 27(9): 783-7
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear

^b Modified Objective Pain Score; score ranges from 0 to 10 (the higher the score, the greater the pain experienced for the child)

^c Derived from a series of questions, answered using a visual analogue scale ('0' for worst, '10' for best)

Bibliographic reference	Cakir E, Uyan ZS, Oktem S, Karakoc F, Ersu R, Karadag B and Dagli E (2008) Flexible bronchoscopy for diagnosis and follow up of childhood endobronchial tuberculosis. <i>Pediatric Infectious Disease Journal</i> 27(9): 783-7
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? yes, although details provided were limited
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? unclear
	Inconsistency
	Groups comparable at baseline? unclear
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 70
Patient characteristics	Inclusion Children

Bibliographic reference	Cakir E, Uyan ZS, Oktem S, Karakoc F, Ersu R, Karadag B and Dagli E (2008) Flexible bronchoscopy for diagnosis and follow up of childhood endobronchial tuberculosis. <i>Pediatric Infectious Disease Journal</i> 27(9): 783-7
	Tuberculosis patients with an inadequate response to antituberculosis treatment, defined as unresolved or progressive clinical and radiographic findings despite 8 weeks of antituberculosis treatment Patients with suspected tuberculosis, defined as children with respiratory symptoms and chest radiograph findings suspicious of tuberculosis, with or without positive tuberculin skin test and history of a household contact with tuberculosis Characteristics of included participants Age (median [interquartile range], months) = 81.5 [13.7–112.5] Gender • male = 38 • female = 32 Symptoms • cough = 62 • sputum = 24 • fever = 13 • dyspnea = 11 Duration of symptoms (mean±SD) = 3.54±3.05 Household contact with active tuberculosis = 31 Positive tuberculin skin test = 38 High erythrocyte sedimentation rate = 34 Tracheobronchial involvement = 33 Mycobacterium tuberculosis isolation = 14
Intervention	Nasogastric aspiration All cases were hospitalized Specimens collected on 3 consecutive days All samples were cultured in Löwenstein-Jensen medium
Comparator	Bronchoalveolar lavage All cases were hospitalized Patients received midazolam and pethidine HCL as premedication and lidocaine was used as topical anesthetic All samples were cultured in Löwenstein-Jensen medium
Location	Istanbul, Turkey
Outcomes measures and effect size	Culture positivity (number positive/total number of participants) • nasogastric aspiration = 5/70

Bibliographic reference	Cakir E, Uyan ZS, Oktem S, Karakoc F, Ersu R, Karadag B and Dagli E (2008) Flexible bronchoscopy for diagnosis and follow up of childhood endobronchial tuberculosis. <i>Pediatric Infectious Disease Journal</i> 27(9): 783-7
	• bronchoalveolar lavage = 7/70
Source of funding	No details provided
Comments	
Abbreviations: SD, standard deviation	

1.1.6 Cakir, 2013

Bibliographic reference	Cakir E, Kut A, Ozkaya E, Gedik AH, Midyat L and Nursoy M (2013) Bronchoscopic evaluation in childhood pulmonary tuberculosis: risk factors of airway involvement and contribution to the bacteriologic diagnosis. <i>Pediatric Infectious Disease Journal</i> 32(8): 921-3
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? unclear
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? unclear
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? unclear
	Inconsistency

Bibliographic reference	Cakir E, Kut A, Ozkaya E, Gedik AH, Midyat L and Nursoy M (2013) Bronchoscopic evaluation in childhood pulmonary tuberculosis: risk factors of airway involvement and contribution to the bacteriologic diagnosis. <i>Pediatric Infectious Disease Journal</i> 32(8): 921-3
	Groups comparable at baseline? yes
	Groups received the same care apart from the intervention(s) studied? yes, although details provided were limited
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 167 participants
Patient characteristics	Inclusion Children with suspected tuberculosis, in accordance with the World Health Organization's TB Standards case definition Characteristics of included participants Age (mean±SD (range), months) = 97.2±48.5 (2 months to 16 years)
	Male = 54% Contact with a known adult index case = 78% Positive tuberculin skin test = 67%
	Primary tuberculosis = 82% Signs and symptoms
	cough = 76%fever = 26%
	weight loss = 20%haemoptysis = 12%
	• wheezing = 5%

Bibliographic reference	Cakir E, Kut A, Ozkaya E, Gedik AH, Midyat L and Nursoy M (2013) Bronchoscopic evaluation in childhood pulmonary tuberculosis: risk factors of airway involvement and contribution to the bacteriologic diagnosis. <i>Pediatric Infectious Disease Journal</i> 32(8): 921-3
	 pulmonary consolidation = 61% atelectasis = 19% cavity = 11% disseminated infiltration = 7% There was no statistical difference between these 2 groups for demographic features, symptoms and radiologic findings
Intervention	Nasogastric aspiration Early morning specimens collected on 3 consecutive days All samples were cultured in Löwenstein-Jensen medium
Comparator	Bronchoalveolar lavage All samples were cultured in Löwenstein-Jensen medium
Location	Istanbul, Turkey
Outcomes measures and effect size	Culture positivity (number positive/total number of patients (%)) • nasogastric aspiration = 54/167 • bronchoalveolar lavage = 48/167 • nasogastric aspiration plus bronchoalveolar lavage = 70/167
Source of funding	No details provided
Comments	
Abbreviations: SD, standard	deviation

1.1.7 Chan, 1994

Bibliographic reference	Chan S, Abadco DL and Steiner P (1994) Role of flexible fiberoptic bronchoscopy in the diagnosis of childhood endobronchial tuberculosis. <i>Pediatric Infectious Disease Journal</i> 13(6): 506-9
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? yes
	Was a case-control design avoided? yes

Bibliographic reference	Chan S, Abadco DL and Steiner P (1994) Role of flexible fiberoptic bronchoscopy in the diagnosis of childhood endobronchial tuberculosis. <i>Pediatric Infectious Disease Journal</i> 13(6): 506-9
	Did the study avoid inappropriate exclusions? unclear
	Participants blinded? unclear
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	n = 36
Patient characteristics	Inclusion Children under 16 years of age admitted to the Children's Medical Centre of Brooklyn for gastric lavage and bronchoalveolar lavage

Bibliographic reference	Chan S, Abadco DL and Steiner P (1994) Role of flexible fiberoptic bronchoscopy in the diagnosis of childhood endobronchial tuberculosis. <i>Pediatric Infectious Disease Journal</i> 13(6): 506-9
	Characteristics of included participants 19 males, 17 females Age (median (range)) = 3.9 years (4 months – 16 years) Symptoms on presentation included fever (n = 20), cough (n = 16) and wheezing (n = 2); 12 patients were asymptomatic
	A history of close contact with tuberculosis in an adult with active tuberculosis was elicited in 12 patients
Intervention	Nasogastric lavage 3 specimens collected on 3 consecutive mornings after an overnight fast Depending on the age of the participant, the child was either allowed to drink 30 to 60 ml of sterile water or it was administered through a nasogastric tube The gastric contents were aspirated through a nasogastric tube and immediately sent for fluorescence microscopy and culture on Löwenstein-Jensen and Middlebrook 7H11 media
Comparator	Bronchoalveolar lavage Single specimen collected on the same day as the gastric lavage Patients received meperidine (2 mg/kg), promethazine (1 mg/kg) and chlorpromazine (1 mg/kg) intramuscularly 30 to 60 minutes before the procedure All patients received supplemental oxygen Topical 2% lidocaine was applied to the nose and larynx as needed The flexible bronchoscope was inserted transnasally, advanced into the trachea and wedged into the most involved area as seen on the chest roentogram or into a subsegment of the right middle lobe if the infiltrate was diffuse After wedging, 5 to 10 ml of sterile nonbacteriostatic 0.9% sodium chloride solution was instilled through the suction channel and subsequently aspirated The specimen was immediately sent for fluorescence microscopy and culture on Löwenstein-Jensen and Middlebrook 7H11 media
Location	New York, US
Outcomes measures and effect size	Culture positivity (number positive participants; note: all participants were considered to have tuberculosis) • nasogastric lavage = 17/36 • bronchoalveolar lavage = 4/36
Source of funding	No details provided

Bibliographic reference	Chan S, Abadco DL and Steiner P (1994) Role of flexible fiberoptic bronchoscopy in the diagnosis of childhood endobronchial tuberculosis. <i>Pediatric Infectious Disease Journal</i> 13(6): 506-9
Comments	

1.1.8 **Hatherill**, 2009

Bibliographic reference	Hatherill M, Hawkridge T, Zar HJ, Whitelaw A, Tameris M, Workman L, Geiter L, Hanekom WA and Hussey G (2009) Induced sputum or gastric lavage for community-based diagnosis of childhood pulmonary tuberculosis? <i>Archives of Disease in Childhood</i> 94(3): 195-201
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? yes, though details provided were limited
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? unclear
	Inconsistency
	Groups comparable at baseline? unclear
	Groups received the same care apart from the intervention(s) studied? yes

Bibliographic reference	Hatherill M, Hawkridge T, Zar HJ, Whitelaw A, Tameris M, Workman L, Geiter L, Hanekom WA and Hussey G (2009) Induced sputum or gastric lavage for community-based diagnosis of childhood pulmonary tuberculosis? <i>Archives of Disease in Childhood</i> 94(3): 195-201
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 1936 specimens
Patient characteristics	Inclusion Children with a tuberculosis contact or compatible symptoms – such as unexplained cough, loss of weight, or failure to thrive – being investigated for suspected pulmonary tuberculosis Characteristics of included participants Age (median [interquartile range], months) = 13 [7–20] History of household tuberculosis exposure = 892 (48%) Signs and symptoms: • failure to thrive = 1045 (56%) • weight loss = 415 (22%) • history of cough of any duration = 1064 (57%), including 743 (40%) with cough for >2 weeks • history of fever = 715 cases (38%) • chest radiographs judged compatible with a diagnosis of pulmonary tuberculosis = 369 (20%) • positive tuberculin skin test = 676 (36%) Positive HIV ELISA = 60 (3.2%), of which 32 (1.7%) were confirmed HIV-infected by PCR
Intervention	Nasogastric lavage Early morning specimens collected on 2 consecutive days after overnight fast Uses 10 ml 0.9% saline and aspiration of 5–10 ml lavage fluid via a nasogastric tube; sputum induction was performed 3–4 hours later Culture was performed using the BACTEC MGIT 960

Bibliographic reference	Hatherill M, Hawkridge T, Zar HJ, Whitelaw A, Tameris M, Workman L, Geiter L, Hanekom WA and Hussey G (2009) Induced sputum or gastric lavage for community-based diagnosis of childhood pulmonary tuberculosis? <i>Archives of Disease in Childhood</i> 94(3): 195-201								
Comparator	Induced sputum								
	Specimens collected on 2 consecutive days								
	saturation, preceded	by administra livered by jet r	ation of 200 pation a	at an oxygen flow of 5 lit	metered do	se inhaler an	ng of pulse and oxygen Id spacer; hypertonic 5% Iwas collected by suctioning		
Location	South Africa								
Outcomes measures and effect size	 Culture positivity (nu nasogastric lavage induced sputum = 1 K statistic = 0.31 Comparative and cu 	= 127/1869 108/1869		` <i>'</i>	llection met	hod/total num	ober of culture positive cases)		
	Comparative and cumulative yields (number cases positive by each collection method/total number of culture positive cases) Culture positive Yield (%) (95% CI)								
	Induced sputum	Canal o p		11010 (70) (0070 01)					
	day 1	73		38 (31 to 45)					
	day 2	51		27 (20 to 33)					
	days 1 and 2	106		55 (48 to 62)					
	Gastric lavage			<u> </u>					
	day 1	80		42 (35 to 49)					
	day 2	75		39 (32 to 46)					
	days 1 and 2	125		66 (59 to 73)					
	Differences in yield between various comparisons of single, cumulative and combined yields among case episodes with a positive <i>M. tuberculosis</i> culture (n=191)								
	Sample type	Number culture positive	Yield (%)	Sample type	Number culture positive	Yield (%)	Difference in yield (%) (95% CI)		
	Single induced sputum (day 1)	73	38	Single gastric lavage (day 1)	80	42	-4 (-15 to 7)		
	Single induced	73	38	Cumulative induced	106	55	-17 (-23 to -11)		

Bibliographic reference		stric lava	age for com				om WA and Hussey G (2009 nary tuberculosis? <i>Archive</i>
	sputum (day 1)	. ,		sputum (days 1 + 2)			
	Single induced sputum (day 1)	73	38	Cumulative gastric lavage (days 1 + 2)	126	66	-28 (-40 to -15)
	Single gastric lavage (day 1)	80	42	Cumulative gastric lavage (days 1 + 2)	126	66	-24 (-31 to -17)
	Single gastric lavage (day 1)	80	42	Cumulative induced sputum (days 1 + 2)	106	55	-13 (-25 to -2)
	Cumulative gastric lavage (days 1 + 2)	126	66	Cumulative induced sputum (days 1 + 2)	106	55	11 (-3 to 24)
	Cumulative gastric lavage (days 1 + 2)	126	66	Combined induced sputum and gastric lavage (day 1)	128	67	-1 (-11 to 9)
	Cumulative induced sputum (days 1 + 2)	106	55	Combined induced sputum and gastric lavage (day 1)	128	67	-12 (-21 to -2)
Source of funding							or Health, Immunopaedia, the Countries Trials Partnership
Comments							
Abbreviations: CI, confidence	interval						

1.1.9 <u>Jiménez, 2013</u>

Bibliographic reference	Jiménez MR, Guillén Martín S, Prieto Tato LM, Cacho Calvo JB, Alvarez García A, Soto Sánchez B and Ramos Amador JT (2013) Induced sputum versus gastric lavage for the diagnosis of pulmonary tuberculosis in children. <i>BMC Infectious Diseases</i> 13(1): 222
Study type	Cross-sectional
Study quality	Study limitations

Bibliographic reference	Jiménez MR, Guillén Martín S, Prieto Tato LM, Cacho Calvo JB, Alvarez García A, Soto Sánchez B and Ramos Amador JT (2013) Induced sputum versus gastric lavage for the diagnosis of pulmonary tuberculosis in children. <i>BMC Infectious Diseases</i> 13(1): 222
	Was a consecutive or random sample of patients enrolled? yes
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? no, excluded participants positive for non-tuberculous mycobacteria
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes

Bibliographic reference	Jiménez MR, Guillén Martín S, Prieto Tato LM, Cacho Calvo JB, Alvarez García A, Soto Sánchez B and Ramos Amador JT (2013) Induced sputum versus gastric lavage for the diagnosis of pulmonary tuberculosis in children. <i>BMC Infectious Diseases</i> 13(1): 222
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 22 participants
	1 participant was excluded for not having provided the collection of samples according to the protocol
	4 patients were excluded because the final diagnosis was not PTB and other microorganisms were identified
	Data available = 17 participants
Patient characteristics	Inclusion
	Children (<15 years old) with suspected pulmonary tuberculosis based on clinical features or radiology
	No evidence of immunosuppression
	Characteristics of included participants
	11 patients (53%) were females
	Median age was 72 months (range 1 month to 14 years of age)
	7 (33%) were ≤5 years of age
	17 patients (80%) were clinically diagnosed of pulmonary tuberculosis, based on a positive tuberculin skin test (Mantoux reaction was >10 mm) and radiological criteria
	An HIV-test (enzyme-linked immunoassay) was performed on only one patient, with negative results
Intervention	Gastric lavage
	3 specimens collected on consecutive days
	Performed early morning on all children, after an overnight fast
	A nasogastric tube was passed and normal saline 20 ml inserted, left for 3 minutes and then aspirated

Bibliographic reference	Jiménez MR, Guillén Martín S, Prieto Tato LM, Cacho Calvo JB, Alvarez García A, Soto Sánchez B and Ramos Amador JT (2013) Induced sputum versus gastric lavage for the diagnosis of pulmonary tuberculosis in children. <i>BMC Infectious Diseases</i> 13(1): 222
	An additional 5–10 ml of normal saline was introduced and aspirated, until a minimum of 20 ml of aspirate was obtained
	Specimens were examined by fluorescence microscopy, nucleic acid amplification tests (COBAS TaqMan, GenoType MTBDRplus assay and Xpert® MTB/RIF introduced step by step into the laboratory procedures), and cultured using Coletsos medium and the BACTEC MGIT 960
Comparator	Induced sputum
	3 specimens collected on consecutive days
	Performed 4 hours after gastric lavage, prior to lunch
	To prevent the risk of bronchospasm induced by hypertonic saline, children were pre-treated with nebulized salbutamol (0.03 ml/kg, maximum 1 ml (1 ml = 5 mg)); subsequently, 5 ml of 5% sterile saline at a flow rate of 5 l per minute was nebulized for 15 minutes, followed by chest percussion on the front and back chest wall
	After this procedure, if spontaneous expectoration was not achieved, sputum was obtained by suctioning through the nasopharynx with a sterile mucus extractor
	Specimens were examined by fluorescence microscopy, NAATs (COBAS TaqMan, GenoType MTBDRplus assay and Xpert® MTB/RIF introduced step by step into the laboratory procedures), and cultured using Coletsos medium and the BACTEC MGIT 960
Location	Madrid, Spain
Outcomes measures and	Culture positivity (number positive/total number participants)
effect size	Cumulative yield for all 3 specimens (days 1 to 3)
	• gastric lavage = 8/17
	• induced sputum = 7/17
	• gastric lavage plus induced sputum = 10/17

Bibliographic reference	Jiménez MR, Guillén Martín S, Prieto Tato LM, Cacho Calvo JB, Alvarez García A, Soto Sánchez B and Ramos Amador JT (2013) Induced sputum versus gastric lavage for the diagnosis of pulmonary tuberculosis in children. <i>BMC Infectious Diseases</i> 13(1): 222
	Cumulative yield for first 2 specimens (days 1 and 2)
	• gastric lavage = 7/17
	• induced sputum = 6/17
	• gastric lavage plus induced sputum = 9/17
	Yield for first specimen (day 1)
	• gastric lavage = 7/17
	• induced sputum = 5/17
	• gastric lavage plus induced sputum = 8/17
	Smear positivity (number positive/total number participants)
	Cumulative yield for all 3 specimens (days 1 to 3)
	• gastric lavage = 1/17
	• induced sputum = 2/17
	NAAT positivity (number positive/total number participants)
	Cumulative yield for all 3 specimens (days 1 to 3)
	• gastric lavage = 2/17
	• induced sputum = 3/17
	Adverse effects – only reported for induced sputum
	Of the 16 patients that showed potential adverse effects to sputum induction (a total of 48 procedures), no serious adverse reactions occurred during or after the procedure

Bibliographic reference	Jiménez MR, Guillén Martín S, Prieto Tato LM, Cacho Calvo JB, Alvarez García A, Soto Sánchez B and Ramos Amador JT (2013) Induced sputum versus gastric lavage for the diagnosis of pulmonary tuberculosis in children. <i>BMC Infectious Diseases</i> 13(1): 222
	The most common adverse events were mild epistaxis in 8 procedures (16.6%), nausea in 3 (6.25%) and increased coughing in 3 (6.25%)
	Only 1 infant had transient hypoxemia in 2 procedures (lowest oxygen saturation 87%) recovered spontaneously
	There were no episodes of bronchospasm
Source of funding	Funded in part by the Foundation for Research and Prevention of AIDS in Spain
Comments	
Abbreviations: NAATs, nucle	eic acid amplification tests

1.1.10 Lloyd, 1968

Bibliographic reference	Lloyd AV (1968) Bacteriological diagnosis of tuberculosis in children: a comparative study of gastric lavage and laryngeal swab methods. <i>East African Medical Journal</i> 45(3): 140-3
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? unclear
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes

Bibliographic reference	Lloyd AV (1968) Bacteriological diagnosis of tuberculosis in children: a comparative study of gastric lavage and laryngeal swab methods. <i>East African Medical Journal</i> 45(3): 140-3
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? unclear
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 60
Patient characteristics	Inclusion
	Children
	Characteristics of included population
	Age ranged from 5 months to 6 years
	5 had miliary tuberculosis, 6 had tuberculous meningitis, 4 had abdominal tuberculosis, 3 had spinal caries and the remainder had pulmonary lesions, either bronchopneumonia, segmental lesions or cavitation
Intervention	Laryngeal swab

Bibliographic reference	Lloyd AV (1968) Bacteriological diagnosis of tuberculosis in children: a comparative study of gastric lavage and laryngeal swab methods. <i>East African Medical Journal</i> 45(3): 140-3
	3 specimens collected, as far as possible, on 3 consecutive days
	The swabs were made of stainless steel wire
	The child was held firmly by its mother and its tongue brought forward with a tongue depressor
	The swab was dipped in sterile water before being inserted into the larynx
	If the child was old enough to understand, he was asked to cough, otherwise the tickling of the swab against the back of the throat was almost always sufficient to produce a cough
	Specimens were cultured on Löwenstein-Jensen medium
Comparator	Gastric lavage
	3 specimens collected, as far as possible, on 3 consecutive days, in the early morning before food was given
	Specimens were cultured on Löwenstein-Jensen medium
Location	Kampala, Uganda
Outcomes measures and	Culture positivity (number positive /total number of cases; note: all participants considered to have tuberculosis)
effect size	Taking into account all 3 specimens
	• gastric lavage = 17/60
	• laryngeal swab = 38/60
	Taking into account first 2 specimens
	• gastric lavage = 4/60
	• laryngeal swab = 15/60
	Taking into account first specimen
	• gastric lavage = 1/60

Bibliographic reference	Lloyd AV (1968) Bacteriological diagnosis of tuberculosis in children: a comparative study of gastric lavage and laryngeal swab methods. <i>East African Medical Journal</i> 45(3): 140-3 • laryngeal swab = 6/60
Source of funding Comments	Supported by grants from Makerere University College and the East African Medical Research Council

1.1.11 **Maciel**, 2010

Bibliographic reference	Maciel EL, Peres RL, do Prado TN, Macedo CR, Palaci M, Vinhas SA, Dietze R, Johnson JL and Struchiner CJ (2010) Saline nebulization before gastric lavage in the diagnosis of pulmonary tuberculosis in children and adolescents. Journal of Tropical Pediatrics 56(6): 458-9
Study type	Randomised controlled trial
Study quality	Study limitations
	Appropriate method of randomisation? unclear
	Adequate allocation concealment? unclear
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded to intervention? unclear
	Investigators blinded to confounding and prognostic factors? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Intent-to-treat principle adhered to? yes

Bibliographic reference	Maciel EL, Peres RL, do Prado TN, Macedo CR, Palaci M, Vinhas SA, Dietze R, Johnson JL and Struchiner CJ (2010) Saline nebulization before gastric lavage in the diagnosis of pulmonary tuberculosis in children and adolescents. <i>Journal of Tropical Pediatrics</i> 56(6): 458-9
	Inconsistency
	Groups comparable at baseline? yes, although details provided were limited
	Groups received the same care apart from the intervention(s) studied? yes, although details provided were limited
	Equal follow-up? yes
	Groups equivalent for intervention completion? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Randomised = 104 participants • gastric lavage plus nebulisation = 36 • gastric lavage alone = 68
Patient characteristics	Inclusion Children with suspected pulmonary tuberculosis based on having a cough for >28 days and meeting one of the following criteria: household contact with a person with tuberculosis; weight loss or failure to gain weight; positive tuberculin skin test; or a chest radiograph with a parenchymal infiltrate, atelectasis, pleural effusion or lymphadenopathy Exclusion Previously treatment for tuberculosis HIV infection Characteristics of included participants Mean age and sex of subjects did not differ between groups

Bibliographic reference	Maciel EL, Peres RL, do Prado TN, Macedo CR, Palaci M, Vinhas SA, Dietze R, Johnson JL and Struchiner CJ (2010) Saline nebulization before gastric lavage in the diagnosis of pulmonary tuberculosis in children and adolescents. <i>Journal of Tropical Pediatrics</i> 56(6): 458-9
Intervention	Gastric lavage plus nebulisation Inhalation of 30 ml of nebulized sterile 3% hypertonic saline using an ultrasonic nebulizer for 30 minutes before gastric lavage After the gastric contents were aspirated and transferred to container with 10% disodium phosphate buffer and processed using standard procedures
Comparator	Gastric lavage alone Gastric contents were aspirated and transferred to container with 10% disodium phosphate buffer and processed using standard procedures
Location	Brazil
Outcomes measures and effect size	Sample volume (mean, ml) • gastric lavage plus nebulisation (n = 36) = 25 • gastric lavage alone (n = 68) = 10
	Culture positivity (number positive/total number of cases) • gastric lavage plus nebulisation = 9/36 • gastric lavage alone = 14/68
Source of funding	Supported by the Tuberculosis Research Unit at Case Western Reserve University, established with funds from the United States National Institute of Allergy and Infectious Diseases, National Institutes of Health, and, in part, by funding for CNPq (National Council for Scientific and Technological Development) and REDE-TB (Brazilian Tuberculosis Research Network)
Comments	

1.1.12 Menon, 2011

Bibliographic reference	Menon PR, Lodha R, Singh U and Kabra SK (2011) A prospective assessment of the role of bronchoscopy and bronchoalveolar lavage in evaluation of children with pulmonary tuberculosis. <i>Journal of Tropical Pediatrics</i> 57(5): 363-7
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes

Bibliographic reference	Menon PR, Lodha R, Singh U and Kabra SK (2011) A prospective assessment of the role of bronchoscopy and bronchoalveolar lavage in evaluation of children with pulmonary tuberculosis. <i>Journal of Tropical Pediatrics</i> 57(5): 363-7
	Did the study avoid inappropriate exclusions? yes
	Participants blinded? unclear
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? unclear
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? unclear
	Equal follow-up? unclear
	Groups comparable for availability of data? unclear
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 52
Patient characteristics	Inclusion

Bibliographic reference	Menon PR, Lodha R, Singh U and Kabra SK (2011) A prospective assessment of the role of bronchoscopy and bronchoalveolar lavage in evaluation of children with pulmonary tuberculosis. <i>Journal of Tropical Pediatrics</i> 57(5): 363-7
	Children <16 years of age with probably pulmonary tuberculosis, based on the presence of cough (≥2 weeks), fever and persistent radiological infiltrates (chest x-ray showing atelectasis, consolidation, cavitations or effusion
	Exclusion
	Antituberculosis prophylaxis or treatment
	HIV infection
	Respiratory failure
	Characteristics of included participants
	Age (mean±SD (range), years) = 7.8±3.93 (0.75–16)
	22 females, 30 males
	38 (73%) of children had cough of >2 weeks
	Positive Mantoux (n = 34) = 27 (65.4%)
	Chest roentography:
	• mediastinal adenopathy = 11
	• cavity = 5
	• consolidation = 13
	• parenchymal infiltrates = 23
Intervention	Nasogastric aspiration
	All participants provided a minimum of 1 specimen
	After an overnight fast of at least 6 hours, nasogastric tube was passed before the child got up and the gastric contents aspirated

Bibliographic reference	Menon PR, Lodha R, Singh U and Kabra SK (2011) A prospective assessment of the role of bronchoscopy and bronchoalveolar lavage in evaluation of children with pulmonary tuberculosis. <i>Journal of Tropical Pediatrics</i> 57(5): 363-7			
	If the aspirate was <10 ml, normal saline (10 ml) was injected in the tube, left for 2 to 3 minutes and then aspirated until at least 10 ml of aspirate was obtained			
	Examined by Ziehl-Neelsen microscopy			
Comparator	Bronchoalveolar lavage			
	All participants provided 1 specimen			
	Performed using a flexible fibreoptic bronchoscope			
	Intravenous midazolam (0.15 mg/kg) and pethidine (1 to 2 mg/kg) were used in all patients			
	Atropine (0.01 mg/kg, at a minimum dose of 0.1 mg) prior to the procedure			
	Lignocaine 2% jelly was used in the nostril, and 1% lignocaine was sprayed over the vocal cords on visualisation			
	The flexible bronchoscope was passed through the roomier nostril, and oxygen was given through the other			
	Lavage was performed with sterile 0.9% saline (2 to 3 ml/kg, at a minimum of 10 ml) from the radiologically affected segment or lesion			
	Examined by Ziehl-Neelsen microscopy			
Location	New Delhi, India			
Outcomes measures and	Smear positivity (number positive/total cases)			
effect size	• nasogastric aspiration = 6/52			
	• bronchoalveolar lavage = 16/52			
	• nasogastric aspiration plus bronchoalveolar lavage = 19/52			
Source of funding	No details provided			

Bibliographic reference	Menon PR, Lodha R, Singh U and Kabra SK (2011) A prospective assessment of the role of bronchoscopy and bronchoalveolar lavage in evaluation of children with pulmonary tuberculosis. <i>Journal of Tropical Pediatrics</i> 57(5): 363-7
Comments	Incomplete data for 2 nd and 3 rd gastric aspirations (n = 22) therefore data not extracted
Abbreviations: SD, standard	deviation

1.1.13 **Mukherjee**, 2013

Bibliographic reference	Mukherjee A, Singh S, Lodha R, Singh V, Hesseling AC, Grewal HM and Kabra SK; Delhi Pediatric TB Study Group (2013) Ambulatory gastric lavages provide better yields of Mycobacterium tuberculosis than induced sputum in children with intrathoracic tuberculosis. <i>Pediatric Infectious Disease Journal</i> 32(12): 1313-7
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? yes
	Participants blinded? unclear
	Individuals administering care blinded? unclear
	Investigators blinded? yes
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes, same day
	Inconsistency

Bibliographic reference	Mukherjee A, Singh S, Lodha R, Singh V, Hesseling AC, Grewal HM and Kabra SK; Delhi Pediatric TB Study Group (2013) Ambulatory gastric lavages provide better yields of Mycobacterium tuberculosis than induced sputum in children with intrathoracic tuberculosis. <i>Pediatric Infectious Disease Journal</i> 32(12): 1313-7
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 403 participants
Patient characteristics	Inclusion
	Children aged 6 months to 15 years with newly diagnosed intrathoracic tuberculosis, based on clinico-radiological criteria as recommended by the Indian Academy of Pediatrics; in the presence of persistent radiological abnormalities, with nonresolution of clinical symptoms and no alternative cause for symptoms and radiological findings, a clinical diagnosis of probable intrathoracic tuberculosis was made <i>Exclusion</i>
	Known HIV infection
	Characteristics of the included population Female = 56.6%
	Age (median [interquartile range], months) = 111 [68–144]
	History of contact with adult tuberculosis case = 150 (37.2%)
	Positive tuberculin skin test = 371 (92.1%)
	Primary pulmonary complex on chest radiograph = 120 (29.8%) Pleural effusion on chest radiograph = 54 (13.4%)
Intervention	Nasogastric aspiration and lavage Specimens collected on 2 consecutive days after overnight fasting of 6–8 hours An appropriate sized feeding tube was inserted through nostril till it reached the stomach; the gastric contents were aspirated

Bibliographic reference	Mukherjee A, Singh S, Lodha R, Singh V, Hesseling AC, Grewal HM and Kabra SK; Delhi Pediatric TB Study Group (2013) Ambulatory gastric lavages provide better yields of Mycobacterium tuberculosis than induced sputum in children with intrathoracic tuberculosis. <i>Pediatric Infectious Disease Journal</i> 32(12): 1313-7 completely followed by a gastric lavage with 5–10 mL of sterile saline Samples were transported to laboratory for processing within 1–2 hours Specimens were examined by Ziehl-Neelsen staining and cultured for 6 weeks on the BACTEC MGIT 960 system			
Comparator	Specimens were examined by Zierli-Neelsen staining and cultured for 6 weeks on the BACTEC MGIT 900 system. Specimens collected on 2 consecutive days after overnight fasting of 6–8 hours; during the same session, and approximately 30 minutes after collecting gastric lavage, induced sputum was also collected from each patient. The child was administered 2 puffs (100 µg/puff) of salbutamol inhalation by metered dose inhaler followed by nebulization with 3 mL of 3% saline over next 15–20 minutes; chest physiotherapy was performed, and then a feeding tube of appropriate size was placed in the nostril till it reached the posterior nasopharyngeal wall, which was then used to aspirate the secretions. The usual volume of sample collected was 1–2 ml. Specimens were examined by Ziehl-Neelsen staining and cultured for 6 weeks on the BACTEC MGIT 960 system.			
Location	Delhi, India			
Outcomes measures and effect size	 Culture positivity (number of positive nasogastric aspiration and lavage = sputum induction plus nasopharyng 	135/403 eal aspiratio Day 1	Day 2	Day 1 or 2
	Nasogastric aspiration and lavage Sputum induction plus nasopharyngeal aspiration	91 48	101 53	135 72
	Smear positivity (number of positive of nasogastric aspiration and lavage = sputum induction plus nasopharyng Nasogastric aspiration and lavage Sputum induction plus nasopharyngeal aspiration Smear and/or culture positivity (number of nasogastric aspiration and lavage = sputum induction plus nasopharyng) K statistic = 0.441	42/403 eal aspiration Day 1 28 14 per of positive 79/403	Day 2 25 16 e cases)	Day 1 or 2 42 23

Bibliographic reference	Mukherjee A, Singh S, Lodha R, Singh V, Hesseling AC, Grewal HM and Kabra SK; Delhi Pediatric TB Study Group (2013) Ambulatory gastric lavages provide better yields of Mycobacterium tuberculosis than induced sputum in children with intrathoracic tuberculosis. <i>Pediatric Infectious Disease Journal</i> 32(12): 1313-7			
	Agreement = 77.7%			
		Day 1	Day 2	Day 1 or 2
	Nasogastric aspiration and lavage	95	103	135
	Sputum induction plus nasopharyngeal aspiration	52	57	79
	Adverse events Both gastric lavage and sputum induction were carried out without any clinically significant adverse events Minor events encountered were 4 (0.5%) episodes of vomiting and 8 (1%) episodes of nasal bleed during gastric lavage; 2 (0.25%) episodes each of vomiting and cough during sputum induction			
Source of funding	No details provided			
Comments				

1.1.14 Oberhelman, 2006

Bibliographic reference	Oberhelman RA, Soto-Castellares G, Caviedes L, Castillo ME, Kissinger P, Moore DA, Evans C and Gilman RH (2006) Improved recovery of Mycobacterium tuberculosis from children using the microscopic observation drug susceptibility method. <i>Pediatrics</i> 118(1):e100-6
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? yes
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? yes, although details provided were limited
	Participants blinded? unclear
	Individuals administering care blinded? unclear
	Investigators blinded? unclear

Bibliographic reference	Oberhelman RA, Soto-Castellares G, Caviedes L, Castillo ME, Kissinger P, Moore DA, Evans C and Gilman RH (2006) Improved recovery of Mycobacterium tuberculosis from children using the microscopic observation drug susceptibility method. <i>Pediatrics</i> 118(1):e100-6
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? unclear
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 165 participants • nasogastric aspirate = 324 specimens • nasopharyngeal aspirate = 319 specimens
Patient characteristics	Inclusion Children aged ≤12 years Clinical suspicion of tuberculosis, defined as a Stegen-Toledo clinical score ≥5 points ^a Absence of antituberculosis therapy Characteristics of included participants Approximately 20% of patients were inpatients, and 80% were outpatients 93 boys and 72 girls

Mean age of all patients was 4.6 years, and the age distribution of each group was as follows: • ≤1 year = 21 (12%) patients • 1 to 5 years = 72 (44%) patients
 >5 years = 72 (44%) patients HIV enzyme-linked immunosorbent assay results were negative in 136 (82%) patients, and the test was declined in 29 (18% patients, none of whom had clinical evidence of HIV infection
Nasogastric aspirate Collected on 2 successive mornings by nasogastric intubation; the volume of the specimens was augmented as needed by injection of sterile water Examined by fluorescence microscopy, and cultured by Microscopic Observation Drug Susceptibility and Löwenstein-Jenser techniques
Nasopharyngeal aspirate Collected daily for 2 consecutive days by insertion of a soft, flexible nasopharyngeal tube into the nasopharynx, lavage with 5 mL of saline, and aspiration of the respiratory secretions into a container with an electrical suction device or hand-held aspirator Examined by fluorescence microscopy, and cultured by Microscopic Observation Drug Susceptibility and Löwenstein-Jenser techniques
Lima, Peru
Löwenstein-Jensen positivity (number positive/total number of cases) • nasogastric aspirate = 11/321 • nasopharyngeal aspirate = 8/314
Microscopic Observation Drug Susceptibility positivity (number positive/total number of cases) • nasogastric aspirate = 19/321 • nasopharyngeal aspirate = 11/314
Fluorescence microscopy positivity (number positive/total number of cases) • nasogastric aspirate = 8/321 • nasopharyngeal aspirate = 4/314
Financially supported by the National Institutes of Health and the National Institute of Allergy and Infectious Diseases

^a Stegen-Toledo criteria (note: although positive *M. tuberculosis* culture is 1 of the Stegen-Toledo clinical criteria, culture results are the primary outcome parameter of the study, and these were not available at enrollment, so this criterion was not used to determine patient eligibility):

Bibliographic reference	Oberhelm (2006) Imp susceptib	pro
Finding	Sco	core
Positive culture	7	
TB granuloma	4	
Positive purified protein derivative >	10 mm 3	
Known contact in the last 2 y	2	
Suggestive radiograph	2	
Suggestive clinical picture (cough >	2 wk) 2	
Clinical criteria		
Highly probable TB: score ≥	7	
Probable TB: score 5-6		
Suspicious TB: score 3-4		
Unlikely TB: score 0-2		

1.1.15 Oberhelman, 2010

Bibliographic reference	Oberhelman RA, Soto-Castellares G, Gilman RH, Caviedes L, Castillo ME, Kolevic L, Del Pino T, Saito M, Salazar-Lindo E, Negron E, Montenegro S, Laguna-Torres VA, Moore DA and Evans CA (2010) Diagnostic approaches for paediatric tuberculosis by use of different specimen types, culture methods, and PCR: a prospective case-control study. <i>Lancet Infectious Diseases</i> 10(9): 612-20
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes, though only because control data not extracted (gastric aspiration not performed in these as procedure too invasive to be justified in healthy individuals)
	Did the study avoid inappropriate exclusions? yes
	Participants blinded? unclear
	Individuals administering care blinded? unclear

Bibliographic reference	Oberhelman RA, Soto-Castellares G, Gilman RH, Caviedes L, Castillo ME, Kolevic L, Del Pino T, Saito M, Salazar-Lindo E, Negron E, Montenegro S, Laguna-Torres VA, Moore DA and Evans CA (2010) Diagnostic approaches for paediatric tuberculosis by use of different specimen types, culture methods, and PCR: a prospective case-control study. <i>Lancet Infectious Diseases</i> 10(9): 612-20
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? unclear
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 218
Patient characteristics	Inclusion
	Clinical evidence suggestive of pulmonary tuberculosis (Stegen-Toledo score ^a ≥ 5 points)
	Age ≤12 years

Bibliographic reference	Oberhelman RA, Soto-Castellares G, Gilman RH, Caviedes L, Castillo ME, Kolevic L, Del Pino T, Saito M, Salazar-Lindo E, Negron E, Montenegro S, Laguna-Torres VA, Moore DA and Evans CA (2010) Diagnostic approaches for paediatric tuberculosis by use of different specimen types, culture methods, and PCR: a prospective case-control study. <i>Lancet Infectious Diseases</i> 10(9): 612-20
	Absence of antituberculosis therapy
	Exclusion
	Children with evidence of HIV infection or AIDS
Intervention	Nasogastric aspirate
	Gastric aspirates were collected on 2 successive early mornings by brief (<10 minute) nasogastric intubation following an overnight fast
	The volume of gastric aspirates was augmented as needed by injecting 5 ml. sterile water and aspirating back
	Examined by IS6110-based PCR, and cultured by Microscopic Observation Drug Susceptibility and Löwenstein-Jensen techniques
Comparator	Nasopharyngeal aspirate
	Nasopharyngeal aspirates were collected daily for 2 days by inserting a soft flexible nasopharyngeal tube into the nasopharynx, lavaging with 5 ml of saline solution, and aspirating with an electrical suction device or hand-held aspirator; the procedure induces a cough and sputum production, which is then aspirated from the nasopharynx
	Examined by IS6110-based PCR, and cultured by Microscopic Observation Drug Susceptibility and Löwenstein- Jensen techniques
Location	Peru
Outcomes measures and effect size	Culture positivity (number of positives/ total cases)
	• nasogastric aspirate = 22/216
	• nasopharyngeal aspirate = 12/215
	PCR positivity (number of positives/ total cases)

Bibliographic reference	Oberhelman RA, Soto-Castellares G, Gilman RH, Caviedes L, Castillo ME, Kolevic L, Del Pino T, Saito M, Salazar-Lindo E, Negron E, Montenegro S, Laguna-Torres VA, Moore DA and Evans CA (2010) Diagnostic approaches for paediatric tuberculosis by use of different specimen types, culture methods, and PCR: a prospective case-control study. <i>Lancet Infectious Diseases</i> 10(9): 612-20
	• nasogastric aspirate = 35/217
	• nasopharyngeal aspirate = 26/218
Source of funding	Supported by the National Institutes of Health, NIAID, IFHAD, FIND and the Wellcome Trust
Comments	

^a Stegen-Toledo criteria (note: although positive *M. tuberculosis* culture is 1 of the Stegen-Toledo clinical criteria, culture results are the primary outcome parameter of the study, and these were not available at enrollment, so this criterion was not used to determine patient eligibility):

Finding	Score
Positive culture	7
TB granuloma	4
Positive purified protein derivative >10 mm	3
Known contact in the last 2 y	2
Suggestive radiograph	
Suggestive clinical picture (cough >2 wk)	
Clinical criteria	
Highly probable TB: score ≥7	
Probable TB: score 5-6	
Suspicious TB: score 3-4	
Unlikely TB: score 0-2	

1.1.16 Owens, 2007

Bibliographic reference	Owens S, Abdel-Rahman IE, Balyejusa S, Musoke P, Cooke RP, Parry CM and Coulter JB (2007) Nasopharyngeal aspiration for diagnosis of pulmonary tuberculosis. <i>Archives of Disease in Childhood</i> 92(8): 693-6
Study type	Cross-sectional
Study quality	Study limitations

Bibliographic reference	Owens S, Abdel-Rahman IE, Balyejusa S, Musoke P, Cooke RP, Parry CM and Coulter JB (2007) Nasopharyngeal aspiration for diagnosis of pulmonary tuberculosis. <i>Archives of Disease in Childhood</i> 92(8): 693-6
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? yes
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? smear positivity, yes; culture positivity, unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	96 eligible for inclusion; 94 participants included A child with AIDS had dyspnoea and pneumonic changes on chest radiograph; he deteriorated during pre-induction with

Bibliographic reference	Owens S, Abdel-Rahman IE, Balyejusa S, Musoke P, Cooke RP, Parry CM and Coulter JB (2007) Nasopharyngeal aspiration for diagnosis of pulmonary tuberculosis. <i>Archives of Disease in Childhood</i> 92(8): 693-6 salbutamol and the procedure was abandoned Another child who became emotionally distressed during nasopharyngeal aspiration developed a brisk epistaxis and the procedure was abandoned Both children were excluded from analysis because their datasets were incomplete Six cultures were contaminated and not included in the paired analysis for culture positivity
Patient characteristics	Inclusion Symptoms suggestive of pulmonary tuberculosis, for example cough for over 2 weeks, weight loss, or severe malnutrition not responding to nutritional rehabilitation Positive tuberculin test (≥10 mm induration, irrespective of the presence of a BCG scar, was regarded as positive; in severely malnourished patients or patients known to have HIV infection, 6–9 mm was accepted) and/or a chest radiograph compatible with tuberculosis (hilar or mediastinal lymphadenopathy, local collapse/consolidation, severe bilateral but asymmetric disease, cavitation or miliary changes) Where the tuberculin test was negative and the chest radiograph was non-specific, the child was given a 2 week course of antibiotics. If symptoms or chest radiograph did not improve, they were then recruited Characteristics of included participants Median (range) age of the study group was 48 (4–144) months 57 (60.6%) were male Median weight-for-height z score (interquartile range) was −1.30 (−2.79 to +0.04) 22.9% were severely malnourished Of 63 children who were tested, 44 (69.8%) were infected with HIV
Intervention	Nasopharyngeal aspiration Single specimen collected; collection performed prior to sputum induction Patients were in the sitting position; a graduated suction catheter was inserted through the nostril into the oropharynx which stimulated a cough reflex; secretions were aspirated mechanically Sputum specimens were digested and decontaminated with NAOH-NALC-NA-citrate before undergoing centrifugation, fluorescence microscopy (the presence of 1 acid-fast bacillus in 100 high-powered microscopy fields defined a positive smear) and culture on Löwenstein Jensen media
Comparator	Sputum induction plus nasopharyngeal aspiration Single specimen collected; collection performed after nasopharyngeal aspiration Salbutamol (500 µg) was nebulised initially for 5 min; this was followed by 15 ml of 3% hypertonic saline for 20 min Nasopharyngeal aspiration was undertaken to obtain the secretions Chest physiotherapy was not undertaken prior to suction Sputum specimens were digested and decontaminated with NAOH-NALC-NA-citrate before undergoing centrifugation,

Bibliographic reference	Owens S, Abdel-Rahman IE, Balyejusa S, Musoke P, Cooke RP, Parry CM and Coulter JB (2007) Nasopharyngeal aspiration for diagnosis of pulmonary tuberculosis. <i>Archives of Disease in Childhood</i> 92(8): 693-6
	fluorescence microscopy (the presence of 1 acid-fast bacillus in 100 high-powered microscopy fields defined a positive smear) and culture on Löwenstein Jensen media
Location	Kampala, Uganda
Outcomes measures and effect size	Culture positivity • nasopharyngeal aspiration = 21/88 • sputum induction plus nasopharyngeal aspiration = 19/88 K statistic = 0.74
	Smear positivity • nasopharyngeal aspiration = 8/94 • sputum induction plus nasopharyngeal aspiration = 9/94 K statistic =0.81
	Adverse events A small number of children had coughing spasms and/or vomiting after sputum induction Some had bloodstained aspirates following both procedures
Source of funding	No details provided
Comments	

1.1.17 Planting, 2014

Bibliographic reference	Planting NS, Visser GL, Nicol MP, Workman L, Isaacs W and Zar HJ (2014) Safety and efficacy of induced sputum in young children hospitalised with suspected pulmonary tuberculosis. <i>International Journal of Tuberculosis and Lung Disease</i> 18(1): 8-12
Study type	Cohort
Study quality	Study limitations
	Method of allocation was unrelated to potential confounding factors? no, based on child's ability to spontaneously produce sputum
	Attempts were made within the design or analysis to balance the comparison groups for potential confounders? no

Bibliographic reference	Planting NS, Visser GL, Nicol MP, Workman L, Isaacs W and Zar HJ (2014) Safety and efficacy of induced sputum in young children hospitalised with suspected pulmonary tuberculosis. <i>International Journal of Tuberculosis and Lung Disease</i> 18(1): 8-12
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Inconsistency
	Groups comparable at baseline? unclear
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	A total of 1270 sputum induction procedures were performed in 690 patients • induced sputum plus nasopharyngeal aspiration = 993 • coughed induced sputum = 264 • unknown = 13
Patient characteristics	Inclusion

Bibliographic reference	Planting NS, Visser GL, Nicol MP, Workman L, Isaacs W and Zar HJ (2014) Safety and efficacy of induced sputum in young children hospitalised with suspected pulmonary tuberculosis. <i>International Journal of Tuberculosis and Lung Disease</i> 18(1): 8-12		
	criteria: household contact kno	wn to be infected with tube s; positive tuberculin skin to reatment or prophylaxis	sis based on chronic cough (>14 days) and one of the following rculosis within the last 3 months; weight loss or failure to gain est; suggestive features on chest radiography
		Total $(N = 690)$ median [IQR] or n (%)	
	Age, months Age group, months <12 12-60 >60	27.3 [13.4 to 64.2] 146 (21.2) 356 (51.6) 188 (27.2)	
	Sex, male HIV infection Infected Non-infected Unknown	368 (53.3) 164 (23.8) 525 (76.1) 1 (0.1)	
	Total IS procedures IS1 IS2 Height-for-age	1270 690 580 -1.3 [-2.3 to -0.3]	
	Malnourished Weight-for-age	175 (25.4) -1.4 [-2.7 to -0.4]	
	Weight/height-for-age Patients presenting with Oxygen saturation, % Receiving supplemental oxygen Respiratory rate, bpm Subcostal recession Wheeze	-0.5 [-1.9 to 0.5] 98 [96 to 99] 33 (4.8) 36 [28 to 45] 115 (16.7) 95 (13.8)	
	IQR = interquartile range; IS = induced s	sputum; bpm = breaths per minute.	
ntervention	Induced sputum plus nasophar	ryngeal aspiration	

Bibliographic reference	Planting NS, Visser GL, Nicol MP, Workman L, Isaacs W and Zar HJ (2014) Safety and efficacy of induced sputum in young children hospitalised with suspected pulmonary tuberculosis. <i>International Journal of Tuberculosis and Lung Disease</i> 18(1): 8-12		
	2 specimens collected Sputum induction was performed after 2 to 3 hours of fasting Children were pre-treated with 200 µg salbutamol via a metered dose inhaler with attached spacer to prevent bronchoconstriction A jet nebuliser attached to oxygen at a flow rate of 5 l/min delivered 5 ml 5% sterile saline solution for 15 minutes Suctioning was performed through the nasopharynx with a sterile mucus extractor Specimens were cultured for 6 weeks on the BACTEC MGIT 960		
Comparator	Coughed induced sputum 2 specimens collected Sputum induction was performed after 2 to 3 hours of fasting Children were pre-treated with 200 µg salbutamol via a metered dose inhaler with attached spacer to prevent bronchoconstriction A jet nebuliser attached to oxygen at a flow rate of 5 l/min delivered 5 ml 5% sterile saline solution for 15 minutes Children were encouraged to cough, and the expectorated sputum sample was collected Specimens were cultured for 6 weeks on the BACTEC MGIT 960		
Location	Cape Town, South Africa		
Outcomes measures and effect size	Culture positivity(number positive/total number of specimens obtained by collection method) • induced sputum plus nasopharyngeal aspiration = 129/993 • coughed induced sputum = 62/264		
	Adverse events Side effects Coughed Suctioned None 259 744 Nose bleed 4 239 Wheeze 3 11 Cough exacerbation 1 3 Vomiting 0 2 Other 0 5		
Source of funding	Funded by the National Institutes for Health, USA, the National Health Laboratory Services Research Trust, the Medical Research Council of South Africa and the National Research Foundation, South Africa		
Comments			

1.1.18 Somu, 1995

Bibliographic reference	Somu N, Swaminathan S, Paramasivan CN, Vijayasekaran D, Chandrabhooshanam A, Vijayan VK and Prabhakar R (1995) Value of bronchoalveolar lavage and gastric lavage in the diagnosis of pulmonary tuberculosis in children. <i>Tubercle and Lung Disease</i> 76(4): 295-9
Study type	Cross-sectional Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? unclear
	Participants blinded? unclear
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes, same day or the following morning
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes

Bibliographic reference	Somu N, Swaminathan S, Paramasivan CN, Vijayasekaran D, Chandrabhooshanam A, Vijayan VK and Prabhakar R (1995) Value of bronchoalveolar lavage and gastric lavage in the diagnosis of pulmonary tuberculosis in children. <i>Tubercle and Lung Disease</i> 76(4): 295-9
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 50 participants
Patient characteristics	Inclusion
	Children (age group 7 months to 12 years) presenting with signs and symptoms suggestive of pulmonary TB who had a radiographic abnormality but were not acutely sick
	Parenchymal lesion on the chest X-ray that did not clear after 3 weeks in spite of antibiotics
	No prior antituberculosis treatment
	Characteristics of included participants
	Mean age = 5.1 years (range 7 months to 12 years)
	27 males and 23 females
	The majority of the patients were malnourished with 33 of the 50 (66%) suffering from second or third degree malnutrition (Gomez classification)
	The commonest radiographic abnormality was a persistent consolidation with or without hilar lymphadenopathy (25 cases), followed by bronchiectatic changes in one or both lower lobes (10 cases); hilar or mediastinal adenopathy alone was seen in 6 cases, 5 children had evidence of cavitation in addition to parenchymal changes, 1 had collapse consolidation and 3 had segmental atelectasis
	The Mantoux test was positive (> 10 ram) in 37 patients and negative in 13 of the 50 children (26%)
	A history of contact with an adult tuberculosis patient was elicited in 21 (42%) children
Intervention	Nasogastric lavage
	Performed early in the morning, after an overnight fast

Bibliographic reference	Somu N, Swaminathan S, Paramasivan CN, Vijayasekaran D, Chandrabhooshanam A, Vijayan VK and Prabhakar R (1995) Value of bronchoalveolar lavage and gastric lavage in the diagnosis of pulmonary tuberculosis in children. <i>Tubercle and Lung Disease</i> 76(4): 295-9
	A nasogastric tube was used to aspirate the gastric contents, after which 5-10 ml normal saline was injected through the nasogastric tube and, again, aspirated
	Specimens were cultured for 6 to 8 weeks on 2 slopes each of Löwenstein-Jensen medium, Löwenstein-Jensen medium with sodium pyruvate and Middlebrooks selective 7HII medium and 2 bottles of selective Kirschner's liquid medium
Comparator	Performed using a flexible fibreoptic bronchoscope The patients were kept fasting overnight Topical anaesthesia with 2% Xylocaine was applied to the nose, larynx and upper airways as the bronchosope was advanced; the amount of Xylocaine used ranged 3-5 ml (60-100 mg) The flexible bronchoscope was inserted transnasally, advanced into the trachea and wedged into the most involved segment as seen on chest X-ray or the nearest segment possible Bronchoalveolar lavage was performed by instilling 2 ml/kg sterile normal saline and subsequently aspirating it back into a specimen trap, using a suction apparatus Specimens were cultured for 6 to 8 weeks on 2 slopes each of Löwenstein-Jensen medium, Löwenstein-Jensen medium with sodium pyruvate and Middlebrooks selective 7HII medium and 2 bottles of selective Kirschner's liquid medium
Location	Madras, India
Outcomes measures and effect size	Culture positivity • nasopharyngeal lavage = 16/50 • bronchoalveolar lavage = 6/50
Source of funding	No details provided
Comments	

1.1.19 Thomas, 2014

Bibliographic reference	Thomas TA, Heysell SK, Moodley P, Montreuil R, Ha X, Friedland G, Bamber SA, Moll AP, Gandhi N, Brant WE, Sturm W and Shah S (2014) Intensified specimen collection to improve tuberculosis diagnosis in children from Rural South Africa, an observational study. <i>BMC Infectious Diseases</i> 14: 11
Study type	Cross-sectional

Bibliographic reference	Thomas TA, Heysell SK, Moodley P, Montreuil R, Ha X, Friedland G, Bamber SA, Moll AP, Gandhi N, Brant WE, Sturm W and Shah S (2014) Intensified specimen collection to improve tuberculosis diagnosis in children from Rural South Africa, an observational study. <i>BMC Infectious Diseases</i> 14: 11
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? yes
	Participants blinded? unclear
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes

Bibliographic reference	Thomas TA, Heysell SK, Moodley P, Montreuil R, Ha X, Friedland G, Bamber SA, Moll AP, Gandhi N, Brant WE, Sturm W and Shah S (2014) Intensified specimen collection to improve tuberculosis diagnosis in children from Rural South Africa, an observational study. <i>BMC Infectious Diseases</i> 14: 11
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 118
Patient characteristics	Inclusion Tuberculosis suspects aged ≥6 months and ≤12 years Subjects were new tuberculosis suspects or had persistent symptoms despite ≥2 months of antituberculosis treatment Patients were eligible if they had at least one of the following: chronic cough, failure to improve after pneumonia treatment, contact with a tuberculosis case, failure to thrive, painless superficial lymphadenopathy, signs of meningitis which were not responsive to antibiotics, or chest radiograph suggestive of tuberculosis Exclusion On antituberculosis treatment for >2 days, or recently defaulted on antituberculosis treatment Characteristics of included participants

	Total	
A	n = 118, (%)	
Age, mean years [SD]	4.7 [3.1]	
Female gender, n (%)	53 (45)	
Mother as primary caregiver	70 (59)	
Either parent deceased	37 (31)	
HIV-infected mother	50 (42)	
Other HIV-infected family member	9 (8)	
TB contact reported	62 (53)	
Drug-resistant TB contact reported	4 (3)	
Contact with "chronic cougher"	19 (16)	
Prior TB	37 (31)	
Prior hospitalization (≥1)	31 (26)	
Malnourished	52 (44)	
Severely malnourished (n, % of malnourished)	42 (81)	
Tested for HIV	99 (84)	
HIV positive	64 (65)	
On HAART, n (% of HIV+)	32 (50)	
CD4%, median [IQR]	16.8 [11.5-22.3]	
Hemoglobin, median g/dL [IQR]	9.9 [8.8-11.2]	
Albumin, median g/L [IQR]	32.0 [28.1-37.0]	
Erythrocyte Sedimentation Rate, median [IQR]	32.0 [14.0-75.0]	
Abnormal chest radiograph consistent with TB	84* (85)	
Enrolled while failing ≥2 mo TB treatment	8 (7)	
Started on TB treatment	59 (50)	
*Of 99 children with chest radiographs available for re		

Bibliographic reference	Thomas TA, Heysell SK, Moodley P, Montreuil R, Ha X, Friedland G, Bamber SA, Moll AP, Gandhi N, Brant WE, Sturm W and Shah S (2014) Intensified specimen collection to improve tuberculosis diagnosis in
	An aliquot was used for fluorescent microscopic sputum examination for acid-fast bacilli and the remainder was split for parallel culture on Middlebrook 7H11 agar plates and automated broth culture via the BACTEC MGIT-960 system (solid cultures were monitored at 21- and 42-days; liquid cultures were monitored continuously for 42 days)
Comparator	Induced or spontaneously produced sputum
	Single sputum induction according to World Health Organisation guidance by the hospital's trained respiratory physiotherapists
	An aliquot was used for fluorescent microscopic sputum examination for acid-fast bacilli and the remainder was split for parallel culture on Middlebrook 7H11 agar plates and automated broth culture via the BACTEC MGIT-960 system (solid cultures were monitored at 21- and 42-days; liquid cultures were monitored continuously for 42 days)
Location	Tugela Ferry, South Africa
Outcomes measures and	Culture positivity (number positive/ total number of cases)
effect size	• nasogastric aspirate = 5/67
	• sputum = 7/67
	• nasogastric aspirate plus sputum = 8/67
	Among the 5 children who were culture-positive by sequential gastric aspirates, the yield from the first gastric aspirate was equivalent to multiple collections
	Adverse effects – only reported for induced sputum
	Sputum induction was well tolerated overall: no child experienced prolonged respiratory distress, one child required brief supplemental oxygen and two had post-procedural vomiting
Source of funding	Supported in part by the Einstein-Montefiore Center for AIDS funded by the National Institutes of Health, the Burroughs Wellcome Fund/American Society of Tropical Medicine & Hygiene, the Howard Hughes Medical

Bibliographic reference	Thomas TA, Heysell SK, Moodley P, Montreuil R, Ha X, Friedland G, Bamber SA, Moll AP, Gandhi N, Brant WE, Sturm W and Shah S (2014) Intensified specimen collection to improve tuberculosis diagnosis in children from Rural South Africa, an observational study. <i>BMC Infectious Diseases</i> 14: 11
	Institute, the US President's Emergency Plan for AIDS Relief and the Doris Duke Charitable Foundation Clinical Scientist Development Award
Comments	

1.1.20 Zar, 2000

Bibliographic reference	Zar HJ, Tannenbaum E, Apolles P, Roux P, Hanslo D and Hussey G (2000) Sputum induction for the diagnosis of pulmonary tuberculosis in infants and young children in an urban setting in South Africa. <i>Archives of Disease in Childhood</i> 82(4): 305-8
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? unclear
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes, sputum induction was performed on enrolment and gastric lavage was performed after a minimum four hour fast

Bibliographic reference	Zar HJ, Tannenbaum E, Apolles P, Roux P, Hanslo D and Hussey G (2000) Sputum induction for the diagnosis of pulmonary tuberculosis in infants and young children in an urban setting in South Africa. <i>Archives of Disease in Childhood</i> 82(4): 305-8
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 142
Patient characteristics	Inclusion
	Children with a primary diagnosis of pneumonia and who were HIV infected, were suspected of having HIV infection or were admitted to the intensive care unit but were not intubated
	Characteristics of included participants

Bibliographic reference	Zar HJ, Tannenbaum E, Apolles P, Roux P, Hanslo D and Hussey G (2000) Sputum induction for the diagnosis of pulmonary tuberculosis in infants and young children in an urban setting in South Africa Archives of Disease in Childhood 82(4): 305-8
	Characteristic Children without TB ($n = 126$) TB ($n = 16$)
	Age (months) 9 (3-21.5) 12 (7-25) Male: female 1.2 2.2 ICU admission, n (%) 18 (14) 2 (12) HIV positive, n (%) 90 (71) 10 (63) TB contact, n (%) 24 (19) 7 (44)* Use of supplemental O ₂ , m(%) 83 (66) 8 (50) Baseline O ₂ saturation in air 94 (90.5-97) 93.5 (88-98) Baseline RR 50 (40-60) 53 (40-63.5)
	Continuous variables expressed as median (25th to 75th percentile). TB, tuberculosis; ICU, intensive care unit; RR, respiratory rate. *p = 0.02.
tervention	Nasogastric lavage Early morning gastric lavage was performed after an overnight fast of at least 4 hours, ideally on 2 or 3 consecutive mornings A nasogastric tube was passed before the child arose and the gastric contents were aspirated
	Normal saline 20 ml was inserted down the tube, left for 2 to 3 minutes and then aspirated; additional 5 to 10 normal saline aliquots were inserted and aspirated until a minimum of 20 ml of aspirate was obtained
	Specimens were cultured for 6 weeks in a BACTEC 12B bottle containing supplemented Middlebrook medium
Comparator	Induced sputum
	Sputum induction was undertaken on the day of enrolment after a 2 to 3 hour fast
	Children were pretreated with 200 µg salbutamol given by a metered dose inhaler with attached spacer to prevent the occurrence of bronchial constriction
	A jet nebuliser attached to oxygen delivered 5 ml of 5% sterile saline for 15 minutes

Bibliographic reference	Zar HJ, Tannenbaum E, Apolles P, Roux P, Hanslo D and Hussey G (2000) Sputum induction for the diagnosis of pulmonary tuberculosis in infants and young children in an urban setting in South Africa. <i>Archives of Disease in Childhood</i> 82(4): 305-8
	Thereafter, physiotherapy techniques – including chest percussion, vibration, shaking and active cycle breathing – were applied
	Sputum was obtained either by expectoration (in children unable to cooperate) or by suctioning through the nasopharynx or oropharynx using a sterile mucus extractor
	Specimens were cultured for 6 weeks in a BACTEC 12B bottle containing supplemented Middlebrook medium
Location	Cape Town, South Africa
Outcomes measures and effect size	Culture positivity • nasogastric lavage = 9/142 • induced sputum = 15/142
Source of funding	Funded by the Medical Research Council of South Africa, the South African Pulmonology Society and the ICH Fund of the Red Cross Children's Hospital
Comments	

1.1.21 Zar, 2005

Bibliographic reference	Zar HJ, Hanslo D, Apolles P, Swingler G and Hussey G (2005) Induced sputum versus gastric lavage for microbiological confirmation of pulmonary tuberculosis in infants and young children: a prospective study. <i>Lancet</i> 365(9454): 130-4
Study type	Cross-sectional
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? unclear
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? yes

Bibliographic reference	Zar HJ, Hanslo D, Apolles P, Swingler G and Hussey G (2005) Induced sputum versus gastric lavage for microbiological confirmation of pulmonary tuberculosis in infants and young children: a prospective study. <i>Lancet</i> 365(9454): 130-4
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes
	Outcomes match the outcomes of interest? yes
Number of patients	Included = 250
Patient characteristics	Inclusion
	Suspected pulmonary tuberculosis on the basis of a chronic cough (more than 28 days) and one of the following

Bibliographic reference	Zar HJ, Hanslo D, Apolles P, Swingler G and Hussey G (2005) Induced sputum versus gastric lavage for microbiological confirmation of pulmonary tuberculosis in infants and young children: a prospective study. <i>Lancet</i> 365(9454): 130-4
	criteria: household contact known to be infected with tuberculosis within the previous 3 months; loss of weight or failure to gain weight within the previous 3 months; positive skin test to purified protein derivative, or chest radiography with parenchymal infiltrate, atelectasis, pleural effusion, or lymphadenopathy
	Exclusion
	Taking treatment for tuberculosis, had completed such treatment within the past 2 weeks, or were taking prophylaxis for this disease
	Had signs of upper airway obstruction
	Had an arterial oxygen saturation less than 92% in room air
	Characteristics of included participants
	141 (56%) were male
	Median age was 13 months (interquartile range 6 to 24)
	Baseline median respiratory rate of children was 56 (interquartile range 40 to 64) breaths per minute
	Median arterial oxygen saturation was 96% (interquartile range 95 to 98%)
	68 (27%) children were receiving supplemental oxygen at the time of sputum induction; 65 via nasal prongs or cannulae and three via headbox oxygen
	30 children (12%) were known to be HIV-infected at enrolment; of the children whose HIV status was unknown, 139 had HIV testing of whom 20 tested positive by PCR, and an additional 45 tested positive by ELISA with clinical features of HIV infection, thus, 95 children (38%) were judged to be HIV infected
Intervention	Nasogastric lavage
	3 specimens
	Early morning gastric lavage after an overnight fast of at least 4 hours
	The first lavage was done the day after enrolment, and 2 more were taken on consecutive mornings

Bibliographic reference	Zar HJ, Hanslo D, Apolles P, Swingler G and Hussey G (2005) Induced sputum versus gastric lavage for microbiological confirmation of pulmonary tuberculosis in infants and young children: a prospective study. <i>Lancet</i> 365(9454): 130-4
	Lavage was done approximately 18 hours after sputum induction
	A nasogastric tube was passed before the child got up and gastric contents aspirated
	If the aspirate was less than 20 ml, 20 ml of normal saline was inserted down the tube, left for 2 to 3 minutes, then aspirated
	Additional 5 to 10 ml samples of normal saline were inserted and aspirated until a minimum of 20 mL of aspirate was obtained
	Samples were cultured singly in a BACTEC 12B bottle containing supplemented Middlebrook medium for 6 weeks
Comparator	Induced sputum
	3 specimens
	Sputum induction was undertaken after a 2 to 3 hour fast
	Children were pretreated with 200 μg salbutamol via metered dose inhaler with attached to prevent bronchoconstriction
	A jet nebuliser that was attached to oxygen at a flow rate of 5 l per minute delivered 5 ml of 5% sterile for 15 minutes; thereafter, chest percussion was done over the anterior and posterior chest wall
	Sputum was obtained by suctioning through the nasopharynx with a sterile mucus extractor
	Sputum induction was done on three consecutive days unless children were discharged, were intubated, or died within this time
	The first specimen was obtained on the day of enrolment; subsequent specimens were obtained on the second and third days after admission, about 6 hours after the early morning gastric lavage
	Samples were cultured singly in a BACTEC 12B bottle containing supplemented Middlebrook medium for 6 weeks

Bibliographic reference	Zar HJ, Hanslo D, Apolles P, Swingler G and Hussey G (2005) Induced sputum versus gastric lavage for microbiological confirmation of pulmonary tuberculosis in infants and young children: a prospective study. <i>Lancet</i> 365(9454): 130-4
Location	Cape Town, South Africa
Outcomes measures and effect size	Culture positivity — 1 st specimen • nasogastric lavage = 19/250 • induced sputum = 37/250 Culture positivity — 3 specimens • nasogastric lavage = 38/250 • induced sputum = 51/250
	Smear positivity – 1st specimen • nasogastric lavage = 8/250 • induced sputum = 19/250 Smear positivity – 3 specimens • nasogastric lavage = 17/250 • induced sputum = 25/250
	Adverse events – reported for sputum induction only No serious adverse reactions attributable to sputum induction occurred during or after the procedure; the most common adverse events were an increase in coughing in 293 procedures (41%), mild epistaxis in 55 (8%), vomiting in three (0·4%), and wheezing that was responsive to an inhaled bronchodilator in two (0·3%) 16 (2%) episodes of transient hypoxia were recorded during sputum induction in which the arterial oxygen saturation dropped below 92%; the lowest oxygen saturation in any child was 88%
Source of funding	Funded by the Medical Research Council, South Africa, and the ICH fund, Red Cross Children's Hospital
Comments	

1.1.22 Zar, 2012

Bibliographic reference	Zar HJ, Workman L, Isaacs W, Munro J, Black F, Eley B, Allen V, Boehme CC, Zemanay W and Nicol MP (2012) Rapid molecular diagnosis of pulmonary tuberculosis in children using nasopharyngeal specimens. <i>Clinical Infectious Diseases</i> 55(8): 1088-95
Study type	Cross-sectional

Bibliographic reference	Zar HJ, Workman L, Isaacs W, Munro J, Black F, Eley B, Allen V, Boehme CC, Zemanay W and Nicol MP (2012) Rapid molecular diagnosis of pulmonary tuberculosis in children using nasopharyngeal specimens. <i>Clinical Infectious Diseases</i> 55(8): 1088-95
Study quality	Study limitations
	Was a consecutive or random sample of patients enrolled? yes
	Was a case-control design avoided? yes
	Did the study avoid inappropriate exclusions? yes
	Participants blinded? no
	Individuals administering care blinded? unclear
	Investigators blinded? unclear
	Appropriate length of follow-up? yes
	Precise definition of outcome? unclear
	Valid and reliable method of outcome measurement? yes
	Was there an appropriate interval between specimen collections? yes, paired specimens
	Inconsistency
	Groups received the same care apart from the intervention(s) studied? yes
	Equal follow-up? yes
	Groups comparable for availability of data? yes
	Indirectness
	Population matches population of interest? yes
	Intervention matches intervention of interest? yes

Bibliographic reference	Zar HJ, Workman L, Isaacs W, Munro J, Black F, Eley B, Allen V, Boehme CC, Zemanay W and Nicol MP (2012) Rapid molecular diagnosis of pulmonary tuberculosis in children using nasopharyngeal specimens. <i>Clinical Infectious Diseases</i> 55(8): 1088-95
	Outcomes match the outcomes of interest? yes
Number of patients	535 had at least one paired induced sputum and nasopharyngeal aspirate
Patient characteristics	Inclusion
	Children under 15 years of age hospitalized with suspected pulmonary tuberculosis
	The reasons for hospitalisation included severe or very severe pneumonia defined by WHO criteria, the need for oxygen or intravenous therapy, or social conditions precluding home care
	Children were eligible if they had a cough for more than 14 days and if one of the following conditions existed: a household tuberculosis contact within the preceding 3 months, weight loss or failure to gain weight within the preceding 3 months, a positive skin test to purified protein derivative, or a chest radiograph suggestive of pulmonary tuberculosis
	Exclusion
	Children who had received tuberculosis drug(s) for longer than 72 hours
	Children without at least one paired IS/NPA specimen
	Children with extrapulmonary tuberculosis
	Characteristics of included participants

		All	
	Age (months; median and IQR)	19.0 (11.2–38.3)	
	Male n (%)	294 (55.0)	
	HIV infection n (%)	117 (21.9)	
	HIV WHO clinical staging n (%)		
	Stage 1	10 (8.6)	
	Stage 2	35 (29.9)	
	Stage 3	42 (35.9)	
	Stage 4	30 (25.6)	
	HIV CDC immune category n (%)		
	None	23 (22.1)	
	Moderate	35 (33.7)	
	Severe	46 (44.2)	
	History of prior tuberculosis N (%)	56 (10.4)	
	Radiological changes suggestive of TB n (%)	333 (67.4)	
	Commenced on anti-tuberculosis treatment n (%)	283 (52.9)	
	Z scores (median and IQR)		
	HAZ	- 1·4 (- 2·5 to - 0·4)	
	WAZ	- 1.4 (- 2.3 to - 0.5)	
	WHZ	− 0·59 (− 1·5–0·3)	
	Malnutrition (WAZ score < - 2) n (%)	68 (15.6)	
	Tuberculin skin-positive n (%)	191 (39.2)	
vention	Nasopharyngeal aspiration		

Bibliographic reference	Zar HJ, Workman L, Isaacs W, Munro J, Black F, Eley B, Allen V, Boehme CC, Zemanay W and Nicol MP (2012) Rapid molecular diagnosis of pulmonary tuberculosis in children using nasopharyngeal specimens. <i>Clinical Infectious Diseases</i> 55(8): 1088-95
	Specimens were examined using the Xpert MTB/RIF assay and cultured for 6 weeks using the BACTEC MGIT 960 system
Comparator	Induced sputum Sputum induction was done at least 30 minutes after a nasopharyngeal aspiration, following a 2–3 hour fast Specimens were examined using the Xpert MTB/RIF assay and cultured for 6 weeks using the BACTEC MGIT 960 system
Location	Cape Town, South Africa
Outcomes measures and effect size	Culture positivity • nasopharyngeal aspiration = 61/535 • induced sputum = 84/535
	Xpert positivity • nasopharyngeal aspiration = 57/535
	• induced sputum = 69/535
Source of funding	Supported by the National Institutes of Health, USA, the National Health Laboratory Services Research Trust, the Medical Research Council of South Africa, and The Wellcome Trust
Comments	