

Review Article

Decontamination of Sputum in the Context of Implementation of Mycobacterial Culture in the Republic of Congo

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Introduction

Tuberculosis is a serious disease. TB is caused by bacteria that spread through the air when a person with contagious TB in their lungs coughs, sneezes, sings, or talks. Pulmonary tuberculosis is still a major health problem in the Republic of Congo and other developing countries despite the major advances in diagnosis and treatment over the past years [1,2]. Sputum smear microscopy is the first-line diagnostic procedure for pulmonary TB in Congo because it is simple, relatively low cost and monitors patient's response to anti-TB treatment. Sputum culture is more sensitive than sputum smear microscopy and though being the gold standard diagnosis for TB [3], remains in limited use due to its cost of implementation, time required to obtain the result and culture contamination [4]. Decontamination is an operation which aims to eliminate, kill or inhibit unwanted microorganisms depending on the objectives set. Only the microorganisms present at the time of the operation are destroyed. Decontamination is partially bacteriostatic, that is to say that under well-defined conditions, the proliferation of microorganisms is momentarily inhibited during the decontamination process.

Culture contamination is a major limitation as it reduces the proportion of interpretable results and diminishes the diagnostic value of culture systems [5]. Factors such as sputum collection method, storage temperature, transport conditions, duration between sample collection and processing, and lab methodology affect contamination rates. The aim of this brief communication is to highlight the main factors linked to contamination through each step from the collection of the sputum sample to its inoculation in culture and which impedes mycobacterial culture establishment in the Congo Table.

Comments

Sputum is mucous that an individual coughs up from deep inside the lungs. It is usually thick, cloudy and sticky. Sputum is not saliva (spit). Saliva comes from your mouth and is thin, clear and watery. The sputum collection must be done very early in the morning, fasting and in the absence of any cigarette intake. These specifications contribute to reducing the effect of contamination of the sample by the bacterial flora of the oral cavity [6].

The multiplication of the common flora bacteria could mask the pathogenic agent (s) responsible for the pulmonary infection. Sputum is thick mucus that is secreted by the lower respiratory tract (bronchi and lungs), which is different from saliva. Teeth should be brushed, the mouth rinsed with a glass of water, two or three deep breaths taken, and finally cough to bring forth the expectoration from the lungs.

Sample collection should be done carefully to ensure that it comes from the lower respiratory tract and not the upper respiratory tract. If the sample contains mainly saliva, the microorganisms identified in culture will not necessarily be responsible for the infection. In addition, the presence of saliva and salivary bacteria in a sputum sample will make it more difficult to identify pathogenic bacteria in the lungs [13]. All samples taken must be sent to the laboratory for mycobacteriological analysis in order to avoid the development of commensal bacteria at the expense of fragile pathogenic bacteria.

The quality of the sputum sample must be checked by the laboratory technician before performing the analysis [14]. In order for the sputum culture to be interpretable, the collection of the sample must be of irreproachable quality [5,15]. A sample of inadequate quality must be rejected and another sample collected [16]. Transport the specimen to the lab the same day it is collected. Within two hours of collection is the preferred time frame for optimal specimen processing.

Indeed, the sample is often contaminated by salivary bacteria (*staphylococci*, *streptococci*, *coryneforms*, *Neisseria*)b [17,18] or commensal bacteria of the respiratory system (that cause no infections under normal conditions, such as *Haemophilus influenzae*, *Streptococcus pneumoniae*, *Staphylococcus aureus*, strict anaerobic bacteria) [19].

Conclusion

The rising rate of contamination of mycobacterial culture shows an urgent need for the improvement of Health center systems in Republic of Congo, patient education, and quality trained personnel at all levels to ensure better specimen collection, handling and processing in order to maintain the viability of the tuberculosis bacilli

Table: Different levels of Sources of Mycobacterial Sputum Contamination.

Test	Conditions	Preservation	Source of contamination
Expectoration	-Well brushed teeth;	- Send the sterile jar containing the sputum within 2 hours following the sample collection; - Protected from light, kept refrigerated.	- Oral flora is a source of contamination for spit [7];
	-Inside of mouth well rinsed with an antiseptic solution;		- Saliva production is a source of contamination [4];
	-Done early in the morning;		- Heat favors contamination [8]
	-Well washed hands;		- Unsupervised collection of sputum by patient [9];
	-Producing mucus from the lungs by coughing heavily.		- Poor Storage and transportation may lead to contamination [10]. - Environment and poorly disinfected laboratory equipment.
Microscopy	-Thoroughly sterilize platinum loop with Bunsen burner flame before use;	-The preservation of the sputum container is put immediately in the refrigerator after staining the slides.	- Poorly sterilized platinum loop [11];
	-Close the sterile jar immediately and store in the refrigerator.		- Open jar exposed to the open air ; - Sputum container left open at room temperature for hours is a source of multiplication of pathogenic microorganisms contained inside the sputum [12].
Decontamination of expectorates	-Sputum should be decontaminated within 3 days before culture;	-Sputum should be stored at refrigerator temperature (+2°C and +8°C) during the 3 day interval.	-A concentration below the standard does not eliminate all contaminants;
	- Final concentration of decontaminating reagent		- the reduced decontamination time does not eliminate all contaminants;
	- sample treatment time is critical		- A non-sterile decontaminant solution will be useless.
	- Use of sterile decontaminating solution		
Culture	-Acceptable contamination for solid media (3-5%)	- Decontaminated expectorates should be cultured immediately;	-The concentration of the decontamination solution must be that recommended for decontamination;
	-Acceptable contamination for liquid media (3-10%)		- Respect of the duration during the process of decontamination must be respected; For example, for NALC-NAOH, a duration of less than 15 minutes would not destroy all the microorganisms contained in the sputum);
	-Good quality reagents (Expiry date)		- Prolonged storage in the refrigerator favours contamination;
	-Sterile culture medium		- Poor storage of decontaminated expectorates in the freezer will favor loss of sample integrity;
	-Sufficient PANTA volume		-Insufficient PANTA volume or expired reagents may favour contamination to >10% (for liquid culture)

with increasing sensitivity while reducing contamination.

References

- Linguissi LS, Mayengue P, Sidibé A, Vouvougui JC, Missontsa M, Madzou-Laboum I, et al. Prevalence of national treatment algorithm defined smear positive pulmonary tuberculosis in HIV positive patients in Brazzaville, Republic of Congo. *BMC Res Notes*. 2014; 7: 578.
- Okemba-Okombi FH, Itoua A, Biniakounou JB, Illoye-Ayet M, Bemba ELP, Ossalé Abacka KB, et al. Apport du GeneXpert dans le diagnostic de la tuberculose pulmonaire : étude préliminaire et perspectives en République du Congo. *Rev Mal Respir*. 2015; 32: A211.
- Magalhães JL de O, Lima JF da C, Araújo AA de, Coutinho IO, Leal NC, Almeida AMP de. Microscopic detection of Mycobacterium tuberculosis in direct or processed sputum smears. *Rev Soc Bras Med Trop*. 2018; 51: 237-239.
- Murphy ME, Phillips PPJ, Mendel CM, Bongard E, Bateson ALC, Hunt R, et al. Spot sputum samples are at least as good as early morning samples for identifying Mycobacterium tuberculosis. *BMC Med*. 2017; 15: 192.
- Reddy M, Gounder S, Reid SA. Tuberculosis diagnostics in Fiji: how reliable is culture? *Public Health Action*. 2014; 4: 184-188.
- Kabore A, Tranchot-Diallo J, Sanou A, Hien H, Daneau G, Gomgnimbou MK, et al. Why oral antiseptic mouth rinsing before sputum collection cannot reduce contamination rate of mycobacterial culture in Burkina-Faso. *Afr Health Sci*. 2019; 19: 1321.
- Jain P, Jain I. Oral Manifestations of Tuberculosis: Step towards Early Diagnosis. *J Clin Diagn Res JCDR*. 2014; 8: ZE18-21.
- Aboubaker Osman D, Garnotel E, Drancourt M. Dry-heat inactivation of "Mycobacterium canettii." *BMC Res Notes*. 2017; 10: 201.
- Chang KC, Leung CC, Yew WW, Tam CM. Supervised and induced sputum among patients with smear-negative pulmonary tuberculosis. *Eur Respir J*. 2008; 31: 1085-1090.
- Tessema B, Beer J, Emmrich F, Sack U, Rodloff AC. Rate of recovery of Mycobacterium tuberculosis from frozen acid-fast-bacillus smear-positive sputum samples subjected to long-term storage in Northwest Ethiopia. *J Clin Microbiol*. 2011; 49: 2557-2561.
- Hunter RA. The routine examination for tubercle bacilli in sputum. *Tubercle*. 1940; 21: 341-359.
- Lagier J-C, Edouard S, Pagnier I, Mediannikov O, Drancourt M, Raoult D. Current and past strategies for bacterial culture in clinical microbiology. *Clin Microbiol Rev*. 2015; 28: 208-236.
- Pragman AA, Berger JP, Williams BJ. Understanding Persistent Bacterial Lung Infections: Clinical Implications Informed by the Biology of the Microbiota and Biofilms. *Clin Pulm Med*. 2016; 23: 57-66.
- Zimba O, Tamuhla T, Basotli J, Letsibogo G, Pals S, Mathebula U, et al. The effect of sputum quality and volume on the yield of bacteriologically-confirmed TB by Xpert MTB/RIF and smear. *Pan Afr Med J*. 2019; 33.
- Datta S, Shah L, Gilman RH, Evans CA. Comparison of sputum collection methods for tuberculosis diagnosis: a systematic review and pairwise and network meta-analysis. *Lancet Glob Health*. 2017; 5: e760-e771.
- Heppele P, Nguete P, Greig J, Bonnet M, Sizaire V. Direct microscopy versus sputum cytology analysis and bleach sedimentation for diagnosis of

- tuberculosis: a prospective diagnostic study. *BMC Infect Dis.* 2010; 10: 276.
17. Cui Z, Zhou Y, Li H, Zhang Y, Zhang S, Tang S, et al. Complex sputum microbial composition in patients with pulmonary tuberculosis. *BMC Microbiol.* 2012; 12: 276.
18. Cardoso-Toset F, Gómez-Laguna J, Amarilla SP, Vela AI, Carrasco L, Fernández-Garayzábal JF, et al. Multi-Etiological Nature of Tuberculosis-Like Lesions in Condemned Pigs at the Slaughterhouse. Cardona P-J, editor. *PLOS ONE.* 2015; 10: e0139130.
19. Cervantes J, Hong B. The gut–lung axis in tuberculosis. *Pathog Dis.* 2017; 75.