INTRODUCTION

Review question / Objective: Tuberculous meningitis (TBM) can lead to serious consequences. Early diagnosis and treatment are very important for TBM, but early diagnosis is still very challenging. This study aims to evaluate the diagnostic accuracy of Xpert MTB/RIF Ultra for TBM using meta-analysis method.

Rationale: The application of Xpert MTB/RIF Ultra in the diagnosis of TBM has its unique advantages.

Condition being studied: TBM is a highly lethal form of extrapulmonary tuberculosis, and although it accounts for a relatively small proportion (1-5%) of new cases of TB, it can kill or severely disable half of all TBM infected people. One of the main reasons for this serious consequence is the failure to diagnose the disease early enough and to treat it properly. Thus, early diagnosis is key to improving the prognosis of TBM. Xpert MTB/RIF Ultra is used in the diagnosis of extrapulmonary tuberculosis, and has achieved good diagnostic efficacy. However, the diagnostic accuracy of Xpert MTB/RIF Ultra on TBM has not been studied well.
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METHODS

Search strategy: #1 "Tuberculosis, Meningeal"[Mesh] OR "Meningeal Tuberculoses" OR "Meningeal Tuberculosis" OR "Tuberculoses, Meningeal" OR "TB Meningitis" OR "TB Meningitides" OR "Tubercular Meningitis" OR "Meningitides, Tubercular" OR "Meningitis, Tubercular" OR "Tubercular Meningitides" OR "Meningitis, Tuberculous" OR "Meningitides, Tuberculous" OR "Tuberculous Meningitides" OR "Tuberculous Meningitis" OR "Tuberculosis Meningitis" OR "Meningitides, Tuberculosis" OR "Meningitis, Tuberculosis" OR "Meningitis, Tuberculosis" OR "Tuberculosis Meningitides" OR "Tuberculous Hypertrophic Pachymeningitis" OR "Hypertrophic Pachymeningitides, Tuberculous" OR "Hypertrophic Pachymeningitis, Tuberculous" OR "Pachymeningitides, Hypertrophic" OR "Pachymeningitis, Tuberculous Hypertrophic" OR "Tuberculous Hypertrophic Pachymeningitides" #2 "Extrapulmonary tuberculosis" OR "Extrapulmonary tuberculosis" #3 "Meningitis"[Mesh] OR Meningitides OR Pachymeningitides OR Pachymentinginities #4 "Cerebrospinal Fluid"[Mesh] OR "Cerebrospinal Fluids" OR "Fluid, Cerebrospinal" OR "Fluids, Cerebrospinal" OR "Cerebro Spinal Fluid" OR "Cerebro Spinal Fluids" OR "Fluid, Cerebro Spinal" OR "Fluid, Cerebro Spinal" OR "Spinal Fluid, Cerebro" OR "Spinal Fluids, Cerebro" #5 #1 OR #2 OR #3 OR #4 #6 "Xpert Ultra" OR "GeneXpert Ultra" #7 #5 AND #6.

Participant or population: Participants using Xpert MTB/RIF Ultra to diagnose TBM.

Intervention: Xpert MTB/RIF Ultra.

Comparator: Comparator test is not an obligatory criteria (single arm study can be enrolled if participants, intervention, outcomes are satisfied because this study will measure the diagnostic accuracy of Xpert MTB/RIF Ultra for TBM.

Study designs to be included: Any study design.

Eligibility criteria: Full-text original researches that assessed the Xpert MTB/RIF Ultra assay for TBM will be included. Clear and appropriate reference standards are defined in researches. True positive (TP), false positive (FP), false negative (FN), and true negative (TN) values for the assay can be extracted or calculated directly from the studies. We will exclude case reports, articles written in languages other than Chinese and English, researches with < 10 specimens, conference reports, and abstracts without full articles.

Information sources: Embase, PubMed, the Cochrane Library, China National Knowledge Infrastructure (CNKI), and the Wanfang database.

Main outcome(s): The main outcome will be measured in terms of sensitivity and the specificity of the diagnostic test of interest. Sensitivity refers to the probability that the index test result will be positive in an infected case. Specificity refers to the probability that the index test result will be negative in a non-infected case.

Data management: Yanqin Shen and Guocan Yu will conduct the search strategies. No language restrictions in our search process. Primary search records will be imported into ENDNOTE X9.2 literature management software, according to eligibility criteria. Two investigators (Yanqin Shen and Guocan Yu) will independently assess the candidate articles by reviewing their titles and abstracts, followed by the full text, for inclusion. Discrepancies between the two investigators will be resolved by discussion with a third investigator (Yazhen Lang). We
will extract data including first author name; publication year; country; TP, FP, FN, and TN values for the assay; reference standard; patient selection method; specimen type; some steps (e.g., homogenization); and condition along with other parameters. The same two investigators will independently extract the necessary information from each of the included articles; we will cross-check the information they obtained. Discrepancies in the two data sets will be settled by a discussion with a third investigator, similar to that used during the literature selection phase. Data from studies against two different reference standards will be treated separately.

Quality assessment / Risk of bias analysis: Based on the two reference standards (CRS and culture), the two investigators will independently divide the studies into two groups and used a revised tool for Quality Assessment of Diagnostic Accuracy Studies (QUADAS-2) to assess study quality separately and the discrepancy between reviewers will be solved by discussion with a third investigator (Yazhen Lang). According to the PRISMA-DTA statement, systematic review and meta-analysis of diagnostic test accuracy studies was not required to assess publication bias.

Strategy of data synthesis: We will first obtain the values corresponding to TP, FP, FN, and TN in each included study, and calculated the estimated pooled sensitivity and specificity of Xpert MTB/RIF Ultra associated with the 95% confidence interval (CI), against CRS or culture, using bivariate random-effects models. Forest plots for sensitivity and specificity will be generated for each study. The areas under summary receiver operating characteristic (SROC) curves (AUC) will be subsequently calculated. I² statistics will be used to assess heterogeneity between the studies and a reference standard. While 0% will indicate no observed heterogeneity, values greater than 50% will be considered to imply substantial heterogeneity. We will explore different types of samples, different patient selection method, decontamination methods, sample conditions, and homogenization as potential sources of heterogeneity, using subgroup and meta-regression analyses. At least four published studies will be required to perform the meta-analysis for predefined variable types. Data from studies against CRS and culture will be analyzed separately. Stata version 15.0 (Stata Corp., College Station, TX, USA) with the midas command packages will be used to generate forest plots of sensitivity and specificity with 95% CI for each study and carry out meta-analyses and meta-regression analyses.

Subgroup analysis: Sensitivity analysis is used to explore the source of heterogeneity when the heterogeneity is obvious.

Sensibility analysis: Sensitivity analysis is used to explore the source of heterogeneity when the heterogeneity is obvious.

Language: No restriction.

Country(ies) involved: China.

Other relevant information: The strength of the body of evidence will be assessed using The Grading of Recommendations Assessment, Development and Evaluation (GRADE) guideline.

Keywords: Diagnostic accuracy, Xpert Ultra, tuberculous meningitis, meta-analysis.

Contributions of each author:
Author 1 - Yanqin Shen - The author drafted the manuscript, searched databases, selected literatures, managed data and assessed quality.
Author 2 - Guocan Yu - The author searched databases, selected literatures, managed data and evaluated quality.
Author 3 - Wuchen Zhao - The author drafted and revised the manuscript.
Author 4 - Yazhen Lang - The author provided statistical expertise, read, feedback and approved the final manuscript.