

Title:

Quality Systems Specialist

## **CERTIFICATE OF ANALYSIS**

This certificate is provided to confirm that the diagnostic kit specified below conforms to the production and testing specifications required by Cepheid's Quality System, in compliance with the US Food and Drug Administration's Quality System Requirements, ISO 13485, European IVD Directive and the Canadian Medical Devices Regulations (CMDR).

Product Name: Xpert	® MTB/RIF					
Cepheid Catalogue Par	rt No.: GXMTB/R	IF-US-10				
Kit Lot No.: 10014538	894					
Cartridge Lot No.: 44	702					
Kit Expiration Date: 2	2026-10-25					
Legal Manufacturer		Manufacturing Facility				
Cepheid		Cepheid	0	Solna	0	Sunnyva
904 Caribbean Drive	TICA	121 N Guild Avenue	_			
Sunnyvale, CA 94089	USA	Lodi, CA 95240		Lodi		
		USA		Loui		
Test Description		Acceptance Criteria		Test Re		
Positive		cted; Rif Resistance not detected		Passe	ed	
Positive Negative	MTB Detec	cted; Rif Resistance not detected  MTB not detected		Passe	ed ed	
Positive Negative	MTB Detec	eted; Rif Resistance not detected  MTB not detected  electronically and therefore valid	without a	Passe	ed ed	
Positive Negative  If checked, this docu	MTB Detection	eted; Rif Resistance not detected  MTB not detected  electronically and therefore valid	without a	Passe	ed ed	
Positive Negative	MTB Detection	eted; Rif Resistance not detected  MTB not detected  electronically and therefore valid	without a	Passe	ed ed	
Positive Negative  If checked, this docu	MTB Detection	eted; Rif Resistance not detected  MTB not detected  electronically and therefore valid	without a	Passe	ed ed	