

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 Devices Regulations.

C	ian Devices Regulations.	, European IVD
Product Name:	Xpert® Xpress CoV-2 plus	
Cepheid Catalogue Par	rt No.: XP3SARS-COV2-10	
Kit Lot No.: 10014	149529	
Cartridge Lot No.:	16504	
Kit Expiration Date:	2025-09-21	
Legal Manufacturer Cepheid 904 Caribbean Drive Sunnyvale, CA 94089 U	Cepheid AB Röntgenvägen 5	nnyvale (Newark () Lodi
Test Description	Acceptance Criteria	Test Result
Negative	SARS-CoV-2 NEGATIVE	Passed
Positive	SARS-CoV-2 POSITIVE	Passed
Robert Fiedler Robert Fiedler (Oct 7, 2024 15:49 GMT+2)	ument is produced electronically and valid without a wo	et signature.
Signature of Quality A	Assurance, Date	

Name: Robert Fiedler

Title: QA Analyst