

## DECLARATION OF CONFORMITY

Manufacturer: Illumina  
5200 Illumina Way  
San Diego, CA 92122  
United States

European Authorized Representative: Illumina Netherlands B. V.  
Freddy van Riemsdijkweg 15  
5657 EE Eindhoven  
The Netherlands

Device Name: **NextSeq 550Dx Instrument**

Device Model/Catalogue Number: **20005715**

Basic UDI-DI: 0081627002NEXTSEQAD


Classification: General IVD

Conformity Assessment Procedure: Annex III of IVDD 98/79/EC Council Directive; Self-Declaration

We, Illumina, declare under our sole responsibility that the *in vitro* Diagnostic Device(s) categories specified above and provided with the CE marking, meet the provisions of the EC Directive 98/79/EC (including amendments issued in the years following) which apply to them.

This declaration is supported by the EC Quality System Certificate(s) according to the provisions of relevant Annex(es) of this Directive. This declaration applies to all devices specified within this Declaration of Conformity and distributed onwards from the signature date below.

Authorized by:



Bryan Schneider  
Associate Director, Regulatory Affairs - HQ

22-APR-2020

Date (DD-MMM-YYYY)