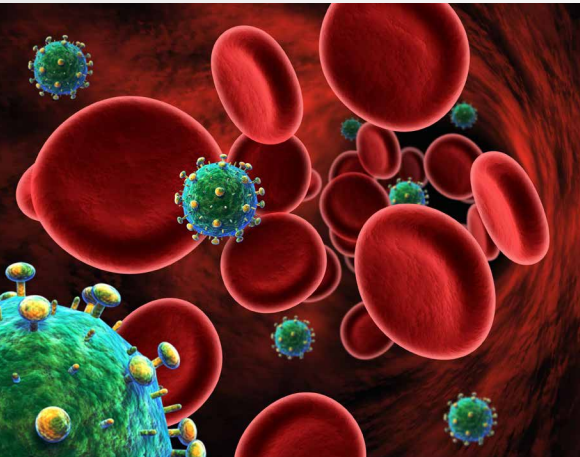




# GeneProof

## GeneProof HIV type 1 (HIV-1) PCR Kit



### HIGH DIAGNOSTIC SPECIFICITY AND SENSITIVITY

- Diagnostic sensitivity was verified on 128 HIV positive plasma samples acquired from Centre for AIDS Reagents (CFAR), National Institute for Biological Standards and Control (NIBSC)
- Diagnostic specificity was verified on 500 HIV negative samples

### DUAL TARGET DETECTION

- Dual targeting prevents detection failure caused by possible mutations inside the HIV-1 genome

### DETECTS A BROAD SPECTRUM OF HIV-1 SUBTYPES

- HIV genotypes A - D, AE, F, AG-GH, Group N, Group O, BF, H, K, CRF03\_AB

### SINGLE TUBE READY-TO-USE MASTER MIX

- Contains all components for PCR amplification
- No additional PCR reagents pipetting necessary
- Reduces the need for qualified laboratory staff
- Ensures reproducibility of the result

### CONTROL OF THE WHOLE DIAGNOSTIC PROCESS

- RNA extraction, reverse transcription and PCR amplification



COMPATIBLE WITH A WIDE  
RANGE OF REAL-TIME PCR  
DEVICES



CERTIFIED  
DIAGNOSTIC TEST



# GeneProof HIV type 1 (HIV-1) PCR Kit

+ GeneProof Hepatitis C Virus (HCV) Diagnostic PCR Kit

+ Hepatitis C Virus (HCV) PCR Kit  
+ HIV type 1 (HIV-1) PCR Kit

+ Hepatitis B Virus (HBV) PCR Kit

<b>TECHNOLOGY</b>	real-time PCR
<b>TYPE OF ANALYSIS</b>	qualitative and quantitative
<b>TARGET SEQUENCE</b>	LTR sequence and GaG gene
<b>ANALYTICAL SPECIFICITY</b>	HIV genotypes A - D, AE, F, AG-GH, Group N, Group O, BF, H, K, CRF03_AB, 100%
<b>ANALYTICAL SENSITIVITY (LOD)</b>	reaches up to 273.971 IU/ml i.e. 153.424 cp/ml with the probability of 95 % (on HIV-1 NIBSC 16/194 using manual extraction GeneProof PathogenFree RNA Isolation Kit) reaches up to 548.121 IU/ml i.e. 306.948 cp/ml with the probability of 95 % (on HIV-1 NIBSC 16/194 using automatic extractor croBEE NA16 Nucleic Acid Extraction System) reaches up to 98.59 IU/ml i.e. 55.21 cp/ml with the probability of 95 % (on Acrometrix HIV-1 Panel IU/ml using manual extraction SpinStar Viral Nucleic Acid Kit 1.0 with SpinStar Pretreatment Solution)
<b>DIAGNOSTIC SPECIFICITY</b>	100 % (CI95 % : 99.10 % - 100 %)
<b>DIAGNOSTIC SENSITIVITY</b>	93.66 % (CI95 % : 87.96 % - 96.88 %)
<b>LINEAR RANGE</b>	10 <sup>9</sup> - 10 <sup>2.5</sup> IU/ml with precision of ± 0.5 log (using manual extraction GeneProof PathogenFree RNA Isolation Kit) 10 <sup>9</sup> - 103 IU/ml with precision of ± 0.5 log (using automatic extractor croBEE NA16 Nucleic Acid Extraction System)
<b>DYNAMIC RANGE</b>	10 <sup>9</sup> - 273.971 IU/ml (using manual extraction GeneProof PathogenFree RNA Isolation Kit) 10 <sup>9</sup> - 548.121 IU/ml (using automatic extractor croBEE NA16 Nucleic Acid Extraction System)
<b>REPORTING UNITS</b>	IU/μl
<b>CONVERSION FACTOR</b>	1 IU = 0.56 cp
<b>METROLOGICAL TRACEABILITY</b>	HIV NIBSC 16/194
<b>EXTRACTION/INHIBITION CONTROL</b>	PCR inhibition and RNA extraction efficiency control (ISEX version)
<b>VALIDATED SPECIMEN</b>	plasma
<b>STORAGE</b>	-20 ± 5 °C
<b>VALIDATED EXTRACTION METHODS</b>	croBEE NA16 Nucleic Acid Extraction System GeneProof PathogenFree RNA Isolation Kit
<b>INSTRUMENTS</b>	croBEE Real-Time PCR System* Applied Biosystems 7300 / 7500 Real-Time PCR System CFX Connect™ / CFX96™/ Dx Real-Time PCR Detection System LightCycler® 480 LineGene 9600 Plus Mic qPCR Cyclers QuantStudio™ 3/5 Real-Time PCR System SLAN® Real-Time PCR System
<b>REQUIRED DETECTION CHANNELS</b>	FAM, HEX
<b>EXTERNAL QUALITY ASSESSMENT</b>	Regularly tested by QCMD and Instand e.V. External Quality Assessment Panels - results at <a href="http://www.geneproof.com">www.geneproof.com</a>
<b>REGULATORY STATUS</b>	CE <sub>1023</sub> IVD

\*Validated instruments

PRODUCT NAME	TECHNOLOGY	ORDER NO.		
		25 REACTIONS	50 REACTIONS	100 REACTIONS
GeneProof HIV type 1 (HIV-1) PCR Kit	real-time PCR	HIV1/ISEX/025	HIV1/ISEX/050	HIV1/ISEX/100