



Declaration of Conformity

Biotage Sweden AB, Box 8, SE-751 03 Uppsala, Sweden

declare under our sole responsibility that the product

Biotage® Extrahera™ LV-200

P/N: 417000

to which this declaration relates are in conformity with the following directives, standards and other normative documents;

Directive 2014/35/EU

Standards used:

Low Voltage Directive (LVD)

EN 61010-1:2010/A1:2019
EN/IEC 61010-2-081:2020

Directive 2014/30/EU

Standards used:

Electro Magnetic Compatibility (EMC)

EN/IEC 61326-1:2013

Directive 2012/19/EU

Standard used:

WEEE

EN 50419:2006

**Directive 2011/65/EU +
2015/863/EU**

Standard used:

RoHS

EN 63000:2018

For Canada and the USA

UL 61010-1:2012 /R:2019-07
UL 61010-2-081:2019

IEC 61010-1:2010, IEC 61010-1:2010/AMD1:2016
IEC 61010-2-081:2019

CAN/CSA-C22.2 No. 61010-1:2012/A1:2018-11
CAN/CSA-C22.2 No. 61010-2-081:2019

CFR 47 FCC part 15 Subpart B. Class A

NTRL Certificate No. US 071091 0039 Rev.00 issued by TÜV SÜD Product Service GmbH

CB Certificate No. DE 3 – 31486 issued by TÜV SÜD Product Service GmbH

Uppsala, 28 April 2021


.....
(Jon Sverre Schanche, Chief Scientific Officer - Biotage)