

Our STN: BL 125792/0 BLA APPROVAL October 17, 2023

Roche Diagnostics Attention: Bin Sun 9115 Hague Road Indianapolis, IN 46250

Dear Bin Sun:

Please refer to your Biologics License Application (BLA) received December 21, 2022 submitted under section 351(a) of the Public Health Service Act (PHS Act) for Elecsys HTLV-I/II.

#### LICENSING

We have approved your BLA for Elecsys HTLV-I/II effective this date. You are hereby authorized to introduce or deliver for introduction into interstate commerce, Elecsys HTLV-I/II under your existing Department of Health and Human Services U.S. License No. 2305.

The Elecsys HTLV-I/II is an in vitro immunoassay for the qualitative detection of antibodies to HTLV-I and HTLV-II in human serum and plasma. Elecsys HTLV-I/II is intended to screen individual human donors, including volunteer donors of whole blood and blood components. The assay is also intended to be used to screen organ, tissue and cell donors, when donor samples are obtained while the donor's heart is still beating. It is not intended for use on cord blood specimens.

The **e**lectro**c**hemi**l**uminescence **i**mmuno**a**ssay "ECLIA" is intended for use with **cobas pro** serology solution equipped with **cobas e** 801 analytical unit.

#### MANUFACTURING LOCATIONS

Under this license, you are approved to manufacture Elecsys HTLV-I/II at your facilities located at (b) (4) Manheim, Germany. You may label your product with the proprietary name Elecsys HTLV-I/II and will market it as approved in your license application.

## **ADVISORY COMMITTEE**

We did not refer your application to the Blood Products Advisory Committee because our review of information submitted in your BLA, including the clinical study design and trial results, did not raise concerns or controversial issues that would have benefited from an advisory committee discussion.

## **DATING PERIOD**

The dating period for Elecsys HTLV-I/II shall be up to 15 months from the date of manufacturing when stored at 2-8°C unopened or for 16 weeks when stored on-board the instrument at (b) (4). The manufacturing date shall be defined in accordance with 21 CFR 610.50.

#### FDA LOT RELEASE

Blind-coded panels will be provided for confirming lot release testing for all manufactured lots. The testing will be performed at Roche Diagnostics. The results of the coded samples will be forwarded to the Division of Biological Standards and Quality Control (DBSQC) through the Center for Biological Evaluation and Review (CBER) Sample Custodian as a component of the Lot Release Protocol. You may not distribute any lots of product until you receive a notification of release from the Director, CBER.

## **BIOLOGICAL PRODUCT DEVIATIONS**

You must submit reports of biological product deviations under 21 CFR 600.14. You should identify and investigate all manufacturing deviations promptly, including those associated with processing, testing, packaging, labeling, storage, holding and distribution. If the deviation involves a distributed product, may affect the safety, purity, or potency of the product, and meets the other criteria in the regulation, you must submit a report on Form FDA 3486 to the Director, Office of Compliance and Biologics Quality, electronically through the eBPDR web application or at the address below. Links for the instructions on completing the electronic form (eBPDR) may be found on CBER's web site at <a href="https://www.fda.gov/vaccines-blood-biologics/report-problem-center-biologics-evaluation-research/biological-product-deviations">https://www.fda.gov/vaccines-blood-biologics/report-problem-center-biologics-evaluation-research/biological-product-deviations</a>:

Food and Drug Administration
Center for Biologics Evaluation and Research
Document Control Center
10903 New Hampshire Ave.
WO71-G112
Silver Spring, MD 20993-0002

## **MANUFACTURING CHANGES**

You must submit information to your BLA for our review and written approval under 21 CFR 601.12 for any changes in, including but not limited to, the manufacturing, testing, packaging or labeling of Elecsys HTLV-I/II, or in the manufacturing facilities.

## **LABELING**

We hereby approve the draft package insert labeling, carton and container labeling submitted under amendment # 18 dated September 29, 2023. This is a reminder that as of September 24, 2014, medical devices that are licensed under the PHS Act are subject to certain provisions of the final Unique Device Identifier (UDI) rule. These provisions include the requirement to provide a UDI on the device label and packages (21 CFR 801.20), format dates on the device label in accordance with 21 CFR 801.18, and submit data to the Global Unique Device Identification Database (GUDID) (21 CFR 830 Subpart E). Additionally, please identify each device identifier implemented for the subject device, and the device identifiers that have been discontinued for the subject device as a labeling change in an annual report consistent with 21 CFR 601.12(f)(3). For more information on these requirements, please see the UDI website at <a href="http://www.fda.gov/udi">http://www.fda.gov/udi</a>.

Please submit all final printed labeling as a PDF electronic copy (eCopy) at the time of use and include implementation information on Form FDA 356h as appropriate.

Two draft copies of the proposed introductory advertising or promotional labeling may be voluntarily submitted for advisory comment with a completed Form FDA 2253 to the Advertising and Promotional Labeling Branch at the following address:

Food and Drug Administration
Center for Biologics Evaluation and Research
Document Control Center
10903 New Hampshire Ave.
WO71-G112
Silver Spring, MD 20993-0002

## **ADVERSE EVENT REPORTING**

You must submit adverse experience reports in accordance with the Medical Device Reporting (MDR) requirements for medical devices (21 CFR 803) as required by 21 CFR 600.80(m)(2). Because your product is characterized as a device as well as a biologic, submit these reports, listing device product code QHM, to the MedWatch System using MedWatch Reporting Form 3500A or an electronic equivalent. Please refer to the *Questions and Answers about eMDR – Electronic Medical Device Reporting – Guidance for Industry, User Facilities and FDA Staff* at <a href="https://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm175805.htm">https://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm175805.htm</a>.

# Page 4 - STN BL 125792/0 - Bin Sun

Required reports are to be submitted to:

Food and Drug Administration Center for Devices and Radiological Health MDR Policy Branch 10903 New Hampshire Avenue WO Bldg. 66, Room 3217 Silver Spring, MD 20993-0002

Sincerely,

Anne Eder, MD, PhD Acting Director Office of Blood Research and Review Center for Biologics Evaluation and Research