BIOPHEN™ Edoxaban Control Low

REF 225401-RUO CI CII 6 vials x 1 mL

BIOPHEN™ Edoxaban Control Plasma

REF 225501-RUO C1 C2 6 vials x 1 mL

FOR RESEARCH USE ONLY. DO NOT USE IN DIAGNOSTIC PROCEDURES.

INTENDED USE:

For quality control of Edoxaban assays, using a quantitative automated method. This kit is for research use only and must not be used for patient diagnosis or treatment.

SUMMARY AND EXPLANATION:

Technical:

These controls are proposed for the quality control of anti-Xa chromogenic assays of Edoxaban in plasma (BIOPHENTM DiXal and BIOPHENTM Heparin LRT, low range / standard range).

REAGENTS:

CI Lyophilized human plasma containing approximately 25 ng/mL of Edoxaban.

CII Lyophilized human plasma containing approximately 80 ng/mL of Edoxaban.

C1 Lyophilized human plasma containing approximately 150 ng/mL of Edoxaban.

C2 Lyophilized human plasma containing approximately 300 ng/mL of Edoxaban.

Control plasmas contain stabilizing agents.

The control concentrations may vary slightly from one batch to another. For the assay, see the exact values indicated on the flyer provided with the kit used.

The product is classified as non-hazardous and is not subject to labeling according to EC Regulation No. 1272/2008 [CLP].

WARNINGS AND PRECAUTIONS:

- Some reagents provided in these kits contain materials of human origin. Whenever human plasma is required for the preparation of these reagents, approved methods are used to test the plasma for the antibodies to HIV 1, HIV 2 and HCV, and for hepatitis B surface antigen, and results are found to be negative. However, no test method can offer complete assurance that infectious agents are absent. Therefore, users of reagents of these types must exercise extreme care in full compliance with safety precautions in the manipulation of these biological materials as if they were infectious.
- Waste should be disposed of in accordance with applicable local regulations.

REAGENT PREPARATION:

Gently remove the freeze-drying stopper, to avoid any product loss when opening the vial.

CI CII C1 C2 Reconstitute the contents of each vial with exactly 1 mL of distilled water.

Shake vigorously until complete dissolution while avoiding formation of foam and load it directly on the analyzer following Application Guide instruction. For manual method, allow to stabilize for 10 minutes at room temperature (18-

25°C), homogenize before use.

This plasmatic reagent can be more or less turbid after reconstitution. This turbidity is mainly due to plasma lipids that, after freeze-drying, become "less" soluble and may form a slight deposit. If necessary, let each vial stabilize 10 minutes at room temperature and shake before use.

STORAGE AND STABILITY:

Unopened reagents should be stored at 2-8°C in their original packaging. Under these conditions, they can be used until the expiry date printed on the kit.

CI CII C1 C2 Reagent stability after reconstitution, free from any contamination or evaporation, and stored closed, is of:

- 7 davs at 2-8°C. ٠
- 60 days frozen at -20°C or less*
- Stability on board of the analyzer: see the specific Application Guide. *Thaw only once, as rapidly as possible at 37°C and use immediately.

REAGENTS AND MATERIALS REQUIRED BUT NOT PROVIDED:

Laboratory material.

TRACEABILITY:

website:

Controls are traceable to internal standard of reference using the LC-MS/MS reference measurement procedure for Edoxaban. Certificate of traceability and uncertainty is available on the HYPHEN BioMed

HYPHEN BioMed

155 rue d'Eragny, 95000 Neuville-sur-Oise, France

155 rue d'Eragny, 95000 Neuville-sur-Oise, France

Tél:+33 (0)1 34 40 65 10 Fax: +33 (0)1 34 48 72 36 www.hyphen-biomed.com info@hyphen-biomed.com

English, revision: 08-2023

Uncertainty			
CI	± 3.5 ng/mL	C1	± 10 ng/mL
CII	± 3.9 ng/mL	C2	± 10 ng/mL

QUALITY CONTROL:

For quality control of Edoxaban assays by chromogenic methods (low range or standard range), with BIOPHEN™ Heparin LRT (221011-RUO, 221013-RUO and 221015-RUO) and BIOPHEN™ DiXal (221030-RUO) kits.

The target values are determined from multi-reagent and multi-instrument tests. The use of quality controls serves to validate method compliance, along with between-series assay homogeneity for a given batch of reagents.

Include the quality controls with each series, as per good laboratory practice, in order to validate the test.

If the controls fall outside of the acceptance range, the series of assays must be invalidated and the analyses repeated. Check all system parameters before repeating the series.

LIMITATIONS:

- If the controls are used under measurement conditions other than those validated by HYPHEN BioMed, the test results may vary. The laboratory is responsible for validating the use of these controls in its own analytical system.
- · Any reagent presenting no limpid appearance or showing signs of contamination must be rejected.

The results obtained should be for research use only and must not be used for patient diagnosis or treatment.

REFERENCES:

- Bathala MS et al. Pharmacokinetics, biotransformation, and mass balance of edoxaban, a selective, direct factor Xa inhibitor, in humans. Drug Metab Dispos. 2012.
- 2. Bounameaux H and Camm AJ. Edoxaban: an update on the new oral direct factor Xa inhibitor. Drugs. 2014.
- Furugohri T et al. DU-176b, a potent and orally active factor Xa inhibitor: in vitro and in vivo pharmacological profiles. J Thromb Haemost. 2008.
 Honda Y and Morishima Y. Thrombin generation induced by tissue factor plus ADP in human platelet rich plasma: A potential new measurement to assess the effect of the substitute and end and the rate of the substitute the rate of a card factor Xa inhibitor advancement and an advancement to assess the affect of the substitute the rate of a card factor Xa inhibitor advancement and advancement to assess the advancement of the substitute the rate of the substitute that and the substitute the rate of the substitute that and the substitute the substitute the substitute the substitute the substitute the concomitant use of an oral factor Xa inhibitor edoxaban and P2Y12 receptor antagonists. Thromb Res. 2015

For customer support or Application Guides, please contact your local provider or distributor (see www.hyphen-biomed.com).

Changes compared to the previous version.

The following symbols may appear on the product labeling:

Product for in-vitro REF Catalogue numbe LOT Batch code RUO research use, only WHO Numerical < x> identification of standard WHO STD Rx i See instructions for use reagent code Temperature limitation Manufacturer Use by Å CONTENTS Ø Reconstitution volume Biological risks Contents See instructions in Method Application Numerical < x> identification of **1**- MA Cx CONTAINS Contains control auide Σ Contains sufficient for EXP Expiration date UNIT Measurement unit Numerical < x> Keep away from TARGET VALUE Target Value CALx identification of sunlight and heat calibrator Contains biological Contains human BIC ۷ UDI Unique Device Identifier material of animal blood or plasma origin derivatives ACCEPTANCE RANGE Acceptance range DANGER Danger WARNING Warning

