

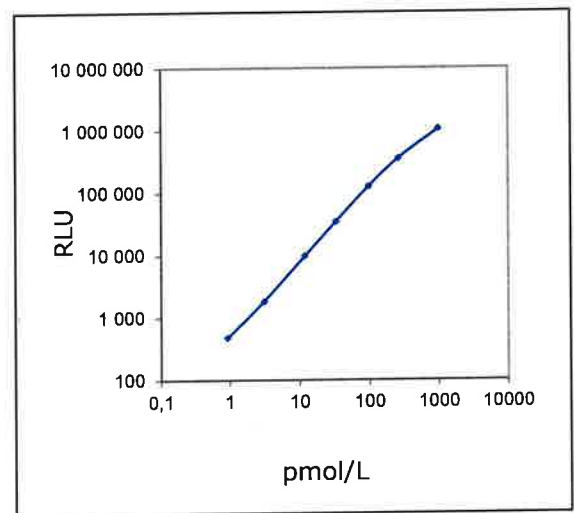
Certificate of Analysis

1. Manufacturer

Merckodia AB, Sylveniusgatan 8A, 754 50 Uppsala, SWEDEN

2. Description

Catalog no: 10-1278-01
 Product: Merckodia Total GLP-1 NL-ELISA
 Lot no: 37246
 Expiry date: 2028-01-31



<i>Component</i>	<i>Art no</i>	<i>Lot no</i>	<i>RLU</i>	<i>Exp. date</i>
Calibrator 0	20-7259	36499	112	2028-02-14
Calibrator 0,928 pmol/L	20-7266	36492	481	2028-03-14
Calibrator 3,09 pmol/L	20-7267	36493	1837	2028-03-14
Calibrator 11,8 pmol/L	20-7268	36494	9784	2028-03-14
Calibrator 32,8 pmol/L	20-7269	36495	34763	2028-03-14
Calibrator 97,1 pmol/L	20-7270	36496	127266	2028-03-14
Calibrator 261 pmol/L	20-7271	36497	358579	2028-03-14
Calibrator 984 pmol/L	20-7299	36498	1052923	2028-03-14
Coated Plate	20-7277	37188		2029-04-13
Enzyme Conjugate 11X	20-7275	37190		2029-04-14
Enzyme Conjugate Buffer	20-7273	36500		2028-03-20
Wash Buffer 21X	20-6746	36958		2032-12-03
Substrate Reagent A	20-7300	36595		2028-02-14
Substrate Reagent B	20-7301	36596		2028-02-14

3. Quality control

Quality control has been performed for lot no 37246 according to standard operating procedures at Mercodia AB, and the product is released based on fulfillment of established acceptance criteria.

4. Calibration

Mercodia Total GLP-1 NL-ELISA is calibrated against an in-house reference preparation of GLP-1 (9-36)amide.

5. Assay method

Test procedure used is according to current Direction for Use for the product and lot.

6. Intended use

Mercodia Total GLP-1 ELISA provides a method for the quantitative determination of Total GLP-1 in EDTA-plasma.

7. Storage and handling

Recommended storage of kit is 2-8°C.

Storage of unused or diluted kit components is stated in the Direction for Use.

8. Hazardous information

Please refer to the Safety Data Sheet for hazard identification.

9. Quality standard documentation

The Mercodia Quality System fulfills the QSR and the requirements for the European CE mark, the Directive 98/79/CE on *in vitro* diagnostic medical devices. Mercodia is ISO 13485 certified, for compliance with the International Quality Management System for medical devices.

10. Names and signatures of certifying officers

Date of analysis:

2026-05-12

Performed by:

Elin Westberg

Signature:

EW

Date of approval:

2026-05-20

Approved by:

Jonas Krick

Signature:

JK