

AUG 10, 2023

Measurement of GLP-1 release in cell supernatant from Hutu-80 enteroendocrine cells via ELISA

r.mezabrovschi¹, rachel.bates¹

¹University College London, Aligning Science Across Parkinson's (ASAP) Collaborative Research Network, Chevy Chase, MD, 20815.



Sofia Koletsi

University College London, University of London

DISCLAIMER

This protocol was adapted from Cat. # EGLP-35K to optimise the seeding density of the assay based on the cell line used (Hutu 80 enteroendocrine cells RRID: CVCL_1301), moreover, cell line used and media conditions used to characterise GLP-1release.

The lowest level of GLP-1 that can be detected by this assay is 2 pM (100ul plasma sample size).

OPEN ACCESS



DOI:

dx.doi.org/10.17504/protocol s.io.j8nlkokm6v5r/v1

Protocol Citation: r.mezabr ovschi, rachel.bates 2023. Measurement of GLP-1 release in cell supernatant from Hutu-80 enteroendocrine cells via ELISA . **protocols.io** https://dx.doi.org/10.17504/protocols.io.j8nlkokm6v5r/v1

License: This is an open access protocol distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

Protocol status: Working We use this protocol and it's working

Created: Jul 26, 2023

ABSTRACT

This protocol describes the method of use for the Glucagon-Like Peptide-1 (Active) ELISA Kit

96-Well Plate (Cat. # EGLP-35K) to measure GLP-1 release (pM) from supernatant. The assay should be run in duplicate.

MATERIALS

GLP-1 (Active) ELISA Plate

Coated with anti-GLP-1 Monoclonal Antibody Quantity: 1 plate

Preparation: Ready to use

Adhesive Plate Sealer

Quantity: 1 Sheet

Preparation: Ready to use

10X Wash Buffer Concentrate

10X concentrate of 10 mM PBS Buffer containing Tween 20 and Sodium Azide.

Quantity: 50 mL

Preparation: Dilute 1:10 with deionized water

GLP-1 (7-36) amide ELISA Standards

Last Modified: Aug 10,

2023

GLP-1 (7-36 amide) in Assay Buffer: 2, 5, 10, 20, 50 and 100 pM Quantity: 1 mL/vial

Preparation: Ready to use

PROTOCOL integer ID:

85525

ELISA GLP-1 (Active) Quality Controls 1 and 2

Various peptides including GLP-1 (7-36 amide) in QC Buffer. Quantity: 1 mL/vial

Preparation: Ready to use

GLP-1 (Active) Assay Buffer

0.05M PBS, pH 6.8, containing proprietary protease inhibitors, with Tween 20, 0.08%

Sodium Azide and 1% BSA.

Quantity: 25 mL

Preparation: Ready to use

GLP-1 (Active) Detection Conjugate

Anti GLP-1-Alkaline Phosphate Conjugate. Quantity: 21 mL

Preparation: Ready to use

Substrate (Light sensitive, avoid unnecessary exposure to light)

Quantity: 10 mg

Preparation: Hydrate in 1 mL deionized water just before use. Use at 1:200 dilution in substrate diluent (e.g. 100 uL hydrated substrate in 20 mL substrate diluent).

Dilute fresh each time just before use.

Substrate Diluent (Light sensitive, avoid unnecessary exposure to light)

Quantity: 21 mL

Preparation: Ready to use

Stop Solution

Quantity: 6 mL

Preparation: Bring to room temperature before use. Mix thoroughly to ensure no

precipitate remains.

Cell Culture Medium

Specific to cell type.

Sterile 24-well plates

Quantity: 2

PBS (1X)

Preparation: Ready to use

Reagents for you experimental media conditions

Varies between experiments.

DPP-1V Inhibitor

Quantity: 10ml (Cat. DPP4-010)

Preparation: Ready to use - to be purchased separately. (Store -20 degrees).

BEFORE START INSTRUCTIONS

All reagents should be warmed to room temperature before proceeding.

Day 1

Plate cells in sterile 24-well plates at 1.5x10^5 per well and leave Overnight in a 37 °C incubator with stable CO2 conditions overnight.

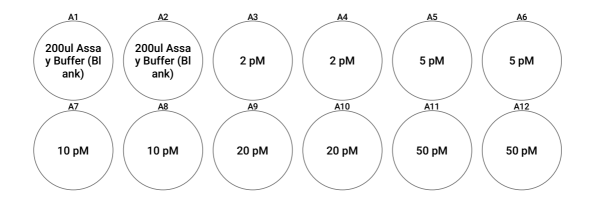
Day 2

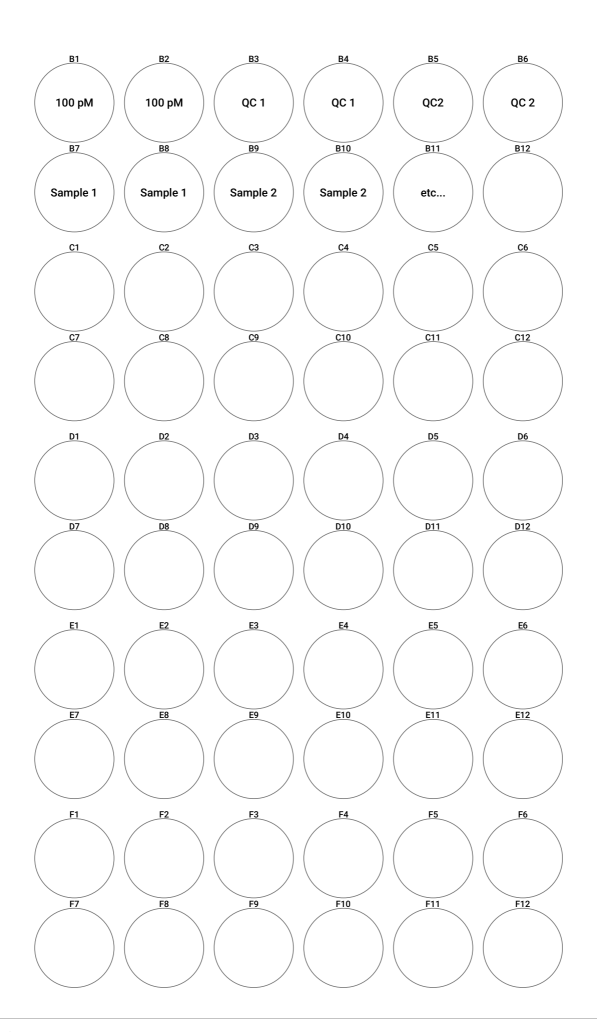
5m

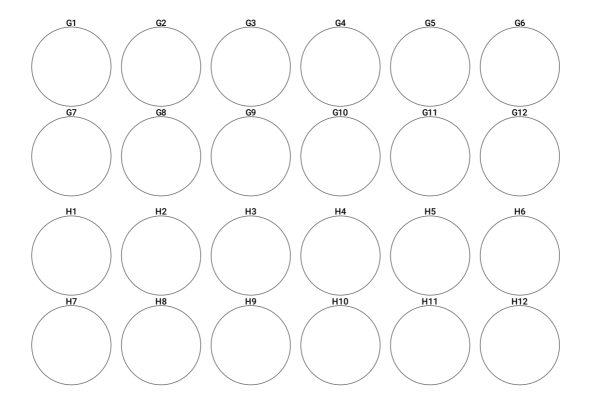
2h

- 2 Check cells are healthy under a microscope.
- In a laminar flow hood aspirate medium and wash with PBS 3 times, adding Δ 500 μ L each time.
- 4 Once the final PBS wash is complete add in Δ 500 μL of the test media made up for your desired experiment (e.g., high glucose media or 2-Deoxy-D-glucose) and place back in the incubator for ৩2:00:00 .
- Make up the **Wash Buffer** using the **10X Wash Buffer Concentrate** and dilute 1:10 with deionised water.
- Once the plate has finished incubating, move over the 4500 µL supernatant into a new sterile 24-well plate and add DPP-IV inhibitor immediately (1:100).to be carried out in a laminar flood hood. The cells may be stored at 45-80 °C for further analysis at a later date.

- In each well of the ELISA plate, add \square 300 μ L of diluted **Wash Buffer** (or in two intervals of \square 150 μ L). Incubate at \square Room temperature for \square 00:05:00 Decant excess buffer and blot with absorbent towels.
- 8 Add Δ 200 μL Assay Buffer to NSB (non-specific binding) wells A1, A2.
- 9 Add 🔼 100 µL Assay Buffer to the remaining wells you wish to load your samples in.
- 10 Add 🗸 100 µL standards in ascending order to wells standards come preprepared.
- Then load I 100 µL of QC1 and QC2 to the plate in seperate wells.







- 13 For good mixing, lightly agitate the plate.
- Cover the plate with plate sealer. Incubate Overnight (20 to 24 hours) at 4 °C .

Day 3

2h 35m

All reagents should be warmed to Room temperature before proceeding.

Decant liquid from plate and tap out excess fluid on absorbent towels.

Wash the plate 5 times with 300 µL pre-diluted Wash Buffer per well with 00:05:00 incubation at Room temperature in Wash Buffer with the fourth wash. Tap out excess buffer on absorbent towels after the fifth wash.

5m

- © 02:00:00 at Room temperature then decant.
- Whilst the plate is incubating dilute the **Substrate.** .
- 18.1 Hydrate in 🔼 1 mL deionized water just before use.
- Use at 1:200 dilution in **Substrate Diluent** (e.g. \bot 100 μ L hydrated substrate in \bot 20 mL substrate diluent). Dilute fresh each time just before use.
- Wash the wells 3 times with $200 \, \mu L$ diluted **Wash Buffer**. Tap out excess buffer on absorbent towels.
- Measure fluorescence on a plate reader at an excitation/emission wavelength of 360/460 every 00:05:00 for a minimum of 00:20:00 .
- If sufficient fluorochrome has been generated, add 50 µL Stop Solution (mix thoroughly to ensure no precipitate remains) to each well in the same order as the Substrate was added. Incubate 00:05:00 at 8 Room temperature in the dark to arrest phosphatase activity.
- Read plate on a fluorescence plate reader with an excitation/emission wavelength of 360/460.

25m

5m