

Standardized, fully automated neat plasma and Mag-Net enrichment workflows enabled by the Evotip Pure

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Highlights

Scalable high-throughput proteomics enabled by Evotip Pure integration on the Biomek i5 liquid handler.

Fully automated sample preparation of up to 576 samples in parallel.

Showcasing four sample preparation workflows for robust analysis of large clinically-relevant sample cohorts.

The need for standardized solutions

A selection of workflows directly integrated with Evotip Pure

As proteomics studies become larger and analyses become more routine, the demand for standardized and scalable solutions is increasing. To address this need, we introduce four fully automated workflows developed for the Biomek i5 Automated liquid handler, seamlessly integrated with Evosep technology to ensure efficient and reproducible end-to-end sample preparation.

Evotip Sample Loading

This workflow uses a specialized prototype sealing adapter enables automated desalting and loading on the Evotip for efficient sample storage prior to downstream LC-MS analysis.

Digestion workflow

The PAC digestion workflow leverages protein aggregation on magnetic beads, followed by direct on-bead digestion of reduced and alkylated protein lysate.

Neat Plasma workflow

Combining steps for reduction and alkylation, this workflow utilizes PAC digestion as a streamlined approach for processing large plasma sample cohorts.

Mag-Net Plasma workflow

For deep plasma proteomics, the Mag-Net workflow combines enrichment of membrane-bound vesicles with PAC digestion.

Designed for large-scale analysis

Enzyme and sample consumption is minimized with online Evotip loading, as peptides are efficiently captured and stored on Evotip Pure allowing utilization of close to 100% of the sample. With efficient deck space usage, 576 samples can be processed simultaneously, resulting in less than two minutes of total processing time per sample for each workflow.

Neat Plasma

Buffers & Beads

Shuffled plates

Mag-Net Plasma

Wash buffer

Trash

Sample tips 4

Sample tips 5

Sample tips 6

Waste

Sample tips 1

Sample tips 2

Sample tips 3

Enzymes

Solvent tips

Sealing mat

Heater/shaker

Magnet

Sample plates 1-6

Solvent A

Workflow dependent

Evotips 4

Evotips 5

Evotips 6

Lids 1-6 for sample plates

Propanol

Evotips 1

Evotips 2

Evotips 3

Average time per sample (minutes)

Plates

1

2

3

4

5

6

3 seconds

Evotip loading

PAC digestion

Neat Plasma

Mag-Net Plasma

Schematic representation of deck layout on the Biomek i5 and an overview of how much time on is spent on average per sample when 1-6 plates are processed with each workflow.

References

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2. Wu CC., Tsantilas KA., Park J., Plubell D., Naicker P., Govender I., Buthelezi S., Stoychev S., Jordaan J., Merrihew G., Huang E., Parker ED., Riffle M., Hoofnagle AN., MacCoss MJ (2023) Mag-Net: Rapid enrichment of membrane-bound particles enables high coverage quantitative analysis of the plasma proteome. BioRxiv., https://www.biorxiv.org/content/10.1101/2023.06.10.544439v1.full.pdf

Seamless integration with Evotip Pure technology

Versatility through modular sample preparation

Automated sample loading on Evotips is the cornerstone of this modular sample preparation strategy. As the final step in each workflow, it facilitates sample clean-up and stable storage on the Evotip until LC-MS analysis. The entire process is fully automated and can process up to 576 samples at the time. Each workflow includes an additional sample processing module for PAC digestion¹, neat plasma or Mag-Net² plasma respectively. These modules are interchangeable, allowing for easy adaptation to various proteomics applications.

Capture & Wash

20 min

Solubilization

15 min

Aggregation

20 min

Wash

10 min

Digestion

16 hours

Evotip loading

5 min

MagNet plasma workflow - 4 µl plasma input

Neat plasma workflow - 1 µl plasma input

PAC Digestion workflow - 1 µg sample input

Evotip loading - 20 µl

Automation without compromise

The combination of Evosep technology and the Biomek i5 liquid handler seamlessly integrates scalable sample preparation with high-throughput LC-MS analysis, ensuring high-quality and reproducible results. All four workflows exhibit excellent performance using the standard 200 SPD method on the timsTOF HT. Overnight digestion at 37°C with trypsin and Lys-C ensures high digestion efficiency. This approach results in deep proteome coverage with high precision for both HeLa and plasma samples.

Digestion efficiency (%)

LC-MS QC HeLa

PAC HeLa

Neat plasma

Mag-Net plasma

Missed cleavages

Zero

One

Two

Protein groups

LC-MS QC HeLa

PAC HeLa

Neat plasma

Mag-Net plasma

Protein group CVs (%)

LC-MS QC HeLa

PAC HeLa

Neat Plasma

Mag-Net Plasma

Comparison of the workflows carried out using 50 ng QC HeLa digest (Automated loading, n=96), 1 µg HeLa lysate (PAC, n=40), 1 µl plasma (Neat, n=40), and 4 µl plasma (Mag-Net, n=40), respectively. All digestions were carried out at 37°C overnight (16 hours). The figures showcase, digestion efficiency, mean identified proteins, and protein CVs, measured on the timsTOF HT.

Preservation of sample and cohort integrity

A single plate containing varying amounts of HeLa input and two types of blanks was processed using the PAC workflow. The consistent summed precursor intensity observed within conditions, along with the negligible intensity in the blanks, confirms the absence of cross-contamination during sample preparation or significant carryover during LC-MS analysis. Additionally, minimal inter plate variation was observed across six plates processed from 1 µg HeLa protein lysate using the PAC workflow and subsequently analyzing resulting 40% peptide digest using the 500 SPD method.

Summed precursor intensity

HeLa 1 µg, n=40

HeLa 500 ng, n=40

Workflow blank, n=8

LC-MS blank, n=8

Plate 1 vs. Plate 1

Plate 1 vs. Plate 4

Plate 1 vs. Plate 2

Plate 1 vs. Plate 5

Plate 1 vs. Plate 3

Plate 1 vs. Plate 6

Pearson R

1.00

0.90

0.80

0.70

Digestion efficiency (%)

Missed cleavages

Zero

One

Two

Summed precursor intensity for HeLa samples with two different input amounts and blanks. Heatmap visualizing Pearson correlation on precursor level for six Biomek i5 processed plates and digestion efficiency across each plate.

Outstanding plasma proteomics

Robust quantification

More than 500 unique proteins were identified from just 1 µl of neat plasma using the 200 SPD method. With the Mag-Net workflow, the dynamic range was successfully compressed yielding 2,700 unique protein identifications with the same 200 SPD method. Additionally, 40 proteins listed in Vesiclepedia were identified in the Mag-Net enriched plasma, with most showing CVs below 20%, validating the method. Plasma is a rich source of potential biomarkers for minimally invasive monitoring, making this biofluid crucial in clinical research. We examined three proteins; The abundant plasma protein (ALB), the EV marker (CD63), and a low abundance cancer biomarker (ALDOA). Both workflows result in good quantification with little variation in the measured abundance within each condition.

Dynamic range plot for neat plasma (n=40) and Mag-Net enriched plasma (n=40). A scatter plot showing all identified proteins across all 40 Mag-Net samples. The dashed line marks 20% CV and proteins listed in Vesiclepedia are highlighted with triangles. A jitter plot showcasing the abundance of ALB, CD63, and ALDOA across 40 replicates of Neat and Mag-Net enriched plasma.

EVOSEP

Evosep One is for General Laboratory Use