US HUPO 2024

BRUKER

Improved dia-PASEF isolation window schemes for proteomics measurements

<u>Shourjo Ghose1</u>, Markus Lubeck2, Stephanie Kaspar-Schoenefeld2, Christoph Krisp2, Andreas Schmidt2 Florian Busch2, Eduardo Carrascosa2, Oliver Raether2, Gary Kruppa3

1 Bruker Scientific LLC, Billerica, MA, 2 Bruker Daltonics GmbH & Co. KG, Bremen, Germany, 3 Bruker s.r.o., Brno, Czech Republic

Introduction

Data-Independent Acquisition (DIA) is widely used for proteomics as it usually outperforms Data-Dependent Acquisition (DDA) for protein identification and quantitation, due to its higher ion usage and reproducibility, resulting from a fixed scheme of rather broad isolation windows. This advantage can be further increased by combining it with trapped ion mobility separation (TIMS), as the additional separation dimension reduces complexity and the sequential elution of condensed ion packages from the TIMS device allows for even more efficient ion usage (dia-PASEF). The two-dimensional mass and mobility space enables method creation with extensively different window schemes.

Results



Methods

Tryptic digest from human cell line K562 (Promega) was diluted with 0.1 % formic acid to a final concentration of 100 ng/µl. To achieve a concentration representing digests of single cells the stock solution was further diluted in 4 steps to a final concentration of 0.250 ng/µl. To prevent peptide losses due to absorption to hydrophobic surfaces, 0.015% n-Dodecyl- β -D-Maltosid (DDM) was added to the dilution buffer. Peptide samples were separated on a nanoElute (Bruker) using 22-minute linear gradients from 5% to 35% acetonitrile in 0.1% formic acid at 250 nl/min. A 25 cm, 75 µm ID column with integrated nanospray emitter (Aurora Ultimate, IonOpticks) was used in combination with a CaptiveSpray ultra source. Data were acquired on a timsTOF Ultra mass spectrometer. dia-PASEF window schemes were initially calculated using the py_diAID1 software tool and further fine-tuned manually (Fig. 1 B). Data were processed with dia-NN 1.8.1 using a library with 560,000 precursors derived from a deeply fractionated K562 sample. All measurements were performed in triplicates; cross-run normalization was switched off.



Figure 2:

Average number of identified precursors and protein groups. dia-NN results from 0.250 ng K562 digest separated using a 22-minute gradient and a selection of the dia-PASEF window schemes and TIMS ramp time combinations described in Figure 2. Number of identified precursors (A) and protein groups (B) are shown. The same data were processed either solely with a library of 560,000 precursors or using an initial search with the same library together with data from a 5ng sample, resulting in a reduced library for the final 2nd -pass search (Match Between Runs, MBR). Precursor (C) and protein groups (D) dia-NN results from 100 ng K562 digest separated with a 22-minute gradient using window scheme and TIMS ramp combinations from Figure 2 with a higher selectivity (higher number of windows and shorter TIMS ramp times).



Figure 1:

considered.

dia-PASEF window schemes (3x8). During an ion mobility scan the quadrupole switches its isolation position so that 3 isolation windows are covered within a single TIMS scan, thus only 8 TIMS scans are required to cover all 24 windows. A) Scheme with a fixed 25 m/z isolation width for all windows. B) Individual isolation widths normalized to precursor density of a spectral library, optimized by the py_diAID software tool1. C) dia-PASEF cycle times for different window schemes in combination with varying TIMS scan/accumulation times. All schemes cover the same m/z-mobility range. A higher number of narrower windows will increase selectivity, but also result in a loss of overall sensitivity, as individual ions will be fragmented less frequently. For the detection of very low abundant ion species longer accumulation of ions prior to detection would be preferred over averaging a multiplicity of sparse signals. However, if the cycle time exceeds a certain

threshold, quantitation will be impaired by undersampling chromatographic peaks,

thus acquisition methods with a cycle time greater than 2s (~FWHM) were not

Conclusion

Highest sensitivity and depth: 5400 protein groups can be identified and quantified with >47,000 precursors from 0.250 ng cell digest; 8500 protein groups with >105,000 precursors from 100 ng.

TIMS enables efficient ion usage and selectivity due to pre-separation of precursor ions into condensed ion packages, that are fragmented consecutively.

A broad precursor range can be covered, while maintaining high acquisition speed without

1Skowronek et.al., Mol Cell Proteomics (2022) 21(9) 100279 https://doi.org/10.1016/j.mcpro.2022.100279









