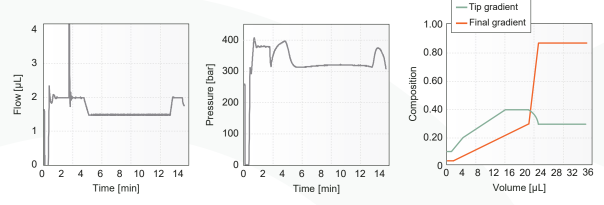
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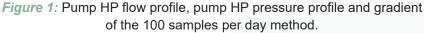
Towards a Standardized Omics Platform with the **100 samples per day** method

1. Introduction

This method has an 11.5 minute gradient and a cycle time of 14.4 minutes. The method is designed for our EV1064 Endurance column,

and we highly recommend to use this column with our range of emitters and spray adapters for optimal performance.

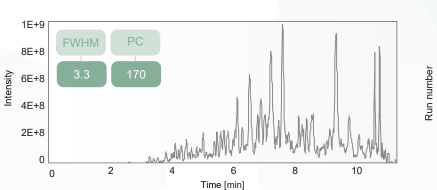




2. Chromatographic performance

By taking advantage of the gradient offset strategy, we maximize the chromatographic

performance. From a HeLa digest analysis, we calculated peak and retention time properties



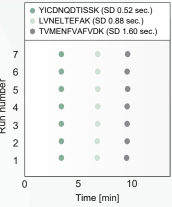


Figure 2: Example base peak chromatogram of 500 ng tryptic HeLa digest and retention time reproducibility of selected peptides across seven consecutive injections.

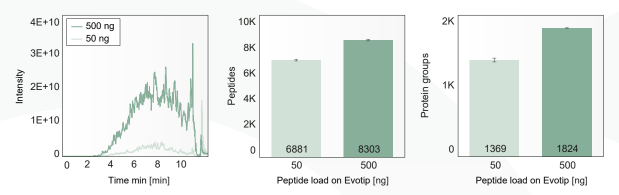


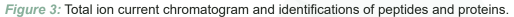
for the eluting peptides. On average, the full peak width at half maximum is 3.3 seconds, resulting in a peak capacity of 170 withing the elution window of 9.3 minutes. We monitored the retention time reproducibility of three

3. Method design

It is important to adjust the MS method to take full advantage of the condensed peptide elution. Therefore, we exploit the fastest scanning mode on an Orbitrap Exploris 480 MS. The MS instrument is operated in data-dependent acquisition mode using a Top12 method. Full MS resolution is set to 60,000 at m/z 200 and full MS AGC target value is 300% with an IT of 45 ms. Mass range is set to 350-1400. AGC target value for fragment spectra is set to 200% and the intensity thres- hold is kept at 2E5. Isolation width is set to 1.3 m/z and normalized different peptides over seven replica injections and find the average standard deviation to be only 1 second indicating highly reproducible retention times (Figure 2).

collision energy to 30%. Peptide match is set to off, and isotope exclusion to on. Former target ions are dynamically excluded for selection for 10 seconds. Higher-energy collision dissociation (HCD) fragment scans are acquired at 28 Hz speed with an injection time of 22 ms using an Orbitrap resolution of 15,000. Digested HeLa peptides are loaded on Evotips in two dilutions, 50 and 500 ng respectively. Technical quadruplicates are measured and analyzed with the Spectromine 2 software.





4. Results and Conclusion

The first peptides elute after 2.4 minutes with the recommended column. With a full peptide elution window of 9.3 minutes, the effective gradient elution is 81%. We identify 8300 peptides with the highest load of 500 ng corresponding to 1800 proteins. (Figure 3). This method is a great "in-between" method, which balances proteome coverage high-throughput with only 2.9 minutes overhead between injections.

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