# **P-I-0146**

# High-throughput high-resolution data-independent acquisition workflow on an Orbitrap Ascend **MultiOmics Tribrid mass spectrometer for accurate label-free quantitation**

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# Abstract

**Purpose**: To develop and assess qualitative and quantitative performance of label-free quantitation (LFQ) with an optimized data-independent acquisition (DIA) method on a Thermo Scientific<sup>™</sup> Orbitrap<sup>™</sup> Ascend Tribrid<sup>™</sup> mass spectrometer using a long (60 min active gradient), short (30 min active gradient) and ultra-high throughput (9 min active gradient) gradient for largescale proteomics analysis.

Methods: DIA workflow investigated in the present study was operated by directly injecting samples onto a 50-cm Thermo Scientific<sup>™</sup> µPAC<sup>™</sup> Neo HPLC column, and the peptides were resolved in a 60-min, 30-min and an ultra-high throughput 9-min gradient operated by a Thermo Scientific<sup>™</sup> Vanquish<sup>™</sup> Neo UHPLC system. The eluted peptides were analyzed on an Orbitrap Ascend Tribrid MS operated in DIA mode.

## Results

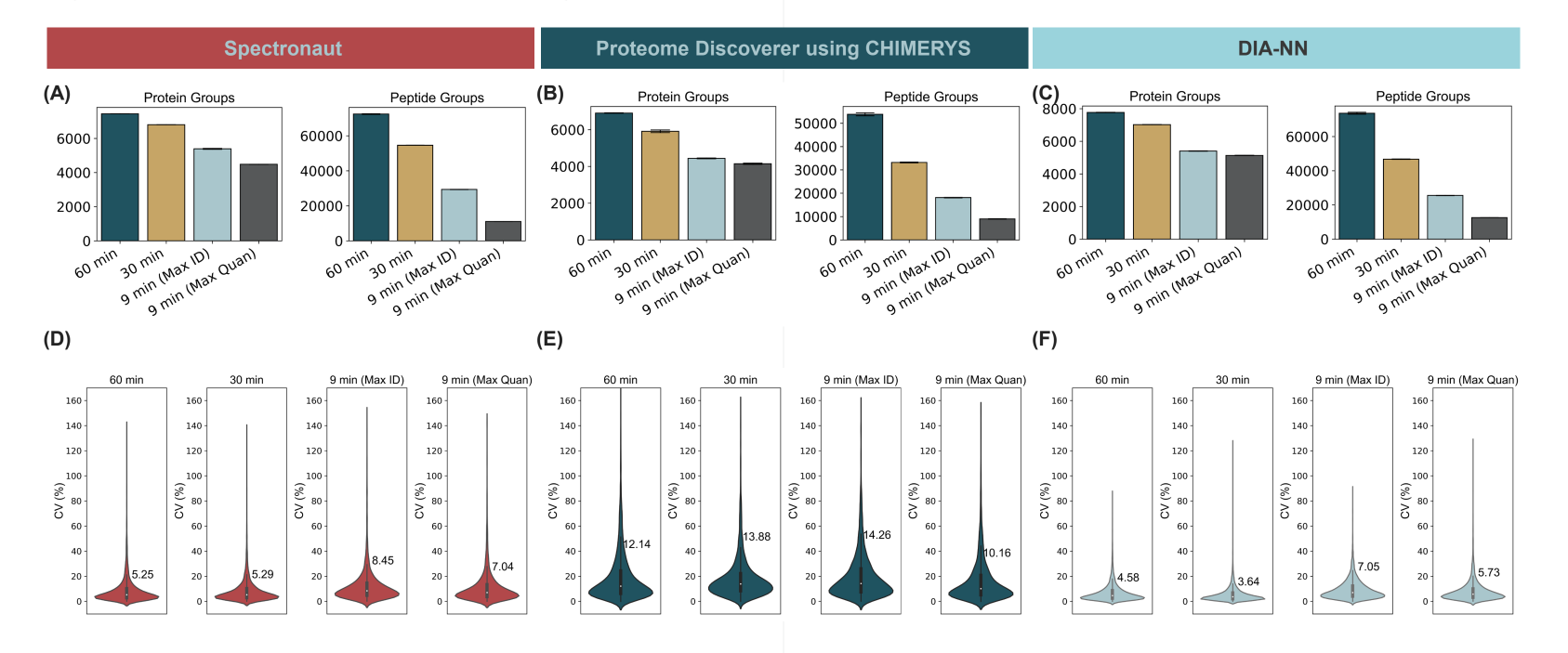
Velocity DIA workflow on Orbitrap Ascend Tribrid mass spectrometers allows for robust protein identification and quantitation

FAIMS Pro interface decreased peptide coverage but increased protein group identifications by an average of 5% (Figure 3). Consequently, the following experiments were all coupled with the FAIMS Pro interface.

With 30 min active gradient, 7,000+ proteins and 47,000+ peptides were identified, along with a protein group CV of approximately 5% (Figure 4), suggesting that the 30 min active gradient method enables relatively high throughput while maximizing identification and quantitative performance. We extended this workflow to 60 min active gradient and successfully identified close to 7,800 proteins and >76,000 peptides, highlighting that deeper proteome coverage can be achieved in the Velocity DIA workflow by using a longer gradient.

# Results

Figure 4. Determination of proteome coverage and quantitation precision of the human proteome.



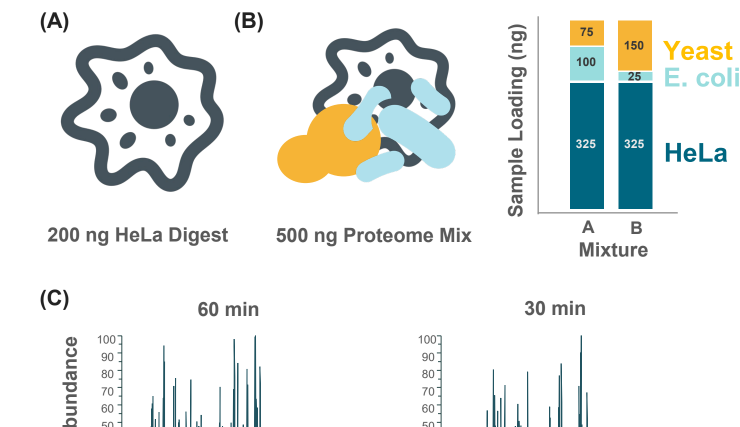
**Results**: Our results demonstrate that Velocity DIA presented herein is a robust and reproducible workflow on the Orbitrap Ascend Tribrid mass spectrometer for accurately quantifying and identifying hundreds to thousands of proteins from single cell-line to complex sample mixtures with a high background of human peptides. The Velocity DIA workflow could be coupled with the optional Thermo Scientific<sup>™</sup> AccelerOme<sup>™</sup> automated sample preparation platform to improve throughput and minimize variations caused by manual sample handling (Figure 1). The LC method and gradient were optimized to fully utilize the capabilities of micropillar array-based separation columns. Similarly, the mass spectrometric method was adopted to account for long (60 min active gradient), short (30 min active gradient), and ultra-high throughput (9 min active gradient) gradient lengths without compromising identification rates at great quantitation accuracy and precision.

## Introduction

Quantitative proteomics is an essential tool for understanding global protein expression and the mechanisms of biological processes and disease states. Statistical significance is improved by decreasing variability in measurements and/ or increasing the sample set. However, increasing throughput means decreasing acquisition time, which often comes at a cost to measurement quality. Therefore, acquisition methods must be extensively optimized and validated to ensure that the data will produce meaningful biological insights.

We also evaluated a 9 min active gradient to enable ultra-high throughput. At such high throughput, there is a tradeoff between depth of coverage and measurement quality. Therefore, we developed two methods to meet different experimental needs: one for maximizing identifications (Max ID) and the other for maximizing quantitative (Max Quan) performance. In the Max ID method, we identify 5,400+ proteins in a 9 min active gradient, highlighting the sensitivity and scanning speed of the Orbitrap mass spectrometer in a high throughput setup. In contrast, the Max Quan method, compromised proteome coverage to provide better quantitative performance, as evidenced by ~6% protein group CV in an ultra-high throughput setup (Figure 4).

Figure 2. Experimental sample and active gradient design of the Velocity DIA workflow for label-free quantitation.



(A-C) Bar charts showing the number of proteins and peptides identified from 200 ng HeLa digests with different active gradients evaluated in the present study. (D-E) Violin plots of all four tested methods reveal high precision of protein quantities in technical replicates. Data was analyzed with Spectronaut (left panel), Proteome Discoverer using CHIMERYS (middle panel) or DIA-NN (right panel).

To understand if a higher load benefits the Velocity DIA workflow in maximizing identification, 500 ng of HeLa digest was analyzed with the 60 min active gradient experiments. The results indicated >8,100 protein groups were identified, with CV being  $\sim 4\%$ (Figure 5). Thus, the ability of the Orbitrap Ascend Tribrid MS platform to accommodate sample loads from picograms to micrograms can be leveraged to further increase the depth and quality of measurements.

Figure 5. A long gradient coupled with a higher sample load affords the superior performance in the Velocity DIA workflow.

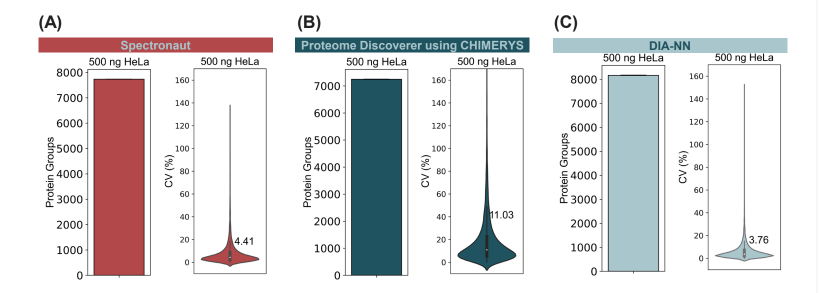
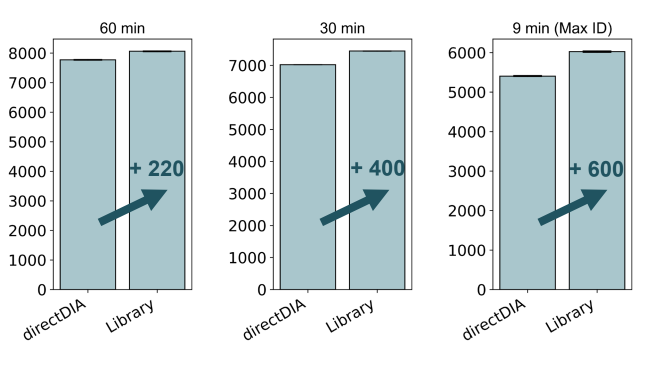


Figure 7. Library search improved proteome coverage.



Bar charts showing the number of protein groups gained through library search by using DIA-NN (v1.8.1).

#### High accuracy and precision of quantitation

Traditional data-dependent analysis (DDA) approaches have been widely employed for LFQ experiments, but they suffer from run-torun inconsistencies due to intensity-based stochastic triggering of precursors, often leading to under sampling especially of lowabundant proteins. Missing values become more likely as sample size increases, and so data independent acquisition (DIA) has emerged as a popular technique for large scale quantitative analyses.

# Materials and methods

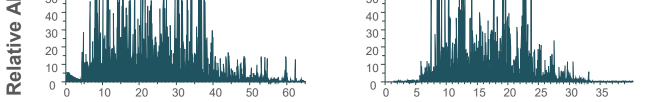
#### Sample preparation

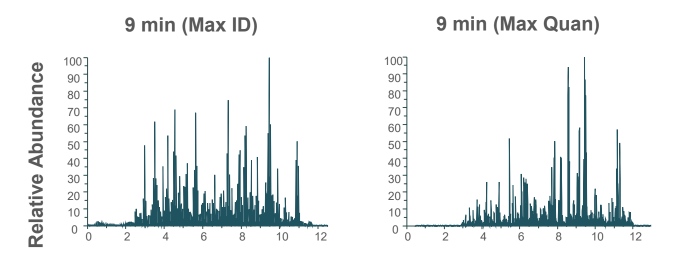
Thermo Scientific<sup>™</sup> Pierce<sup>™</sup> HeLa Protein Digest Standard, Waters E. coli MassPREP Standard and Promega Mass Spec-Compatible Yeast Protein Digest were dissolved in 0.1% formic acid (FA) with 30 seconds of vortexing. For the three-proteome mix, E. coli peptide digest and yeast peptide digest were added to a fixed amount of HeLa digest (325 ng) at amounts of 100 ng to 25 ng, and 75 ng to 150 ng, respectively, yielding an E. coli peptide ratio of 1:4 and a yeast peptide ratio of 1:0.5.

## Test method(s)

HeLa digest and three-proteome mixtures were loaded onto a 50 cm µPAC Neo HPLC column and separated at a 350 nL/min flow rate in direct injection mode using a Vanquish Neo UHPLC system over 9 min, 30 min, and 60 mins LC gradients, respectively, before being transferred into the Orbitrap Ascend Tribrid mass spectrometer (Figure 1 and 2).

Source parameters, including spray voltage and ion transfer tube temperature, are tunable parameters and must be optimized for the individual setup.

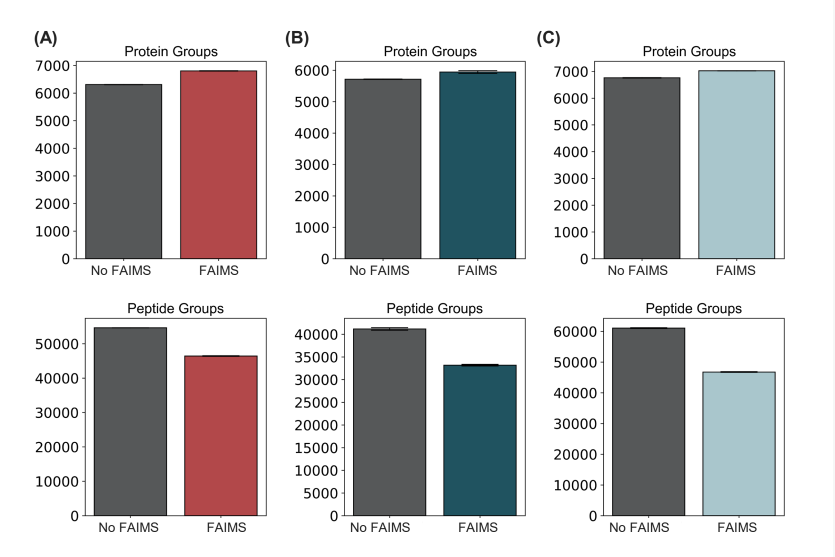




#### **Retention Time (min)**

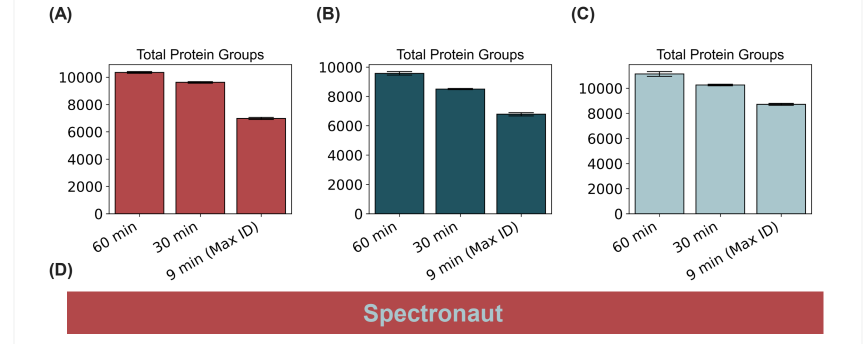
(A) 200 ng HeLa digest to access quantitation precision and proteome coverage of human samples. (B) The three-proteome mix contains a medium human background of 325 ng HeLa peptides together with yeast and E. coli peptides digested in ratios of 0.5:1 and 1:4, respectively. The mixtures have been chosen because they closely mimic biological samples with larger or smaller protein expression changes. (C) Three different active gradient lengths, including 60 min, 30 min, and 9 min, were selected to develop the Velocity DIA workflow for different throughput needs. For the ultra-high throughput 9 min gradient setup, two methods were developed to obtain deep proteome coverage (Max ID) or excellent quantitation performance (Max Quan).

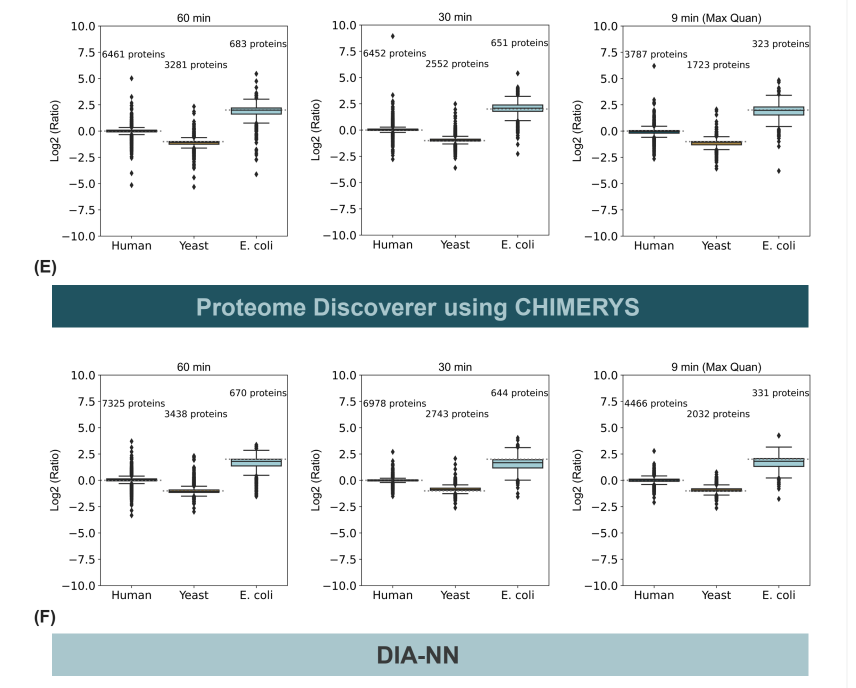
#### Figure 3. Evaluation of the FAIMS Pro interface in the Velocity **DIA workflow.**



Bar chart and Violin plots showing the deeper proteome coverage and high precision of protein quantitation from 500 ng of HeLa digest in the 60 min method. Data was analyzed with (A) Spectronaut, (B) DIA-NN and (C) CHIMERYS on Proteome Discoverer.

## Figure 6. Determination of microbial and human protein abundance ratios in a three-proteome mixture.





In addition to protein identification, quantitative data is necessary to study biomarkers and get insights into biological pathways. The Velocity DIA workflow yielded excellent quantitative accuracy across a wide dynamic range with median values extremely close to the theoretical ratios, as well as a narrow distribution of all data points around the median values, indicating high quantitative accuracy and precision of the workflow (**Figure 6**).

## Evaluating data processing software in DIA analysis

Library-based search in DIA proteomics may enhance proteome coverage and improve peptide identification in certain scenarios. To evaluate if Velocity DIA data can benefit from library search, we generate a general library by using Pulsar on Spectronaut (v18) from single shots of 12 cell lines acquired on Orbitrap Astral mass spectrometer. The raw files were re-processed with library-based search on DIA-NN (v1.8.1). In line with the benefit of library-based search, we observed a 3-10% increase in protein identification with more benefits for shorter gradients (Figure 7).

## Conclusions

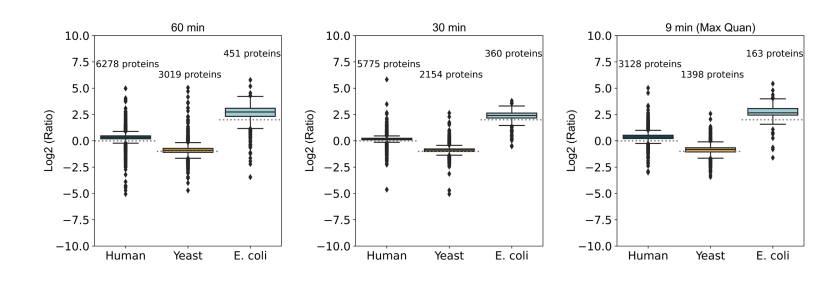
The high-resolution DIA workflow for label-free quantitation setup on an Orbitrap Ascend Tribrid mass spectrometer coupled to a Vanquish Neo UHPLC system running with a 50 cm µPAC Neo UHPLC column was shown to fulfill the following performance criteria:

- Deep (7000+ protein groups) proteome coverage from 200 ng of HeLa standard.
- Excellent quantitative accuracy and precision for small amounts of bacterial and fungal proteomes from challenging sample matrices.
- Sample throughput and quality of the obtained data while achieving high proteome coverage.

## Data analysis

Acquired data has been processed by Spectronaut software (Biognosys, v18) using a directDIA approach, DIA-NN (v1.8.1) or Thermo Scientific<sup>™</sup> Proteome Discoverer<sup>™</sup> software with CHIMERYS<sup>™</sup> intelligent search algorithm by MSAID.

Box plots depicting the number of proteins and peptides identified from 200 ng of HeLa digest in the 30 min method with the FAIMS Pro interface at a CV of -45 V. Data were analyzed with (A) Spectronaut, (B) Proteome Discoverer software using CHIMERYS or (C) DIA-NN.



(A-C) Bar plots showing the total protein groups identified from three proteome mix in different active gradients. Data was analyzed with Spectronaut (left), Proteome Discoverer using CHIMERYS (middle) or DIA-NN (right). (D-F) Whisker box plots of protein abundance ratios of all three species demonstrate excellent quantitation accuracy by being consistent with the theoretical ratios (gray dotted line).

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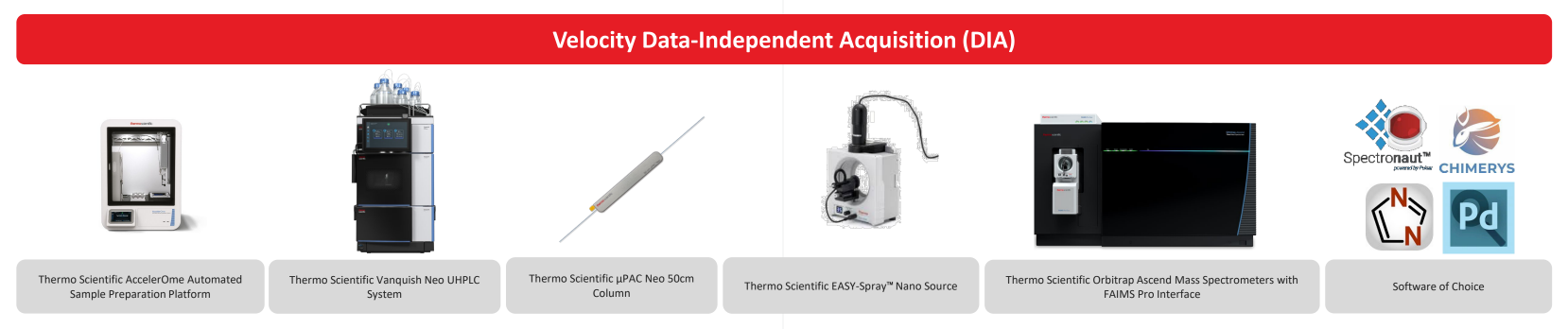
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## Figure 1. Complete "end-to-end" Velocity DIA workflow



Velocity DIA workflow on Orbitrap Ascend Tribrid MS together with the optional AccelerOme automated sample preparation platform.