Discovery to Validation: Transition from SWATH® Acquisition to Targeted MRM Analysis for Quantitative Proteomics Pipeline

Using the TripleTOF® 5600+ System and QTRAP® 6500 System

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In their efforts to accurately quantify and verify putative biomarkers or other proteins of interest, researchers often move from discovery techniques, such as shotgun proteomics, to targeted assays in order to obtain accurate quantitation of analytes in complex mixtures. MRM (Multiple Reaction Monitoring) is the gold standard for performing this type of targeted analysis, providing high sensitivity and selectivity. However, there is a limit to the number of MRMs that can be monitored in a single injection. Data independent acquisition is rapidly growing as a key technique for more comprehensive quantitation.

SWATH® Acquisition brings together data independent acquisition and targeted data processing to deliver high quality quantitative data with extremely high multiplexing. The quantitative statistics of SWATH Acquisition have been demonstrated to be of very high quality, approaching those of MRM assays on a triple quadrupole. While the MRM approach on a high sensitivity QTRAP® system is still the method of choice for high throughput analysis of a smaller subset of targets and to obtain the best quantitative statistics for the lowest abundant species; SWATH Acquisition offers comprehensive sample coverage with only a small compromise in quantitative precision.

In this study, the workflow of performing SWATH Acquisition for comprehensive quantitation followed by MRM analysis on a subset of targets is demonstrated and the results quality of the two techniques compared.

Key Aspects of Transitioning from SWATH® Acquisition to MRM Analysis

- SWATH Acquisition on the TripleTOF® 5600+ System
  - Full scan MS/MS of all detectable ions is acquired, enabling post-acquisition interrogation of data
  - No upfront assay development is required
  - Comprehensive high quality quantification with confirmation of identity on every detectable protein and peptide

- MRM analysis on a QTRAP® 6500 System
  - LINAC® collision cell technology in both TripleTOF and QTRAP systems means that SWATH acquisition data can be directly transitioned into an MRM assay on the QTRAP 6500 system providing seamless pipeline (Figure 1)

- Higher sensitivity and specificity MRM assays on a subset of proteins and peptides for validation
- Use additional specificity and sensitivity to accelerate assays using faster gradients or higher flow rates

Figure 1. Comprehensive Quantitation using SWATH Acquisition followed by High Sensitivity MRM Analysis. SWATH acquisition on a TripleTOF® 5600+ system provides a powerful new approach for discovery and verification experiments. Transition to MRM analysis for high throughput validation on QTRAP® systems is straightforward.
Experimental

**Sample Preparation:** A tryptic digest of E.coli sample was analyzed with a 1 µg load on column.

**Chromatography:** The samples were analyzed using the Eksigent ekspert™ nanoLC 425 System combined with the chiPLC® System in Trap-Elute mode. The samples were first loaded on the chiPLC trap (200 µm x 0.5 mm ChromXP™ C18-CL, 3 µm, 300 Å) and washed for 10 mins at 2 µL/min. Then, an elution gradient of 5-35% acetonitrile (0.1% formic acid) in 60 mins was used on a nano chiPLC column (75 µm x 15 cm ChromXP C18-CL, 3 µm, 300 Å).

**Mass Spectrometry:** SWATH® Acquisition was performed using a TripleTOF® 5600+ system interfaced to the NanoSpray® Source. The acquisition strategies used was 25Da Q1 windows with 100 msec accumulation time across a mass range of 400-1000 m/z. A set of protein/peptides of interest across an abundance range obtained from SWATH acquisition data were then chosen for further MRM analysis. Protein/peptide data were loaded into Skyline for MRM assay development on the QTRAP® 6500 System. For low intensity proteins with only a single peptide quantified, *in silico* MRM transitions of other peptides from the same protein were added to the assay, and targeted using MIDAS™ Workflow for sequence confirmation. Replicates were run for both acquisition strategies.

**Data Processing:** SWATH Acquisition data was processed using the PeakView® Software and the SWATH Acquisition MicroApp. MRM data analysis was performed using MultiQuant™ Software and data were compared to SWATH acquisition data using Excel tools.

![High Reproducibility SWATH® Acquisition](image)

**Figure 2:** Quantitative Analysis of SWATH Acquisition Replicates. (Top) Fragment XICs with peak areas across ~3.5 orders dynamic range were quantified. (Middle) High reproducibility was achieved on very high numbers of peptides and transitions across the intensity range of peak areas. High intensity peaks showed median CVs as low 2%, with many low intensity peaks still showing good reproducibility of 15%. (Bottom) Cumulative %CV curves show that ~90% of the transition level data has CVs better than 20% and the peptide and protein level data is even better.

**Figure 3.** Skyline for MRM Assay Development. A subset of proteins from the SWATH acquisition data were loaded into Skyline software for MRM assay development. Previous MS/MS results were used as a reference, then fragment ion and collision energy optimization was performed. On the right is the MS/MS spectrum of the highlighted E.coli peptide along with the MRM transitions for the final assay.

**High Reproducibility SWATH® Acquisition**

SWATH Acquisition provides an excellent technique to broadly interrogate many proteins and peptides in a sample with very good quantitative quality (Figure 2). Because it is a data independent acquisition strategy, all data are collected using a single acquisition method, with no upfront method development required. Data are then interrogated post-acquisition to select the best peptides and transition for quantitation. High quality data was obtained with SWATH acquisition across five replicate injections (Figure 2). Across the whole dataset, ~90% of peptide...
transitions had CVs less than 20%. When assessed relative to the transition peak areas, the highest peak intensities provided the best reproducibility as expected, but even the lowest intensity peaks showed excellent reproducibility.

**Transition Subset of Proteins to MRM Analysis**

Once a subset of proteins and peptides are found to be of interest and need to be investigated further, they can easily be transitioned to an MRM assay for a higher throughput, higher sensitivity assay. In this case, Skyline was used to develop the MRM assay on a subset of the lower abundance peptides (Figure 3). As both the TripleTOF® and QTRAP® systems utilize similar collision cell technology, the MS/MS data obtained from the SWATH® acquisition can be mined to directly inform the building of robust MRM assays.

**Comparing Results Quality**

As the same peptides were analyzed by the two techniques, it provides the opportunity to compare the results quality. Most of the SWATH Acquisition data has very low CVs for the replicate analysis, as seen in Figure 2. If the SWATH data is analyzed considering peak intensity (Figure 4, bottom), it is observed that the reproducibility is high across the abundance range of the proteins analyzed, but shows a small decay as the peak intensity decreases. The same analysis was done for the MRM data (after performing an intensity correction on the data to account for the higher sensitivity). One can see that the SWATH acquisition data is very similar to the MRM data quality across the abundance range, except for the lowest abundance peaks where the added MRM specificity and sensitivity is advantageous.
The CVs obtained for individual peptides that were analyzed in both techniques were compared and plotted in Figure 4 (top). Again, many of the peptides showed very similar CVs between the two techniques showing the high quantitative quality obtained by both techniques. However, some peptides showed improved quantitation using MRM, such as the peptide example highlighted with the orange arrow. The data for this peptide is displayed in Figure 5 (bottom) and shows that for the low abundance transitions, moving to the targeted MRM assay provided improved data quality. This is expected as the QTRAP 6500 system is more sensitive than the TripleTOF® 5600+ system, and is operating with a tighter Q1 window for added specificity (improved S/N, see Figure 5).

Scale and Completeness of Data

Another important aspect of quantitation is the completeness of measurements across all samples - a key advantage of targeted quantitation. It is highly desirable to have a good measurement for every peptide and protein targeted in every sample analyzed. Figure 6 provides a detailed analysis of the data completeness observed across the 5 replicates for the SWATH® acquisition experiment. The blue text show that ~19,500 fragment ions were extracted and 78% of targeted fragments had good peak areas and provided a peak group score of 1% FDR (Figure 6, top). Once fragment ions are summed to the peptide and protein level, the data completeness approaches 95% or higher. The MRM assay was developed to focus on the lower abundance peptides, resulting in a smaller number of peptides targeted (175 peptides), but with a higher success rate of ~91% at the transition level in this case.

Conclusions

High quantitative reproducibility on very large numbers of peptides and proteins can be obtained using SWATH Acquisition on the TripleTOF® 5600+ System. By targeting a smaller subset of peptides using MRM analysis on a QTRAP® 6500 System, a further improvement in quantitative quality can be achieved, especially on for lower abundance proteins. This work demonstrates the ability to perform initial quantification (verification) on large numbers of proteins using SWATH Acquisition, then easily transition to an improved targeted MRM assay for accelerated quantification.

References

1. SWATH® Acquisition - Comprehensive Quantification with Qualitative Confirmation using the TripleTOF® 5600+ System. SCIEX Technical RUO-MKT-02-2878-A.
2. Exploring the Sensitivity Differences for Peptide Quantification in the Low Flow Rate Regime - nanoLC™ 400 System for High Performance Nanoflow and Microflow LC. SCIEX technical note RUO-MKT-02-3252-A.
4. In order to overlay the %CV curves for both instruments, a correction factor was applied to adjust the X-axis for the signal intensity difference between the two instruments (~10x).