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Enhancement in MALDI-TOF MS analysis of the low molecular weight human serum proteome

Christine L. Gatlin
William & Mary

Dariya I. Malyarenko
William & Mary

Krista Y. White

Christopher E. Wilkins

O. John Semmes

See next page for additional authors

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Enhancement in MALDI-TOF MS analysis of the low molecular weight human serum proteome

The combination in advances in MALDI-TOF MS instrumentation[1] and serum sample preparation techniques[2–4] has led to the emergence of MALDI serum protein expression profiling as a promising tool for biomarker discovery.[5] However, three factors still pose significant challenges for MALDI profiling of serum: the limited mass window of MALDI, serum protein complexity,[6] and analytical reproducibility.[7,8] MALDI has a small mass preference due to the ionization efficiency of intact protein molecules and the focusing of flight of the ions. In the linear mode, TOF analyzers have limited sensitivity for masses above 20 kDa. To address this, we previously reported the enhancement of MALDI broad mass range detection of protein signals.[9,10] Spanning a 100-kDa mass range, we improved the sensitivity of detection by an order of magnitude through the combined optimization of sample preparation, instrument parameters and data processing procedures.

The complexity of the blood proteome is very high, with protein concentrations differing by up to ten orders of magnitude.[10] This large dynamic range exceeds current proteomic analytical capabilities; thus, analysis of easily prepared subproteomes of serum or plasma is essential. Here, we shift our focus back to the enhancement of MALDI analysis in the low mass regime. The low molecular weight (LMW) subcomponent of serum promises to be a rich source of undiscovered biomarkers, as biological processes give rise to a plethora of proteolytic protein fragments.[11] Currently, there is no consensus on what constitutes the mass limits of this derivative proteome (also termed peptidome). Hence, ≪15 kDa is often cited in the literature based on a serum MALDI study using 20 MWCO filters.[12]

To date, small native protein/peptide mass measurements have been mainly conducted in reflectron mode (≈4 kDa)[13] and linear mode up to 10 kDa.[14] However, no systematic comparison has been described for a combination of preparation steps: spotting, filtering and matrix choice, to optimize the performance. A rigorous reproducibility study for different preparation strategies is likewise missing, which prevents the extension of the technique to clinical proteomics applications. The purpose of this work was to provide such a systematic comparison, using rigorous performance metrics to optimize sample preparation for LMW serum proteome profiling by MALDI-TOF MS analysis up to 20 kDa. We explored a combination of MALDI sample preparation and spotting methods. Procedures that gave the best results included: (1) MALDI spotting with a thin-layer (TL) technique using sinapinic acid on a grond steel plate and (2) centrifugal ultrafiltration with 50,000 MWCO filters in the presence of 2% TFA.[9,10]

For the dried droplet (DD) method, matrices were dissolved in 50% ACN, 0.1% TFA, (5 mg/ml CHCA or saturated SA). Samples were mixed with matrix in the ratio of 1:5, then 1 µl was spotted onto an AnchorChip or ground steel targets and dried droplet versus ground steel targets and dried droplet versus TL spotting methods. The following six MALDI spotting methods were compared in detail (target–matrix–spotting technique):

- Anchorchip–CHCA–dried droplet.
- Steel–CHCA–dried droplet.
- Steel–SA–dried droplet.
- Steel–CHCA–thin layer.
- Steel–SA–thin layer.

For the dried droplet (DD) method, matrices were dissolved in 50% ACN, 0.1% TFA, (5 mg/ml CHCA or saturated SA). Samples were mixed with matrix in the ratio of 1:5, then 1 µl was spotted onto an AnchorChip or ground steel plate. For our modified TL method, 0.3 µl of matrix solution (CHCA saturated in MeOH or SA saturated in EtOH) was deposited onto the ground steel target as a seed layer. Protein sample is mixed with matrix solution as above, followed by 1 µl deposited onto the seed layer.

PS1 contains a good spread of ion signals in the range of 3–20 kDa and was used to compare the different MALDI spotting methods. The PS1 mixture contains insulin (5734 Da), ubiquitin (8565 Da), cytochrome c (12,361 Da) and myoglobin (16,952 Da). PS1 was spotted ten times for each preparation. A quick visual inspection showed the TL method to be far superior to the more common DD method in the 3–20 kDa range. An advantage of the TL method is the very homogeneous size of microcrystals.[15] On deposition of the seed layer, the solution quickly was prepared to compare MALDI profiling in the LMW range (3–20 kDa). Details are given in the following section.

Mass spectra were acquired in linear and reflectron positive ion mode using an Ultraflex III™ MALDI-TOF/TOF mass spectrometer (Bruker Daltonics). The instrument is equipped with a smartbeam™ laser, and acquisition laser power was optimized using the PS1 calibration mixture before the collection of sample data. Instrument settings, optimized for mass range m/z 0–20,000, were as follows: ion source 1 = 25.0 kV, ion source 2 = 23.7 kV, lens voltage = 6.0 kV, pulsed ion extraction time = 200 ns, matrix suppression mass cut off = 1500, ADC offset = 50, pre-amplifier filter bandwidth = high, digitizer sampling frequency = 500 MHz. All spectra were generated by averaging 1000 shots from 10 nonoverlapping positions (100 shots/position).

Results and Discussion
MALDI spotting
Clinical profiling of body fluids by MALDI-MS is highly influenced by the choice of MALDI spotting and sample preparation techniques. Our goal was to determine an optimal method for MALDI profiling of LMW serum protein/peptide ions, based on analytical performance measurements of resolution, sensitivity, reproducibility and broad mass range coverage. For sample/matrix crystal preparation, α-cyano-4-hydroxy-cinnamic acid (CHCA) and sinapinic acid (SA) are most commonly used. Resolution and sensitivity comparisons of these two matrices are well documented in the literature; however, reproducibility studies are limited. Because reproducibility is one of the most critical elements in profiling studies, our main focus was to optimize reproducibility over the LMW serum mass range (=20 kDa), without undermining other performance factors. We evaluated a variety of matrices, sample/matrix ratios, ionic liquid additives, target plates and spotting methods. We narrowed down the principal comparisons to CHCA versus SA matrix (with no additives); Bruker AnchorChip (600 µm) versus ground steel targets and dried droplet versus TL spotting methods. The following six MALDI spotting methods were compared in detail (target–matrix–spotting technique):

- Anchorchip–CHCA–dried droplet.
- Steel–CHCA–dried droplet.
- Steel–SA–dried droplet.
- Steel–CHCA–thin layer.
- Steel–SA–thin layer.

Experimental Procedures
All chemicals and solvents were purchased from Sigma–Aldrich (St. Louis, MO, USA) unless otherwise noted. Protein standard 1 (PS1), MALDI matrices and targets and ClinProt MB-C3 magnetic beads (C3) were obtained from Bruker Daltonics (Leipzig, Germany). Amicon-4 and Amicon-0.5 Ultra centrifugal filter devices were obtained from Millipore (Billerica, MA, USA). A quality control (QC) serum sample was prepared by pooling human serum from a controlled normal group as previously described.[16] Six different MALDI target preparations and 23 different serum fractionations were prepared to compare MALDI profiling in the LMW range (3–20 kDa). Details are given in the following section.

Mass spectra were acquired in linear and reflectron positive ion mode using an Ultraflex III™ MALDI-TOF/TOF mass spectrometer (Bruker Daltonics). The instrument is equipped with a smartbeam™ laser, and acquisition laser power was optimized using the PS1 calibration mixture before the collection of sample data. Instrument settings, optimized for mass range m/z 0–20,000, were as follows: ion source 1 = 25.0 kV, ion source 2 = 23.7 kV, lens voltage = 6.0 kV, pulsed ion extraction time = 200 ns, matrix suppression mass cut off = 1500, ADC offset = 50, pre-amplifier filter bandwidth = high, digitizer sampling frequency = 500 MHz. All spectra were generated by averaging 1000 shots from 10 nonoverlapping positions (100 shots/position).

Correspondence to: Christine L. Gatlin, National Human Genome Research Institute/National Institutes of Health, 5635 Fishers Lane, Bethesda, MD 20892, USA. E-mail: christine.gatlin@nih.gov
spreads and evaporates almost instantaneously (<2 s) leaving an ultra-thin uniform coating. The method generally yields higher resolution spectra, and the detection limit is increased compared with the DD method.

Of the six preparations, three were then statistically compared: CHCA–DD on AnchorChip, CHCA–TL on steel plate and SA–TL on steel plate. Signal processing was performed on raw PS1 spectra to enhance signal to noise (SNR). Processing algorithms, described by Malyarenko et al., included analytical model baseline subtraction, integrative down-sampling optimal linear filtering, pedestal removal, peak detection and alignment. Noise-level spectra were estimated by finding the standard deviation of the noise in the down-sampled spectra and adjusting for the expected effect of filtering. Poisson dependence on baseline amplitude was included to account for the observed dark current amplification for early TOF. Integrated down-sampled signal intensities were compared to exclude the influence of the originally different peak widths. Implementation details and Matlab toolbox for signal processing are available from matlabcentral/fileexchange/24469. Metrics for 9 m/z peaks common to each of the three preparations included: measured ion intensity, estimated noise, SNR, normalized intensity, normalized SNR and %CV.

Figure 1 shows the mean raw spectra for each of the three preparations. The top spectrum (SA spotted with the TL method on ground steel) proved to be of highest quality based on SNR and reproducibility. Compared with CHCA on AnchorChip, SA on steel produced 3.5× greater average signal intensity and 10× greater average SNR (based on the average values for all peaks and replicates for each sample preparation); whereas TL methods on steel produced similar ion signals for both CHCA and SA. Noise was greatly reduced with SA (1/3.5) giving an overall 3× increase in SNR for SA over CHCA on steel. Comparing individual ion signals in Fig. 1 shows that five of the nine ions have higher signals for CHCA on steel than for SA on steel. However, only two of these ions (ubiquitin 3+ and 2+) have higher SNR (with only 1.5× increase).

In the literature, the utilization of TL preps have focused generally on increasing signal and reproducibility of peptides (m/z <4000) using CHCA or higher mass proteins (m/z >10 000) using SA. However, our interest is in the middle range – m/z 3000–20 000 – where LMW serum protein peaks are commonly found. As Fig. 1 demonstrates, we determined SA is a better matrix over CHCA in our modified TL method, due to the reduced noise, higher SNR and resolution that SA offers in the 3–20 kDa range.

Coefficients of variation (%CV) were calculated for experimental replicates for each m/z peak based on normalized intensity and normalized SNR. Table 1 presents the %CV for each peak and for each surface preparation. The average %CVs for a given preparation are shown at the bottom of the table. Reproducibility was comparable whether
Table 1. %CVs for PS1 ions (ten replicate average) for the studied MALDI preparations

<table>
<thead>
<tr>
<th>Peak m/z (Da)</th>
<th>Matrix/surface</th>
<th>CHCA on anchor dried droplet</th>
<th>CHCA on steel thin layer</th>
<th>SA on steel thin layer</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Intensity</td>
<td>SNR</td>
<td>Intensity</td>
</tr>
<tr>
<td>2855 Ub+3</td>
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<td>17.0</td>
<td>16.4</td>
<td>7.8</td>
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<tr>
<td>2867 In+2</td>
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<td>16.3</td>
<td>16.9</td>
<td>9.9</td>
</tr>
<tr>
<td>4283 Ub+2</td>
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<td>6.0</td>
<td>5.9</td>
<td>5.2</td>
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<td>7.7</td>
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<td>7.5</td>
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<td>8.0</td>
<td>7.1</td>
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</table>

Ave %CV 12.5 12.3 9.8 9.8 9.5 9.4

Table 2. Within and between samples %CVs based on replicate spotting of eight aliquots of QC serum processed by ultrafiltration (%CV values calculated for 16 peaks normalized by SNR)

<table>
<thead>
<tr>
<th>Peak m/z (Da)</th>
<th>QC Aliquot</th>
<th>A (8 reps)</th>
<th>D (9 reps)</th>
<th>G (10 reps)</th>
<th>H (10 reps)</th>
<th>J (10 reps)</th>
<th>K (10 reps)</th>
<th>M (10 reps)</th>
<th>N (9 reps)</th>
<th>Range %CV (within sample)</th>
<th>Between sample %CV a</th>
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<tr>
<td>2371</td>
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<td>24.1</td>
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<td>18.5</td>
<td>8.8</td>
<td>5.8–35.1</td>
<td>42.4</td>
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</table>

Within sample ave %CV 22.8 11.8 8.6 10.6 10.4 13.4 12.7 8.5–22.8 25.2

| a | based on 74 replicates |

mean CV : 12.3% within sample
mean CV : 25.2% between samples

using normalized intensity or SNR. All preps gave good within-day reproducibility; however, the TL prep with SA on steel was slightly better (%CV = 9.5). Our previous studies for reproducibility of calibration spectra indicated that under carefully preserved instrumental settings, the %CV does not exceed 10% in the course of several months. This confirmed that the major contribution to the variability of MS signals comes from the sample preparation step rather than the instrument or time of the study.

Ultrafiltration

Human QC serum samples were used with Millipore Amicon ultrafiltration devices to optimize MALDI profiling of the LMW serum fraction. Devices were used according to the manufacturer’s recommendations. Twenty-three comparisons were made depending on: (1) MWCO: 3, 10, 30, 50 and 100 K, (2) collecting filtrate or retentate, (3) filter size: 0.5 or 4 ml, (4) double filter combinations, (5) denaturation and (5) MB-C3 purification. A list of these combination preparations can be found in the Supporting Information. All samples were MALDI spotted using the TL method with SA on ground steel.

For the MWCO, we found that serum filtrate from the 50K filter gave the best profile in the 3–20 kDa range. This was enhanced by pre-filtration protein denaturation with 2% TFA and post-filtration C3 magnetic bead desalting. Other filtrations that gave complimentary profiles (but less overall coverage in the 3–20 kDa range) included the 10K retentate and combination 50K filtrate followed by 3K retentate (data not shown). The filter size did not matter unless double filtrations were performed (0.5 ml filter for the second filtration).

Figure 2 shows MALDI spectra of processed serum. Figure 2(a) is a typical profile of C3 magnetic bead processed QC serum. Figure 2(b) is 50K filtered QC serum followed by C3 bead cleanup. In comparison, noise is greatly reduced for the filtered sample which enhanced peaks in the m/z 2000–6000 range. Adding upfront 2% TFA denaturation (Fig. 2(c)), noise is further reduced with enhanced peaks in the m/z 7000–10 000 range.
The reproducibility of the two-step serum preparation was determined based on variation in peak intensities and normalized SNP. The systematic reproducibility study was performed only for the best preparation method (Fig. 2(c)), since other methods did not provide the desired signal gain over the full LMW range. Eight separate aliquots of QC serum (eight biological replicates) underwent 50 kDa centrifugal ultrafiltration and C3 magnetic bead purification. Prepared aliquots were spotted 10 times (10 experimental replicates) on a MALDI target. Raw QC spectra were processed using the same procedures (with slightly different parameters) as the PS1 spectra. Sixteen m/z peaks were selected for comparison. From the data, we found that intensity CVs were not correlated to the within samples spectra. Sixteen experimental replicates on a MALDI target. Raw QC spectra were processed magnetic bead purification. Prepared aliquots were spotted 10 times (10 over the full LMW range. Eight separate aliquots of QC serum (eight reproducibility study was performed only for the best preparation method [16,18,28,29]. Of greater note, reproducibility of 25% is not very noteworthy, it is within the range of CVs between 13 and 47%, with an average of 25%. While this sample-to-sample variability is greatly enhanced in the LMW serum protein range of LMW protein/peptide signatures since all the potential markers to date correspond to proteolytic fragments of the most abundant plasma proteins. However, our ultrafiltration/MALDI method extensively improves peptide/protein peak intensities and reproducibility in the 3–20 kDa range. Our preliminary results (data not shown) comparing prostate cancer to normal serum samples indicate significant peak differences in the LMW range. For future work, we plan to conduct a larger LMW serum clinical profiling study of prostate cancer-related samples using our denaturing ultrafiltration technique.

Acknowledgements

This investigation was supported by NIH grant CA126118 and its ARRA supplement from the Advanced Proteomics Platforms and Computational Sciences Program of the National Cancer Institute. We thank Drs Eugene Tracy, William Cooke and Dennis Manos of the College of William and Mary for comments, and Dr Lisa Cazares of Eastern Virginia Medical School for experimental support.

Yours,

CHRISTINE L. GATLIN,a,b,c KRYSTAL Y. WHITE,b MAUREEN B. TRACY,c CHRISTOPHER E. WILKINS,b O. JOHN SEMMES,b JULIUS O. NYALWIDHE,b RICHARD R. DRAKEb AND DARIYA I. MALYARENKOa a Department of Applied Science, College of William and Mary, Williamsburg, VA 23187-8795, USA b Center for Biomedical Proteomics, Eastern Virginia Medical School, Norfolk, VA 23501, USA c William and Mary Research Institute, College of William and Mary, Williamsburg, VA 23187-8795, USA

Supporting information

Supporting information may be found in the online version of this article.

References


