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## Identification and Characterization of Human Proteoforms by **Top-Down LC-21 Tesla FT-ICR Mass Spectrometry**

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## Abstract

Successful high-throughput characterization of intact proteins from complex biological samples by mass spectrometry requires instrumentation capable of high mass resolving power, mass accuracy, sensitivity, and spectral acquisition rate. These limitations often necessitate the performance of hundreds of LC-MS/MS experiments to obtain reasonable coverage of the targeted proteome, which is still typically limited to molecular weights below 30 kDa. The National High Magnetic Field Laboratory (NHMFL) recently installed a 21 T FT-ICR mass spectrometer, which is part of the NHMFL FT-ICR User Facility and available to all qualified users. Here we demonstrate top-

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Supporting Information

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Figure S-1, Silver-stained gel image of DLD-1 A protein fractions. Figure S-2, Differential protein and proteoform identification by CID and ETD. Figure S-3, Score and MW Distributions of DLD-1 B Proteins and Proteoforms. Figure S-4, Score and MW Distributions of DLD-1 C Proteins and Proteoforms. Table S-1, Number of unique proteins and proteoforms identified at 1% FDR from three biological replicates. Table S-5, List of fragments matched to putative sequence of p23 (UniProt P13693) by TDPortal (shown in Figure 2C). (PDF) Tables S-2, S-3, and S-4, Proteoforms identified at 1% FDR with C-scores of 40 or better from each biological replicate (DLD-1 A, B, and C, respectively). (XLSX)

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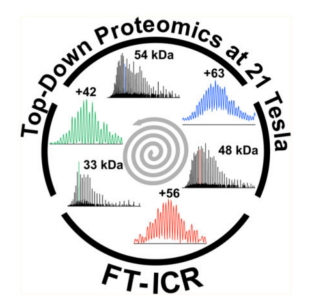
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The authors declare the following competing financial interest(s): Some authors are involved with software commercialization activities

All raw spectrum files (.raw), search engine files (.tdreport), and the flat file (.txt) used for the creation of the database against which data were searched have been uploaded to the UCSD MassIVE repository with the identifier/username MSV000079978 and can be accessed here: ftp://massive.ucsd.edu/MSV000079978/. The TDViewer application, which allows one to view the contents of the .tdreport files, is available for download at http://topdownviewer.northwestern.edu/

down LC-21 T FT-ICR MS/MS of intact proteins derived from human colorectal cancer cell lysate. We identified a combined total of 684 unique protein entries observed as 3238 unique proteoforms at a 1% false discovery rate, based on rapid, data-dependent acquisition of collision-induced and electron-transfer dissociation tandem mass spectra from just 40 LC–MS/MS experiments. Our identifications included 372 proteoforms with molecular weights over 30 kDa detected at isotopic resolution, which substantially extends the accessible mass range for high-throughput top-down LC–MS/MS.

## Graphical abstract



#### Keywords

FT-ICR; 21 tesla; top-down proteomics; FTMS

### 1. INTRODUCTION

Regulation of nearly every cellular process is directly linked to protein primary structure. Comprehensive knowledge of primary structure cannot be derived from the genome because translation of mRNA into protein does not dictate the composition of the final protein complement. Alternative splicing, single nucleotide polymorphisms, endogenous protein cleavages, and the vast number of ways in which any single protein can be post-translationally modified expand the number of theoretically possible proteoforms to well over a billion.<sup>1,2</sup> Each proteoform has the innate potential to impact and affect different biological outcomes including phenotypes of health and disease. Mass spectrometry (MS) is an invaluable source of protein structural information, including amino acid sequence and identification and site-localization of post-translational modifications (PTMs).

The ultimate proteomics platform must be capable of unequivocal differentiation of closely related proteoforms, which requires intact analysis (i.e., top-down). However, top-down

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proteomics typically suffers from slow spectral acquisition rate, low signal-to-noise ratio (S/N), inefficient protein fragmentation, and increased spectral complexity.<sup>3</sup> Tandem mass spectra typically contain many overlapping fragments that require high mass resolving power (RP) and spectral averaging to improve S/N. These obstacles have proven difficult for the burgeoning field of top-down proteomics, particularly for proteins larger than 30 kDa.

Fourier-transform ion cyclotron resonance (FT-ICR) MS offers the highest achievable mass RP and mass accuracy of any mass analyzer,<sup>4</sup> which is especially important for sequence characterization of intact proteins and site-localization of PTMs. We recently described the design and initial performance of the first 21 tesla (T) FT-ICR mass spectrometer.<sup>5</sup> At 21 T, increased ion cyclotron frequency provides higher mass RP at faster scan rates. The instrument design includes an external quadrupole ion trap (multipole storage device, MSD) located between the Velos ion trap and the ICR cell,<sup>6</sup> which is used to store multiple accumulations of analyte precursor or fragment ions prior to high-resolution mass analysis in the ICR cell.<sup>7</sup> The use of multiple ion accumulations improves spectral S/N more rapidly than spectral averaging and facilitates acquisition of multiple high-quality tandem mass spectra (MS/MS) on a time scale that is more compatible with chromatography.<sup>8</sup> The frontend of the instrument is equipped with an electric discharge-based chemical ionization source that is used to create reagents for gas-phase ion/ion reactions such as electron-transfer dissociation (ETD).<sup>9,10</sup> Electron-transfer and collision-induced dissociation (CID) techniques combine to produce hundreds of fragment ions, facilitating sequence determination and site-specific characterization of PTMs.

Here we demonstrate top-down proteomic analysis of human colorectal cancer cell lysate by liquid chromatography (LC)–MS/MS on a 21 T FT-ICR mass spectrometer. We identify hundreds of proteins expressed as thousands of unique proteoforms at a 1% false discovery rate (FDR), based on rapid, data-dependent acquisition of CID and ETD fragment ion spectra. Finally, we report significant progress toward the extension of high-throughput top-down proteomics to proteins larger than 30 kDa while maintaining isotopic resolution throughout the analysis.

## 2. EXPERIMENTAL PROCEDURES

#### 2.1. Cell Culture

DLD-1 parental (KRas wt/G13D) human colorectal cancer cells (HD PAR-086, Horizon Discovery Limited, Cambridge, U.K.) were grown at 37 °C and 5% CO<sub>2</sub> in RPMI 1640 (Corning-Mediatech, Manassas, VA) supplemented with 10% fetal bovine serum (Sigma-Aldrich, St. Louis, MO) and 0.5% penicillin/streptomycin (Corning-Mediatech). Cells were trypsinized in 0.5% trypsin-EDTA (Life Technologies, Carlsbad, CA), resuspended in serum-free RPMI 1640, counted by hemocytometer, and washed twice in 10 mL of ice-cold 1× Dulbecco's PBS (Life Technologies) prior to centrifugation at 200*g* for 10 min at 4 °C to remove residual media and serum proteins. Cell pellets, each comprising  $2 \times 10^7$  cells, were stored dry at -80 °C prior to lysis and protein quantitation.

#### 2.2. Sample Preparation

Cell pellets were thawed on ice and resuspended in 1 mL of 20 mM Tris, pH 7.5, containing 100 mM sodium chloride, 1% (w/v) N-lauroylsarcosine, and 1× final concentration of HALT protease/phosphatase inhibitor cocktail (EDTA-free) (Thermo Fisher Scientific, San Jose, CA). Lysates were incubated on ice for 20 min. Magnesium chloride was added to a final concentration of 1 mM, followed by 750 units of benzonase nuclease (Sigma-Aldrich). Lysates were incubated at 37 °C for 20 min, chilled on ice, and centrifuged at 16 800g for 15 min at 4 °C to pellet cellular debris. Total lysate protein concentration was determined by microplate BCA assay (Thermo Fisher Scientific). Following protein quantitation, 400  $\mu$ g of protein from each lysate was precipitated in acetone and incubated at -80 °C overnight. Pellets were reconstituted in 100  $\mu$ L of 1% (w/v) SDS containing 50  $\mu$ M DTT and 1× Trisacetate sample buffer (Expedeon, San Diego, CA). Samples were incubated at 95 °C for 5 min and centrifuged at 16 800g for 10 min at room temperature to pellet any remaining debris. Supernatants were loaded into a 10% acrylamide monomer (%T) gel-eluted liquidfraction entrapment electrophoresis (GELFrEE) cartridge and resolved into 12 fractions according to the manufacturer's protocol (GELFrEE 8100 Fractionation System, Expedeon). Aliquots (10 µL) from each GELFrEE fraction were resolved by SDS-PAGE and visualized by silver nitrate stain<sup>11</sup> to evaluate total protein content (example shown in Figure S-1). Eluted fractions were stored at -80 °C. Directly prior to LC-MS/MS analysis, fractions were precipitated with a mixture of methanol, chloroform, and water to remove SDS.<sup>12</sup> After the final MeOH wash, pellets were immediately reconstituted in 50 µL of ice-cold HPLC Solvent A (0.3% formic acid and 5% acetonitrile (v/v) in water; all MS grade) with gentle pipetting.

#### 2.3. Liquid Chromatography

Reconstituted protein fractions were optionally diluted up to 5-fold in ice-cold HPLC solvent A (based on silver nitrate stain intensity) and analyzed by reverse-phase LC-MS/MS. For each injection, 5 µL was loaded onto an in-house-fabricated 360 µm o.d. × 150  $\mu$ m i.d. fused-silica microcapillary trap column packed 2 to 3 cm with PLRP-S resin (5  $\mu$ m particle, 1000 Å pore, Agilent Technologies, Palo Alto, CA) or with Poroshell 300-SB C8 resin (5 µm particle, 300 Å pore, Agilent Technologies). The nano-HPLC system (ACOUITY M-Class, Waters, Milford, MA) was operated at 2.5 µL/min for loading onto the trap column and washed with 95% A for 10 min. Separation was achieved on an in-housefabricated 360  $\mu$ m o.d.  $\times$  75  $\mu$ m i.d. fused-silica microcapillary analytical column packed 15 cm with PLRP-S or C8 resin (same as corresponding trap columns). For Sample Set 1, all fractions were eluted at 0.3  $\mu$ L/min using a gradient of 5–20% B in 5 min, 20–40% B in 20 min, 40-60% B in 40 min, 60-75% B in 15 min, and 75-95% B in 5 min (85 min total length). For Sample Set 2, fractions 5–8 were eluted with a gradient of 5–20% B in 5 min, 20-35% B in 20 min, 35-60% B in 75 min, 60-75% B in 15 min, and 75-95% B in 5 min (120 min total length). The gradients utilized solvent A, 0.3% formic acid and 5% acetonitrile in water, and solvent B, 47.5% acetonitrile, 47.5% 2-propanol, 4.7% water, and 0.3% formic acid (% all expressed as v/v). Following separation, samples were directly ionized by microelectrospray ionization using a 15  $\mu$ m fused-silica PicoTip (New Objective, Woburn, MA) emitter, which was packed with 2-5 mm PLRP-S resin to minimize the formation of bubbles and promote stable ESI.

#### 2.4. Mass Spectrometry

The instrument was operated in data-dependent mode using Xcalibur software (Thermo Fisher Scientific). Precursor (MS1) and product (MS2) ion spectra were collected in the ICR mass analyzer at 21 T. For Sample Set 1 (DLD-1 A), instrument parameters were set as follows: For MS1 spectra –  $RP = 150\ 000$  at  $m/z\ 400\ (\sim 381\ ms\ transient\ duration);\ 1E6$ automatic gain control (AGC) target; 3 transients (microscans,  $\mu$ S) summed per spectrum. The RP was chosen as the minimum required for isotopic resolution of proteins up to  $\sim 60$ kDa to maximize spectral acquisition rate and number of detected proteins and proteoforms. For MS2 spectra –  $RP = 150\ 000$  at  $m/z\ 400$ ; 7.5E5 AGC target; 4 fragment ion fills of the MSD; CID activation used 41% normalized collision, 10 ms activation period energy and 0.4q, 15 m/z isolation window; dynamic exclusion with a repeat count of 1, repeat duration of 240 s and an exclusion duration of 240 s. For Sample Set 2 (DLD-1 B and C), replicate injections alternated between CID and ETD fragmentation. Instrument parameters were as described above with the following exceptions: For fractions 1-4, ETD MS2 spectra - 2E5 precursor AGC target; 4E5 ETD reagent (fluoranthene) AGC target; 10 fragment ion fills of the MSD; 20 ms reaction period. For fractions 5–8, MS1 spectra –  $RP = 300\ 000$  at  $m/z\ 400$ (~762 ms transient duration); 1E6 AGC target; 3 ion fills of the MSD; 2  $\mu$ S. For fractions 5– 8, MS2 spectra -12 ms ETD reaction period; 12 ETD fragment ion fills of the MSD; CID activation used 0.25q. Total data acquisition times were as follows: DLD-1 A fractions 1–8, 880 min (110 min/run, 8 LC-MS/MS runs); DLD-1 B and C, fractions 1-4, 1760 min (110 min/run, 16 LC-MS/MS runs); DLD-1 B and C, fractions 5-8, 2320 min (145 min/run, 16 LC-MS/MS runs). In sum, the data took 82.7 h to acquire.

#### 2.5. Data Analysis

Raw data were submitted to the National Resource for Translational and Developmental Proteomics (Northwestern University, Evanston, IL) for processing on the TDPortal highperformance computing environment at Northwestern University (available for academic collaborators here: http://nrtdp.northwestern.edu/tdportal-request/). Instrument data were then processed using an in-house-developed spectral averaging strategy that involved averaging 30 s windows of reduced profile MS1 data prior to converting data to centroid and use of Thermo Fisher's Xtract deconvolution algorithm to decharge and deisotope the spectra. For all analyses, a standardized three-pronged search strategy was employed against a database of  $\sim 1 \times 10^7$  candidate proteoforms created from the 2016 04 release of the Swiss-Prot human proteome. The search strategy used a tree made of three modes as defined for ProSight PTM 2.0<sup>13</sup>: first a narrow absolute mass search (with an MS1 tolerance of 2.2 Da and 10 ppm tolerance for MS2), then a biomarker search (akin to a no-enzyme type search in peptide data analysis; MS1 and MS2 tolerance of 10 ppm), and finally a wide absolute mass search (MS1 tolerance of 200 Da and 10 ppm tolerance for MS2, with  $\Delta m$ mode activated to find unexpected modifications). Data derived from each biological replicate were analyzed separately. Searches for all raw files were completed in 46.9 h.

Estimations of FDR instantaneous *q*-values at the protein entry and proteoform level were performed using an in-house-developed target-decoy system to be detailed in a separate manuscript. In brief, local FDR calculations were performed by the method previously introduced in ref <sup>14</sup>, which involved searching a scrambled database with the MS1 and MS2

data. All resulting hits were taken as incorrect; the distribution of incorrect hits was then used to estimate the null distribution of the scoring metrics. The probability of a forward hit receiving the observed score, or better, due to chance alone was calculated from this null distribution and then combined with conservative corrections for multiple tests<sup>15</sup> to accurately estimate FDR with minimum influence from unknown dependencies. Global FDR calculations were calculated by pooling the local FDR results from multiple search strategies and employing the "pick best" approach.<sup>16</sup> Note that an FDR is determined at multiple levels (e.g., protein and proteoform), so by default we chose to report only those proteoforms that map to protein entries scoring at an FDR of 1% or better, such that each reported proteoform asserted to be present links to a protein entry. Post-search results were analyzed using the TDViewer version 0.9.0.10 (http://topdownviewer.northwestern.edu). Lists of identified proteins and proteoforms at 1% FDR were exported into Microsoft Excel and either collated into Venn diagrams with Venny 2.1 (http://bioinfogp.cnb.csic.es/tools/ venny/) or compared within histograms generated by Microsoft Excel 2013. Specific examples highlighted below were manually validated. The .tdReport files, containing lists of identified protein entries and proteoforms, their respective observed sequence coverage, qvalues, C-scores, and other statistics, as well as all .raw files and the .txt file used in the creation of the search database are available for download from the MassIVE repository with identifier/username MSV000079978 (ftp://massive.ucsd.edu/MSV000079978/).

## 3. RESULTS AND DISCUSSION

Our primary aim was to determine the ability of the LC 21 T FT-ICR MS/MS platform to identify and characterize intact proteins and proteoforms from complex biological samples. We pursued a prototypical sample of whole-cell lysate generated from a well-characterized human colorectal cancer cell line<sup>17,18</sup> and fractionated into discrete molecular weight (MW) ranges by use of GELFrEE at 10% T.<sup>19</sup> GELFrEE fractions 1–8, containing proteins between approximately 5 and 50 kDa (Figure S-1), were collected from three biological replicates (hereafter designated as DLD-1 A, B, and C) and divided into sample sets for two experiments: (1) analysis of DLD-1 GELFrEE fractions by single-injection, top-down LC-MS/MS at 21 T (Sample Set 1, DLD-1 A) and (2) MS2 fragmentation method comparison (Sample Set 2, DLD-1 B and C). All samples were subjected to reverse-phase LC-MS/MS on a 21 T FT-ICR mass spectrometer with the goal of maximum protein identification and proteoform characterization in the minimum possible time. Data were searched concurrently against forward and decoy databases on the TDPortal with an average time requirement of 16 h per .raw file; the number of proteins and proteoforms identified in each biological replicate is shown in Table S-1. The C-score, a recently introduced metric that quantifies confidence in proteoform identification and characterization, can be interpreted as follows: Those C-scores below 3 indicate a proteoform that has been neither confidently identified nor characterized, those C-scores between 3 and 40 indicate a proteoform that has been confidently identified but not fully characterized, and those C-scores above 40 indicate a proteoform that has been confidently identified and extensively characterized.<sup>20</sup> Proteoforms identified with C-scores of 40 or better from DLD-1 A, B, and C are listed in Tables S-2, S-3, and S-4, respectively.

#### 3.1. Single Injections of GELFrEE Fractions Analyzed at 21 T

For Sample Set 1, single injections of DLD-1 A GELFrEE fractions 1–8 were subjected to LC-MS/MS analysis utilizing an 85 min LC gradient and top 2 data-dependent CID fragmentation. The resulting eight .raw files were searched against the database of candidate human proteoforms, which resulted in the identification of 580 unique proteins (defined here by UniProt accession numbers) observed as 1820 unique proteoforms (defined here by Proteoform Record, PFR; Consortium for Top-Down Proteomics Proteoform Repository http://repository.topdownproteomics.org/) at 1% FDR. The number of proteoforms identified per injection ranged from 141 to 593 (Figure 1A). Figure 1B shows the distribution of instantaneous q-values<sup>21</sup> for unique proteins (with a median q-value of  $5 \times 10^{-19}$ ), and Figure 1C shows the distribution of C-scores<sup>20</sup> for unique proteoforms identified from Sample Set 1. A total of 792 proteoforms (44%) had C-scores of 40 or above and 445 (24%) had C-scores between 3 and 40, indicating that a total of 1237 proteoforms were identified and at least partially characterized from a single GELFrEE separation of 400  $\mu$ g of wholecell lysate protein analyzed by eight LC-MS/MS runs. Of the 1820 proteoforms reported, 1360 matched to a proteoform in the database within 2.2 Da. The remaining 460 proteoforms were identified with mass shifts corresponding to multiple undocumented PTMs, artificial adducts, or cleavage events.

Representative data obtained from a single DLD-1 A GELFrEE fraction (fraction 4) are shown in Figure 2. The total ion chromatogram (TIC) is shown (Figure 2A) along with single-scan MS1 spectra, which show the charge-state distributions of proteins eluting over 10 selected peaks. The protein identities were manually validated, and global q-values and C-scores are given. The data acquisition period ranged from 1.67 to 3.17 s/spectrum for MS1 spectra, which was collected as the sum of three 381 ms transient acquisitions (3  $\mu$ S) to increase S/N for accurate deconvolution and further analysis.<sup>22</sup> CID MS2 elapsed scan period ranged from 1 to 3 s and was collected as a single 381 ms transient (1  $\mu$ S) following four fragment ion fills of the MSD. Maximum injection period was set to 500 ms for both scan types; up to 2 s of the quoted scan period was used solely for precursor ion accumulation. A single duty cycle (1 MS1, 2 MS2) was typically between 3 and 6 s. A total of 2422 spectra were acquired across the chromatogram shown (10–100 min). A total of 248 unique UniProt entries (observed as 561 proteoforms) were detected at a 1% FDR in this single experiment. A histogram giving the MW distribution of observed proteoforms is displayed in Figure 2B, which is consistent with the expected MW range based on the stained gel shown in Figure S-1.

A single-scan CID MS2 spectrum (1.23 s elapsed scan time) of a protein eluting at ~41 min (indicated in green in the chromatogram) is shown in Figure 2C. The precursor ion was [M +16H]<sup>16+</sup> with S/N of ~41:1 (range for all charge states was 102:1 to 5:1). This protein was detected with ~2.1 ppm intact mass error following deconvolution and was identified as translationally-controlled tumor protein (p23, UniProt P13693, ~20 kDa) with no modifications. A total of 54 fragment ions were matched to the putative sequence with an RMS error of 0.72 ppm (10 ppm allow tolerance; Figure 2D). The most abundant isotopologues in each multiplet matched to the putative sequence had S/N ratios from ~486:1 to ~5:1 and are given in Table S-5 along with observed RP and mass error. Taken

together, these data exemplify the wealth of intact protein and proteoform information that can be extracted from a single injection of a complex biological sample analyzed by LC-FT-ICR MS/MS at 21 T.

#### 3.2. CID and ETD Fragmentation Analyzed at 21 T

In addition to collision-induced (ion trap CID/beam-CID) fragmentation, the 21 T FT-ICR mass spectrometer is equipped with a front-end reagent ion source that is used to ionize fluoranthene for ETD fragmentation in the high pressure cell of a Velos Pro dual cell ion trap assembly.<sup>10</sup> For sample set 2 (DLD-1 B and C), we compared the number of proteins and proteoforms identified by CID versus ETD fragmentation (performed as technical duplicates). Note that ETD is typically carried out with a smaller precursor ion population than CID because precursor ions are confined to the smaller, rear section of the trap as reagent ion is introduced.<sup>9,23</sup> The use of small precursor ion population combined with the charge-destructive nature of ETD and increased sequence coverage afforded by ETD<sup>24,25</sup> dilutes the remaining ion signal of any single fragment ion. Such reduced signal necessitates the use of more fragment ion fills of the MSD to achieve S/N comparable to CID MS2 spectra. Therefore, 10–12 fragment ion fills were used for ETD MS2. At a 500 ms maximum injection time per fragment ion fill, acquisition of one ETD spectrum was observed to take up to 9 s, as opposed to just 3 s for CID. Up to approximately 3-fold lower duty cycle was observed, with fewer spectra collected overall.

Figure 3 summarizes the results from Sample Set 2; four injections each (two biological replicates, DLD-1 B and C, two technical replicates - CID/ETD) of GELFrEE fractions 1–8 (32 total LC–MS/MS runs). A combined total of 538 unique proteins and 2476 proteoforms were identified at an FDR of 1% by the TDPortal-supported searches. Venn diagrams comparing the proteins and proteoforms identified within each biological replicate (DLD-1 B or C) by CID or ETD fragmentation can be seen in Figure S-2. Instantaneous *q*-value distributions for the proteins as well as C-score and MW distributions for the proteoforms identified within DLD-1 B or C can be seen in Figures S-3 and S-4. For the DLD-1 B and C datasets, 75 and 72% of the unique proteins were detected by both CID and ETD, with CID identifying slightly more protein entries than ETD. At the proteoform level, only 36 and 35%, respectively, were identified by both fragmentation techniques, and CID identified up to 228 (13%) more unique proteoforms than ETD. We believe that these discrepancies are due to both the longer time required to obtain ETD MS2 spectra and to the complexity of the samples studied, i.e., that more proteoforms remain to be discovered within each sample.

Representative MS2 data from Sample Set 2 are shown in Figure 4. Single-scan CID and ETD MS2 spectra are shown for  $[M+26H]^{26+}$  (1.5 s scan acquisition) and  $[M+29H]^{29+}$  (3.3 s scan acquisition), respectively, of the 20 kDa protein peptidyl-prolyl *cis–trans* isomerase B (UniProt P23284, PFR 1364), taken from two replicate injections of fraction 5 (DLD-1 C). Zoom insets are shown and fragment ions are labeled. In both cases, the protein was detected with  $\mathcal{L}$  ppm intact mass error and identified with high confidence (*q*-value  $\leq 5 \times 10^{-49}$ ). Note the greater degree of sequence information from the ETD MS2 spectrum (51% of possible bond cleavages) compared with the CID MS2 spectrum (21% of possible bond cleavages). As can be seen in Figures S-3 and S-4, C-scores for proteoforms identified by

ETD fragmentation skewed higher than those identified by CID. ETD fragmentation also resulted in higher overall median *q*-values for proteins identified within DLD-1 B ( $3 \times 10^{-17}$ ) and DLD-1 C ( $3 \times 10^{-20}$ ) compared with those identified by CID fragmentation ( $1 \times 10^{-13}$  and  $2 \times 10^{-14}$ , respectively). This is not surprising because ETD tends to fragment large, highly charged precursors more efficiently than CID, resulting in higher sequence coverage and enabling more precise PTM site localization.<sup>25,26</sup>

#### 3.3. Analysis of Larger Proteins

More than half of predicted human proteins are >30 kDa,  $^{14}$  so it is crucial that top-down methods improve at higher mass. We expect that the high-mass RP and dynamic range afforded by 21 T FT-ICR MS will push the boundaries of protein identification and characterization to well above 30 kDa. Toward that goal, we used DLD-1 cell lysate as a test case. For Sample Set 1 (DLD-1 A), the observed molecular weight distribution of unique proteoforms identified at 1% FDR is shown in Figure 5A. A total of 228 proteoforms with MW greater than 30 kDa were identified at 1% FDR; this represents ~13% of the total number of unique proteoforms within that sample set. All proteoforms larger than 30 kDa were observed in fractions 5–8 alone (four total injections for DLD-1 A). Confidence metrics for proteoforms identified at 1% FDR are shown in Figure 5B, in which C-scores (red) and log (q-values) (blue) are plotted against proteoform MW. As expected, confidence metrics for proteoforms larger than 30 kDa were lower than those for smaller proteoforms, which we believe can be at least partially attributed to the need for better chromatographic resolution of larger proteins. A large number of proteins were observed to coelute during analyses of fractions 7 and 8, resulting in very low (<10) S/N in the MS1 spectra and chimeric MS2 spectra, which confounded subsequent data analysis. We plan to investigate the use of chromatographic methods better optimized for the larger proteins that we expect to observe in fractions 9–12 (~50–100 kDa).

Typical results for proteins larger than 30 kDa are shown in Figure 6. MS1 data depicting charge-state distributions for three proteins are shown (Figure 6B) along with zoom insets that show selected isotopic distributions (right). Two of the three distributions shown were resolved in a single MS1 scan (3  $\mu$ S, green and red). Resolution and charge-state determination for the 54 kDa protein required post-FT spectral averaging (average of 20 scans). We were able to determine these protein identities via *de novo* sequencing from the corresponding CID MS2 spectra (example shown in Figure 6C). All results were consistent with the identifications returned by subsequent database searching via TDPortal.

## 4. CONCLUSIONS

The 21 T FT-ICR instrument at the National High Magnetic Field Laboratory is particularly well equipped for high-throughput top-down proteomics due to the high RP obtained per unit time and the use of an external MSD to perform multiple precursor or fragment ion fills prior to high-resolution mass analysis. Multiple fills provide a necessary boost in S/N, which, when combined with the transient acquisition rate and RP achieved at 21 T, enable acquisition of high-quality MS2 spectra that facilitate intact protein identification on a chromatographic time scale. Here we demonstrate high-throughput proteoform identification

by LC-FT-ICR MS/MS at 21 T. Initial experiments produced unparalleled results on the basis of number of identified proteins and proteoforms (580 UniProt entries expressed as 1820 unique proteoforms at 1% FDR for DLD-1 A) per total number of injections (eight individual LC–MS/MS runs) compared with previous top-down proteomics studies.<sup>14,27,28</sup> For example, the largest previous top-down proteomics study focused on proteins <30 kDa and achieved 1063 unique proteins at 1% FDR from 423 LC–MS/MS runs.<sup>27,29</sup> Here >50% of the coverage (on the protein entry level) was achieved in <2% of the number of comparable LC runs, demonstrating that LC-21 T FT-ICR MS/MS is the current state of the art method for high-throughput top-down proteomics.

We also compared CID and ETD in terms of optimal scan rate, number of proteoform identifications, and characterization confidence. Here we demonstrate that ETD outperformed CID with regard to confidence in protein identification and proteoform characterization (defined here with the scoring metrics, *q*-value, and C-score, respectively), which is consistent with prior studies.<sup>14,24,26,30,31</sup> However, a smaller precursor ion population is used,<sup>9,23</sup> and total ion current is diluted among more fragment ion channels, which requires more fragment ion fills of the MSD prior to high-resolution mass analysis. Ultimately, this results in a longer acquisition period required for ETD MS2 spectra. We observed that CID outperformed ETD with regard to total number of proteoforms identified. This, we presume, is due to a combination of faster spectral acquisition rates and higher S/N observed in CID MS/MS spectra, as total ion current tends to be diluted through fewer, more specific fragment ion channels.<sup>25,26,32,33</sup> In the future, we intend to explore ways to increase the initial precursor ion population for ETD<sup>23</sup> to lessen the need for more fragment ion fills and increase spectral acquisition rate. We have also implemented ultraviolet photodissociation<sup>34</sup> within the ICR cell and plan to assess its utility for high-throughput proteoform identification and characterization.

Analysis of intact proteins becomes more challenging as protein mass increases. Ion current is distributed among more charge states, isotopologues, adducts, and fragment ion channels.<sup>22</sup> Additionally, space-charge capacity in the linear trap scales inversely with m/ $z^{35-37}$  and data analysis is more difficult. These challenges are reflected by the focus of several recent studies on proteins <30 kDa or on the analysis of whole cell lysates derived from prokaryotes, which typically have proteomes comprising a lower molecular weight range.<sup>29,38–41</sup> We were able to identify 372 proteoforms larger than 30 kDa at 1% FDR with only 20 LC injections (GELFrEE fractions 5-8, all bioreplicates, duplicates removed), setting new efficiency targets for isotopically resolved proteoform analysis. Furthermore, we believe that we can improve performance by optimizing the chromatographic separation of proteins larger than 30 kDa. We expect that improved sample resolution and optimal MS2 transient summing should combine with the high mass accuracy and ultrahigh RP achieved at 21 T to further improve the number of identifications for proteins up to or exceeding 60 kDa on a time scale that remains compatible with chromatography. Increasing the average MW range of the human proteome available to investigation by high-throughput top-down proteomics by even 20 kDa could facilitate the discovery of potentially thousands of new proteoforms, a significant portion of which might have direct clinical relevance in human disease or become dysregulated in human cancers.

## **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

## Acknowledgments

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## ABBREVIATIONS

MS	mass spectrometry
PTM	post-translational modification
S/N	signal-to-noise ratio
RP	resolving power
FT-ICR	Fourier-transform ion cyclotron resonance
Т	tesla
MSD	multipole storage device
MS/MS	tandem mass spectrometry
ETD	electron-transfer dissociation
CID	collision-induced dissociation
LC	liquid chromatography
FDR	false discovery rate
%T	total acrylamide and bis-acrylamide monomer in g/100 mL
GELFrEE	gel-eluted liquid fraction entrapment electrophoresis
MS1	precursor mass spectrum
MS2	fragment ion spectrum
AGC	automatic gain control
μS	microscans
q	ion trap CID excitation q value
MW	molecular weight

**PFR** proteoform record number

TIC total ion chromatogram

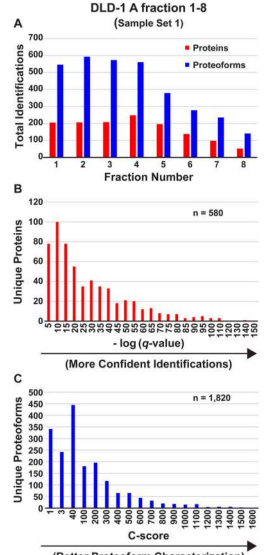
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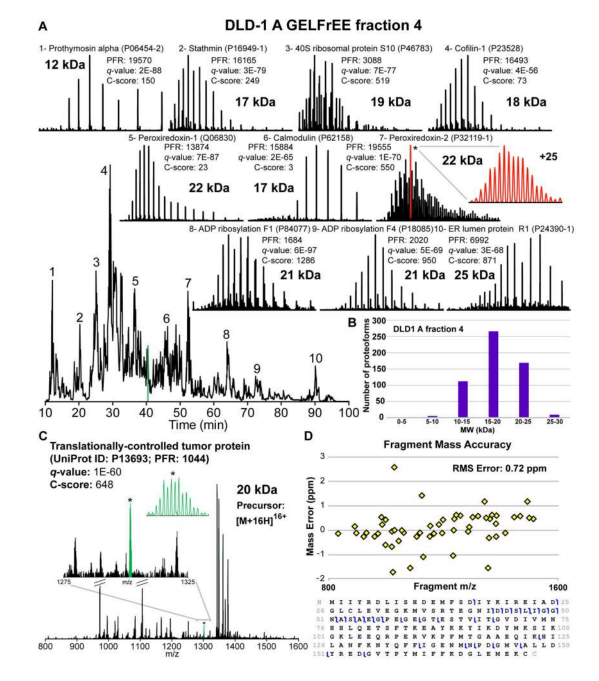
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#### (Better Proteoform Characterization)

#### Figure 1.

Protein and proteoform identification and scoring metric distributions. Data from Sample Set 1 (DLD-1 A). (A) Total number of proteins (red) and proteoforms (blue) identified at 1% FDR per single LC–MS/MS injection of each of the eight 10% GELFrEE fractions of whole cell lysate. (B) –log (*q*-value) distribution for the 580 unique proteins (unique UniProt accession numbers) identified at 1% FDR. (C) C-score distribution for the corresponding 1820 unique proteoforms. For (B) and (C), each bin comprises the sum of identifications with scores between that bin and the one to the left, save for the lowest, which comprises the sum of the identifications between that number and zero.

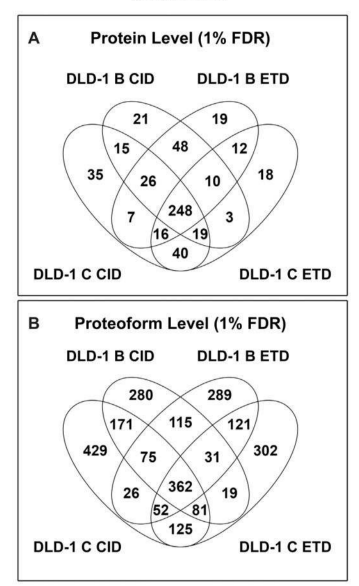


#### Figure 2.

LC–MS/MS of a single injection of GELFrEE fraction at 21 T. Data from DLD-1 A (Sample Set 1) GELFrEE fraction 4. (A) Total ion chromatogram (TIC). Ten example identifications are indicated (with UniProt and PFR numbers) along with single-scan broadband mass spectra, *q*-values, and C-scores. (B) Histogram depicting the molecular weight distribution of proteoforms identified in the fraction; 248 unique UniProt entries (observed as 561 proteoforms) were detected at a 1% FDR in this single LC–MS/MS analysis. (C) Single-scan CID MS2 spectrum taken at ~41 min (green label in TIC shown in panel A) identified as translationally controlled tumor protein, p23. (D) Top: fragment mass error (ppm) versus

m/z and sequence coverage obtained from (C). Bottom: 54 fragments were matched to the putative sequence within a 10 ppm mass tolerance (RMS error 0.72 ppm, S/N  $\approx$  500–5:1), yielding  $\sim$ 17% sequence coverage.

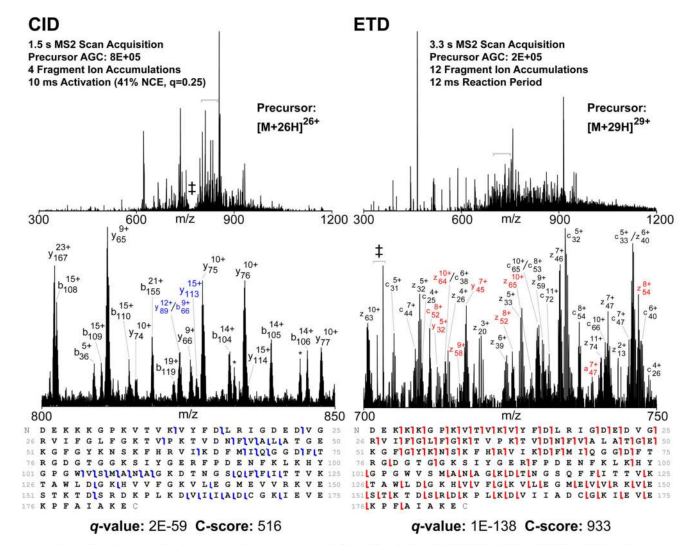




#### Figure 3.

Differential protein and proteoform identification by CID and ETD. (A) Venn diagram showing the proteins (unique UniProt accession numbers) identified within Sample Set 2 (DLD-1 B and C, GELFrEE fractions 1–8). 248 identified proteins were observed in all four replicate analyses, but CID and ETD identified distinct subsets of unique proteins. (B) Venn diagram of proteoforms (unique PFRs) identified within Sample Set 2. 362 proteoforms were observed in all four replicate analyses, but CID and ETD resulted in the identification of discrete subsets of unique proteoforms.

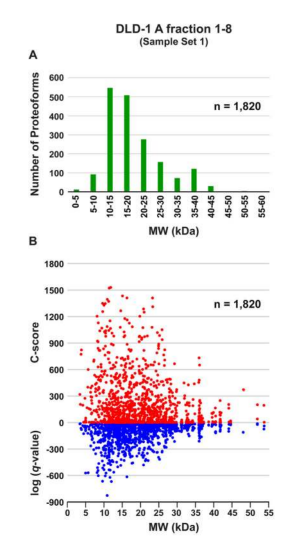
Anderson et al.



Peptidyl-prolyl cis-trans isomerase B (UniProt ID: P23284; PFR: 1364; 20 kDa)

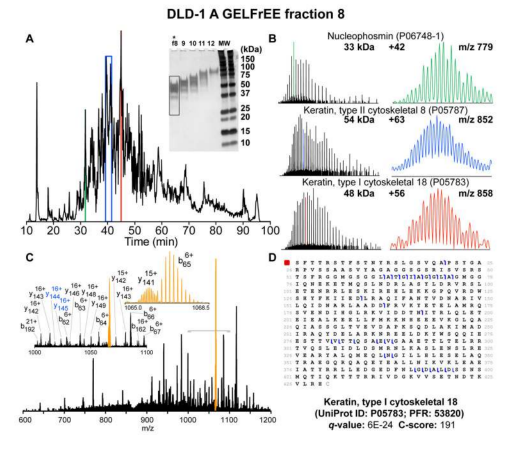
#### Figure 4.

CID and ETD MS2 spectra of peptidyl-prolyl *cis–trans* isomerase B. (Top) Single-scan CID (Left) and ETD (Right) MS2 spectra of a precursor identified as peptidyl-prolyl *cis–trans* isomerase B (UniProt P23284 residues 34–216, PFR 1354). Expanded views (Center) of 50 m/z regions (denoted by gray brackets) reveal the wealth of identified fragments. Colored fragment labels indicate multiplets identified by manual inspection only. \* = neutral loss of water from nearby fragment;  $\ddagger$  = coisolated chemical noise/activation window. Sequence coverage maps and confidence metrics determined by TDPortal are given (Bottom).



#### Figure 5.

Proteoform molecular weight distribution and scoring metrics as a function of molecular weight. Data from Sample Set 1 (DLD-1 A). (A) Molecular weight distribution of proteoforms identified at 1% FDR. 1820 proteoforms were identified at 1% FDR from just 8 LC–MS/MS runs (1 injection each of GELFrEE fractions 1–8). Proteoforms are grouped by monoisotopic mass (kDa). (B) Scatter plot depicting C-score (red) and log (*q*-value) (blue) as a function of proteoform molecular weight.



#### Figure 6.

Resolved isotopic distributions of proteins with MW > 30 kDa. (A) Total ion chromatogram obtained by LC–MS/MS of DLD-1 A GELFrEE fraction 8. (B) Single-scan mass spectra show protein charge-state distributions (650–1500 *m/z*, Left) for three chromatographic peaks and zoom insets of single charge states with resolved isotopic distributions (Right). Resolution and charge-state determination for the 54 kDa protein required post-FT spectral averaging of 20 scans. (C) CID MS2 spectrum of  $[M+55H]^{55+}$  ions of keratin type 1 cytoskeletal 18 protein. Zoom insets provide expanded views of the region indicated with gray bracket. Colored fragment labels indicate multiplets identified by manual inspection only. (D) Sequence coverage map and confidence metrics for the identification of keratin type 1 by TDPortal.