Quantitative analysis of low-abundance peptides in HeLa cell cytoplasm by targeted liquid chromatography/mass spectrometry and stable isotope dilution: emphasising the distinction between peptide detection and peptide identification

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We present the application of a targeted liquid chromatography/mass spectrometry (LC/MS) approach developed on a linear ion trap for the evaluation of the abundance of cytoplasmic proteins from a HeLa cell extract. Using a standard data-dependent approach, we identified some specific peptides from this extract which were also commercially available in their AQUA form (use for absolute quantitation). For some of the peptides, we observed a non-linear response between the intensity and the added quantity which was then fitted using a quadratic fit. All AQUA peptides spiked into a mix of 3 μg of the HeLa cell digest extract were detected down to 16 fmol. We placed an emphasis on peptide detection which, in this study, is performed using a combination of properties such as three specific Q3-like ion signatures (for a given Q1-like selection) and co-elution with the AQUA peptide counterparts. Detecting a peptide without necessarily identifying it using a search engine imposes less constraint in terms of tandem mass (MS/MS) spectra purity. An example is shown where a peptide is detected using those criteria but could not be identified by Mascot due to its lower abundance. To complement this observation, we used a cross-correlation analysis approach in order to separate two populations of MS/MS fragments based on differences in their elution patterns. Such an approach opens the door to new strategies to analyse lower intensity peptide fragments. An in silico analysis of the human trypsinosome allows the evaluation of how unique are the sets of features that we are using for peptide detection. Copyright © 2010 John Wiley & Sons, Ltd.

A major issue of using a data-dependent approach in liquid chromatography/mass spectrometry (LC/MS) mode which is based on selecting the most intense peaks at a given time is its limitation to cope with screening complex samples characterised by large dynamic range. Recent improvements in different mass spectrometric instruments, either in their data acquisition speed or in the dynamic range they can tackle, have contributed largely to encourage large-scale studies.¹⁻⁴

Significant increase in proteome coverage has been achieved by combining the use of improved mass spectrometry instruments with various fractionation strategies such as protein or peptide fractionation and gas-phase fractionation.⁵⁻⁷ However, the gain in the number of proteins identified and ideally quantified is done at the expense of longer LC/MS acquisitions and tedious data analysis. Such a situation imposes an important constraint for designing more elaborate studies. Some of the resulting drawbacks are the difficulty in characterising large amounts of biological replicates, to design complex time course studies and to simply probe a given proteome under stress with various perturbations.

Other approaches involve targeting specific peptides using prediction tools.⁸⁻¹¹ Recently, there has been a resurgence of established MS-based approaches such as the multiple reaction monitoring (mrm) and selected reaction monitoring (srm) scanning modes to detect lower abundance species.⁴⁻¹²⁻¹⁴ Several bioinformatics tools have been recently developed in order to help to choose sets of adequate transitions.¹⁵,¹⁶

These approaches have been adapted to modern proteomic constraints, i.e. the need to identify a peptide using a search engine (e.g. MIDAS) or simply adapting MS scanning methods to different types of instrument like an ion trap.¹⁸ The advantage of using an ion trap to fragment a given mass range allows one to acquire continuously in time precursor ion spectra that can be subsequently used for identification using a search engine. Moreover, the basepeaks of specific fragment/transition ions can be extracted and then reconstituted to give the equivalent of a srn/mrm
signal post-acquisition. Although a MIDAS approach has certain advantages, it allows one to identify only fragmented peptides that generate the most intense signal at a given time. It is a limiting factor which impedes the detection of lower abundance peptides.

In this study, we use an adaptation of the targeted-MS developed initially by Arnott et al.\(^8\) by combining the product of three fragment transitions in order to improve the selectivity of the pseudo-mrm assay. From a data-dependent analysis of the HeLa cell cytoplasm extract digest, we identified a few peptides which were also commercially available in their AQUA form. A peptide detection method based on a selected combination of properties such as their mass-to-charge ratio, their retention time and the combination of the profile intensity of some of the MS/MS fragments is presented and validated with a pool of 1295 identified peptides.

Such an approach allows us to detect low-abundance peptides which are either not intense enough at any time to generate a defined MS/MS fragmentation pattern or are simply masked by more abundant species impeding their identification by a conventional search engine. The use of the combination of properties increases the level of detection when peptide separation is limited. An example is shown of a peptide which is not identified by Mascot but the elution profile of its several fragments can be extracted and separated from a stronger contaminant. By using a cross-correlation analysis approach, it is possible to separate at least two populations of fragments from various origins eluting with a difference of approximately 8 s.

### EXPERIMENTAL

#### Materials

Acetonitrile and water used for LC/MS/MS analysis or sample preparation were of HPLC quality (Fisher, UK). Formic acid (FA) was Suprapure (98–100%, Merck, Darmstadt, Germany) and trifluoroacetic acid (TFA) was 99% purity sequencing grade (Sigma, UK). All peptides and AQUA peptides were purchased from Sigma UK.

All other chemicals used in the preparation of the samples were of reagent grade or better, unless specified. Sequencing-grade modified porcine trypsin was purchased from Promega (Roche, UK). All protein and peptide standards were of >95% purity from Sigma (St. Louis, MO, USA). All connector fittings were from Upchurch Scientific or Valco (purchased from RESTEK UK). HeLa cell cytoplasmic extracts were obtained commercially from Dundee Cell Products Ltd. (UK). The cell count estimated to 2.75e7 cell/mg (±10%) was measured using a hemocytometer and cell viability was checked using trypan blue.

#### Sample preparation

A small fraction of the HeLa cell cytoplasm extract was quantified using a Bradford assay and the overall cell extract was estimated to 1 mg. The extract was diluted in water up to 300 μL and protease inhibitors were added according to the supplier (Roche, UK). To the extract, we added 125 μL urea (8 M), 25 μL ammonium bicarbonate (1 M), 25 μL dithiothreithol (200 mM), left it to stand for 30 min at room temperature and then added 25 μL iodoacetamide (500 mM) and 20 μg trypsin, and left this to digest overnight. Samples were split into aliquots and stored at –80°C.

#### High-performance liquid chromatography (HPLC) and mass spectrometry

Micro-HPLC/MS/MS analyses were performed using an online system consisting of a micro-pump Agilent 1200 binary HPLC system (Agilent, UK) coupled to a hybrid LTQ-Orbitrap XL instrument (Thermo-Fisher, UK). The LTQ was controlled through Xcalibur 2.07 and LTQ Orbitrap XL MS2.4SPI. Capillary Picotip columns (10 cm × 360 μm o.d. × 75 μm i.d.) with a 15 μm tip opening and fitted with a borosilicate frit were purchased from New Objective (Presearch, UK). Fused-silica tubing was purchased from Composite Metal (UK).

The reversed-phase bulk material used in this study was 5 μm Pursuit C18 obtained from Varian (UK). Buffer A was 97.5% H₂O, 2.5% acetonitrile, 0.1% FA, and buffer B was 90% acetonitrile, 10% H₂O, 0.025% TFA, 0.1% FA. The HPLC system was used in low consumption mode to reduce solvent consumption. For all measurements, 8 μL of sample was injected using an Agilent micro-WPS autosampler at 5 μL/min. After sample loading, the flow rate across the column was reduced to approximately 100–200 nL/min using a vented column arrangement.\(^19,20\)

Samples were analysed on either a 2 h gradient for data-dependent analysis (1survey scan in FT mode at 60 k resolution followed by 3 MS/MS in IT mode) or 1 h gradient for targeted MS/MS in IT mode only. For the 1 h, the solvent gradient program was as follows: 0% buffer B (0–6 min), 0–5% buffer B (6–8 min), 5–15% buffer B (8–18 min), 15–35% buffer B (18–40 min), 35–100% buffer B (40–48 min), followed by 100% buffer B for 9 min and back to 0% buffer B for 3 min. The timing for the 2 h was simply doubling of the steps for the 1 h gradient.

Prior to each LC/MS micro-LC/MS/MS analysis, a column/pre-column wash and conditioning step consisting of 1 h gradient of 0–100% buffer B over 20 min followed by an isocratic conditioning step at 0% buffer B over 40 min was performed. The MS instrument transmission was optimised using mostly the +1 charge state of the standard peptide MRFA (Sigma, UK), in static nanospray mode (Proxeon, Denmark). The ESI voltage was set at 1.7 kV and the interface temperature set at 200°C for all measurements.

In data-dependent mode, the MS acquisition settings were as follows: a single FT scan at a resolution of 60k in profile mode (400–2000 Th) was first performed without the lock mass function, followed by three data-dependent MS/MS scans of the three most intense ions in LTQ centroid mode with a window of 2 Th. Selected MS/MS precursor ion data were excluded for 180 s and the exclusion list was set at 500 for a narrow isolation width of 0.05 Th. The maximum fill time and target value were 200 ms, 200 ms, 1 × 10⁶ and 1 × 10⁴ ions for the FT survey scan and IT MS², respectively.

All targeted MS/MS experiments were performed in the ion trap mode with 60 min LC/MS gradient. Up to ten MS/MS transitions were followed without the need to introduce different segments in the MS method. A narrow isolation width of 3 Th for each transition was considered.
Data processing

Mascot Generic Format (MGF) input files were generated with the EXTRACT_MSN tool (Bioworks 3.3, ThermoQuest Corp., San Jose, CA), and merged with the precursor grouping option disabled. Spectra containing less than ten fragment data points were discarded. MS/MS data were searched using Mascot version 2.2 (Matrix Science Ltd., UK) against a human subset of IPI UniProt database (EBI) (March 2008) version 3.42 with 72340 sequences.

All basic Mascot searches were performed using a maximum missed-cut value of 1, mass modifications can be found on-line.21 These are a variable methionine oxidation modification (+15.994915 Da), S,T,Y phosphorylation (+79.966331), and fixed cysteine carbamidomethylation modification (+57.021464 Da). The precursor mass tolerance was set to 12 ppm and a MS/MS tolerance of 0.4 Th was considered. The significance threshold p was set at 0.05 using the MudPIT scoring scheme. The false discovery rate based on a decoy database was below 2%. Extraction of the semi-quantitative emPAI factor22 used in this study was the output value obtained directly from Mascot.

Specific searches on the targeted MS/MS experiments were also performed. They were done in order to identify the targeted peptide as well as potential contaminants having a similar mass range or retention time. In those specific cases we used a narrow isolation width for MS/MS around 3 Th and 0.4 Th for the MS/MS fragment. Semi-trypsin was selected in order to increase the chance to identify some of the potential contaminants. For those particular searches, we considered only the highest ranking peptide hits identified at a specific retention time (RT) (close to the elution time of the AQUA peptide form). To confirm the targeted peptide identification, we use the RT information measured for the AQUA peptide form.

Extraction of the pseudo-mrm signal was performed post-acquisition as follows. For a given peptide a series of three specific fragments are selected. The choice of those fragments is defined empirically and roughly they are: high intensity fragments, at least one fragment has to be of a mass higher than the selected precursor ion, mostly b and y ions. We selected the same fragment type from the targeted peptide and the AQUA form. Exploration of the best combination of three fragments was done using the AQUA peptide at different concentrations and tested by trial and error with the criterion that it should generate a unique signal with the best limit of detection. Confirmation of the fragment assignment was done using Protein prospector from UCSF. The three selected fragments were then extracted as basepeak from the filtered specific MS/MS spectrum and exported directly into Excel where, for a selected mass range, the product of the three transitions was performed for each time point.

Cross-correlation analysis

The raw time series data consists of three columns: time, mass-to-charge ratio (m/z) and intensity. The data are filtered according to mass such that for each specific fragment one obtains the variation of intensity with time. The mass filter is set to ±0.5 Th about the expected m/z for a particular fragment. This procedure is repeated for 12 selected fragments: C1 to C6 (refers to the fragment associated with the contaminant) and P1 to P6 (refers to the fragment associated to the targeted peptide). To compute the correlation coefficient for two different fragments, one first eliminates data entries for times which are present in one peptide time series but not in the other and then computes the desired coefficient using the Excel function correl(array1,array2).

Protein organelle localisation was obtained from GoMiner.23 In the case of a small subset of proteins either not recognised or without any localisation assigned, we manually curated this set using Genecard24 as well as the Harvester.25

In silico analysis

A subset of peptides identified from a 1 h run by Mascot was exported into a csv format where a few parameters were extracted such as the sequence information, score, mass and the time at which each peptide was identified. A hydrophobic scale based on peptide sequence was then built in a similar way as previously described.8 Each peptide fragment assignment and intensity was obtained after data was parsed into MaxQuant (version 1.0.13.8). An experimental subset of 1295 peptides was used and compared to an in silico human trypsinosome. This experimental subset of 1295 peptides corresponds to all peptides from Mascot having at least a score of 20 and that was also retained in MaxQuant. The in silico analysis of the human trypsinosome was done using R (version 2.9.1). For each peptide we calculated their ratio mass/charge as 2+ and 3+, their hydrophobicity RTcalc, and we generate their theoretical b and y fragment ions.

An overall description of the experimental design in this study is presented in Fig. 1. Two different HeLa cell cytoplasm extract digests were prepared and characterised by LC/MS. The first one was done to identify peptides which are available in their AQUA form. On the second digest, the number of cell equivalents was preliminary determined prior to digest and we performed the quantitation using AQUA peptides.

RESULTS AND DISCUSSION

Experimental analysis

HeLa cell cytoplasm extracts analysis and AQUA peptide characterisation

The HeLa cell cytoplasm extract was trypsinised using a standard in-solution digest method and analysed by LC/MS using a data-dependent approach. The protein composition of the cytoplasm was analysed by LC/MS as described in the Experimental section and searched against an IPI database using Mascot. The false discovery rate using a decoy database search was below 2% for a p value of 0.05. For an initial digest, a total of 825 proteins were identified under those conditions. Under more stringent conditions; i.e. two peptides per protein having at least a Mascot score of 20, a total of 625 proteins were identified. The list of generated proteins was analysed using GoMiner for protein localisation (see Table 1). Of the 625 proteins from this digest 1, 80% (500 proteins) were reported to have a cytoplasm localisation; the
second higher fraction was for the nucleus with 10–14.5% and a similar observation for the second cytoplasm extract digest from HeLa cell. This observation regarding protein origin confirms that the cytoplasm extract we are using in this study is a good representation of the proteins that can be found in this organelle.

From the list of identified peptides from this first HeLa cell cytoplasm extract digest we have identified a small subset of peptides which are also already commercially available in their AQUA peptide forms. Having determined a precise mass accuracy for each peptide by the Orbitrap increases the level of confidence for choosing the corresponding AQUA peptides.

A total of seven commercially available AQUA peptides were identified. Two of them were rejected for the following reasons. The sequence IGPLGLsPK associated to 60S ribosomal protein L12 (RPL12 or IPI00024933) is available in its phosphorylated form which is probably not a major component in our study. The sequence TNQIGTVNDR associated to IPI00024660 isoform 1 of uncharacterised protein KIAA0174 (protein rank 821) was identified with only one peptide and was then not retained due to its relatively low abundance and also for a poor-quality MS/MS spectra.

Of the five available AQUA peptides, the sequence FYEEVHDLER, associated to NAP1L1 nucleosome assembly protein 1-like 1 (IPI00023860), ranked 137 out of 825 proteins, was identified and kept. The MS/MS spectrum associated with this sequence has a Mascot score of 26, but the protein was identified with at least eight other different peptides. Both the sequences TPAQFADDEL and GTDVNVFNTILTTR associated to ANXA1 annexin A1 ranked 21 out of 825 and finally the sequences GALQNII-PASTGAAK and LVINGNPITIFQER associated to GAPDH
glyceraldehyde-3-phosphate dehydrogenase IPI00219018 ranked 15 out of 825 were kept. All these three proteins were identified with more than one peptide in digest 1.

Although the instrument of choice for a targeted mrm assay is a hybrid triple quadrupole linear ion trap type of instrument, various studies have been conducted using an ion trap.\textsuperscript{18,26} The method consists of acquiring product ion spectra of a targeted peptide referring to Q1-like selection. The Q3-like signal is reconstituted post-acquisition. This approach allows one to choose the optimum Q3-like signal after LC/MS acquisition as well as to combine various specific Q3-like transitions in order to improve the uniqueness of the assay.

The signal intensity for an isolation width of 3 Th around 710.4 Th (Q1 like) associated with the AQUA peptide GALQNIIPASTGAAK spiked at 16 fmol into the cytoplasm HeLa cell extract digest is illustrated in Fig. 2. There are three different basepeak chromatogram curves for different Q3-like selection as detailed in the legend of Fig. 2. For all the three selected fragments, the basepeak signals show various transitions; however, one signal is present in all three curves around 37.3 min N with S/N varying between 10 for Q3 like of 597.3 Th to a S/N ratio of 90 for 823.5 Th. The pseudo-mrm signal shown in Fig. 2(D) is the result of the intensity product at a given time of the three signals measured in Figs. 2(A), 2(B) and 2(C). However, in this specific case, the final product is relatively close to the signal detected in Fig. 2(B). The intensity product of the three curves slightly increases the specificity of the peptide signal detection.

Table 2 lists information about the selected peptides and their AQUA counterparts. All the AQUA peptides were also analysed individually in static nanospray in order to ensure that we are indeed characterising the right peptide. An example is shown in Supplementary Fig. S1 (see Supporting Information), regarding the sequence FYEEVHDLER. All of the analysed fragmented AQUA peptides in MS/MS were confirmed using Protein Prospector. Their fragmentation patterns were compared to the counterpart natural peptides; in addition, we monitored their optimum charge state for maximum intensity. As also illustrated in Table 2 for both the AQUA and natural forms of a given peptide, we chose the same three fragment, ‘Q3 like’, in order to acquire a comparable pseudo-mrm signal for both counterparts.

**Pseudo-mrm quantitation assay on HeLa cell cytoplasm extract**

Various quantities of the AQUA peptide mix were added to the HeLa cell cytoplasm extract digest 2 (see Experimental section). For this study, we used 6600 and 32000 cell equivalents for the HeLa cell cytoplasm extract. The injected quantity of the AQUA peptides mix was 0, 16, 48, 160 and 480 fmol and all experiments were performed in duplicate. As illustrated in Supplementary Fig. S1 (see Supporting Information), extraction of the normalised pseudo-mrm signal for both the natural form of the peptide and its AQUA counterpart for a given LC/MS run co-elute with no discernible time lag confirming that we are characterising the right peptide through the use of the AQUA peptide form. The relationship between the pseudo-mrm signal as a function of the different AQUA peptide quantities was examined in duplicate. A recapitulative summary of this information is presented in Table 3.

Two out of the five AQUA peptides do not have a pseudo-mrm linear response in relation to the added quantity (see also Fig. 3(B)). For the lower concentration of the AQUA peptide LVINGNPITIFQER at 812.5 Th a smaller intensity response is observed and seems to increase significantly from 160 fmol and upwards.

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**Table 1. Number of protein identified in the cytoplasm extract and their putative localisation**

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<tr>
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<th>Digest 1</th>
<th>Digest 2</th>
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<tr>
<td>total protein ID</td>
<td>629 (825)</td>
<td>607 (823)</td>
</tr>
<tr>
<td>% cytoplasm</td>
<td>80.0</td>
<td>76.6</td>
</tr>
<tr>
<td>% nuclear</td>
<td>10.7</td>
<td>14.5</td>
</tr>
<tr>
<td>% plasma membrane</td>
<td>0.8</td>
<td>0.8</td>
</tr>
<tr>
<td>% other organelle</td>
<td>3.7</td>
<td>4.8</td>
</tr>
<tr>
<td>% unknown</td>
<td>5.0</td>
<td>3.2</td>
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</table>

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**Figure 2.** Signal intensity for an isolation width of 3 Th around 710.4 Th (Q1 like) associated with the AQUA peptide GALQNIIPASTGAAK spiked at 20 fmol into a cytoplasm HeLa cell extract digest. Three different basepeak chromatogram curves for different ‘Q3-like’ selection: (A) basepeak for 597.3 Th (b6), (B) basepeak for 823.5 Th (y9), (C) basepeak for 936.5 Th (y10), and (D) intensity product (A* B*C)^0.333. The peak at 37.3 min associated with this peptide is not the most intense ion for a 1 h LC/MS run in (A) and (C). The intensity product of the three curves increases the specificity of the AQUA peptide signal in (D).
Probing the underlying reasons behind the non-linear response for peptides LVINGNPITIFQER and GTDVNVFNTILTTR is beyond the scope of this paper. There is definitely an important difference between the pseudo-mrm assay done on an ion trap compared to a real-mrm assay performed on a triple quadrupole mass spectrometer. The observed non-linearity between the MS signal and the quantity for some peptides could be due to the fact that the signal intensity in an ion trap is calculated from the time it takes to fill the trap up to a target value. One major difference observed for these two peptides compared to the other peptides is at lower quantity (48 fmol), even at their maximum intensity, the target value for filling the ion trap is not reached (result not shown). A different strategy would be to reduce the ion target value and increase the number of microscans but we have to expect that the dynamic range for an ion trap for such an assay is lower than for a triple quadrupole mass spectrometer. In order to minimise the

<table>
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<tr>
<th>Protein ID</th>
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<th>Selected sequence</th>
<th>Mascot score</th>
<th>Δmass Da</th>
<th>Peptide form</th>
<th>Pseudo-mrm transition</th>
<th>RT</th>
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<td></td>
<td></td>
<td>AQUA form</td>
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<td>-0.0005</td>
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<td>2</td>
<td>673.8 778.4 907.5 1036.5</td>
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</table>

Figure 3. Variable amount of the AQUA peptide from 0, 16, 48, 160 and 480 fmol of (A) the AQUA form TPAQFDADELRL at 636.8 Th spiked in 132 000 cell equivalents of HeLa cell cytoplasm extract (natural form in open circle at 631.8 Th) and (B) LVINGNPITIFQER at 812.5 Th spiked in 6600 cell equivalents of HeLa cell cytoplasm extract (natural form at 807.4 Th). The LC gradient was 60 min and all the LC/MS runs were performed in duplicate. (C, D) Log-log scales of (A) and (B), respectively.
impact of this specific peptide non-linearity observed in our study, we prepared standard curves for each of the AQUA peptides added to a specific amount of HeLa cell extract. As described by Howells and Sauer, we used a second-order polynomial function in order to quantify the relationship between the intensity and the AQUA peptide quantity. With the exception of those two peptides that generated a non-linear response, the three other peptides generated good linear correlation (see Fig. 3(A) and Table 3). A non-linear response can have multiple origins, a saturation generated good linear correlation (see Fig. 3(A) and Table 3). generated a non-linear response, the three other peptides which cannot be deciphered by a search engine.

Estimation of the quantity present for the three proteins in the HeLa cell extracts are also presented in Table 3 in term of copies per cell. Two out of the three proteins were quantified with two peptides (i.e. GAPDH and ANXA1). For both proteins, GAPDH and ANXA1, the error estimation for each peptide is significantly smaller than the difference in the protein quantity estimation from both peptides which can be associated to variation in the digestion efficiency or peptide identification confidence. However, using such an approach increases significantly the identification confidence. Similarly, we acquired MS/MS data in continuous mode on a peptide using a protein search engine such as Mascot, a specific intensity of the mrm signal and associated to a given peptide has to generate the most intense peaks and therefore limits the detection of lower abundance species in complex mixtures. It is common in such mixtures to have several peptides having close masses and hydrophobicity which cannot be deciphered by a search engine.

For example, we can identify the protein NAP1L1 through the measured peptide FYEEVHDLER in digest 2. This is achieved by analyzing the sequence and comparing it with the database. For instance, ANXA1 and NAP1L1 are less represented in digest 2. Hence, the emPAI factor is calculated for each protein. The three proteins cover quite well the dynamic range of the emPAI factor. In the upper right corner of Fig. 4(B) a trend is observed regarding the calculated copy number per cell for the three proteins and the emPAI factor. We are aware that a more precise quantitative trend will be more meaningful using a larger set of AQUA peptides for the quantitation.

Since ANXA1 and NAP1L1 are less represented in digest 2, two out of the five selected peptides were not directly identified by Mascot using a data-dependent approach on the second digest (i.e. peptides FYEEVHDLER and TPAQFDADELR). In the targeted mode where precursor ion scans were acquired continuously for all peptides and their AQUA counterparts, we were able to easily identify the peptide TPAQFDADELR using Mascot; however, Mascot failed to identify the sequence FYEEVHDLER in digest 2.

**Peptide detection using a combination of specific properties**

Using a data-dependent mode allows the identification of the most intense peaks in a mixture. Using a targeted approach such as MIDAS, where a single MS/MS event is triggered at a specific intensity of the mrm signal and associated to a peptide using a protein search engine such as Mascot, increases significantly the identification confidence. Similarly, we acquired MS/MS data in continuous mode on a small subset of peptides and using Mascot we increased the identification confidence. However, using such an approach where detection is based on peptide identification has some limitations. The fragmentation pattern associated with a given peptide has to generate the most intense peaks and therefore limits the detection of lower abundance species in complex mixtures. It is common in such mixtures to have several peptides having close masses and hydrophobicity which cannot be deciphered by a search engine.

For example, we can identify the protein NAP1L1 through several peptides but we cannot identify the sequence FYEEVHDLER using Mascot in digest 2. The fragmentation of the two AQUA forms and natural sequence FYEEVHDLER shown in Supplementary Fig. S1 (see Supporting Information) was extracted from digest 1 and clearly shows a good fit between the AQUA and natural forms of this peptide which was also confirmed by Mascot with a score of 26. However, it is not possible to identify the sequence in digest 2.
but, as illustrated in Fig. 5, we can detect the peptides through the combination of two properties: (1) a specific Q1-like narrow isolation width, and (2) a pseudo-mrm peak eluting at a specific time confirmed by its AQUA counterpart.

The pseudo-mrm signal associated to FYEEVHDLER at 668.32 Th with a peak at 33.28 min can be seen in Fig. 5(A). In the same figure, for a Q1-like isolation width of 668.32 Th, another component is detected (red curve) and associated to a contaminant exhibiting a slight lag in the elution time compared to the peptide FYEEVHDLER (peak at 30.42 min). Also superimposed on this figure is the AQUA form of FYEEVHDLER which co-elutes with its natural counterpart (black dotted curve). The three different peaks have been normalised for comparison purposes in Fig. 5(B).

The MS/MS spectrum shown in Fig. 5(C) was recorded at the maximum of the transition for the peptide FYEEVHDLE. On the same MS/MS spectra, acquired at 30.28 min, the main contribution from both peptides are highlighted (C for the fragment associated to the contaminant VFGQSGAGNNWAK and P for the targeted peptide FYEEVHDLER). Under those conditions Mascot cannot identify the targeted peptide FYEEVHDLER as the generated MS/MS fragment at 668.3 Th is a mix of at least two different components even if the fragment intensities associated with the targeted peptides are at maximum intensity. In Fig. 5, the table on the right shows some of the peak assignments for both the contaminant and the sequence FYEEVHDLER.

The contaminant was identified as a peptide from the tubulin beta chain with a Mascot score of 54 based on a semi-tryptic search. This semi-tryptic peptide was not present in digest 1 which made the identification of FYEEVHDLER easier in that case. As illustrated in Fig. 5 in the left table, the identified contaminant VFGQSGAGNNWAK had both a close mass and hydrophobicity to the targeted peptide which renders LC/MS separation difficult. This leads to the situation where one cannot identify the targeted peptide using Mascot due to a mixed fragmentation pattern although a pseudo-mrm signal for both species is relatively strong and distinct.

In order to differentiate both peptide groups (i.e. the targeted peptide and its AQUA form from the contaminant), we extracted six known MS/MS transitions associated to each peptide to determine how we can statistically combine or separate the contaminant peptide from the targeted sequence and its AQUA form based on their difference in retention time (RT) (see value from the table in Fig. 5).

In Table 4, we have performed a Student’s test on the elution time distribution for the targeted peptide, the AQUA form and the contaminant. According to this test both forms of the targeted peptide and its AQUA form are eluting at a similar time and are distinct enough from the elution time of the contaminant. The test highlights that the three peptides can be separated into two groups. Note that although the fragmentation pattern associated with the targeted peptide FYEEVHDLER could not be identified using Mascot, we could detect it using a pseudo-mrm approach based on three MS/MS transitions as well as with the RT correlation of six MS/MS transitions.

Cross-correlation analysis of MS/MS for a 3 min time window
Another approach can be used to group or separate sets of fragments based on a slight variation in their elution profile.
Figure 5. MS/MS fragmentation of the 668.8 Th with a 3 Th width window between 29 and 32 min. (A) Highlighted in black and red on the MS/MS spectra fragments associated with the targeted peptides (low intensity FYEEVHDLER) and a stronger contaminant semi-tryptic peptide from the tubulin beta chain: VFGQSGAGNNWAK. The two peptides shared a close mass as well as a similar calculated hydrophobicity on a 1 h scale (for calculation details, see Supporting Information) as illustrated in the left table. On the MS/MS trace in (C), the targeted peptides show low intensity fragments compared to the contaminant, thus impeding its identification using a standard search engine. However, the product of three transitions illustrated in (B) show that the targeted peptide (continuous black line) co-elutes with its AQUA form (dashed) and the contaminant exhibits a slight shift in the elution pattern. The three selected Q3-like fragments for the FYEEVHDLER peptide were y6, y7 and y8 and for the contaminant VFGQSGAGNNWAK y9, y10 and b12.
Such approaches have been used with other types of spectroscopy like infrared, Raman or NMR, and referred to as two-dimensional correlation spectroscopy obtained by cross-correlation analysis. Such a tool has been previously used to detect similarity across different spectra in order to define an optimal set of transitions to represent a peptide by an MRM type of assay. Applied to our particular case, group of fragments that co-elute will generate a correlation signal. In Fig. 6 the time window is limited to 3 min (from 29 to 32 min) and only 12 fragments are interrogated (6 from the Table 4. Student test on the RT distribution for the natural and AQUA form of FYEEVHDLER and the contaminant VFGQSGAGNNWAK

<table>
<thead>
<tr>
<th>Peptides comparison</th>
<th>Degree of freedom</th>
<th>$T_{theor}$ $\alpha = 0.05$</th>
<th>$T_{calc}$</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 vs. 2</td>
<td>10</td>
<td>2.23</td>
<td>1.92</td>
<td>Acceptance of the equality hypothesis of the RT$_{max}$ value for an $\alpha = 0.05$</td>
</tr>
<tr>
<td>1 vs. 3</td>
<td>10</td>
<td>2.23</td>
<td>11.77</td>
<td>Rejection of the equality hypothesis of the RT$_{max}$ for an $\alpha = 0.05$</td>
</tr>
<tr>
<td>2 vs. 3</td>
<td>10</td>
<td>2.23</td>
<td>15.88</td>
<td>Rejection of the equality hypothesis of the RT$_{max}$ for an $\alpha = 0.05$</td>
</tr>
</tbody>
</table>

1 FYEEVHDLER
2 FYEEVHDLER (AQUA form)
3 VFGQSGAGNNWAK (contamination)

Note: Hypothesis: Normal distribution of RT$_{max}$

Figure 6. Cross-correlation analysis for selected sets of identified fragments for a Q1-like of 668.8 Th between 29 and 32 min time window. (A) Table showing the magnitude of the correlation coefficient for the time series of the selected fragment pairs and (B) colour density plot of the correlation coefficient data shown in the table above.
targeted peptides P1 to P6 and 6 from the identified contamination C1 to C6. Ideally, such a procedure can be extended and optimised to longer time windows in an automated manner where each of the fragments are interrogated against each other and clustered based on their correlation factor. However, such a global analysis is beyond the scope of this work.

The average cross-correlation pairing extracted from Fig. 6(A) for P-P fragments was $0.77 \pm 0.23$ while for the C-C fragments the cross-correlation was $0.93 \pm 0.06$. A lower cross-correlation for a C-P pair of $0.36 \pm 0.17$ has been measured. A more detailed analysis of those cross-correlations is illustrated in Fig. 6(B) where the dark green zone is associated with strong correlation while the white area implies lack of correlation. The fragment p3 (associated with the y4 ion from the targeted peptide) has in general a lower correlation with the rest of the P fragments whilst the fragment p5 (ion y7 at 897.4 Th for the targeted peptide) shows some level of cross-correlation with the fragments c3, c4, c5 and c6. A more in-depth analysis of the fragmentation pattern of the contaminant peptides VFGQSGAGNNWAK highlights the possibility of a fragment with multiple loss of $b_{10}$-$H_2$O-$NH_3$ at approx. 897.4 Th which could explain this higher correlation value between p5 and the C group.

**In silico human proteome digest and interrogation based on using the concept of combination of properties**

The concept of combining mass and retention time as a tool to interrogate different proteomes has been described previously. In Supplementary Fig. S3 (see Supporting Information), we illustrate a global representation of the tryptic digest for the human proteome defined in the following space: mass/charge as a function of the calculated elution time. The elution time was calculated using the measured 1 h hydrophobic scale (see Supplementary Fig. S4, Supporting Information). The space coverage is quite similar to that previously reported. Short peptides are more dominant with an elution time centered around 30 min in our case.

As the retention time and a given mass are not enough to identify a given peptide, we introduce the use of at least three specific MS/MS fragments. A total of 1295 peptides identified from a 1 h LC/MS run were interrogated against the human trypsinosome.

For each of the experimental 1295 peptides we extracted their mass/charge as a double and triple charge, the time where they were selected for MS/MS (close to the RT), and we also extracted their identified MS/MS fragments. We use the combination of RT and mass/charge ratio to identify in the overall in silico trypsinosome how many peptides could also fit those criteria. We use this set of RT $\pm 8.2$ min (95% confidence for peptide elution time prediction) and sharing an $m/z$ value within a window of $\pm 1.5$ Th (either as a 2$^+$ or 3$^+$) with the experimental set of peptides. Such an $m/z$ window is similar to the one used in our pseudo-mrm assay.

An additional constraint was also imposed to complete the set of combinations of properties each set of paired experimental and in silico peptides should have: any three $y$ or $b$ fragments in common. The pairing result is shown in Fig. 7(A). On average each experimental peptide under these conditions fits at least 20 of the in silico peptides. The lack of specificity in this case is due amongst other things to a shared similarity at the level of the low mass fragment ($b_1, b_2, b_3$ or $y_1, y_2, y_3$) which are not specific enough as a unique feature; such an observation is detailed in Sherman et al.

In Fig. 7(B) we use a more stringent criterion, i.e. the most intense experimental $b/y$ ion where at least one of the fragments has a mass/charge ratio higher than the precursor ion and we eliminate any fragment ion with shorter sequence than $b_3$ or $y_3$. Those were also the criteria used for the pseudo-mrm experiments in this study. In Fig. 7(B) of the pool of 1295 peptides 61 peptides under these conditions did not return any sequence and only three more peptides were incorrectly identified which means that more than 95% of the peptides were correctly identified. Of the 1295 peptides, a total of 1027 peptides generate a unique corresponding good sequence which represents approximately 80% of them and an average of 1.26 in silico peptide per experimental one. As a consequence only 20% of the peptides generate more than one possibility with these criteria of $\pm 1.5$ Th mass/charge range, a calculated hydrophobicity of $\pm 8.2$ min and the above set of rules to select three MS/MS fragments.

Additionally, such an approach can be used to increase the specificity of the selected peptide having unique features not from a sequence point of view but rather from its physicochemical behaviour. As an example, the sequence FQIATVTEK shared with at least 30 other tryptic peptides a similar hydrophobicity, mass/charge and high stringent fragmentation pattern.

By using a basic in silico analysis of peptide properties we show that it is possible to correctly identify a peptide with a minimal level of information such as its mass, its hydrophobicity (expressed in elution time RT), and three specific

![Figure 7](Image 320x173 to 560x357)

**Figure 7.** Number of associated in silico peptides out of the experimental pool of 1295 peptides in bin of 1. (A) The sets of similar features are an $m/z$ of $\pm 1.5$ Th, a calculated hydrophobicity of $\pm 8.2$ min and at least three MS/MS fragments in common. (B) The conditions are similar except that the rule for similar fragments is more stringent; we looked at the most intense experimental $b/y$ ion where at least one of the fragments has a mass/charge ratio higher than the precursor ion and we eliminate any fragment ion shorter than $b_3$ or $y_3$. Those were the criteria used for the pseudo-mrm experiments performed in this study.
b/y fragments. In this interrogation mode we did not include the use of an AQUA peptide which adds another level of confirmation (i.e. co-elution can be monitored in the range of a few seconds) and reduces drastically the need to confirm peptide hits by a search engine.

CONCLUSIONS
In this study we have used a linear ion trap in a pseudo-mrm mode as an effective tool for targeted absolute quantitative proteomics. We also present a new way of generating a pseudo-mrm signal which is performed post-acquisition by combining the intensity product of at least three fragment transitions in order to increase the specificity of the signal.

Using different approaches we have shown that it is possible to detect a given peptide without necessarily identifying it using a search engine such as Mascot. Peptide identification using Mascot implies that at a given time its fragments generate a signal that is high enough to be the dominant one. However, such a situation limits the dynamic range of peptide detectability. Definition of a set of unique features that can be used to detect a given peptide is therefore attractive. In this study, we used a combination of various features such as a given retention time (similar to its AQUA form), and co-elution of three fragments for a small Q1-like window. We interrogate in a relatively simple manner the human proteome in order to validate the uniqueness of combining specific properties. From the set of 1295 peptides only 61 peptides did not generate a correct in silico match which can be due to the range of the calculated hydrophobicity (a window of ±8.2 min generates an interval of 95% confidence). Interestingly, more than 80% of the peptide generate a single hit which helps support the fact that peptide detection based only on specific combination of properties is possible without the need to use a search engine like Mascot.

We also explored the use of cross-correlation analysis in order to highlight the elution pattern of a group of peptide fragments from two different peptides sharing similar mass and hydrophobicity. We showed that it is possible to clearly separate most of the fragments from both peptides which are co-eluting with 8 s difference. In this study, we used two different approaches in order to assess the mixed population of peptide fragments; t-test and cross-correlation analysis. Both methods confirm that we have a mixed population of fragments. The visual aspect of the cross-correlation analysis approach allows rapid detection of some potential outliers such as multiple fragments from different peptides but having a similar mass. Another interesting aspect of the cross-correlation analysis approach is related to the fact that it can be used to decipher the various fragments without any prior knowledge of their origin, in contrast to the t-test. Cross-correlation analysis as a tool to segregate fragments based on a slight difference in their elution pattern could potentially be applied to larger mass and time ranges, although this will definitely present a computational challenge.

SUPPORTING INFORMATION
Additional supporting information may be found in the online version of this article.

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