



Identification of phosphorylated human peptides by accurate mass measurement alone

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ABSTRACT

At sufficiently high mass accuracy, it is possible to distinguish phosphorylated from unmodified peptides by mass measurement alone. We examine the feasibility of that idea, tested against a library of all possible *in silico* tryptic digest peptides from the human proteome database. The overlaps between *in silico* tryptic digest phosphopeptides generated from known phosphorylated proteins (1–12 sites) and all possible unmodified human peptides are considered for assumed mass error ranges of ± 10 , ± 50 , ± 100 , ± 1000 , and $\pm 10,000$ ppb. We find that for mass error ± 50 ppb, 95% of all phosphorylated human tryptic peptides can be distinguished from nonmodified peptides by accurate mass alone throughout the entire nominal mass range. We discuss the prospect of on-line LC MS/MS to identify phosphopeptide precursor ions in MS1 for selected dissociation in MS2 to identify the peptide and site(s) of phosphorylation.

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1. Introduction

Phosphorylation is a reversible modification affecting both the folding and function of proteins and regulation of many cellular processes such as cell cycle, growth, apoptosis, and differentiation [1,2]. Among the commonly occurring 20 amino acids, serine, threonine, tyrosine, histidine, arginine, lysine, cysteine, glutamic acid, and aspartic acid residues may be phosphorylated. However, phosphorylation of the hydroxyl group of serine, threonine or tyrosine residues by forming a phosphate ester bond is most frequently observed, with relative occurrence of 90% (pS), 10% (pT), and 0.05% (pY) [3]. Identification of the number and site(s) of phosphorylation is thus a major aspect of proteomics [4].

1.1. MS/MS for identifying the presence and sequence location of phosphorylation in peptides

Mass spectrometry is playing an increasingly important role in the characterization of posttranslational modifications, including protein phosphorylation [5,6]. A simple way to identify the presence of a phosphopeptide by MS is from proteolytic peptide mass spectra before and after phosphatase treatment, revealing an 80 Da mass reduction (HPO_3) for each phosphorylation [7,8], but is limited to analysis of abundant peptides.

Most modern methods for phosphoproteome analysis rely on the use of MS/MS techniques: for example, from precursor ion and neutral loss scans by triple quadrupole mass spectrometry [9–13]. For parent ion scanning, the first quadrupole (Q1) scans through the entire m/z range, precursor ions are subjected to collision-induced dissociation in the second quadrupole (Q2), and the third quadrupole (Q3) monitors the specific fragment ion characteristic of the modification of interest, e.g., the loss of PO_3^- (79 Da) for a negative phosphopeptide ion. In the neutral loss scan, Q1 and Q3 are scanned simultaneously over two different m/z ranges with the (fixed) difference corresponding to the m/z value of the appropriate neutral (e.g., 98 Da for H_3PO_4) that is lost in Q2 [1]. These methods can identify the presence (but not the sequence location) of phosphorylation. Moreover, efficiency of the method decreases as sample complexity increases [14].

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Electron capture dissociation (ECD) and electron transfer dissociation (ETD) techniques mainly generate c/z^+ ions by cleavage of the N–C $_{\alpha}$ backbone bond and are particularly useful for analysis of large multiply charged peptides and for identification and localization of posttranslational modifications. Both methods typically provide more extensive sequence coverage and retain labile posttranslational modifications relative to slow-heating dissociation methods (e.g., collision-induced dissociation (CID), infrared multiphoton dissociation), thereby enabling more complete sequencing of peptides and more precise assignment of phosphorylation sites simultaneously in a single MS/MS experiment [15–17]. For example, in a global phosphoproteome analysis, ETD identified 60% more phosphopeptides than CID, with an average of 40% more fragment ions to facilitate localization of phosphorylation sites. With combined ETD and CID, more than 80% of the known phosphorylation sites in more than 1000 phosphorylated peptides were identified [18].

Another problem is that phosphorylation occurs at low stoichiometry and/or for a protein with low expression level versus nonphosphorylated proteins found *in vivo*. Moreover, for a complex mixture containing many other unmodified peptides, phosphopeptides can exhibit low relative abundance due to more efficient ionization of other species. Although that effect is less pronounced for negative ions, their low fragmentation efficiency (e.g., electron detachment dissociation, negative electron capture dissociation) can preclude phosphorylation identification [1]. Therefore, because current MS methods do not distinguish between phosphorylated and non-phosphorylated precursor ions *before* MS/MS, it is typically necessary to resort to other methods to pre-select phosphorylated peptides: e.g., protease-specific digestion, phospho-enrichment steps, and/or LC separation.

1.2. Phosphopeptide enrichment strategies

Immobilized metal-ion affinity chromatography (IMAC) is the most frequently used technique for enrichment by electrostatic interaction between a negatively charged phosphate and positively charged metal ion (e.g., Fe³⁺, Ga³⁺, or Al³⁺) bound to the stationary phase via linkers [19], and is compatible with subsequent separation and detection in LC-ESI-MS/MS [20]. However, nonspecific binding of peptides containing acidic amino acids such as Glu and Asp limits selectivity. Moreover, minor alteration of pH, ionic strength, and/or organic solvent composition can significantly affect the selectivity of the IMAC stationary phase [1]. Esterification of carboxylic acids to methyl esters by use of HCl-saturated, dried methanol can eliminate the nonspecific binding of acidic peptides in subsequent IMAC enrichment, but at the cost of additional analysis time, sample loss, and increased sample complexity due to incomplete esterification and side reactions, etc. [1,21].

Alternatively, strong cation exchange chromatography can separate singly charged positive tryptic digest phosphopeptides from doubly positively charged nonphosphopeptides at acidic pH (~2.7). However, specificity is low [22]. Other enrichment methods include TiO₂ column chromatography, ZrO₂ microtips, chemical modification of the phosphate by an affinity tag, calcium phosphate precipitation, etc. [23–26]. In all of the above techniques, interference from abundant, nonspecific peptides complicates MS/MS, especially when coupled with on-line LC separation.

1.3. Phosphate identification based on mass defect

Mass defect is the difference between exact mass and the nearest-integer (nominal) mass. The atomic mass defects for ¹²C, ¹H, ¹⁶O, ¹⁴N, ³¹P, and ³²S are shown in Table 1. The limited elemental compositions (C, H, O, N and S) of 20 common amino acids localize the possible mass defects for unmodified peptides to only

Table 1

The accurate atomic masses and corresponding mass defects for five elements found in amino acids [35]. Note the large negative mass defect for ³¹P.

Elements	Accurate mass (Da)	Mass defect (Da)
¹ H	1.007825032	+0.007825032
¹² C	12.000000000	+0.000000000
¹⁴ N	14.003074005	+0.003074005
¹⁶ O	15.994914622	–0.005085378
³¹ P	30.973761487	–0.026238513
³² S	31.972070690	–0.027929310

approximately one-third of each 1-Dalton mass range. Moreover, peptide masses are discrete, so that amino acid composition can be uniquely inferred from sufficiently resolved and accurate mass measurement [27,28]. The introduction of another element with a large mass defect into the elemental compositions can shift a modified peptide ion into an otherwise empty mass spectral segment, for unique identification by accurate mass measurement. Originally introduced to provide non-overlapping (typically fluorinated) mass calibrants [29], mass defect labeling of peptides has been used to identify protein serine (pS) and tyrosine (pY) phosphorylation [27,30–33]. Here, we point out that, by virtue of its large mass defect (–0.0262 Da, see Table 1), the presence of ³¹P in a peptide ion can in principle be identified by sufficiently accurate mass measurement alone.

2. Methods

Tryptic peptides were generated *in silico* from the ProSightPC human protein database [34]. Two types of N-terminal modifications (acetylation and methionine cleavage) were considered, and each digested peptide was allowed one missed tryptic cleavage site. To calculate the overlaps of unmodified and phosphorylated peptides at each nominal mass, a database consisting of the *in silico* trypsin-digested unmodified human proteome and a database containing *in silico* tryptic digest phosphopeptides were generated from human proteins phosphorylated at known positions. Each database includes unique elemental composition and corresponding accurate monoisotopic neutral mass for each peptide. ¹H, ¹²C, ¹⁴N, ¹⁶O, ³¹P, and ³²S atomic mass values (see Table 1 [35]) were used to compute the monoisotopic neutral mass for each elemental composition.

3. Results and discussion

3.1. Phosphorylated vs. non-phosphorylated human peptides; elemental composition vs. amino acid composition

The total entries, non-redundant (i.e., not counting separately isomeric peptides with identical elemental composition but different sequence or amino acid composition—see below) entries, and neutral mass ranges for both phosphorylated and nonphosphorylated human tryptic digest peptide databases are given in Table 2. There are 73,407 phosphorylated and 956,594 nonphosphorylated (non-redundant) peptides with masses ranging from 580–10,088 Da and 500–10,040 Da. As many as 12 phosphorylation sites have been found on a single database peptide, whereas ~34% of observed phosphorylated peptides are modified at two or more sites: ~65.7%, 22.0%, 7.8% and 4.5% containing, one, two, three, and greater than three phosphates (see Fig. 1).

Apart from sequence isomers (e.g., AlaGlyHis vs. GlyHisAla), peptides of a given elemental composition may arise from multiple isomeric amino acid compositions (e.g., Met & Ala vs. Val & Cys) [36]. Although positional isomers have different heat of formation, an energy difference of ($E_2 - E_1$) = 1 eV corresponds to a mass difference of ~10^{–9} Da according to Einstein's mass-energy

Table 2

Total entries, non-redundant entries, and neutral mass range for phosphorylated and nonphosphorylated human tryptic digest peptides.

Phosphopeptide database (N-terminal acetylation and methionine cleavage)			
No. of phosphorylation modification sites	Total entries	Non-redundant entries	Neutral mass range (Da)
1	80357	48265	580–10088
2	40415	16090	662–10127
3	23228	5709	772–10069
4	13712	2074	1040–10077
5	7873	799	1225–9742
6	4222	290	1554–9189
7	2043	111	2071–9269
8	860	46	2151–9248
9	299	17	2231–9328
10	78	3	3805–5799
11	13	2	3885–5879
12	1	1	3965
Total	173101	73407	

Nonphosphopeptide database (N-terminal acetylation and methionine cleavage)		
Total entries	Non-redundant entries	Neutral mass entries range (Da)
2307173	956594	500–10040

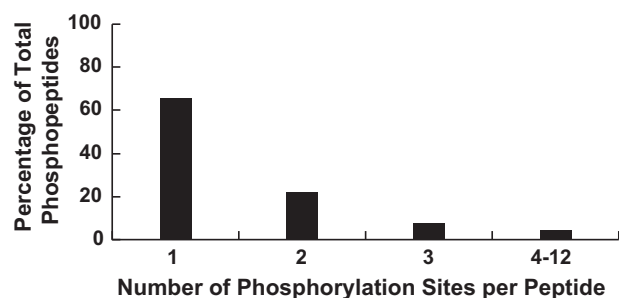


Fig. 1. Distribution of phosphates among all previously identified human phosphopeptides.

relation, $(E_2 - E_1) = (m_2 - m_1)c^2$, a mass difference not accessible by any existing mass analyzer [36,37]. Both the number of amino acid compositions and unique elemental compositions at different nominal masses are shown in Fig. 2 (human non-phosphopeptides) and Fig. 3 (human phosphopeptides). The number of isomers at each nominal mass is readily obtained from difference between amino acid compositions and unique elemental compositions. Note that 2038 of the 2109 amino acid compositions of unmodified peptides of nominal mass 730 Da (i.e., 97%) are isomers, requiring MS/MS for identification.

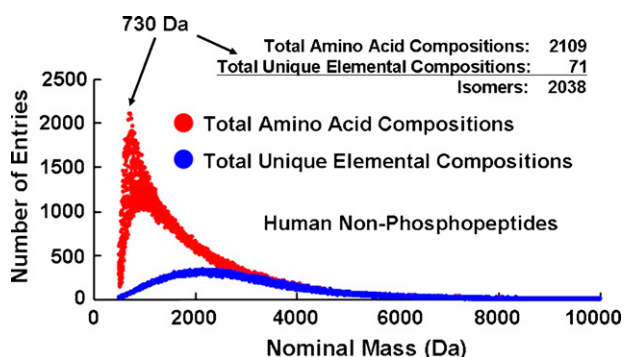


Fig. 2. Amino acid compositions and unique elemental compositions at each nominal mass for the human non-phosphopeptide database.

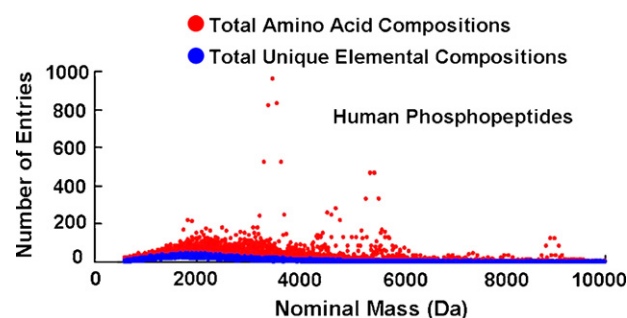


Fig. 3. Amino acid compositions and unique elemental compositions at each nominal mass for the human phosphopeptide database (1–12 phosphorylations per peptide).

3.2. Mass accuracy required to identify phosphopeptides by mass alone

Fig. 4 shows the possible mass distributions of phosphorylated and nonphosphorylated human (neutral) peptides across a 0.05 Da segment of the mass range at a mass accuracy (i.e., mass bin width) to within ± 0.1 ppm. For that small mass segment, it is clear that every phosphopeptide can be distinguished from every nonphosphopeptide. By comparison, Conrads et al. previously showed that at 0.1 ppm mass measurement accuracy, more than 80% of yeast phosphopeptides of 2000 Da nominal mass can be identified directly from their masses [38]. To establish the mass measurement accuracy required to distinguish phosphopeptides from nonphosphopeptides at various nominal masses, Spengler et al. recently assembled a database of theoretically possible peptides up to 2000 Da and calculated the fraction of mass-distinguishable mono- and di-phosphorylated peptides from unmodified peptides, for assumed mass error of 100 and 1000 ppb [39]. Here, we restrict our database to the human proteome (but do not limit the mass range), and then consider the overlap between experimentally confirmed peptides phosphorylated at 1–12 sites and all possible unmodified human peptides for assumed mass error values ranging from 1 to 10,000 ppb.

We now consider the feasibility of distinguishing phosphorylated from unphosphorylated precursor peptide ions in the first stage of an MS/MS experiment, so as to be able to select only phosphopeptides for subsequent fragmentation in the second MS/MS stage. Fig. 5 shows the percentage of overlaps between phospho- and non-phosphopeptides at every nominal mass up to 10,000 Da, for each of five mass measurement error ranges. Although the

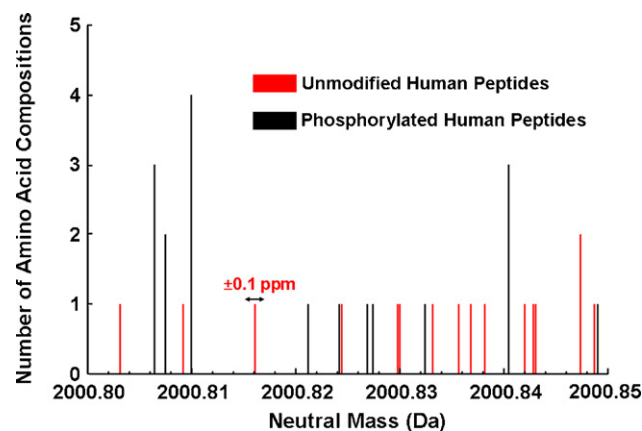


Fig. 4. Histogram of phosphorylated and nonphosphorylated neutral human peptides (counting sequence isomers) across a 0.05 Da mass segment at nominal mass 2000 Da, at a mass accuracy to within ± 0.1 ppm.

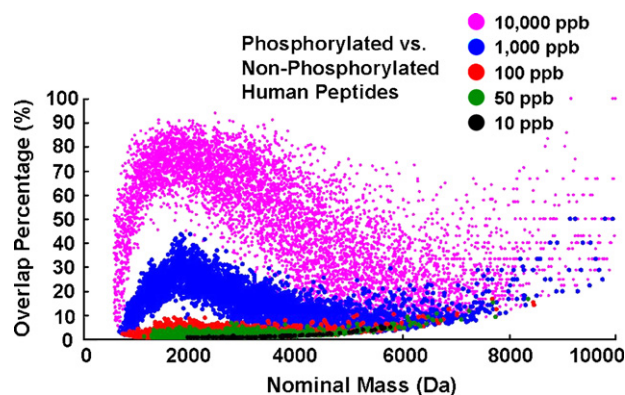


Fig. 5. Human phosphopeptide vs. nonphosphopeptide mass overlap likelihood for each of five mass measurement errors (10–10,000 ppb) at nominal masses up to 10,000 Da.

overlap percentage is less than 50% throughout the entire mass range even for a mass measurement error range of ± 1000 ppb, overlap drops to less than 5% of the nominal masses, for a mass measurement error range of ± 50 ppb, with no overlaps below 1148 Da. Stated another way, Fig. 6 shows the lowest nominal mass at which overlap first appears, starting at 580 Da for mass error of ± 10 ppm and increasing to 1967 Da for ± 0.01 ppm. Note that $\sim 83\%$, 52% , 29% , 6% , and 0.3% of total phosphopeptides (173,101) overlap with at least one nonphosphopeptide at respective mass measurement error of $\pm 10,000$, $\pm 1,000$, ± 100 , ± 50 , and ± 10 ppb. Consideration of possible interferants from other modified peptides such as sulfation, nitrosylation, and ubiquitination (each type is from *in silico* tryptic digest of corresponding known human modified proteins) does not change the overlap results between phosphopeptides and nonphosphopeptides at each mass error, due to the relatively few occurrences for each modified peptide: e.g., 1702 total entries (469 nonredundant) for sulfated peptides at up to 7 sites, 48 total entries (42 nonredundant) for nitrosylated peptides at up to 3 sites, and 606 total entries (577 nonredundant) for ubiquitinated peptides (leaving only a GG tag on each peptide after tryptic digest) at up to 2 sites.

Finally, Fig. 7 shows that even at 1 ppb mass accuracy, there are still two phosphopeptide/non-phosphopeptide pairs of unique elemental compositions at nominal masses 3885 and 4443 Da that cannot be distinguished, corresponding to the mass doublets, $C_{45}H_{62}N_{19}$ vs. $O_{33}P_{11}$ (~ 1.3 μ Da) and $C_{20}H_{17}O_1S_1P_1$ vs. N_{24} (~ 3.5 μ Da). The mass accuracy required to resolve each pair, 0.3 ppb and 0.8 ppb, is not currently attainable.

3.3. Feasibility of identifying phosphopeptides by mass measurement alone

The best prospect for realizing the identification of phosphopeptides is with Fourier transform ion cyclotron resonance mass

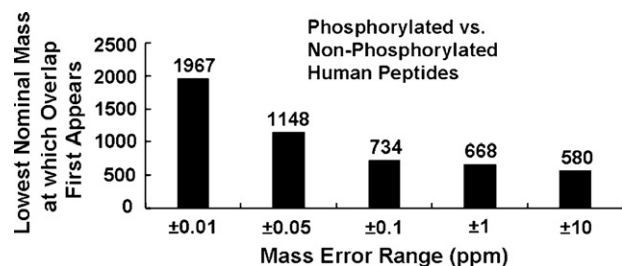


Fig. 6. The lowest nominal mass at which overlap between human phospho- and non-phosphopeptides, for each of five mass measurement error ranges.

Nominal Mass (Da)	Elemental Composition	Accurate Mass (Da)	Minimum Mass Error Range (ppb)
3885	$C_{181}H_{273}N_{49}O_{47}$	3885.0478482	± 0.32
	$C_{136}H_{211}N_{30}O_{80}P_{11}$	3885.0478494	
4443	$C_{176}H_{279}N_{71}O_{66}$	4443.0658043	± 0.80
	$C_{196}H_{296}N_{47}O_{67}S_1P_1$	4443.0658008	

3885 Da ($C_{45}H_{62}N_{19}$ vs. $O_{33}P_{11}$) ~ 1.3 μ Da Mass Difference
YSPTSPTYPSPpTpSPKYpSPpTpSPpTpSPpTpSPK (11 Isomers), vs.
TWVRNAEFSFHEALTHLLHSHLLPEVFTLATLR (No Isomers),

4443 Da ($C_{20}H_{17}O_1S_1P_1$ vs. N_{24}) ~ 3.5 μ Da Mass Difference
PLNRTLpSMSSPLGLEDEFDLENAVLFVAVEVANK (1 Isomer), vs.
AEQQQQQQQQQQQHQQQQQQQQQQQQQHLR (No Isomers),

Fig. 7. Elemental compositions, accurate masses, and amino acid sequences of the (only) two phosphopeptide/non-phosphopeptide pairs not resolved at ± 1 ppb mass measurement error range.

spectrometry, which provides ten-fold higher mass resolution and mass measurement accuracy than other mass analyzers [37]. For example, we have recently achieved mass measurement rms error of <50 ppb for a constantly infused petroleum crude oil sample with thousands of resolved peaks in a single mass spectrum [40]. For peptide mixtures analyzed by LC/MS, mass resolving power and mass accuracy are necessarily lower, due to the need to acquire each precursor mass spectrum in ~ 1 s. However, with recently introduced phase correction to yield absorption-mode display [41], improved ICR cell designs (Tolmachev, Nikolaev), and high magnetic field [42], it should be possible to achieve mass measurement accuracy to within ~ 50 ppb up to 1000 Da sufficient to identify phosphopeptides unambiguously up to ~ 1200 Da. It is worth noting that current instruments typically perform MS/MS only for the (say) five most abundant precursor ions, and therefore fail to access phosphopeptides present at lower abundance. Being able to identify the phosphopeptides *before* MS/MS should thus significantly improve their detection efficiency (and subsequent sequence location by MS/MS).

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