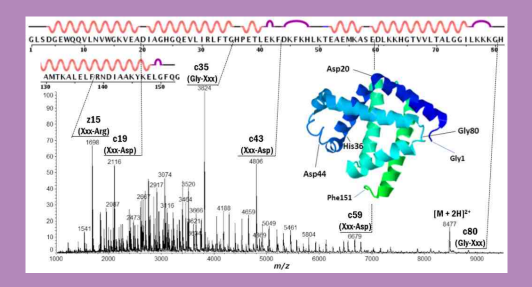


Mass Spectrometry Application in Biology

Edited by

Greg Gorman

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Greg Gorman (Ed.)

Mass Spectrometry Application in Biology



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Preface

The explosive growth in scientific research over the past few decades has been extraordinary and has provided a basis for merging of individual aspects of what was once distinct scientific disciplines into more of a systems based approach. Mass spectrometry, once a specialized tool of analytical chemists has now become a vital instrument for scientist in many disciplines in exploring how complex multi-component systems function and how they are impacted by external elements. The application of mass spectrometry has advanced our knowledge in many areas of science, but perhaps one area in particular where astonishing advances have been observed are the collective group of biological sciences. This has been due in large part to improvements in ionization techniques, sensitivity, mass resolution, and data analysis which have allowed mass spectrometry to be applied to complex biological systems. Application of modern instrumentation capable of high mass resolution are routinely used in in the study of drug metabolism and position specific post-translation modifications from both in vitro and in vivo explorations that are critical to the discovery and development of new drug therapies. Mass spectrometry has also been a critical analytical tool in the omics explosion. Various segment configurations of mass analyzer stages resulting in hybrid instruments including time-of-flight (TOF), quadrupoles (Q), and linear ion traps (LIT) coupled with an array of ionization techniques such as matrix assisted laser desorption/ionization (MALDI), electrospray ionization (ESI) etc. have given scientists the ability to generate critical data in this field. Modern instrument configurations such as Q-TOF, LIT-TOF and TOF-TOF allow for identification and quantitative analysis of proteins, peptides generated from protein digests, and lipids producing expansive data in proteomics and lipidomics. Likewise this same instrumentation can generate data on the complex aspects of metabolism in the field of metabolomics. A bit more recently, mass spectrometry has been used to develop a molecular imagining technology which provides a means to conduct in situ analysis of thin tissue sections. In this application mass spectrometry is used to produce images of the spatial distribution of molecules of interest within a tissue sample. This application known as "imaging MS" generates a view of biological processes from the distributions of molecules within the sample while maintaining intact histological features. The technique is applicable to various molecules including proteins, peptides, lipids, drugs, and metabolites.

The aim of this book is to provide a glimpse of some of the many applications of the advancing field of mass spectrometry and its application to complex biological problems. The topics presented here touch on many of the areas listed above will hopefully serve as a catalyst to generate new ideas for the continued innovative application of mass spectrometry to the fields of medicine, biology and beyond.

Greg Gorman Samford University Guest Editor

Phosphorylation Stoichiometries of Human Eukaryotic Initiation Factors

Armann Andaya, Nancy Villa, Weitao Jia, Christopher S. Fraser and Julie A. Leary

Abstract: Eukaryotic translation initiation factors are the principal molecular effectors regulating the process converting nucleic acid to functional protein. Commonly referred to as eIFs (eukaryotic initiation factors), this suite of proteins is comprised of at least 25 individual subunits that function in a coordinated, regulated, manner during mRNA translation. Multiple facets of eIF regulation have yet to be elucidated; however, many of the necessary protein factors are phosphorylated. Herein, we have isolated, identified and quantified phosphosites from eIF2, eIF3, and eIF4G generated from log phase grown HeLa cell lysates. Our investigation is the first study to globally quantify eIF phosphosites and illustrates differences in abundance of phosphorylation between the residues of each factor. Thus, identification of those phosphosites that exhibit either high or low levels of phosphorylation under log phase growing conditions may aid researchers to concentrate their investigative efforts to specific phosphosites that potentially harbor important regulatory mechanisms germane to mRNA translation.

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1. Introduction

The process of converting nucleic acid to functional protein, a process involving the precise spatial and temporal arrangement of proteins, protein complexes, and nucleic acids, is known as translation [1–3]. Initiation, elongation, termination, and recycling constitute the four chronological phases of mRNA translation with initiation bearing the greatest regulation. The principal molecules that regulate initiation for eukaryotes are the eukaryotic initiation factors (eIFs), a family which is comprised of 12 members encompassing at least 25 individual proteins [1]. One of the most common means of regulating proteins involves phosphorylation, a well-established post-translational modification. Herein, we report our global investigation of phosphorylation quantification for three essential eukaryotic initiation factors, eIF2, eIF3, and eIF4G.

At the onset of eukaryotic translation initiation, the 40S ribosome binds several factors including eIF2 and eIF3 to form the 43S preinitiation complex (PIC). eIF2 (a heterotrimer consisting of the proteins eIF2α, eIF2β, and eIF2γ) binds as a ternary complex (TC) comprised of eIF2, the initiator methionyl tRNA (Met-tRNA_i), and GTP (guanosine triphosphate). The PIC can then bind to nascent messenger RNA (mRNA) through the direct interaction of eIF3 and eIF4G, a member of the eIF4F cap-binding complex. The largest of the eIFs, eIF3 is comprised of thirteen distinct subunits with a holoprotein mass of approximately 800 kDa. Numerous studies implicate the holoprotein eIF3 and its many subunits as an essential factor during translation initiation: It is the central assembly on which other factors, proteins, and nucleic acids bind [4–10]. The factor eIF4F is also an essential

protein to translation initiation. It is comprised of three other eIFs: eIF4A, eIF4E, and the largest of the three, eIF4G with a mass of 176 kDa. eIF4G functions as a scaffolding protein onto which other factors, such as eIF3, eIF4A, and eIF4E functionally interact [11–13].

Although each subunit has a defined function during translation initiation, the regulatory mechanisms governing initiation have yet to be entirely solved. Protein phosphorylation, albeit a well-established post-translational modification, is still not completely understood, particularly in eIFs, as regards target specificity. However, the resulting effects once a protein is phosphorylated can be dramatic as that of ser-51 phosphorylation on eIF2 α , which has been extensively studied and serves as a prime example of the effect phosphorylation has on the initiation process.

During initiation, eIF2 TC hydrolyzes GTP then releases Met-tRNA_i following start codon recognition. eIF2B, a GTP exchange factor (GEF) for eIF2, exchanges GDP (guanosine diphosphate) for GTP, which then promotes binding of a new Met-tRNA_i for further rounds of translation. Phosphorylation of eIF2 α at ser-51 reduces the dissociation rate of eIF2 from eIF2B, effectively sequestering eIF2B and preventing eIF2 TC regeneration and thus globally repressing the rate of mRNA translation [14–18]. The largest protein of the three subunits, eIF2 γ , functions as a scaffolding protein on which the remaining subunits bind. We recently identified eight novel sites of phosphorylation on eIF2 γ and demonstrated the potential *in vitro* effects of eIF2 γ phosphorylation via protein kinase C (PKC) [19].

In addition to the identification of novel phosphosites, determining those levels of phosphorylation on specific residues allows researchers to define potentially important phosphosites, thereby distinguishing those sites from potentially less biologically meaningful ones [20–26]. As with eIF2α, phosphorylation at ser-51 becomes more pronounced under conditions of cellular stress which demonstrates how the fluidity of the phosphorylation stoichiometry reveals intrinsic mechanisms implicated in regulating phosphorylation in response to cellular cues [14–18]. Establishment of phosphosite stoichiometry under specific biological conditions may help to focus ensuing investigations for deciphering biologically important phosphosites.

While identification of novel phosphosites is the essential first step in the eventual evaluation of their biological impact, follow-up studies, even on a few novel sites of phosphorylation, often presents a task too burdensome for subsequent in-depth investigations. Researchers identify phosphosites for further study through evaluation of their structural location. Phosphosites residing in structurally salient locations such as established binding sites, binding pockets, or areas of significant secondary, tertiary, and/or quaternary structure, are considered desirable candidates for further evaluation. However, without such informative data, further investigations into newly discovered phosphosites are often avoided.

In order to assess the amount of phosphorylation occurring at specific residues, we recently published a mass spectrometry method proficient at measuring phosphosite stoichiometry [27]. Our method relies on the measurement of dephosphorylation of phosphopeptides accomplished chemically via cerium oxide [28]. We have further optimized cerium oxide's capacity for dephosphorylating peptides and have subsequently developed a method to measure the amount of dephosphorylation via tandem mass tags (TMT) [25]. Use of the isobaric TMT reporter ions allows for assessment of phosphorylation stoichiometry. We have verified the use of this quantification

protocol and have shown its efficacy in measuring the stoichiometry of previously established phosphosites. One of the subunits of eIF3, eIF3h, has one previously identified phosphorylation site, ser-183 [29], and we have now determined its level of phosphorylation to be 70% in log phase grown HeLa cells (data provided and discussed in the results section). This observation coincides with the critical *in vivo* effects of eIF3h's ser-183 phosphorylation during malignant transformation of NIH 3T3 cells [27,30].

This report is the first to highlight the phosphorylation levels of three heavily phosphorylated eukaryotic initiation factors. Additionally, our quantification analysis is specific to the analysis of cells grown at optimal conditions and thus underscores the importance of variability possible with these phosphosites under different environmental circumstances. Knowledge gained from this study provides a platform for future investigations not only for phosphorylated proteins, but also for the inherent variability of specific phosphosites within the protein itself. Hence, the purpose of this study is to add to the growing pool of knowledge of these factors and more importantly, to initiate an investigation primarily aimed at quantified phosphorylated residues and their implication on translation initiation. Future studies into the regulation of these factors may be based on the findings within this study.

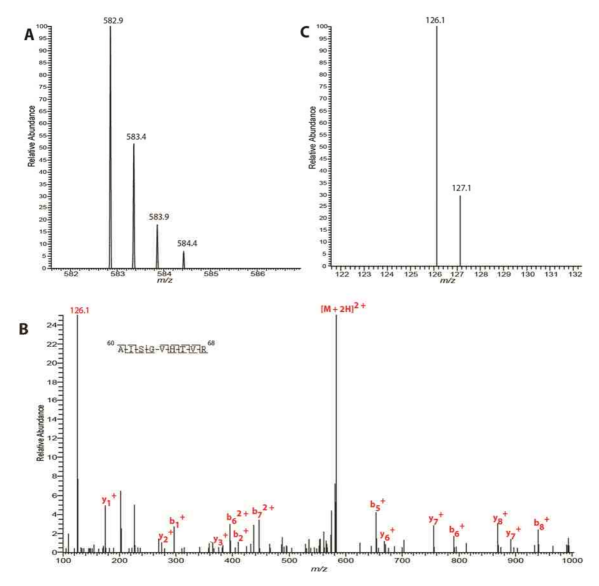
2. Results and Discussion

2.1. Quantification of Phosphorylation for Eukaryotic Initiation Factor 2 (eIF2)

We isolated eIF2 from HeLa cell lysate in order to quantify its level of phosphorylation. The factor was purified from HeLa cells under optimal growth conditions (log phase growth). As eIF2 is a heterotrimer with a molecular mass of 126 kDa, we analyzed the two factors that have been previously reported as phosphorylated within HeLa cells, eIF2 β and eIF2 γ . While eIF2 α has been heavily studied at its principally phosphorylated residue of ser-51, quantification of that residue does not lead to new information or developments, which is the principal aim of this study. Thus, we focused our efforts at quantifying the numerous sites on the remaining two subunits of eIF2 in order to gain further insight into their function and regulation.

Recently, we published the identification of eight sites of phosphorylation, seven of which were novel and reside on the largest subunit, eIF2 γ , the core subunit of the heterotrimer [19]. All identified phosphosites were quantified. On eIF2 γ , numerous functional domains exist dedicated to specific tasks during eukaryotic translation. Threonine 66, a phosphorylated residue, is located within the switch-1 region of the protein, and in this study, we observed its level of phosphorylation at 71% (Figure 1). Two sites adjacent to one another, ser-55 and thr-56, reside directly in the nucleotide binding pocket of eIF2 γ and our quantitative phosphoanalysis revealed the levels of phosphorylation to be 85% for both ser-55 and thr-56 combined (Supplementary Information). Proximal to the *C*-terminal end of the protein, ser-412 and thr-413 exhibit phosphorylation at 7% while ser-418 and thr-435 have phosphorylation at 70% and 60%, respectively (Supplementary Information). Lastly, thr-109's phosphorylation level was observed at 30% and this residue sits adjacent to the zinc binding domain (Supplementary Information).

Figure 1. Quantification of phosphorylation on Thr-66 on eIF2 γ . (**A**) Precursor ion mass scan of the [M + 2H]²⁺ ion is shown; (**B**) MS/MS spectra of m/z ion 582.9 illustrating indicative b- and y-ions for peptide spanning residues 60–68 of eIF2 γ ; and (**C**) Zoomed in view of the m/z region containing the TMT (tandem mass tags) reporter ions. Calculation of the reporter ion ratio reveals a phosphorylation level of 70.5% for Thr-66.



In contrast to eIF2 γ , eIF2 β has a lower abundance of phosphorylated residues within HeLa cells. Nevertheless, we quantified these sites and observed levels of 18% for thr-31 but near or less than 5% for the other reported eIF2 β phosphosites (Table 1).

Table 1. Three of the factors, eIF2, eIF3, and eIF4G were analyzed as to their quantification of phosphorylation levels within log phase grown HeLa cells. Percentages of the phosphosites are shown.

Protein	Subunit	Residue Phosphorylated	% Phosphorylation
		Thr-31	18
		Ser-67	<5
	β	Ser-105	<5
		Thr-111	<5
		Ser-158	8
		Ser-55	85
eIF2		Thr-56	85
		Thr-66	71
	24	Thr-109	30
	γ	Ser-412	7
		Thr-413	7
		Ser-418	70
-		Thr-435	66
		Ser-881	84
	0	Ser-1198	92
	a	Ser-1336	18
		Ser-1364	36
		Ser-83	85
	b	Ser-85	85
eIF3	U	Ser-119	70
		Ser-125	70
	c	Thr-524	95
	~	Thr-41	31
	g	Ser-42	31
	h	Ser-183	89
	j	Thr-109	88
	-	Thr-647	65
		Ser-1028	5
		Ser-1077	15
		Ser-1092	40
		Ser-1144	45
		Ser-1147	45
eIF4G		Ser-1185	70
CII'4U	-	Ser-1187	70
		Ser-1209	50
		Thr-1211	50
		Ser-1231	75
		Thr-1425	62
		Ser-1430	62
		Ser-1596	<5

2.2. Quantification of Phosphorylation for Eukaryotic Initiation Factor 3 (eIF3)

Although the exact structure and placement of each subunit within the tridodecamer of eIF3 has yet to be fully solved, the number of phosphorylations derived from log phase HeLa cells is 29 [29]. We thus sought to quantify the levels of each of these 29 sites of phosphorylation. Given that each subunit is a distinct protein, the varying levels of phosphorylation from subunit to subunit or within each subunit should not be surprising.

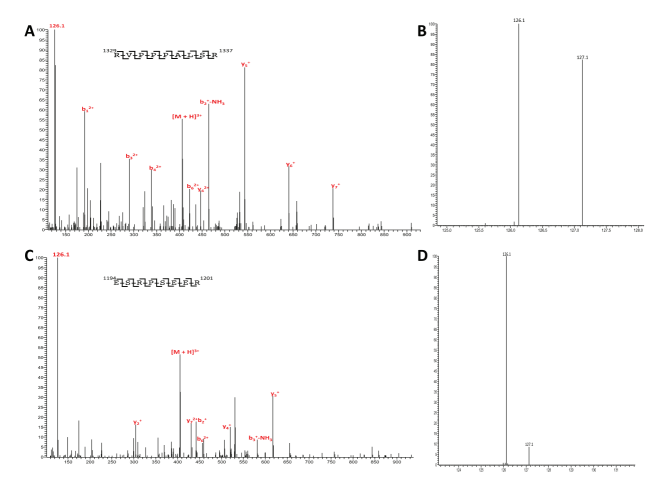
The core subunits of eIF3 are widely believed to be eIF3a, eIF3b, and eIF3c, the three largest of the thirteen subunits. These three subunits house 21 of the 29 sites of phosphorylation for eIF3, 15 of which were accurately quantified. The largest subunit, eIF3a, has a varied landscape of phosphorylation ranging from 18% for ser-1336 to a high of 92% for ser-1198 (Figure 2). Intriguing are the two phosphorylated residues at the *C*-terminal end (or *C*-Terminus) of the protein, ser-1336 and ser-1364. While ser-1336 has a phosphorylation of 18%, ser-1364 has a level of 36% (Supplementary Information). The exact crystal structure of human eIF3a has yet to be solved, but from this study, we see that the *C*-terminal phosphorylated residues are lower in phosphorylation levels than that for ser-1198 at 92% and for that of ser-881 at 84%. This rather high level of phosphorylation for these residues may implicate functional significance for eIF3 quaternary structure given that the *C*-terminal phosphorylated residues possess a lower level of phosphorylation.

The second largest eIF3 subunit, eIF3b, has 7 sites of phosphorylation. Unfortunately, we did not attain full sequence coverage for all proteins analyzed which lead to a lack of quantification for some previously identified phosphosites. This was due in part to either the absence of a lysine or arginine within the vicinity of the phosphosite in question or an inadequacy of sufficient identifiable b- and y-ions. However, as with eIF3a's ser-1198 and ser-881, the levels of phosphorylation for eIF3b were relatively large ranging from 70% for both ser-119 and ser-125, and 85% for both ser-83 and ser-85 (Supplementary Information). Again, implications in eIF3 quaternary structure for these high levels of phosphorylation will be investigated further.

The third largest purported eIF3 core subunit is eIF3c. Again, due to the same limitations as previously mentioned, the *N*-terminal phosphorylated residues of ser-9, ser-11, ser-13, ser-15, ser-16, ser-18, and ser-39 could not be quantified. But as with subunits eIF3a and eIF3b, higher levels of phosphorylation were observed for thr-524 at 95% (Supplementary Information).

Regarding the subunit eIF3h, we have already published a report detailing its relatively high level of phosphorylation from log phase HeLa cells at 70% and within the current study, 89% [27]. This high level of phosphorylation is in direct correlation with biological data showing that mutation of this residue to alanine produced a decrease in growth rate of NIH-3T3 cells [30]. Residues not quantified for eIF3 due to previously mentioned experimental limitations were eIF3f (ser-258) and the *N*-terminal residues of eIF3j (ser-11, ser-13, and ser-20). A level of 31% phosphorylation was observed for eIF3g's two phosphorylated residues, thr-41 and ser-42, and a high level of phosphorylation comparable to that of eIF3h was observed for thr-109 on eIF3j (Table 1 and Supplementary Information).

Figure 2. Quantification of phosphorylation on eIF3. (**A**) MS/MS spectra of m/z ion 406.6, z = 3. Peptide spans residues 1329–1337 for eIF3a; (**B**) Zoomed in view of the m/z region containing the TMT reporter ions. Calculation of the reporter ion ratio reveals a phosphorylation level of 18% for Ser-1336; (**C**) MS/MS spectra of m/z ion 405.5, z = 3. Peptide spans residues 1194–1201 for eIF3a; and (**D**) Zoomed in view of the m/z region containing the TMT reporter ions. Calculation of the reporter ion ratio reveals a phosphorylation level of 92% for Ser-1198.



2.3. Quantification of Phosphorylation for Eukaryotic Initiation Factor 4G (eIF4G)

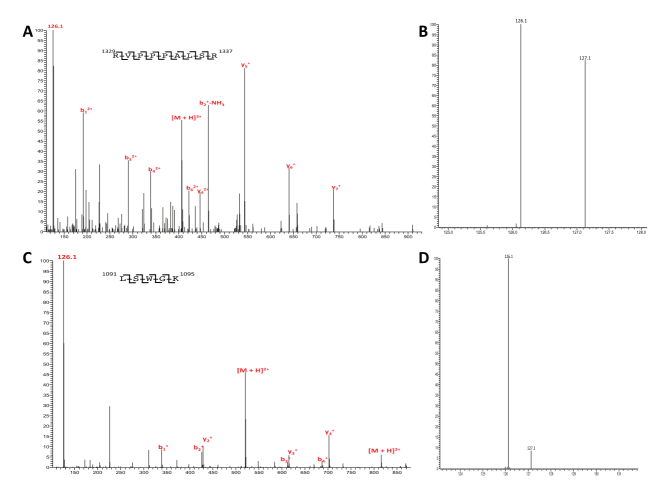
A large scaffolding protein, eIF4G commonly occurs with varying isoforms that are dependent on the location of the start codon. Nonetheless, the principal isoform of the protein is comprised of 1599 amino acids with a molecular mass of 176 kDa. Prior to translation initiation, eIF4G serves as a docking site to which other proteins bind prior to forming larger complexes necessary for the translation process. Known proteins that bind to eIF4G are poly A binding protein (PABP), the kinase Mnk1, and initiation factors eIF4E, eIF4A, and eIF3. The phosphorylation of eIF4G from HeLa cells has been well documented with evidence of up to 20 sites of phosphorylation corresponding to eIF4G derived from HeLa cells have been identified. However, as with all other eIFs, the quantification of this phosphorylation has yet to be reported.

Currently, considerable investigation has focused on the portion of eIF4G that has been mapped to bind to eIF3 [31]. This portion, corresponding to residues 1015 to 1105, contains three sites of

phosphorylation identified from HeLa cells, ser-1028, ser-1077, and ser-1092. Quantification of each site of phosphorylation reveals 5%, 15%, and 40% for ser-1028, ser-1077, and ser-1092, respectively (Figure 3). The only other site of phosphorylation that maps to a known binding region on eIF4G is located at the *C*-terminus of the protein and is ser-1596 that corresponds to the Mnk1 binding region. Quantification of its phosphorylation reveals less than 5% levels of phosphorylation.

As for the remaining known sites of phosphorylation for eIF4G from HeLa cells, insufficiently scored peptides prevented our quantitative analysis of phosphosites residing in the *N*-terminal portion of the protein. However, high levels of phosphorylation were observed for thr-647 at 65%. Regions near but distinct from the eIF3 binding site had phosphorylation levels at or near 50%. The peptide mapped to ser-1144 and ser-1147 had a combined level of 45% and the peptide mapped to ser-1209 and ser-1211 was at 50%. The peptide mapping to ser-1185 and ser-1187 had a level of 70% and the peptide mapping to ser-1231 had a level of 75%. Peptides mapping in close proximity to the Mnk1 binding region, thr-1425 and ser-1430 were quantified at phosphorylation levels of 62%.

Figure 3. Quantification of phosphorylation on eIF4G. (**A**) MS/MS spectra of m/z ion 992.6, z = 2. Peptide spans residues 1072–1085; (**B**) Zoomed in view of the m/z region containing the TMT reporter ions. Calculation of the reporter ion ratio reveals a phosphorylation level of 15% for Ser-1077; (**C**) MS/MS spectra of m/z ion 520.8, z = 2. Peptide spans residues 1091–1095; and (**D**) Zoomed in view of the m/z region containing the TMT reporter ions. Calculation of the reporter ion ratio reveals a phosphorylation level of 40% for Ser-1092.



The entire process of translation initiation requires a concerted and choreographed effort from numerous proteins to effectively translate the nascent mRNA into functional protein. Although the process of translation may be separated into the four distinct phases of initiation, elongation, termination, and recycling, much of the regulation is centralized to the initiation phase. Translation initiation is the process by which the 40S ribosome, bound to Met-tRNA_i and several initiation factors, binds to the 5' end of mRNA then scans in the 5' to 3' direction until the initiation codon is recognized. Recruitment of Met-tRNA_i requires formation of a ternary complex also containing eIF2 and GTP, while mRNA recruitment to the ribosome complex requires binding of eIF4G and eIF3.

As the formation of the ternary complex is critical to translation, a better understanding of the major protein component, eIF2, would help define this process. A closer look into the function of eIF2 reveals that the heterotrimer consists of three subunits, eIF2 α and eIF2 β bound to eIF2 γ but not each other. Classically, eIF2 α 's phosphorylation on ser-51 converts the eIF2 heterotrimer into an inhibitor of eIF2B, the heteropentamer responsible for exchanging GDP for GTP on eIF2. Since the phosphorylation of this one residue on eIF2 α has a profound impact on the overall process of translation, we hypothesized that the other subunits of eIF2 may also contribute in some part to the overall process of translation.

Our phosphoanalysis of eIF2 centered on known phosphosites for both eIF2 β and eIF2 γ derived from HeLa cells. In our previous report for eIF2 γ , one of the novel phosphosites identified was thr-66 [19]. Further investigations of this site were conducted based on previous structural studies placing it within eIF2 γ 's switch 1 region [32,33]. As eIF2 forms a ternary complex with met-tRNAi and GTP, the switch 1 region of the molecule undergoes a conformational change in order to allow for complexation [34,35]. Our previous report revealed not only phosphorylation of thr-66, but also established PKC as a kinase with the ability to phosphorylate thr-66 *in vitro*. In the current study, we observed that for cells propagated under log-phase growth conditions, thr-66 on eIF2 γ is 71%. Given that a majority of thr-66 is phosphorylated for a cancer cell line under optimal conditions, future discernment of the physiological role of this phosphosite should be of paramount importance. Logically, phosphorylation within this region of the molecule may have a significant impact on the protein's overall structure and consequently its influence on the conformational change experienced upon GTP binding to eIF2 γ .

Two other structurally noteworthy phosphorylation sites are ser-55 and thr-56 [36,37]. Both of these sites reside directly within the nucleotide binding pocket of eIF2 γ . As eIF2 γ represents the core protein of the eIF2 heterotrimer, it binds directly to eIF2 α and eIF2 β , yet the α and β subunits do not bind to each other [38,39]. Two-thirds of the ternary complex forms upon GTP binding to the heterotrimer, specifically, to eIF2 γ . In its inactive form, GDP is bound along with a magnesium molecule in eIF2 γ 's binding pocket. Our quantitative phosphoanalysis revealed that the level of phosphorylation is approximately 85% for both ser-55 and thr-56 combined. As with most phosphorylation quantification protocols, phosphosites residing on the same tryptic peptide are difficult to quantify individually [40]. Nonetheless, a level of 85% phosphorylation within the nucleotide binding pocket raises some intriguing possibilities. Phosphorylation in this binding pocket may potentially present itself as a regulatory mechanism eliciting steric strain, which could prevent binding of an additional nucleotide. Such a high level of phosphorylation within this binding pocket

most likely influences the interplay of magnesium and GTP binding. The temporal aspects of kinase and/or phosphatase interplay with GTP and magnesium binding necessitate future investigations. The possibility of enzyme-aided phosphorylation diminishes with such a small binding pocket thus raising questions of this event possibly occurring prior to protein folding. Clearly, future studies are needed to determine the effects of such high levels of phosphorylation within an important region of the protein.

In contrast to those phosphosites near the binding pocket and within the switch 1 region, phosphosites associated with the C-terminal portion of the protein, ser-412 and thr-413 have levels at 7% while ser-418 and thr-435 are at or near 70%. Thus, from the current study, we can see the dynamics of the phosphorylation landscape unfold. Residues 412 and 413 reside on the outer face of the molecule and despite the fact that they may be more readily available to kinases as compared to residues 55, 56, and 66, (which lie on the interior of the molecule), their levels of phosphorylation are significantly smaller. Future investigations will determine the significance of these phosphorylation stoichiometries, but for now, we can speculate that phosphorylation does not play a major role in the binding to other nucleotides and/or proteins. The same appears not to be true for either ser-418 or thr-435. Again, future studies will determine these residues' phosphorylation significance, but undoubtedly, different physiological circumstances and pressures exist on the phosphorylated residues of the C-terminal of eIF2 γ . Such a variation of phosphorylation may imply a structural necessity adopted by the C-terminus during the binding processes. It would be interesting to observe phosphorylation differences not only under different growth conditions, but also when one of the four C-terminal sites is mutated and how this effect translates to the overall phosphorylation landscape across the C-terminal region.

Lastly, thr-109's phosphorylation level at 30% seems to neither weaken nor strengthen the importance of this phosphosite to the overall physiology of eIF2. However, as stated in our previous report, this residue does reside near a zinc binding domain. The electrostatics involved with the zinc cation and the negative charge inherent in a phosphate moiety could be responsible for an interaction that may determine the proper positioning of the zinc ion. Proteins containing zinc binding motifs have been well documented in their binding affinity for nucleotides. As eIF2 binds to met-tRNA_i and is part of the ternary complex that shuttles this polynucleotide to the awaiting mRNA during protein translation initiation, the interplay of the phosphate on thr-109 and the zinc ion may have a meaningful impact on protein translation.

Considerable more research is required to decipher the significance of eIF2 β phosphosite structural regions. Previous targeted investigations into eIF2 β have established an interaction of the protein with the kinase CK2 and suggest numerous physiological pressures affecting the dynamics of eIF2 β phosphorylation warranting further investigation [41]. Nonetheless, our report represents the first measurements of the levels of quantification of these phosphosites. Aside from ser-51 on eIF2 α , the phosphosites of eIF2 β and eIF2 γ are not only dynamic, but may house important functions necessary for translation initiation [42].

Along with eIF2, eIF3 is a necessary initiation factor as it binds to many of the numerous eIFs including eIF4A, eIF1, eIF1A, eIF2, and eIF4G. A critical function of eIF3 is prevention of premature association to the 60S ribosome by binding to the 40S ribosome along with other initiation

factors. Although its function and structure have been previously investigated, to date eIF3 has neither a definitive crystal structure nor have any of its phosphorylation sites been thoroughly investigated. Recent studies have shown the structure of the tridodecamer as a 5 lobed entity comprised of a core with five protrusions [43]. The exact structure of each of the lobes has yet to be fully determined, but mass spectrometry along with reconstitution of a recombinant tridodecamer has indicated the core subunits, eIF3a, eIF3b, and eIF3c, as being the three largest [10,44,45]. Interestingly, these same three subunits contain 21 of the 29 possible phosphorylation sites for eIF3. All three of the purported core subunits have high levels of phosphorylation. As the quaternary structure of eIF3 may rely heavily on the integrity of its core proteins, a high level of phosphorylation, especially from log phase grown cells, suggests an increasing role of electrostatic interactions that possibly strengthen the integrity of the entire complex. Levels of phosphorylation were not as elevated for the other phosphorylated subunits except for that of eIF3j. However, eIF3j has been known to disassociate from the other 12 subunits of eIF3 thus a high level of phosphorylation for thr-109 may be indicative of an increased affinity to the holoprotein during translation [7,8]. Future investigations will help determine the nature of each of these phosphosites.

The final protein whose phosphorylation was quantified in this study was eIF4G. A classic example of a scaffolding protein, much like the core proteins of eIF3, phosphorylation was present at its highest level for residues residues residues in known areas of protein binding. Three phosphorylated residues lie within the eIF3 binding region of eIF4G and all three exhibited high levels of phosphorylation. Other residues that also showed high levels of phosphorylation are residues between the eIF3 and eIF4A binding regions. Although future investigations will decipher the role of these specific phosphosites, phosphosite ser-1187 is known in HEK293 cells to function as a substrate for the kinase PKCα and modulates Mnk1 binding [30]. Again, phosphorylated residues implicated in binding appear to have the highest levels of phosphorylation.

3. Experimental Section

3.1. Purification of eIFs from HeLa Cell Lysate

All proteins for this study, eIF2, eIF3, and eIF4G were enriched from HeLa cell lysate prior to nano-LC-MS/MS according to previously published protocols [19,29]. Briefly, HeLa cell lysate (from approximately 30 L cells) was quickly thawed at 37 °C supplemented in a mixture including 10% glycerin, 1 mM EDTA, 1 mM EGTA, 50 mM NaF, 50 mM beta-glycerol phosphate, 10 mM benzamidine, 1 mM DTT, and 1× protease inhibitor mixture (Roche, Basel, Switzerland). After stirring for 10 min at 4 °C, the mixture was centrifuged at $20,000 \times g$ for 20 min at 4 °C. To the resulting supernatant, KCl was added to a final concentration of 450 mM followed by centrifugation in a Beckman Ti-45 rotor for 4 h at 4 °C at 45,000 rpm. The middle two-thirds of the supernatant was carefully removed and stirred at 4 °C while saturated ammonium sulfate was added to a final concentration of 40%. After stirring on ice for 1 h, the suspension was centrifuged at $20,000 \times g$ for 10 min at 4 °C and the pellet (referred to as the A cut) was frozen for future use. To the remaining supernatant, ammonium sulfate was added to a final concentration of 70%. After stirring for 1 h, the mixture was centrifuged at $20,000 \times g$ for 10 min at 4 °C. The pellet (the B cut) was resuspended in

50 mL of buffer A (20 mM HEPES, pH 7.5, 10% glycerol, 1 mM EDTA, 1 mM EGTA, 50 mM NaF, 50 mM β-glycerol phosphate, 10 mM benzamidine, 1 mM DTT) containing 50 mM KCl and dialyzed in two liters of the same buffer for 2.5 h at 4 °C. Following dialysis, the lysate was passed through a 0.2 µm syringe filter and loaded onto a MonoQ (10/10) column (GE Healthcare, Sunnyvale, CA, USA). The column was eluted with a linear gradient of 100 to 500 mM KCl in buffer A at 2 mL/min with 3 mL fractions collected. The fractions that contained the protein of interest (either eIF2, eIF3, or eIF4G) were then pooled, dialyzed against buffer A containing 100 mM KCl for 2.5 h at 4 °C, and loaded onto a MonoS (10/10) column (GE Healthcare, Sunnyvale, CA, USA). The same gradient as that of the MonoQ column was applied and fractions from the MonoS column were analyzed using SDS-PAGE in a similar fashion. Fractions containing the protein of interest were pooled and loaded onto a hydroxyapatite column made in-house using commercial hydroxyapatite (Calbiochem, San Diego, CA, USA). Elution was performed using a linear gradient from 0% to 100% 0.5 M potassium phosphate buffer at pH 7.5. Fractions were again analyzed via SDS-PAGE in a similar manner to those eluting from either the MonoQ or MonoS columns. Fractions containing the now purified protein were pooled and concentrated using Amicon Ultra filtration devices with a MWCO of 10,000 Da to yield a final concentration between 1 and 2 mg/mL.

3.2. Tryptic Digestion of eIFs with Subsequent Tandem Mass Tag (TMT) Labeling

Each eIF protein was digested with trypsin following the same protocol. Approximately 1 μ mol of either eIF2, eIF3, or eIF4G was first reduced at 56 °C for 45 min in 5.5 mM DTT final concentration followed by alkylation for one hour in the dark with iodoacetamide (IAA) added to a final concentration of 10 mM. Trypsin was added at a final enzyme:substrate mass ratio of 1:50 and digestion carried out overnight at 37 °C. The reaction was quenched by flash freezing in liquid nitrogen and the digest was lyophilized. Prior to TMT labeling, each lyophilized sample was dissolved into 100 μ L of 2% acetonitrile supplemented with 0.1% trifluoroacetic acid.

All eIF peptides were prepared for TMT labeling following our published protocol [27]. Briefly, each eIF peptide mixture was divided equally; one population was treated with cerium oxide nanopowder (Sigma-Aldrich, St. Louis, MO, USA) to dephosphorylate the phosphopeptides, the other half was left untreated. Subsequent to dephosphorylation, the dephosphorylated samples were labeled with TMT-126. The remaining untreated half was labeled with TMT-127. Samples were then combined after labeling prior to nano-LC-MS/MS analysis.

3.3. Nano-LC-MS/MS of eIF Proteins

All samples underwent complete labeling and were combined as outlined in our protocol [19,27] prior to nano-LC-MS/MS analysis using an LTQ-Orbitrap XL (Thermo Fisher, San Jose, CA, USA) mass spectrometer equipped with an ADVANCE ion max source (Michrom Bioresources Inc., Auburn, CA, USA), a Surveyor MS pump (Thermo Fisher, San Jose, CA, USA), and a microautosampler (Thermo Fisher, San Jose, CA, USA). Samples were loaded onto the column for 30 min at 2% solvent B (0.1% (v/v) formic acid in acetonitrile) with 98% solvent A at a flow rate of 750 nL/min. Peptides were eluted off the column at 750 nL/min using the following gradient: 2%–10% solvent B for 5 min, 10%–35% solvent B for 65 min, 35%–70% solvent B for 5 min, 35%–70% solvent B for

5 min, 70%–90% solvent B for 5 min, 90% solvent B for 5 min, then reversed to 2% solvent B for 10 min. All ions with TMT labels were analyzed via higher energy collision dissociation (HCD), all others with collision-induced dissociation (CID). Settings for the LTQ-Orbitrap XL were set as follows: data-dependent scan of the top 5 most abundant ions, minimum signal threshold of 55,000, collision energy set to 35%, resolution set to 30,000, dynamic exclusion set to 60 s, repeat count set to 2, and repeat duration set to 30 s. All RAW data were deposited directly into SEQUEST Bioworks 3.3.1 (Thermo Fisher, San Jose, CA, USA) and manually validated. Searches were conducted against an in-house developed database consisting of 584 proteins containing the sequences of all known mammalian eukaryotic initiation factors as well as their reversed sequences. Common contaminants including human keratins, porcine trypsin, bovine serum albumin, bovine beta-casein, as well as their reversed sequences were also included. We performed searches with tryptic specificity and allowed for three missed cleavages at a tolerance of 20 ppm in MS mode and 0.2 Da in MS² mode. Possible structure modifications included for consideration were N-terminal and lysine TMT labeling, methionine oxidation, carbamidomethylation of cysteine, and serine, threonine, and tyrosine phosphorylation. Peptides with Xcorr scores greater than 2.5 were considered for further manual validation. All TMT ratios were measured individually and experiments were carried out in triplicate.

4. Conclusions

This study represents the first global attempt to elucidate the levels of phosphorylation for three integral protein complexes in the translation initiation pathway (Table 1). Prior studies have revealed the identification of new and novel phosphosites within proteins. In this current investigation, we have provided quantification of phosphosites that may aid in further research of these important sites of regulation. As phosphorylation is a highly dynamic modification itself involving enzymatic addition and removal, levels of phosphorylation may be indicative of a more subtle mechanism of regulation incapable of being observed by identification of novel phosphosites alone. We have concentrated on those sites within this investigation that exhibit high levels of phosphorylation; however, those that display lower levels should not be ignored as gain or absence of function at those residues may be less sensitive to the absolute level of their phosphorylation. Phosphorylation levels appear to be at their highest when the residues are within a region known to be involved in binding to other proteins, metal ions, or nucleic acids. Phosphomimetic studies will ultimately unravel the significance of these phosphosites, which may be critical to protein structure and function as well as to overall influence on the translation initiation process.

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Author Contributions

A.A.: designed the study, performed experiments, isolated and provided eIF2 and eIF3, interpreted data, and wrote the manuscript; N.V.: isolated and provided eIF4G, assisted in writing of

the manuscript; W.J.: assisted in interpretation of mass spectra; C.S.F.: isolated and provided eIF4G, assisted in design of the study; J.A.L.: supervised the project, assisted in design of the study, and was involved in data interpretation and writing of the manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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Direct Analysis of hCG\u00e3cf Glycosylation in Normal and Aberrant Pregnancy by Matrix-Assisted Laser Desorption/Ionization Time-of-Flight Mass Spectrometry

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Abstract: The analysis of human chorionic gonadotropin (hCG) in clinical chemistry laboratories by specific immunoassay is well established. However, changes in glycosylation are not as easily assayed and yet alterations in hCG glycosylation is associated with abnormal pregnancy. hCGβ-core fragment (hCGβcf) was isolated from the urine of women, pregnant with normal, molar and hyperemesis gravidarum pregnancies. Each sample was subjected to matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI TOF MS) analysis following dithiothreitol (DTT) reduction and fingerprint spectra of peptide hCGβ 6–40 were analyzed. Samples were variably glycosylated, where most structures were small, core and largely mono-antennary. Larger single bi-antennary and mixtures of larger mono-antennary and bi-antennary moieties were also observed in some samples. Larger glycoforms were more abundant in the abnormal pregnancies and tri-antennary carbohydrate moieties were only observed in the samples from molar and hyperemesis gravidarum pregnancies. Given that such spectral profiling differences may be characteristic, development of small sample preparation for mass spectral analysis of hCG may lead to a simpler and faster approach to glycostructural analysis and potentially a novel clinical diagnostic test.

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1. Introduction

In a post-genomic era the importance of proteoforms has come to the fore [1], and it is the subtleties of the proteoform that underlay many pathologies not yet characterized at a genetic level. This is not simply splice variants but the form a protein takes within a functional cellular and physiological system. Critical to clinical functionality of a coded protein are its post translational modifications, e.g., pre- and pro-peptide cleavage, phosphorylation and glycosylation. Detection and relative quantification of particular proteoforms will form the bases of new biomarker discovery and not necessarily simple measurement of any given mass of protein [2].

The detection of human chorionic gonadotropin (hCG) is used extensively in obstetrics and gynecology for the detection and monitoring of pregnancy. The hormone is an $\alpha\beta$ hetero-dimeric glycoprotein with eight glycosylation sites, comprising four *N*-linked oligosaccharides and four *O*-linked oligosaccharides. Two *N*-linked oligosaccharides are attached to each of the subunit polypeptide chains by β -*N*-glycosidic bonds to asparagine residues. These moieties share the same basic structural characteristics: *N*-acetylglucosamine (GlcNAc) is attached to an asparagine residue followed by another GlcNAc, mannose, and two more branches of mannose. This is the

mono-antennary pentasaccharide core with the remaining components being variable [3–5]. The O-linked oligosaccharides are attached by α -O-glycosidic bonds onto serine residues of the β -subunit carboxyl terminal peptide [6–9].

Carbohydrate heterogeneity has been extensively reported for the free β-subunit of hCG (hCGβ) with variable mono-, bi-, and tri-antennary carbohydrate structures being found in normal and abnormal pregnancies, as well as in gestational trophoblastic disease and in particular choriocarcinoma and early pregnancy [10–14]. In general, a greater proportion of tri-antennary oligosaccharide structures are usually indicative of abnormalities in pregnancy, while bi-antennary forms account for the majority of structures found in normal pregnancy [13].

hCG is excreted intact into the urine, as documented by extensive implementation of urinary pregnancy testing. However, hCG is also degraded in liver and kidneys and a large proportion of immunoreactive hCG in the urine is attributed to this urinary degradation product of the hCG β subunit hCG β -core fragment (hCG β cf). The carbohydrate structures of the hCG β cf have been studied independently [15,16] and the molecule is composed of peptides, β 6–40 and β 55–92, connected by four disulfide bridges. It retains many of the antigenic determinants of the original hCG β molecule prior to metabolism, which occurs primarily in the kidney [17]. The β 6–40 polypeptide chain contains the two hCG β *N*-linked carbohydrate moieties, although the oligosaccharides are truncated due to metabolism. Urinary hCG β cf can be isolated with relatively straightforward procedures [15] from a simple urine sample and offers a convenient way of providing insights into glycosylation of the hCG β subunit and therefore the hCG from which it was derived [18]. This presents an opportunity to indirectly study pregnancy disorders known to exhibit glycoform variants of hCG.

Matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI TOF MS) is a technique that can be used for the determination of the mass of macromolecules, originally developed by Karas & Hillenkamp [19]. MALDI TOF MS can be used in the characterization of glycopeptides [20] and/or oligosaccharides that are released from glycoproteins with the use of enzymatic digestion [21–23]. Dithiothreitol (DTT) can also be used in situations where disulfide linkages are present and can reduce the mass of peptides bringing them into relatively optimum resolution for this mass spectrometer.

In the case of hCGβcf, the amino acids β 55–92 are linked to β 6–40 from the original β-subunit in hCG. After disulfide reduction, these two peptides along with glycosylation moieties can be analyzed by MALDI TOF MS and oligosaccharide masses calculated by subtraction of the peptide mass of the β 6–40 chain from the observed peak mass of each glycoform. Carbohydrate heterogeneity has been reported on hCGβcf and a population of mono- and bi-antennary structures has been proposed by various studies [24–27]. Using a MALDI TOF MS technique we have previously shown that the remaining oligosaccharide structures found on hCGβcf do not possess sialic acid and the extent to which those structures are truncated prior to urinary excretion as hCGβcf [28]. This made it possible to analyze glycosylation moieties whilst still attached to the peptide, thus eliminating the need for glycosidase digestion. However, this previous work was conducted on a pooled sample preparation and there has, as yet, been no report of hCGβcf glycosylation patterns from individual patients. In order to provide some insights in hCGβcf

glycosylation in aberrant pregnancies, we used the same MALDI TOF MS technique to analyze hCGβcf isolated from individual patient samples with normal pregnancy or conditions such as molar pregnancy and hyperemesis gravidarum.

2. Results and Discussion

2.1. Mass Spectral Profiles

hCG β cf purified from pregnancy urine samples (normal, molar and hyperemesis gravidarium) subjected to MALDI TOF MS generated mass spectra for hCG β cf displaying a broad peak between m/z 8700 and 10,700, as published previously [28]. On reduction of the disulfide linkages using DTT, this broad peak was replaced by a set of lower molecular weight peaks (Figure 1). A peak at m/z 3950 was seen in the spectra from hCG β cf samples N2 β cf and HG β cf (Figure 2b,e). Common to all samples was the peak at m/z 4156.8, corresponding to the non-glycosylated hCG β cf peptide β 55–92 (Figure 1).

2.2. Determination of Glycostructures

Prediction of the glyco-structures that resulted in the remaining peaks was achieved by the subtraction of the corresponding mass of the primary amino acid sequence of β 6–40 from the observed m/z values corresponding to the glycosylated isoforms (Figure 2 and Table 1). Despite the fact that the exact predicted mass of the hCG β cf asparagine-linked carbohydrate moieties were not observed directly, the low percentage errors between the observed and expected mass match of the peaks acquired show that it is likely that these glycoforms were detected. The proposed carbohydrate moieties identified from the mass spectra are shown in Figure 2. Each of the five pregnancy samples contained between 8 and 11 out of the 25 glycosylated forms of β 6–40 identified in this set of samples (Figure 2 and Table 1).

Figure 1. Matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI TOF MS) of human chorionic gonadotropin β-core fragment (hCGβcf) treated with dithiothreitol (DTT). hCGβcf purified from pregnancy urine samples; Normal (**A,B**), Molar (**C,D**) and Hyperemesis Gravidarium (**E**). Disulfide linkages were reduced using DTT. The indicated peak at m/z 4156.8 (β 55–92) appears in all samples and represents the unglycosylated peptide of beta-core. Arrowed peak (↓) only appears in samples N2βcf and HGβcf and indicates a fragment smaller than β 55–92 and as such is likely to be β 6–40 with minimal or no-glycosylation. All remaining peaks are attributed β 6–40 glycopeptides and described in Table 1.

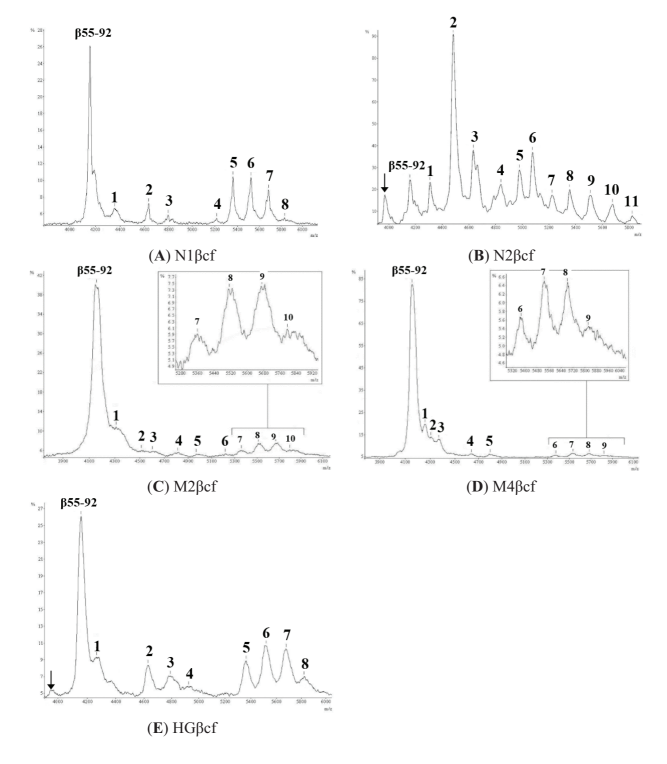


Figure 2. Oligosaccharide structures of hCGβcf. Structures identified in samples used in this study. The information for each structure includes; structure letter, schematic and molecular weight (Da). \blacksquare , GlcNAc (221.2 Da); \circ , mannose (180.2 Da); Δ , Fucose (164.2 Da); \bullet , Galactose (180.2 Da).

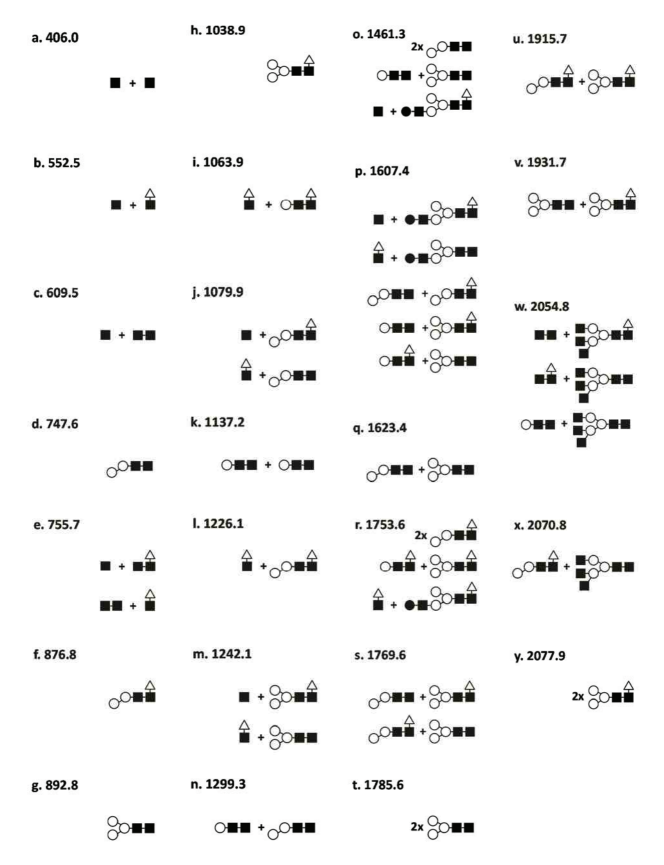


Table 1. Identifying MALDI TOF MS peaks. For each peak in each sample; an inferred oligosaccharide (CHO) mass was calculated and best fit structure assigned (Figure 2). The theoretical mass of the glycopeptides (β 6–40 plus CHO moiety) was then calculated as percentage fit (mass match) to the observed peak mass. The calculated relative abundance of each observed peak represents the proportion of area under the curve for the mass spectral range (m/z 4200–6000) for that peak/glycopeptides (% abundance).

Dools	Observed Mass	Predicted Carbohydrate	Predicted Mass (Da)	% Mass	%				
Peak	$(m/z) [M + H]^+$	Structure (Figure 2)	of Glycopeptide	Match	Abundance				
N1bcf									
1	4353.8	С	4361.9	0.9982	16.2				
2	4634.7	f	4629.2	1.0012	5.1				
3	4798.8	h	4790.8	1.0017	4.4				
4	5220.8	0	5213.7	1.0014	3.2				
5	5366.7	p	5359.8	1.0013	18.1				
6	5529.1	S	5522.0	1.0013	22.8				
7	5689.8	V	5684.1	1.0010	17.8				
8	5840.6	у	5830.3	1.0018	12.4				
			N2bcf						
1	4307.7	b	4277.9	1.007	4.7				
2	4477.6	d	4483.0	1.0012	35.2				
3	4630.9	g	4630.9	1.0030	17.3				
4	4837.4	j	4832.3	0.9989	6.6				
5	4976.6	1	4978.5	1.0004	5.3				
6	5074.3	n	5051.4	0.9955	7.5				
7	5219.7	0	5213.7	0.9899	3.3				
8	5352.6	p	5359.8	1.0013	5.9				
9	5504.9	r	5506.0	1.0002	7.3				
10	5672.8	u	5568.1	0.9992	5.3				
11	5820.9	X	5823.2	1.0004	1.6				
		I	M2bcf						
1	4310	ь	4304.9	0.9988	24				
2	4515.2	e	4508.1	0.9984	0.3				
3	4601.7	f	4628.4	1.0058	0.6				
4	4805.8	i	4816.3	1.0022	13.1				
5	4966.4	m	4994.5	1.0056	9.8				
6	5219.1	0	5213.7	0.9970	1.7				
7	5359.5	p	5359.8	1.0001	9.8				
8	5518.9	S	5522.0	1.0006	13.4				
9	5673	u	5668.1	0.9991	15.3				
10	5796.1	W	5807.2	1.0019	12				

Table 1. Cont.

Peak	Observed Mass	Predicted Carbohydrate	Predicted Mass (Da)	% Mass	%
геак	$(m/z) [M + H]^+$	Structure (Figure 2) of Glycopeptic		Match	Abundance
	M		/I4bcf		
1	4256.3	a	4158.4	0.9770	25.6
2	4305.2	b	4304.9	0.9999	16.2
3	4369	c	4361.9	0.9984	24.5
4	4639.1	f	4629.2	0.9979	6.1
5	4801.5	h	4791.3	0.9979	8.1
6	5378.9	q	5375.8	0.9994	4.1
7	5535.7	t	5538.0	1.0004	7.3
8	5686.9	V	5684.1	0.9995	5.7
9	5824.9	X	5823.2	0.9997	2.4
		H	IGbef		
1	4265.4	b	4304.4	1.0091	12.2
2	4628.3	f	4629.2	1.0002	9.2
3	4790.2	h	4791.3	1.0002	8.9
4	4929.4	k	4889.6	0.9919	3.1
5	5361.6	p	5359.8	0.9997	13.5
6	5516.8	S	5522.0	1.0079	21.7
7	5675.7	u	5568.1	0.9987	14.9
8	5820.2	X	5823.2	1.0005	16.5

2.3. Relative Abundance of Glycoforms

The most commonly detected glycol-structure found in 4 of 5 of the samples were Figure 2 structures b (m/z 552.5), f (m/z 876.8), p (m/z 1607.4); and in 3 of 5 samples h (m/z 1038.9), o (m/z 1461.3), s (m/z 1769.6) and v (m/z 1915.7). Collectively structures b (m/z 552.5), p (m/z 1607.4), and s (m/z 1769.6) represent a third of the peak abundance of all the spectra.

The incidence of the remaining glyco-structures was low as was the abundance of the mass spectra generated for the urine samples from normal pregnancies; sample N1 β cf had one unique peak at m/z 2077.9 (carbohydrate structure y in Figure 2) and sample N2 β cf four- structures: d (m/z 747.6), g (m/z 892.8), j (m/z 1079.9) and l (m/z 1226.1). Peaks corresponding to structures e (m/z 755.7), i (m/z 1063.9), m (m/z 1242.1) and w (m/z 2054.8) were present only in the hCG β cf purified from M2 β cf, whilst the spectra for the second molar pregnancy urine sample M4 β cf displayed peaks representing structure q (m/z 1623.4) and t (m/z 1769.6). Interestingly the hCG β cf preparation from the hyperemesis gravidarium pregnancy urine did not reveal any unique glycoforms.

Fucose at 1–6 of the basal GlcNac was a common retained feature of the residual glycosylation moieties, occurring in 16 of the 25 identified structures and in terms of abundance could be accounted for in 76% of the peak areas of the combined samples.

The glyco-structures that contributed to the greatest proportion of samples are; N1 β cf—s (m/z 1769.6) (22.8%); N2 β cf—d (m/z 747.6) (35.2%), M2 β cf—b (m/z 552.5) (24%), M4 β cf—a (m/z 406.0) (24.5%) and HG β cf—s (m/z 1769.6) (21.7%). Mono-antennary structures (m/z 406–1226.1) and bi-antennary structures (m/z 892.8–2077.9) were found in all samples. Tri-antennary carbohydrate

moieties w (m/z 2054.8) and x (m/z 2070.8) were only detected in molar pregnancy-M2 β cf and Hyperemesis gravidarum-HG β cf samples.

2.4. Discussion

HCG is produced by placental trophoblast cells and is a glycoprotein hormone in the diagnosis of pregnancy testing and in the detection of cancer. It would be a significant improvement on current methods to develop a rapid and reliable analytical technique for the characterisation of peptide and carbohydrate portions of hCG rather than a simple quantification of serum or urine levels. By differentiating between those hCG moieties present and with the development of analytical peptide standards for hCG, progress can really be made in identifying hCG glyco-variants as specific clinical diagnostic markers. As such the utilisation of mass spectrometry for the detection and characterisation of hCG would provide an additional diagnostic tool in both the monitoring of pregnancy and cancer.

This study examined the structural heterogeneity of hCG\u03b3cf from individuals with normal pregnancy, hydatidiform mole and hyperemesis gravidarum. MALDI TOF MS was used to analyse hCG\u00e4cf isolated from individual pregnancy urine samples. Reduction of hCG\u00e4cf purified from normal pregnancy urine resulted in the separation of the two peptides; non-glycosylated (β 55–92) and glycosylated (β 6–40) chains corresponding to the mass spectral peaks at m/z 4156.8 and 5840.6 respectively (See Figure 1). In addition to the peaks attributed to the non-glycosylated and the glycosylated peptide and its glycoforms (discussed extensively below) hCG\u00bbcf mass spectra from samples N2 β cf and HG β cf displayed a peak at m/z 3950. This peak was not detected in the pooled urine samples from multiple pregnancies from our previous study. The m/z value of this species is too high to attribute the peak to a non-glycosylated β 6–40, we speculate therefore that this is a glycoform of β 6–40 with a carbohydrate moiety of approximate molecular mass 197.6 Da. Studies of the biosynthesis of N-linked sugar chains have demonstrated that a common core of 3 mannose residues forming two branches of GlcNAc₂ (Man₃GlcNAc₂) is transferred en bloc to the polypeptide chain and that the removal of portions of this unit and addition of other sugar residues occur during subsequent processing [29,30]. This processing of the N-linked oligosaccharide and also the attachment of the sugars to specific serine or threonine residues takes place in the Golgi apparatus [31]. In the first instance it is possible to suggest that this peak is due solely to the attachment of either galactose or mannose directly on the β 6–40, as there molecular weights are both 180.2 Da. However, in line with the mechanism by which N-linked carbohydrates are processed, it may be that this peak represents β 6–40 with one GlcNAc (m/z 221.21) suggesting that this oligosaccharide may have been removed during processing and modified no further.

The remaining mass spectral peaks are attributed to the multiple glycosylated forms of the peptide β 6–40. Absolute quantification of the relative amounts of each carbohydrate moiety was not possible using this method, one of the perceived restrictions of MALDI TOF MS is its inability to quantitate from spectra. However, we applied a semi-quantitative approach by determining the areas under the peaks of the reduced peptides, similar to that used for data generated by HPLC. These results suggest that hCG is *N*-linked hyperglycosylated to a greater extent in disease and abnormal pregnancy as has been previously described [13,14] and that these glycosylation moiety variation structures are

reflected through to the pattern and abundance of urinary metabolite hCG\u00e3cf glycoforms. This combined finding suggests a possible use of hCG\u00e3cf glycoform analysis by MALDI TOF MS or other methodologies as a novel marker of these diseases.

However, before the use of MALDI TOF MS, as described here, is a clinical reality several technical problems need to be overcome: The first is that we examined purified hCG\u00b3cf originating from large pools of collected urine. This volume collection alone renders this approach in-practical for routine clinical analysis purposes. Micro-scale enrichment columns (akin to Zip TipsTM) may be needed to process, in both terms of analyte concentration and purity, the much smaller volume urine samples available/collected for large sample sets of clinical samples to be logistically (and economical) analysed by this proposed approach. Secondly, and as referred to above, a major criticism of MALDI TOF MS is that it is not quantifiable. That is the y-axis is a relative intensity within a profile and not directly proportional to the various amounts of given molecules present in the sample, i.e., the molecules that ionize easily give more intense signals compared to molecules that might be more abundant but do not ionise easily, and therefore give weaker intensity signals. This reduces the value of MALDI TOF MS spectral data; but to partially overcome this we have adopted a normalisation approach in order to render peak intensities axis comparable between sample spectra. Thus, we transformed the y-axis values to a percentage of the spectral region being compared. It has yet to be seen if such a simple processing approach is sufficiently robust to be reproducible when comparing large numbers of samples in a clinical diagnostic situation.

There is some debate as to whether the carbohydrate composition of hCG β cf in pregnancy urine can be directly correlated to that of the parental hCG β subunit. The results from this study are in line with literature that suggests that carbohydrate heterogeneity has been found in hCG β in both normal and abnormal pregnancies and that this remains in the terminal urinary degradation product hCG β cf [13,28]. Other studies suggest that hCG β cf glycoforms are very different from that of the hCG β subunit, proposing the presence of shortened asparagine-linked oligosaccharides on hCG β cf that had generally been metabolised to their pentasaccharide cores as well as smaller sugars [24–27]. One such study reported that 22%–44% of the hCG β cf failed to show binding ability to Concavalin A, which according to the authors is as a consequence of having no sugar molecules [27]. In the previous study, our group have shown that the hCG β cf glycosylated peptide β 6–40 is never completely trimmed of oligosaccharides and that there is only one non-glycosylated hCG β cf peptide, β 55–92 at m/z 4156.8 after reduction of hCG β cf with DTT [28]. This is also true for the samples in the current study. This discrepancy in the literature may be due to the difference in hCG β cf preparations; the source or the methods used for its purification and characterisation.

In our earlier study [28] hCG β cf was purified from pooled normal pregnancy urine samples and isolated by sequential size exclusion. In the present study the method of purification was ion exchange chromatography and in this case each sample; normal, molar and hyperemesis gravidarum was processed and analysed individually similar to that performed by Elliott *et al.* [13]. In fact some of the preparations used here were prepared alongside this study and as such can be compared directly to M4 and M2 hCG described therein. The largest oligosaccharides (m/z 2420.3 and 2598.4) detected in the pooled urine hCG β cf population previously were not detected in this cohort of patients in which the largest carbohydrate moiety was identified at m/z 2077.9. Previous studies in normal

pregnancy suggest that by the 10th week of gestation some tri-antennary forms fall to less than 10% of total hCG, indicating that hCG tri-antennary glycoforms, including hyperglycosylated hCG are only seen in significant proportions earlier in pregnancy [13,32,33]. As the samples collected for this study were from the gestational period 7 to 13 weeks, it is possible that the proportion of hCG β cf composed of tri-antennary sugars in the urine samples from the weeks before gestation week 10 were diluted or cleared and in each of the samples this structure occurs in such low concentrations as to be undetectable by this technique.

Our previous study [28] proposed that there was a general absence of galactose with the pooled samples even though in one of the spectra a structure was observed that correlated to a single peak $(m/z \ 1607.5)$ [28]. Peaks corresponding to this same glycoform (structure I, Figure 2) have been identified in the present study in all samples except M1 β cf. Carbohydrate structures j, k, l, m and n (Figure 2) which have been attributed to peaks in the samples analysed for this study contain, in some isoforms, a galactose residue. In the literature the galactose content of hCG β cf has been reported differently. In some studies involving carbohydrate analysis after acid hydrolysis or the conversion of sugars to glycamines, small amounts of galactose has been detected [15,27]. In contrast to this, other groups have found hCG β cf N-linked sugars lacking galactose [24,26]. MALDI TOF MS analysis of the samples in this study has highlighted peaks which contribute significantly to the overall spectrum that cannot be correlated directly with currently identified carbohydrate moieties. It is tempting to speculate about the presence of additional hCG β cf peptide variants as has been suggested previously and their potential involvement in pregnancy and pregnancy associated disorders [14].

3. Materials and Methods

3.1. Biological Samples

Urine samples from five individual pregnancies were used in this study: two were complete molar pregnancies (M2βcf, M4βcf; *i.e.*, moles existed *in utero* when the urine sample was taken), one hyperemesis gravidarum (HGβcf), and two from apparently normal uncomplicated pregnancies (N1βcf, N2βcf). Because hCG reaches its highest levels in urine during the 10th week of pregnancy, all samples were obtained between the 7th and the 13th week of gestation, therefore allowing a 3-week window on either side of the hCG peak; 3 to 5 L of urine were collected continuously from each individual over several days. M2βcf and M4βcf were collected and stored (-80 °C) previously (and intact hCG extracted, the structure of which was reported earlier) [11]. The other samples were collected and purified at the University Of New Mexico School Of Medicine (Albuquerque, MN, USA) following full consent from pregnant women and ethical approval for the study was granted by the OB/GYN departmental research ethics committee.

3.2. Sample Treatments

Proteins were precipitated from urine, initially with acetone (acetone:urine = 2:1 (v:v)) (Merck, Nottingham, UK) overnight at 4 °C according to methods described previously [34]. The precipitate was collected by centrifugation, and re-dissolved in a minimum amount of distilled-deionized water

and re-precipitated with ethanol (ethanol:sample = 9:1 (v:v)) (Merck) overnight at 4 °C. The resulting precipitate was collected by centrifugation, air-dried to remove excess ethanol, re-dissolved in a minimum amount of distilled-deionized water, and dialyzed against 0.05 M ammonium bicarbonate.

Samples M2 β cf and M4 β cf were initially fractionated by size exclusion chromatography on an S-200 Sephacryl column (Pharmacia, Piscataway, NJ, USA). The hCG β cf content of each fraction was then determined by specific immunoassay [35]. These samples were co-purified along with intact hCG, some of which were later characterized [13]. The hCG β cf fractions were lyophilized and stored at -80 °C.

All samples were then fractionated on a DEAE-Sepharose ion exchange column [36]. One hundred and thirty milliliter of DEAE-Sepharose CL-6B (Pharmacia, Piscataway, NJ, USA) was packed into an XK26 column (Pharmacia) (26×245 mm) at a flow rate of 1.5 mL/min. The void volume (V_0) was calculated by detection of changes in salt concentration using silver nitrate precipitation, after the elution buffer was changed from 0.1 to 1 M ammonium bicarbonate ($V_0 = 144$ mL). The column was then equilibrated with 2 L of 0.1 M ammonium bicarbonate buffer (pH 7.1) (Sigma-Aldrich, St. Louis, MO, USA) and kept at 4 °C at all times in order to prevent protein degradation by bacterial enzymes. Individual samples were loaded onto the column and the column was eluted with 200 mL of stepwise increases in ammonium bicarbonate buffer (pH 7.1) starting with 0.1 M at a flow rate of 1.5 mL/min, followed by 200 mL of 0.15 M, 200 mL of 0.2 M, 200 mL of 0.25 M and finally 200 mL of 1 M ammonium bicarbonate buffer. The eluent was collected as 9 mL fractions and the concentration of hCG β cf in each fraction was determined by enzyme-linked immunosorbent assay (ELISA).

3.3. hCG\(\beta\)cf Enzyme-Linked Immunosorbent Assay (ELISA)

The assay utilized a monoclonal antibody INN-hCG-106 against the β 11 epitope on hCGβcf as the capture antibody [37]. The S504 polyclonal antibody [38] was used as a primary detection antibody and a donkey-anti-sheep-HRP monoclonal (Jackson Immunoresearch Inc., West Grove, PA, USA) was used as a secondary detection antibody. All fractions with hCGβcf immunoreactivity were pooled and their hCGβcf levels were determined once again.

3.4. Matrix-Assisted Laser Desorption/Ionization Time-of-Flight Mass Spectrometry (MALDI TOF MS)

Post DEAE fractionation, samples were lyophilized against liquid nitrogen in order to remove buffers prior to mass spectrometric analysis. After two freeze-dry/rehydration cycles, the protein was re-dissolved in a minimum amount of distilled-deionized water.

3.5. Whole Molecule hCG\(\beta\)cf Analysis (Non-Reduced)

One micro-liter of sample was applied to a stainless steel MALDI TOF MS target and allowed to dry and crystallize at room temperature. 0.6 μ L of sinapinic acid (20 mg/mL⁻¹) (Sigma-Aldrich) in acetonitrile (Merck) and 0.1% trifluoroacetic acid (Merck) was applied on top of the sample and allowed to dry prior to mass spectrometric analysis.

3.6. Dithiothreitol (DTT)-Treated hCGβcf Analysis (Reduced)

Five micro-liters of neat sample was incubated with 5 μ L of 100 mM DTT (Sigma-Aldrich) in 100 mM ammonium bicarbonate for 1 h at room temperature. Sample and matrix were then applied on the MALDI TOF MS target as described above.

A pulsed nitrogen laser ($\lambda_{max} = 337$ nm) was used to desorb ions from the sample, which were accelerated by a 20 kV electrical field down a 0.5 m linear tube and detected by a micro-channel plate detector. The detector was digitized at a sampling rate of 500 MHz. Spectra were generated by summing 20–30 laser shots by using a Finnigan LASERMAT 2000 instrument (Thermo-Finnigan, Waltham, MA, USA).

Mass calibration was assigned using horse heart cytochrome C (20 pmol/ μ L "on target") as an external calibrant (two point calibration at [M + H]⁺ = 12,361 Da and [M + 2H]²⁺ = 6181 Da) for spectral analysis of whole hCG β cf. For spectra analysis of DTT-reduced hCG β cf, the non-glycosylated peptide of hCG β cf (β 55–92) was used as an internal calibrant (one point calibration at [M + H]⁺ = 4156.8 Da; calculated from its given primary sequence). A 0.5% error during peak mass allocation was allowed for, as this was typical in the linear mode for the MALDI instrument used.

3.7. Treatment of Spectra

In order to determine the masses of the carbohydrate moieties, a previously described method was used [28,39]: reduced hCG β cf spectra were calibrated by using the β 55–92 non-glycosylated peptide as described above. The inferred masses were determined by subtracting the mass of the glycosylated peptide of hCG β cf (β 6–40), which was calculated from its given primary sequence at a mass of 3752.4 Da. The carbohydrate content of each peak was then determined by sequential subtraction of the masses of individual sugar residues [28]. An error \leq 0.25% was allowed between observed and predicted carbohydrate masses.

The percentage represented by each of the peaks in individual spectra was also calculated by using the following formula:

%Area = [(Peak height from baseline \times Peak width at ½ height) \times 100] \div Σ of Spectrum Peak Area

4. Conclusions

In conclusion, hCGβcf hyperglycosylation due to tri-antennary glycoforms was found to be the highest in the urine from women with molar and hyperemesis gravidarum pregnancies compared to the samples from normal pregnancy. Although such molecules are subject to metabolic processing, this supports previously published data from Elliott *et al.*, which has shown that hCG is *N*-linked hyperglycosylated to a greater extent in disease and abnormal pregnancy. Although a very high percentage of tri-antennary glycoforms were seen on the hCGβ subunit in abnormal pregnancy in that study [13], such distinct hyperglycosylation has not previously been seen as clearly in hCGβcf. The MALDI TOF MS technique described here, although not definitive is considerably simpler and faster than conventional approaches to glycostructural analysis and presents a potential novel approach to provide additional clinical information. Chromatographic purification prior to MALDI TOF MS analysis is still laborious; however, this may become unnecessary when coupled with

affinity capture MALDI techniques as described by Neubert *et al.* [40] and in turn this may lead to more rapid analysis of multiple patients from spot urine samples.

The application of mass spectrometry in the analysis of glycosylation proteforms is developing rapidly. Glycomics, as demonstrated in Manfred Wuhrer's recent review, is now entering the clinical diagnostic arena [41] and, as a result, international searchable databases specifically addressing glycosylation patterns are emerging [42].

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Author Contributions

S.A.B. and L.A.C. carried out sample preparation and purification. R.K.I. analyzed the MS data. S.A.B. and L.A.C. prepared the clinical sample information. R.K.I. and S.A.B. designed the experiment and drafted the manuscript. All authors read and approved the final manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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Quantitative Proteomics to Characterize Specific Histone H2A Proteolysis in Chronic Lymphocytic Leukemia and the Myeloid THP-1 Cell Line

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Abstract: Proteome studies on hematological malignancies contribute to the understanding of the disease mechanism and to the identification of new biomarker candidates. With the isobaric tag for relative and absolute quantitation (iTRAQ) method we analyzed the protein expression between B-cells of healthy people and chronic lymphocytic leukemia (CLL) B-cells. CLL is the most common lymphoid cancer of the blood and is characterized by a variable clinical course. By comparing samples of patients with an aggressive *vs.* indolent disease, we identified a limited list of differentially regulated proteins. The enhanced sensitivity attributed to the iTRAQ labels led to the discovery of a previously reported but still not clarified proteolytic product of histone H2A (cH2A) which we further investigated in light of the suggested functional properties of this modification. In the exploratory proteome study the Histone H2A peptide was up-regulated in CLL samples but a more specific and sensitive screening of a larger patient cohort indicated that cH2A is of myeloid origin. Our subsequent quantitative analysis led to a more profound characterization of the clipping in acute monocytic leukemia THP-1 cells subjected to induced differentiation.

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1. Introduction

Proteomics approaches are often trailing genetic studies but are essential in the multi-disciplinary field of hematological research. As opposed to, e.g., mRNA microarray data, there is a better understanding of which proteins are actually expressed, although seeing the forest for the trees in long lists of protein identifications remains challenging [1,2]. In neoplastic hematology, protein studies have contributed to the elucidation of the disease mechanism, defined prognostic or therapeutic biomarkers and clarified previously reported uncharacterized phenomena [2]. By analyzing body fluids, cell lines, and tissues with quantitative high throughput mass spectrometry techniques complementary biological insights in hematopoietic malignancies can be generated. Uncovering relevant posttranslational modifications (PTMs), such as phosphorylations and proteolytical cleavages, might be associated with specific disease stages and could, hence, be informative on the biology of the disease [3].

The most common adult hematopoietic malignancy is chronic lymphocytic leukemia (CLL), a disease characterized by a widely variable median survival. After the initial staging of the patient, the risk of progression is defined by a set of genetic and protein based laboratory assays. A well-established

prognostic marker is the mutational status of the immunoglobulin heavy chain (*IGVH*) genes encoding for the B-cell antigen-binding domain: CLL patients who have B-cells with unmutated (UM, >98% germ line identity) *IGVH* genes have an unfavorable outcome, whereas mutated (M) *IGHV* genes predict a more indolent course [4]. Surrogate markers on the protein level, such as the Zeta-chain-associated protein kinase (ZAP) 70 and CD38, are more easily applicable in the clinical practice although CD38 is considered to have less predictive value since discordancy with gene status is commonly observed and problems with standardization occur. ZAP70 expression however, is still used in the research on CLL pathogenesis and is considered as an independent biomarker [5]. Hence, for determining disease progression and survival of CLL patients, specific genetic markers, e.g., chromosomal aberrations, such as 13q14 deletion, are of increasing importance [6]. More recently, also genome sequencing, miRNA expression profiling and methylome studies are starting to offer new insights in the disease onset and progression. By the same token, specific epigenetic modifications together with protein alterations became valuable targets in leukemia research due to their reversible character and thus potential in therapy [7].

To extend the knowledge on CLL pathology and to identify new biomarker candidates, we applied quantitative mass spectrometry strategies to target the lower abundant proteins and peptides on patient and control samples [3]. The expressional differences between isolated age-matched healthy B-cells and CLL B-cells clearly showed that the morphological differences inherent to cancerous cells challenge disease marker discovery. Our comparative proteome analysis of UM and M CLL B-cells however, revealed that remarkably, only a limited amount of the identified proteins was differentially expressed between patients with a different outcome. For both the UM and M patient group, known up-regulations of proteins contributing to cell proliferation were corroborated [8,9].

Analysis of the iTRAQ (isobaric tag for relative and absolute quantitation) data at the peptide level surfaced an interesting aberrant proteolytic product of a histone protein: clipping of the histone H2A C-tail. The specific clipping of histone H2A after V₁₁₄ (cH2A) was previously reported in leukemia and leukemia cell lines and is catalyzed by the so-called "H2A specific protease" (H2Asp) [10–14]. Recently, we identified the enzyme Neutrophil Elastase (NE) as an important candidate for the identity of the H2Asp [15]. Even though proteolysis is often not considered as a regulated PTM and the importance of protein degradation in biological functions is frequently unclear, new technologies have started to unravel the critical role of clipping in cellular homeostasis and disease [16,17]. In some reports, histone clipping has even been suggested to be a functional modification with epigenetic potential [18,19]. More specifically, cH2A caught our attention as H2A is the only histone with a C-terminus protruding out of the nucleosomal core and the clipping site is localized at the entry and exit points of the DNA [20]. In general, histone modifications help in determining the heritable transcriptional state and lineage commitment development in normal B-cells [21]. Consequently, we persisted in the investigation of the H2A clipping in CLL as disruption of the histone code is suggested to drive hematopoietic cells in lymphomagenesis [22]. Here, we initially observed that H2A clipping was more abundant in CLL patients compared to healthy controls while no differences were found between M and UM. However, we showed this was not due to the disease itself, since this clipping seemed to be associated with the amount of myeloid cells present in the predominantly lymphoid samples. To further unfold the actual role of histone H2A proteolysis, we examined this cH2A clipping during induced differentiation of myeloid THP-1 cells into macrophages through quantitative mass spectrometry [12]. We concluded that synchronization

of the THP-1 cells before the stimulation abrogates the temporally uprise of H2A clipping which has initially been observed at the onset of the differentiation.

2. Results

2.1. Differential Protein Expression between Healthy B-Cells and Chronic Lymphocytic Leukemia (CLL) B-Cells of M⁻ and UM⁺ Staged Patients

The difference in protein expression between the B-cells of patients with indolent mutated ZAP70⁻ (M⁻) CLL, the aggressive unmutated ZAP70⁺ (UM⁺) CLL and healthy 50+ donors was quantified using iTRAQ. The six runs were merged into a dataset which comprised 18,014 MS/MS (mass spectrometry) queries, yielding 7473 annotated peptides which were derived from 536 unique proteins (Table S1a). For the ratios included in the analysis, in total 107 proteins differed significantly between groups (Table S1b). Among the leukemia samples, only 22 proteins were distinct between M⁻ and UM⁺ as opposed to 70 between healthy and both CLL samples. One label in each run comprised the same pool of all CLL samples, which was included to simplify inter-run comparison and to increase sensitivity.

Functional clustering annotation of the protein lists from differentially expressed proteins between healthy and leukemia B-cells, showed that most up-regulated proteins in the leukemia samples (both UM⁺ and M⁻) are involved in mRNA processing, implying an increased transcriptional activity in cancer cells (Table S1c). Compared to healthy, most of the proteins that are down-regulated in the leukemia cells are predominantly connected to actin binding and cellular localization [23]. A cellular component analysis confirmed that 64% of the proteins up-regulated in CLL are categorized as nuclear whereas down-regulated proteins are primarily located in the cytosol (60%) (Table S1c) [24]. These results suggest morphological differences between the healthy and cancerous cells rather than true molecular aberrancies.

In the limited list of proteins specifically overexpressed in the aggressive UM⁺ compared to M⁻ two proteins are involved in ATP-binding, one is a ribosomal subunit and the H2B and Prohibitin proteins are involved in chromosomal organization (Table 1). For proteins significantly down-regulated in UM⁺/M⁻, the Interleukin enhancer-binding factor 3 and Bcl-2-associated transcription factor 1 are associated with DNA but most proteins are involved in metabolic mechanisms or are cytoskeletal.

2.2. Quantitative Mass Spectrometry and Western Blot Analysis on CLL Samples Endorsed the Myeloid Origin of H2A Clipping

Additional in-depth analysis at the peptide level, surfaced one remarkably aberrant modification in all the samples of the described iTRAQ analysis: clipping of histone H2A after V₁₁₄ (Figure 1). The peptide VTIAQGGVLPNIQAV (m/z 740.4, charge 2+) was the only semi-tryptic peptide out of the >7400 annotated MS/MS spectra that was identified in all six runs. The annotation of the peptide was confirmed by *de novo* sequencing on the MS/MS spectrum (Figure S1). These results highlight how iTRAQ chemistry contributes to a better annotation of semi-tryptic peptides by enhancing the sensitivity due to the multiplexing and increased *b*-ion formation.

Figure 1. Sequence of histone H2A. Sequence from Uniprot [25]. The VTIAQGGVLPNIQAV peptide (m/z 740.4, charge 2+) was the only semi-tryptic peptide identified in all six runs. * indicates the clipping site after V₁₁₄ (here described as amino acid 115 since methionine, generated by the start codon, is the first amino acid in the Uniprot sequence); Bold underlined sequences are the same as the isotopically labeled absolute quantification (AQUA) peptides, used for the subsequent specific cH2A quantitation. The bold double underlined sequence is the Western blot epitope.

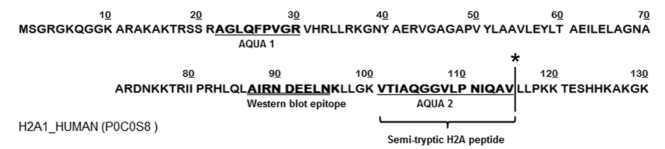
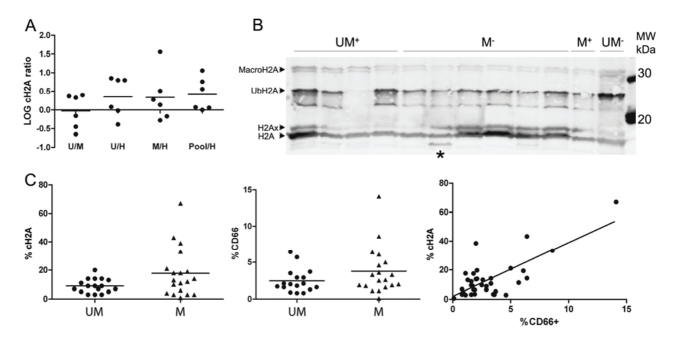


Table 1. Proteins which are differentially expressed between UM⁺ and M⁻ Chronic Lymphocytic Leukemia (CLL) B-cells. The table lists the protein name, Uniprot_ID, average UM+/M ratio, the number of iTRAQ samples where the protein was identified (n), the p-value of the performed t-test and the protein function (derived from UniProt GO).

•					
$UM^+ \nu s. M^-$	Protein Name	Identified Protein	UM*/M- Average Ratio	n p-Value	Function (GO)
	Serine/arginine-rich splicing factor	SRSF1_HUMAN	0.61	4 2.77×10^{-4}	RNA binding
		SRSF3_HUMAN	0.77	$5 1.30 \times 10^{-2}$	RNA binding
		SRSF7_HUMAN	0.90	3 2.70×10^{-2}	RNA binding
_ J /	Bcl-2-associated transcription factor	BCLF1_HUMAN	69.0	3 3.84×10^{-2}	DNA binding, induction of apoptosis
I / ₊ J /	40S ribosomal protein	RS21_HUMAN	0.70	3 3.64×10^{-2}	Ribosomal subunit
NΩ		RS28_HUMAN	0.67	4 1.47×10^{-2}	Ribosomal subunit
ui l	Stress-induced-phosphoprotein	STIP1_HUMAN	0.73	3 4.68×10^{-2}	Golgi apparatus
əte	Protein S100-A6	S10A6_HUMAN	0.75	$5 2.24 \times 10^{-2}$	Calcium ion binding
gng	Heat shock cognate 71 kDa protein	HSP7C_HUMAN	0.79	5 1.44×10^{-2}	ATP binding
ı-re	Matrin-3	MATR3_HUMAN	0.80	$3 1.85 \times 10^{-2}$	RNA binding
IMO	Ezrin	EZRI_HUMAN	0.84	4 3.32×10^{-2}	Actin filament
De	Transitional endoplasmic reticulum ATPase	TERA_HUMAN	0.84	$5 1.13 \times 10^{-2}$	ATP binding
	Interleukin enhancer-binding factor	ILF3_HUMAN	0.87	3 5.81×10^{-3}	DNA binding
	Heterogeneous nuclear ribonucleoprotein	HNRPK_HUMAN	0.88	$6 1.38 \times 10^{-3}$	ATP binding, RNA binding
	Heterogeneous nuclear ribonucleoprotein	ROA1_HUMAN	0.89	$5 1.15 \times 10^{-2}$	mRNA splicing
	40S ribosomal protein	RS13_HUMAN	1.27	$5 ext{ 4.81} imes 10^{-2}$	Ribosomal subunit
uị	Prohibitin	PHB_HUMAN	1.35	4 3.72×10^{-2}	DNA replication
M_ M	Endoplasmin	ENPL_HUMAN	1.80	$3 2.05 \times 10^{-2}$	ATP binding
il ⁺ IV	ATP synthase subunit beta, mitochondrial	ATPB_HUMAN	1.93	$3 3.72 \times 10^{-2}$	ATP binding
o-re Ul	Histone H2B	H2B1C_HUMAN	2.59	3 1.24×10^{-2}	DNA binding
ďΩ		H2B1D_HUMAN	1.10	3 1.84×10^{-2}	DNA binding
		H2B10_HUMAN	1.18	3 2.97×10^{-2}	DNA binding

For quantitation of this fragment, we compensated for abovementioned morphological differences by normalizing the reporter intensities of the clipping fragments to the average of the whole H2A protein. The relative amount of H2A that is cleaved after V₁₁₄ was found to be on average higher in leukemia cells compared to healthy B-cells as seen by the log-transformed ratios. Particularly, all six ratios of the pool of UM⁺ and M⁻ samples were consistently positive opposed to healthy (Figure 2A).

Figure 2. Quantitative mass spectrometry and Western blot analysis on CLL samples revealed that histone H2A clipping is from myeloid origin. (A) The iTRAQ ratios of the cH2A peptide (normalized to the average of the H2A protein to compensate for morphological differences) hint towards an increased abundance in CLL compared to healthy B-cells. The individual log ratios of all six runs are presented as dots, the average ratios as the horizontal bars. Histone H2A (cH2A) is up-regulated in the samples of the leukemia pool compared to the samples of the healthy B-cells in all six runs; (B) The histone extracts from 12 of the 36 samples (From left to right: patient samples UM⁺ 8, 9, 10 and 11; M⁻ 9, 10, 11, 12, 13 and 14; M⁺ 5 and UM⁻ 6). Western blot with an H2A antibody against the epitope depicted in Figure 1 detects the H2A variants H2Ax, Ubiquitinated H2A and macroH2A. The band under ubiquitinated H2A could not be identified. cH2A was only faintly detected, except for one sample (*); (C) Specific cH2A screening with AQUA peptides and flow cytometry data revealed the myeloid characteristic of the clipping. Left panel: %cH2A differs significantly (p = 0.049)between CLL patients with a distinct mutational status (UM: 11× UM⁺ & 6× UM⁻, M: $14 \times M^- \& 5 \times M^+$); **Middle panel**: Although not significant (p = 0.15), the %CD66b suggested a similar correlation with the mutational status; **Right panel**: Relation between %cH2a and %CD66b. Spearman's Rho correlation between CD66b⁺ and %cH2A was significant at the 0.01 level (Spearman's Rho correlation coefficient: 0.439; p = 0.007; Data: Table S2b).



The consequences of proteolytic PTMs are ubiquitously underappreciated so we persisted in the more specific investigation of cH2A in a larger patient population due to the biomarker potential of this previously reported modification (Scheme of workflow: Figure S2). Immunodetection of H2A on histone extracts visualized cH2A only in one sample (Figure 2B, SYPRO staining: Figure S3). The flow cytometry data showed that this particular sample had the highest amount of granulocytes (expressed as %CD66b⁺ cells) in the screened patient population. In line with this, direct comparison of %cH2A with the more sensitive and specific AQUA (absolute quantification) approach suggested higher clipping in the M compared to UM CLL B-cells (Figure 2C, left panel) where a similar relationship between the amount of granulocytes cells and the respective mutational status was observed (Figure 3C, middle panel). Indeed, a statistical analysis of the possible correlation between the amounts of V₁₁₄ clipped H2A and all cellular markers investigated for each sample confirmed a significant correlation between the amount of CD66b⁺ cells present in the samples and %cH2A (Figure 3C, right panel) (Details patient screening are listed in Table S2). Even though all steps were performed at 4 °C in the presence of protease inhibitors, we observed that H2A clipping can actually be caused in vitro to some extent as we observed some residual clipping activity on spiked-in biotinylated H2A (data not shown).

2.3. Characterization of cH2A in THP-1 Cells

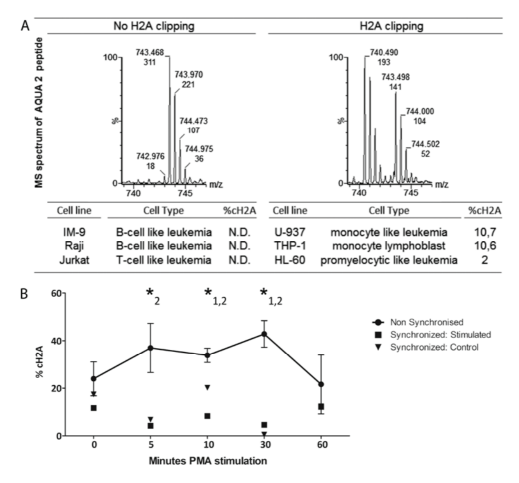
To further corroborate the myeloid nature of cH2A we cultured human leukemia cell lines of different origin and prepared histone extracts when a density of approximately $1 \times 10^6/\text{mL}$ was achieved. In all the investigated lymphatic cell lines, no clipping of H2A was detected. For the myeloid cells, $\pm 10\%$ cH2A was measured in the monocyte-like THP-1 and U-937 cell lines and, although not unambiguously, $\pm 2\%$ in the promyelocytic HL-60 cells (Figure 3A).

Next, we applied our sensitive and specific AQUA approach to specifically quantify H2AV₁₁₄ clipping in a previously reported model in which Ohkawa *et al.* describe a temporal increase of truncated H2A proteins during stimulated differentiation of acute monocytic leukemia THP-1 cells into macrophages using both phorbol 12-myristate 13-acetate (PMA) and retinoic acid (RA) [13]. For three biological replicates, %cH2A was specifically quantified in histone extracts, isolated at different time-points shortly after PMA supplementation (Figure 3B). The clipping at $T_{\rm PMA0}$ was significantly different than at $T_{\rm PMA~10}$ (p=0.045) and $T_{\rm PMA60}$ (p=0.011). Equally as to $T_{\rm PMA~30}$ (p=0.0075). These results confirm that H2A clipping at V₁₁₄ indeed ascends and sequentially decreases upon THP-1 differentiation.

Finally, since we hypothesized that the variation might be due to differences in cell cycle synchronization, THP-1 cells were synchronized by double thymidine block, as the more open chromatin structure during cell division could very well be more susceptible to proteolytical activity during cell lysis. After synchronization, cells were either left untreated or were subjected to differentiation and cH2A was subsequently quantified by AQUA screening on histone extracts, obtained from different time points (Figure 3B). %cH2A was significantly lower in both the control $(p = 6.09 \times 10^{-4})$ and stimulated $(p = 9.18 \times 10^{-5})$ THP-1 cells compared to the unsynchronized cells. Further, no difference was observed between the synchronized stimulated and control cells (p = 0.22).

Sampling 48 h after the start of the differentiation indicated that the amount of clipped H2A was further reduced in both cell lines (data not shown).

Figure 3. Specific cH2A quantitation in myeloid cell lines. (**A**) Example of an MS spectrum of the AQUA 2 peptide if H2A V₁₁₄ clipping is absent (**left**) or present (**right**). The analysis of different cell lines confirms the myeloid characteristic of cH2A in the U-937, THP-1 and HL-60 cells. H2A clipping is not present in the investigated lymphatic cell lines and could not be unambiguously detected in the promyelocytic HL-60 cell line; (**B**) THP-1 cells stimulated with PMA show a transient H2A clipping pattern. Three biological replicates display the high variance. *1: significant differences of %cH2A between $T_{\rm PMA0}$ and the other data points; *2: the equivalent for $T_{\rm PMA60}$. The %cH2A of synchronized stimulated and control THP-1 cells, presented respectively as squares and triangles, is lower than the %cH2A of non-synchronized cells.



3. Discussion

Quantitative proteomics on patient samples and on leukemia cell lines can help to define new biomarkers and insights into the pathogenesis of lymphoid and hematopoietic neoplasms [2,3]. In our comparative proteome study between healthy and CLL B-cells of patients with different prognosis, we implemented two quantitative label-based mass spectrometry methods: iTRAQ and AQUA. The isobaric character of the iTRAQ labels allows multiplexing different samples, resulting in an

increased signal and lower sample complexity [26]. We included both CLL samples of patients with a different disease prognosis and healthy B-cells in the analysis to obtain deeper insights in disease biology. A substantial part of the observed significant differences in relative protein expression between healthy and CLL B-cells is primarily due to differences in morphology between healthy and neoplastic cells [27]. Cancerous B-cells are indeed characterized by a larger nucleus and a denser cytoplasm, which we affirmed here by functional grouping and cellular component analysis of the up- and down-regulated proteins. We caution for the interpretation of proteomics data when healthy and malignant cells differ strongly in their morphology, an important restriction that is not always validated prior to relative comparison in proteomics approaches. From a clinical perspective however, aberrant ratios between patient groups with UM⁺, who have a bad prognosis and poor overall survival range and the less aggressive M⁻, are more relevant. Notably, less than 5% of the identified proteins were significantly different between these two groups. In this list of proteins with a log ratio significantly different from zero, defining distinctions between M⁻ and UM⁺, one of the most remarkable candidates is the Bcl-2-associated transcription factor 1 ($p = 3.84 \times 10^{-2}$). The Bcl-2 family regulates apoptosis and is an established hallmark in CLL as aberrant expression of Bcl-2 proteins causes apoptosis resistance of CLL B-lymphocytes. Although not all the Bcl-2 proteins correlate consistently with known CLL biomarkers, several Bcl-2-anatagonists are in clinical trials for CLL treatment [8].

The use of the iTRAQ label and the subsequent manual analysis of the results at the peptide level together surfaced another fascinating finding: a semi-tryptic peptide derived from histone H2A clipped at V₁₁₄. Of all the trypsin-based mass spectrometry experiments uploaded in the PRIDE database, this fragment was only annotated once (accession: 10,528) [28]. Remarkably, we identified the peptide in all six runs. This could be explained by the contribution of the iTRAQ labels which are covalently bound to the peptide *N*-termini, generating intense b-series and consequently contributing to an in increased peptide score [26]. cH2A is generated by the removal of fifteen amino acids from the carboxy-terminal end of the intact H2A molecule after V₁₁₄, coordinately removing K119 which is an important site of mono-ubiquitination [29–33]. cH2A had been described over 35 years ago as a product of H2A specific protease (H2Asp) activity [14] and shortly afterward, other groups observed similar cleavage patterns in extracts from both myeloid and lymphatic leukemia's [10,11]. Since then, this clipping event was only referenced on a few occasions.

Although the biological function of protein degradation is largely unknown, proteolysis is an important category of PTMs, e.g., 5%–10% of all drug targets are proteases [16]. Truncation of histone tails has already been linked to cell differentiation and the *C*-tail of H2A is known to be important in cellular homeostasis and chromatin biology [18,20]. We recently identified the myeloid enzyme NE as being a prime candidate to fulfill the reaction mediated by the H2Asp but could not clarify if cH2A formation is involved in an epigenetic process or is rather a consequence of NET formation [15]. In healthy hematopoietic cells we only observed H2A clipping in cells of myeloid origin but the references to clipped H2A found in literature are mainly in the context of leukemia [10–13]. We thus persisted in a more detailed analysis of this modification and examined if cH2A formation is an epigenetic hallmark of CLL.

The preliminary iTRAQ results indicated a greater average abundance of the cH2A peptide in leukemia samples compared to the healthy controls. As validation is required, we specifically quantified cH2A in histone extracts derived from CLL B-cells of 36 clinically staged patients by applying the isotopic synthesized AQUA peptides that allow to target specific known H2A peptides, present in low concentrations [26]. We could this time not define any direct connection between %cH2A and any known disease marker. However, the amount of granulocytes in the samples did correlate with %cH2A, which corroborates the myeloid character of cH2A and is in line with the identification of NE. As complete inhibition of proteases is known to be very challenging, as also seen by us in a spike-in experiment of biotinylated H2A (not shown), it is difficult to define how much, if any, cH2A is endogenously generated [34].

On the other hand, the reported transient clipping of the H2A *C*-tail during the induced differentiation of THP-1 promonocytes, implies a potential biological role of H2A processing in the hematopoietic development of cells from myeloid origin [13]. Cells may for instance have mechanisms to control histone degradation for re-establishing the epigenetic marks on their tails in the proliferating state [35]. However, although our AQUA results demonstrate a brief uprise in specific H2A V₁₁₄ clipping upon PMA or RA stimulation, synchronization of the THP-1 cells before the stimulation abrogated such trend. Histone clipping seems to correlate with the myeloid cell cycle and as our results suggest, cH2A fluctuation is more likely caused by possible differences in cell cycle stage synchronization, rather than PMA or RA induced differentiation. Instead of being a regulated mechanism, we hypothesize that the high degree of variation found in these experiments probably is due to the more open chromatin structure during cell division rendering histones more susceptible to proteolytical activity.

4. Experimental Section

4.1. Cells and Reagents

Phosphate buffered saline (PBS), media, L-glutamine, Fetal bovine serum (FBS), penicillin/streptomycin, Dynabeads and SYPRO Ruby were from Life Technologies (San Diego, CA, USA), ammonium bicarbonate (ABC), sodium dodecyl sulfate (SDS), *N*-cyclohexyl-3-aminopropanesulfonic acid (CAPS) and Tween-20 from Millipore (Billerica, MA, USA). ReadyPrep sequential extraction kit was from BioRad (Hercules, CA, USA) and Vivaspin-2 columns from Sartorius (Göttingen, Germany). The Recombinant human H2A (M2502S) was obtained from New England Biolabs (Ipswich, MA, USA) and bovine histone extract (cat. no. 223565) from Roche (Basel, Switzerland). All other reagents were purchased from Sigma Aldrich (St. Louis, MO, USA) unless described otherwise. Raji, Jurkat and HL-60 cells were cultured in Dulbecco's Modified Eagle Medium and IM-9, U-937 and THP-1 cells in RPMI-1640 medium, both enriched with 2% (*w/v*) L-glutamine, 10% (*w/v*) FBS and 50 IU/mL penicillin/streptomycin. To achieve synchronization at the late G1– early S phase, 2 mM thymidine was added over two intervals of 12–16 h, with an incubation in non-thymidine containing RMPI-1640 medium for 8 h in between 50 ng/mL PMA was added to the THP-1 medium for the differentiation of the non-synchronized cells and 1 μM RA for the differentiation of the synchronized cells.

4.2. Patient Samples

For the iTRAQ analysis, whole blood of six UM⁺ and six M⁻ clinically staged CLL patients was obtained from Ghent University Hospital, Department of Hematology. Informed consent was given according to the requirements of the Ethics Committee of the Ghent University Hospital. PBMCs were isolated from a Ficoll-Paque (GE Healthcare, Waukesha, WI, USA) density gradient, B-cells were purified using Dynabeads Untouched Human B (Life Technologies, San Diego, CA, USA). Correspondingly, control samples were isolated from healthy volunteers aged 50+. For the high throughput screening of cH2A, samples were obtained from the UCSD CLL Research Consortium (CRC). Immediately after thawing, cells were washed twice with cold PBS containing 1 mM phenylmethanesulfonyl fluoride (PMSF) and protease inhibitor cocktail (Roche, Basel, Switzerland). CD5⁺CD19⁺ cells consistently made out more than 85% of the lymphocyte population as seen by flow cytometry.

4.3. Flow Cytometry

For each measurement 2 × 10⁵ cells were washed twice at 4 °C with PBS 1% Bovine Serum Albumin (BSA) and analyzed using a Cytomics FC500 flow cytometer (Becton Dickinson Immunocytometry Systems, San Jose, CA, USA) with monoclonal antibodies (mAbs) antibodies from BD-Biosciences (Franklin Lakes, NJ, USA): Isotype controls, anti-CD5 (PECy5), anti-CD19 (FITC), anti-CD33 (PECy5), anti-CD66b (FITC) and anti-Annexin V (FITC). The synchronization of THP-1 cells was monitored with Propidium Iodide staining.

4.4. Cell Lysis and Histone Isolation

All steps were performed at 4 °C. To obtain a complete cell lysate for the iTRAQ analysis, cells were washed twice with PBS, pelleted and resuspended in the ReadyPrep sequential extraction buffer 1 at 5 × 10⁶ cells/mL, supplemented with protease inhibitor cocktail (Roche), 1 mM PMSF, 10 μ L 200 mM Tributylphosphine (Biorad), 20 μ L phosphatase inhibitor cocktail 1 and 2 and 1 μ L 250-units/ μ L benzonase. After sonication and centrifugation at 1500 rpm for 5 min, the proteins in the supernatant were transferred to a new eppendorf. The obtained pellet was resuspended in buffer 3 from the extraction kit and sonicated for 10 min. After centrifugation at 1500 rpm for 5 min the supernatant was pooled with the previous extract. Detergents, inhibitors and urea were removed by washing twice with Milli-Q water on a Vivaspin-2 column.

For the histone extracts, harvested cells were washed twice in PBS containing 1 mM PMSF, and protease inhibitor cocktail. 10^7 cells/mL were resuspended in Triton extraction buffer (PBS containing 0.5% (v/v) Triton $100\times$, 1 mM PMSF and protease inhibitor cocktail) and lysed by gentle stirring. Pelleted nuclei were subsequently washed in PBS containing 1 mM PMSF and proteinase inhibitor cocktail. Histones were extracted overnight after benzonase treatment of the sonicated nuclei by acid extraction: incubation in 250 μ L 0.2 M HCl at 4 °C with gentle stirring. Precipitated proteins were pelleted and the supernatant containing the histones was dried and stored at -20 °C until further use. The Bradford Coomassie Assay determined the protein content of all samples.

4.5. Western Blot Analysis

An amount of 3 µg of a dried histone extract was suspended in Laemmli buffer and run on a 15% PreCast Gel (Biorad) for 30 min at 150 V and 60 min at 200 V and subsequently transferred to a nitrocellulose membrane in a 10 mM CAPS buffer with 20% MeOH (Merck, New York, NY, USA). The remaining proteins in the gel were visualized after overnight Sypro Ruby staining. For Western blot, all steps were performed at room temperature with intermediate washing steps in 0.3% Tween-20 in PBS. The Histone H2A (LS-C24265, LifeSpan BioScience, Seattle, WA, USA) antibody incubation was performed overnight in a 1:1000 dilution in PBS 1% BSA, followed by 1 h incubation in the same buffer, with a stabilized horseradish-peroxidase (HRP)-conjugated goat anti-rabbit immunoglobulin G (Pierce, Rockford, IL, USA). The Supersignal West Dura Extended Duration Substrate (Pierce) was applied to perform the chemiluminescence and both the gel and Western blot membrane were visualized with a VersaDoc Imaging System.

4.6. Quantitative Mass Spectrometry Analysis

All digests were performed according to the iTRAQ (ABSciex, Foster City, CA, USA) reagent Kit guidelines, as was the given labeling itself. For the iTRAQ analysis, six 4plex runs each encompassed 4 × 100 µg of the protein lysates from CLL B-cells from UM⁺, M⁻ and B-cells from healthy donors aged 50⁺. The fourth label was used for an additional pool of all 12 CLL samples for inter-run comparison and to increase the number of identifications. Technical variation was also minimized by reversing the labeling order. All six samples were first fractionated off line into 12 fractions on a Poros 10S strong-cation exchange (SCX) column (300 μm i.d. × 15 cm, ABSciex, Foster City, CA, USA) for subsequent nano reversed phase liquid chromatography (Dionex U3000, Dionex, Chelmsford, MA, USA) separation on a gradient specifically optimized for sample content of each fraction [36]. All other samples were separated on a 70 min organic gradient from 4%–100% buffer B (80% (v/v) acetonitrile (Millipore, Billerica, MA, USA) in 0.1% (v/v) FA). All samples were analyzed on an Electrospray ionization—Q-TOF Premier (Waters, Wilmslow, UK). Data was searched against the SwissProt (541,954 sequence entries) database using Mascot 2.3 and additionally manually interpreted with Mascot Distiller (Matrix Science, London, UK). Proteins were included for the analysis if a proteotypic peptide had a Mascot score above 45 in at least 3 runs. For iTRAQ quantitation, ratios were normalized based on summed intensities and only proteins recurring in at least three out of six different runs were withheld by an automated in-house approach. Gene ontology (GO) analysis was performed on the significantly up- or down-regulated proteins with Uniprot, DAVID and the WEB-based GEne SeT AnaLysis Toolkit (WEBGESTAT) [23–25].

For specific cH2A screening, a mix of the isobaric peptides (AQUA1, Thermo, Waltham, MA, USA) AQUA2, Sigma Aldrich (St. Louis, MO, USA) was added right before MS analysis. The tryptic H2A *N*-terminal peptide R.AGLQFPVGR.V (*m*/*z* 475.7 was used to quantify the total amount of histone H2A present in the sample (AQUA 1). Specific clipping is quantified by means of the semi-tryptic peptide K.VTIAQGGVLPNIQAV.L (*m*/*z* 743.4) (AQUA2). This is the same sequence as was found during the iTRAQ analysis. To 1 µg of HE, 10 pmol of AQUA1 and 1 pmol of AQUA 2 was spiked right before the MS analysis [15]. For the quantitation, the total ion current (TIC) from

both AQUA peptides was obtained from a single MS-scan acquired on the top of the two extracted-ion chromatograms. An example of the MS spectra of the two AQUA peptides is given at the right site of the accolade in Figure S2. To confirm the efficiency of the histone extraction, the raw data of the data-directed analysis was equally searched against the SwissProt database using Mascot 2.3 (Matrix Sciences, London, UK).

4.7. Statistical Analysis

For the iTRAQ analysis, ratios were log transformed and averaged. Statistical analysis was performed by means of a homoscedastic two-tailed t-test to inspect which log ratios were significantly different from zero (p = 0.05). The average ratio of a protein indicated whether a protein is up- or down-regulated between two samples. The same t-tests were performed to test if %cH2A and %CD66b was different between the M and UM patients. For the comprehensive analysis of the AQUA screening, the relationship between cH2A and the other variables was determined by the nonparametric Spearman's Rho correlation test wherein the ZAP and the mutational status were analyzed as dichotomized values (+ or -) with the SPSS Statistics 20 software (Endicott, NY, USA). To corroborate if the clipping temporally increased during the differentiation of the THP-1 cells, one-tailed t-tests were applied to compare each data point with the %cH2A at t-test was used to validate if the %cH2A between the synchronized stimulated and control cells was analogous.

5. Conclusions

We particularly emphasize both the pitfalls and the benefits of applying quantitative proteomic strategies in hematological research. Although morphological differences hamper proteome-wide comparison of healthy and leukemia B-cells, we resurfaced a previously described clipping product of the H2A *C*-tail. A more profound characterization in CLL and THP-1 samples indicates that H2A V₁₁₄ clipping occurs in hematopoietic cells of the myeloid lineage. Although the process of histone clipping has considerable epigenetic potential, many questions about the relevance of specific histone proteolysis remain.

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Author Contributions

P.G. wrote the manuscript and performed the bulk of the described laboratory and mass spectrometry experiments together with the subsequent data analysis. L.V. performed the THP-1 synchronization and associated flow monitoring and shared her expertise on cell culturing and overall histone research. K.V.S. helped with/optimized the electrophoresis and Western blot analyses and

reviewed the writing. S.L. assisted with the design of the iTRAQ analysis and performed the statistics of the CLL screening. F.V.N. wrote the semi-automated data analysis tool and contributed to the interpretation of the data. F.O. provided the healthy and CLL patient samples for the iTRAQ analysis and supervised the determination of the mutational status of the IGVH genes and the ZAP70 expression. T.K. provided the CLL patient samples for the AQUA analysis and supervised the determination of the mutational status of the IGVH genes and the ZAP70 expression. M.D. performed the iTRAQ analysis and assisted with the subsequent experiments, data analysis and the writing of the manuscript. D.D. supervised this work.

Conflicts of Interest

The authors declared no conflict of interest.

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Radiation-Induced Changes in Serum Lipidome of Head and Neck Cancer Patients

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Abstract: Cancer radiotherapy (RT) induces response of the whole patient's body that could be detected at the blood level. We aimed to identify changes induced in serum lipidome during RT and characterize their association with doses and volumes of irradiated tissue. Sixty-six patients treated with conformal RT because of head and neck cancer were enrolled in the study. Blood samples were collected before, during and about one month after the end of RT. Lipid extracts were analyzed using MALDI-oa-ToF mass spectrometry in positive ionization mode. The major changes were observed when pre-treatment and within-treatment samples were compared. Levels of several identified phosphatidylcholines, including (PC34), (PC36) and (PC38) variants, and lysophosphatidylcholines, including (LPC16) and (LPC18) variants, were first significantly decreased and then increased in post-treatment samples. Intensities of changes were correlated with doses of radiation received by patients. Of note, such correlations were more frequent when low-to-medium doses of radiation delivered during conformal RT to large volumes of normal tissues were analyzed. Additionally, some radiation-induced changes in serum lipidome were associated with toxicity of the treatment. Obtained results indicated the involvement of choline-related signaling and potential biological importance of exposure to clinically low/medium doses of radiation in patient's body response to radiation.

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1. Introduction

Metabolomics, an emerging field of the "omics" sciences, has a capacity to deliver essential information about small biomolecules (<1 kDa) that are end-products of all cellular processes. Lipidomics, which deals with dynamic changes of cellular lipids and their derivatives, is one of the most complex areas of metabolomics [1]. More than 500 different lipid species was reported to be present in human plasma specimens [2]. The most abundant category of lipids are glycerophospholipids (phospholipids; PLs). PLs are both key components of biological membranes and important players in different cellular mechanisms [3,4]. Derivatives of PLs are important signaling molecules involved in regulation of proliferation and apoptosis [5,6]. Of note, metabolism of phosphatidylcholines (PCs) and other PLs is significantly disturbed in cancer cells, hence elevated serum levels of their precursors (e.g., choline) and derivatives (e.g., lysophosphatidylcholines, LPCs) are promising cancer markers [7]. Changes in level of choline-containing lipids were observed in

malignant tumors during anti-cancer therapy [8]. Metabolism and blood levels of PLs changed also after exposure to ionizing radiation [9,10]. Although such effects have only been studied in animal models until now, they indicated applicability of serum phospholipid profiles in assessment of radiation exposure.

Radiotherapy (RT), either alone or in combination with chemotherapy, is an effective treatment of different types of cancer allowing preservation of structure and function of a target organ. The effects of ionizing radiation concerns damage induced not only in cancer cells, but also in adjacent normal tissue. Conformal methods of radiotherapy, like intensity-modulated radiation therapy (IMRT), were developed to allow precise delivery of high radiation doses to a tumor volume, minimizing the dose delivered to surrounding normal tissues [11]. This technique is being used most extensively in treatment of tumors located near critical structures, such as head and neck cancers [12]. IMRT is accomplished by application of many non-coplanar radiation fields that markedly extends the volume of normal tissues being exposed to low doses of radiation, for which biological relevance is not clear at the moment [13]. Radiation-induced damage of normal tissues could lead to acute and/or late injury reactions, which in extreme cases might significantly affect patient's comfort and effectiveness of the treatment. For this reason planning and monitoring of radiotherapy would be greatly facilitated if molecular markers of individual response to radiation were available in the clinical practice. In addition, molecular markers of exposure to ionizing radiation would have a great applicability in the epidemiology field and for exposure assessment after radiation accidents [14].

Local irradiation during cancer radiotherapy induces patient's whole body response that could be detected at the level of blood components. Markers of human exposure to ionizing radiation have been searched in blood cells using different genetic and genomics approaches [15–18]. Mass spectrometry-based proteomics approaches have been also explored, which allowed identifying of radiotherapy-related changes in serum proteome of cancer patients [19,20]. More recently, it has been shown that IMRT-induced changes in the low-molecular-weight fraction of serum proteome of head and neck cancer patients were affected by clinically irrelevant doses of radiation delivered to large volumes of normal tissues [21]. Here we aimed to extend the analysis of radiotherapy-related changes and radiation dose-effects on the lipid component of serum. MALDI-oa-ToF profiling was applied for the first time to search for radiation-induced changes in human serum lipidome. The positive mode of MALDI ionization was selected in order to favor the analysis of choline-based compounds and other phospholipids, which already have been proposed as potential markers of the response to radiation and anti-cancer treatment [8,10].

2. Results

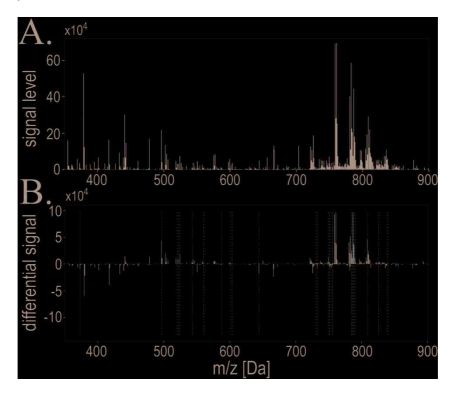
2.1. Exposure to Radiation during Radiotherapy Induced Changes in the Serum Lipidome Profiles

In the analyzed mass range 350–900 Da 842 spectral components (i.e., lipid species with their isotope variants) common for all mass profiles were detected (an average mass profile is presented on Figure 1A). In order to find radiotherapy-related changes individual differential spectra were computed paired with respect to consecutive time points (i.e., changes A Δ B, B Δ C and A Δ C), and then the statistical significance of differences in component's abundances was estimated (Figure 1B shows resulting differential spectra). Several spectral components changed their abundances significantly between compared time points (FDR < 5% was selected as a statistical significance threshold), which are listed in Table 1 (complete data regarding all registered components are presented in Supplementary Table S1). We observed that major changes occurred between pre-treatment and within-treatment samples (the $A\Delta B$ change), where 27 spectral components (lipid species) changed their abundance with high level of statistical significance (FDR < 5%). When within-treatment samples were compared with post-treatment samples (the BΔC change), 14 spectral components showed significantly changed abundance. However, abundances of only three spectral components remained different at high level of statistical significance when pre-treatment and post-treatment samples were compared (the $A\Delta C$ change). Of note, we observed that seven spectral components significantly differentiated samples B from both samples A and samples C (registered m/z values = 520.36, 522.39, 603.68, 749.51, 760.63, 786.64 and 788.65 Da). Moreover, one spectral component (m/z value = 751.47 Da) differentiated samples A both from samples B and samples C. Half of the differentiating components were identified with respect to their lipid class (see Table 1), almost all of them being phospholipids containing the choline "head": phosphatidylcholines (PC; 10 compounds), lysophosphatidylcholines (LPC; 4 compounds) and sphingomyelines (SM; 2 compounds).

Table 1. Spectral components that changed abundances significantly between analyzed time points. Shown here is the registered m/z value, significant change in abundance, real pattern of changes, cluster number (hypothetical pattern of changes) and identification (lipid class and length of fatty acyl chains) of analyzed spectral components (*i.e.*, lipid species); components of isotopic envelope were excluded from analysis.

Ion mass[w/z]	Significant change	Pattern of	Cluster	Linid aloss identification
Ion mass[m/z]	(FDR < 5%)	changes	number	Lipid class identification
373.08	$A\Delta B$	A < B > C	#4	not assigned
496.36	$A\Delta B$	A > B < C	#2	$LPC(16:0) + H^{+}$
520.36	ΑΔΒ;ΒΔС	A > B < C	#2	$LPC(18:2) + H^{+}$
522.39	ΑΔΒ;ΒΔС	A > B < C	#2	$LPC(18:1) + H^{+}$
524.38	$A\Delta B$	A > B < C	#2	$LPC(18:0) + H^{+}$
543.39	$A\Delta B$	A > B = C	#2	not assigned
560.28	$A\Delta B$	A > B = C	#1	[Vitamin D3 adduct] + H ⁺
564.64	$B\Delta C$	A = B > C	#4	$Cer(36:2) + H^+$
587.33	$A\Delta B$	A < B > C	#4	not assigned
601.12	$A\Delta B$	A < B > C	#4	not assigned
603.68	ΑΔΒ;ΒΔС	A < B < C	#6	not assigned
644.11	$A\Delta B$	A < B > C	#4	not assigned
703.58	$B\Delta C$	A = B < C	#6	$SM(34:1) + H^+$
721.49	ВДС	A > B < C	#2	not assigned
726.53	$A\Delta C$	A = B < C	#6	not assigned
730.62	$A\Delta B$	A < B = C	#3	$PC(32:2) + H^+$
732.47	ВДС	A > B < C	#2	not assigned
732.63	$A\Delta B$	A < B = C	#3	$PC(32:1) + H^{+}$
749.51	ΑΔΒ;ΒΔС	A > B < C	#2	not assigned
751.47	ΑΔΒ;ΑΔС	A > B = C	#1	not assigned
751.61	$A\Delta B$	A < B = C	#4	not assigned
755.42	$A\Delta B$	A > B = C	#1	not assigned
755.63	$A\Delta B$	A < B = C	#3	$SM(38:3) + H^+$
758.61	ВДС	A > B < C	#2	$PC(34:2) + H^{+}$
760.63	ΑΔΒ;ΒΔС	A > B < C	#2	$PC(34:1) + H^{+}$
762.63	$B\Delta C$	A > B < C	#2	$PC(34:0) + H^{+}$
767.47	ВДС	A > B < C	#2	not assigned
777.33	$A\Delta C$	A = B > C	#5	not assigned
784.62	$A\Delta B$	A > B < C	#2	$PC(36:3) + H^+$
786.64	ΑΔΒ;ΒΔС	A > B < C	#2	$PC(36:2) + H^{+}$
786.94	$A\Delta B$	A > B < C	#2	not assigned
788.65	ΑΔΒ;ΒΔС	A > B < C	#2	$PC(36:1) + H^{+}$
790.65	ΑΔΒ	A > B < C	#2	$PC(36:0) + H^{+}$
808.62	$A\Delta B$	A > B < C	#2	$PC(38:5) + H^{+}$
825.58	$A\Delta B$	A > B = C	#2	not assigned
839.50	$A\Delta B$	A > B = C	#4	not assigned

Figure 1. Mass profiles of serum lipids were affected during radiotherapy. (Panel **A**): Averaged mass spectrum of serum lipids registered in the 350–900 Da range for pre-treatment samples (**A**); (Panel **B**): Averaged differential spectrum for pre-treatment and within-treatment samples (A Δ B); components that changed their abundances significantly (FDR < 5%) are marked with red lines.

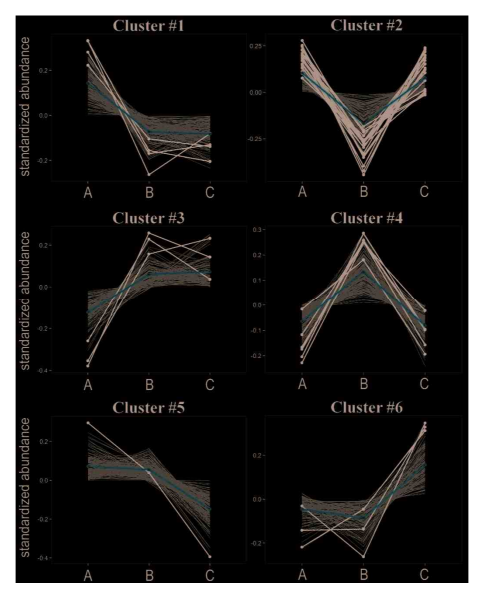


2.2. Radiotherapy-Related Changes in Lipidome Profiles Showed Different Time-Course Patterns

To identify different patterns of radiotherapy-related changes an unsupervised cluster analysis was performed. We identified six different hypothetical patterns of changes (clusters) characterized in Table 2 and depicted in Figure 2 (in case of a few spectral components where differences between time points were not statistically significant observed patterns of changes were not strictly coherent with cluster characteristics); detailed data are presented in Supplementary Table S2. Identified clusters could be further divided into three groups with two "mirrored" clusters in each, where reverse changes were observed: #1 [A > B = C] and #3 [A < B = C], #2 [A > B < C] and #4[A < B > C], #5 [A = B > C] and #6 [A = B < C]. Of note, the second group (i.e., clusters #2 and #4) where "earlier" changes (A Δ B) were compensated by "later" changes (B Δ C) was the most numerous (about 70% of all detected components). Furthermore, cluster #2 [A > B < C] contained the majority of differentiating components, that changed abundances significantly between consecutive time points (19 out of 36 "significant" components, see Table 1). This indicated that pattern of changes where "earlier" changes were reversed/compensated by "later" changes was the most common feature of lipidome profiles in serum from irradiated patients. Of note, the majority of "differentiating" LPCs and PCs belonged to cluster #2, and their serum levels decreased significantly during radiotherapy and then increased afterwards; these included LPC(18:2), LPC(18:1), PC(34:1), PC(36:2) and

PC(36:1) (Figure 3). On the other hand PCs containing 32 carbons (32:2 and 32:1) and SM(38:3) significantly increased their levels during radiotherapy (cluster #3) (see Table 1).

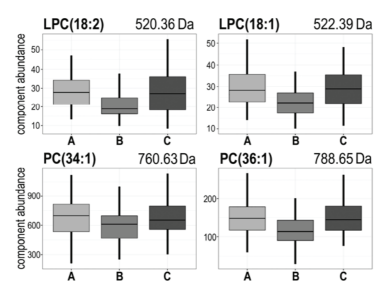
Figure 2. Radiation induced changes followed different patterns. Presented are characteristics of six identified clusters of components with similar time-courses of changes; marked are average profiles for each cluster (red lines) and components that changed abundance significantly (FDR < 5%; solid black lines); other components belonging to each cluster are marked with grey lines.



Cluster	Pattern of change *	Number of components	Number of differentiating components **
#1	A > B = C	147	4
#2	A > B < C	129	19
#3	A < B = C	121	3
#4	A < B > C	170	6
#5	A = B > C	160	1
#6	A = B < C	115	3

Table 2. Characteristics of identified clusters of spectral components.

Figure 3. The abundance of several choline-containing phospholipids decreased markedly during radiotherapy and increased afterward. Presented are examples of lysophosphatidylcholines: LPC(18:2) [m/z = 520.36 Da] and LPC(18:1) [m/z = 522.39 Da], and phosphatidylcholines: PC(34:1) [m/z = 760.63 Da] and PC(36:1) [m/z = 788.65 Da]. Boxplots show: minimum, lower quartile, median, upper quartile and maximum values (outliers were removed from the plots for perspicuity).



2.3. Radiotherapy-Related Changes in Serum Lipidome Were Associated with Doses of Radiation and Volumes of Irradiated Tissues

In the next step of the study we searched for association between features of serum lipidome (*i.e.*, changes in abundance of particular lipidome components) and doses of ionizing radiation received by patients (doses accumulated until a time point corresponding to the collection of sample B in case of the A Δ B changes and total doses in case of the A Δ C and B Δ C changes). Correlations were identified between specific features of the serum lipidome and either the total (maximum) dose received by gross tumor volume (GTV), volume of the patient's body irradiated at different (smaller) doses or dose delivered to different volume of tissue. Numbers of serum lipidome components, for which changes in abundances correlated with maximum GTV doses are shown in Table 3 (p < 0.05

^{*} Pattern of change refers to the dominant characteristics of change in the specified cluster (with some not significant deviations from the pattern within the cluster); ** Components which abundances changed significantly between consecutive time points (FDR < 5%).

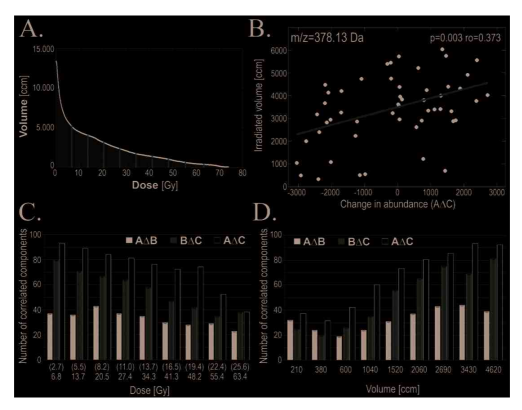
was selected as a statistical significance threshold). We found the highest number of identified correlations was observed in case of AΔC changes (60), yet clear association between maximum GTV doses and features of lipidome were detected also for the AAB and BAC changes (44 and 37 components, respectively). The maximum radiation doses (up to 72 Gy) were delivered only to tumor and its adjacent margins (usually 100-200 ccm), while tissue irradiated with lower doses represent much higher volumes (e.g., about 4000 ccm irradiated with 10 Gy). Hence, we searched for correlations between features of serum lipidome and volumes of tissues (including normal tissues irradiated upon IMRT treatment) irradiated with different doses (including "low" or "clinically irrelevant" doses); see Figure 4A,B. Figure 4C shows the numbers of lipidome components, which changes in abundance correlated with volume of tissue irradiated at different doses (p = 0.05 was selected as statistical significance threshold). We observed that association between lipidome features and dose-volume effects were the most frequent in case of larger tissue volumes irradiated with clinically low-to-medium doses (i.e., less than 20 Gy in case of the A Δ B change and less than 40 Gy in case of the B Δ C and A Δ C changes; which corresponded to dose fractions below 1 Gy). Additionally, when reverse analysis was performed and serum lipidome features were correlated with doses delivered to a given volume similar results were obtained—majority of correlations were observed for high volumes irradiated with low doses (Figure 4D). Detailed data on correlation of serum lipidome features with volumes irradiated at a given dose or with doses delivered to a given volume are presented in Supplementary Tables S3–S5. Our results clearly indicated that radiotherapy-related changes in serum lipidome profiles depended on doses of delivered radiation, and that low-to-medium doses delivered to large volumes of normal tissue could affect observed changes.

Table 3. Serum lipidome features associated with radiation doses and acute radiation toxicity.

Change	GTV-D	AMR	Examples of components [m/z] *
$A\Delta B$	44	36	473.11; 514.21; 590.61; 872.42
$B\Delta C$	37	41	583.61; 669.64
$A\Delta C$	60	35	614.38; 641.33; 649.43; 655.65; 673.62; 765.64; 803.71; 886.88

^{*} Components for which abundances correlated with both doses of radiation and radiation toxicity (p < 0.05).

Figure 4. Dose-volume effects in serum lipidome changes. (Panel **A**): Averaged Dose-Volume Histogram; doses corresponding to deciles of the area under curve of the histogram are marked with red lines; (Panel **B**): Correlations between volume of tissue irradiated with 13.7 Gy and changes in abundance of the m/z = 378.13 Da component in pre- and post-treatment samples (C-A; arbitrary units); Numbers of serum lipidome features correlating with tissue volumes irradiated at a given dose of radiation (Panel **C**) or doses of radiation delivered to a given tissue volume (Panel **D**). Shown here are the AΔB (black bars), BΔC (grey bars) and AΔC (empty bars) changes; p = 0.05 was selected as a statistical significance threshold (doses in parentheses refer to AΔB changes).



2.4. Radiotherapy-Related Changes in Serum Lipidome Were Associated with Radiation Toxicity

Finally, we searched for potential association of serum lipidome features with toxicity of the treatment. The clinically relevant response of normal tissues to toxicity of radiation was assessed using a modified Dische system [22] relying on the intensity of the acute mucosal reaction. The maximum AMR intensity correlates with both maximum GTV dose and volume of normal tissues irradiated with "intermediate" doses (about 0.8–1 Gy per fraction), which was documented in another study based on very similar group of head and neck cancer patients [21]. Here we searched for association between the early radiation toxicity and radiation-induced changes of serum lipidome features. We found correlation between changes in abundance of several lipidome components and the maximum AMR intensity: about 40 lipidome features correlated with the AMR for each of analyzed time-courses (Table 3). Of note, several serum lipidome features associated with the maximum AMR intensity also correlated with volumes of tissues irradiated at given doses of radiation (or radiation doses delivered to a given tissue volume). These features are listed in Table 3. Hence, we concluded

that radiotherapy-related changes in the serum lipidome were associated with dose-related toxicity of radiation.

3. Discussion

To our knowledge, this is the first paper to analyze the response of human organism to ionizing radiation due to local cancer irradiation performed at the level of blood lipidome. The main changes in abundances of lipid serum components were observed between pre-treatment samples and samples collected during radiotherapy (the AAB changes). Unsupervised cluster analysis revealed that major group of lipids (70% of registered spectral components) consisted of species, for which radiation-induced changes observed during radiotherapy were reversed/compensated in the post-treatment samples collected 1–2 months after the end of radiotherapy. A minor group of lipids (20% of registered spectral components) consisted of species, where radiation-induced changes detected during radiotherapy remained not reversed/compensated during the follow-up. As a consequence, only a few lipid species showed significant differences when their pre-exposure and post-exposure levels were compared (the $A\Delta C$ changes). This observation indicated that in case of majority of serum lipids their return to the initial pre-exposure steady-state level was efficient enough during 1–2 months after the end of radiation treatment. Of note, when radiotherapy-related changes in serum proteome profiles were analyzed in a very similar group of patients, the major changes were observed in post-treatment samples collected 1–2 months after the end of radiotherapy (corresponding to the A Δ C and B Δ C changes). Such serum proteome changes apparently reflected escalation of radiation toxicity (acute mucosal reaction) and its subsequent healing during the follow-up [21]. Here, we show that radiotherapy-related changes in serum lipidome profiles are apparently "faster" compared to changes observed in the low-molecular-weight fraction of serum proteome. In fact, most radiation-induced changes in serum lipidome could be reversed within 1–2 months after completion of radiotherapy, while similar changes in serum proteome could be detected several months after the treatment. This indicated that changes in lipidome and proteome profiles observed in cancer patients treated with radiotherapy might reflect different radiation-induced mechanisms.

Lipid class identification (by MS/MS and/or annotation of registered *m/z* values at LipidMaps database) allowed annotation of the majority (85%) of lipids revealing radiation-induced changes as choline-based phospholipids. High extent of lipids referring to this type apparently resulted from both chosen conditions of serum extraction, which favored glycerophospholipids, sphingolipids, sterols and prenols [2], and positive mode of MALDI ionization, which narrowed the ionization of glycerophospholipids to neutral (zwitterionic) representatives, such as phosphatidylocholines and phosphatidylethanolamines [23]. The majority of PLs identified in this work, including different PCs and LPCs as well as SM(34:1), belong to the most abundant in their classes measured in human plasma samples [2]; SM(38:3) and Cer(36:2) are less common species. Phosphatidylocholines are the main building blocks of membrane bilayers and in plasma they are mostly located in high density lipoproteins (HDL). Decreased levels of PCs in serum of irradiated patients may be explained by their rapid turnover in stressed/damaged cells, which resulted in an increased PC's uptake from the blood. In addition to their main function as a membrane constituent, PCs have a role in signaling

through the generation of LPCs (by phospholipase A₂ enzymes), SMs (by SM synthase), phosphatidic acid (PA; by phospholipase D enzymes) and/or diacylglycerols (DAG; by phospholipase C enzymes). From this point of view, significant down-regulation of major serum PCs observed during radiotherapy might be relevant for increasing capability of cell signaling pathways depending on PC-derived compounds. LPCs are reported to be the major bioactive lipid component of oxidized low density lipoproteins (LDL) and therefore mainly responsible for their pro-inflammatory functions [24]. Down-regulation of LPCs in blood was significantly correlated with activated inflammatory status in many cancer types [25]. Radiotherapy-related down-regulation of LPCs apparently indicated association between inflammatory processes and whole body response to radiation, which was previously documented at the level of serum proteome [21]. Another important class of signaling lipids derived from PCs and LPCs upon action of phospholipase D enzymes are lysophosphatidic acids (LPAs). The most prominent LPA functions include stimulation of cell proliferation, cell survival, and tumor cell invasion [26]. Down regulation of both PCs and LPCs may be therefore explained by the increased formation of LPAs. Another potential mechanism explaining down-regulation of PCs and LPCs involves the disruptive action of reactive oxygen species (ROS) appearing in high levels in irradiated tissues and causing the degradation of these lipids [27]. In contrast to PCs and LPCs, which indicated decreased levels during radiotherapy and were compensated during the follow-up, both identified sphingomyelines showed significant radiation-related up-regulation only: SM(38:3) during earlier stage of the treatment while SM(34:1) during later stage of the treatment or subsequent follow-up. New SM molecules were most probably generated from degraded PC compounds by SM synthase (this transferase utilizes a choline "head" from PC) and suitable ceramide molecules, which was coherent with observed down-regulation of Cer(36:2). SMs can be hydrolyzed back to ceramides by SMase action. This balance between sphingomyeline production and degradation is a key factor in SM-related apoptotic signaling, and generation of ceramides from SMs' degradation was reported to influence both the rate and form of cell death [28].

Although the model presented here is rather complicated and could be affected by many different processes ongoing in the patient's organism, one could expect that accumulation and subsequent healing of radiation-induced damage, such as acute mucosal reaction, would have the major influence on general therapy-related changes assessed at the level of serum lipidome. This expectation is supported by observed association of radiotherapy-related serum lipidome features with doses of radiation delivered to normal tissues and intensity of radiation-induced acute mucosal reaction. Although correlations identified here between particular lipidome components and different parameters reflecting radiation doses and toxicity possess moderate statistical power when analyzed separately, reliable conclusions could be drawn based on the general patterns of observed association. Of note, collected data indicated that low-to-medium doses delivered to large volumes of normal tissues during IMRT (considered as "therapeutically irrelevant") significantly affected whole body response observed at the level of serum lipidome. These observations are consistent with results of our earlier study, where similar association between dose-volume effects and features of the low-molecular-weight fraction of serum proteome has been observed in similar group of head and neck cancer patients [21]. The data indicated collectively, that a whole body response to the local cancer irradiation could be detected at the level of both serum proteome and lipidome. However, the

majority of radiation-induced changes in abundances of serum lipids returned to pre-exposure steady-state levels within a relatively short time after the treatment, while changes in serum proteome could be detected even several months after irradiation.

4. Experimental Section

4.1. Characteristics of the Patients

Sixty-six patients with head and neck squamous cell carcinoma (HNSCC) were enrolled in this study. All of them were Caucasians (64 men) 45–82 years old (median age 63 years); 82% of them were current smokers and 86% alcohol consumers. Cancer was located mainly in larynx (45 pts.), but also in oropharynx (15 pts.) or hypopharynx (6 pts.). The primary tumor stage was scored as: T1 (21%), T2 (44%), T3 (26%) and T4 (9%); 68% of N0. All patients were subjected to IMRT using 6 MeV photons with 1.8 Gy daily fraction doses according to the continuous accelerated irradiation scheme (CAIR) [29]). Total radiation doses delivered to gross tumor volume (GTV) were in the range of 61.2–72 Gy (median 66.6 Gy). Neither surgery nor induction/concomitant chemotherapy was applied to patients enrolled in this study. Three consecutive blood samples (5 mL) were collected from each patient: pre-treatment sample A (within one week before RT; 66 pts.); within-treatment sample B (10–22 days after the start of RT, median 15 days; 66 pts.) and post-treatment sample C (23–59 days after the end of RT, median 36 days; 56 pts.). The acute mucosal reaction (AMR) was estimated using the modified Dische score system [22] every 3–5 days during the radiotherapy. The study was approved by the appropriate Ethics Committee and all participants provided informed consent indicating their conscious and voluntary participation.

4.2. Preparation of the Samples

Blood was collected into a 5 mL Vacutainer Tube (Becton Dickinson, Franklin Lakes, NJ, USA), incubated for 30 min RT to allow clotting, and then centrifuged at $1000 \times g$ for 10 min to remove the clot. The serum was aliquoted and stored at -70 °C until extraction. Total lipids were extracted according to modified Folch method [30]. In brief, 25 μ L of serum was mixed with 350 μ L of 1:1 methanol/chloroform mixture (v/v) containing antioxidants: 0.01% (w/v) 2,6-di-*tert*-butyl-4-methylphenol and 0.005% (w/v) retinol. The solution was vortex-mixed for 0.5 min and incubated for 30 min at 20 °C. Then 80 μ L of water was added to the mixture, vortex-mixed for another 0.5 min and centrifuged (5 min, $10,000 \times g$). Chloroform phase (the lower one) was kept and stored at -70 °C until mass spectrometry analysis (within three weeks).

4.3. MALDI Mass Spectrometry Analysis

Samples was analyzed using MALDI quadrupole/orthogonal acceleration ToF (oa-ToF) high-definition MS (HDMS) SYNAPT G2-HDMSTM system (Waters, Manchester, UK) equipped with the 355 nm Nd:YAG laser. First, 0.5 μ L of sample was mixed directly on stainless steel target plate with 0.5 μ L of 30 mg/mL of α -cyano-4-hydroxycinnamic acid (CHCA) matrix (Bruker Daltonics, Billerica, MA, USA) dissolved in 70% (ν / ν) acetonitrile containing 0.1% (ν / ν)

trifluoroacetic acid; each sample was analyzed in triplicate (*i.e.*, placed on three different spots). Mass spectra were recorded using the positive ion mode in the 350–900 Da range with resolution of 10,000 FWHM. Spectra were calibrated with a standard solution of polyethylene glycol (PEG), and m/z scales were adjusted after acquisition using the PEG signal at m/z 701.3935 as a lock mass and centroided prior to the generation of accurate mass peak lists. Samples were spotted and analyzed in a random sequence to avoid "batch effect".

4.4. Processing of the Mass Spectra

The initial preprocessing of spectra including alignment, detection of outlier profiles (using Dixon's Q test), averaging of three technical replicas, additional alignment of averaged individual spectra (*i.e.*, averaged technical replicas), baseline removal and normalization of the total ion current (TIC) was performed according to procedures considering to be standard in the mass spectrometry field [31]. Preprocessed spectra were transferred to Spectrolyzer software suite (v.1.0, MedicWave AB, Halmstad, Sweden; [32]) for peak detection and binning (peak clustering) analysis. The processing steps performed in Spectrolyzer software were also consistent with the standard procedures used for spectral data processing [33,34].

4.5. Testing for Differentiating Spectral Components

For each of the 842 spectral components (spectral peaks) statistical significance of differences in abundance between different time points (*i.e.*, samples A, B and C) was estimated using appropriate tools available in R statistical software (see [35]). Individual differential spectra were computed paired with respect to consecutive time points (A–B, B–C and A–C), which resulted in 66 samples for comparison of A vs. B, and 56 samples for comparison A vs. C and B vs. C. To verify whether observed differences in abundances were significant, the Wilcoxon signed rank test was used (with the null hypothesis that the median value of intensities in the differential spectrum is equal to zero). To account for multiple comparisons the Storeys approach [36] that allows for FDR (false discovery rate) control was used. Statistically significant components involved also features that were identified manually as isotopes of other compounds; these components were rejected from the final list of specific components intended for identification.

4.6. Identification of Differentiating Components

Spectral components showing significant differences between analyzed time points (FDR < 5%) were analyzed by MS/MS in order to identify lipid class and length of fatty acyl chains. PCs and SM classes were recognized in MS/MS based on characteristic 184.1 Da phosphocholine fragmentation ion, while LPCs based on both 184.1 Da phosphocholine and 104.1 choline fragmentation ions Additionally, other spectral components were annotated at the LipidMaps database [37] based on their registered m/z values; mass tolerance 0.1 Da and no limit for category/class was applied. Compounds that were not identified experimentally (due to the low abundance of precursor or productions) were regarded as identified only if a single unique lipid record was return from the database search.

4.7. Analysis of Patterns of Changes

To investigate the general patterns of changes in abundances of spectral components between compared time points averaged "time courses" were computed based on individual time courses. Data standardization (centering and scaling separately for each of the spectral component) was performed to account for wide differences in abundance ranges observed for distinct components. Cluster analysis was performed using Partitioning Around Medoids (PAM) method, which is a classical algorithm of unsupervised analysis widely used for similar problems [38]. For a given number of clusters (k) the PAM finds k representative objects (so called medoids) that are most different from each other and assigns all the remaining objects to the most similar of the representatives. The similarity of the objects being an input for the PAM was computed based on correlation between average time courses. In order to determine the optimal k number and assess the quality of clustering results, an average Silhouette Width (SW) criterion was used [39], which revealed a six-cluster solution as the optimum.

4.8. Correlation of Component's Abundance with Radiation Parameters

Correlations between changes in abundance of spectral components and parameters reflecting absorbed doses of radiation, as well as maximal intensities of AMR, were analyzed using the Spearman's rank correlation coefficient. Total radiation dose absorbed by patient's body was estimated from the individual dose-volume histogram generated during the treatment planning. For the analysis of dose/volume effect we selected the doses corresponding to deciles of the area under the curve of the averaged dose-volume histogram (for details see [21]).

5. Conclusions

This study demonstrates for the first time the massive involvement of choline-based lipid serum components in the response of humans to ionizing radiation. Significant change in LPCs' levels suggests activation of inflammatory processes, while disturbances in levels of sphingomyelines and ceramides indicate involvement of apoptotic pathways. Additionally, correlations of lipidome changes with low and moderate radiation doses call attention to the biological relevance of "therapeutically irrelevant" doses during IMRT.

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Author Contributions

KJ—Performed and interpreted experiments, identified lipid class and length of fatty acyl chains of statistically significant lipid components, prepared the final manuscript, MP—Interpreted MS

results, MR—Interpreted MS results, AZ—Performed testing for differentiating spectral components, AS—Performed analysis of patterns of changes, JP—Performed analyses of correlation of component's abundance with radiation parameters, MM—Performed analyses of correlation of component's abundance with radiation parameters, TR—Collected and interpreted clinical data, KS—Collected and interpreted clinical data, MRC—Interpreted MS results, PW—Designed the project, designed and interpreted experiments, prepared the final manuscript. All authors read and approved the final manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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Radiosensitization of Human Leukemic HL-60 Cells by ATR Kinase Inhibitor (VE-821): Phosphoproteomic Analysis

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Abstract: DNA damaging agents such as ionizing radiation or chemotherapy are frequently used in oncology. DNA damage response (DDR)—triggered by radiation-induced double strand breaks—is orchestrated mainly by three Phosphatidylinositol 3-kinase-related kinases (PIKKs): Ataxia teleangiectasia mutated (ATM), DNA-dependent protein kinase (DNA-PK) and ATM and Rad3-related kinase (ATR). Their activation promotes cell-cycle arrest and facilitates DNA damage repair, resulting in radioresistance. Recently developed specific ATR inhibitor, VE-821 (3-amino-6-(4-(methylsulfonyl)phenyl)-N-phenylpyrazine-2-carboxamide), has been reported to have a significant radio- and chemo-sensitizing effect delimited to cancer cells (largely p53-deficient) without affecting normal cells. In this study, we employed SILAC-based quantitative phosphoproteomics to describe the mechanism of the radiosensitizing effect of VE-821 in human promyelocytic leukemic cells HL-60 (p53-negative). Hydrophilic interaction liquid chromatography (HILIC)-prefractionation with TiO₂-enrichment and nano-liquid chromatography—tandem mass spectrometry (LC-MS/MS) analysis revealed 9834 phosphorylation sites. Proteins with differentially up-/down-regulated phosphorylation were mostly localized in the nucleus and were involved in cellular processes such as DDR, all phases of the cell cycle, and cell division. Moreover, sequence motif analysis revealed significant changes in the activities of kinases involved in these processes. Taken together, our data indicates that ATR kinase has multiple roles in response to DNA damage throughout the cell cycle and that its inhibitor VE-821 is a potent radiosensitizing agent for p53-negative HL-60 cells.

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1. Introduction

One of the treatment modalities in oncology is radiotherapy. It often employs chemical agents increasing sensitivity towards ionizing radiation (IR). IR induces the most deleterious lesions of DNA, double strand breaks (DSB), and their repair is regulated by ataxia telangiectasia-mutated kinase (ATM), DNA-dependent protein kinase (DNA-PK), and ATM and Rad3-related kinase (ATR). While activation of ATM and DNA-PK is triggered by DNA double stranded breaks, ATR kinase responds to a broad spectrum of agents inducing single stranded DNA [1]. ATR acts primarily in S- and G2-phases and responds to replication and genotoxic stress, however, a recent report has shown that ATR is activated also in irradiated G1 phase cells [2].

In 2011, a specific inhibitor of ATR, VE-821 (3-amino-6-(4-(methylsulfonyl)phenyl)-N-phenylpyrazine-2-carboxamide), was developed [3]. Its excellent selectivity in regards to sensitization of cancer cells towards various types of DNA damaging agents leaving healthy cells

unaffected has been recently reported [4–7]. VE-821 selectivity is based on the concept that (i) more than 50% of cancer cells have lost their G1-phase checkpoint for example due to p53 mutation/deletion and thus rely on S- and G2-checkpoints; which are known to be regulated by ATR and (ii) cancer cells with activated oncogenes generate replication stress at much higher levels than normal cells, thus activating ATR [8,9]. This hypothesis is based on well-established "classical" methodology in molecular biology comprising western-blotting, confocal microscopy, flow-cytometry and so forth. However, this approach is often focused on one or two particular mechanisms not reflecting the complexity of cellular signaling pathways. Therefore it fails to give a comprehensive view of the mechanisms of radiosensitization by ATR inhibition in the context of cancer cells.

In the DNA damage response as well as in the other molecular processes (such as transcriptional and translational regulation, proliferation, differentiation, apoptosis, cell survival and many others) phosphorylation frequently initiates and propagates signal transduction pathways [10]. Therefore we decided to apply a phosphoproteomic approach in order to study the effect of specific inhibition of ATR by the small molecule inhibitor VE-821.

Our goal was to describe the changes in the phosphoproteome in radiosensitized tumor cells lacking functional protein p53. Although mass spectrometry of phosphopeptides obtained from tryptic protein digests has become a powerful tool for characterization of phosphoproteins involved in cellular processes, there is an inevitable part of the protocol: phosphopeptide enrichment. It compensates the low abundance, insufficient ionization, and suppression effects of non-phosphorylated peptides [11]. Hence, we optimized the metal oxide affinity chromatography (MOAC) enrichment using titanium dioxide and employed it in our experiments [12].

We have previously compared the effects of inhibitors of ATM (KU55933) and ATR kinases (VE-821) on the radiosensitization of human promyelocytic leukemia cells (HL-60), lacking functional protein p53. The inhibition of ATR by its specific inhibitor VE-821 resulted in a more pronounced radiosensitizing effect in HL-60 cells compared to the inhibition of ATM. In contrast to KU55933, the VE-821 treatment prevented HL-60 cells from undergoing G2 cell cycle arrest [13].

In this paper we characterize the radiosensitizing effect of VE-821 from a new perspective and we report the mechanisms and signaling pathways involved in the processes triggered by IR in p53-negative leukemic cells.

2. Results

Our goal was to describe the changes in the phosphoproteome in radiosensitized tumor cells, since in the DNA damage response (as well as in other molecular processes) phosphorylation frequently initiates and propagates signal transduction pathways. We aimed to characterize the effect of specific inhibition of ATR kinase by VE-821 and to report the mechanisms and signaling pathways involved in the processes triggered by IR in p53-negative human promyelocytic leukemic cells HL-60 that are ATR-dependent. To reach our goal, we employed SILAC-based quantitative phosphoproteomics together with metal oxide affinity chromatography using TiO₂ microparticles to specifically enrich for phosphorylated peptides. To increase the number of identified and quantified phosphorylation sites, peptides were prefractionated by hydrophilic interaction liquid chromatography (HILIC) chromatography prior to the enrichment. Additionally, to make sure that the changes in the

phosphoproteome are not based on changes of protein abundance caused by alterations in gene expression or protein degradation that could occur because of the relatively long incubation time (1 h after irradiation, and 1.5 h after VE-821 treatment), we also analyzed a non-phosphorylated complement of phosphorylated peptides obtained from flow-through fractions of the phosphopeptide enrichment.

2.1. Overall Phosphoproteomic Analysis Reveals Thousands of Phosphorylation Sites

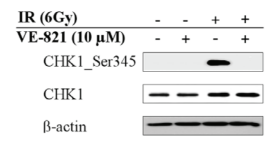
In summary, we identified 9834 phosphorylation sites from 3210 protein groups, among them 4809 were quantified in all three experimental replicates. The phosphosite ratios were shown to correlate well between the replicates (Pearson correlation coefficient was between 0.8 and 0.87). Moreover, the correlation between the normalized H/L ratios of phosphopeptides and the normalized protein H/L ratios was also calculated (the calculation was based on 1941 phosphorylation sites from 557 proteins), and the low correlation coefficient value (R = 0.0162) indicated that the quantified changes in the phosphoproteome were presumably not caused by changes in the protein abundance. The overview of all identified phosphorylation sites is given in the supplementary (Table S1).

To determine whether a phosphorylation site exhibits an appropriate alteration in abundance in all of the replicates to be considered as differentially regulated after the treatment, we employed the Global rank test (GRT) with the non-parametric estimate of the false discovery rate (FDR; see material and methods). At the 1% FDR level, 336 phosphorylation sites (260 proteins) were evaluated as differentially up-regulated, whereas 202 phosphorylation sites (168 proteins) were differentially down-regulated (some of the proteins contained both down- and up-regulated phosphorylation sites). Since a kinase inhibitor was applied to modulate the response to IR induced DNA damage, the number of down-regulated sites would be expected to be larger than the number of up-regulated sites. However, the number of up-regulated sites increased and possibly reflected relatively long time period that passed from the initial treatment, and allowed the cells to eventually trigger a compensatory phosphorylation response in order to overcome the inhibitory effect of VE-821.

2.2. Combination of Ionizing Radiation and VE-821 Treatment Abrogated Phosphorylation of Checkpoint Kinase-1 on Ser345

In order to confirm the inhibitory effect of VE-821 on ATR in our model, we assessed phosphorylation of checkpoint kinase-1 (CHK1) on Ser345. Since it reports active and functional ATR, this particular direct downstream target of ATR was found to be phosphorylated upon irradiation. As shown in Figure 1, CHK1 phosphorylation was abrogated after VE-821 pre-incubation.

Figure 1. In order to confirm the inhibitory effect of VE-821 on ATM and Rad3-related kinase (ATR) we assessed phosphorylation of CHK1 on Ser345 (a direct downstream target of ATR) and we found this particular phosphorylation to be inhibited. Beta-actin was used as a gel loading control. The representative blots from at least three independent experiments are shown.



2.3. Most of the Regulated Phosphoproteins Were Localized in the Nucleus and Related to Mitosis, Cell Cycle Regulation, DNA Damage Response and Gene Expression

To functionally and spatially interpret the list of 396 phosphorylated proteins resulting from the GRT, Gene Ontology (GO) annotation was performed using the ConsensusPathDB, over-representation analysis "web tool" [14,15] and the summary of the results is given in Figure 2. The analysis of cellular component GO terms (level 4) revealed that most of the proteins were localized in the nucleus, many of them included in chromatin, localized at the centromeric region of a chromosome, replication fork or as a part of histone acetyltransferase or deacetylase complexes. Nevertheless, there was also a considerable fraction of proteins annotated to be a part of cytosol. The most over-represented molecular functions (level 4) of regulated phosphoproteins were binding of both nucleic acids (RNA binding and DNA binding), transcriptional co-activator and co-repressor activities, as well as DNA ligase or topoisomerase II activities. The list of overrepresented biological process terms (level 4) contained processes related to mitosis (nuclear envelope disassembly, sister chromatid cohesion, spindle localization or cytokinesis), cell cycle regulation (cell cycle progression, cell cycle phase transition, cell cycle checkpoint, regulation of transcription involved in G1/S transition of the mitotic cell cycle, cell cycle DNA replication), cellular response to DNA damage stimulus (base- and nucleotide-excision repair, double-strand break repair) or gene expression (mRNA transport, histone acetylation).

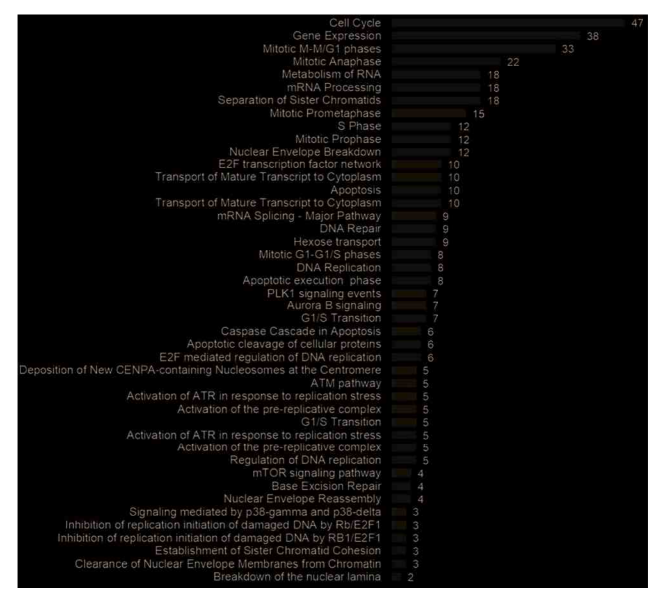
Figure 2. Selected results of Gene Ontology terms over-representation analysis (FDR < 0.01): (**A**) Overrepresented Gene Ontology (GO) Biological process level 4 terms; (**B**) Overrepresented GO Cellular component level 4 terms; (**C**) Overrepresented GO Molecular function level 4 terms. *x*-axis contains the number of proteins involved in a particular pathway that were found differentially phosphorylated in our study.



2.4. The Pathways Affected by Co-Treatment Were Involved in Cell Cycle Progression, Stimulus-Based Changes in Gene Expression, DDR and Apoptosis

To assess which signaling pathways contained differentially phosphorylated proteins and thus could be considered as triggered or modulated by ATR inhibition, we performed a pathway over-representation analysis using the same web-tool as in the case of GO enrichment, however, we addressed the signaling pathways stored in the Reactome pathway database [16] and Pathway Interaction Database (PID; [17]). The overrepresented pathways are shown in Figure 3. It is apparent that most of the pathways were related to different phases of the cell cycle and transitions between them, stimulus-based changes in gene expression, DNA damage repair and DNA damage-induced programmed cell death.

Figure 3. Selected results of Reactome pathways (blue bars) and Pathway Interaction Database (PID) pathways (orange bars) over-representation analysis (FDR < 0.01): x-axis shows the number of proteins involved in particular pathways that were differentially phosphorylated in our study.



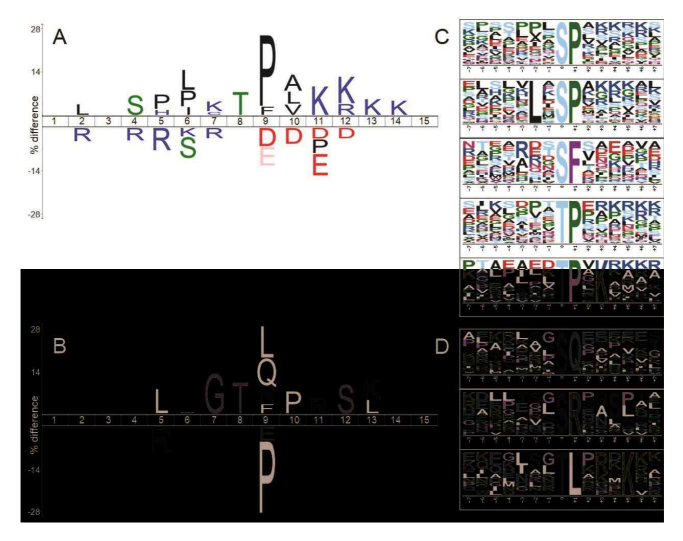
2.5. Significantly Down-Regulated Phosphorylation Sites Comprised SP/TP Motifs, while SQ/TQ Were amongst Up-Regulated Ones

To analyze and visualize sequence motifs that were enriched among significantly regulated class I phosphorylation sites (see material and methods), we utilized the iceLogo tool [18] and the motif-x algorithm [19,20]. Amino acid sequences surrounding either up- or down-regulated phosphorylation sites were tested against a background reference set composed of sequences surrounding all phosphorylation sites revealed in our study that reached the minimal localization probability 0.75. Results of these analyses are shown in Figure 4.

Figure 4A,C depicts that among up-regulated sites the proline-directed and basic motifs were overrepresented whereas acidic motifs were significantly underrepresented (typical e.g., for casein

kinases). SP/TP motifs followed by basic amino acids are known to be typical consensus motifs of cyclin-dependent kinases (CDKs), mitogen-activated protein kinases (MAPKs) or glycogen synthase kinase 3 (GSK-3). On the other hand, SP/TP motifs were underrepresented amongst significantly down-regulated phosphorylation sites (Figure 4B,D) while SQ/TQ motifs (typical for DNA repair enzymes), TL, SR, and GT motifs were significantly overrepresented.

Figure 4. Sequence motifs analyses and visualization performed using iceLogo tool (\mathbf{A} , \mathbf{B}) and motif-x algorithm (\mathbf{C} , \mathbf{D}). The amino acid sequences of significantly differentially up- or down-regulated phosphorylation sites (\mathbf{A} , \mathbf{C} or \mathbf{B} , \mathbf{D} respectively) were analyzed against a statistical background comprising all class I sites revealed in our study. In \mathbf{A} and \mathbf{B} , amino acids that were more frequently observed in the proximity of a regulated phosphorylation site are indicated over the middle line, whereas the amino acids with lower frequency are indicated below the line; phosphorylated amino acid is located at position 8; \mathbf{C} and \mathbf{D} depict sequence motifs extracted by motif-x algorithm at a significance level of p < 0.00003 (which approximately corresponds to a q-value of 0.01 after Bonferroni correction for multiple hypothesis testing).



2.6. Kinase Activity Analyses Predicts Various Kinase Substrates to Be Regulated

There are several tools available to predict kinase-specific phosphorylation sites. Since each suffers its own limitations, the employment of different prediction algorithms seems to be a profitable approach to describe the alterations of intracellular signaling. Based on this assumption, at first we applied a very common approach relying on simple assigning of consensus motifs downloaded from HPRD (Human proteome reference database) to phosphorylated sequences. Consecutively, we employed two kinase predictors that apply consensus motif scoring together with the network context of kinases and their potential substrate, *i.e.*, NetworKIN 3 [21] and iGPS [22], differing in their motif scoring algorithms. The results are depicted in Figure 5. Only those kinases that reached the significance level of p < 0.05 are reported. The position score indicates the degree of regulation.

Figure 5. Analysis of kinases activity. Kinase-substrate relations predicted by iGPS (**A**) NetworKIN 3; and (**B**) linear motifs analysis derived from motifs stored in Human proteome reference database (HPRD) database; (**C**) The color shade corresponds to the value of the "position score" that indicates if the ratios (normalized H/L ratios) of sites phosphorylated by a particular kinase tend to be larger (0 to 1) or lower (-1 to 0) than the rest of the data (p < 0.05), *i.e.*, green color of a region in a heatmap indicates an increased activity of a particular kinase after IR and VE-821 co-treatment, red color indicates trend of decreased activity.



Our analyses confirmed the decrease in activity of the group of enzymes sharing the SQ/TQ motif (*i.e.*, ATM, ATR and DNA-PK). The remaining down-regulated kinases composed a quite heterogenic group of protein kinases including Casein kinase 2 (CK2), Protein kinase B (AKT) a subgroup of Protein kinases C (PKCs) or various isoforms of Pyruvate dehydrogenase kinase (PDK).

On the contrary, the increase in kinases with SP/TP specificity (*i.e.*, CDKSs, MAPKs, and GSK-3 *etc.*) was revealed. Additional groups of kinases that were indicated to be up-regulated mostly comprised kinases that were involved in mitosis, e.g., Polo-like kinase 1 (PLK1), Never in mitosis A-related kinase 2 (NEK2) or Dual specificity protein kinase TTK (TTK).

3. Discussion

DNA damage response (DDR) is an essential mechanism to maintain the genome integrity of cells. Moreover, its functionality implicates the efficiency of radio- or chemotherapeutic agents administered during cancer therapy and thus underlies cell resistance to genotoxic stress in normal as well as in cancer cells. Therefore, the therapeutic inhibition of DNA repair enzymes represents a promising strategy how to increase sensitivity towards genotoxic insults in cancer cells. One of the repair enzymes is the ATR kinase that is considered to primarily act in the S and G2 phases of the cell cycle and suppress the "physiological" replication stress as well as genotoxic stress induced by agents such as hypoxia, UV- and ionizing radiation, and hydroxyurea (HU) [8].

A recent discovery of the first selective inhibitors of ATR kinase facilitated investigation of the cellular functions of ATR and launched the development of new potential anti-cancer drugs targeting this kinase to sensitize cancer cells to chemo- or radiotherapy [3]. Reaper *et al.* confirmed that ATR inhibition using VE-821 that belongs to the group of 3-amino-6-pyrazines may offer great promise in cancer treatment. In concordance with previous results suggesting that disruption in ATM-p53 pathway should enhance the sensitivity of cells to the disruption of the ATR pathway [23], VE-821 was shown to selectively induce cytotoxicity in cancer cells without affecting normal cells. Treatment with 20 μM VE-821 alone induced irreversible growth arrest and apoptosis in a large fraction of different cancer cell lines that were p53-deficient (human colorectal carcinoma cell lines HT23, HT29 or human malignant melanoma cell line HT144) or p53-proficient but with a defective ATM-mediated pathway (human colorectal carcinoma cells HCT116). On the other hand, growth arrest induced by VE-821 in normal human lung fibroblasts (HFL-1) was fully reversible with minimal death [4].

In our previous study, we have confirmed the radiosensitizing effects of VE-821 in relatively radioresistant p53 negative cells of promyelocytic leukemia (HL-60). The inhibition of ATR (10 µM VE-821) in combination with IR (3 Gy) resulted in pronounced decreases in clonogenic survival and significant changes in the proportions of cells in S and G2 phases of the cell cycle as compared to irradiated control cells [13]. In our recent work, we employed the phosphoproteomic approach to describe the changes in intracellular signaling caused by radiosensitization in more detail. Our goal was to characterize signaling pathways triggered by irradiation and modulated by inhibition of an essential DNA repair enzyme ATR kinase. We aimed to reveal which kinases showed alterations in their activities in the presence of the ATR inhibitor VE-821 and thus were probably functionally linked to ATR-mediated response upon genotoxic stress. Moreover, deeper insight into processes

triggered in cancer cells sensitive to ATR inhibition may in the future lead to identification of potential markers of sensitivity to VE-821 treatment.

Both "motif-network" kinase analyses confirmed an expected decrease in the activity of the ATR kinase that was inhibited in order to radiosensitize the HL-60 cells. However, a decrease in ATM kinase activity was also evaluated to be statistically significant. Since it was reported, that the activity of the ATM kinase is not affected by VE-821 in 10 µM concentration ([4,13]; assessed by monitoring of CHK2 Thr68 phosphorylation which is a widely accepted specific marker of ATM activity), we account it to the decrease in phosphorylation of the consensus SQ/TQ motif which is shared between ATR, ATM and also DNA-PK. Additionally, ATM and ATR kinases share overlapping substrate specificity and hence the addition of protein-protein interaction scoring into the analysis could not improve the scoring algorithm to further distinguish between them. Interestingly, the activity of CHK1, which is a direct downstream target of ATR, was revealed to be significantly decreased in the HPRD motifs analysis (p-value lower than 0.05), however the activity shift was evaluated to be lower than expected after ATR/CHK1 pathway disruption (position score of only -0.07, where position score of 0 indicates non-regulated activity and -1 strongly downshifted activity). Moreover, Ser296 autophosphorylation of CHK1 and importantly Ser216 phosphorylation of CDC25C (which is an accepted marker of functional CHK1 dependent signaling) were not significantly altered in our work. On the other hand, we detected down-regulation of phosphorylation at Ser345 by immunoblotting, which serves to localize CHK1 to the nucleus following checkpoint activation [24]. Another important CHK1 phosphorylation at Ser317 was not detected in our phosphoproteomic analysis. Altogether, these findings underline the important concepts that have been reported recently: (i) the abrogation of different phosphorylation sites at CHK1—and thus the ATR/CHK1 pathway-by VE-821—is cell line—and treatment-dependent [7]; (ii) the radiosensitizing mechanism of VE-821 cannot be attributed merely to the abrogation of the ATR/CHK1 pathway [2,7].

All analyses further confirmed an increase in the activity of CDKs (SP/TP consensus motifs) that are inhibited by DNA repair enzymes in response to genotoxic stress in order to induce the cell cycle arrest to allow the repair of the damaged DNA lesions. It has been reported previously, that p53-negative HL-60 cells irradiated by lower doses of IR (up to 10 Gy) cannot undergo the p53-dependent G1 cell cycle arrest and therefore transiently accumulate in S phase, arrest at G2/M cell cycle checkpoint and eventually die by a so-called postmitotic apoptosis [24]. In the presence of VE-821, the CDKs inhibitory effect of ATR kinase (that is normally activated by replication and genotoxic stress in S and the G2 phase of the cell cycle) diminished, which was proven by the cell cycle analysis; the proportion of cells in G2 phase significantly decreased when compared to irradiated cells indicating that this cell population proceeded into mitosis [13]. Together, these anticipated trends in ATR and CDKs activities revealed by bioinformatic processing of our data, proved their biological reliability and relevance in addition to calculated statistical significance.

IR induces various types of DNA damage in irradiated cells including base damage, DNA crosslinks, single strand breaks (SSBs) and importantly, the most deleterious double strand breaks (DSBs). While ATM is activated exclusively in response to DSBs, ATR responds to DSBs as well as to SSBs [25], however the response of ATR to DSBs that are not part of the replication process is ATM-dependent [26]. GO biological process (GOBP) over-representation analysis revealed

a significantly overrepresented term "cellular response to DNA damage stimulus" that included 37 proteins with significantly changed phosphorylation measured in our experiments. Moreover, GOBP terms such as "base-excision repair", "nucleotide-excision repair" or "double-strand break repair" were also significantly enriched. Overrepresented pathways comprised for example "Activation of ATR in response to replication stress" from Reactome database or "ATM pathway" from PID.

Naturally, the most desirable effect of ATR inhibition in cancer treatment is increased cell death. Our data indicated, that phosphorylation changes in signaling pathways involved in apoptosis were apparent 1 h after irradiation, which can be illustrated by results of pathways overrepresentation analysis (e.g., "Apoptosis", "Apoptotic cleavage of cellular proteins"). Again, increased apoptosis induction in HL-60 cells after combined treatment with IR (3 Gy) and VE-821 has been reported previously [13].

The disruption of the G2/M cell cycle checkpoint by VE-821 can be monitored by the cell cycle analysis conducted by flow-cytometry 24 h after irradiation as a significant decrease of G2 phase cells in comparison to a control group of cells irradiated without VE-821 and was published in our previous work [13]. In our recent experiment, we observed multiple over-represented GOBP level 4 terms related to mitotic processes (e.g., "mitosis", "centromere complex assembly", "cytokinesis", "sister chromatid cohesion", or "spindle localization") together with GO Cellular Component (GOCC) terms revealing the localization of some of the phosphorylated proteins at centromeric regions of condensed chromosomes ("condensed chromosome kinetochore") which occur during mitosis. Some of the pathways related to cell division were also significantly over-represented, comprising for example "Mitotic M-M/G1 phases" or "Separation of sister chromatids" from the Reactome database, and "Aurora B signaling" or "PLK1 signaling events" from PID. Moreover, multiple protein kinases with known functionality in mitotic entry, spindle checkpoints, regulation of chromosome segregation and cytokinesis, such as PLK1 or NEK2, were more activated in the presence of VE-821 and both of them are known to be inhibited in an ATM-dependent manner in response to IR [27-30]. Notably, the regulation of PLK1 by ATR kinase after UV-but not gamma-irradiation has been very recently described [31].

In addition to these expected markers of G2/M checkpoint disruption, we further focused on the mechanisms involved in the G1/S transition and their possible modulation by ATR kinase. The direct involvement of ATR in regulation of transcription at G1/S transition after replication stress (*i.e.*, genotoxic stress in S-phase; caused by HU-treatment) has been reported recently [32]. Our results indicate the role of ATR in these processes after IR-induced DNA damage since a substantial portion of regulated phosphoproteins revealed in our study are involved in over-represented Reactome pathways such as: "Mitotic G1-G1/S phases", "G1/S Transition", "E2F mediated regulation of DNA replication", and "Inhibition of replication initiation of damaged DNA by RB1/E2F1".

The transcriptional regulators at G1/S comprise multiple proteins with different functions and thus are often grouped into four categories: activators (E2F1, E2F2, and E2F3), repressors (E2F4, E2F5, E2F6, E2F7, and E2F8), inhibitors (RB) and co-repressors (RBL1 and RBL2). Besides transcriptional regulation (e.g., E2F1 induces transcription of E2F6, E2F7, and E2F8 which in turn function as a negative feedback loop), phosphorylation mediated by CDK-Cyclin complexes, DNA repair kinases

(ATM, ATR), and checkpoint kinases (CHK1 and CHK2) also have essential roles in regulation of G1/S [32]. Regarding the activators, cells exposed to IR in G1 phase were shown to accumulate E2F1 in an ATM-dependent manner (E2F1 stabilizing phosphorylation), which led to induction of S-phase and subsequently to apoptosis [33]. On the other hand, E2F6 repressor inhibition by ATR-CHK1-phosphorylation in response to replication stress (HU-treatment) resulted in pronounced E2F-dependent G1/S transcription, which in this particular scenario led to cell survival [32]. In summary, these two phosphorylation-mediated contradictory roles of the same transcriptional mechanism highlight the importance of phosphorylation in tight regulation of this process. Since we identified some significantly regulated phosphorylation sites of these proteins after ATR inhibition, we suggest that ATR kinase may contribute considerably to regulation of G1/S after gamma-irradiation at least in regards to HL-60 cells. Importantly, while the role of phosphorylation in modulation of pocket proteins (RB, RBL1, RBL2) activity is a well-described mechanism, the impact of posttranslational modifications on "atypical repressors" E2F7 and E2F8 functionality have not been described yet. We found two significantly increased phosphorylation sites in E2F8 and E2F4 repressors after ATR inhibition, however, further experiments are necessary to validate these phosphorylations as biologically relevant, which is our next goal.

Moreover, proteins involved in the pre-replicative complex (pre-RC) formation were significantly modified by phosphorylation after VE-821 treatment (e.g., "Activation of the pre-replicative complex" in over-represented Reactome pathways). Pre-RC formation and licensing, occurs in the G1 phase and the RC is activated at the G1/S transition by phosphorylation of its components, mainly ORCs (Origin recognition complex subunits) and MCMs (minichromosome maintenance complex subunits) complexes, CDC6, and CDT1, to initiate DNA-replication and thus progress into S phase [34]. We observed multiple phosphorylation sites with increased phosphorylation in ORCs and MCM proteins; most of them contained the SP/TP consensus motif typical for CDKs. Previously, ATR has been described to suppress origin firing in response to replication stress [9], which was confirmed here in the context of IR-induced DNA damage.

In conclusion, our analyses conducted in asynchronous p53 negative cells allowed us to observe a comprehensive impact of ATR kinase inhibition on cellular response to IR-induced genotoxic stress in all phases of the cell cycle. While the described work was conducted on a single cell line and the data need further validation, we found that ATR kinase inhibition modulated the mechanisms at the G1/S transition, impacted the intra-S-checkpoint, disrupted the G2/M checkpoint and altered the activity of kinases involved in mitosis.

4. Experimental Section

4.1. Cell Culture and Culture Conditions

HL-60 cells were obtained from the European Collection of Animal Cells Cultures (Porton Down, Salisbury, UK). Cells were cultured in Iscove's modified Dulbecco's media (IMDM) for SILAC containing 20% dialyzed fetal calf serum, 2 mM glutamine, 100 UI/mL penicillin, and 0.1 mg/mL streptomycin (all purchased from Sigma-Aldrich, St. Louis, MO, USA) at 37 °C, under controlled 5% CO₂ and humidified atmosphere. Media were further supplemented with either unlabeled L-lysine

(100 mg/L, K0) and L-arginine (84 mg/L, R0) or equimolar amounts of L-¹³C₆-lysine (K6) and L-¹³C₆-arginine (R6; isotopicaly labelled amino acids were purchased from Invitrogen, Carlsbad, CA, USA). L-proline (300 mg/L) was added to avoid metabolic conversion of arginine to proline [35]. For complete incorporation of labelled amino acids, cells were cultured for at least 6 doublings [36]. Cell counts were assessed by a hemocytometer and the cell membrane integrity was determined by the Trypan Blue exclusion technique (Sigma-Aldrich, St. Louis, MO, USA).

4.2. Gamma-Ray Irradiation

The selective inhibitor of ATR kinase, 3-amino-6-(4-(methylsulfonyl)phenyl)-*N*-phenylpyrazine-2-carboxamide (VE-821, APIs Chemical Co., Ltd., Shanghai, China) was dissolved in DMSO in 10 mM aliquots and stored at -80 °C. Thirty min before irradiation, the inhibitor was added to the "heavy" cells (K6/R6) at concentration of 10 μM, the "light" cells (K0/R0) were treated with DMSO, whose final concentration in culture was lower than 0.1% to avoid the DMSO-induced differentiation of HL-60 cells. Both groups were irradiated by the dose of 6 Gy using a ⁶⁰Co gamma-ray source (VF, Cerna Hora, Czech Republic) with a dose rate 0.5 Gy/min. After irradiation, the flasks were placed in an incubator. Three experimental replicates were analyzed.

4.3. Cell Lysis and Protein Digest

One hour after irradiation, the cells were washed with cold PBS and lysed as was published [37]: the cells were resuspended in ice-cold lysis buffer (50 mM ammonium bicarbonate, 1% sodium deoxycholate, phosphatase inhibitor cocktail 2). The lysate was immediately placed into boiling water bath and after 5 min incubation the samples were cooled to room temperature. To decrease viscosity, bensonase nuclease (2.5 U/µL) and MgCl₂ (1.5 mM) were added to samples. The lysate was then clarified by centrifugation and the protein concentration was measured by bicinchoninic acid assay. Sample volumes corresponding to 1.75 mg of "light" proteins and 1.75 mg of "heavy" proteins were pooled together, reduced with dithiothreitol, alkylated with iodoacetamide and digested O/N with trypsin at an enzyme-to-substrate ratio of 1:60 (sequence grade modified trypsin, Promega Corporation, Madison, WI, USA). Sodium deoxycholate was then extracted by ethyl acetate [38] and peptides were desalted via 500 mg Supelco C18 SPE cartridges (Supelco Analyticals, Bellefonte, PA, USA). All other chemicals for cell lysis and protein digestion were purchased from Sigma-Aldrich.

4.4. Electrophoresis and Western Blotting

One hour after irradiation by 6 Gy, the HL-60 cells were washed with PBS and lysed. Whole cell extracts were prepared by lysis in 500 μL of lysis buffer (137 mM NaCl; 10% glycerol; 1% *n*-octyl-β-glucopyranoside; 50 mM NaF; 20 mM Tris, pH = 8; 1 mM Na₃VO₄; 1 tablet of protein inhibitors CompleteTM Mini, Roche, Manheim, Germany). The lysates containing equal amount of protein (30 μg) were loaded onto a 12% SDS polyacrylamide gel. After electrophoresis, proteins were transferred to a polyvinylidene difluoride membrane (BioRad, Hercules, CA, USA), and hybridized with an appropriate antibody: anti-CHK1 and anti-CHK1 phosphorylated at serine 345

(1:500) from Cell Signaling (Danvers, MA, USA); anti- β -actin (1:2000) from Sigma. After washing, the blots were incubated with secondary peroxidase-conjugated antibody (Dako, Glostrup, Denmark) and the signal was developed with ECL detection kit (BM Chemiluminescence-POD, Roche) by exposure to a film.

4.5. Hydrophilic Interaction Liquid Chromatography Fractionation and Enrichment of Phosphopeptides

Dried peptide samples were fractionated by HILIC chromatography according to a protocol that has been published previously [39] using the 4.6 × 25 cm TSKgel[®] (Tosoh Biosciences, Stuttgart, Germany) Amide-80 HR 5 μm particle column with the TSKgel® Amide-80 HR 5 μm 4.6 × 1 cm guard column operated with Waters Separations Module 2695 at 0.5 mL/min. Briefly, 3.5 mg of evaporated samples were reconstituted in 80% B (98% acetonitrile (ACN)/0.1% trifluoroacetic acid (TFA), mobile phase A consisted of 2% acetonitrile with 0.1% TFA) and loaded onto the HILIC column. Peptides were then separated by a gradient of A over B from 80% to 60% B in 40 min and from 60% to 0% B in 5 min. Across the gradient, 22 fractions were collected (2×2 and 20×1 mL) from each replicate. Each fraction was then enriched for phosphopeptides using titanium dioxide chromatography [40]. At first, each fraction was supplemented with TFA and glutamic acid to reach final concentrations of 2% TFA and 100 mM glutamic acid. Titanium dioxide particles (Titansphere® 5 μm particles, GL Sciences, Torrance, CA, USA) were suspended in the loading solution (65% acetonitrile, 2% TFA, 100 mM glutamic acid) and a particular volume of titanium dioxide suspension depending on an expected amount of peptides and phosphopeptides in a particular fraction was added to each sample microtube. Microparticles with bound phosphopeptides were then washed with 200 μL of loading solution, 200 μL of washing solution 1 (65% acetonitrile with 0.5% TFA), 200 μL of washing solution 2 (65% acetonitrile with 0.1% TFA) and 100 μL of washing solution 2. Phosphopetides were then eluted by 150 µL of elution solution (20% acetonitrile/NH₄OH, pH 11.5) in two sequential elutions. Late fractions were subjected to the second enrichment. Eluates from the first and second enrichment were pooled together, acidified with 100% formic acid and placed in a SpeedVac until all crystals of ammonium formate were evaporated.

4.6. Mass Spectrometric Analysis

On-line LC-MS analyses were performed on Thermo Scientific Dionex UltimateTM 3000 RSLCnano system (Thermo Scientific, Bremen, Germany) coupled through Nanospray Flex ion source with Q Exactive mass spectrometer (Thermo Scientific, Bremen, Germany). TiO₂-enriched HILIC fractions were dissolved in 18 μL of 2% ACN/0.05% TFA and 2 μL were injected into RSLCnano system. Peptides were loaded on capillary trap column (C18 PepMap100, 3 μm, 100 A, 0.075 × 20 mm; Dionex) by 2% ACN/0.05% TFA mobile phase at flow rate 5 μL/min for 5 min and then eluted and separated on capillary column (C18 PepMap RSLC, 2 μm, 100 A, 0.075 × 150 mm; Dionex). Elution was carried out by step linear gradient of mobile phase B (80% ACN/0.1% FA) over mobile phase A (0.1% FA); from 4% to 36% B in 19 min and from 36% to 55% B in 6 min at flow rate 300 nL/min. Temperature of the column was 40 °C and eluent was monitored at 215 nm

during the separation. Spraying voltage was 1.7 kV and heated capillary temperature was 220 °C. The mass spectrometer was operated in the positive ion mode performing survey MS (range 300 to 1800 m/z) and data-dependent MS/MS scans performed on the six most intense precursors with dynamic exclusion window of 40 s. MS scans were acquired with 70,000 resolution at 200 m/z from 1×10^6 accumulated charges (maximum fill time was 100 ms). The lock mass at m/z 445.12003 ([(C₂H₆SiO)₆ + H]⁺) was used for internal calibration of mass spectra. Intensity threshold for triggering MS/MS was set at 1×10^5 for ions with $z \ge 2$ with a 3 Da isolation window. Precursor ions were accumulated with AGC of 1×10^5 (maximum fill time was 100 ms) and normalized collisional energy for HCD fragmentation was 27 units. MS/MS spectra were acquired with 17,500 resolution (at 200 m/z).

4.7. Data Processing

The raw files were processed with MaxQuant software version 1.3.0.5 [41]. Peak lists were searched against the human UniProt database (release February 2014) using Andromeda search engine [42]. Minimum peptide length was set to six amino acids and two missed cleavages were allowed. Carbamidomethylation of cysteines was set as fixed modification while oxidation of methionine, protein N-terminal acetylation, deamidation of glutamine and arginine and phosphorylation of serine, threonine, and tyrosine residues were used as variable modifications. Additionally, appropriate SILAC labels were selected (Arg6, Lys6). A mass tolerance of 6 and 20 ppm was allowed for MS and MS² peaks, respectively. Only proteins, peptides and phosphorylation sites with FDR lower than 0.01 were accepted. For protein quantification only unmodified peptides, peptides oxidized at methionine residues, acetylated at *N*-terminus, or deamidated were accepted, both razor and unique peptides were used for the calculation of protein *H/L* ratios.

4.8. Bioinformatic Analysis

Contaminants and reversed hits were removed before further data processing and data were further manually inspected. The GRT was used to find differentially regulated phosphorylation sites. Phosphorylation sites quantified in all three replicates were included in GRT and the FDR was estimated non-parametrically as described by Zhou *et al.* [43]. The significance level of GRT was set to FDR < 0.01.

GO and signaling pathways over-representation analyses were performed using ConsensusPathDB, over-representation analysis web tool [14,15]. Only those proteins containing differentially regulated phosphorylation sites evaluated in GRT were included in the analyses. Significance was estimated using hypergeometric testing with Benjamini-Hochberg FDR correction (FDR < 0.01). All three GO classes (molecular function, cellular component and biological process) were statistically tested as well as the enrichment of proteins involved in signaling pathways stored in the Reactome pathway database [16] and in the PID [17]. The background reference set for the statistical analysis comprised all ConsensusPathDB entities: (i) annotated with a UniProt identifier that were included in at least one signaling pathway or (ii) that were annotated with a GO category which comprised at least one protein from the "regulated" dataset.

To analyze and visualize sequence motifs surrounding phosphorylation sites identified and quantified in our study, we employed the iceLogo tool and motif-x algorithm. In both motif analyses, the amino acid sequences (± 7 residues) surrounding either significantly up- or down-regulated phosphorylation sites (determined using GRT) were tested against a background reference set composed of sequences surrounding all phosphorylation sites revealed in our study that reached the minimal localization probability 0.75 (*i.e.*, "class I phosphosites" [44]). In the iceLogo analysis the significance level was set to p < 0.01. Using motif-x, the significantly enriched linear motifs were extracted. Search parameters were set to at least 10 occurrences of a particular motif and the significance level of p < 0.00003 (which approximately corresponds to a q-value of 0.01 after Bonferroni correction for multiple hypothesis testing).

The changes in kinase activities were evaluated using three different tools for the predictions of phosphorylation site-specific kinases together with the evaluation of significance based on the Wilcoxon-Mann-Whitney test ("1D annotation enrichment" tool available in Perseus software, version 1.4.0.20 [45]). Input data were filtered to contain only those class I phosphorylation sites that were quantified in at least 2 of 3 replicates. In the first analysis, the consensus sequence motifs downloaded from the HPRD database were simply matched to sequences motifs of phosphorylation sites identified in our study (using "Add linear motifs" tool integrated in Perseus program). To further specify kinase predictions by combining motif scoring together with contextual motif (*i.e.*, protein-protein interaction scoring downloaded from the STRING database [46]) we employed two freely available predictors NetworKIN 3 [21] and iGPS [22]. Kinase-substrate relations predicted by NetworKIN 3 algorithm were further filtered according to the "NetworKIN score" (>2). For iGPS predictions, the "high" significance threshold was chosen (*i.e.*, FPR of 2% for S/T kinases and FPR of 4% for Y kinases). The significance threshold for the Wilcoxon-Mann-Whitney test was set to p < 0.05.

5. Conclusions

Defects in the DSB-induced DDR such as ATM and/or p53 deletion/mutation are common in human tumors, occur in up to 70% of cancer cells [47,48] and have been proposed to enable the proliferation of cancer cells with DNA lesions [49]. Our previous study suggested that in a p53-negative environment, the most effective inhibition involves ATR rather than ATM and/or DNA-PK [13]. This work provides for the first time a complex insight into the mechanism of radiosensitization of the p53-deficient model cell line (HL-60, human promyelocytic leukemia) by the highly specific ATR inhibitor, VE-821. We show here a powerful phosphoproteomic approach for investigation of the role of ATR-mediated signaling in the context of radiosensitization of cancer cells. We found that significantly regulated phosphorylation sites were involved in cellular processes related to mitosis, cell cycle regulation, and DNA repair. Furthermore, understanding these processes by applying the presented strategy might facilitate characterization of biomarkers for patients susceptible to selective ATR inhibition. Although the experiments presented here were performed only in one cell line, this study provides valuable information regarding complex ATR-mediated signaling changes, and should be further validated in other cell lines as well as in primary tumor

samples from patients, which is our next goal. Our data underline the concept of radiosensitization in tumor cells as a promising new way to substantially increase efficacy of current cancer therapy.

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Author Contributions

Aleš Tichý, Barbora Šalovská, and Martina Řezáčová conceived and designed the experiments; Barbora Šalovská performed cell cultivation, SILAC labeling, cell treatment, lysis, protein digest and phopsphopeptide enrichment; Kamila Ďurišová and Jiřina Vávrová performed Western-blotting experiments; Barbora Šalovská, Ivo Fabrik and Marek Link performed HILIC prefractionation and MS analysis; Barbora Šalovská performed data processing and bioinformatical analysis; Barbora Šalovská and Aleš Tichý wrote and revised the manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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Development of Laser Ionization Techniques for Evaluation of the Effect of Cancer Drugs Using Imaging Mass Spectrometry

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Abstract: Recently, combined therapy using chemotherapy and photodynamic therapy (PDT) has been proposed as a means of improving treatment outcomes. In order to evaluate the efficacy of combined therapy, it is necessary to determine the distribution of the anticancer drug and the photosensitizer. We investigated the use of imaging mass spectrometry (IMS) to simultaneously observe the distributions of an anticancer drug and photosensitizer administered to cancer cells. In particular, we sought to increase the sensitivity of detection of the anticancer drug docetaxel and the photosensitizer protoporphyrin IX (PpIX) by optimizing the ionization-assisting reagents. When we used a matrix consisting of equal weights of a zeolite (NaY5.6) and a conventional organic matrix (6-aza-2-thiothymine) in matrix-assisted laser desorption/ionization, the signal intensity of the sodium-adducted ion of docetaxel (administered at $100 \, \mu M$) increased about 13-fold. Moreover, we detected docetaxel with the zeolite matrix using the droplet method, and detected PpIX by fluorescence and IMS with α -cyano-4-hydroxycinnamic acid (CHCA) using the spray method.

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1. Introduction

In chemotherapy for cancer, the curative effect of an anticancer drug decreases with time because the cancer cells acquire drug resistance [1,2]. For example, docetaxel is often used to treat prostate cancers, but the response rate of docetaxel is not high (44.2% when used with prednisolone [3]), and the drug-resistant cells that arise are refractory to treatment. Consequently, new methods for treating drug-resistant tumors are urgently needed. One proposal for increasing the efficacy of treatment involves combining chemotherapy with photodynamic therapy (PDT). In PDT, the patient is treated with a photosensitizer that selectively accumulates in the tumor tissue, and then the tumor is targeted with a laser or other light source in order to excite the photosensitizer. However, current techniques for PDT do not exert a sufficient curative effect [4,5]. When chemotherapy and PDT are combined, the anticancer drug-resistant cells are killed by PDT, resulting in treatment that is more effective than PDT alone [6,7].

In order to evaluate the efficacy of combined therapy, it is important to determine the distributions of the anticancer drug and photosensitizer. Autoradiography is the standard means for determining drug distributions, but conventional drug imaging by autoradiography cannot be used to simultaneously observe multiple drugs [8]. Such simultaneous observation can be achieved using imaging mass spectrometry (IMS) of drugs, their metabolites, and other molecules in the biological tissue. However, the major disadvantage of the IMS is that its detection sensitivity is inferior to that

of autoradiography. Moreover, a large difference in the ionization efficiencies of the drugs of interest can result in ion suppression (*i.e.*, obstruction of the ionization of the drug with lower ionization efficiency).

To resolve this problem, we are investigating ways to use IMS to simultaneously observe the distributions of anticancer drug and photosensitizer administered to cancer cells. In IMS, matrix-assisted laser desorption/ionization (MALDI) is often used as the ionization method because of its sensitive solid sampling and soft ionization technique, which has been extensively applied to the analysis of macromolecules and low-molecular weight compounds [9–13]. The sensitivity of detection of a drug of interest by MALDI can be greatly enhanced by the use of ionization-assisting reagents. The purpose of this research was to increase the sensitivity of detection for the anticancer drug and photosensitizer administered to human prostate cancer cells, first by optimizing the ionization-assisting reagents used in the process, and then by investigating a new technique for simultaneous imaging of the anticancer drug and photosensitizer.

2. Results and Discussion

2.1. Ionization of Docetaxel Using Zeolite Matrix

With the goal of enhancing the sensitivity of detection for anticancer drugs and photosensitizers, we attempted to optimize the ionization-assisting reagents used in the detection process. To this end, we investigated four conventional matrices: α-cyano-4-hydroxycinnamic acid (CHCA), 2,5-dihydroxy benzoic acid (DHB), 6-aza-2-thiothymine (ATT), and 4-nitroaniline (4NA). Figure 1 shows the mass spectra obtained from a 1-μL spot of 100 μM docetaxel using each matrix. Protonated, sodium-adducted, and potassium-adducted ions of docetaxel were detected. The peak of the sodium-adducted ion of docetaxel was most intense when 4NA was used as the matrix, but the ionization efficiency was not sufficient to allow detection of docetaxel administered to human prostate cancer cells (PC-3). To enhance the detection sensitivity further, we investigated the use of an additive dissolved in the matrix solvent, e.g., sodium acetate or potassium acetate, because the peak intensity of the sodium-adducted ion of docetaxel was the highest among the three peaks we detected. Figure 2 shows the average peak intensities of the sodium-adducted and potassium-adducted ions of docetaxel when sodium acetate or potassium acetate were added to the matrix solvent. Although the peak intensity increased about 4-fold, the ionization efficiency was still not sufficient to allow detection of docetaxel administered to PC-3 cells.

Figure 1. Typical mass spectra obtained from a the spot of the docetaxel using (a) α-cyano-4-hydroxycinnamic acid (CHCA); (b) 2,5-dihydroxy benzoic acid (DHB); (c) 4-nitroaniline (4NA); and (d) 6-aza-2-thiothymine (ATT). M indicates docetaxel.

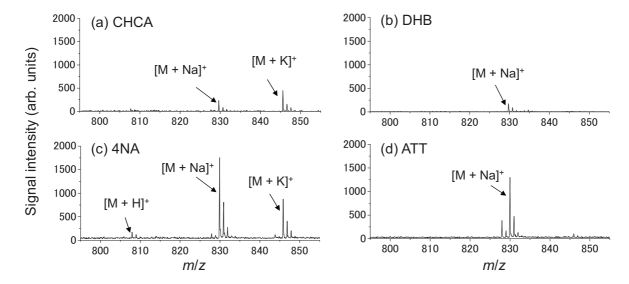
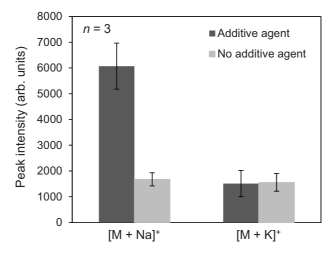


Figure 2. Relationship between the signal intensities of $[M + Na]^+$ and $[M + K]^+$. Black and gray bars mean that additive agent or no additive agent, respectively. M indicated docetaxel.



Next, in order to further enhance the detection sensitivity of docetaxel, we compared conventional matrix to a mixture of conventional matrix and zeolite matrix, a new ionization-assisting reagent that has been reported to increase the sensitivity of detection [14]. For this experiment, we selected NaY5.6 zeolite because it contains the sodium ion in its structure; as noted above, the peak intensity of sodium-adducted docetaxel was elevated when sodium acetate was added to the matrix solvent. Figure 3 shows typical mass spectra obtained from a spot of 100 µM docetaxel using the conventional matrix (ATT or 4NA) either alone or in combination with a zeolite matrix (NaY5.6). The peak intensity of the sodium-adducted ion [M + Na]⁺ was increased. Figure 4 shows the relationship between the signal-to-noise (S/N) ratio of the docetaxel peak in the presence of zeolite matrix at each concentration and laser pulse energy. Zeolite matrix prepared at 5 mg/mL was the most suitable for

ionization of docetaxel on a metal sample plate; at a zeolite matrix concentration of 1 or 3 mg/mL, docetaxel could not be detected. Figure 5 shows the average peak intensity of the sodium-adducted ion of docetaxel with 4NA or ATT, in the presence or absence of zeolite matrix. When zeolite matrix was used with ATT, the detection sensitivity of the sodium-adducted docetaxel peak increased 13-fold. By contrast, when zeolite matrix was used with 4NA, the detection sensitivity was not increased. Using the Voyager DE-PRO TOF instrument, the detection limit of docetaxel using zeolite matrix was 1 μ M. However, another MALDI-TOF instrument, the Ultraflex III, could obtain a peak at a lower concentration, 0.1 μ M. Thus, docetaxel and PpIX in the PC-3 cells could easily be detected using the conventional MALDI matrix and a modern time of flight (TOF) instrument.

Figure 3. Typical mass spectra obtained from docetaxel using (a) 4NA; (b) 4NA/NaY; (c) ATT; (d) ATT/NaY with no additive agents. M indicates docetaxel.

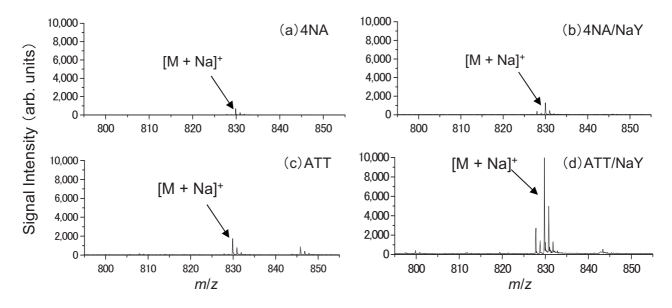


Figure 4. Relationship between S/N ratio of the docetaxel peak using zeolite matrix with no additive agents, at the indicated concentrations and laser pulse energies.

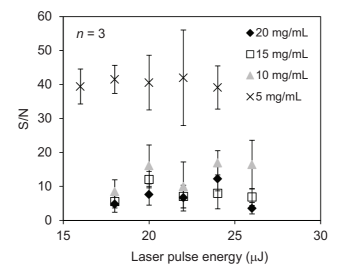
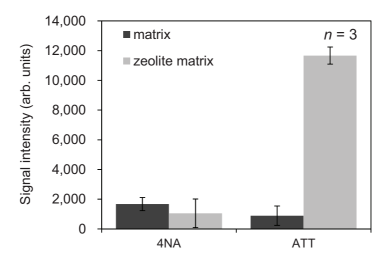


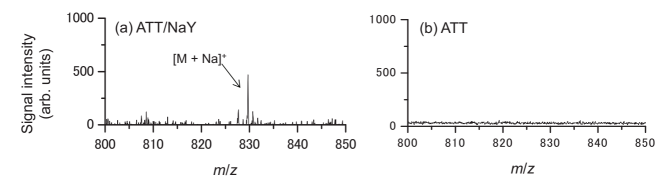
Figure 5. The average of peak intensity of the sodium-adducted ion of docetaxel using zeolite matrix with 4NA or ATT with no additive agents.



2.2. Detection of Docetaxel Administered to PC-3 Cells

As a first step toward acquiring images of docetaxel in prostate cancer tissue, we attempted to detect docetaxel administered to PC-3 cells using the zeolite matrix, ATT/NaY. Figure 6 shows typical mass spectra obtained from PC-3 cells administered 100 μ M docetaxel using (a) zeolite matrix or (b) conventional matrix using the droplet method. Only in condition (a), the sodium-adducted ion of docetaxel [M + Na]⁺ was detected from a spot of PC-3 cells. Figure 7 shows the S/N ratio of the [M + Na]⁺ peak at concentrations of 0, 10, and 100 μ M. [M + Na]⁺ was detected from PC-3 cells administered 10 μ M docetaxel. In this method, we observed the "sweet spot" effect. Therefore, each mass spectrum used for analysis was the average of 10 mass spectra.

Figure 6. Typical mass spectra obtained from the prostate cancer cells (PC-3) cells administered 100 μ M docetaxel using (a) the zeolite matrix; ATT/NaY or (b) the conventional matrix, ATT.



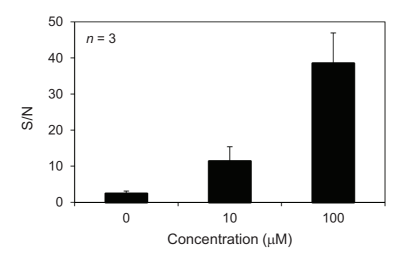
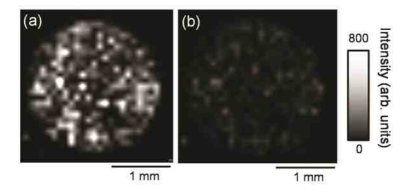


Figure 7. S/N ratio of the peak of $[M + Na]^+$ peak at 0, 10 and 100 μ M docetaxel.

2.3. Imaging of Cancer Drugs Administered to PC-3 Cells

The use of the zeolite matrix enhanced the ionization efficiency of docetaxel sufficiently to allow detection of the drug in cancer cells. Next, we investigated ways to use IMS to simultaneously observe the distributions of anticancer drug and photosensitizer administered to cancer cells. First, we attempted IMS of docetaxel and PpIX administered to PC-3 with the zeolite matrix using the droplet method. Figure 8 shows the ion image of (a) m/z 829.5 (sodium-adducted docetaxel) and (b) m/z 563.1 (protonated PpIX) obtained from cultured PC-3 administered cells treated with 50 μ M docetaxel and 10 μ M PpIX, obtained with the zeolite matrix, ATT/NaY using the droplet method. In previous studies, the peak intensity of docetaxel was reduced when conventional matrices were used. In this study, however, docetaxel administered to cancer cells could be detected using zeolite matrix, whereas PpIX could not be detected in the same samples.

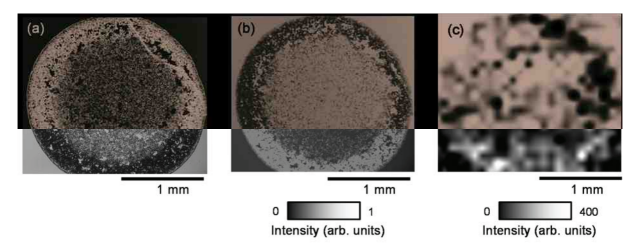
Figure 8. Ion image of (a) m/z 829.5 [D + Na]⁺; and (b) m/z 563.3 [P + H]⁺ obtained from PC-3 cells administered 50 μ M docetaxel and 10 μ M protoporphyrin IX (PpIX) using the zeolite matrix, ATT/NaY with the droplet method.



2.4. Imaging of PpIX Administered to PC-3 Cells

Although docetaxel administered to PC-3 cells could be detected with the zeolite matrix, PpIX could not be detected by IMS in the same samples. To address this issue, we investigated a new combined method. Initially, we had acquired optical and fluorescence images of the PC-3 cells, and also detected docetaxel by IMS. The new method was developed to allow acquisition of the ion image by IMS after acquisition of the fluorescence and optical images. Figure 9 shows (a) optical bright field image, (b) fluorescence image, and (c) ion image measured with CHCA using the spray method. All images were obtained from the same cultured PC-3 cells administered 10 µM PpIX. Under these conditions, PpIX could be detected, and the distribution of PpIX in the ion image approximately corresponded to that in the fluorescence image. In MALDI imaging, docetaxel and PpIX could not be detected in the same sample. However, using the new method, it was possible to obtain both fluorescence images and IMS from the same cells. Thus, using this method, docetaxel and PpIX could be detected in the same sample, by MALDI imaging and fluorescence imaging, respectively. We will seek to confirm this in a future study with cellular scale using a stigmatic imaging mass spectrometer, MULTUM-IMG2 [15,16].

Figure 9. (a) Optical bright field image; (b) fluorescence image; and (c) ion image measured with CHCA using the spray method, obtained from the same PC-3 cells administered $10 \mu M$ PpIX.



3. Experimental Section

3.1. Chemical and Reagents

Docetaxel is frequently used to treat prostate cancer. Docetaxel trihydrate was purchased from Wako (Osaka, Japan). PpIX was purchased from SIGMA-Aldrich (Tokyo, Japan). CHCA, DHB, 4NA, and ATT, used as the matrix, were purchased from Sigma-Aldrich (Tokyo, Japan). Acetonitrile and methanol (SIGMA-Aldrich, Tokyo, Japan) were used as the matrix solvent. The zeolite, NaY5.6, was supplied by the Catalysis Society of Japan (Tokyo, Japan).

3.2. Cell Culture

The cell line used was the hormone-antagonistic human prostate cancer cell line PC-3. Cells were cultured in Dulbecco's modified Eagle's medium (D-MEM, D6046, SIGMA-Aldrich, Tokyo, Japan) containing 10% fetal bovine serum (FBS, S1820, Biowest SAS, Nuaillé, France) and 100 units/mL each of penicillin and streptomycin (T4049, SIGMA-Aldrich, Tokyo, Japan). Cells were incubated at 37 °C in an atmosphere containing 5% CO₂.

3.3. Mass Spectrometer

All experiments were carried out in positive mode with reflectron-mode using Voyager DE-PRO TOF mass spectrometer (Applied Biosystems, Foster City, CA, USA) equipped with a 355-nm third-harmonic Nd:YAG laser (GAIA II 30-T, Rayture Systems Co., Ltd., Tokyo, Japan). Instrument parameters on the reflectron-mode of Voyager DE-PRO were as follows: accelerating voltage, +20 kV; voltage for the extraction grid, +13.6 kV; voltage for the guide wire, 0 V; extraction delay time, 100 ns.

3.4. Sample Preparation

For the conventional matrices, CHCA was dissolved in 1:1 (v/v) acetonitrile:water (v/v) at 10 mg/mL, and DHB, 4NA, and ATT were dissolved in 1:1 (ν/ν) methanol:water at 30, 5, and 10 mg/mL, respectively. Sodium acetate or potassium acetate was used as an additive agent and dissolved in the matrix solvent at 10 mM. For the zeolite matrices, 4NA or ATT was mixed with an equal weight of NaY5.6 zeolite with a mortar for over 10 min, and then suspended in 1:1 (v/v)methanol:water. To optimize the matrix for ionization of docetaxel, initially a 1-µL drop of docetaxel was spotted onto a metal sample plate, and then dried under vacuum; the each matrix was placed on dried spot of docetaxel. In this experiment, either indium tin oxide (ITO)-coated glass slide (#237001, Bruker, Billerica, MA, USA) or a metal plate (4347686, Applied Biosystems, Foster City, CA, USA) was used as the sample plate. From experiments described in Sections 2.1–2.3, the metal sample plate was used. For the experiment described in Section 2.4, the ITO-coated glass slide was used to acquire the optical images. To compensate for the low conductivity of the glass slide, which caused a charge increase relative to the metal sample plate, the ITO glass slide was coated with gold at a thickness of 10 nm using an ion-sputtering device (E-1010, HITACHI, Tokyo, Japan). Then, PC-3 cells were spotted onto the ITO glass slide, and an optical bright field image and fluorescence image were acquired after the spot was dried under vacuum. Before matrix application, Gold was again coated at a thickness of 10 nm. Matrix was applied using an airbrush (PS-153: 3500 PRO SPRY Mk-2, Mr. hobby, GSI Creos Co., Tokyo, Japan). Because it was possible that the metal nozzle would be degraded by the acidity of the matrix, reducing the intensities of the signals of interest, the nozzle was replaced with a microchip (DIAMOND TIP 0.1–10 μL, F171100, Gilson, Middleton, WI, USA) and a glass tube (Calibrated Pipets, 2-000-001-90, Drummond Scientific Co., Broomall, PA, USA).

3.5. Drug Administrations into Cells

Docetaxel and PpIX were dissolved in dimethyl sulfoxide (DMSO, D4540, SIGMA-Aldrich, Tokyo, Japan). The solution of docetaxel or PpIX was diluted 100-fold in medium containing 10% fetal bovine serum or not, respectively. PC-3 cells were incubated for 3 h in the medium containing each drug. After each drug administration, the cells were washed once with phosphate-buffered saline (PBS, D8537, SIGMA-Aldrich, Tokyo, Japan), trypsinized, resuspended in PBS, and centrifuged. After removal of the supernatant, a suspension of PC-3 cells at 1.0×10^7 cells/mL was prepared by addition of distilled water.

3.6. Microscopy

The optical bright field and fluorescence images of PpIX administered to PC-3 cells were acquired using an inverted fluorescence microscope (LEITZ DMIRB, Leica Microscope and System GmbH, Wetzlar, Germany) equipped with a mercury lamp (02651000, IREM, Borgone Susa, Italy) as the excitation light source. Exposure time was set at 20 s.

4. Conclusions

In order to evaluate the efficacy of combined therapy, it is important to determine the distributions of the anticancer drug and photosensitizer. The purpose of this research was to enhance the sensitivities of detection of anticancer drug and photosensitizer administered to PC-3 human prostate cancer cells by improving the ionization assisting reagents used in the process. When we used a matrix consisting of equal weights of zeolite (NaY5.6) and the conventional organic matrix 6-aza-2-thiothymine (ATT) in matrix-assisted laser desorption/ionization, the signal intensity of the sodium-adducted ion of docetaxel (administered at 100 µM) increased about 13-fold. Moreover, docetaxel administered to PC-3 cells could be detected with zeolite matrix using the droplet method. Although the ion image of docetaxel and PpIX could not be acquired simultaneously, PpIX administered to PC-3 cells could be detected in a fluorescence image, after which we could successfully detect PpIX in the same samples by IMS with CHCA using the spray method. These results demonstrate that it is possible to detect docetaxel in IMS using a zeolite matrix, following the detection of PpIX by fluorescence imaging.

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Author Contributions

Hiroki Kannen and Hisanao Hazama carried out the analysis and interpretation of data, the collection and assembly of data and drafted the manuscript. Yasufumi Kaneda and Tatsuya Fujino

designed the conception of this study from medical and chemical sides, respectively. Kunio Awazu supervised this work. All authors read and approved the final manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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Continuous Flow Atmospheric Pressure Laser Desorption/Ionization Using a 6-7-µm-Band Mid-Infrared Tunable Laser for Biomolecular Mass Spectrometry

Ryuji Hiraguchi, Hisanao Hazama, Kenichirou Senoo, Yukinori Yahata, Katsuyoshi Masuda and Kunio Awazu

Abstract: A continuous flow atmospheric pressure laser desorption/ionization technique using a porous stainless steel probe and a 6–7- μ m-band mid-infrared tunable laser was developed. This ion source is capable of direct ionization from a continuous flow with a high temporal stability. The 6–7- μ m wavelength region corresponds to the characteristic absorption bands of various molecular vibration modes, including O–H, C=O, CH₃ and C–N bonds. Consequently, many organic compounds and solvents, including water, have characteristic absorption peaks in this region. This ion source requires no additional matrix, and utilizes water or acetonitrile as the solvent matrix at several absorption peak wavelengths (6.05 and 7.27 μ m, respectively). The distribution of multiply-charged peptide ions is extremely sensitive to the temperature of the heated capillary, which is the inlet of the mass spectrometer. This ionization technique has potential for the interface of liquid chromatography/mass spectrometry (LC/MS).

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1. Introduction

Currently, electrospray ionization (ESI) [1,2] and matrix-assisted laser desorption/ionization (MALDI) [3,4] are the two major ionization methods for large biomolecules. Although these methods are used in combination with other separation techniques (e.g., liquid chromatography) for proteome analysis, they have some limitations. ESI can continuously ionize flowing liquid samples, but ionization is unstable for low volatility solvents, such as buffer solutions. On the other hand, conventional MALDI is limited to dry samples, because the MALDI process is conducted under a high vacuum, and co-crystallization of the analyte and aromatic matrix leads to spatial inhomogeneity and a low reproducibility. In fact, conventional soft ionization methods for biological polymers cannot produce a highly stable ionization with low volatility solvents. Consequently, novel ionization techniques are necessary for high-throughput and reproducible proteome analysis.

To overcome the above issues, the solvent, the state in conventional ionization methods (ESI and MALDI) and various original ambient ionization methods that can simplify or omit the sample preparation process have been intensively investigated. A few methods have been reported at atmospheric pressure. Ionization under atmospheric pressure can tolerate various sample states, because the samples do not have to be dry. Additionally, electrospray laser desorption ionization (ELDI) [5] generates gas phase ions when an untreated tissue sample is irradiated with an ultraviolet

(UV) laser under atmospheric pressure (AP), and the charged liquid droplets by the electrospray act to desorbed neutral molecules and ions. Unlike ELDI, matrix-assisted laser desorption electrospray ionization (MALDESI) [6] increases the ionization efficiency by adding an aromatic matrix. Laser ablation electrospray ionization (LAESI) [7] enables lipids from untreated biological samples to be directly ionized using the intensive absorption of an infrared (IR) laser by a tissue and electrospray, while modified ESI techniques without laser ablation/desorption have been developed for surface analysis. Desorption electrospray ionization (DESI), which was initially reported by Cooks *et al.*, enables proteins, peptides, *etc.*, to be directly analyzed from the surfaces of tissues and plants under ambient conditions [8]. DESI can also be used with thin-layer chromatography [9]. Probe electrospray ionization (PESI), which was developed by Hiraoka *et al.*, has a sampling system where a solid needle moves along a vertical axis by a motor-driven system [10]. PESI has the potential to analyze proteins directly from salt/urea-contaminated solutions [11].

However, these ambient ionization methods using laser desorption cannot ionize a liquid state sample directly. To realize high-throughput proteomics with a high reproducibility, an ambient laser ionization method for liquid state samples with low volatility solvents is desired.

Although conventional MALDI has been performed using UV lasers, IR-MALDI [12–14] has been investigated, because it may utilize more diverse matrices; many organic compounds with strong characteristic absorption peaks in the mid-IR wavelength range have been proposed as new matrices.

On the other hand, attempts to conduct the MALDI process under atmospheric pressure have been executed in order to expand the range of the analytical protocol. Laiko *et al.* presented an AP-MALDI technique coupled with an orthogonal acceleration time-of-flight (oaTOF) mass spectrometer and an ion trap mass spectrometer [15–17]. AP-MALDI has a better softness than conventional vacuum MALDI due to the collisional cooling of the expanding plume. In addition, alternating the surrounding condition to atmospheric pressure may expand AP-MALDI to include samples containing volatile compounds. In fact, online analysis of a solution sample by AP-MALDI has been demonstrated [18,19].

One of the most promising IR matrices is water, because it is abundant, environmentally benign and has a strong absorption in the mid-IR region. In 2002, Laiko *et al.* demonstrated the atmospheric pressure ionization of peptides in an aqueous solution using a Yb:YAG laser-pumped optical parametric oscillator (OPO) with a fixed wavelength of 3 µm [20]. Their work strongly suggests that utilizing a solvent as a light-absorbing matrix can expand the target samples in atmospheric pressure infrared matrix-assisted laser desorption ionization (AP-IR-MALDI) to include natural (aqueous) solutions. AP-IR-MALDI using a mid-IR OPO with a 2.9-µm wavelength has been applied to online liquid chromatography/mass spectrometry (LC/MS) analysis of the tryptic digest of bovine serum albumin [21]. Several works have employed 3-µm-band lasers, including OPO pumped by a Nd:YAG laser and an Er:YAG laser with a 2.94-µm wavelength for AP-IR-MALDI. This wavelength corresponds to the absorption peak of the O–H stretching vibration of liquid water.

The mid-IR wavelength region corresponds to the characteristic absorption bands of various molecular vibration modes, including O–H, C=O, CH₃ and C–N bonds. Consequently, many organic compounds and solvents, including water, have characteristic absorption peaks in this region (e.g., acetonitrile, methanol, acetone and several buffer solutions, such as a phosphate buffer, acetic acid

buffer and Tris buffer). For example, the O–H bending vibration for liquid water has an absorption peak at a wavelength of 6.07 μ m. Therefore, 6–7- μ m-band mid-IR lasers should be able to utilize diverse solvents as a matrix compared to 3- μ m-band lasers.

In fact, a 6-μm-band mid-infrared tunable laser using difference-frequency generation (DFG), which utilizes energy absorption in the C=O stretching region, has been applied to vacuum MALDI [22], and the softness for labile molecules (e.g., polysulfated oligosaccharides or polysialylated gangliosides) appears to exceed those of all other UV MALDI methods [23]. Simultaneous irradiations of a UV laser and 6-μm-band mid-IR free electron laser (FEL) enable protein samples containing a denaturant at a high concentration to be analyzed [24]. Awazu *et al.* have demonstrated a promising technique of IR-MALDI using a DFG laser utilizing various compounds (e.g., urea) as a matrix [25]. Although a 6–7-μm-band mid-infrared laser may utilize organic compounds, as well as various other solvents as a matrix when the MALDI process is conducted at atmospheric pressure, an AP-IR-MALDI method using a 6–7-μm-band infrared laser has yet to be reported.

In this study, we developed a new atmospheric pressure laser ionization method using a novel 6–7-µm-band mid-IR tunable laser. Our method allows the mass spectra of peptides to be directly measured from the continuous flow of several solutions.

2. Results and Discussion

2.1. Temporal Stability of Continuous Flow (CF) Ionization of Peptides

We initially examined the adequacy of the temporal stability of the ion signal intensity. Figure 1 shows a typical result of continuous ionization with a $5-\mu L/min$ sample flow and laser irradiation at 6.05 or 7.27 μm . The singly protonated ion $[M+H]^+$ of angiotensin II is predominantly observed. In addition, the extracted ion chromatogram of $[M+H]^+$ obtained from 50 mass spectra shows a superior stability of the ionization process compared with conventional MALDI using a solid matrix (Figure 2). The variability of the ion signal intensity has a standard deviation of 12.7%. On the other hand, the DFG laser has a variation of up to 10% in the output energy. In addition, the temporal relationship between laser pulses and injection time of the ion trap may be responsible for these variations, because the lasers were operated without synchronization. Thus, the sequential ion signal from the solution at a surface near the frit is responsible for the continuous flow ionization. This result suggests that this technique may be a novel interface in LC and MS. On the other hand, the efficiency of sample desorption depends strongly on the parameters of the laser and solvent. Thus, we then investigated the relationships between the laser wavelength, pulse energy, absorption coefficient of the solvent and ionization efficiency.

Figure 1. Typical mass spectra from angiotensin II dissolved in an 80% acetonitrile aqueous solution upon irradiation with a mid-infrared laser at wavelengths of (a) $6.05 \mu m$ and (b) $7.27 \mu m$. The electric potential of the frit and the capillary temperature were set to 2.5 kV and $270 \,^{\circ}\text{C}$, respectively.

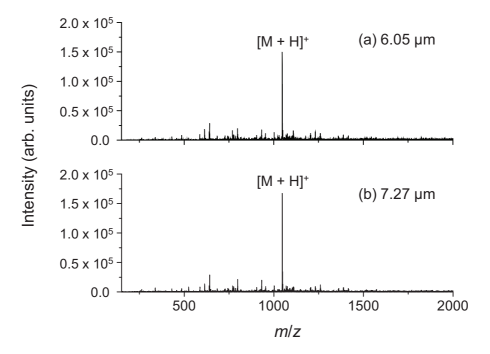
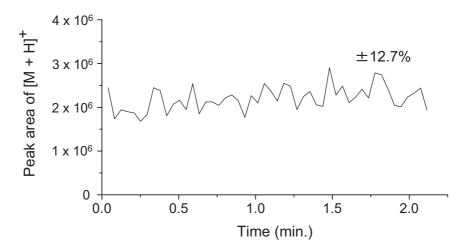


Figure 2. Extracted ion chromatogram of the protonated ion $[M + H]^+$ of angiotensin II. The wavelength of the DFG laser is set at 6.05 μ m, and the pulse energy is 400 μ J. Each data point was extracted from the raw mass spectra, which were averaged from three micro-scans. The electric potential of the frit and the capillary temperature were set to 2.5 kV and 270 °C, respectively.



2.2. Wavelength Dependence of the Ion Signal Intensity

The laser wavelength must be properly selected for efficient and subsequent ion yield due to the characteristic absorption of the solvent in the mid-infrared region. Figure 3 shows the absorption spectra of a typical mobile-phase in reversed phase liquid chromatography, which includes water, acetonitrile and a small amount of formic acid. The spectral shape of the mixed solvent depends on the mixing ratio. The O–H bending vibration around 6 μ m is due to water, whereas the CH₃ symmetric and degenerate bending vibration modes around 7.3 and 6.9 μ m, respectively, are due to acetonitrile. Previous reports about laser ionization without additional matrix elements depended on the strong absorption of water [20,21]. Herein, we tried to utilize a matrix containing water and acetonitrile using the mid-infrared tunable laser, which covers a wide wavelength range of 6–7 μ m.

Figure 4 shows the relationship between the ion signal intensity of $[M + H]^+$ and the IR absorption spectrum. Mainly two local maxima are observed in the plot of the ion signal intensity at the respective absorption peak wavelengths of water and acetonitrile. In addition, these peak wavelengths are very similar to the case of an aqueous solution on the sample plate. These results indicate that acetonitrile can act as a matrix by using a laser with the wavelength that corresponds to the matrix absorption peak.

In general, ESI is taken as one of the most useful techniques for the production of molecular ions also in LC/MS. In the gradient analysis in which the mixing ratio of the organic solvent in an aqueous solution is elevated as the analysis proceeds, an excessive concentration of acetonitrile decreases the ionization efficiency of ESI. On the other hand, our novel method using a mid-infrared laser can utilize both water and acetonitrile as the "ionization support agent". Therefore, the appropriate wavelength can be selected based on the eluent condition of HPLC in LC/MS with this ionization technique. Even in the case of gradient LC/MS, the center wavelength of the peaks is almost constant; even the mixing ratio is changed, as shown in Figure 3. Thus, gradient LC/MS should be possible using a fixed laser wavelength.

Figure 3. IR absorption spectra of the acetonitrile mixture used in reversed-phase liquid chromatography.

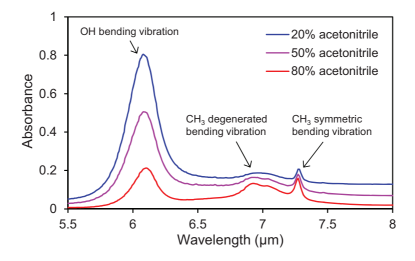
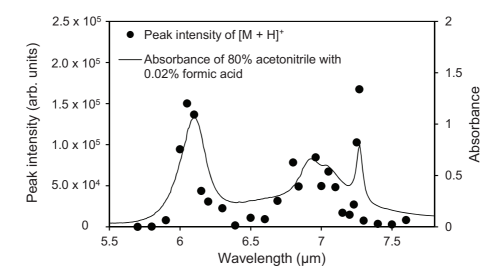


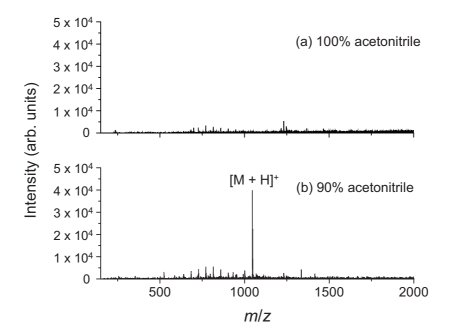
Figure 4. Wavelength dependence of the peak intensity of $[M + H]^+$ of angiotensin II from an 80% acetonitrile aqueous solution and the IR absorption spectrum of the 80% acetonitrile aqueous solution. The electric potential of the frit and the capillary temperature were set to 2.5 kV and 270 °C, respectively.



2.3. Dependence of the Ionization Efficiency on the Mixing Ratio of Water and Organic Solvent

It has been previously suggested that laser absorption at 7.27 µm by acetonitrile contributes to the ionization of a dissolved peptide sample. On the other hand, the residual water solvent contained in the dried matrix crystal has been reported to influence the ionization processes in UV-MALDI [26]. Thus, we investigated whether water is necessary for the laser wavelength corresponding to the specific absorption peak of acetonitrile. Figure 5 shows typical mass spectra obtained from angiotensin II dissolved in 100% acetonitrile and a 90% acetonitrile aqueous solution. [M + H]⁺ of angiotensin II is clearly observed for the 90% acetonitrile containing 10% water, but not for 100% acetonitrile, suggesting that the presence of water is essential, regardless of whether the laser wavelength is fixed at the absorption peak of another solvent (acetonitrile in this case).

Figure 5. Typical mass spectra of angiotensin II dissolved in (a) 100% acetonitrile and (b) a 90% acetonitrile aqueous solution upon irradiation with a mid-IR laser at a wavelength of 7.27 μm, which corresponds to the absorption peak of the CH₃ symmetric bending vibration mode in acetonitrile. The electric potential of the frit and the capillary temperature were set to 2.5 kV and 270 °C, respectively.



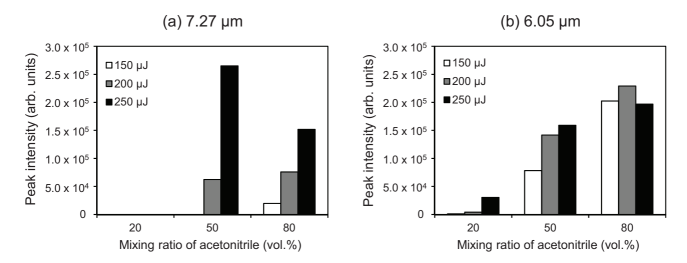
The absorption coefficients of the solvent at several wavelengths depend on the mixing ratios of the solvents. A larger mixing ratio of water leads to a stronger absorption peak in the 6- μ m range, which corresponds to the O–H bending vibration mode. Water also has a broadband absorption over the 6- μ m peak. The decrease in the absorption intensity at 7.27 μ m, which corresponds to the CH₃ symmetric bending vibration mode in acetonitrile, is almost countered by increasing the broadband absorption of water. Therefore, laser absorption by the solvent matrix may become constant even if the mixing ratio of the solvent (the eluent of HPLC) changes temporally according to the gradient program.

In conventional LC/MS using an ESI source, the fluctuating ionization efficiency due to the solvent mixing ratio makes quantitative analysis difficult. Hence, we investigated the dependence of the solvent mixing ratio on the ionization efficiency when using a 7.27- μ m laser wavelength. Figure 6 shows the relationship between the ion signal intensity of $[M + H]^+$ and three mixing ratios of water and acetonitrile. The ion signal intensities depend greatly on both the mixing ratio and the laser pulse energy.

We also investigated the influence of the solvent mixing ratio on the ion efficiency for the wavelength corresponding to water absorption (6.05 μ m). Similarly, the mixing ratio affects the ion signal intensity of [M + H]⁺ (Figure 6). The 80% acetonitrile aqueous solution gives the strongest ion signal of [M + H]⁺ despite having the weakest absorption at 6.05 μ m (Figure 3), indicating that the absorption intensity is not the only factor contributing to the ionization process. It is possible that the volatility and surface tension of the solvent may influence the desorption efficiency. Water has a relatively high surface tension and a lower volatility compared to other organic solvent. If the surface

tension and volatility influence the ionization process, desolvation would produce ions. All mass spectrometers with an atmospheric pressure ion source have a heated capillary or a skimmer to promote desolvation from charged droplets containing sample molecules. Thus, the capillary temperature may have a considerable effect on ion production.

Figure 6. Relationships between the ion signal intensity of $[M + H]^+$ of angiotensin II and the mixing ratio of acetonitrile in an aqueous solvent with laser wavelengths of (a) 7.27 µm and (b) 6.05 µm. The electric potential of the frit and the capillary temperature were set to 2.5 kV and 270 °C, respectively.



2.4. Relationship between the Generation of Multiply Charged Ions and the Desolvation Temperature

The above results suggest that the desolvation temperature and ion production are related. Thus, we investigated the relationship between the temperature of the heated capillary in an ion trap mass spectrometer (LCQ Classic, Thermo Finnigan, CA, USA) and the production of multiply-charged ions of peptides. Mass spectra were obtained from a mixture of three peptides (angiotensin II, P₁₄R and ACTH (adrenocorticotropic hormone) Fragment 18–39) dissolved in an 80% acetonitrile aqueous solution with 0.01% formic acid.

In previous research using the AP-IR-MALDI of peptides and proteins [27], multiply-charged ions were produced instead of singly-charged ions, which are chiefly observed in UV-MALDI. In addition, it was recently reported that liquid AP-UV-MALDI enables stable ion yields of multiply-charged ions of peptides and proteins [28].

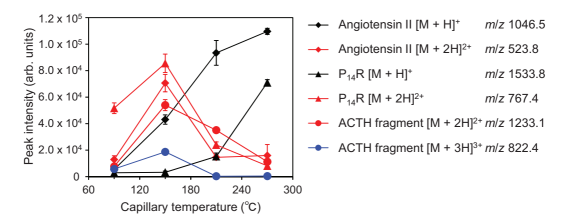
This ionization method produces multiply-charged ions, whose distribution drastically depends on the capillary temperature (Figure 7). For a capillary temperature of 270 °C, the mass spectrum contains mainly singly-protonated ions $[M + H]^+$, but the intensities of doubly-charged ion $[M + 2H]^{2+}$ and triply-charged ion $[M + 3H]^{3+}$ increase as the temperature decreases. These results indicate that the ion production process in 6–7- μ m atmospheric pressure laser desorption/ionization using a solvent matrix occurs inside the heated capillary and not at the laser irradiation point. Usually, in UV-MALDI, ions are produced in or form a dense plume containing the matrix and analytes in

the gas phase. Thus, the mechanisms of the ionization process in conventional MALDI and 6–7-μm atmospheric pressure laser desorption/ionization using a solvent matrix should differ.

On the other hand, the distribution of the ion valences produced by an electrospray shifts to the high-valence side, according to the increment of the vaporization rate from charged droplets [29,30]. On the other hand, a higher vaporization rate from charged droplets causes the formation of more highly charged ions in electrospray ionization [29,30]. In ESI, multiply-charged ions are produced due to the rapid desolvation from the charged droplets and the condensation of charge. Conversely, slow desolvation may result in poorly charged ions. ESI and continuous flow AP-IR-MALDI atmospheric pressure laser desorption/ionization are completely opposite in this regard. However, both have a commonality: they need the desolvation from charged droplets, which is not required for conventional MALDI. However, the mechanism is unclear at this time, and a more detailed study is necessary.

These investigations suggested that the laser might induce not only the desorption of a charged droplet, but ionization. On the other hand, for example, simply heating is one of the choices for just desorption. However, probe heating preferentially would desorb highly volatile compounds. As a result, non-volatile compounds remain on the probe. In this regard, the use of laser desorption could provide an important benefit that every compounds in the solution would be desorbed regardless of the volatility.

Figure 7. Relationship between the signal intensity of the protonated peptide ions and the temperature of the heated capillary in the mass spectrometer. The electric potential of the frit was set to 2.5 kV.



3. Experimental Section

3.1. Mid-Infrared Tunable Laser Using Difference-Frequency Generation (DFG)

A mid-IR tunable laser using difference-frequency generation (DFG) was used for ionization [31]. The DFG laser was developed by Kawasaki Heavy Industries, Ltd. (Kobe, Hyogo, Japan), and RIKEN (Wako, Saitama, Japan). Laser pulses from a Nd:YAG laser with a wavelength λ_1 of 1.064 µm and a tunable Cr:forsterite laser with a wavelength λ_2 of 1.19–1.32 µm were synchronized and mixed in two nonlinear optical crystals (AgGaS₂). Consequently, the mid-IR output of DFG with

a wavelength $\lambda_{DFG} = (1/\lambda_1 - 1/\lambda_2)^{-1}$ was obtained. This laser system had a tunable wavelength range of 5.50–10.00 µm, and wavelength tuning with a minimum interval of 0.01 µm was automatically controlled by a computer. The laser pulse width was about 5 ns, and the pulse repetition rate was 10 Hz. In the experiments, the laser pulse energy ranged between 250 and 400 µJ at the sample surface.

3.2. Ion Trap Mass Spectrometer with an Atmospheric Pressure Laser Ion Source

All experiments were conducted with an ion trap mass spectrometer (LCQ Classic, Thermo Finnigan, CA, USA) integrated with a self-assembled continuous flow atmospheric pressure laser ion source.

The sample probe was a porous stainless steel substance called a "frit" (Figure 8). A frit has been used with a conventional ionization technique using a neutral beam and some viscous matrices, called fast atom bombardment (FAB) [32,33]. These conditions realized continuous flow ionization, which was used as the interface of LC/MS [34–36]. In addition, a frit has applied to a modified MALDI technique using an infrared laser, named continuous flow IR-MALDI [37]. Although ionization by frit-FAB or continuous flow IR-MALDI was conducted under a high vacuum, in this work, the frit was subjected to atmospheric pressure in continuous flow laser ionization.

Figure 8. (a) Photograph of the frit probe and (b) scanning electron microscope image of the frit surface.

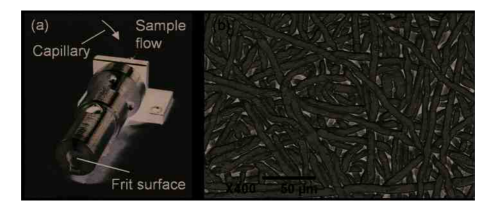


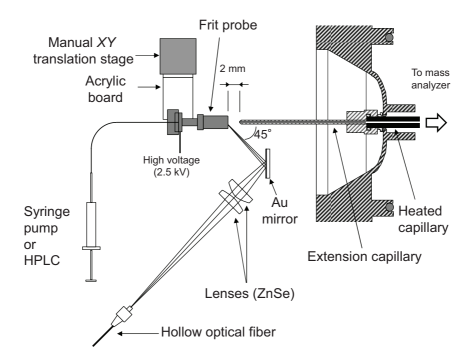
Figure 9 schematically depicts the measurement system. The output of the DFG laser was introduced into a hollow optical fiber with an inner diameter of 700 µm using an off-axis parabolic mirror to deliver the DFG laser to the ion source. The laser light from the hollow optical fiber was focused onto the sample plate using two ZnSe plano-convex lenses. The incident angle of the laser against the surface of the frit was about 45°. To calculate laser fluences on the frit surface, the knife-edge method measured the focused laser spot size, which was approximately 0.16 mm².

An extension capillary was connected to the heated capillary of the mass spectrometer to bring the ion inlet close to the frit surface. The length and inner diameter of the extension capillary were 80 and 0.6 mm, respectively, and the temperature of the heated capillary was set to 270 °C. The frit probe mounted on a manual XY translation stage was located 2 mm from the tip of the extension capillary. It is supposed that the potential of the frit probe should have an influence on the ion production. However, the effect of the sample plate voltage on the atmospheric ionization using

a 3-µm-band mid-infrared laser has already been reported elsewhere [38]. Thus, a static high voltage of 2.5 kV was applied to the frit probe by connecting the voltage source originally used for the ESI nozzle [39].

The ion injection time of the ion trap was set to 300 ms, and ions were produced by the irradiation of the DFG laser in the meantime. A raw mass spectrum was acquired by three micro-scans of the ion trap, and the mass spectra shown in Figures 1 and 5 were obtained from the averaging of 50 raw mass spectra.

Figure 9. Schematic of the ion source for continuous flow atmospheric pressure laser/desorption ionization using a frit probe integrated with the ion trap mass spectrometer.



3.3. Materials and Methods

Human angiotensin II (A8846), P₁₄R (synthetic) (P2613) and human ACTH Fragment 18–39 (A8346) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Distilled water (049-16787) was purchased from Wako Pure Chemical Industries Ltd. (Osaka, Japan). All were used as received. These peptides were dissolved in 20%–90% acetonitrile aqueous solutions with 0.005%–0.04% formic acid (11-0780-5, Katayama Chemical Industries Co., Ltd., Osaka, Japan) at a concentration of 10 pmol/μL. Otherwise, a peptide was dissolved in 100% acetonitrile.

Sample solutions were delivered to the frit probe via a syringe with a 250- μ L capacity (Unimetrics, IL, USA) and an LCQ classic syringe pump. The sample flow rate was controlled in the range of 3–20 μ L/min. A sample solution was pumped at a constant flow rate during laser irradiation.

The mass spectra shown in Figures 1 and 5 were averaged 50 times for only the reduction of noise. The temporal changes in the ion signal intensity were investigated by plotting the intensities in the respective spectra. The heated capillary was set at 90, 150, 210 or 270 °C.

4. Conclusions

The continuous flow atmospheric pressure laser ion source using a frit and a 6–7- μ m-band mid-infrared tunable laser is capable of direct ionization from a continuous flow with a high temporal stability. This modified ion source requires no additional matrix and utilizes water or acetonitrile as the solvent matrix at several absorption peak wavelengths (6.05 and 7.27 μ m).

In addition, the effects of the solvent mixing ratio on the ionization efficiency were investigated. The ion signal intensity depends on the mixing ratio at wavelengths of 6.05 and 7.27 μ m. In the case of 6.05 μ m, the ion signal intensity obtained from the 20% acetonitrile aqueous solution is lower than that from the 50% and 80% acetonitrile aqueous solutions despite their relatively high absorbance at 6.05 μ m, indicating that not only the intensity of laser absorption, but also the volatility and surface tension may affect the desorption efficiency.

The distribution of multiply-charged peptide ions is extremely sensitive to the temperature of the heated capillary, which is the inlet of the mass spectrometer. This suggests that ions are produced from the charged droplet after desorption of the solvent matrices (water or acetonitrile). The solvent matrices used in this work were broadly used as mobile-phase in reversed-phase liquid chromatography. Thus, this ionization method has the potential for the interface of LC/MS as an alternative for ESI.

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Author Contributions

The authors' contributions were as follows: Ryuji Hiraguchi and Hisanao Hazama designed and conducted the experiments; Kenichirou Senoo, Yukinori Yahata, Hisanao Hazama and Ryuji Hiraguchi constructed the experimental systems including the ion source; Katsuyoshi Masuda and Kunio Awazu supervised this work and suggested the experiments 2.2 and 2.3; Ryuji Hiraguchi and Hisanao Hazama prepared and edited the manuscript. All authors read and approved the final manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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Access of Hydrogen-Radicals to the Peptide-Backbone as a Measure for Estimating the Flexibility of Proteins Using Matrix-Assisted Laser Desorption/Ionization Mass Spectrometry

Mitsuo Takayama, Keishiro Nagoshi, Ryunosuke Iimuro and Kazuma Inatomi

Abstract: A factor for estimating the flexibility of proteins is described that uses a cleavage method of "in-source decay (ISD)" coupled with matrix-assisted laser desorption/ionization mass spectrometry (MALDI MS). The MALDI-ISD spectra of bovine serum albumin (BSA), myoglobin and thioredoxin show discontinuous intense ion peaks originating from one-side preferential cleavage at the N-C α bond of Xxx-Asp, Xxx-Asn, Xxx-Cys and Gly-Xxx residues. Consistent with these observations, Asp, Asn and Gly residues are also identified by other flexibility measures such as B-factor, turn preference, protection and fluorescence decay factors, while Asp, Asn, Cys and Gly residues are identified by turn preference factor based on X-ray crystallography. The results suggest that protein molecules embedded in/on MALDI matrix crystals partly maintain α -helix and that the reason some of the residues are more susceptible to ISD (Asp, Asn, Cys and Gly) and others less so (Ile and Val) is because of accessibility of the peptide backbone to hydrogen-radicals from matrix molecules. The hydrogen-radical accessibility in MALDI-ISD could therefore be adopted as a factor for measuring protein flexibility.

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1. Introduction

Mass spectrometry (MS) is an indispensable tool for analyzing biological molecules such as protein, nucleic acid, saccharides, hormones, lipid and neurotransmitters in living organisms. In particular, matrix-assisted laser desorption/ionization (MALDI) [1,2] and electrospray ionization (ESI) [3,4] have become widely recognized as powerful soft ionization methods for identifying the expressed proteome. Although both MALDI MS and ESI MS have outstanding abilities that allow identification of proteins due to their rapidity and high sensitivity, it is more difficult to glean the kind of information about secondary and tertiary structures that can be determined by X-ray crystallography and nuclear magnetic resonance (NMR) spectroscopy. However, recent reports about the relationships between MS and NMR data of cytochrome c suggest that MALDI [5,6] and ESI [7] might be useful for estimating flexible regions of proteins.

One of the recent interests in protein science is the concept of flexibility by means of the motion of backbone and/or side chains. It is believed that the flexibility of protein molecules relates to several functions such as post-translational modifications and drug interactions. The concept of intrinsically disordered proteins has been recognized as key in clarifying functions with respect to intermolecular

interactions with other proteins, nucleic acids and drug chemicals [8]. The flexibility of protein molecules is estimated by several measures such as the B-factor [9,10], the turn preference factor [11,12], the protection factor [13] and the fluorescence decay factor [14]. The B-factor and turn preference factor are based on X-ray crystallography in the crystal phase, while the protection factor is based on the hydrogen/deuterium exchange (HDX) reaction in NMR and the fluorescence decay factor can be obtained from protein molecules in solution. The B-factor which relates to side chain mobility suggests that Asp, Asn, Gly, Pro, Lys, Glu, Gln and Ser residues have a more flexible nature than other residues [15]. The turn preference factor indicates that Asp, Asn, Gly, Pro, Cys and Ser residues do not favor intra-molecular hydrogen bonded secondary structures such as α -helix and β -sheet [11,12]. The flexible residues estimated by the protection factor from NMR-HDX experiments of ubiquitin and cytochrome c have been reported to be Asp, Asn, Gly, Lys, Thr, Ile and Met [13,16]. The fluorescence decay factor shows that Asp, Asn, Gly, Ser and Ala are more flexible than the other residues [14]. Although the residues estimated are intrinsic to that method used, it is interesting that Asp, Asn and Gly residues are common to all the methods, independent of the principle and experimental conditions such as crystal and solution phase.

We recently reported that Asp, Asn, Gly and Cys residues in equine cytochrome c susceptible to in-source decay (ISD) coupled with MALDI-ISD were partly consistent with them being in flexible regions as estimated by the protection factor in NMR-HDX experiments [6,17]. This suggested that MALDI-ISD could be used as a method for estimating the flexibility of proteins. The MALDI-ISD experiments were performed by irradiating the surface of the matrix crystals, in which were embedded analyte protein molecules, with ultraviolet (UV) pulsed laser photons under vacuum conditions. The irradiation of the crystal phase with UV-laser photons results in ablation (to generate a dense-gas plume), proton transfer (to generate analyte ions), and hydrogen-radical transfer (to produce the ISD reaction). ISD is a radical-initiated cleavage at the N-Cα bond of the peptide backbone, and the processes can be divided into two steps: (a) the formation of hypervalent radical species of protein molecules (Scheme 1a) and (b) prompt cleavage (within several tens of nanoseconds in the ion source) to generate ISD fragments c, z', z-matrix and w (Scheme 1b,c) [18,19]. The formation of the ISD fragments can be rationalized by so-called "Takayama's model" [19]. According to the model, the hydrogen-radicals are generated from active-hydrogens (-OH and -NH₂) of the matrix molecules excited with UV photons [20-22]. However, reasons why the specific residues of Asp, Asn, Gly and Cys are more susceptible to ISD than other residues and why the one-side preferential cleavage at the N-Cα bond of the residues (Xxx-Asp, Xxx-Asp, Gly-Xxx and Xxx-Cys) occurs have not yet been rationalized, in spite of a related previous report [23]. It is of particular interest to consider the influence of secondary structures of proteins on the ISD fragment abundance, although it is difficult to obtain any evidence for the presence of α -helix and β -sheet for a given protein in the MALDI matrix crystal state and/or MS gas-phase conditions.

Scheme 1. (a) Interaction between matrix active-hydrogens and carbonyl oxygens of the peptide backbone, hydrogen transfer from matrix to the backbone and the formation of hypervalent radical species of protein. (b) Pathways of N-Cα Bond Cleavage of the Backbone to generate Fragment c/z• and c•/z Pairs and the Nomenclature of ISD (in-source decay) Fragments and (c) Formation of ISD Fragments w, z' and z-ANL.

Here we describe relationships between discontinuous intense peaks in the ISD spectra of analyte proteins and secondary structures based on the X-ray and NMR data. The proteins used are bovine serum albumin (BSA), equine myoglobin and human thioredoxin, for which there are primary, secondary and tertiary structures as determined by X-ray crystallography and NMR spectroscopy, and are recorded in the protein data bank (PDB). The ISD spectra showing fragment peaks of c-, z'- and w-ions are discussed from definite secondary structures as judged by means of X-ray and NMR data. The reactions of N-Cα bond cleavage and one-side preferential cleavage at the N-Cα bond of Xxx-Asp/Asn/Cys and Gly-Xxx are discussed on the basis of *ab initio* calculations. The results indicate that the one-side preferential N-Cα bond cleavage may be due to hydrogen bonding

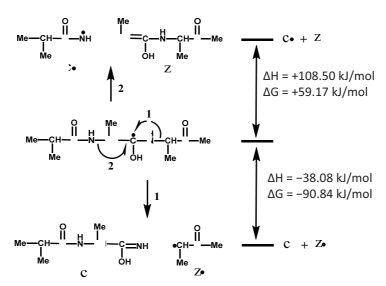
interactions between matrix active-hydrogen(s) and the carbonyl oxygens of the backbone of protein molecules embedded in/on the matrix crystal. Furthermore, it maybe that the protein molecules embedded in/on matrix crystals may partially maintain their helix structure, and that the carbonyl oxygens of helix-free backbone regions of the protein molecules embedded in/on matrix crystals are exposed to the matrix molecules and are easily accessed by matrix active-hydrogens. It is demonstrated that hydrogen-radical accessibility to the peptide-backbone is a factor for estimating the flexibility of protein and is compatible with the turn preferential factor for the flexible amino acid residues Asp, Asn, Gly and Cys.

2. Results and Discussion

2.1. Preferential Binding of Hydrogen-Radicals Determining One-Side Preferential Cleavage at N-Ca Bond of Peptide Backbone

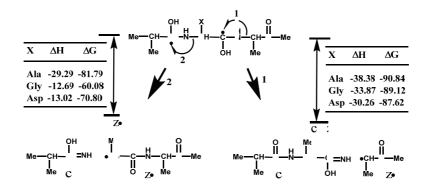
It is important to divide the overall ISD process into two parts, (a) the formation of hypervalent radical species II (Scheme 1a), and (b) the generation of ISD fragments (Scheme 1b,c). For the fragment formation in Scheme 1b, it needs be confirmed which reaction is energetically favorable for the formation of either the c/z•-pair (reaction 1) or the c•/z-pair (reaction 2) in Scheme 1b (II). In order to avoid any intra-molecular hydrogen bonding in the analyte, here we use a methylated model tri-peptide (Me-Ala-Ala-Me) for *ab initio* calculations. The calculations were performed for a radical species by which radical site was located on the carbonyl carbon at Ala2 residue (Scheme 2). The calculation results indicate that reaction 1 is overwhelmingly favorable over reaction 2, as shown in Scheme 2. Further, the results support a fact that the ISD spectra never give c-ANL fragment ions [6,19] originated from a radical fragment c• (IIIb in Scheme 1b), while z-ANL fragment ions originated from a radical fragment z• can be observed in the ISD spectra [6,19]. We can therefore employ reaction 1 for the formation and nomenclature of ISD fragments (Scheme 1b).

Scheme 2. Thermochemical Values ΔG (kJ/mol) and ΔH (kJ/mol) for the Formation of ISD Fragments c/z• and c•/z pairs obtained from *ab initio* Calculations.



To estimate the one-side preferential cleavage at the N-C α bond of n-th residue Xxx-Xn or Xn-Xxx, *ab initio* calculations were performed for the model peptide Me-Ala-X-Ala-Me (X = Ala, Gly, Asp). Both calculated enthalpy ΔH and Gibbs energy ΔG showed that *C*-terminal side cleavage (Xn-Xxx) is more favorable than the *N*-terminal side (Xxx-Xn), as shown in Scheme 3. The result supports the preferential cleavage at the N-C α bond of Gly-Xxx, but does not explain the even cleavage of Xxx-Ala and Ala-Xxx and the preferential cleavage at Xxx-Asp. On the basis of the calculations, it is suggested that the one-side preferential cleavage is not governed by the energetics of reactions 1 and 2 in Scheme 3. It seems reasonable to consider that the one-side preferential cleavage is attributable to preferential binding of hydrogen-radicals to the *N*-terminal side carbonyl oxygens (CO-NH-Asp/Asn/Cys) and *C*-terminal side carbonyl oxygens (Gly-CO). This indicates that accessibility of hydrogen-radicals to the backbone carbonyl oxygens of analyte proteins is essential for the preferential cleavage leading to the observation of discontinuous intense ISD fragment ions. It is of importance to note that the backbone regions easily accessible by hydrogen-radicals or matrix molecules have to be lying in flexible secondary structures.

Scheme 3. Thermochemical Values ΔH (kJ/mol) and ΔG (kJ/mol) for the Formation of c/z•. Pairs Originating from Cleavages at the N-C α bond of Both Sides of n-th Residues Xxx-X and X-Xxx.



2.2. Discontinuous Intense ISD (In-Source Decay) Ion Peaks Reflect Helix-Free Regions of Protein

Positive ion MALDI-ISD spectra of bovine serum albumin (BSA) and equine apo-myoglobin obtained with 5-amino-1-naphthol (5,1-ANL) as a matrix (which is a suitable matrix for MALDI-ISD experiments with protein [23,24]) are shown in Figures 1 and 2, respectively. The ISD spectra showed remarkably discontinuous intense peaks assigned as the fragments c-, z'- and w-ions (Table 1). The ionization (protonation and deprotonation) of the ISD fragments takes place independent of the ISD processes [25], and the ISD fragment ions are usually observed as singly-charged ions, *i.e.*, c-ion = $[c + H]^+$ and $[c - H]^-$, z'-ion = $[z' + H]^+$ and $[z' - H]^-$, z-matrix ion = $[z-matrix + H]^+$ and $[z-matrix - H]^-$, and w-ion = $[w + H]^+$ and $[w - H]^-$. The primary structures and ISD fragment ions (intensity) observed for the analyte proteins used here are presented in Supplementary Information 1 and 2, respectively.

Table 1. Discontinuous Intense Fragment Ions and Cleavage Sites observed in Positive Ion ISD Spectra of Intact Proteins BSA, Myoglobin and Thioredoxin.

Protein	Fragment Ion and Cleavage Residues
BSA	c12 (Xxx-Asp), c15 (Gly-Xxx), c33 (Xxx-Cys)
Myoglobin	z'15 (Xxx-Arg), c19 (Xxx-Asp), c35 (Gly-Xxx),
	c43 (Xxx-Asp), c59 (Xxx-Asp), c80 (Gly-Xxx)
Thioredoxin	c3 (Lys-Xxx), c19 (Gly-Xxx), c25 (Xxx-Asp), c31 (Xxx-Cys),
	c34 (Xxx-Cys), c61 (Xxx-Cys), c68 (Xxx-Cys), c72 (Xxx-Cys),
	w33 (Xxx-Cys), w37 (Xxx-Cys), w44 (Xxx-Cys), w71 (Xxx-Cys), w74 (Xxx-Cys)

Figure 1. Positive ion matrix-assisted laser desorption/ionization in-source decay (MALDI-ISD) spectrum of bovine serum albumin (Mr 66430.3) obtained with 5-amino-1-naphthol matrix. The inset represents tertiary structure obtained by X-ray crystallography (PDB: 4F5S). The upper line represents primary and secondary structure (wave: α -helix, convex: turn, straight: bend or empty).

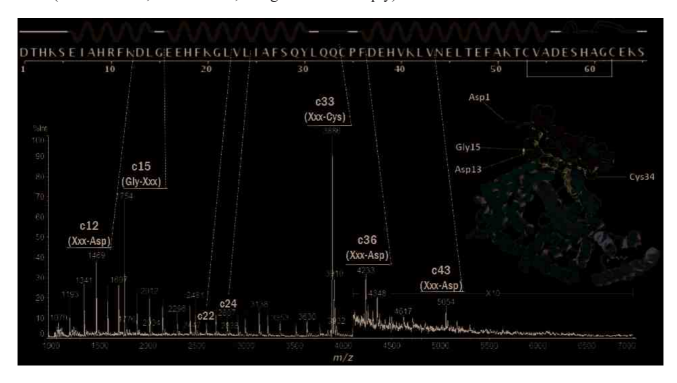
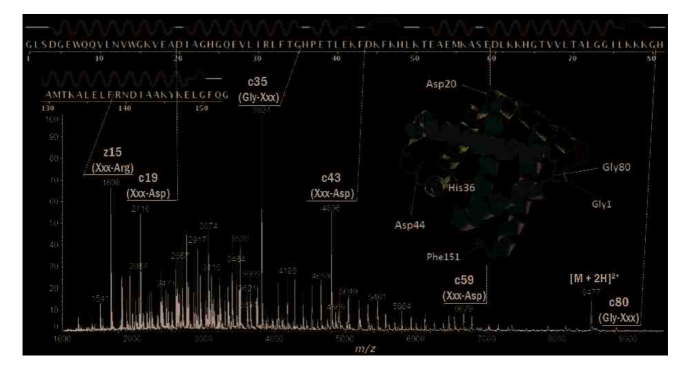
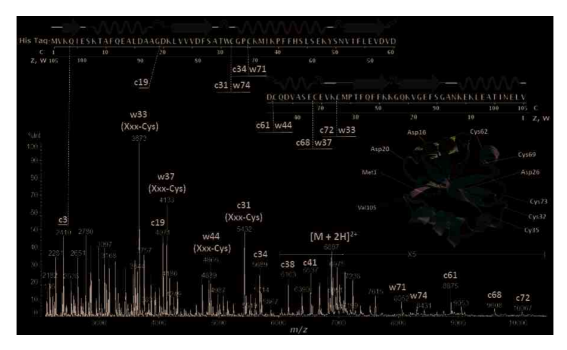


Figure 2. Positive ion MALDI-ISD spectrum of equine apo-myoglobin (Mr 16951.4) obtained with 5-amino-1-naphthol matrix. The inset represents tertiary structure obtained by X-ray crystallography (PDB: 2FRF). The upper line represents primary and secondary structure (wave: α -helix, convex: turn, straight: bend or empty). Adapted from [18] with permissions from Springer, copyright 2014.



It is important to note here that the peak abundance of the ISD fragment ions is dependent upon two factors: (A) the preferential cleavage at the N-Cα bond of the peptide backbone; and (B) the presence of basic residues (Arg, His, Lys and *N*-terminus) or acidic residues (Glu, Asp and *C*-terminus) to give positive or negative ions. The factor B refers to the ionization efficiency *I* (ions/molecules) for the formation of analyte and fragment ions. It is well known that the presence of basic amino acid residues (Arg, Lys, His) and the *N*-terminus generally increases the abundance of the analyte and fragment ions [26,27]. A very intense peak corresponding to *C*-terminal side ISD ion z'15 in Figure 2 can be explained by the appearance of Arg15 residue from *C*-terminus. In the ISD spectrum of thioredoxin (Figure 3), the *N*-terminal side intense c3 ion corresponding to the HisTag-MVK (Met-Val-Lys) may be due to the presence of the Lys3 residue. However, other discontinuous intense peaks observed in Figures 1–3 cannot be explained by the charge sites (factor B).

Figure 3. Positive ion MALDI-ISD spectrum of human thioredoxin with HisTag (Mr 13,769.6) obtained with 5-amino-1-naphthol matrix. The inset represents tertiary structure obtained by X-ray crystallography (PDB (protein data bank): 1AUC). The upper line represents primary and secondary structure (wave: α-helix, arrow: β-strand, convex: turn, straight: bend or empty).



With respect to factor A, the access of active-hydrogens of matrix molecules to carbonyl oxygens of the backbone is influenced by some of the physical characteristics of a given protein, such as intra-molecular hydrogen-bonded secondary structures (α -helix and β -sheet) and bulky side chains resulting in steric hindrance that inhibits the access of the hydrogen-radicals. It may be expected that the presence of α -helix and β -sheet and the bulky side chains may inhibit any preliminary hydrogen-bonding between matrix active-hydrogens and backbone carbonyl oxygens. Although it is difficult to estimate the presence of α-helix and β-sheet in protein molecules embedded in/on the MALDI matrix crystals, the observation of remarkably discontinuous intense peaks for c15 and c33 ions in the ISD spectrum of BSA (Figure 1) strongly suggests that the helix segments as determined by X-ray crystallography are maintained, even when BSA molecules are lying in the matrix crystal. However, the observation of a lot of c-ions (m/z 1000–5000) indicates that the helix segments are incompletely maintained in the matrix crystal phase. Furthermore, the ISD spectrum of myoglobin also showed discontinuous intense peaks of c19, c35, c43 and c80 originating from cleavage at the N-Cα bond of Xxx-Asp or Gly-Xxx residues lying in helix-free regions. Recently, we reported that positive ion collision-induced dissociation (CID) of c35 and c43 ions generated from MALDI-ISD resulted in a remarkable product ion y23 originating from the cleavage at the peptide bond between Asp20-Ile21 residues lying in a helix-free region between two long helical domains [18], while the negative ion CID did not produce any readable product ions. This suggested that gas-phase fragment ions c35 and c43 generated from the MALDI-ISD of myoglobin maintain α-helix structure. It should be noted here that the secondary structures determined by X-ray and NMR of proteins (myoglobin (PDB: 2FRF (X-ray), 1MYF (NMR)), cytochrome c (PDB: 1HRC (X-ray),

1AKK (NMR)) and thioredoxin (PDB: 1AUC (X-ray), 3TRX (NMR))) are similar to each other. This indicates that secondary structures of the proteins are energetically stable in both crystal and solution phases. From this, it may be deduced that protein molecules embedded in/on matrix crystals partly maintain their secondary structures as determined by X-ray and NMR. Regarding this, here we have to pay attention to the denaturation of protein molecules which might be caused through solution preparation and crystallization. The denaturation of protein molecules would occur via two processes such as unfolding of native tertiary structures and destroy of secondary structures made of intra-molecular hydrogen bonding. The excess amounts of solvent and matrix molecules in MALDI sample solutions might cause the denaturation via inter-molecular interactions. It should be noted however that relatively long α-helix structures made of multi-hydrogen bondings are stable by a cooperative effect even when acids were added [28]. So, it seems to be reasonable to assume that protein molecules embedded in/on MALDI matrix crystals partly maintain their secondary structures.

Thioredoxin contains five Cys residues which do not form disulfide bonds. The ISD spectrum showed discontinuous intense peaks of c3 (Lys-Xxx), c19 (Gly-Xxx), c25 (Xxx-Asp), c31 (Xxx-Cys), c34 (Xxx-Cys), c61 (Xxx-Cys), c68 (Xxx-Cys), c72 (Xxx-Cys), w33 (Xxx-Cys), w37 (Xxx-Cys), w44 (Xxx-Cys), w71 (Xxx-Cys) and w74 (Xxx-Cys). The most notable characteristic in the ISD of thioredoxin is cleavage at the N-Cα bond of Xxx-Cys residues to generate w-ions accompanied by the loss of the SH (thiol) group from the side chain (Scheme 1b). The Cys residues in thioredoxin lie in the helix- and sheet-free regions, with the exception of Cys35 (upper in Figure 3). The one-side preferential cleavage at the N-terminal side N-Cα bond of Cys residues (Xxx-Cys) is also observed in the ISD spectrum of BSA (Figure 1). This suggests that the N-terminal side of Cys (CO-NH-Cys) exposes its carbonyl oxygen to the matrix active-hydrogens more than that on the C-terminal side of Cys residues (Cys-CO). A further characteristic of the ISD of thioredoxin is the formation of a c/w-ion pair originating from N-Cα bond cleavage of Xxx-Cys residues, i.e., c31/w74, c34/w71, c61/w44, c68/w37 and c72/w33. The sulfur atom has a capability of the binding of electron or unpaired electron, and the Cys residue tends to appear in turns and to disrupt α -helices. It may therefore be expected that Cys residues expose the adjacent backbone region to matrix molecules resulting in preliminary hydrogen-bonding between backbone carbonyl oxygens or the sulfur atom and matrix active-hydrogens in the MALDI matrix crystal embedded protein molecules.

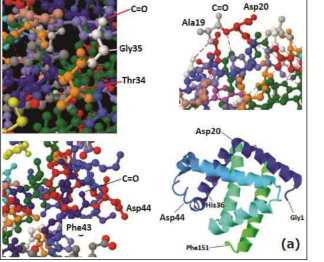
2.3. Accessibility of Carbonyl Oxygens of the Backbone to Hydrogen-Radicals

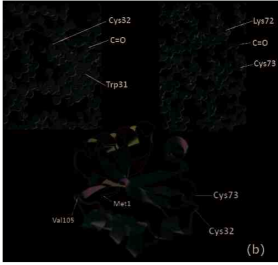
For the one-side preferential N-C α bond cleavage that leads to the formation of abundant c-, z'- and w-ions, it can be considered that the hydrogen radicals or matrix active-hydrogens interact with the *N*-terminal side carbonyl oxygens of Xxx-Asp/Asn/Cys and the *C*-terminal side carbonyl oxygens of Gly-Xxx residues. This indicates that those carbonyl oxygens are free from intra-molecular hydrogen bonded secondary structures such as α -helix and β -sheet. With respect to this, it is important to take into account the secondary structure preference of amino acids [11,12]. According to the report of Chou *et al.* [11], Asn, Gly, Pro, Asp, Ser and Cys residues are preferred in turn structure. In particular, Asn, Cys and Asp residues appear at the start of β -turn regions with high probability. It should be noted further that Gly, Cys, Ser, Pro and Tyr residues tend to destroy helix structure. The preferential cleavage of Xxx-Asp, Xxx-Asn, Xxx-Cys and Gly-Xxx residues as

ascertained here is consistent with the criteria for turn preference of amino acids [11,12], with the exception of the Pro residue. The Pro residue gives incomplete cleavage in ISD, because of its ring structure. As expected, the turn regions lie on the surface of protein molecules, so that the turn preferred residues tend to expose their backbone and side-chains to the environment including matrix molecules in MALDI crystal phase. In other words, the turn preferred residues are highly accessible to the matrix molecules in MALDI crystal phase. Therefore, the one-side preferential N-Cα bond cleavage of Xxx-Asp/Asn/Cys and Gly-Xxx residues in the MALDI-ISD experiments here may be due to high accessibility of the carbonyl oxygens located in the Xxx regions to hydrogen-radicals or matrix active-hydrogens.

To help illustrate the point Figure 4 shows the sites of carbonyl groups at Xxx-Asp20, Xxx-Asp44 and Gly35-Xxx residues obtained from the X-ray data of myoglobin (PDB:2FRF). The carbonyl groups shown here have oxygen atoms exposed to the surroundings without intra-molecular hydrogen bonding. The carbonyl groups at Xxx-Cys32 and Xxx-Cys73 residues of thioredoxin (PDB: 1AUC) also have their oxygen atoms exposed to the surroundings (Figure 4b).

Figure 4. The sites of carbonyl groups at (a) Xxx-Asp20, Xxx-Asp44 and Gly35-Xxx residues obtained from the X-ray crystallography structure of myoglobin (PDB: 2FRF); (b) Xxx-Cys32 and Xxx-Cys73 residues of thioredoxin (PDB: 1AUC).





It is of importance to point out here that Ile and Val residues are more insensitive to ISD than the other amino acids. The peak abundance of c22 (Xxx-Val) and c24 (Xxx-Ile) in Figure 1 and c4 (Xxx-Ile) and c22-c24 (Xxx-Val23-Val24) in Figure 3 is very much lower than that of other peaks. This indicates that the *N*-terminal side carbonyl oxygen of the Ile and Val residues are prevented from forming interactions with matrix molecules. This inhibition may be due to a steric hindrance of bulky side chains of the Ile and Val residues (Scheme 4). It would be expected that such steric hindrance would result in low accessibility of hydrogen-radicals for the carbonyl oxygens of the backbone. This indicates that the access of hydrogen-radicals or matrix active-hydrogens to the backbone carbonyl oxygens of protein molecules is a principal factor in determining the peak abundance of ISD fragment ions.

Scheme 4. Steric hindrance by Ile and Val residues to form preliminary hydrogen-Bonding between matrix active-hydrogens and carbonyl oxygens of the backbone. The arrow represents hydrogen-bonding.

2.4. Discussion

Although it is difficult to estimate the secondary structure and conformation of protein molecules embedded in/on MALDI matrix crystals, the high susceptibility of Asp, Asn, Cys and Gly residues and the low susceptibility of Ile and Val residues to ISD may be rationalized from the standpoint of the accessibility of the peptide backbone by hydrogen-radicals or matrix active-hydrogens. The high susceptibility of Asp, Asn, Cys and Gly residues is attributable to the high accessibility of the backbone by hydrogen-radicals or matrix active-hydrogens in flexible turn regions, while the low susceptibility of Ile and Val residues can be attributed to steric hindrance which prevents the backbone carbonyl oxygens from interactions with matrix molecules.

We can compare the flexible amino acid residues as judged by means of the hydrogen-radical accessibility as presented here with those estimated by other factors representing the flexibility of proteins, such as the B-factor, the turn preference factor, the protection factor and the fluorescence decay factor, as summarized in Table 2. It is of interest that Asp, Asn and Gly residues are identified as flexible by all the measures of flexibility despite the different principles used in the different measures. It is especially, noteworthy that Asp, Asn, Gly and Cys residues are common residues as identified by turn preference by X-ray data and hydrogen-radical accessibility by MALD-ISD data. This suggests that protein molecules embedded in/on matrix crystals partly maintain their secondary structures such as α -helix, and that the hydrogen-radical access to the backbone carbonyl oxygens could be adopted as a factor for measuring protein flexibility.

Table 2. Amino Acid Residues estimated from the B-Factor, the Turn Preference Factor, the Protection Factor, the Fluorescence Factor and the Hydrogen-Radical Accessibility Factor that represent Flexibility of Protein.

Factor for Protein Flexibility	Flexible Amino Acid Residue
B-factor [10]	Asp, Asn, Gly, Pro, Lys, Glu, Gln, Ser
Turn preference [11]	Asp, Asn, Gly, Cys, Pro, Ser
Protection [13]	Asp, Asn, Gly, Lys, Thr, Ile, Met
Fluorescence decay [14]	Asp, Asn, Gly, Ser, Ala
Hydrogen-radical accessibility	Asp, Asn, Gly, Cys

3. Materials and Methods

3.1. Chemicals

The matrix material 5-amino-1-naphthol (5,1-ANL) was purchased from Tokyo Chemical Industry (Tokyo, Japan). Acetonitrile was purchased from Wako Pure Chemicals (Osaka, Japan). Water used in all experiments was purified using a MilliQ water purification system from Millipore (Billerica, MA, USA). Bovine serum albumin (*M*r 66,430.3), equine apo-myoglobin (*M*r 16,951.4) and human thioredoxin with His-Tag (GSSHHHHHHHSSGLVPRGSH) (*M*r 13,769.6) were purchased from Sigma-Aldrich (St. Louis, MO, USA). All reagents were used without further purification.

3.2. Mass Spectrometry

MALDI-TOF mass spectra were acquired on a time-of-flight mass spectrometer AXIMA-CFR (Shimadzu, Kyoto, Japan) equipped with a nitrogen laser (337 nm wavelength) operating at a pulse rate of 10 Hz. The pulse width of the laser was 4 ns. The laser spot size on the target substrate was ca. 100 µm in diameter. The ions generated by MALDI were accelerated using 20 kV with delayed extraction. The analyzer was operated in linear mode and the ions were detected using a secondary electron multiplier. A total of 500 shots were accumulated for each mass spectrum acquisition. The reproducibility of all ISD spectra were confirmed by the peak intensity patterns for several runs using the raster function installed on the AXIMA-CFR mass spectrometer.

3.3. Sample Preparation

Analyte for the MALDI-TOF MS experiments was dissolved in water at a concentration of 20 pmol/ μ L. The matrix material was dissolved in water/acetonitrile (3:7, ν/ν). The matrix and analyte solutions were prepared without any additives such as acetic acid. A sample solution was prepared by mixing a volume of 10 μ L of analyte solution with a volume of 10 μ L of matrix solution. A volume of 1.0 μ L of the sample solution was deposited onto a stainless-steel MALDI plate and the solvents were removed by allowing evaporation in air at room temperature.

3.4. Ab Initio Calculations

The initial structures of the ISD fragments, *i.e.*, closed shell fragments c and z and open shell radical fragments c• and z• for model tri-peptide Me-Ala-X-Ala-Me (X = Ala, Gly, Asp), were made by a CS Chem3D Ultra (CambridgeSoft, Cambridge, MA, USA). The input files were minimized under semi-empirical MO (PM7) run through Winmoster interface using MOPAC2010. The output files were minimized under *ab initio* methods with the DFT UB3LYP level of theory and 6-31 + G (d) basis set. All *ab initio* calculations were performed with the Gaussian 09 suite of programs installed in an HP EliteBook 8460w (Nippon Hewlett-Packard, Tokyo, Japan) with an i7-2630 QM CPU Q720@1.60GHz (Intel, CA, USA). For every optimized structure vibrational analysis was performed and the heat of formation ΔH and Gibbs free energy ΔG were obtained.

4. Conclusions

Although it had been believed that mass spectrometry (MS) is in a disadvantageous position for obtaining information about flexibility or secondary and tertiary structures of protein molecules, here we present a factor for estimating the flexible amino acid residues in intact proteins by using a method of in-source decay (ISD) coupled with MALDI MS. The discontinuous intense peaks observed in MALDI-ISD spectra of BSA, myoglobin and thioredoxin originate from one-side preferential cleavage at the N-Cα bond of Xxx-Asp, Xxx-Asp, Xxx-Cys and Gly-Xxx residues lying in turn regions determined by X-ray crystallography. The flexible residues Asp, Asn, Cys and Gly estimated with MALDI-ISD were compared with the residues estimated from other measures of protein flexibility, such as the B-factor, turn preference, protection and fluorescence decay factors. The comparison shows that Asp, Asn and Gly residues were common to all the measures, and that Asp, Asn, Cys and Gly residues were identified by the ISD based method described here and by turn preference factor based on the X-ray crystallography. The results obtained suggest that protein molecules embedded in/on MALDI matrix crystals partly maintain the α-helix. Furthermore, the reasons that Asp, Asn, Cys and Gly residues are more susceptible and Ile and Val residues less susceptible to ISD can be rationalized in terms of the accessibility of matrix hydrogen-radicals to the backbone carbonyl oxygens. This implies that the hydrogen-radical accessibility in the MALDI-ISD experiments could be adopted as a factor for measuring protein flexibility.

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Author Contributions

M.T.—Designed and interpreted all experimental data, prepared the final manuscript, K.N.—Performed MS experiments and analyzed MS spectra, R.I.—Performed ab initio calculations, K.I.—Performed MS experiments. All authors read and approved the final manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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Legionella dumoffii Utilizes Exogenous Choline for Phosphatidylcholine Synthesis

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Abstract: Phosphatidycholine (PC) is the major membrane-forming phospholipid in eukaryotes but it has been found in only a limited number of prokaryotes. Bacteria synthesize PC via the phospholipid N-methylation pathway (Pmt) or via the phosphatidylcholine synthase pathway (Pcs) or both. Here, we demonstrated that Legionella dumoffii has the ability to utilize exogenous choline for phosphatidylcholine (PC) synthesis when bacteria grow in the presence of choline. The Pcs seems to be a primary pathway for synthesis of this phospholipid in L. dumoffii. Structurally different PC species were distributed in the outer and inner membranes. As shown by the LC/ESI-MS analyses, PC15:0/15:0, PC16:0/15:0, and PC17:0/17:1 were identified in the outer membrane and PC14:0/16:0, PC16:0/17:1, and PC20:0/15:0 in the inner membrane. L. dumoffii pcsA gene encoding phosphatidylcholine synthase revealed the highest sequence identity to pcsA of L. bozemanae (82%) and L. longbeachae (81%) and lower identity to pcsA of L. drancourtii (78%) and L. pneumophila (71%). The level of TNF-α in THP1-differentiated cells induced by live and temperature-killed L. dumoffii cultured on a medium supplemented with choline was assessed. Live L. dumoffii bacteria cultured on the choline-supplemented medium induced TNF-α three-fold less efficiently than cells grown on the non-supplemented medium. There is an evident effect of PC modification, which impairs the macrophage inflammatory response.

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1. Introduction

Legionella are Gram-negative bacilli that are highly successful in colonizing natural and artificial aquatic environments. Dissemination in the environment is facilitated by their characteristic biphasic lifestyle; Legionella may adapt to distinct intracellular and aquatic environments by alternating between a "replicative" and a "virulent" form in response to growth conditions. In water systems, Legionella infects and replicates within protozoa, colonize surfaces, and grow in biofilms [1]. Bacteria enclosed in water-air aerosol are inhaled into the lower respiratory tract and subsequently engulfed by enteric pulmonary macrophages. The capability of Legionella of intracellular proliferation in immune cells designed to kill bacteria and using them as their host cell is crucial for development of pneumonia known as Legionnaires' disease. Currently, the family Legionellaceae is composed of 58 species isolated from environmental sources, but 21 of them have been isolated from humans [2,3]. Among Legionella species that cause human pneumonia, L. pneumophila is the most common causative agent, while L. dumoffii is the fourth [4]. Pneumonia caused by L. dumoffii is rapidly progressive and fulminant owing to the ability of this bacterium to invade and proliferate in

human alveolar epithelial cells [5]. The disease is often fatal, especially in immunocompromised patients. *L. dumoffii* has been isolated from pericarditis, prosthetic valve endocarditis, and septic arthritis, which indicates that *L. dumoffii* is responsible for extrapulmonary infections [6,7].

The way of bacterial penetration into the host cell and factors indispensable for settling the specific microniche, i.e., the digestive vacuole, are dependent on virulence factors released from the cell, unique properties of surface components, and the ability to utilize host metabolites. Crucially for the biogenesis and maintenance of the bacterial replicative vacuole, L. pneumophila uses a type IV secretion system (Dot/Icm) to deliver a large number of effector proteins to the host cell [8]. The coordinate actions of the bacterial effectors allow Legionella to subvert innate immune response and evade host destruction. Moreover, the components of the cell envelope: proteins, peptidoglycan, lipopolysaccharide (LPS), and phospholipids participate in the highly specific Legionella-host interactions. Legionella phospholipids are characterized by a high phosphatidylcholine (PC) content, which is untypical of bacteria and specific for a narrow group of pathogenic and symbiotic microbes whose life cycle is strictly associated with eukaryotic cells. PC is a major component of eukaryotic cell membranes and plays a significant role in signal transduction. The high PC content in intracellular membranes of pathogens, such as Legionella, makes the cells of the microbes similar to the host cells. Bacteria synthesize PC via two different routes, i.e., the phospholipid N-methylation (Pmt) or the phosphatidylcholine synthase (Pcs) pathway. In the Pmt pathway, phosphatidylethanolamine is methylated three times to yield PC in reactions catalyzed by one or several phospholipid N-methyltransferases (PMTs). In the Pcs pathway, choline is condensed directly with CDP-diacylglyceride to form PC in a reaction catalyzed by a bacterium-specific Pcs enzyme [9]. Since choline is not a biosynthetic product of prokaryotes, the Pcs pathway is probably a direct sensor of environmental conditions, using choline availability as an indicator of the status of the location in which the bacterium is found. It has been shown that L. pneumophila and L. bozemanae are capable of utilisation of exogenous choline for PC synthesis [10,11]. Apart from the structural role, the exact function of PC in Legionella cells remains unexplained. However, PC-deficient mutants of L. pneumophila exhibited attenuated virulence and increased susceptibility to macrophage-mediated killing. These defects were attributed to reduced bacterial binding to macrophages and a poorly functioning Dot/Icm system. In the process of binding to macrophages, L. pneumophila uses the platelet-activating factor receptor (PAF receptor), which harbors the same glycerophosphocholine head group as PC. Due to this structural similarity, PC is required for efficient binding of L. pneumophila to macrophages via the PAF receptor [12].

In response to infection caused by *Legionella*, macrophages produce inflammatory cytokines, such as interleukin 6 (IL-6), interleukin 1α (IL- 1α), interleukin 1β (IL- 1β), interleukin 12 (IL-12), interferon γ (INF γ), and tumor necrosis factor α (TNF- α) [13]. Among these cytokines, TNF- α appears to be pivotal for activation of phagocytes and resolution of pneumonic infection [14,15]. Treatment of rat alveolar macrophages with TNF- α resulted in decreased intracellular growth of *L. pneumophila* [16]. In turn, inhibition of endogenous TNF- α activity via TNF- α -neutralizing antibodies resulted in enhanced growth of *L. pneumophila* in the mouse lung [17]. However, little is known about how the pathogen PC influences the induction of proinflammatory cytokines in the host.

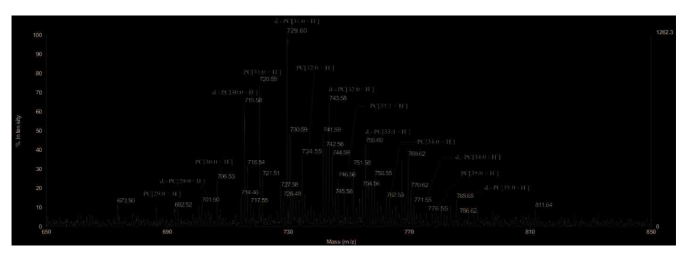
The aim of our study was to investigate the ability of L. dumoffii to utilize exogenous choline for PC synthesis using Matrix-Assisted Laser-Desorption/Ionization (MALDI)-Time of Flight (TOF) MALDI/TOF and Liquid Chromatography Coupled with the Mass Spectrometry Technique Using the Electrospray Ionization Technique (LC/ESI-MS) techniques. We also wanted to determine whether the bacteria use the Pcs pathway for PC synthesis by identification of a pcsA gene encoding phosphatidylcholine synthase. Next, the correlations between the PC species content in the L. dumoffii membranes and the level of TNF- α produced by human macrophages were investigated.

2. Results

2.1. Utilization of Exogenous Choline by L. dumoffii

In order to study the ability of *L. dumoffii* to use exogenous choline for PC synthesis, the bacteria were grown on BCYE medium supplemented with deuterium-labeled choline. Lipids isolated from the cells were analyzed using MALDI-TOF mass spectrometry with a CHCA matrix (in the reflectron mode). The positive ionization MALDI-TOF spectrum contained a cluster of molecular ions at *m/z*: 692.52–785.65 corresponding to PC species (Figure 1). Within the cluster, protonated ions were identified at *m/z* 692.52 PC[29:0 + H]⁺, 706.53 PC[30:0 + H]⁺, 720.55 PC[31:0 + H]⁺, 734.55 PC[32:0 + H]⁺, 746.56 PC[33:1 + H]⁺, 762.53 PC[34:0 + H]⁺, 776.55 PC[35:0 + H]⁺, and corresponding ions with mass higher by 9 at *m/z* 701.50 d₉-PC[29:0 + H]⁺, 715.58 d₉-PC[30:0 + H]⁺, 729.60 d₉-PC[31:0 + H]⁺, 743.58 d₉-PC[32:0 + H]⁺, 755.60 d₉-PC[33:1 + H]⁺, 771.55 d₉-PC[34:0 + H]⁺, 785.65 d₉-PC[35:0 + H]⁺. Deuterium-labeled PC[31:0 + H]⁺ was observed as the dominant constituent of the extract. The presence of labeled PCs in the MALDI-TOF spectrum showed that *L. dumoffii* was able to utilize exogenous choline for PC synthesis.

Figure 1. Positive ion mode MALDI-TOF spectrum of the PCs of *L. dumoffii* cultured on the medium with labeled choline: m/z = 692.52-785.65. All peaks were marked according to their m/z ratio.

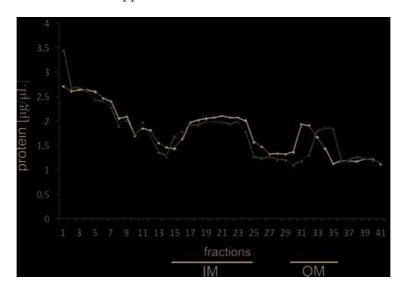


2.2. Membrane Localization of PC

2.2.1. Membrane Fractionation

To determine PC localization in the *L. dumoffii* membranes, separation of outer and inner membranes from bacteria cultured with and without labeled choline was performed by sucrose density gradient ultracentrifugation. The protein concentration in the individual fractions showed two major peaks with maxima in fractions 20–22, 31, and 32 for the choline non-supplemented bacteria. In the case of choline-supplemented bacteria, the maxima were detected in fractions 19–23 and 33–35 (Figure 2).

Figure 2. Separation of *L. dumoffii* inner (IM) and outer membrane (OM) by discontinuous sucrose density gradient centrifugation. Fractions of 1 mL were collected from the top of the gradient and assayed for the presence of protein ($\mu g/\mu L$). Black line—for bacteria cultured on medium non-supplemented with choline; Blue line—for bacteria cultured on medium supplemented with choline.



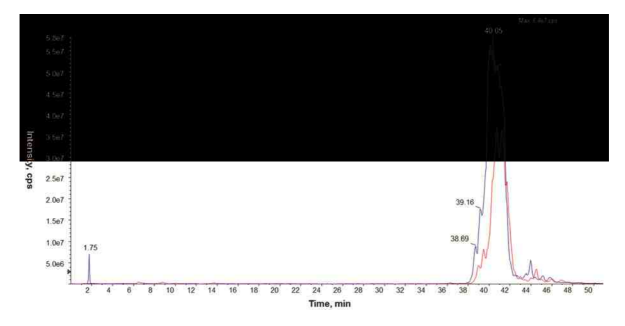
To assign the peaks to a membrane compartment, NADH oxidase and esterase activities (as characteristic for the inner and outer membrane, respectively) were determined. NADH oxidase activity was concentrated in pooled fractions corresponding to the inner membrane (fractions 16–24 for choline non-supplemented bacteria and 15–24 for bacteria cultured with choline). NADH oxidase activity was 228 μmol min⁻¹·mL⁻¹. Esterase activity was concentrated in pooled fractions represented by the outer membrane (30–34 for bacteria cultured without choline and 32–35 for choline-supplemented bacteria). The NADH oxidase and esterase activities detected confirmed the efficiency of fractionation.

2.2.2. Localization of PC in the Outer and Inner Membranes by LC/ESI-MS Analysis

In the *L. dumoffii* membranes (inner and outer), unlabeled and labeled PC species were identified by the LC/ESI-MS technique using the precursor ion mode (PI) and neutral loss scan (NL) in the positive ion mode. The characteristic fragment of PC compounds at m/z 184 corresponding to the

polar head group ($C_5H_{15}NPO_4^+$) was employed to determine unlabeled PC using the PI mode, while d₉-PC compounds were identified by a diagnostic ion of the head group at m/z 193 (Figure 3).

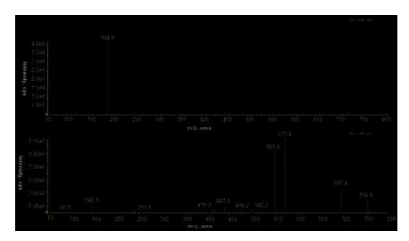
Figure 3. Chromatogram MS obtained with the Precursor Ion Mode for unlabeled (blue; m/z 184) and labeled (red; m/z 193) PC molecular species, recorded under NP LC/MS conditions (phase A—hexane/isopropanol (3:2, v/v) and phase B—an isopropanol/hexane/5 mM aqua solution of ammonium acetate (38:56:5, v/v/v). The following elution program was employed: from 53% B to 80% B for 23 min, 80% B maintained for 4 min, 80% B to 100% B for 9 min, and 100% B maintained for 14 min. The flow rate was 1 mL/min) in the positive ion MS mode; the lipid extract was isolated from the inner cell membrane of L. dumoffii cultured on labeled choline.



In the MS spectra of the analyzed PC compounds, $[M + H]^+$ and $[M + Na]^+$ ions were observed in the positive ion mode while in the negative ion mode—peaks corresponding to $[M + CH_3COO]^-$ adducts.

Fragmentation of the PC standard (1,2-dipalmitoyl-sn-glycero-3-phosphocholine; Figure 4) showed that the fragmentation of the sodiated PC molecule gave more information about the structure of the investigated compound. In the CID spectrum of the $[M + Na]^+$ ion of the PC standard peaks at m/z 441.0, 478.2, and 500.2 reflected the presence of palmitic acid in the PC molecule.

Figure 4. CID spectrum of the PC standard (1,2-dipalmitoyl-sn-glycero-3-phosphocholine) of protonated (top) and sodiated (below) molecules, recorded under NP LC/MS conditions (phase A—hexane/isopropanol (3:2, v/v) and phase B—an isopropanol/hexane/5 mM aqua solution of ammonium acetate (38:56:5, v/v/v). The following elution program was employed: from 53% B to 80% B for 23 min, 80% B maintained for 4 min, 80% B to 100% B for 9 min and 100% B maintained for 14 min. The flow rate was 1 mL/min) in the positive ion MS mode.

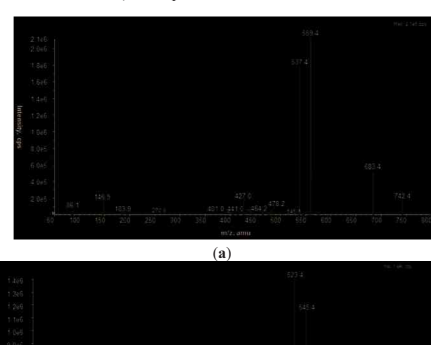


Next, neutral loss scanning was applied as an alternative method for the precursor ion mode to identify sodiated PC species in the mixture of lipids isolated from *L. dumoffii* membranes. PC compounds were found in the positive ion mode by the loss of the polar head group with a molecular weight at 183 Da (unlabeled) and 192 Da (d9-labeled). According to Hsu and Turk, this phenomenon occurs in two steps through elimination of trimethylamine (59 Da), leading to the loss of C5H14NPO4 (183 Da) from the PC structure [18]. Using the Precursor Ion and Neutral Loss scanning method, numerous PC species were found in the lipid extract obtained from *L. dumoffii* cell membranes. The PI mode seems to be more sensitive than the NL scanning technique for both unlabeled and labeled PC species. On the other hand, the NL method facilitated identification of sodiated PC molecular species in the mixture, which gave more informative CID spectra in the positive ion mode in the fragmentation process.

The presence of unlabeled and labeled PC molecules in *L. dumoffii* cell membranes was also confirmed by the CID spectra of sodiated PC ions obtained in the positive ion mode. For bacteria cultured on the medium without labeled choline, the major PCs with the following composition of fatty acids: 16:0/15:0, 17:0/15:0, and 16:0/17:1 (or cyclic 17:0, indistinguishable under the present mass spectrometry conditions) were determined in the inner membrane on the basis of the fragmentation spectra of peaks at m/z 742.4, 756.4, and 768.4 (Figure 5a). In the outer membrane, the major species of PC were diacyl 15:0/15:0, 14:0/16:0 (m/z 728.4), 16:0/17:1 (or cyclic 17:0) (m/z 768.8), and 17:0/17:1 (or cyclic 17:0) (m/z 782.8) (Figure 5b). In the CID spectrum of 16:0/15:0 PC shown in Figure 5a, loss of trimethylamine (59 Da) was observed. The peaks at m/z 559.4 and 537.4 correspond to the elimination of the PC head group giving ions [M + Na - 183]⁺ and [M + Na - 205]⁺, respectively. The fragment ions reflecting the presence of pentadecanoic (15:0) and palmitic (16:0) acids in the PC structure were found at m/z 405.2, 427.2, 464.2, 478.2, and 500.3.

The fatty acids were eliminated as neutral molecules (m/z 427.2) or as sodium salts (m/z 405.2, 464.2, 478.2) from the $[M + Na]^+$ and $[M + Na - N(CH_3)_3]^+$ ions in the fragmentation pathway of PC molecular species in the positive ion mode.

Figure 5. (a) CID spectrum of unlabeled 16:0/15:0 PC molecular species identified in the lipid mixture isolated from the inner membrane (*L. dumoffii* cultivated on the medium without labeled choline), recorded under NP LC/MS conditions (phase A—hexane/isopropanol (3:2, v/v) and phase B—an isopropanol/hexane/5 mM aqua solution of ammonium acetate (38:56:5, v/v/v). The following elution program was employed: from 53% B to 80% B for 23 min, 80% B maintained for 4 min, 80% B to 100% B for 9 min and 100% B maintained for 14 min. The flow rate was 1 mL/min) in the positive ion MS mode; (b) CID spectrum of unlabeled PC molecular species with FA combination: 14:0/16:0 and 15:0/15:0 identified in the lipid mixture isolated from the outer membrane (*L. dumoffii* cultivated on the medium without labeled choline), recorded under NP LC/MS conditions (phase A—hexane/isopropanol (3:2, v/v) and phase B—an isopropanol/hexane/5 mM aqua solution of ammonium acetate (38:56:5, v/v/v). The following elution program was employed: from 53% B to 80% B for 23 min, 80% B maintained for 4 min, 80% B to 100% B for 9 min and 100% B maintained for 14 min. The flow rate was 1 mL/min) in the positive ion MS mode.



In the case of bacteria cultured on the medium with labeled choline, d₉-labeled PC molecular species were identified in the inner and outer membranes and their structures were established as shown in Table 1.

Table 1. The major d₉-labeled phosphatidycholine (PC) species determined by Liquid Chromatography Coupled with the Mass Spectrometry Technique Using the Electrospray Ionization Technique (LC/ESI-MS) in the inner and outer membrane of *L. dumoffii* cells.

/-	PC structure, Sn-1/Sn-2 ^a		
m/z	Inner membrane	Outer membrane	
737.8	d ₉ -PC15:0/15:0	d ₉ -PC15:0/15:0	
	d ₉ -PC14:0/16:0		
751.9	-	d9-PC16:0/15:0	
777.8	d ₉ -PC16:0/17:1	-	
791.9	d ₉ -PC17:0/17:1	d9-PC17:0/17:1	
807.9	d ₉ -PC20:0/15:0	-	
819.9	d ₉ -PC20:0/16:1	-	

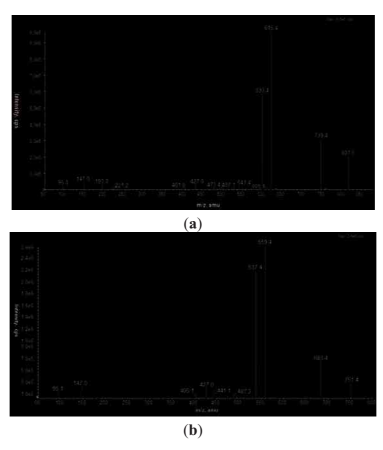
^a 17:1 or cyclic 17:0, indistinguishable under the present mass spectrometry conditions.

Figure 6 presents the fragmentation spectra of individual d₉-labeled PC molecular species found in the inner and outer cell membranes recorded in the positive ion mode. In Figure 6a, the peaks at m/z 739.4, 615.4, and 593.4 corresponding to $[M + Na - N(CD_3)_3]^+$, $[M + Na - 192]^+$, and $[M + Na - 214]^+$ confirmed the presence of the deuterated PC head group in the PC compound. Loss of the 20:0 and 15:0 fatty acids as RCOOH and RCOONa was observed in the CID spectrum. A similar fragmentation process was also observed in the CID spectra shown in Figure 6b, where the 16:0/15:0 PC molecular species was determined. In addition to the labeled PCs, unlabeled PCs m/z 706.9, 720.8, 734.9, 748.7, 762.9, 776.7, and 790.7 were identified in the inner and outer membranes.

To compare the amount of the unlabeled PC phospholipid class in *L. dumoffii* cell membranes (outer and inner), the Multiple Reaction Monitoring (MRM) mass spectrometry mode was employed. The MRM mode is defined as a sensitive and a selective mass spectrometry technique. In the first step of the technique, an ion of interest was selected and subjected to the fragmentation process. Then, only predefined and characteristic fragment ions were detected.

In our studies, the MRM pairs were prepared on the basis of fragmentation spectra of peaks at m/z 764.9 and 778.9 corresponding to the most intense PC signals in the negative ion mode in both cell membranes. The [M + CH₃COO]⁻ ion and the characteristic fragment ions [M + CH₃COO – 74]⁻ and [RCOO]⁻ of PCs obtained in the fragmentation process under the negative ion mode were chosen as MRM data (764.9/690.4, 764.9/241.2, and 778.9/704.4, 778.9/241.2). In this experiment, the MRM peak areas were compared and they showed higher values for the inner cell membranes (for the inner membrane: 764.9/690.4, peak area—1.02 × 10⁶, 764.9/241.2—1.47 × 10⁶; 778.9/704.4—2.08 × 10⁶, 778.9/241.1—3.18 × 10⁶; for the outer membrane: 764.9/690.4—4.85 × 10⁵, 764.9/241.2—6.47 × 10⁵, 778.9/704.4—8.57 × 10⁵, 778.9/241.1—1.31 × 10⁶). This indicated that the inner cell membrane seems to be richer in the PC phospholipid class.

Figure 6. (a) CID spectrum of labeled 20:0/15:0 PC molecular species identified in the lipid mixture isolated from the inner membrane (*L. dumoffii* cultivated on the medium with labeled choline), recorded under NP LC/MS conditions (phase A—hexane/isopropanol (3:2, v/v) and phase B—an isopropanol/hexane/5 mM aqua solution of ammonium acetate (38:56:5, v/v/v). The following elution program was employed: from 53% B to 80% B for 23 min, 80% B maintained for 4 min, 80% B to 100% B for 9 min and 100% B maintained for 14 min. The flow rate was 1 mL/min) in the positive ion MS mode; (b) CID spectrum of labeled 16:0/15:0 PC molecular species identified in the lipid mixture isolated from the outer membrane (*L. dumoffii* cultivated on the medium with labeled choline), recorded under NP LC/MS conditions (phase A—hexane/isopropanol (3:2, v/v) and phase B—an isopropanol/hexane/5 mM aqua solution of ammonium acetate (38:56:5, v/v/v). The following elution program was employed: from 53% B to 80% B for 23 min, 80% B maintained for 4 min, 80% B to 100% B for 9 min and 100% B maintained for 14 min. The flow rate was 1 mL/min) in the positive ion MS mode.

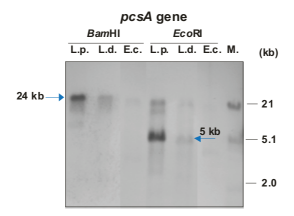


2.3. Identification of the pcsA Gene in the L. dumoffii Genome

Our biochemical analyses concerning PC synthesis in *L. dumoffii* indicated that this compound is produced in a one-step pathway, suggesting that a gene encoding phosphatidylcholine synthase is present in the genome of this bacterium. In order to identify the *pcsA* gene in *L. dumoffii*, Southern hybridization under low-stringency conditions was performed using genomic DNAs from *L. dumoffii*, *L. pneumophila*, and *E. coli* (as a negative control) digested with *Bam*HI and *Eco*RI. A

DIG-labeled DNA fragment containing the *pcsA* of *L. pneumophila* was used as a probe (Figure 7). As a result, a homologue of *pcsA* was found in the *L. dumoffii* genome, although the intensity of detected signals was significantly lower in comparison to the signals obtained for *L. pneumophila*. The *pcsA* probe hybridised to a 24-kb-long *Bam*HI fragment and a 5-kb-long *Eco*RI fragment of *L. dumoffii*. In the case of *L. pneumophila*, strong positive signals were observed for the 24-kb *Bam*HI fragment and the 6-kb *Eco*RI fragment, respectively. On the contrary, only a slight unspecific signal was observed in hybridization with *E. coli* genomic DNA, confirming absence of a *pcsA* homolog in this genome (Figure 7).

Figure 7. Identification of the *pcsA* gene in the *L. dumoffii* genome by Southern hybridization using a probe containing the *pcsA* gene of *L. pneumophila*. Hybridization was performed with genomic DNAs of *L. dumoffii* (L.d.), *L. pneumophila* (L.p.), and *E. coli* (E.c.) digested with *Bam*HI and *Eco*RI as described in Materials and Methods. M—molecular size standard (λ DNA digested with *Hind*III and *Eco*RI).



In conclusion, these results showed that the *L. dumoffii* genome contained a *pcsA* gene; however, the low intensity of the observed signal suggested a low sequence identity between *L. pneumophila pcsA* and *L. dumoffii* homolog. This observation is in agreement with literature data that described high genetic and phenotypic diversity between *L. pneumophila* and other *Legionella* species [19].

To determine the nucleotide sequence of *L. dumoffii pcsA*, a set of degenerate primers was designed based on genomic sequences of a few *L. pneumophila* strains, *L. longbeachae* D-4968 and *L. drancourtii* LLAP12. Using these primers, a 900-bp-long PCR product of *L. dumoffii* was obtained and sequenced. In the assessment of the genetic relatedness between *L. dumoffii pcsA* and this gene in other sequenced *Legionella* species, the highest sequence identity to *L. bozemanae pcsA* (82%) and *L. longbeachae* (81%) was found. *L. dumoffii pcsA* showed lower sequence identity to *pcsA* of *L. drancourtii* (78%) and *L. pneumophila* ATCC 33155 (71%). The comparison of the *pcsA* sequence indicated a high level of diversity among the *Legionella* species, especially in 5' and 3' ends of *pcsA* genes. *L. dumoffii pcsA* encodes putative 254-aa-long phosphatidylcholine synthase, whose length was identical or very similar to the other *Legionella* PcsA proteins: *L. drancourtii* (254 aa), *L. longbeachae* (253 aa), and *L. pneumophila* (255 aa). In the upstream of the *L. dumoffii pcsA* coding sequence (–135 to –86 bp), a promoter sequence of high potential activity (*p* = 0.99) was identified (5'-actatttttatgtttttattttttgataaaacaacaatagtttagccca-3'). In addition, 6 nt upstream of the ATG

translation start codon, a potential ribosome-binding site (AGGA) was found. The PcsA of *L. dumoffii* showed the highest sequence identity to the *L. bozemanae* homolog (90% identity and 96% similarity). Lower identity of this protein was established to PcsA of *L. longbeachae* (87% identity and 94% similarity), *L. drancourtii* (79% identity and 90% similarity), and *L. pneumophila* (74% identity and 84% similarity). The most conservative amino acid sequences were found in the central regions of PcsA proteins that are probably-functional domains of the enzyme, whereas their *N*- and *C*-termini are substantially more divergent.

2.4. The Cytotoxic Effect of L. dumoffii on THP-1 Cells

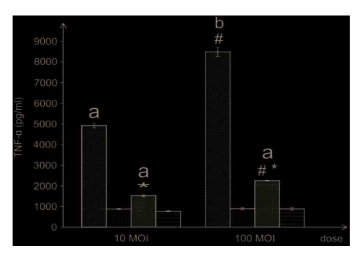
To determine the infectious doses of *L. dumoffii* for human macrophages (THP-1 cell line), the cytotoxic effect of different bacteria concentrations (expressed as the multiplicity of infection—MOI) was assessed with the MTT (3-(4,5 dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide) method. The viability of the THP-1 differentiated cells incubated with *L. dumoffii* for 4 h at a MOI of 10, 50, 100, and 200 was found to be similar to that of the control cells (cell viability $103.05\% \pm 7.4\%$, $105.5\% \pm 8.0\%$, $101.6\% \pm 7.8\%$, $96.2\% \pm 7.2\%$, respectively). After 24-h incubation of THP-1 cells with the bacteria at a MOI of 100 and 200, a slight decrease in cell viability was observed (cell viability $99.2\% \pm 7.2\%$, $95.4\% \pm 7.4\%$, respectively). The analysis of the cytotoxic effect of *L. dumoffii* on THP-1 cells was also performed for the same doses of *L. dumoffii* cultured on choline-supplemented medium. No statistically significant differences of the toxic effect on the THP-1 cells were found between the bacteria cultured with and without choline. Based on these results and literature data [20,21], *L. dumoffii* at a MOI of 10 and 100 were chosen to study the TNF- α induction in the THP-1 differentiated cells.

2.5. TNF-α Induction by L. dumoffii in the THP-1 Differentiated Cells

2.5.1. TNF-α Induction in the THP-1 Differentiated Cells by *L. dumoffii* Cultured on Choline-Supplemented and Non-Supplemented Medium

Levels of human TNF- α in the supernatants from the THP-1 experimental cultures were measured by an enzyme-linked immunosorbent assay (ELISA) after 4-h incubation of cells with live and temperature-killed *L. dumoffii* bacteria at the MOI of 10 or 100. To study the influence of choline on TNF- α production, both live and temperature-treated bacteria were obtained from BCYE medium supplemented with choline and without this compound. Regardless of the culture conditions, dose-dependent production of TNF- α was observed in all the experiments. However, statistically significant differences in obtained results were only noted when we considered the experiments with live bacteria. Bacteria cultured on the choline-supplemented medium, regardless of their concentration, induced TNF- α production at significantly lower level ($p \le 0.05$). However, inside study groups—live bacteria cultured with or without choline supplementation—level of TNF- α depended on bacteria concentration and was significantly higher when 100 MOI bacteria were used. The temperature-killed bacteria induced TNF- α at a considerably statistically significant lower level in comparison to live bacteria and presence of choline in the bacterial culture medium did not change the level of cytokine production (Figure 8).

Figure 8. Effects of live and temperature-treated L. dumoffii cells cultured on the choline-supplemented medium and without choline on the ability to induce TNF- α production in THP-1 differentiated cells. Data represent the mean \pm S.D. of five independent experiments. Control THP-1 differentiated cells (PMA-activated): TNF- α 22.6 \pm 0.6 (pg/mL); Control non-activated THP-1 cells: TNF- α 18.32 \pm 0.9 (pg/mL). # Significantly different from the lower bacteria dose, $p \le 0.05$; * Statistically significant in comparison to bacteria cultured on the choline non-supplemented medium, $p \le 0.05$ (ANOVA and Tukey's post hoc test). Statistically significant in comparison with similar experiments preformed with temperature-treated bacteria ($^a p \le 0.05$, $^b p \le 0.01$) c (ANOVA and Tukey's post hoc test) —live L. dumoffii; —temperature-treated L. dumoffii; —temperature-treated medium; —temperature-treated L. dumoffii cultured on choline-supplemented medium.

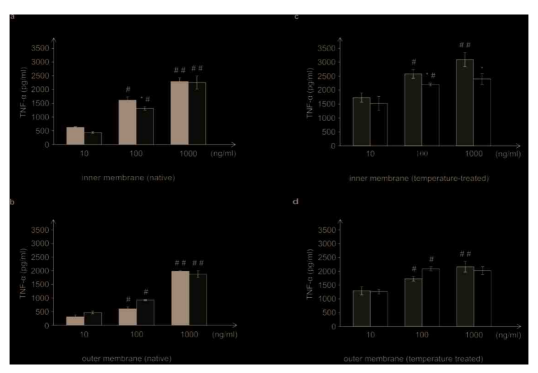


2.5.2. TNF- α Induction by the Outer and Inner Membrane of *L. dumoffii* in the THP-1 Differentiated Cells

Inner and outer membranes (10, 100, and 1000 ng/mL concentrations) isolated from *L. dumoffii* cultured with and without choline were used for TNF- α induction (Figure 9). Both the inner and outer temperature-treated membranes induced TNF- α production in each dose and in a dose-dependent manner more efficiently than the temperature non-treated membranes. A comparison of the ability of the membranes to induce TNF- α showed that the inner membrane induced TNF- α more efficiently than the outer membrane, irrespective of the doses used and temperature treatment or no treatment.

Choline supplementation influenced TNF- α production in a different manner depending on the membrane type. The inner membrane isolated from choline-supplemented bacteria induced TNF- α less efficiently than the inner membrane of bacteria cultured without choline. However, only in the presence of 100 ng/mL of these membranes, the decrease in the TNF- α production was statistically significant (Figure 9A,C). In turn, the same concentration of the outer membranes of choline-supplemented bacteria (both, non- and temperature-treated) caused opposite effect—slightly increased the level of TNF- α in comparison with the outer membranes from bacteria cultured without choline (Figure 9B,D).

Figure 9. TNF- α (pg/mL) induction in the THP-1 differentiated cells by the outer and inner membrane of *L. dumoffii* (**a,b,c,d**). Control THP-1 differentiated cells (PMA-activated): TNF- α 22.6 \pm 0.60 (pg/mL); Control non-activated THP-1 cells: TNF- α 18.32 \pm 0.90 (pg/mL). Data represent the mean \pm S.D. of five independent experiments; #—Significantly different from the lower (10 ng/mL) membrane concentration, $p \le 0.05$; ##—Significantly different from the lower (100 ng/mL) membrane concentration, $p \le 0.05$; *—Statistically significant in comparison to bacteria cultured on the choline non-supplemented medium, $p \le 0.05$ (ANOVA and Tukey's *post hoc* test). ——membrane from bacteria cultured on the choline-supplemented medium. ——membrane from bacteria cultured on the choline-supplemented medium. ——temperature-treated membrane from bacteria cultured on the choline non-supplemented medium. ——temperature-treated membrane from bacteria cultured on the choline non-supplemented medium. ——temperature-treated membrane from bacteria cultured on the choline-supplemented medium. —temperature-treated membrane from bacteria cultured on the choline-supplemented medium.



2.6. In Vitro Infection of Differentiated THP-1 Cells with L. dumoffii

To compare the levels of L. dumoffii internalization by differentiated THP1 cells, bacteria cultured on the medium with and without choline were incubated with macrophages for 2 h. Next, bacteria that were not phagocytised by macrophages were killed by gentamycin treatment. Only intracellular bacteria released after macrophage lysis were seeded onto the BCYE agar medium. The number of colonies was higher for bacteria cultured with choline before the internalization process (171.8 \pm 4.2 for the choline-grown bacteria, 143.2 ± 3.8 for the non-choline grown bacteria). This investigation indicates that choline-grown bacteria undergo association by macrophages more readily than non-choline grown bacteria. However, the results were not statistically significant. L. dumoffii did not replicate in differentiated THP1 cells irrespective of the bacterial culture conditions (data not shown).

3. Discussion

Our previous and current investigations showed that *L. lytica*, *L. bozemanae*, and *L. dumoffii* form PC [11,22]. In this study, the ability of *L. dumoffii* to synthesize PC in a choline-dependent manner was investigated. It was evidenced that the bacteria used exogenous choline for PC synthesis via the Pcs pathway. The main PC d9-PC[31:0 + H]⁺, identified on the MALDI/TOF spectrum and synthesised via this pathway, was confirmed by LC/ESI-MS as PC 16:0/15:0, also identified in *L. bozemanae*. Both species exhibit blue-white fluorescence; and the presence of PC 16:0/15:0 is an additional chemotaxonomic feature that classifies both species as representatives of the genus *Fluoribacter* in the *Legionellaceae* family [23]. The Pcs pathway in *L. dumoffii* was confirmed by identification of the *pcsA* gene encoding the phosphatidylcholine synthase. All *Legionella* spp. genomes characterized so far contain the *pcsA* gene encoding phosphatidylcholine synthase, suggesting a significant function of this protein in their metabolism. Previously, we have identified the *pcsA* gene in *L. bozemanae* and found that PC is effectively produced in a one-step pathway, in which the PcsA enzyme is engaged [11]. Likewise, the *L. dumoffii* PcsA presented in this study and *L. bozemanae* PcsA exhibited the highest sequence amino acids identity to the *L. longbeachae* PcsA, which confirmed genetic relatedness of these three *Legionella* species.

The presence of unlabeled PC in the MALDI-TOF spectrum obtained from lipids of bacteria cultured on choline has indicated that *L. dumoffii* forms PC also by triple PE methylation. LC/ESI-MS analysis of PC species present in both membranes labeled with a deuterated precursor allowed us to distinguish PC species synthesized from the CDP-choline pathway and the PE methylation pathway. Our previous investigations showed that *L. bozemanae* produced PC via two independent pathways PmtA and Pcs [11]. Similarly, both PC synthesis pathways function in *L. pneumophila*. The pathway in which *Legionella* spp. utilize exogenous choline is dominant and seems to be more energetically efficient [10]. However, in the genetic experiments that were conducted (hybridization, PCR analyses using several degenerative primers and amplicon sequencing), we were unable to identify a *pmtA* homologue in the *L. dumoffii* genome (data not shown). Both PC synthesis pathways have also been reported in legume endosymbionts (*Rhizobium leguminosarum*, *Sinorhizobium meliloti*, *Bradyrhizobium japonicum*) and in the plant pathogen *Agrobacterium tumefaciens*. In human pathogens such as *Pseudomonas aeruginosa*, *Brucella melitensis*, and *Borrelia burgdorferi*, PC is produced only via the Pcs pathway [24].

Several studies have shown that the PC of bacterial membranes can be important to host-associated bacteria in pathogenesis and symbiosis. Our results have indicated that bacteria cultured on choline undergo internalization by THP1 macrophages more readily than bacteria cultured on non-choline-supplemented medium. The increase in internalization in the case of bacteria cultured on choline was not statistically significant, which may suggest a low level of participation of PC species in this process, although other papers indicate that some bacterial PC derivatives play a direct role in association with macrophages [25].

PC-deficient mutants of *B. japonicum* and *S. meliloti* were characterized by substantially reduced symbiosis with their plant hosts [26,27]. PC is indispensable for the plant pathogen *A. tumefaciens* to assembly T4SS components, important factors in formation of plant crown-gall tumors [28,29].

A *Brucella abortus pcs* mutant exhibited an altered cell envelope; therefore, it did not establish a replication niche inside the macrophages. Additionally, it showed a severe virulence defect in a murine model of infection [30,31].

A PC-deficient *P. aeruginosa* mutant exhibited the same level of sensitivity to antibiotics and antimicrobial peptides as wild strains. PC deficiency did not change the mobility and capability of biofilm formation on an abiotic surface. However, PC may have a specific role in the interaction with eukaryotic hosts, e.g., it might aid in assembly or localization of specific proteins in *P. aeruginosa* [32].

L. dumoffii cultured on choline-supplemented medium exhibited altered sensitivity to Galleria mellonella antimicrobial defense factors such as defensin and apoLp-III [33]. Replacement of PE with PC induced concurrent structural and functional changes in the ABC multidrug exporter of Lactococcus brevis [34].

The membrane fraction experiment and LC/ESI-MS analyses suggest that PC in *L. dumoffii* is present in the outer and inner membranes. In *L. pneumophila* and *P. aeruginosa*, PC was also localized in both membrane compartments [35,36]. MRM analysis showed that the amount of PC in the inner membrane of *L. dumoffii* was higher than in the outer membrane. Moreover, there were differences in the structure of the PC species present in bacteria cultured with and without choline. These differences might be important for interactions of bacteria with host cells.

Several lines of evidence indicate that PC is able to modulate the inflammatory functions of monocytic cells. Tonks et al. showed that PC 16:0/16:0 significantly inhibited TNF-α release from the human monocytic cell line MonoMac-6 in a dose-dependent manner. In contrast, PC 20:4/16:0 did not reduce TNF-α, which indicated that regulation of the inflammatory response was associated with the composition of fatty acids forming PC [37]. In our study, live L. dumoffii bacteria cultured on the choline-supplemented medium, induced three times less TNF-α than the cells grown on the non-supplemented medium. Similarly, the inner (temperature-treated and -untreated) membranes isolated from bacteria cultured on choline, applied at almost all the doses, induced a decreased level of TNF-α, in comparison to bacteria cultured without choline. Bacteria grown on choline incorporate into their inner membranes PC species with longer fatty acids than bacteria cultured without choline, which may have a significant effect on the lower induction of TNF- α level. In comparison to live bacteria, temperature-treated bacteria induced significantly lower TNF-α level. It is connected with the fact, that in live Legionella HSP, flagellin, and LPS are the main inducer of TNF-α. Temperature heating (90 °C, 20 min) causes HSP and flagellin degradation, therefore still present LPS is only inducer of this cytokine. This may be similar to the case of L. pneumophila LPS [38], although it is not known what the efficiency of L. dumoffii LPS in the induction of TNF-α is, since no such investigations have been carried out and the structure of L. dumoffii LPS is not known.

In the case of the 10- and 100-ng/mL concentrations of the outer membrane of live bacteria cultured on choline, TNF- α induction was higher than in the case of bacteria without choline; however, these differences were not statistically significant. It has been shown that lipopolysaccharide (LPS) of Gram-negative bacteria, a well-known TNF- α inducer located in the outer membrane, had an influence on the level of the cytokine as well. However, *L. pneumophila* LPS is about 1000 times less potent in its ability to induce pro-inflammatory cytokines (TNF- α , IL-1 β , IL-6, IL-8) in Mono Mac 6 cells than the LPS of the *Enterobacteriaceae* members [21].

Cao *et al.* showed that addition of PC18:2/18:2 into the culture of Kupffer cells significantly reduced LPS-stimulated TNF- α generation [39]. We did not observe such an effect. Probably, the PC structure has a significant impact on the level of induction of this cytokine.

4. Experimental Section

4.1. Bacterial Strain and Growth Conditions

L. dumoffii strain ATCC 33279, L. pneumophila serotype 3 ATCC 33155 were cultured on buffered charcoal-yeast extract (BCYE) agar plates, which contained a Legionella CYE agar base (Oxoid, Basingstoke, Hampshire, UK) supplemented with the Growth Supplement SR0110A (ACES buffer/potassium hydroxide, ferric pyrophosphate, L-cysteine HCl, α-ketoglutarate; Oxoid, Basingstoke, Hampshire, UK) for three days at 37 °C in a humid atmosphere and 5% CO2 [40]. L. dumoffii were also cultivated on this medium enriched with 100 μg·mL⁻¹ of choline-trimethyl-do chloride (Sigma-Aldrich, Steinheim, Germany). The bacteria collected from this medium were washed three times with water by intensive vortexing and centrifugation at 8000× g for 10 min. Bacteria cultured with and without choline were killed by 90 °C for 20 min. The efficiency of temperature inactivation of bacteria was checked by streaking the bacteria on BCYE plates. Escherichia coli DH5α strain was grown in Luria-Bertani (LB) medium at 37 °C [41].

4.2. Fractionation of L. dumoffii Cultured on BCYE Medium with and without Choline

The bacteria were collected from 12 BCYE plates supplemented with labeled choline and 12 non-supplemented BCYE plates. They were washed twice in saline in order to remove the remaining medium. Cell membrane isolation was performed essentially as described by Hindahl and Iglewski [35].

The cells were washed twice with cold 10 mM HEPES (N-2-hydroxyethylpiperazine-N-2-ethanesulfonic acid; Sigma-Aldrich, Steinheim, Germany) buffer (pH 7.4) and centrifuged at 8000× g, 15 min in 4 °C. The cell pellets were suspended in 15 mL of 10 mM HEPES buffer containing 20% sucrose (w/v) and incubated with DNase (0.3 mg) (Sigma-Aldrich, Steinheim, Germany) and RNase (0.3 mg) (Sigma-Aldrich, Steinheim, Germany) at 37 °C for 30 min 0.8-mL suspensions were lysed by three passages through a French press (SLM-Amico Instruments, Thermo Spectronic, Rochester, NY, USA) at 18000 lb/in². After centrifugation for 20 min at $1000 \times g$, 4 °C performed to remove cell debris and undisrupted cells, total membrane fractions were collected by centrifugation for 60 min at $100,000 \times g$, 4 °C (SW 32Ti rotor, Beckman Coulter, Brea, CA, USA). Next, they were washed twice in cold 10 mM HEPES buffer by centrifugation at $100,000 \times g$ for 1 h, 4 °C, the pelleted membranes were suspended in 2.5 mL of 10 mM HEPES buffer and layered onto a seven-step sucrose gradient.

4.3. Isolation of the Outer and Inner Membranes

Two milliliters of each cell membrane suspension were loaded on the top of a discontinuous gradient prepared by combining sucrose solutions of the following concentrations: 6 mL 70%, 9 mL 64%, 8 mL 58%, 5 mL 52%, 4 mL 48%, 3 mL 42%, 3 mL 36% (w/v) in 10 mM HEPES buffer,

pH 7.5. The gradient was centrifuged at $114,000 \times g$, 20 h, 4 °C (SW 32Ti rotor, Beckman Coulter, Brea, CA, USA), and 1 mL fractions were subsequently collected from the top of the centrifugation tube. The protein content was determined in the individual fractions and the fractions from the upper band of the gradient and the lower band of the gradient were collected separately. Next, the upper and lower fractions pooled separately were suspended in 10 mM HEPES buffer and centrifuged at $54,000 \times g$ for 1 h (MLA 80, Optima MAX-XP, Beckman Coulter, Brea, CA, USA). The membrane pellets were washed three times in cold deionized water and centrifuged at $54,000 \times g$ for 1 h (MLA 80, Optima MAX-XP, Beckman Coulter, Brea, CA, USA). Next, they were diluted in 250 µL of water (MQ, Millipore, Billerica, MA, USA) for further analyses, *i.e.*, enzyme assays and lipid isolation.

The efficacy of the membrane isolation procedure was confirmed by measuring the activity of NADH oxidase [42] and esterase. The concentration of protein in the fractions was determined using the Bradford method and bovine serum albumin as a standard [43].

4.4. Enzyme Assays

As marker enzymes for the inner and outer membrane, activities of NADH oxidase and esterase, respectively, were determined by spectrophotometric analysis of the absorbance decrease at 340 nm for NADH oxidase and the absorbance increase at 405 nm for esterase. For NADH oxidase, 62 μL of the reaction mixture (50 mM Tris/HCl, pH 7.5, 0.2 mM DTT, 0.12 mM NADH) (Sigma-Aldrich, Steinheim, Germany) was incubated for 15 min at 37 °C with 8 μL of each fraction in a microtitre plate. For esterase activity, 63 mg of *p*-nitrophenyl acetate (Sigma-Aldrich, Steinheim, Germany) was dissolved in 10 mL of ethanol. One milliliter of this solution was slowly added to 100 mL of distilled water. Ten microliters of each fraction were added to 90 μL of the substrate solution, incubated for 10 min at 25 °C, and absorbance at 405 nm was recorded. Fractions with the NADH oxidase and esterase activities representing the inner and outer membrane fractions were lyophilized and weighed.

4.5. Isolation of Lipids

Lipids were isolated from bacterial cells cultivated on BCYE medium enriched with labeled choline using the Bligh and Dyer (1959) method: chloroform/methanol (1:2 v/v) [44]. Lipids were analyzed with MALDI/TOF mass spectrometry.

Lipids from the inner and outer membranes prepared from bacteria grown on the choline-supplemented and non-supplemented medium were extracted using the same extraction method. The dried organic phase was then purified with a mixture of hexane/isopropanol (3:2, v/v). The extracts were dried under nitrogen before weighing and then dissolved in chloroform for further LC/ESI-MS analysis.

4.6. Matrix-Assisted Laser-Desorption/Ionization (MALDI)-Time of Flight (TOF) Mass Spectrometry

MALDI-TOF mass spectrometry analysis was performed on a Voyager-Elite instrument (PE Biosystems, Foster City, CA, USA) using delayed extraction in the reflectron mode. The dry

lipid extract was dissolved in a CHCl₃/CH₃OH mixture (2:1, v/v). The sample constituents mixed with 0.5 M aniline salt of α -cyano-4-hydroxycinnamic acid (CHCA) as a matrix were desorbed and ionized with a nitrogen laser at an extraction voltage of 20 kV. Angiotensin was used as an internal standard. Each spectrum was the average of about 256 laser shots.

4.7. Liquid Chromatography Coupled with the Mass Spectrometry Technique Using the Electrospray Ionization Technique (LC/ESI-MS)

The LC/ESI-MS analyses were performed on a Prominence LC-20 (Shimadzu, Kyoto, Japan) liquid chromatograph coupled with a tandem mass spectrometer 4000 Q TRAP (Applied Biosystems Inc., Foster City, CA, USA), equipped with an electrospray (ESI) ion source (TurboIonSpray, Applied Biosystems Inc., Foster City, CA, USA) and a triple quadrupole/linear ion trap mass analyzer. The separation of the phospholipid mixture was carried out by normal phase chromatography using a 4.6×150 mm Zorbax SIL RX column (Agilent Technologies, Palo Alto, CA, USA). Hexane/isopropanol (3:2, v/v) was used as solvent A and an isopropanol/hexane/5 mM aqua solution of ammonium acetate (38:56:5, v/v/v) as solvent B. The following elution program was employed: from 53% B to 80% B for 23 min, 80% B maintained for 4 min, 80% B to 100% B for 9 min and 100% B maintained for 14 min. The flow rate was 1 mL/min. The lipid extracts were dissolved in phase A.

To identify PC species, the precursor ion mode (PI) and neutral loss scan (NL) mass spectrometry techniques were employed in the positive ion mode. The measurements were performed using an electrospray ion source (ESI) with the following parameters: ion spray voltage (IS)—5500 V, declustering potential (DP)—40 V, and entrance potential (EP)—10 V. Nitrogen was used as a curtain (CUR 20 psi), a nebulizer (GS1 50 psi), and a collision gas. The source temperature was set at 250 °C. The collision-induced dissociation (CID) spectra were obtained in the positive and the negative ion mode using collision energy of 50 eV.

The Multiple Reaction Monitoring (MRM) technique performed in the negative ion mode was employed to compare the PC amounts in the analyzed cell membranes. The most abundant PC molecular species at m/z 778.9 (16:0/15:0) and 764.9 (15:0/15:0) were chosen in this experiment. Based on the fragmentation spectra of the selected peaks in the negative ion mode, the following MRM pairs were used: 764.9/690.4, 764.9/241.2, and 778.9/704.4, 778.9/241.2.

Chemicals for the LC/MS system (hexane, isopropanol, acetonitrile LC grade) were obtained from Merck (Darmstadt, Germany), and ammonium acetate from Sigma-Aldrich Fluka (Steinheim, Germany).

4.8. DNA Methods, PCR Amplification, and Southern Hybridization

Restriction enzyme digestion, agarose gel electrophoresis, DNA labeling, and Southern hybridization were used for genomic DNA isolation [41]. To amplify the DNA fragment containing *pcsA* of *L. pneumophila* serotype 3 (ATCC 33155), the PCR reaction was performed using 100 ng of genomic DNA, REDTaq ReadyMix (Sigma-Aldrich, Steinheim, Germany) and 0.2 μM of each forward (pcsF: 5'-CTCTAGGATCCGTAATGAATCCAATAAA-3') and reverse (pcsR: 5'-CATAAATTGGATCCAAACTCAATCTTTATTAT-3') primers in a 50-μL final volume.

The PCR reaction was performed with the following temperature profile: initial denaturation at 94 °C for 4 min, 30 cycles of denaturation 94 °C for 1 min, annealing at 48 °C for 40 s, extension at 72 °C for 60 s, and final extension at 72 °C for 4 min. This PCR fragment was labeled using the non-radioactive DIG DNA Labeling and Detection kit according to the manufacturer's instruction (Roche Applied Science, Penzberg, Germany). The 800-bp-long DIG-labeled amplicon with the pcsA gene was used as a probe in the Southern hybridization. Additionally, DIG-labeled DNA of phage λ digested with EcoRI and HindIII restriction enzymes were used as a molecular size marker. Genomic DNA from L. pneumophila serotype 3, L. dumoffii, and E. coli DH5 α (used as a negative control) were digested with BamHI and EcoRI enzymes, separated by 0.7% agarose gel electrophoresis, and blotted. In addition, λ DNA digested with both EcoRI and HindIII enzymes was used as molecular markers. The hybridization experiments were performed at reduced-stringency conditions at 37 °C using 20% formamide in pre-hybridization and hybridization solutions as described previously [11].

To amplify and sequence the *L. dumoffii pcsA* gene, a set of degenerate primers has been designed based on the genomic sequences of *L. pneumophila* strain Philadelphia 1, *L. longbeachae* D-4968, and *L. drancourtii* LLAP12. Primers pcsLd-F (5'-ACTTTTKATWATYGATRMTATTTT-3') and pcsLd-R (5'-TAATCATWAAADABYCAAAGTCTAT-3') allowed amplification of the longest 900-bp fragment containing *pcsA* gene. The PCR reactions were performed with the temperature profile described above, except the annealing temperature that was decreased to 43 °C. The amplified DNA fragment was purified on the columns (A&A Biotechnology, Gdynia, Poland) and sequenced using the BigDye terminator cycle sequencing kit (Applied Biosystems, Inc., Foster City, CA, USA) and the ABI Prism 310 sequencer. Database searches were done with the BLAST and FASTA programs available at the National Center for Biotechnology Information (Bethesda, MD, USA) and the European Bioinformatics Institute (Hinxton, UK). The sequence of *L. dumoffii pcsA* obtained in this study has been deposited in NCBI GenBank under accession number KC197708. Promoter prediction in the upstream region of *L. dumoffii pcsA* was done using BDGP Neural Network Promoter Prediction (fruitfly.org).

4.9. THP-1 Cell Culture

The human acute monocytic leukemia *cell line* (THP-1) was obtained from the European Collection of Cell Cultures (Cat No. 88081201). Cells were cultured at $(0.5-7) \times 10^5$ cells/mL in RPMI 1640 supplemented with 10% heat-inactivated fetal calf serum (FCS), 10 mM HEPES, 2 mM glutamine, 100 U/mL penicillin, and 100 μ g/mL streptomycin. The cells cultured in tissue culture flasks (Falcon, Bedford, MA, USA) were incubated at 37 °C in a humidified atmosphere of 5% CO₂. The culture media, antibiotics, and FCS were purchased from Sigma-Aldrich (Steinheim, Germany).

4.10. THP-1 Cell Differentiation

THP-1 were seeded onto 24-well plastic plates (Nunc, Roskilde, Denmark) at a density of 5×10^5 cells/well in RPMI 1640 supplemented with 10% FCS, and treated with a final concentration of 50 ng/mL phorbol 12-myristate 13-acetate (PMA) (Sigma-Aldrich, Steinheim, Germany) for three

days to induce maturation toward adherent macrophage-like cells. Subsequently, unattached cells were removed and after three-time washing adherent THP-1 cells were cultured in medium without PMA for three consecutive days with daily fresh medium change.

4.11. THP-1 Cell Viability Assay

The viability of THP-1 cells exposed to $L.\ dumoffii$ bacteria was determined by the MTT assay, in which the yellow tetrazolium salt 3-(4,5 dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide (MTT) is metabolized by viable cells to purple formazan crystals. The cells at a density of 5×10^4 cells/well were seeded onto 96-well plates (Nunc, Roskilde, Denmark) and cultured in 10% RPMI 1640 for 24 h. Next, the medium was replaced with a fresh one with addition of 2% FCS and different concentrations of $L.\ dumoffii$ at a MOI of 10, 50, 100, and 200. The treated cells were maintained in a humidified CO₂-incubator at 37 °C for 4 and 24 h. Subsequently, MTT solution (25 μ L of 5 mg/mL in PBS) (Sigma-Aldrich, Steinheim, Germany) was added to each well. The cells were incubated at 37 °C for 3 h, and 100 μ L of SDS in 0.01 M HCl was added to dissolve the formazan crystals during overnight incubation. The controls included native (non-treated) cells and the medium alone. The spectrophotometric absorbance was measured at 570 nm wavelength using a VICTOR X4 Multilabel Plate Reader (Perkin Elmer, Waltham, MA, USA). The data are presented as percentage of control cell viability.

4.12. In Vitro Infection of Differentiated THP-1 with L. dumoffii—Control of Internalization and Intracellular Growth

Differentiated THP-1 cells (as described in 4.10.) were infected with 10 MOI of live *L. dumoffii* cultured on choline-supplemented and non-supplemented medium. After incubation for 2 h at 37 °C, 5% CO₂, nonphagocytized bacteria were killed by the addition of 100 mg of gentamicin/mL for 1 h. Next, supernatants were removed and the macrophages were washed three-times with PBS. 1 mL of sterile distilled water (for bacterial internalization study) or 1 mL of RPMI 1640 supplemented with 10% FCS (without antibiotics) was added and the macrophages were incubated for 24, 48, and 72 h (intracellular bacterial growth study).

4.12.1. Bacteria Internalization Assay

Cells suspended in 1 mL of sterile distilled water were disrupted by aspiration through a 25-gauge needle and then series of 10-fold dilutions were made. Subsequently, 0.1 mL of each dilution was inoculated onto BCYE agar and colonies of culturable *L. dumoffii* were counted after three days of incubation at 37 °C, 5% CO₂.

4.12.2. Intracellular Bacteria Growth Assay

After 24, 48, and 72 h of cell culture incubation, supernatants were collected into sterile tubes, centrifuged (8000 rpm/min for 10 min), and washed with sterile distilled water. One milliliter of sterile distilled water was added to THP-1 cells, which were disrupted as described above and pooled

with centrifuged pellet of the respective supernatant. 0.1 mL of series of 10-fold dilutions of each sample were inoculated onto BCYE agar and incubated as above.

Formation of colonies was determined in triplicate for at least two independent experiments.

4.13. Induction of TNF-α with L. dumoffii in THP-1 Cells

The differentiated macrophages were treated with live or dead L. dumoffii for 4 h at a MOI = 10 or 100. In another experiment, the macrophages were treated with different concentrations (10–1000 ng/mL) of L. dumoffii outer or inner membranes non-treated and treated with temperature (90 °C, 20 min).

Additional controls were performed: non-treated macrophages, non-activated THP-1 cells.

In all experiments, the bacteria had been previously cultured on medium with or without addition of choline (choline-trimethyl-d₉ chloride, Sigma-Aldrich, Steinheim, Germany). After incubation for 4 h at 37 °C, 5% CO₂, cell culture supernatants were collected and frozen immediately at –80 °C for further TNF-α determination. TNF-α level was measured by the ELISA method using a commercial kit from R&D Systems according to the *manufacturer's instructions* (*R&D Systems Inc., Minneapolis, MN, USA*). All experiments were conducted in five independent replicates.

4.14. Statistics

Values are expressed as mean \pm S.D. Results were statistically evaluated using two-way ANOVA and Tukey's *post hoc* tests (Statistica software ver. 6.0, StatSoft Inc., Tulsa, OK, USA, 2001). p values of \leq 0.05 were considered significant.

5. Conclusions

We have demonstrated that $L.\ dumoffii$ has an ability to utilize extracellular choline in the Pcs pathway, which may be a dominant pathway, as indicated by the presence of a strong promoter for the phosphatidylcholine synthase gene. The use of the LC/ESI-MS technique allowed us to provide evidence that the PCs synthesized by $L.\ dumoffii$ are localized in both the inner and outer membranes, but they are structurally different. This technique allowed identification of the PC species in total lipid extracts isolated from live bacteria without any chemical modification, thereby showing the physiological state of the cells. The structural differences in the PC species affect the induction of TNF- α —a key player in immuno-inflammatory response. In the presence of choline, the bacteria exhibited a reduced capability of TNF- α induction. The reduction of inflammatory response caused by PC species might collaborate to evade the immune system, thus, allowing $L.\ dumoffii$ to establish in the host. The identification of selective inhibitors of phosphatidylcholine synthase may contribute to development of specific adjuvant therapy in treatment of legionellosis.

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Conflicts of Interest

The authors declare no conflict of interest.

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Recent Advances in Bacteria Identification by Matrix-Assisted Laser Desorption/Ionization Mass Spectrometry Using Nanomaterials as Affinity Probes

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Abstract: Identifying trace amounts of bacteria rapidly, accurately, selectively, and with high sensitivity is important to ensuring the safety of food and diagnosing infectious bacterial diseases. Microbial diseases constitute the major cause of death in many developing and developed countries of the world. The early detection of pathogenic bacteria is crucial in preventing, treating, and containing the spread of infections, and there is an urgent requirement for sensitive, specific, and accurate diagnostic tests. Matrix-assisted laser desorption/ionization mass spectrometry (MALDI-MS) is an extremely selective and sensitive analytical tool that can be used to characterize different species of pathogenic bacteria. Various functionalized or unmodified nanomaterials can be used as affinity probes to capture and concentrate microorganisms. Recent developments in bacterial detection using nanomaterials-assisted MALDI-MS approaches are highlighted in this article. A comprehensive table listing MALDI-MS approaches for identifying pathogenic bacteria, categorized by the nanomaterials used, is provided.

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1. Introduction

Worldwide, infectious diseases cause nearly 40% of the total 50 million deaths annually [1]. According to the World Health Organization, microbial hazards are the primary concern [2] because microbial diseases are a major cause of death in many developing and developed countries of the world [3,4]. Therefore, the development of rapid, accurate, and sensitive methods for bacterial identification is important for the clinical diagnosis, efficient treatment and prevention of diseases, environmental monitoring and food safety [5–8]. In clinical laboratories, bacterial identification is typically based on phenotypic tests, including Gram staining, culture and growth characteristics, and biochemical patterns. A number of methods are currently employed to detect and identify pathogenic agents, and these mainly rely on specific microbiological and biochemical identification methods [9–11]. These methods include culturing the microbes and counting the bacterial colonies, immunology-based methods, antigen-antibody interaction methods, and the polymerase chain reaction method, which involves DNA analysis. These methods can be sensitive and inexpensive, and can provide both qualitative and quantitative information about the test microorganisms; however, they are often time-consuming and laborious because each involves a pathogen amplification step. At present, most bacteria can be identified between a few hours to 1-2 days using these methods, with slow-growing microorganisms requiring additional time or supplementary tests [12]. Consequently, there is an urgent requirement for developing a rapid, sensitive, and selective detection method for such pathogens to treat individuals at risk, to improve public health surveillance and epidemiology, which is essential for ensuring the safety of food supplies, and to diagnose infectious bacterial diseases accurately.

There are challenges associated with identifying various types of pathogenic bacteria in a wide range of samples. Matrix-assisted laser desorption/ionization mass spectrometry (MALDI-MS) has been used to analyze various biomolecules, including peptides, proteins, DNA, RNA, oligonucleotides, oligosaccharides, and polymers [13–16]. This approach was first introduced by Tanaka and Karas in the late 1980s [17,18], and is a soft ionization method that provides mass spectra of the analytes with a minimum amount of fragmentation. MALDI-MS has a number of advantages over conventional methods including ease of operation, providing structural information of molecules with high throughput, speed, sensitivity, accuracy, and reproducibility [19,20]. Therefore, it has become a powerful tool for rapid characterization, differentiation, and identification of microorganism species [21–27]. For example, the mass-spectral profiling of whole cells can indicate the presence of unique biomarkers that can serve as the basis for identifying microbes [28,29]. In general, mass spectra of microbes isolated from a sample may contain unique patterns that can be automatically matched with spectra in a well-established reference library of microorganisms that have been characterized using appropriate sample preparation protocols. Matching the spectra allow the microbes to be identified as well as evaluated.

A sufficient number of bacterial cells (typically ~10⁴ cells per well) are required to generate detectable MALDI-MS ion signals. However, samples obtained from infectious biological fluids or food poisoning samples are difficult to characterize directly by MALDI-MS because the ions generated from the bacterial cells may be seriously suppressed by the complex sample matrices. This led to the idea of using nanoparticles as affinity probes, to enhance the ability of MALDI-MS to detect bacteria [30,31]. Nanoparticles provide a high surface to volume ratio, giving them high binding and capture efficiencies for bacteria. Affinity separation approaches are methods of selectively concentrating trace amounts of bacteria from complex biological and food samples before they are characterized using MALDI-MS. When inorganic nanoparticles are used in MALDI-MS, instead of organic matrices, the method is called surface-assisted laser desorption and ionization MS (SALDI-MS) [32–36]. SALDI-MS was originally proposed by Sunner and Chen as early as 1995, and graphite particles were originally used as ion emitters [37]. This method was called SALDI-MS to emphasize that the surfaces and surface structures are critical to not only sample preparation but also desorption and ionization processes [38]. Numerous types of nanoparticles such as gold (Au) nanoparticles [39-41], silver (Ag) nanoparticles [42,43], magnetic nanoparticles [44,45], titanium dioxide (TiO₂) nanoparticles [46,47], carbon nanotubes [48,49], carbon nanoparticles [50], nanodiamonds [51], and graphene and graphene oxide [52] have been successfully used as matrices in SALDI-MS. The nanomaterials used in SALDI-MS play similar roles to the organic matrices used in MALDI-MS, absorbing energy from the laser irradiating them and efficiently transferring the energy to the analytes, causing the analytes to be desorbed and ionized [30]. The method provides several advantages including lower background noise in the low mass region, high surface areas, simple sample preparation, flexibility in sample desorption under different conditions, and high UV

absorptivity [34]. Nanoparticles can also act as affinity probes, making it easy to concentrate the analytes, and offering good sensitivity and reproducibility [34].

In this review article, we focused on the overview of the recent advancements in the use of nanoparticles as affinity probes to enhance the detection sensitivity and selectivity of bacteria using MALDI-MS. Several examples of successful MALDI-MS approaches for detecting pathogenic bacteria have been provided to illustrate the advantages of this approach with respect to simplicity, sensitivity, and reproducibility. Furthermore, this article also provides some examples for the identification of bacteria in real samples using nanomaterials-assisted MALDI-MS approaches.

2. Bacterial Identification Using MALDI-MS

MALDI-MS is a very sensitive method where a single bacteria colony is sufficient for analysis, while other methods typically require culturing or enrichment of bacteria to obtain sufficient materials. Therefore, in clinical microbiological laboratories, the MALDI-MS is increasingly used for bacterial identification through the determination of the exact molecular masses of numerous peptides and small proteins, many of which are ribosomal. Conventional biochemical differentiation methods [24] have already been replaced by MALDI-MS. Because MALDI-MS is primarily applicable for analyzing clonal isolates, cultivation of the microorganism is still required. Moreover, for accurate identification, MALDI-MS can be used directly on the clinical samples that contain very few bacteria for accurate identification. In 2010, Ferreira et al. [53] introduced a MALDI-MS method for direct analysis of urine samples (4 mL) and observed that the inoculum level in the samples must be greater than 10⁵ cfu/mL (colony-forming unit/mL). In 2010, a protocol for direct analysis of blood was introduced by Stevenson et al. [54], who separate bacteria from the red blood cells and plasma proteins via several centrifugation steps. A total of 212 positive cultures representing 32 genera and 60 species or groups were examined. Besides urine and blood, Barreiro et al. [55] inoculated pasteurized and homogenized samples of whole milk with the bacterial loads of 10³-10⁸ cfu. SepsityperTM Kit (Bruker, Billerica, MA, USA) was used to for the testing milk sample and then analyzed by the Bruker BioTyper database. For a slightly contaminated (10⁴ cfu/mL bacteria) milk sample, bacterial identification could be performed after initial incubation at 37 °C for 4 h. The detection limits for bacteria were in the range of 10⁶–10⁷ cfu/mL.

3. Nanoparticles Used as Affinity Probes

Nanoparticles are clusters of a few hundred to a few thousand atoms, and range from 1 to 100 nm in diameter. The chemical and physical properties of nanoparticles depend on their surfaces; therefore, these properties are highly dependent on the sizes, shapes, and compositions of the nanoparticles [56–58]. Nanoparticles have high surface-to-volume ratios, and those with excellent optical, magnetic, and electronic properties have been employed in sensing, imaging, catalysis, electronics, optics, and optoelectronics applications [59–65]. Nanoparticles can play an important role in determining the sensitivity of MALDI-MS and provide a high surface-to-volume ratio to give a high binding efficiency for bacteria. The affinity separation approach has been used to attempt to selectively concentrate trace amounts of bacteria from biological and food samples. Nanoparticles

(functionalized or unmodified) that have been used as affinity probes to increase the sensitivity of MALDI-MS for detecting microbes are summarized in Table 1.

Table 1. Nanomaterials used as affinity probes in MALDI-MS.

Nanomaterials	Functionalized molecule	Pathogen	Application	LOD (cfu/mL)	Ref.
Fe ₃ O ₄ NPs	IgG	S. aureus; S. saprophyticus		3.0×10^{5}	[66]
Fe ₃ O ₄ NPs	IgG	S. saprophyticus	Urine	3.0×10^{7}	[66]
Fe ₃ O ₄ NPs	Vancomycin	S. aureus; S. saprophyticus	Urine	7.8×10^4 ; 7.4×10^4	[67]
Fe ₃ O ₄ NPs	Vancomycin	B. cereus; E. faecium; S. aureus	Tap water, reservoir water	5.0×10^{2}	[68]
Ag NPs		E. coli; S. marcescen		N.D.	[69]
Ag NPs		B. lactis; L. acidophilus; S. thermophilus; L. bulgaricus	Yogurt	N.D.	[70]
Ag NPs		L. acidophilus; B. longum; L. bulgaricus; S. thermophilus	Yogurt	N.D.	[70]
CdS QDs		E. coli		N.D.	[71]
CdS QDs		S. cerevisiae; C. utilis		N.D.	[72]
CdS QDs	Chitosan	P. aeruginosa; S. aureus		2.0×10^2 ; 1.5×10^2	[73]
Pt NPs	Mixed with ionic liquid (1-butyl-3- methylimidazolium hexafluorophosphate)	E. coli; S. marcescens		10 ⁶	[74]
Pt NPs	IgG	B. thuringiensis; B. subtilis	Rhizospheric soil and root	N.D.	[75]
Pt NPs	IgG	S. marcescens; E. coli		10^{5}	[76]
AuNCs	Lysozyme	E. coli; K. pneumoniae; P. aeruginosa; pandrug-resistant A. baumannii; S. aureus; E. faecalis; vancomycin-resistant E. faecalis	Fetal bovine serum; urine	N.D.; 10 ⁶	[77]
Graphene magnetic nanosheets	Chitosan	P. aeruginosa; S. aureus	Blood	6.0×10^2 ; 5.0×10^2	[78]

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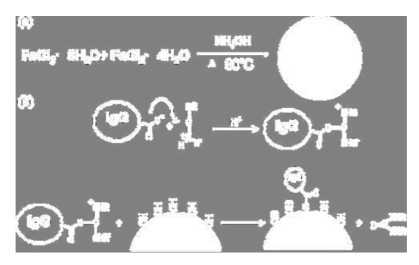
Nanomaterials	Functionalized molecule	Pathogen	Application	LOD (cfu/mL)	Ref.
NiO NPs		E. coli		10^{7}	[79]
TiO ₂ NPs		S. aureus	Human nasal passage	N.D.	[80]
ZnO NPs		E. coli		N.D.	[81]
Ag, Pt, NiO, TiO ₂ , ZnO NPs		S. aureus; P. saeruginosa		N.D.	[82]

Ref., Reference; Ag, silver; AuNCs, gold nanoclusters; CdS, cadmium sulfide; IgG, immunoglobulin; LOD, limit of detection; N.D., not determined; NiO, nickel oxide; NPs, nanoparticles; Pt, platinum; QDs, quantum dots; TiO₂, titanum dioxide; ZnO, zinc oxide.

3.1. Magnetic Nanoparticles

Ho et al. [66] immobilized human immunoglobulin (IgG) onto the surfaces of magnetic Fe₃O₄ nanoparticles through covalent bonding (Figure 1). The functionalized magnetic nanoparticles were used as affinity probes to selectively concentrate pathogens, such as *Staphylococcus aureus* (*S. aureus*) and *Staphylococcus saprophyticus* (*S. saprophyticus*), from sample solutions. The bacteria were then characterized using MALDI-MS. The lowest bacterial concentration detected in an aqueous sample solution (0.5 mL) was 3×10^5 cfu/mL, for both *S. aureus* and *S. saprophyticus*, and the lowest detectable *S. saprophyticus* concentration in a urine sample was 3×10^7 cfu/mL.

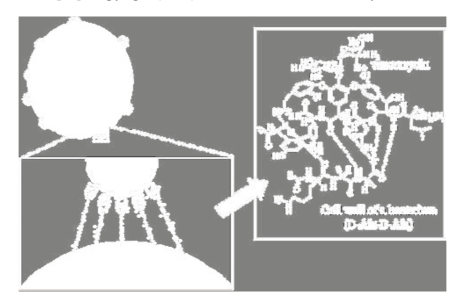
Figure 1. Synthetic route for immobilizing immunoglobulin (IgG) onto the surfaces of Fe₃O₄ magnetic nanoparticles. Reprinted with permission from [66]. Copyright (2014) American Chemical Society.



Vancomycin-modified 11 nm magnetic (Fe₃O₄) nanoparticles were used as affinity probes to selectively bind to the surface walls of Gram-positive bacteria (*S. aureus* and *S. saprophyticus*), as shown in Figure 2, allowing the bacteria to then be directly characterized using MALDI-MS [67]. Vancomycin is one of the most potent antibiotics, and has a high specificity for the *D*-Alanine (Ala) (*D*-Ala) moieties on the cell walls of Gram-positive bacteria. The lowest cell concentrations that

could be detected in a urine sample (3 mL) were 7.4×10^4 cfu/mL for *S. aureus* and 7.8×10^4 cfu/mL for *S. saprophyticus*.

Figure 2. Cartoon illustrations of the proposed method for anchoring vancomycin-immobilized magnetic nanoparticles onto the surface of a Gram-positive bacterial cell and the binding of vancomycin to the terminal of *D*-Alanine (*D*-Ala)–*D*-Ala units of the peptides on the cell wall of a Gram-positive bacterium. Reprinted with permission from [67]. Copyright (2014) American Chemical Society.



IgG- and vancomycin-modified magnetic nanoparticles have been demonstrated to exhibit effective affinities for selectively concentrating traces of bacteria from the sample solutions. However, because interferences from the urine matrix affect the binding capacity of these nanoprobes, further improvements are required to reduce the matrix effects in the analysis of biological samples.

A combination of membrane filtration and vancomycin-modified magnetic (Fe₃O₄) 15–20 nm nanoparticles has been used to selectively concentrate Gram-positive bacteria from tap water and reservoir water, allowing the bacteria to be rapidly analyzed using whole-cell MALDI-MS [68]. The capture efficiency for Gram-positive bacteria using these vancomycin-modified magnetic nanoparticles was 26.7%–33.3%, and the analysis time was approximately 2 h. This approach enhanced the sensitivity of the method by a factor of approximately 6×10^4 , giving a limit of detection of 5×10^2 cfu/mL for *Bacillus cereus* (*B. cereus*), *Enterococcus faecium* (*E. faecium*), and *S. aureus* in water samples (2 L).

3.2. Silver (Ag) Nanoparticles

The bifunctional properties of Ag nanoparticles allowed them to be used as affinity probes for *Escherichia coli* (*E. coli*) and *Serratia marcescens* (*S. marcescens*), by Gopal *et al.* [69], to increase the sensitivity of MALDI-MS when characterizing the bacteria. The critical concentration of affinity probes for Ag nanoparticles was 1 mL/L in the case of *E. coli* and 0.5 mL/L in the case of *S. marcescens*. Ag nanoparticle concentrations higher than these values showed pronounced bactericidal activities.

The same research group also observed that an ionic solution (CrO₄²⁻) and 0.035 mg of Ag nanoparticles could be used to capture yogurt bacteria (*Bifidobacterium lactis* (*B. lactis*), *Lactobacillus acidophilus* (*L. acidophilus*), *Streptococcus thermophilus* (*S. thermophilus*), and *Lactobacillus bulgaricus* (*L. bulgaricus*) from AB yogurt and *L. acidophilus*, *Bifidobacterium longum* (*B. longum*), *L. bulgaricus*, and *S. thermophilus* from Lin yogurt), improving the sensitivity achieved for detecting bacteria in yogurt samples [70]. This method has demonstrated a rapid, selective and sensitive means of bacterial detection using MALDI-MS for food microbiology.

3.3. Cadmium Sulfide (CdS) Quantum Dots (QDs)

Gopal *et al.* [71] has reported that CdS QDs can degrade the extracellular polysaccharides of *E. coli* cells when using MALDI-MS. Adding 20 μL/L of CdS QDs was observed to enhance the extracellular polymeric substance (EPS) peaks using an incubation time of up to 3 h. The authors confirmed that CdS QDs can function as more than just affinity probes, being able to degrade EPSs. CdS QDs can, therefore, be used to inactivate pathogenic *E. coli* and also inhibit the growth of *E. coli* biofilms.

Manikandan and Wu [72] observed that CdS QDs (10 mg/L) with particle sizes of 1–7 nm performed fungicidal roles and functioned as protein signal enhancement probes in the MALDI-MS analysis of the fungi *Saccharomyces cerevisiae* (*S. cerevisiae*) and *Candida utilis* (*C. utilis*). From their MALDI-MS results, the authors proposed the mechanism involving the CdS QDs interacting with the EPSs and removing small molecules from the EPS layers. The MALDI-MS protein signals were enhanced at all of the CdS QD concentrations that were tested (10–30 mg).

Chitosan-modified CdS QDs have been used as effective bacterial biosensors because of the strong affinities between chitosan molecules and bacterial membranes [73]. In that study, *Pseudomonas aeruginosa* (*P. aeruginosa*) and *S. aureus* cells were detected at low concentrations, 200 and 150 cfu/mL, respectively, after an extremely short time (1 min). MALDI-MS and transmission electron microscopy were used to confirm the interactions and the biocompatibility of the chitosan-modified CdS QDs with bacterial cells.

3.4. Platinum (Pt) Nanoparticles

Ahmad and Wu [74] employed a single drop microextraction technique, using an ionic liquid (1-butyl-3-methylimidazolium hexafluorophosphate) drop mixed with Pt nanoparticles, to extract bacterial proteins from aqueous samples to characterize pathogenic bacteria using MALDI-MS. This approach is based on surface changes in the ionic liquid and the membrane proteins of the bacteria, and it was successfully used to detect *E. coli* and *S. marcescens* at concentrations as low as 10^6 cfu/mL.

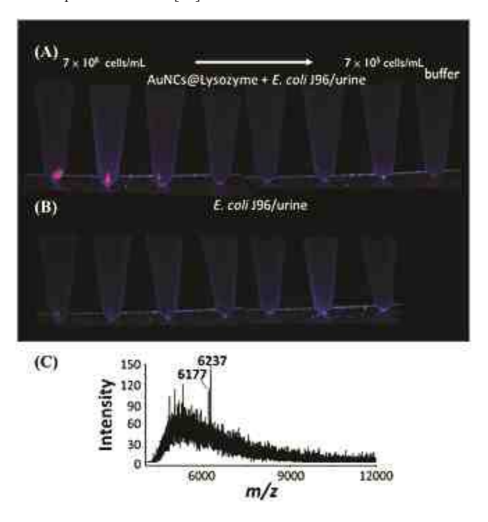
A rapid method for detecting bacteria associated with plants by an on-particle ionization and enrichment approach using IgG-functionalized Pt nanoparticle-assisted MALDI-MS was reported by Ahmad *et al.* [75]. The approach was successfully used to detect *Bacillus thuringiensis* (*B. thuringiensis*) and *Bacillus subtilis* (*B. subtilis*) isolated from rhizospheric soil and carrot plant roots. This study proved that bacteria can be directly detected even at low concentrations.

A rapid and sensitive approach to studying interactions between an affinity probe and a bacterial wall was introduced by Ahmad and Wu [76]. IgG was immobilized on Pt nanoparticles and MALDI-MS was used to detect the specific surface proteins of the bacteria *S. marcescens* and *E. coli*. This approach enabled the rapid detection of bacterial proteins, at a high resolution and with good sensitivity, without the need for tedious washing and separation procedures, and can be used to detect approximately 10⁵ cfu/mL of *S. marcescens* and *E. coli*.

3.5. Other Nanomaterials

Chan et al. [77] demonstrated that pathogenic bacteria, including E. coli, Klebsiella pneumonia (K. pneumoniae), P. aeruginosa, pandrug-resistant Acinetobacter baumannii (A. baumannii), S. aureus, E. faecalis, and vancomycin-resistant E. faecalis, can be concentrated by lysozyme-encapsulated gold nanoclusters (AuNCs) that photoluminesce red, and distinguished by the results combining MALDI-MS and principal component analysis. Figure 3A shows photographs of sample tubes after the lysozyme-AuNCs were used as probes for E. coli J96 in urine samples containing different concentrations of the E. coli. Photographs of the control experiment sample tubes are shown in Figure 3B for comparison. Figure 3C shows the MALDI-MS spectra of the conjugates containing E. coli J96, in which the peaks at m/z 6177 and 6237 correspond to E. coli J96. The lowest E. coli concentration that could be detected using this approach was approximately 106 cfu/mL. The advantages of this method include speed (without cell culturing) and simplicity, and it can be used as universal affinity probes for Gram-positive/negative and antibiotic-resistant bacteria.

Figure 3. Photographs obtained by vortex-mixing (**A**) the lysozyme-AuNCs with *E. coli* J96 at different cell concentrations and (**B**) *E. coli* J96 alone for 1 h at different cell concentrations, followed by centrifugation at 3500 rpm for 5 min. The samples were prepared in urine that was diluted 50-fold with PBS solution (pH 7.4) containing BSA (~10 μM). The photographs were taken under illumination of UV light ($\lambda_{max} = 365$ nm); (**C**) Examination of the limit of detection. Matrix-assisted laser desorption/ionization mass spectrometry (MALDI-MS) obtained after using the lysozyme-AuNCs (1.36 mg/mL, 0.1 mL) as affinity probes to concentrate target species from a urine sample (0.90 mL) containing *E. coli* J96 (1.59 × 10⁶ cells/mL) for 1 h. The urine sample was diluted 50-fold with PBS solution (pH 7.4) containing BSA (~10 μM) prior to bacterial spiking. Reprinted with permission from [77].



Abdelhamid and Wu [78] demonstrated that multifunctional graphene magnetic nanosheets modified with chitosan (GMCS) can be used in MALDI-MS for the sensitive detection of pathogenic bacteria (*P. aeruginosa* and *S. aureus*). The GMCS were observed to act as efficient separation and preconcentration nanoprobes for SALDI and enhance the ionization of bacterial biomolecules. GMCS have been used in the direct detection of low concentrations of *P. aeruginosa* and *S. aureus* in blood samples, demonstrating their practical applicability. This approach offers many advantages such as robustness, simplicity, and the capability for fluorescence based real-sample monitoring.

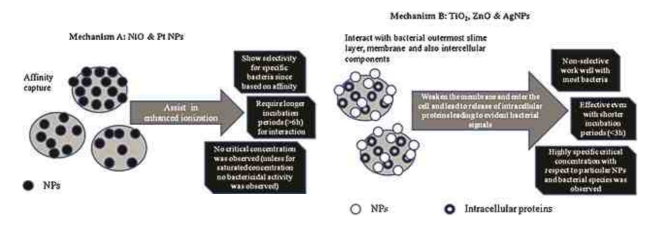
The heat stress response of E. coli (at 10^7 cfu/mL) at different temperatures has been studied using nickel oxide (NiO) nanoparticle-assisted MALDI-MS by Hasan $et\ al$. [79]. MALDI-MS was successfully used to detect 10 kDa chaperonin proteins produced by E. coli under heat stress at temperatures between 40 and 80 °C in the absence or presence of NiO nanoparticles. Dramatic decreases in the viability of E. coli in the presence of NiO were confirmed from the MALDI-MS results. This technique is a rapid, sensitive, and efficient approach for bacterial detection under extremely harsh conditions.

Gopal *et al.* [80] demonstrated that *S. aureus* isolated from the human nasal passage can be directly detected using MALDI-MS assisted by TiO₂ nanoparticles, without any culturing steps or sample pretreatment being required. TiO₂ nanoparticles were used to enhance the bacterial signals in the direct MALDI-MS analysis.

MALDI-MS has been used to evaluate bactericidal activity, by detecting proteins produced because of the inactivation of *E. coli* cells by zinc oxide (ZnO) nanoparticles [81]. The results showed that at concentrations of 1 and 5 g/L ZnO nanoparticles can be used as affinity probes to improve the signal intensities in the MS spectra. The significant differences in the spectral patterns confirmed that MALDI-MS was successfully used to evaluate the bactericidal activity of ZnO nanoparticles.

Gopal *et al.* [82] proposed mechanisms for the interactions between five nanoparticles (Ag, NiO, Pt, TiO₂, and ZnO) and two bacteria (*S. aureus* and *P. aeruginosa*) from studies using transmission electron microscopy, ultra spectrometry, and MALDI-MS. Two mechanisms (Figure 4) were proposed for the interactions: (1) Mechanism A was proposed for Pt and NiO nanoparticles, the function of which is based on their affinities for bacterial walls; and (2) Mechanism B was proposed for bactericidal nanoparticles, such as TiO₂, ZnO, and Ag nanoparticles.

Figure 4. Schematic diagram showing the mechanisms (Mechanism A and Mechanism B) for interactions of five nanoparticles with two pathogenic bacteria postulated in the study. Reprinted with permission from [82].



4. Conclusions

MALDI-MS is an emerging analytical tool for detecting and identifying microorganisms. It offers high sensitivity, simple sample preparation processes, low sample consumption volumes, and the possibility of automated and high-throughput analyses. In this review, we have described several MALDI-MS approaches for detecting pathogenic bacteria using nanomaterials (such as AuNCs, Ag, magnetic, and Pt nanoparticles and CdS QDs) as affinity probes. The nanomaterials described here act as concentration probes for the selective capture of unique biomarkers from microorganisms, and as surfaces to absorb energy from the laser irradiation, thereby inducing desorption and ionization of the analytes.

As mentioned above, the most important advantage of the affinity-based nanoparobe methods is their ability to selectively concentrate and purify microorganisms from complex samples, such as urine and blood, and allow the further identification of microorganisms without microbial culturing using MALDI-MS. For the nanomaterials-assisted MALDI-MS, Direct analysis of microorganisms at low microbial levels can be performed using the nanomaterials-based MALDI-MS. Numerous nanomaterials have been demonstrated to be useful as affinity probes for targeting bacteria. However, some nanoparticles such as Ag, TiO₂ and ZnO, also exhibit bactericidal activity, and therefore might not be good affinity probes at higher nanoparticles concentrations. Controlling of the nanoparticle concentration will be a key factor. All these nanomaterials-assisted MALDI-MS methods also encounter challenges with respect to the enrichment of unknown target bacterial species from the urine, blood, and cerebrospinal fluid. Thus, a limiting factor in MALDI-MS analysis is insufficient database entries. The addition of certain species to the database has been demonstrate significantly improve MALDI-MS precision in bacterial identification.

The broad adoption of nanomaterials-assisted MALDI-MS methods for bacterial identification will require a substantial improvement in performance compared with the existing methods, such as conventional MALDI-MS and biochemical tests. Thus, the standardization of terminology is required. Advances in nanomaterials-assisted MALDI-MS methods will support the simple and accurate means of bacterial identification for food safety, environmental monitoring and clinical diagnosis.

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Conflicts of Interest

The author declares no conflict of interest.

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MALDI Q-TOF CID MS for Diagnostic Ion Screening of Human Milk Oligosaccharide Samples

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Abstract: Human milk oligosaccharides (HMO) represent the bioactive components of human milk, influencing the infant's gastrointestinal microflora and immune system. Structurally, they represent a highly complex class of analyte, where the main core oligosaccharide structures are built from galactose and N-acetylglucosamine, linked by 1-3 or 1-4 glycosidic linkages and potentially modified with fucose and sialic acid residues. The core structures can be linear or branched. Additional structural complexity in samples can be induced by endogenous exoglycosidase activity or chemical procedures during the sample preparation. Here, we show that using matrix-assisted laser desorption/ionization (MALDI) quadrupole-time-of-flight (Q-TOF) collision-induced dissociation (CID) as a fast screening method, diagnostic structural information about single oligosaccharide components present in a complex mixture can be obtained. According to sequencing data on 14 out of 22 parent ions detected in a single high molecular weight oligosaccharide chromatographic fraction, 20 different oligosaccharide structure types, corresponding to over 30 isomeric oligosaccharide structures and over 100 possible HMO isomers when biosynthetic linkage variations were taken into account, were postulated. For MS/MS data analysis, we used the de novo sequencing approach using diagnostic ion analysis on reduced oligosaccharides by following known biosynthetic rules. Using this approach, de novo characterization has been achieved also for the structures, which could not have been predicted.

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1. Introduction

Human milk oligosaccharides (HMO) are the third most abundant type of component in human milk [1]. Their core structure is biosynthesized by adding *N*-acetyllactosamine units to the single lactose core at the reducing end. Apart from variable branching patterns, even a more significant layer of structural diversity is provided by extensive fucosylation. Recent studies have shown that HMO consist of around 200 oligosaccharide structures, including both neutral and negatively charged structures [2–4]. Their most understood function is the interaction with infant's gut microflora, stimulating the growth of probiotic bacteria [5,6] and providing immunological benefits to the infant [7]. Based on animal studies, a role of sialylated oligosaccharides in brain development has been postulated [8,9]. In recent studies, milk oligosaccharide patterns from women delivering preterm and at term were compared [10], focusing on differences in fucosylation patterns and lacto-*N*-tetraose (LNT) abundance. In spite of significant advances in the analytical techniques used for oligosaccharide structure elucidation over the past two decades [2–4,11–19], all individual components of HMO cannot be fully revealed.

In addition to the complex nature of HMO biosynthesis, additional variety in HMO structures can arise from chemical procedures carried out during sample preparation steps or due to biological enzymatic activity. To these belongs also the chemical procedure of mild acid hydrolysis in order to remove fucose and sialic acid residues [20]. This may be necessary to simplify a detailed structural analysis of the oligosaccharide core structure. It has also been found that sugars on the nonreducing terminus of the oligosaccharide are susceptible to cleavage by glycosidases in breast and during the storage of milk, although this degradation was found to be modest [21]. Other experimental parameters and biological specificities of the milk donor may influence HMO composition. For targeted (data-dependent acquisition or DDA) MS analysis methods, preliminary characterization of the sample is necessary. Matrix-assisted laser desorption/ionization (MALDI) MS is the method of choice for oligosaccharide screening, in particular in the context of high-throughput options. In most cases, MALDI-MS fingerprints of glycan mixtures display a singly charged ionic signal per component, which tends to simplify spectral interpretation in comparison to electrospray ionization (ESI). For most experiments conducted on glycans, ubiquitous sodium ions from glass and other sources attach to glycan molecules to form [M + Na]⁺ ions. For direct measurements on neutral glycans, [M + Na]⁺ abundances (peak area) may be compared in a semi-quantitative way within the same sample [22]. Combined with collision-induced dissociation (CID) available on quadrupole-time-of-flight (Q-TOF) mass spectrometers, it can quickly provide sufficient information for subsequent, more detailed MS analyses for a large number of precursor ions.

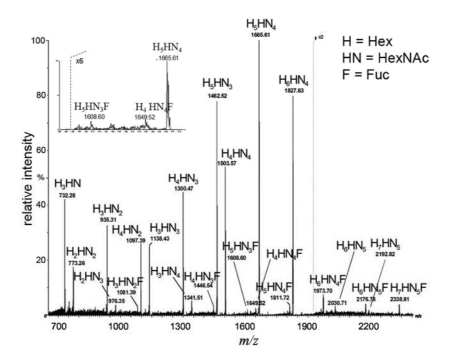
For large molecular weight HMO, which can possess more complex branching sites, as well as multiple fucosylation sites, the structures are predominantly difficult to define, also due to isobaric mixtures. This is frequently solved by the application of liquid chromatography (LC) protocols, which allow isomer separation prior to MS analysis [2,3]. More direct is the MALDI Q-TOF CID *de novo* approach. We show that the already known oligosaccharide structures can be assigned according to specific diagnostic ions in combination with the HMO biosynthetic rules, but also, novel structure types can be proposed *de novo*. The additional advantages of the MALDI Q-TOF MS analysis are the sensitivity, speed, high dynamic range and ability to analyse data in more detail on the same sample spot once the preliminary data analysis is completed.

2. Results and Discussion

2.1. MALDI TOF (Matrix-Assisted Laser Desorption/Ionization Time-of-Flight) Mapping of Complex Oligosaccharide Mixtures

The MALDI TOF map of the sample containing human milk oligosaccharides is shown in Figure 1. Twenty two different ions were detected that could be assigned according to monosaccharide compositions (in terms of the building blocks, Hex (H, hexose), HexNAc (HN, *N*-acetylhexosamine) and dHex (F, deoxyhexose)). The oligosaccharides ranged from tetra- to dodeca-saccharides, including eight fucosylated structures. The MALDI spectrum was acquired from the sample without further purification procedures as an initial oligosaccharide mapping experiment. The most intense ions represent octa-, nona- and deca-saccharides. The undeca- and dodeca-saccharide-related ions were at a lower intensity. Tetra- to hepta-saccharide series were detected, as well, with similar relative abundances.

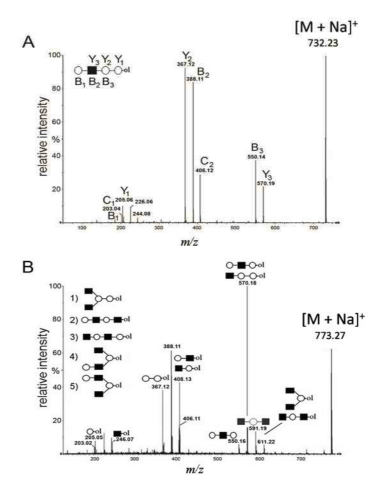
Figure 1. The (+) matrix-assisted laser desorption/ionization (MALDI) quadrupole-time-of-flight (Q-TOF) spectrum of the sample containing human milk oligosaccharide alditols. The m/z values represent $[M + Na]^+$ ions.



2.2. De Novo MALDI Q-TOF CID Data Analysis Using Diagnostic Ions

The fragmentation spectra (MS/MS) were assigned using the theoretical cleavage ions according to Domon and Costello [23]. In Figure 2A, the CID spectrum of the basic tetrasaccharide precursor ion at m/z = 732 is shown. Due to the reduced reducing end, it was possible to unambiguously assign Y₁₋₃ ions for the full sequence information, along with B₁₋₃ ions (m/z = 185, 388 and 550), in agreement with the known biosynthetic core structure of human milk oligosaccharides (HMO) belonging to LNT and LNnT tetrasaccharides. No intra-ring fragmentation was observed under these conditions.

Figure 2. The (+) MALDI Q-TOF collision-induced dissociation (CID) spectra of two low mass oligosaccharide alditol ions from the mixture (Figure 1). (**A**) Tetrasaccharide alditol lacto-*N*-tetraose (732.23); and (**B**) isobaric mixture of tetrasaccharide alditols (773.27) of the sum composition Hex₂HexNAc₂.



In Figure 2B, the CID spectrum of parent ion at m/z = 773 assigned to H₂HN₂ is shown. The cleavage ions at m/z = 205, 246, 367, 408, 550, 570, 591 and 611 are postulated to be the partial structures shown. From the ion at m/z = 367, Structure 1 could be postulated. For the ion at m/z = 408, two tetrasaccharide structures (2 and 3) are possible, according to HMO biosynthetic rules. Structure 2 corresponds to the ion at m/z = 246. Structure 3 was not confirmed, due to the ambiguity of fragment ions at m/z = 205 and 570, which could also be generated from Structure 1. It is important to note that both Structure 2 and 3 could have been built upon enzymatic or chemical hydrolysis of HMO. Furthermore, both Structures 4 and 5 could give rise to fragment ions at m/z = 408 and 611.

In Figure 2, MS analysis of oligosaccharides when isomeric mixtures are present is presented. For manual *de novo* tandem MS data analysis, the pragmatic first step is the assignment of diagnostic ions (Table 1). Accordingly, the fragment ion at m/z = 367 as a diagnostic one is relevant for Structure 1, indicating the presence of the lactose core on the reducing end. The fragment ion at m/z = 246 is diagnostic for the presence of HexNAc on the reducing end, typical for the truncated structure. The diagnostic ion at m/z = 408 in combination with other diagnostic ions, such as at m/z = 205 (indicating the presence of a hexose on the reducing end) and at m/z = 246 (indicating the presence of an N-acetylhexosamine on the reducing end), represents the "conditional" diagnostic ion, because

additional information was necessary to fully assign its structure. In the case of isomeric mixtures, this information can be probed with MS³ experiments on appropriate instrument types for full structural elucidation. A list of diagnostic and conditional diagnostic ions used in this study is shown in Table 1. Ions at m/z = 246, 367 and 794 in Table 1 are diagnostic ones. Additionally, two examples of non-diagnostic ions are given, illustrating that no significant information can be deduced from them, alone or in combination with other fragment ions.

Table 1. Diagnostic and conditional diagnostic ions found and monosaccharide symbols used in this study. ■ denotes *N*-acetylglucosamine, ■ galactose and ● glucose.

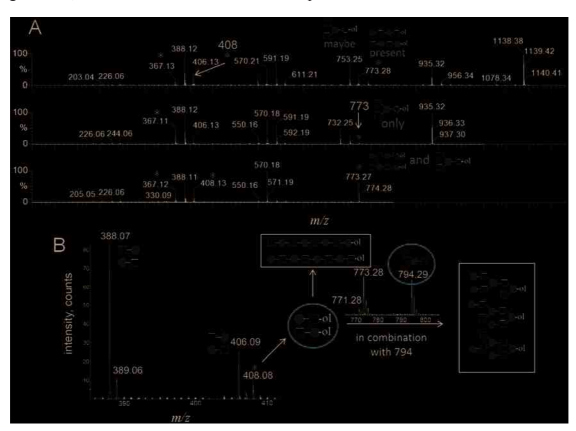
Type of ions	m/z	Туре	Structure
non-diagnostic ions	753	B ₄ or internal B/Y fragment ion	
non-diagnostic ions	771	C ₄ or internal C/Z fragment ion	
	205	hexose at reducing end	○-ol ○-ol
diagnostic ions	246	HexNAc at reducing end (non-biosynthetic)	E-ol
angueste tene	367	contains biosynthetic core lactose unit	ol
	794	branching at $\text{Hex}_{n\geq 4}$	D>1
	408	non-biosynthetic reducing end	no 246
	408	structure	no 205
	611	non-biosynthetic reducing end	no 246
conditional diagnostic ions	011	structure	no 205
	732	Y4 with core lactose—linear	no 773 and
	134	oligosaccharide	no 794
	773	branching at Hex ₂	yes 367 and
		(biosynthetic structure)	no 408

Efficient fragmentation has been obtained by MALDI Q-TOF CID (Micromass, Manchester, UK) from major parent ions at m/z = 732, 773, 935, 1097, 1138, 1300, 1462, 1503, 1665 and 1827. The parent ion at m/z = 1341 fragmented poorly. The parent ion at m/z = 1503, was further tested along low-intensity parent ions at m/z = 1608, 1973 and 2338 on an AB SCIEX QSTAR® Pulsar i instrument (AB SCIEX, Toronto, ON, Canada).

In Figure 3A, the fragmentation spectra of parent ions at m/z = 773, 935 and 1138 are depicted. A structural assignment has been carried out using the diagnostic ion at m/z = 367 (blue asterisk) and conditional diagnostic ions at m/z = 408 (red asterisk) and 773 (green asterisk) (see Table 2). For the parent ion at m/z = 935, the assignment to two possible isomeric pentasaccharide structures is ambiguous. Based on the diagnostic ion at m/z = 367 and the conditional diagnostic ion at m/z = 773 (which in the

absence of the conditional diagnostic ion at m/z = 408 and the presence of the diagnostic ion at m/z = 367 indicates the presence of a branching point at Hex₂ (see Table 2)), the presence of a single tetrasaccharide pattern (structure type) could be postulated. However, the location of the final galactose residue cannot be deduced from our data. Furthermore, the presence of a linear pentasaccharide structure cannot be proven in the presence of branched structure(s). Finally, in the case of the CID of parent ion at m/z = 1138, we detected all three ion types at m/z = 367, 408 and 773. According to the diagnostic ion at m/z = 794, a branching point is at Hex₄. A branching point at Hex₂ (supported by the presence of the conditional diagnostic ion at m/z = 773) could not be deduced due to the presence of another conditional diagnostic ion at m/z = 408.

Figure 3. The assignment of CID MS data using diagnostic (367 and 794) and conditional diagnostic (408 and 773) fragment ions. (**A**) Ions at m/z = 773, 935 and 1138; and (**B**) zoom-in of the mass range of 385–410 and 770–800 Da. Possible structures are all in accord with the fragmentation pattern. \blacksquare denotes *N*-acetylglucosamine, \bigcirc galactose and \bigcirc glucose. *, * and * denote ions relevant to the explanation in the text.



possible, due to galactose 1-3 and 1-4 linkages, and most of them are further fucosylated in vivo with fucose 1-2, 1-3 or 1-4. Table 2. Confirmed and proposed oligosaccharide alditol structures along with respective diagnostic ions. Additional branched isomers are N-acetylglucosamine, ■ galactose, ● glucose and ▲ fucose. * structures that could be neither confirmed nor excluded. * present in very small amounts.

Monosaccharide				;
composition	Biosynthetic	Truncated with lactose core	Truncated without lactose core	Diagnostic ions
H ₃ HN (732)	10-0-0-1	ı	•	367
H ₂ HN ₂ (773)	1	10-0-01	01	246, 367, 408
H ₃ HN ₂ (935)	ı		1	367, 773
H_2HN_3 (976)	•	_	lo	1
H_4HN_2 (1097)	lo	-	-	367
H ₃ HN ₃ (1138)	ı	10-0-01*	ol	246, 367, 408, 773, 794
H ₄ HN ₃ (1300)	1		1	367, 773, 794
$\mathrm{H_{3}HN_{4}}\ (1341)$		[o-o-o]		773, 794
H_5HN_3 (1462)	10-7			367

Table 2. Cont.

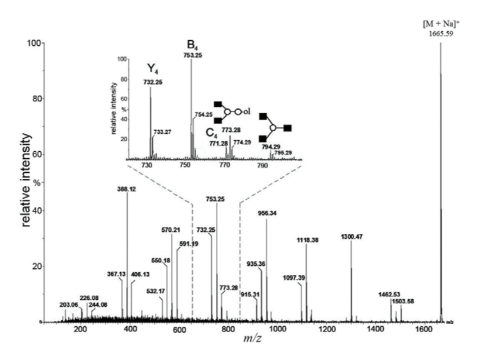
Monosaccharide composition	Biosynthetic	Truncated with lactose core	Truncated without lactose core	Diagnostic ions
H ₄ HN ₄ (1503)	1	10-0-01	lo	367, 408, 773, 794
H ₅ HN ₄ (1665)	1		1	367, 773, 794
H_6HN_4 (1827)	lo	-	1	367
H ₅ HN ₃ F (1609)	A	1	-	878
H ₆ HN ₄ F (1973)	A	1	•	1243
H ₇ HN ₅ F (2338)	* 0-5	ı		1243 ?

In Figure 3B, the inserts of the MALDI Q-TOF CID spectrum of the parent ion at m/z = 1503 are shown, obtained by using the "enhance all" function on the AB SCIEX QSTAR® Pulsar i (see the Experimental Section). In this experiment, the conditional diagnostic ion at m/z = 408 was detected (left insert) and allowed one to deduce the presence of two further oligosaccharide structures in the m/z = 1503 parent ion mixture. Furthermore, in combination with the diagnostic ion at m/z = 794, the existence of three additional truncated isomeric structures could be proposed (if we restrict our analysis to the reported HMO structures, up to lacto-N-decaose), but even more isomeric structures are possible, when Type I and Type II branches are considered [19]. Therefore, at least five novel structure types could be added to the list of possible isomeric octasaccharide structures present in the parent ion at m/z = 1503, most of them as truncated isomers.

2.3. Analysis of Branched Higher Molecular Weight Oligosaccharides

In Figures 2 and 3, we showed how the existence of truncated structures can exponentially increase the number of theoretically possible isomers present in the isobaric parent ion mixture. In Figure 4, the MALDI Q-TOF CID spectrum of the parent ion at m/z = 1665 and the conditional diagnostic ion at m/z = 408 was not present, but the positive structural evidence was obtained by two important fragment ions, the conditional diagnostic ion at m/z = 773 and the diagnostic ion at m/z = 794. The diagnostic ion at m/z = 367 together with the conditional diagnostic ion at m/z = 773 provides evidence of branched nonasaccharide structures with a branching point at Hex₂. The diagnostic ion at m/z = 794 indicates the presence of branched nonasaccharide structures with a branching point at Hex₄. By applying biosynthetic rules and restricting our analysis to the reported HMO structures (up to lacto-N-decaose), only three out of 13 theoretical branched structure types remain possible.

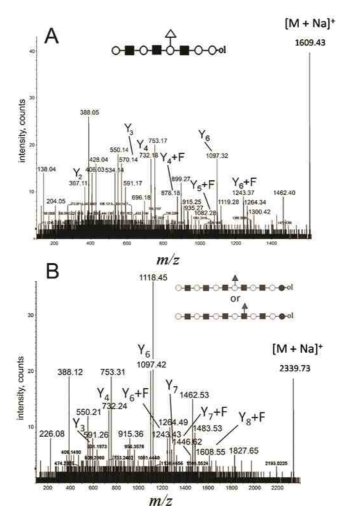
Figure 4. Fragmentation spectrum of the parent ion at m/z = 1665.59. The analysis of fragment ions revealed the presence of isobaric nonasaccharide alditols.



2.4. MALDI Q-TOF CID of Low-Intensity and High Molecular Weight Parent Ions

The ability to sequence high molecular weight and low-intensity precursor ions with MALDI Q-TOF CID is illustrated in Figure 5, where the CID spectra of parent ions at m/z = 1608 (Figure 5A) and 2338 (Figure 5B) are shown. The precursor ion at m/z = 1608 is of low abundance, and the precursor ion at m/z = 2338 is the highest m/z ion in the primary mixture (Figure 1); both are fucosylated. Both low abundant precursor ions delivered on the AB SCIEX QSTAR® Pulsar i, with the "enhance all" function enabled, good evidence and sufficient sequence information to deduce linear octasaccharide and dodecasaccharide HMO chains. Furthermore, a significant amount of fucosylated product ions were detected. We were able to clearly assign the fucosylation site to Hex4 in the case of fucosylated octasaccharide (Figure 5A). In the case of fucosylated dodecasaccharide (Figure 5B), fragment ions corresponding to either Y_5 or $Y_5 + F$ ions were not detectable, while the $Y_{6-8} + F$ ion series was visible. Accordingly, the position of the fucose residue in the dodecasaccharide HMO structure was not assigned, indicating the limits of sensitivity at sequencing low intensity, high molecular weight oligosaccharides.

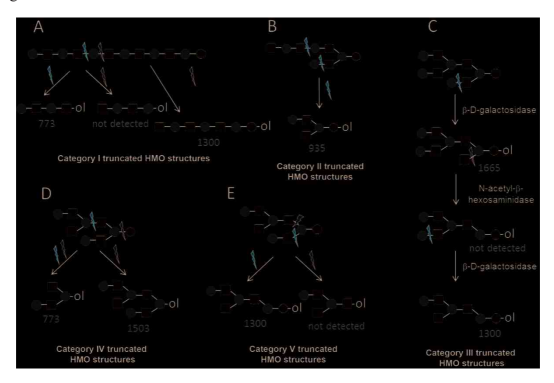
Figure 5. The potentials and limits of fragmentation data using low abundant ions at high m/z values. (A) Parent ion at m/z = 1608.43; and (B) parent ion at m/z = 2338.73. The spectra were acquired on the AB SCIEX QSTAR® Pulsar i (see the Experimental Section).



2.5. Analysis of HMO Truncated Structures

A number of oligosaccharides have been identified as truncated structures by their MS/MS patterns. In Figure 6, the possible mechanism of the generation of such structures is presented. The proposed reaction schemes in five practical categories (I–V) for linear and branched species is illustrated, following single or multiple truncation, based on fragmentation data and the biosynthetic considerations of the elongation of the lactose core by *N*-acetyllactosamine units in a linear or branched fashion. Most evident are oligosaccharide structures with an odd monosaccharide composition, which could possibly arise also by the exoglycosidase cleavages [6].

Figure 6. The formation of truncated oligosaccharide structures in the human milk oligosaccharide (HMO) fraction. ↑ and depict various possible hydrolysis events, each of which, alone (**A**–**D** right, **E**) or combined (**D** left), yield different oligosaccharide structure types shown in the figure. ■ denotes *N*-acetylglucosamine, ○ galactose and ● glucose.



Category I (Figure 6A) is the simplest type of hydrolysis. The resulting oligosaccharides would give rise to a fragment ion at m/z = 408. Furthermore, these hydrolytic products could all give rise to odd-numbered linear oligosaccharide structures. Similar to Category I, the same hydrolytic events could occur on branched oligosaccharide structures, giving rise to odd-numbered Category II branched oligosaccharide structures (Figure 6B). Category III represents possible exoglycosidase events, described previously [6]. The sequential trimming of monosaccharide units on the nonreducing termini can yield structures, such as H_5HN_4 (m/z = 1665), which is the most abundant component in our sample (see Figure 1). In the case of the nonasaccharide, the trimming of another terminal galactose residue seems favourable, leading to respective H_4HN_4 structures, while the trimming of a N-acetylglucosamine (GlcNAc) residue seems unfavourable or leads to a quickly

degradable product. Category IV hydrolysis depicts the type of structures arising from hydrolytic events on the nonreducing end of glucose. All parent ion mixtures that gave rise to the fragment ion at m/z = 611 could theoretically contain these type of structures (parent ions at m/z = 773, 1138 and 1503), although they seem to be present in very low amounts. Category V hydrolysis shows the remaining two theoretical hydrolytic fragments. We detected some of these types of fragments unambiguously in parent ions at m/z = 1138 and 1300, albeit they were present only near the detection limit in the latter.

2.6. Fragmentation of the Ion at m/z = 1341

The assignment of the parent ion at m/z = 1341.51 is based on the fragment ions at m/z = 794 and 773 (spectrum not shown). This type of branching pattern has been described before, but is still considered novel [2,24], perhaps due to its incomplete characterization and rare observance in the published literature. One novel and five additional possible HMO structures were postulated recently to be based on this structure [2]. This structure type has been indeed resolved previously by FAB (fast atom bombardment) MS and NMR spectroscopy on an example of lacto-N-decaose in 1988 [20], as reviewed recently [25]. Having proven the existence of this branching pattern in our sample and HMO mixtures, in general, it is indicated as a possible isomer in the isobaric parent ion mixtures containing the diagnostic fragment ion at m/z = 794. It would be interesting to prove the existence of these isomers in the high molecular weight HMO fractions and to determine the relative abundance of this branching pattern in HMO in general. One line of study in this direction has been carried out by Amano *et al.* [19], in which the decaose structure has been best described so far in terms of its fucosylation patterns.

2.7. General Conclusions about Identified HMO Structures

After having analysed all the CID data, it became possible to summarize the data (Table 2), as well as to categorize the postulated structures, depending on the monosaccharide composition of oligosaccharides (Table 3). Accordingly, all HMO with the general formula $H_{n+1}HN_n$ were identified as linear. The odd-numbered HMO with the general formula $H_{n+1}HN_n$ were all truncated on the nonreducing termini only. The HMO with the general formula H_nHN_n were most diverse. Unlike in the first two groups, this group always contained the fragment ion at m/z = 408, indicative of truncation on the reducing termini. They also had multiple truncations on the non-reducing termini. Surprisingly, they are not the most abundant structures in the spectrum and are only about half as abundant as the previous two groups. Finally, the HMO of the general formula $H_{n-1}HN_n$ were of very low abundance. They also yielded the m/z = 408 fragment ion, although it could not be detected for H_3HN_4 , most likely due to the low intensity of fragment ions obtained in the CID spectrum. Monofucosylated parent ions corresponding to all these groups were detected, except for the $H_{n-1}HN_n$ group. Further and more detailed studies will be carried out to study these phenomena. In particular, quantitative analysis should provide important insight about the abundance of individual HMO isomers.

Table 3. Categorization of HMO structures identified in this study based on their monosaccharide composition. ■ denotes *N*-acetylglucosamine, ② galactose and ③ glucose. * structures that could be neither confirmed nor excluded. * present in very small amounts.

Monosaccharide composition	HMO structure types identified
$H_{n+2}HN_n$	
$H_{n+1}HN_n$	
$\mathrm{H_{n}HN_{n}}$	
$H_{n-1}HN_n$	

3. Experimental Section

A pool of human milk was obtained from Donor 0, a Le^a non-secretor. The high molecular weight chromatographic fraction was obtained by gel permeation Biogel P-4 chromatography and submitted to mild hydrolysis to remove fucose and sialic acid, as described in Bruntz *et al.* [20]. The oligosaccharide fraction was reduced with NaBH₄ and analysed using MALDI MS in the positive ion mode.

2,5-Dihydroxybenzoic acid (DHB) (Sigma, Steinheim, Germany) was used as a MALDI matrix. For MALDI TOF analysis, $\gamma=20$ mg/mL, dissolved in H2O/ACN (acetonitrile) 1:1 (ν/ν), was used. For MALDI Q-TOF analysis, $\gamma=80$ mg/mL in H2O/ACN 70%:30% (ν/ν) was used. The oligosaccharide sample was dissolved in ddH2O, and 1 μ L of the sample was mixed with 1 μ L of matrix solution on the MALDI target and allowed to dry. MALDI TOF MS experiment was performed on a TofSpec 2E instrument (Micromass, Manchester, UK), equipped with a LeCroy (Teledyne Lecroy Inc., New York, NY, USA) digitizer LSA1000 with a 2 GHz acquisition rate. The majority of the MALDI Q-TOF CID experiments were conducted on a prototype Micromass instrument. Additional MALDI Q-TOF CID experiments were performed on a commercial instrument, AB SCIEX QSTAR® Pulsar i.

In general, there are differences in experimental parameters between these three instruments, as axial (MALDI TOF) and orthogonal (MALDI Q-TOF) instruments have different modes of operation. For MALDI TOF MS, the minimum laser fluence that allowed stable signal collection was used, in order to increase the spectral resolution, to minimize possible detector saturation and to minimize in-source and post-source decay (ISD and PSD). For the prototype MALDI Q-TOF instrument from Micromass, as well as the commercial AB SCIEX QSTAR® Pulsar i, no

sample-to-sample signal optimization was required, as the instrument default method was robust enough. The practical difference in the mode of operation between MALDI TOF and MALDI Q-TOF instruments is the high laser fluence requirement for MALDI Q-TOF instruments. This is technically necessary because orthogonal instruments are less efficient in the ion transport than axial instruments, and therefore, more ions need to be produced within the oMALDI (orthogonal MALDI) source. Two practical consequences of higher laser fluence used in our hands were: (i) the ability to use higher $\gamma(DHB)$ as a matrix solution on MALDI Q-TOF instruments; and (ii) the use of DHB as a matrix was required, as other matrices used in our laboratory did not give any signal on a MALDI Q-TOF instrument. Despite these technical and experimental differences, the MS1 spectra on all three instruments were very similar. However, in our hands, the MALDI TOF oligosaccharide map of the analysed chromatographic fraction acquired on the Micromass TofSpec 2E was of slightly higher quality in terms of signal intensity and is, thus, presented here.

MS/MS spectra were obtained by collisionally-induced dissociation (CID) using Ar as a collision gas on MALDI Q-TOF instruments under conditions like low-energy CID. The collision energy that produced the optimal fragmentation pattern across the whole m/z range for a given parent ion was adjusted to be proportional to the parent ion m/z value. The AB SCIEX QSTAR® Pulsar i instrument has a unique additional mode of operation in MS/MS mode, in which it can amplify the intensity of the detected fragment ions by using a "pulse" function option, which allows the user to select which m/z range in the CID spectrum needs to be "enhanced" (in terms of fragment ion intensity). When this option is chosen, the built-in control software automatically recalculates "pulsing" windows, which determine which fragment ion packets are sent to the TOF analyser more often, thus increasing their relative intensities in the final CID spectrum. The "enhance all" option in the control software allows enhanced product ion detection over the whole m/z range of the CID spectrum. This option in the recording of the CID spectra of low-intensity parent ions was functional (Figure 5), as well as in the recording of diagnostic low-intensity fragment ions (Figure 3B).

4. Conclusions

De novo analysis of tandem MS data is described as an option for an in-depth view on already known or new structures. Since many molecular ions represented isomeric mixtures of oligosaccharides, a rationale for the assignment based on the biosynthetic pathway was applied. Oligosaccharide structures not found in databases were proposed for assignment according to specific diagnostic ions detected in fragmentation spectra. In the mixture of linear and branched HMO structures within a single precursor ion m/z value, the isomeric components were assigned using a diagnostic ion analysis technique. A minor amount of fucosylated oligosaccharides was found, where fucosylation occurred on a single site. In summary, four non-fucosylated linear HMO and three monofucosylated linear HMO were characterized. About twenty branched structure types were proposed, as in agreement with MS/MS data, known HMO structures and hydrolytic patterns. These structures are furthermore isomeric with many possible biosynthetic HMOs and, when taking into account the 1-3 and 1-4 galactose linkage isomers, over a hundred HMO structures can be assumed to be present in the analysed chromatographic fraction. The summarized results are depicted in Tables 2 and 3. Truncated structures without glucose are omitted for clarity.

Although we were able to determine the presence of isomeric structures in our sample, our data also illustrate the limitations of MALDI Q-TOF CID technique in their full characterization. It is important to note that mass spectrometry in general can overcome these limitations in several ways: (i) by using an additional liquid chromatography (LC) or capillary electrophoresis (CE) separation step prior to MS analysis; (ii) by using high-energy CID to generate cross-ring cleavages; and (iii) by using MSⁿ fragmentation techniques. Additional LC separation step prior to MALDI Q-TOF MS/MS analysis can be done in off-line mode [26]. In on-line mode, the applicability of LC-ESI-MS/MS [2,3,17] and CE-LIF (laser induced fluorescence)-MSⁿ [16] in the analysis of HMO has been well documented. Unlike low-energy CID, used in MALDI Q-TOF instruments, high-energy CID used in MALDI TOF-TOF instruments typically yields cross-ring cleavages, which can carry information about the branching pattern of the parent ion [14,15]. Finally, the MSⁿ technique has been successfully applied to the analysis of HMO [19]. In general, different MS instrumentation setups are required for obtaining the maximum amount of structural information about the analysed sample and these often require either more time for analysis (LC-MS, CE-MS, MSⁿ), a derivatization step prior to MS analysis (CE-MS, MSⁿ) [16,19,27] and more time or additional software tools to analyse the large amount of data these techniques yield [2,18]. Traditionally, MALDI MS has been advantageous in the high-throughput MS1 screening of biological samples, and with modern instrumentation allowing the acquisition of high-quality MALDI MS/MS data, this property of MALDI MS can be extended to the high-throughput screening of tandem MS data, as well.

The advantages of MALDI Q-TOF CID for structural elucidation of complex oligosaccharides are the precursor ion selection properties, the ability to sequence very low abundant parent ions and the speed of the MS/MS data acquisition. The majority of MS1 signals could be sequenced with the prototype Micromass MALDI Q-TOF instrument, and all detectable signals could be sequenced with the commercial AB SCIEX MALDI Q-TOF instrument. This technical advancement alone allows for a huge jump in the data throughput of a glycomic research laboratory, where MALDI MS is the method of choice for all preliminary experiments related to the structural characterization of complex oligosaccharide mixtures. Finally, the advantages of the MALDI Q-TOF MS/MS indicate its applicability to high-throughput sample analysis, requiring new software tools for faster data analysis [18,28].

Future studies on high molecular weight HMO should provide better insight on biologically interesting data, like the quantitation of isomers and the reexamination of sequences, which indicate the biosynthetic pathways involved. Quantitative studies are necessary in order to determine the relative abundance of particular isomers, *i.e.*, are they all present in equal, random or more selective amounts, where the presence of certain isomers can be neglected.

Acknowledgments

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Author Contributions

M.J. performed experiments on the TofSpec 2E and QStar Pulsar i instruments, analysed the data and wrote the manuscript. R.T.W. designed the Micromass prototype MALDI Q-TOF instrument and performed the related experiments in this work. He also contributed to the Experimental Section. G.P. developed the isolation procedure for HMO and prepared the sample for analysis. J.P.K. designed the experiment and wrote the manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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Interleukin-6 Receptor rs7529229 T/C Polymorphism Is Associated with Left Main Coronary Artery Disease Phenotype in a Chinese Population

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Abstract: Left main coronary artery disease (LMCAD) is a particular severe phenotype of coronary artery disease (CAD) and heritability. Interleukin (IL) may play important roles in the pathogenesis of CAD. Although several single nucleotide polymorphisms (SNPs) identified in IL related genes have been evaluated for their roles in inflammatory diseases and CAD predisposition, the investigations between genetic variants and CAD phenotype are limited. We hypothesized that some of these gene SNPs may contribute to LMCAD phenotype susceptibility compared with more peripheral coronary artery disease (MPCAD). In a hospital-based case-only study, we studied IL-1A rs1800587 C/T, IL-1B rs16944 G/A, IL-6 rs1800796 C/G, IL-6R rs7529229 T/C, IL-8 rs4073 T/A, IL-10 rs1800872 A/C, and IL-10 rs1800896 A/G SNPs in 402 LMCAD patients and 804 MPCAD patients in a Chinese population. Genotyping was done using matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS) and ligation detection reaction (LDR) method. When the IL-6R rs7529229 TT homozygote genotype was used as the reference group, the CC or TC/CC genotypes were associated with the increased risk for LMCAD (CC vs. TT, adjusted odds ratio(OR) = 1.46, 95% confidence interval (CI) = 1.02-2.11, p = 0.042; CC + TC vs. TT, adjusted OR = 1.31, 95% CI = 1.02-1.69, p = 0.037). None of the other six SNPs achieved any significant differences between LMCAD and MPCAD. The present study suggests that IL-6R rs7529229 T/C functional SNP may contribute to the risk of LMCAD in a Chinese population. However, our results were limited. Validation by a larger study from a more diverse ethnic population is needed.

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1. Introduction

Coronary artery disease (CAD) is the leading cause of mortality and morbidity in many countries including China [1,2]. Multifaceted phenotypes of CAD (number of involved vessels, location of lesions, severity of diameter narrowing, and morphology of lesions) have different mechanisms. Left main coronary artery disease (LMCAD) is a particular severe phenotype of CAD because it is associated with higher risk of fatal cardiovascular events [3]. Thus, any information in detecting the asymptomatic relatives of these patients can be useful for primary prevention.

LMCAD was heritability and frequently shared by siblings with CAD [4]. Fischer *et al.* [5] reported a stronger genetic component in LMCAD phenotype. However, Kolovou *et al.* [6] found no significant difference in cholesteryl ester transfer protein (CETP) allele frequency or genotype distribution among

LMCAD and more peripheral coronary artery disease (MPCAD) patients. Thus, more investigations are needed in detecting association between genetic variants and CAD phenotype.

Inflammation is important in the initiation, progression, and clinical outcome of CAD [7,8]. Interleukin (IL) plays a key role in the inflammatory response, immune regulation and development of CAD. IL-1 family (including IL-1A, IL-1B *et al.*) plays a major role in inflammation by affecting antigen recognition patterns and lymphocyte function [9]. IL-6 is a multi-functional cytokine involved in various contradictory processes, a high circulating concentration of IL-6 is associated with increased risk of CAD [10]. IL-6 binds to IL-6R and then activates an intracellular signaling cascade leading to the inflammatory response [11]. IL-8 plays a key role in the development of atherosclerotic plaques [12]. IL-10 has anti-inflammatory and immunosuppressive effects by decreasing the production of pro-inflammatory mediators, exerts important protective effects on atherosclerotic lesion development [13].

Recently, a link between CAD genetic susceptibility and the response to inflammatory signaling has been established [8,14]. Growing evidence for heritability of pro-inflammatory state suggests that individual genetic background also modulate the development of CAD and its magnitude [15,16]. Despite well-described associations between inflammatory-related genetic variation and susceptibility to CAD, there is a paucity of data regarding to the phenotype of CAD.

The identification of novel genetic variants for use in assessing early risk of LMCAD have potentially important clinical implications, such as identifying high-risk individuals and adapting therapeutic management to the individual's genetic make-up. Single nucleotide polymorphisms (SNPs) strongly influence the plasma levels and biological activity of the corresponding proteins. On the basis of the biological and pathologic significance of IL, we performed a genetic association study on the SNPs of *IL-1A*, *IL-1B*, *IL-6*, *IL-6R*, *IL-8* and *IL-10*.

The objective of this investigation was to evaluate the association between *IL* SNPs and LMCAD susceptibility compare with MPCAD in a hospital-based case-only study. We performed genotyping and analyses for the seven SNPs in a cohort of 1206 CAD patients (402 LMCAD patients and 804 MPCAD patients) in a Chinese population.

2. Results and Discussion

2.1. Characteristics of the Study Population

The demographic and clinical characteristics of all subjects are summarized in Table 1. Among 1206 DNA samples, the seven SNPs were successful ranging from 98.59% to 99.92% (Table 2). The mean age for LMCAD patients (62.24 \pm 8.66) is significantly higher than for MPCAD patients (60.14 \pm 8.96), p < 0.001 (Table 1). There are no significant difference for sex and mean body mass index (BMI), family history of CAD, previous smoker, hypertension, hyperlipidemia, diabetes mellitus, mean ejection fraction, circumflex branch of left coronary artery system, right coronary artery system, off-pump coronary artery bypass grafting (OPCAB) or conventional coronary artery bypass grafting (cCABG) for LMCAD patients and MPCAD patients (Table 1). LMCAD patients are more likely to have three disease territories than MPCAD patients (93.3% vs. 87.7%, p = 0.003) (Table 1).

2.2. Associations between the Seven SNPs and Risk of LMCAD and MPCAD

The genotype frequencies of the *IL-6R* rs7529229 T/C polymorphism were 34.1% (TT), 49.0% (TC) and 16.9% (CC) in LMCAD patients, and 40.4% (TT), 45.9% (TC) and 13.7% (CC) in MPCAD patients (p = 0.027) (Table 3). When the *IL-6R* rs7529229 TT homozygote genotype was used as the reference group, the CC genotype was associated with the increased risk for LMCAD (CC vs. TT, adjusted OR = 1.46, 95% CI = 1.02–2.11, p = 0.042). The TC genotype was not associated with the risk for LMCAD (TC vs. TT, adjusted OR = 1.26, 95% CI = 0.97–1.65, p = 0.088). In the dominant model, when the *IL-6R* rs7529229 TT genotype was used as the reference group, the TC/CC genotypes were associated with the increased risk for LMCAD (TC/CC vs. TT, adjusted OR = 1.31, 95% CI = 1.02–1.69, p = 0.037). The polymorphism was not associated with the risk for LMCAD in recessive genetic models (CC vs. TT/TC, adjusted OR = 1.29, 95% CI = 0.92–1.79, p = 0.140).

None of the other six polymorphisms achieved a significant difference in the genotype distributions between cases and controls. The six polymorphisms were not associated with the risk for LMCAD/MPCAD both in homozygote comparison, heterogeneity comparison, dominant genetic model and recessive genetic models (Table 3).

Table 1. Patient Demographics and Risk Factors with coronary artery disease (CAD) (Left main coronary artery disease (LMCAD) and more peripheral coronary artery disease (MPCAD)).

Variable	LMCAD b	MPCAD ^c	<i>p</i> -Value	All CAD
v ariable	(n = 402)	(n = 804)	p-value	(n = 1206)
Mean age, y	62.24 (±8.66)	60.14 (±8.96)	< 0.001	60.84 (±8.91)
Woman, %	67 (16.7)	170 (21.1)	0.065	237 (19.7)
Mean BMI ^a , kg/m ²	25.55 (±3.16)	25.75 (±3.13)	0.299	25.68 (±3.14)
Family history of CAD, %	21 (5.2)	32 (4.0)	0.323	53 (4.4)
Previous smoker, %	211 (52.5)	422 (52.5)	1.000	633 (52.5)
Hypertension, %	261 (64.9)	530 (65.9)	0.732	791 (65.6)
Hyperlipidemia, %	275 (68.4)	573 (71.3)	0.290	848 (70.4)
Diabetes mellitus, %	125 (31.1)	271 (33.7)	0.363	396 (32.8)
Mean ejection fraction, %	60.25 (±7.96)	59.55 (±8.76)	0.177	59.78 (±8.50)
Disease territories, %				
1–2	27 (6.7)	99 (12.3)	0.003	126 (10.4)
3	375 (93.3)	705 (87.7)	0.003	1080 (89.6)
LMCAD, %	402 (100)	0 (0)	_	402 (33.3)
Anterior descending artery system, %	386 (96.0)	793 (98.6)	0.004	1179 (97.8)
Circumflex branch of left coronary artery system, %	373 (92.8)	744 (92.5)	0.876	1117 (92.6)
Right coronary artery system, %	374 (93.0)	751 (93.4)	0.807	1125 (93.3)
OPCAB/cCABG ^d	209/193	438/366	0.414	647/559

^a BMI: body mass index; ^b LMCAD: left main coronary artery disease; ^c MPCAD: more peripheral coronary artery disease; Bold values are statistically significant (p < 0.05); ^d OPCAB: off-pump coronary artery bypass grafting; cCABG: conventional coronary artery bypass grafting.

Table 2. Primary information for seven genotyped single nucleotide polymorphisms (SNPs).

Genotyped SNPs	Chr a	Regulome DB Score ^b	TFBS c	Splicing (ESE or ESS)	Location	MAF ^d for Chinese in Database	% Genotyping Value
<i>IL-1A</i> : rs1800587 C/T	2	5	Y	Y	5' UTR	0.073	99.92
<i>IL-1B</i> : rs16944 G/A	2	1f	Y	_	5' near gene	0.453	99.83
<i>IL-6</i> : rs1800796 C/G	7	4	Y	_	ncRNA	0.233	99.75
<i>IL-6R</i> : rs7529229 T/C	1	2b	_	_	intron	0.442	98.59
<i>IL-8</i> : rs4073 T/A	4	2b	Y	_	5' near gene	0.389	98.84
<i>IL-10</i> : rs1800872 A/C	1	5	Y	_	5' near gene	0.238	99.83
<i>IL-10</i> : rs1800896 A/G	1	6	Y	_	5' near gene	0.059	99.83

^a Chr, chromosome; ^b http://www.regulomedb.org/; ^c TFBS: Transcription Factor Binding Site; ^d MAF: minor allele frequency.

Table 3. Main effects of SNPs on LMCAD risk.

	Genotyping	Genotyping (AA/AB/BB) a	AB vs. AA	BB vs. AA	BB vs. (AA + AB)	(BB + AB) vs. AA	
Genotyped SNPs	LMCAD $(n = 402)$	\mathbf{MPCAD} $(n = 804)$	Adjusted OR (95% CI); p-Value	p Trend			
<i>IL-1A</i> : rs1800587 C/T	331/67/3	645/145/14	0.91 (0.66–1.26);	0.42 (0.12–1.42);	0.43 (0.12–1.50);	0.87 (0.64–1.19);	0.314
<i>IL-1B</i> : rs16944 G/A	104/203/95	212/397/193	1.04 (0.78–1.39); 0.794	1.01 (0.72–1.42); 0.974	0.98 (0.74–1.30); 0.890	1.03 (0.78–1.35); 0.841	0.948
<i>IL-6</i> : rs1800796 C/G	182/174/46	358/348/95	0.97 (0.75–1.25); 0.797	0.96 (0.65–1.43); 0.856	0.98 (0.67–1.43); 0.916	0.97 (0.76–1.23); 0.782	0.970
IL-6R: rs7529229 T/C	135/194/67	320/364/109	1.26 (0.97–1.65); 0.088	1.46 (1.02–2.11); 0.042	1.29 (0.92–1.79); 0.140	1.31 (1.02–1.69); 0.037	0.080
<i>IL-8</i> : rs4073 T/A	127/198/71	280/375/141	1.18 (0.90–1.56); 0.226	1.14 (0.80–1.62); 0.482	1.03 (0.75–1.41); 0.862	1.17 (0.91–1.52); 0.231	0.545
<i>IL-10</i> : rs1800872 A/C	155/197/49	336/371/96	1.15 (0.89–1.49); 0.277	1.08 (0.73–1.60); 0.716	1.00 (0.69–1.44); 0.982	1.14 (0.89–1.46); 0.305	0.558
<i>IL-10</i> : rs1800896 A/G	330/69/2	639/154/10	0.86 (0.63–1.18); 0.350	0.38 (0.08–1.76); 0.217	0.39 (0.09–1.81); 0.230	0.83 (0.61–1.13); 0.242	0.316

 a AA/AB/BB means homozygote, heterozygote and mutated homozygote; Bold values are statistically significant (p < 0.05); Adjusted for age and sex.

3. Experimental Section

3.1. Ethical Approval of the Study Protocol

The study protocol was approved by the Review Board of Peking Union Medical College (Beijing, China). We have complied with the World Medical Association Declaration of Helsinki regarding ethical conduct of research involving human subjects and/or animals. All patients provided written informed consent to be involved in the study.

3.2. Study Subjects

This study involved 1206 patients with CAD for the purpose undergoing coronary artery bypass grafting (CABG). Patients were consecutively recruited from the Cardiovascular Institute and Fuwai Hospital, Chinese Academy of Medical Sciences and Peking Union Medical College (Beijing, China) between December 2007 and December 2008. All patients were diagnosed using angiography which was scored systematically by an experienced interventional cardiologist. CAD patients were defined as having angiographic coronary stenosis of at least 50% lumen reduction. LMCAD was defined as at least 50% stenosis by visual assessment in the left main (LM) vessel, including ostial stenosis. While lesions compromising the lumen by ≥50% further from LM were defined as MPCAD [6,17].

All subjects were genetically unrelated ethnic Han Chinese. All data were collected by trained clinical research staff and were subsequently double entered into computer databases. Baseline information on personal and clinical characteristics was complete for all 1206 patients involved in the study.

3.3. Candidate Genes and SNPs Selection

Six candidate genes (*IL-1A*, *IL-1B*, *IL-6*, *IL-6R*, *IL-8* and *IL-10*) involved in the pathogenesis of inflammation and CAD were selected a priori based on previous transcription profiling in humans, pathway analysis, association studies reported in the literature, and expert opinion [18,19]. Seven typical SNPs, *IL-1A* rs1800587 C/T, *IL-1B* rs16944 G/A, *IL-6* rs1800796 C/G, *IL-6R* rs7529229 T/C, *IL-8* rs4073 T/A, *IL-10* rs1800872 A/C, and *IL-10* rs1800896 A/G, were subsequently selected in these candidate genes, based on function importance [20,21].

3.4. Isolation of DNA and Genotyping by Matrix-Assisted Laser Desorption/Ionization Time-of-Flight Mass Spectrometry (MALDI-TOF MS) and Ligation Detection Reaction (LDR)

Blood samples were collected using vacutainers and transferred to test tubes containing ethylenediamine tetra-acetic acid (EDTA). Genomic DNA was isolated from the lymphocytes of whole blood samples using the Wizard Genomic DNA Purification Kit (Promega, Madison, WI, USA). For quality control, all sample DNAs were conducted polymerase chain reaction (PCR), analyzed on a 3% agarose gel and visualized by ethidium bromide staining. *IL-1A* rs1800587 C/T, *IL-1B* rs16944 G/A, *IL-6* rs1800796 C/G, *IL-10* rs1800872 A/C and *IL-10* rs1800896 A/G genotyping was done by MALDI-TOF MS support from CapitalBio Corporation (Beijing, China),

using the MassARRAY system (Sequenom, San Diego, CA, USA) as previously described [22]. LMCAD and MPCAD at a proportion of ≈1:2 were assayed. Completed genotyping reactions were spotted onto a 384-well spectroCHIP (Sequenom) using a MassARRAY Nanodispenser (Sequenom), and analyzed by MALDI-TOF-MS (Sequenom, San Diego, CA, USA). Genotype calling was done in real time with MassARRAY RT software (version 3.1; Sequenom), and analyzed using MassARRAY Typer software (version 4.0; Sequenom). For *IL-6R* rs7529229 T/C and *IL-8* rs4073 T/A, genotyping study was performed using the LDR method [23,24], with technical support from Shanghai Biowing Applied Biotechnology Company (Shanghai, China).

3.5. Statistical Analyses

Differences in demographics, variables, and genotypes of the seven SNPs were evaluated using a chi-squared test. The associations between the seven SNPs and risk of CAD phenotype were estimated by computing odds ratios (ORs) and 95% confidence intervals (CIs) using logistic regression analyses, and by using crude ORs and adjusted ORs. Statistical differences with p < 0.05 were considered significant, and all statistical analyses were done with SAS software (version 9.1.3; SAS Institute, Cary, NC, USA).

4. Conclusions

In this study, we found that the *IL-6R* rs7529229 T/C polymorphism may increase the risk of LMCAD compared with MPCAD. The CC genotype is associated with higher risk of LMCAD phenotype. None of the other six polymorphisms showed any overall predisposition to LMCAD phenotype susceptibility. To the best of our knowledge, this study provides the first evidence linking variation in the *IL-6R* gene and LMCAD phenotype risk in a Chinese population.

This finding is supported by studies displaying a higher heritability of LMCAD. Capodanno *et al.* [25] investigated the epidemiology and the clinical impact of different anatomical phenotypes of the LM coronary artery, and Iwasaki *et al.* [26] investigated the distribution of coronary atherosclerosis in patients with CAD. Their findings suggested that LM phenotypes are more likely to present with atherosclerotic disease and significant stenosis and are particularly heritable [25,26].

Our results showed that *IL-6R* rs7529229 CC genotype as well as C allele was more prevalent in the LMCAD group than in the MPCAD control group. *IL-6R* rs7529229 T/C polymorphism was associated with increased risk of LMCAD, which may lead to LMCAD by enhancing inflammation. However, it was not found to be statistically significant in other six candidate gene variants.

IL-6 is an important pleiotropic cytokine that has a broad range of humoral and cellular immune properties relating to inflammation, tissue injury and contributes to the clinical evolution of CAD [27]. IL-6 exerts its biological activities through the IL-6R. The *IL-6R* gene is located on human chromosome 1q21, a region that previous studies have reported to be linked to metabolic syndrome, type 2 diabetes and atrial fibrillation [28–30]. The human *IL-6R* gene is highly polymorphic and there is considerable variation in its expression between individuals. A number of SNPs have been reported and genetic variants in *IL-6R* gene are associated with several different kinds of diseases, including CAD [31]. *IL-6R* rs7529229, in which the polymorphism is localized to a functional domain of

the receptor protein, is a T/C variation in the *IL-6R* gene (intronic) on human chromosome 1. In 40 studies including up to 133449 individuals, mendelian randomization analyses revealed that *IL-6R* rs7529229 T/C marking a non-synonymous *IL-6R* variant (rs8192284; p.Asp358Ala), was associated with increased circulating log IL-6 concentration as well as reduced C-reactive protein and fibrinogen concentrations [32]. This suggests that *IL-6R* rs7529229 variant is strongly associated with CAD and cardiovascular events.

In our study, Using Power and Sample Size Calculation (PS, version 3.0, 2009), considering IL-6R rs7529229 T/C mutant alleles in the LMCAD group, OR, LMCAD samples and MPCAD samples, the power of our analysis ($\alpha = 0.05$) was 0.857 in 396 LMCAD cases and 793 MPCAD cases with adjusted OR = 1.46. This suggests that SNP of IL-6R gene seems to also have a causal role in development of LMCAD. Thus, the findings indicate that IL-6R rs7529229 polymorphism may have potential importance in screening individuals at high risk for developing LMCAD, and targeting of IL-6R could provide a novel therapeutic approach for the prevention of LMCAD.

Several limitations of the present study need to be addressed. First, this was a hospital-based study; selection bias was unavoidable; Second, the polymorphisms we investigated, based on their functional considerations, may not offer a comprehensive view of the genetic variability. Further fine-mapping analysis of *IL* genes or high-density whole genome genetic analyses evaluating different CAD phenotypes might give further insights to the pathophysiologic mechanisms underlying LMCAD; Third, a single case—case study is not sufficient to fully interpret the relationship between *IL-6R* rs7529229 T/C polymorphism and susceptibility to LMCAD because of the relatively small number of patients with LMCAD. Replication studies with larger numbers of subjects are necessary to confirm our findings; Finally, we did not evaluate plasma IL-6 levels and the function of *IL-6R* rs7529229 T/C, which restricted our analyses.

Despite the limitations, the present study provided strong evidence that *IL-6R* rs7529229 T/C functional polymorphism may contribute to the risk of LMCAD. However, our results were obtained from a moderate-sized sample, and therefore this is a preliminary conclusion. Validation by a larger study from a more diverse ethnic population is needed to confirm these findings.

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Conflicts of Interest

The authors declare no conflict of interest.

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Investigation into Variation of Endogenous Metabolites in Bone Marrow Cells and Plasma in C3H/He Mice Exposed to Benzene

Rongli Sun, Juan Zhang, Lihong Yin and Yuepu Pu

Abstract: Benzene is identified as a carcinogen. Continued exposure of benzene may eventually lead to damage to the bone marrow, accompanied by pancytopenia, aplastic anemia or leukemia. This paper explores the variations of endogenous metabolites to provide possible clues for the molecular mechanism of benzene-induced hematotoxicity. Liquid chromatography coupled with time of flight-mass spectrometry (LC-TOF-MS) and principal component analysis (PCA) was applied to investigate the variation of endogenous metabolites in bone marrow cells and plasma of male C3H/He mice. The mice were injected subcutaneously with benzene (0, 300, 600 mg/day) once daily for seven days. The body weights, relative organ weights, blood parameters and bone marrow smears were also analyzed. The results indicated that benzene caused disturbances in the metabolism of oxidation of fatty acids and essential amino acids (lysine, phenylalanine and tyrosine) in bone marrow cells. Moreover, fatty acid oxidation was also disturbed in plasma and thus might be a common disturbed metabolic pathway induced by benzene in multiple organs. This study aims to investigate the underlying molecular mechanisms involved in benzene hematotoxicity, especially in bone marrow cells.

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1. Introduction

Benzene is an important industrial chemical widely used in the production of many products and is also a component of cigarette smoke, gasoline, crude oil and automobile emissions [1]. The hematopoietic system is the most critically affected target tissue following exposure to benzene in humans and animals. Benzene is an established cause of acute myeloid leukemia (AML), myelodysplastic syndromes (MDS), and very likely also lymphocytic leukemias and non-Hodgkin lymphoma (NHL) in humans [2–4]. Benzene was identified as an environmental carcinogen in 1982 [5] and placed in the Group 1 human carcinogen category in 1987 by the International Agency for Research on Cancer [6].

The major adverse health effect from exposure to benzene is hematotoxicity. Benzene can cause a decrease in the three major circulating cell types: platelets (thrombocytopenia), red blood cells (anemia) and white blood cells (leukopenia); and an increase in red cell mean corpuscular volume [7,8]. Sustained exposure may result in continued marrow depression involving multiple cell lineages. This multi-lineage depression of blood counts is also known as pancytopenia [9]. Continued exposure may eventually lead to damage to the bone marrow, accompanied by pancytopenia or aplastic anemia. Benzene metabolism is inherently complex [10], and its secondary metabolism

occurs in the bone marrow [11,12]. It is clear that bone marrow is the most critical target organ for benzene metabolites, and both progenitor cells and stromal cells in bone marrow have been considered to be potential targets of benzene hematotoxicity. There are a number of mechanistic studies in the literature that can help us to understand the primary mode of action for benzene, and significant progress has been made in this area. These studies describe effects of benzene such as chromosomal aberrations [13], covalent binding [14], and gene mutations [15]; as well as newly identified mechanisms that include alterations in gene expression [16], oxidative stress [17], epigenetic regulation [18], and immune suppression [19]. In addition, a number of biomonitoring studies have estimated internal benzene exposure for humans, and these have identified and quantified benzene or its biological adducts in blood, urine or expired air; urinary biomarkers include S-phenylmercapturic acid (SPMA), trans, trans-muconic acid (ttMA), phenol, catechol and hydroquinone. Although there is a wealth of epidemiologic data regarding benzene in humans and animals, exposure and toxicology data on benzene, and the mechanism of action for the hematotoxic effects of benzene are not completely understood; and assessment of internally based metabolites responsible for these effects are not currently available.

A systems biology approach to disease-related biology is revolutionizing our knowledge of the cellular pathways and gene networks that underlie the onset and progression of disease, and their associated pharmacological treatments [20]. The study of metabonomics depends upon the production of global metabolite profiles that enable diagnostic changes in the concentrations, or proportions, of low-molecular-weight organic metabolites in samples (such as biofluids and organ extracts) to be assayed. Such investigations thus generate metabolic phenotypes (metabotypes [21]), and, by studying these, it may be possible to identify target organ response to a specific toxicant [22], to assess the toxicity of candidate chemical agents, and to gain new insights into the mechanisms of toxicity of xenobiotics [23]. Kristin et al. carried out metabolomic analyses of stem cell samples from peripheral blood collected from a cohort of patients before hematopoietic cell transplantation, and the results suggested that the development of therapy-related myelodysplasia syndrome (t-MDS) was associated with dysfunctions in cellular metabolic pathways, including those involved in the metabolism of alanine, aspartate, glyoxylate, dicarboxylate, and phenylalanine; the citrate acid cycle; and aminoacyl-t-RNA biosynthesis [24]. A recent study suggested that the acridone derivative, 2-aminoacetamido-10-(3,5-dimethoxy)-benzyl-9(10H)-acridone hydrochloride, altered metabolism of fatty acids, nucleosides, amino acids, glycerophospholipid, and glutathione; it also induced oxidative stress-mediated apoptosis in CCRF-CEM leukemia cells [25]. Metabonomic approaches also enable identification of predictive markers and biomarkers of disease progression. Huang et al. used metabonomic profiling to identify a putative specific biomarker pattern in urine as a noninvasive bladder cancer (BC) detection strategy, and found carnitine C9:1 and component I (in a combined biomarker pattern) with a high sensitivity and specificity that allowed discrimination of bladder cancer patients [26]. One of the major analytical techniques used for global metabolic profiling at this time is mass spectrometry. Mass spectrometry (MS) occupies a major role in holistic metabolite profiling due to its sensitivity and widespread availability. Liquid chromatography (LC-MS) is currently the most widely used mass spectrometry technology, especially in the life science and bioanalytical sectors, due to its ability to separate and detect a wide range of molecules [27].

In our previous study [28], the results indicated that pathways of purine, spermidine, fatty acids, tryptophan, and peptide metabolism were disturbed in benzene-exposed mice; but this study was restricted to urine, which meant that relevant information regarding the hematopoietic system and interactions among compartments was lost. Compared with biofluids, bone marrow (as a target organ) more directly reflects the pathophysiologic state of disease processes induced by benzene. Endogenous metabolites in plasma reflects systemic metabolic effects associated with benzene exposure, while bone marrow cell metabolites analyses enable a more precise investigation of local metabolic changes. However, there are presently no studies reported regarding benzene-induced metabolic changes in bone marrow cells or plasma.

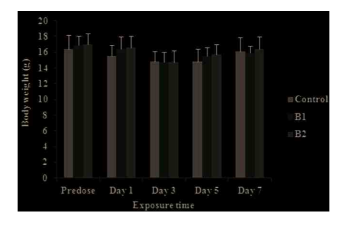
In the present study, we used an integrated approach that entailed metabonomic analyses (based upon HPLC-TOF-MS) of bone marrow cells and plasma to discern changes in the respective metabolomes and the interactions between the two compartments. Our aims were to study the variation of endogenous metabolites involved in benzene toxicity from metabonomic information derived from bone marrow, and to identify specific endogenous metabolites in plasma as potential biomarkers of benzene's toxic hematopoietic effects.

2. Results and Discussion

2.1. Body Weights and Relative Organ Weights

The mice in benzene 2 group (receiving 600 mg/kg b.w.) manifested some irritability and lethargy after benzene exposure for seven consecutive days. There were no significant differences in the body weights of mice at any of the time intervals analyzed (Figure 1) (p > 0.05), suggesting that the toxicity induced by benzene was insufficient to cause observable body weight changes. The relative organ (liver, spleen, lung, and kidney) weights of mice are presented in Table 1. There was a significant decrease in relative lung weights in benzene 2 group mice on the seventh day of benzene exposure. In addition, there was a significant decrease in relative spleen weights in both benzene 1 and 2 groups.

Figure 1. Body weight of mice at each time point and dose. Each bar represents means \pm standard deviation (SD) from one-way ANOVA.



Group	Relative liver weight	Relative spleen weight	Relative lung weight	Relative kidney weight
Control	6.01 ± 0.32	0.36 ± 0.05	0.69 ± 0.10	1.63 ± 0.14
Benzene 1	6.38 ± 0.31	0.27 ± 0.09 *	0.67 ± 0.08	1.65 ± 0.15
Benzene 2	6.54 ± 0.48	$0.25 \pm 0.07 *$	0.56 + 0.02 *	1.69 ± 0.06

Table 1. Relative organ weights of male C3H/He mice on benzene exposure day 7.

2.2. Blood Parameters and Bone Marrow Smear

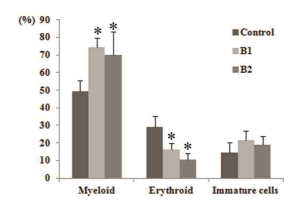
The parameters of peripheral blood and bone marrow smears were investigated to assess the hematotoxicity of benzene. As shown in Table 2, a significant decrease in RBC number and hemoglobin (Hgb) concentration occurred in both groups of mice following exposure to benzene for seven days. No effect on WBC number was found, which may be due to the relatively short time of exposure. The platelet (pit) was reduced in the benzene-exposed groups, but not to a statistically significant extent. The bone marrow smears were made to observe the extent of nucleated cell proliferation and cell morphology (Figure 2). The results showed significant myeloid hyperplasia and a marked reduction of erythroidin benzene groups, while no significant difference was observed in the ratio of immature cells in benzene exposure groups. These observations suggest that benzene exposure leads to hematotoxicity.

Table 2. Blood parameters in male C3H/He mice on benzene exposure day 7.

Group	WBC $(10^9/L)$	$RBC (10^{12}/L)$	Hgb (g/L)	Pit (10 ⁹ /L)
Control	4.05 ± 0.65	7.98 ± 0.39	137.20 ± 5.76	364.25 ± 60.50
Benzene 1	3.87 ± 1.06	7.30 ± 0.14 *	126.67 ± 3.50 *	322.00 ± 107.53
Benzene 2	4.48 ± 0.96	7.32 ± 0.42 *	127.60 ± 7.50 *	282.2 ± 50.01

^{*} significant difference compared with control group (p < 0.05).

Figure 2. Bone marrow smear examination in male C3H/He mice following 7 days of benzene exposure. * significant difference compared with control group (p < 0.05).



^{*} significant difference compared with control group (p < 0.05).

2.3. LC-MS Fingerprinting of Mouse Bone Marrow Cells and Plasma

Typical HPLC-MS total ion current (TIC) chromatograms of mouse bone marrow cell and plasma samples on day seven taken from the control and benzene-exposed groups are shown in Figures 3 and 4. As shown in Figure 3, there were significant visual differences in the TIC among the groups, especially from 1 to 3 min. The difference between the control group and benzene-exposed groups was more apparent than that between the two dosed groups. A similar metabonomic profile difference was also observed in the TIC of plasma (Figure 4).

Figure 3. Total ion chromatograms (TICS) of bone marrow cell samples obtained from the control group (**C**), and benzene 1 (**B1**) and benzene 2 groups (**B2**) of male C3H/He mice following 7 days of benzene exposure, using LC/MS (positive mode).

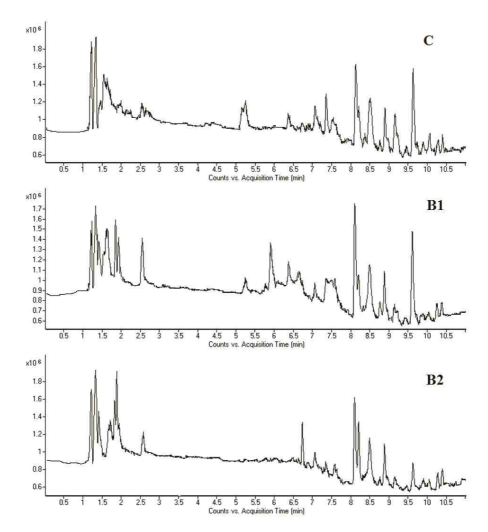
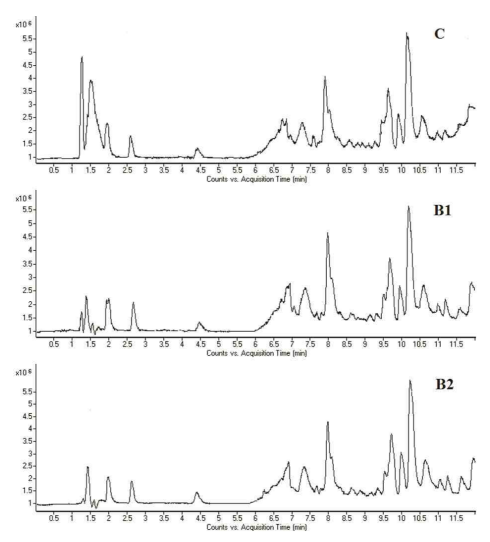


Figure 4. Total ion chromatograms (TICS) of plasma samples obtained from the control group (**C**), and benzene 1 (**B1**) and benzene 2 groups (**B2**) of male C3H/He mice following 7 days of benzene exposure, using LC/MS (positive mode).

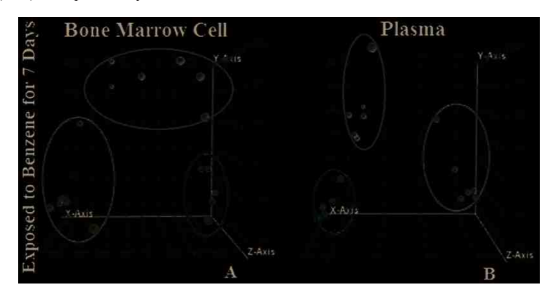


2.4. Principal Component Analysis and Discovery of Changed Endogenous Metabolites

We determined metabolites that were responsible for the changes illustrated above by using one-way ANOVA (p < 0.05, fold-change ≥ 2). Principal Component Analysis (PCA) was used to further select biomarkers that could discriminate between groups. PCA (an unsupervised method), is quite useful in distinguishing the several thousand biochemical endpoints retrieved from each sample. In the PCA score plots, each spot represents a metabonomic sample and each assembly of samples expresses a unique metabolic pattern at different time points. This analysis was successful in this experiment because more than 80% of the variability was explained using four components. PCA score plots derived from levels of 16 metabolites from bone marrow cells showed marked differences between the control and benzene-exposed mice on day seven. Similar results can be found in the PCA score map derived from plasma metabolites (Figure 5). The results showed that benzene exposure induced significant changes in 16 metabolites in bone marrow cells and 25 metabolites in

plasma, with some metabolites changed in more than one compartment. These metabolites were considered to be potential biomarkers of benzene action.

Figure 5. 3D PCA score plot of the metabolic profiles of bone marrow cells (**A**) and plasma (**B**) in the control group (red), benzene 1 group (brown), and benzene 2 group (blue) on exposure day 7.



2.5. Identification of Changed Endogenous Metabolites

Herein, we took the ion at m/z 166 ([M+H]⁺) as an example to illustrate the identification process. First, the corresponding quasi-molecular ion peak was found according to the retention time in the extracted ion chromatogram (EIC) of m/z 166 (Figure 6A). The accurate mass of the quasi-molecular ion was found as m/z 166.0863, and C₉H₁₁NO₂ was calculated as the most probable molecular formula using Agilent MassHunter software. Then, we conducted its fragmentation by tandem MS. Three major fragment ions were found at m/z 103.0545 and 120.0806, which represent the fragments of [C₈H₇]⁺ and [C₈H₁₀N]⁺, respectively. With the aforementioned information, we searched the freely accessible databases HMDB (http://www.hmdb.ca), METLIN (http://metlin.scripps.edu) and KEGG (http://www.kegg.jp). Finally, considering elemental composition, fragmentation patterns and chromatographic retention behavior, the m/z of 166 was identified as L-phenylalanine, which was then validated using a standard (Figure 6B,C). Likewise, other biomarkers have been identified and are listed in Table 3. However, the remaining biomarkers (data not shown) were unidentifiable due to insufficient intensity for the MS/MS experiments, or due to restrictions in the current metabolite databases.

Figure 6. Identification of a selected marker (m/z 166). (**A**) Extracted ion chromatogram (EIC) of m/z 166; (**B**) MS/MS spectrum of the ion; (**C**) MS/MS spectrum of a commercial standard L-phenylalanine. The collision energy was 20 V.

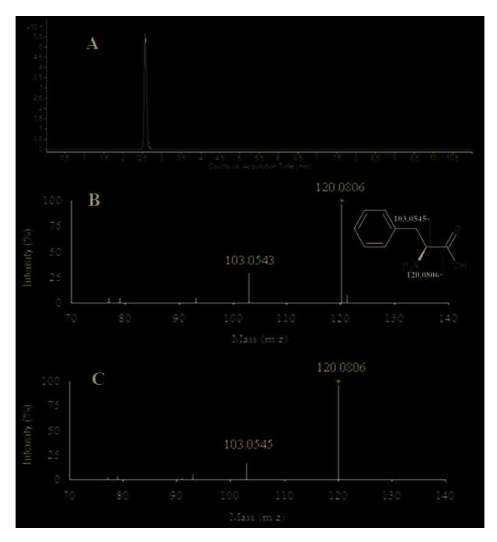


Table 3. Identified endogenous biomarkers in bone marrow cell and plasma on exposure day 7.

Compartmen t	m/z	RT (Retention time)	Trend	<i>p</i> -value	Metabolites	Related pathway
	204.123	1.54	\downarrow	9.30×10^{-4}	L-Acetylcarnitine	Oxidation of Fatty Acids
	165.054 6	1.681	1	5.14×10^{-3}	p-Coumaric acid	Unknown
Bone Marrow Cell	182.081	1.668	↑	3.97×10^{-4}	L-tyrosine	Tyrosine/Phenylalanine and Tyrosine Metabolism, Catecholamine Biosynthesis
	166.086 3	2.548	↑	1.62×10^{-3}	L-Phenylalanine	Phenylalanine and Tyrosine Metabolism
	147.116 8	9.205	\downarrow	1.65×10^{-4}	Lysine	Lysine Degradation, Biotin Metabolism, Carnitine Synthesis
	192.064 4	1.2775	\downarrow	1.92×10^{-6}	5- Hydroxyindoleacetic acid	Tryptophan metabolism
	112.086 9	1.3	\downarrow	2.06×10^{-4}	Histamine	Histidine metabolism
Dlagma	156.076 5	1.321	↑	1.08×10^{-5}	L-Histidine	Histidine Metabolism, Ammonia Recycling Transcription/Translation
Plasma	126.102 6	1.322	\downarrow	1.32×10^{-4}	N-Methylhistamine	Histidine metabolism
	204.123	1.58	\downarrow	6.01×10^{-5}	L-Acetylcarnitine	Oxidation of Fatty Acids
	130.049	2.177	1	2.88×10^{-6}	Pyrrolidonecarboxylic acid	Gamma-glutamyl cycle
	400.342 1	9.195	\downarrow	3.11×10^{-7}	Palmitoylcarnitine	Fatty acid Metabolism

^a Change trend of benzene exposure mice vs. control mice. Variations compared to control samples: \uparrow , indicates relative increase in signal; \downarrow , relative decrease in signal (p < 0.05).

2.6. Biological Significance of Endogenous Metabolites Alternations in Bone Marrow Cells and Plasma

Metabolic profiling contributes diagnostic information and presents mechanistic insights into the biochemical effects of toxins [29]. The variation of endogenous metabolites in bone marrow may be indicative of benzene's toxic hematopoietic mechanisms, and the fluctuation may then be observed in plasma. Since blood is circulated around the body, the endogenous metabolites alternations in multi-organs induced by benzene can also be reflected in the blood. This may be helpful in identifying potential biomarkers of toxic effects that would relate to adverse health effects, and that could be monitored in blood. Therefore, in this study, we analyzed benzene-induced endogenous metabolites changes in bone marrow cells and plasma.

2.6.1. Significance of Changed Endogenous Metabolites in Bone Marrow Cells Induced by Benzene

Benzene is hematotoxic and leukemogenic in humans and induces bone marrow suppression in rodents. In recent years, research into the metabolic pathways involved in the renewal and differentiation of HSC and hematopoietic system diseases has intensified. In this study, we uncovered and identified five potential biomarkers in bone marrow cells, and expect to use these to further investigate the hematotoxic mechanisms of benzene. Our results showed in bone marrow cells increases in the levels of phenylalanine and tyrosine, and a decrease in L-lysine, three of the essential amino acids. In addition, a significant lowering of acetylcarnitine was found in bone marrow cells as compared to controls.

A recent study found an increased level of phenylalanine in stem cells from peripheral blood of t-MDS/AML patients, suggesting that there may exist an alteration in mitochondrial activity in these patients relative to controls [24]. In our study, the significant increase in phenylalanine levels in bone marrow cells might then be a manifestation of mitochondrial dysfunction. Phenylalanine is needed for the synthesis of protein, melanin and tyrosine. The increased bone marrow cell levels of phenylalanine may enhance the metabolic pathway from phenylalanine to tyrosine, resulting in a high concentration of bone marrow cell tyrosine. Higher serum levels of phenylalanine and tyrosine were observed in AML patients, which might be the result of enhanced degradation of proteins from the host experiencing a cancerous condition, and these two amino acids are needed for gluconeogenesis and for catabolism to provide intermediates for the tricarboxylic acid cycle (TCA) cycle [30]. It is possible that the toxicity to bone marrow with benzene exposure may be involved in the TCA cycle in mitochondria. The signaling pathway associated with increased phenylalanine and tyrosine may be one of the vital mechanisms intrinsic to benzene-induced hematotoxicity.

L-lysine is an essential amino acid and exerts antifibrinolytic activity by inhibition of fibrinolysis, and exerts a protective effect on platelets [31]. Lysine acetyltransferases were reported to play a key role in leukemogenesis and interact with Runx1 (or AML1), one of the most frequent targets of chromosomal translocations in leukemia [32]. Carnitine is synthesized from lysine residues in existing proteins, and then used to further synthesize acetylcarnitine via carnitine palmitoyl transferase I. Acetylcarnitine is an acetic acid ester of carnitine that facilitates movement of acetyl CoA into the matrices of mammalian mitochondria during the oxidation of fatty acids [33], and it has been observed that lower levels of acetylcarnitine are found in the blood of AML patients [30]. Ito et al. showed that fatty acid oxidation was associated with hematopoietic stem cell proliferation and differentiation, which determines whether they undergo symmetric or asymmetric cell division [34]. Therefore, decreases in bone marrow cell lysine and acetylcarnitine levels are likely associated with a down-regulation of carnitine synthesis, which then disturbs oxidation of fatty acids in hematopoietic stem cells that are exposed to benzene. In addition, significant changes in the levels of these compounds in bone marrow were traced in plasma, and we found specific metabolites that related to hematopoietic toxicity of benzene. It is worth noting that only acetylcarnitine levels in plasma decreased, consistent with the changes in bone marrow, indicating that lower levels of acetylcarnitine in plasma might be indicative of hematotoxic effects of benzene.

2.6.2. Significance of Changed Endogenous Metabolites in Plasma Induced by Benzene

The levels of five significantly changed metabolites in bone marrow were also traceable in plasma. Only a reduction in acetylcarnitine was found in plasma. In addition, benzene exposure caused an elevation of L-histidine and pyrrolidone carboxylic acid, concomitant with decreases in 5-hydroxyindoleacetic acid, histamine, *N*-methylhistamine, and palmitoyl-L-carnitine in plasma (Table 3). These significantly changed endogenous metabolites could be used to illustrate the multiple-organ toxicity induced by benzene.

Benzene was demonstrated to cause a disturbance in histidine-related metabolism, including a significant decline in histamine and *N*-methylhistamine, concomitant with an elevation in histidine. The high plasma histidine in the benzene-exposed groups may be caused by a diminished activity of histidine decarboxylase (HDC) [35], which results in inhibiting decarboxylation of histidine to histamine in benzene-exposed mice. Histamine is an amine derived by enzymatic decarboxylation of histidine, which plays a pivotal role in a number of processes, including inflammation, allergic reactions, gastric acid secretion and neurotransmission [36]. Histamine was reported to inhibit production of reactive oxygen species (ROS) in CML cells via the H₂-receptor (H₂R) [37]. A salt of histamine, histamine dihydrochloride (HDC), is used as a drug for the prevention of relapse in patients diagnosed with AML [38,39]. Recently, Aurelius *et al.* found that HDC acted on H₂R expressed by leukemia cells to reduce ROS formation, which might impact the effectiveness of histamine-based immunotherapy [40]. Phenolic metabolites of benzene accumulate in the bone marrow where myeloperoxidase and other peroxidases convert them to reactive semiquinones and quinines [41], which can further lead to the formation of ROS [42]. Histamine might thereby be consumed in order to respond to ROS generation induced by benzene.

Palmitoylcarnitine is a long-chain acyl fatty acid derivative ester of carnitine that facilitates the transfer of long-chain fatty acids from the cytoplasm into mitochondria during the oxidation of fatty acids. Palmitoylcarnitine was shown to stimulate the activity of caspases 3, 7 and 8, and the level of this long-chain acylcarnitine increased during apoptosis [43]. Ibuki *et al.* reported that benzene metabolites induced an anti-apoptotic effect, and that the effect was mainly due to the production of ROS by benzene metabolites (*p*-benzoquinone and hydroquinone) that inhibited caspase-3 activation. Inhibition of apoptosis, aberrantly prolonging cell survival, may contribute to cancer by facilitating the creation of mutations and by allowing a permissive environment for genetic instability [44]. Thus, the decreased plasma palmitoylcarnitine levels in the benzene-exposed groups may be related to disturbed fatty acid metabolism and the suppression of apoptosis by inhibiting caspase activation.

5-Hydroxyindoleacetic acid (5-HIAA) is a breakdown product of serotonin and levels of these substances may be measured in plasma to monitor progression of diseases such as carcinoid tumors [45] and pulmonary hypertension [46]. In the present study, plasma 5-HIAA was found to be notably reduced in benzene-exposed mice, and this may due to benzene toxicity.

Pyrrolidonecarboxylic acid can be irreversibly converted to glutamate, which is used to generate glutamine. Peng *et al.* [47] investigated amino acid concentrations during induction and preconsolidation therapy in cerebrospinal fluid (CSF) of children with lymphoblastic leukemia

(ALL) with or without CNS involvement. However, they did not find any significantly changed amino acid levels except for higher baseline glutamine levels, indicative of a greater risk for CNS leukemia. Although increased levels of glutamine were not detected in the plasma of mice exposed to benzene, the increased pyrrolidonecarboxylic acid might indirectly reflect an early influence of benzene on glutamate metabolism.

3. Experimental Section

3.1. Chemicals and Reagents

Benzene was purchased from Sigma Co. (St. Louis, MO, USA); and corn oil from COFCO (Beijing, China). Ultrapure water (18.2 MO) was prepared with a Milli-Q water purification system (Millipore, Bedford, MA, USA). LC/MS grade methanol and acetonitrile were purchased from Spain Scharlau, Ltd. (Barcelona, Spain); and analytical grade formic acid was supplied by Dikma Corp. (Richmond, NY, USA).

3.2. Ethics Statement

This study was carried out in strict accordance with the recommendations of the Guide for the Care and Use of Laboratory Animals of the State Committee of Science and Technology of the People's Republic of China. The protocol of experiments was reviewed and approved by the Research Ethics Committee of the Southeast University (approval number: 20130027). Animals were maintained and experiments were conducted in accordance with the Institutional Animal Care and Use Committee of Southeast University.

3.3. Animals and Treatments

Eighteen male C3H/He mice (aged 4 weeks, and weighing 17.11 ± 1.03 g) obtained from Wei Tong Li Hua Laboratory Animal Co. Ltd. (Beijing, China) were acclimatized for one week in the Specific-Pathogen Free (SPF) animal facility prior to administration of substances. Animals were maintained under a 12-h light/12-h dark cycle at a temperature of 25 ± 2 °C with a relative humidity of 45%–65%. Animals had *ad libitum* access to a certified standard diet and to drinking water, and were divided randomly into a control group (vehicle, oil; n = 6), benzene group 1 (B1: 300 mg/kg b.w.; n = 6) and benzene group 2 (B2: 600 mg/kg b.w.; n = 6). Mice were injected subcutaneously with either corn oil or a benzene-corn oil mixture once daily for seven consecutive days. The aim of these doses is to examine the corresponding hematotoxicity of benzene. The route of benzene administration (injection, s.c.) was to allow us better control of benzene dosages [48,49].

The body weight of each mouse was recorded every other day during exposure periods. The mice were sacrificed on the seventh day of exposure. The mice were anesthetized with pelltobarbitalum natricum, blood was collected, and then were sacrificed by decapitation. Liver, spleen, lung, and kidneys were excised and weighed. Relative organ weight was calculated as the ratio between organ weight and body weight. Bone marrow cells were flushed from one tibia using a 26-gauge needle to make smears.

3.4. Collection of Plasma and Bone Marrow Cell Samples

The plasma and bone marrow cell samples of mice were collected after being exposed to benzene for seven days. Plasma was extracted from whole blood at 3000 rpm for 10 min at room temperature. After acquiring mouse femurs and tibias, the marrow cavities were washed with a 26-gauge needle (on ice). Mouse bone marrow cells were collected after centrifugation at $300 \times g$ for 10 min at 4 °C. To 5×10^6 bone marrow cells, 1 mL of quenching solution (iced, 0.9% [w/v] NaCl) was quickly added. All cell samples were centrifuged at $1000 \times g$ for 1 min. Cell pellets were resuspended in ice-cold 50% aqueous acetonitrile, vortexed, and incubated on ice for 10 min. The extracts were dried in a SpeedVac, and the dried cell extracts were resuspended in 500 μ L water. All obtained samples were frozen immediately and stored at -80 °C until analysis.

3.5. Sample Preparation and HPLC/MS Analysis

Plasma samples were thawed at room temperature and then centrifuged at $13,000 \times g$ for 15 min at 4 °C. Each 100 µL aliquot of plasma was mixed with 300 µL of methanol and vortexed to allow for protein precipitation. After centrifugation at 13,000× g for 15 min at 4 °C, the combined supernatants were transferred to the auto-sample vials. The bone marrow cells were thawed and could be analyzed directly. For the metabonomic studies, aliquots of 1 µL of each sample were injected into a ZORBAX Eclipse Plus C18 column (3.00 mm × 100 mm × 1.8 μm, Agilent, Santa Clara, CA, USA) using an Agilent 6224 TOF LC-MS system (Agilent). The mobile phase was 0.1% formic acid in water (A) and 0.1% formic acid in acetonitrile (B). The optimized HPLC elution conditions were: (a) plasma: 0–1 min, 5% B; 1–3.5 min, 5%–80% B; 3.5–10 min, 80%–95% B; 10–12 min, 95% B; 12-12.5 min, 5% B; (b) bone marrow cell: 0-0.3 min, 5% B; 0.3-7 min, 5%-95% B; 7-9 min, 95% B; 9-10 min, 5% B. The flow rate was 0.4 mL/min. The column and autosample were maintained at 35 and 4 °C, respectively. The positive ion mode was used for the mass detection. The source parameters were set as follows: drying gas flow rate, 9 L/min; gas temperature, 350 °C; pressure of nebulizer gas, 40 psig; Vcap, 4000 V; fragmentor, 150 V; skimmer, 60 V; and scan range, m/z 50–1000. The tune mixture solution (Agilent) was employed as the lock mass (m/z = 121.050873, 922.009798) at a flow rate of 30 µL/min, via a lock spray interface for accurate mass measurement. To confirm the identity of the metabolites obtained after the non-targeted analysis (MS analysis and database search), a LC/6530 Q-TOF-MS (Agilent) was used. The MS/MS analysis was acquired in targeted MS/MS mode with collision energy from 10, 20 and 40 V; and a scan rate of 1 (MS/MS) scans/s.

3.6. Data Processing

The Masshunter Data Analysis Software (Ver B.02.01, Agilent Technologies, Barcelona, Spain) was used to analyze results; and the Masshunter Qualitative Analysis Software (Agilent Technologies) was used to obtain the molecular features of the samples, representing different, co-migrating ionic species of a given molecular entity using the Molecular Feature Extractor (MFE) algorithm. Finally, the Masshunter Mass Profiler Professional Software (Ver B.02.02, Agilent Technologies) was used to perform a non-targeted metabolomic analysis of the extracted features.

Samples with a minimal absolute abundance of 2000 counts and with a minimum of 2 ions were selected. Multiple charge states were not considered. Compounds from different samples were aligned using a RT window of $0.2\% \pm 0.15$ min and a mass window of 10 ppm \pm 2.0 mDa. Only common features (found in at least 75% of the samples of the same condition) were analyzed, correcting for individual bias. Data for PCA analysis were obtained using this software. As a classic unsupervised method (no prior knowledge concerning groups or tendencies within the data sets was necessary) of pattern recognition, PCA was expected to discern through statistical protocols several distinct variables for use as potential biomarkers.

3.7. Statistics

Statistical analyses for non-targeted metabonomics analyses were performed using the Mass Profiler Professional Software (Agilent Technologies). The exact masses with significant differences in abundance were determined using a one-way analysis of variance (ANOVA); fold-change >2 was considered to be significant at p < 0.05, and was searched against various databases (METLIN, HMDB, LIPID MAPS and KEGG). Otherwise, statistical calculations were performed using the SPSS 15.0 software (SPSS, Chicago, IL, USA). Multiple comparisons were analyzed using one-way ANOVA. Statistical significance was established at a level of p < 0.05.

4. Conclusions

In conclusion, we applied an LC-MS-based metabonomics approach to investigate benzene-induced toxicity in male C3H/He mice. The combined experimental results of metabonomics, relative organ weights, blood parameters and bone marrow smears indicated benzene-induced hematotoxicity. The obvious metabolic alterations in mouse bone marrow cells and plasma indicated that benzene exposure disrupted metabolism of essential amino acids (lysine, phenylalanine and tyrosine) in bone marrow cells; resulting in benzene-induced hematotoxicity. Benzene also caused disturbance in the metabolism of fatty acids oxidation. The decreased acetylcarnitine in plasma was commensurate with that in bone marrow cells; suggesting that acetylcarnitine in plasma is very likely an appropriate biomarker of benzene hematotoxicity. Our work offers a new clue for further clarification of the mechanism(s) involved in benzene-induced toxicity via studying variation of endogenous metabolites.

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Conflicts of Interest

The authors declare no conflict of interest.

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Phosphosite Mapping of HIP-55 Protein in Mammalian Cells

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Abstract: In the present study, hematopoietic progenitor kinase 1 (HPK1)-interacting protein of 55 kDa (*HIP-55*) protein was over-expressed in HEK293 cells, which was genetically attached with 6x His tag. The protein was purified by nickel-charged resin and was then subjected to tryptic digestion. The phosphorylated peptides within the *HIP-55* protein were enriched by TiO₂ affinity chromatography, followed by mass spectrometry analysis. Fourteen phosphorylation sites along the primary structure of *HIP-55* protein were identified, most of which had not been previously reported. Our results indicate that bio-mass spectrometry coupled with manual interpretation can be used to successfully identify the phosphorylation modification in *HIP-55* protein in HEK293 cells.

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1. Introduction

Protein phosphorylation is a fundamental type of post-translational modification, which plays a significant role in a wide range of cellular processes. Reversible phosphorylation results in a conformational change in the structure of many enzymes, receptors and adaptor proteins, triggering cellular signaling transduction, and modulating protein function, stability, interaction and localization. [1,2] Phosphorylation usually occurs on serine, threonine and tyrosine in eukaryotic proteins. [3] Phosphorylation on serine is the most common, followed by threonine and tyrosine. Determining protein phosphorylation sites is often the first step in the elucidation of a biological mechanism. Within a protein, phosphorylation can occur on several amino acids. The different phosphorylation sites mediate different biological processes. Therefore, the identification of *in vivo* phosphorylated sites of proteins is extremely important for understanding biological function and processes.

HIP-55, also called SH3P7, mAbp1 and DBNL, is a multi-domain adaptor protein, with an actin-binding domain at its N-terminus and an SH3 domain at its C-terminus [4]. HIP-55 acts as adaptor protein in many cellular processes such as cell signaling transduction and receptor endocytosis [5,6]. HIP-55 also plays important roles in T-cell proliferation, immune responses and the development of cerebellar architecture [7–10]. Some important functions of the HIP-55 protein are reported as mediated and regulated by its phosphorylation. HIP-55 was identified as a tyrosine kinase substrate using anti-phosphotyrosine antibodies [11]. HIP-55 is phosphorylated by Syk, Lyn, and Blk and further links antigen receptor signaling to components of the cytoskeleton [12]. HIP-55 is also identified as a novel MELK substrate and is important for stem-cell characteristics and invasiveness [13]. Furthermore, src-mediated phosphorylation of HIP-55 regulates podosome rosette formation in transformed fibroblasts [14]. Clearly, the HIP-55 protein is involved in many signal transduction processes; thus, investigation of its phosphorylation modifications is important to better understand the role of HIP-55 protein in the precise regulation of signal transduction.

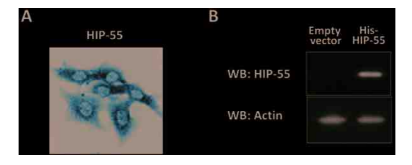
Traditionally, three approaches are used to determine phosphorylation sites: the bioinformatics approach, the biochemical approach, and the genetics approach. In the last decade, advancements in mass spectrometry have redefined conventional biochemical approaches for the identification of phosphorylation sites. Mass spectrometry (MS) has become currently the most powerful technique for analysis of phosphorylation sites [15,16]. In the present study, mass spectrometry combined with phosphopeptide enrichment techniques were employed to identify the phosphorylation sites of *HIP-55* protein in mammalian cell. *HIP-55* protein carrying a tag of six histidines was over-expressed in HEK293 cells, which was enriched by Ni-NTA resin. Reverse capillary high efficiency liquid chromatography (HPLC) coupled with mass spectrometry was employed to profile phosphorylation modifications in the purified *HIP-55* protein. Several novel sites of phosphorylation in *HIP-55* protein were identified.

2. Results and Discussion

2.1. Expression and Purification of His-Tagged HIP-55 Protein

HIP-55, as an adaptor protein, was found to be expressed in various mammalian cells. The distribution of HIP-55 was examined with an immunofluorescence assay. The result showed that HIP-55 was observed throughout the cytosol and appeared enriched in the perinuclear area (Figure 1A). pDEST-His-HIP-55 plasmid with full length HIP-55 gene attached to a six-histidine tag were transfected into HEK293 cells which were maintained in DMEM containing 10% fetal bovine serum for 36 h at 37 °C. After the medium was removed, cells were washed with PBS three times. Then the cells were harvested and lyzed. HIP-55 protein was purified with Ni-NTA resin and further precipitated with chilled acetone to remove high amounts of salts such as guanidine HCl and imidazole. The protein pellet was collected and dissolved in SDS-containing buffer. The purified HIP-55 protein was checked by western blot analysis as indicated in Figure 1B.

Figure 1. Expression and purification of *HIP-55* protein. (**A**) Cells were fixed and labeled with antibodies to *HIP-55*. Subcellular location of *HIP-55* was shown by laser scanning confocal microscopy; (**B**) After His-pull-down, the overexpression of *HIP-55* in the HEK293 cell line was detected with Western blot assay by *HIP-55* antibody. Actin was used as a loading control.



2.2. Identification of HIP-55 Protein by MS

The eluted proteins from Ni-NTA resin were desalted by acetone precipitation and then subjected to tryptic digestion. Prior to the analysis of phosphorylation, the peptide mixture was directly subjected to LC-MS/MS analysis, and the obtained data were inputted into a protein database for searching. Results showed a series of tryptic peptides from *HIP-55* protein with a coverage rate of 45.1%, in which only one phosphopeptide (LRS*PFLQK) was identified when Ser, Thr and Tyr phosphorylations were considered during the database searching. Figure 2 shows the distribution of peptides in the identification of *HIP-55* protein.

Figure 2. Primary structure of *HIP-55* protein. The tryptic peptides identified by LC-MS/MS analysis are highlighted.

```
MAANLSRNGP ALQEAYVRVV TEKSPTDWAL FTYEGNSNDI RVAGTGEGGL EEMVEELNSG

KVMYAFCRVK DPNSGLPKFV LINWTGEGVN DVRKGACASH VSTMASFLKG AHVTINAR AE

EDVEPECIME KVAKASGANY SFHKESGRFQ DVGPQAPVGS VYQKTNAVSE IKRVGKDSFW

AKAEKEEENR RLEEKRRAEE AQRQLEQERR ERELREAARR EQRYQEQGGE ASPQRTWEQQ

QEVVSRNRNE QESAVHPREI FKQKERAMST TSISSPQPGK LRSPFLQKQL TQPETHFGRE

PAAAISRPRA DLPAEEPAPS TPPCLVQAEE EAVYEEPPEQ ETFYEQPPLV QQQGAGSEHI

DHHIQGQGLS GQGLCARALY DYQAADDTEI SFDPENLITG IEVIDEGWWR GYGPDGHFGM

FPANYVELIE
```

2.3. MS Analysis of Phosphorylation Sites of HIP-55 Protein

By enrichment of phosphopeptides from tryptic peptides mixture with TiO₂, a total of fourteen phosphopeptides from *HIP-55* protein were identified by LC-MS/MS analysis and database searching, including the one previously detected without TiO₂ enrichment (Table 1).

A few spectra were given below as examples for the identification of some of these phosphorylation sites in *HIP-55* protein.

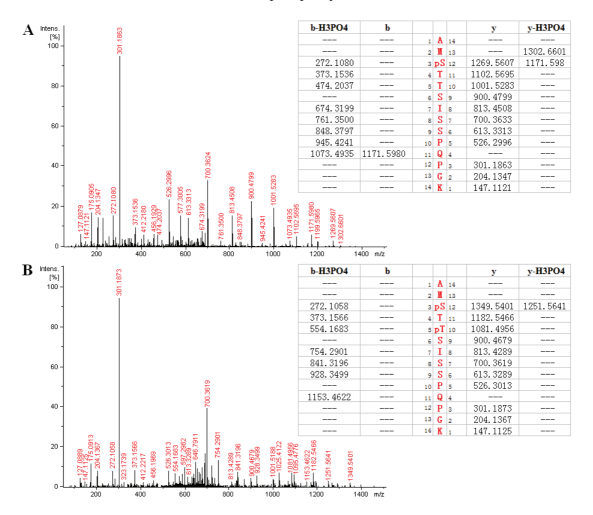
Table 1. Identification of phosphopeptides within *HIP-55* protein over-expressed in HEK293 cells.

Phosphopeptides	Experimental m/z (mono)	Theoretical m/z (mono)	Charge	XCorr	Position in protein
GACAS *HVSTMASFLK	795.353	795.346	2	1.98	95–109
GACAS *HVSTM *ASFLK	803.351	803.344	2	1.94	95-109
GAHVT *INAR	509.744	509.745	2	2.37	110-118
ASGANY *SFHKES *GR	835.828	835.822	2	2.21	135–148
TNAVS *EIK	471.224	471.221	2	3.06	165–172
VGKDS *FWAK	559.253	559.258	2	2.57	174–182
REQRY *QEQGGEAS *PQR	1039.924	1039.926	2	1.94	220-235
AMS *TTSISSPQPGK	736.333	736.329	2	3.63	267–280
AMS *TT *SISSPQPGK	776.318	776.312	2	2.56	267–280
AM *S *TTSISSPQPGK	744.324	744.326	2	3.23	267–280
LRS *PFLQK	534.785	534.784	2	2.41	281-288
QLT *QPETHFGR	697.321	697.317	2	2.01	289-299
QLT *QPET *HFGR	737.304	737.300	2	1.98	289-299
EPAAAIS *RPR	574.289	574.285	2	2.49	300–309

NOTE: T *, S * and Y * refer to the phosphorylated form of threonine (Thr, T), serine (Ser, S) and tyrosine (Tyr, Y), respectively; M * refers to the oxidized methionine (Met, M).

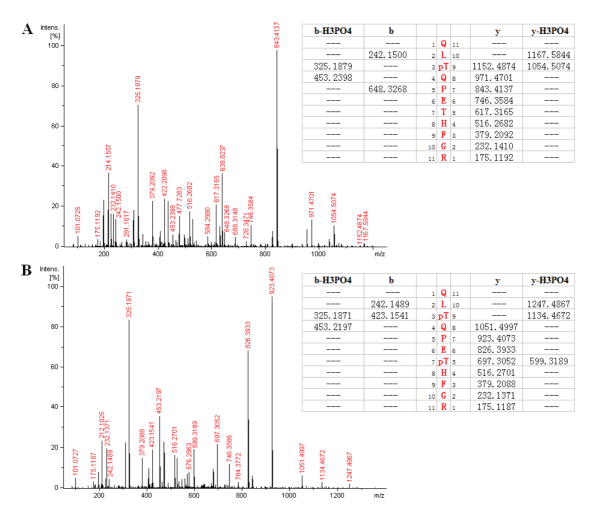
Figure 3A showed the MS/MS spectrum of a doubly-charged molecular ion peak with m/z at 736.333. Database searching identified a phosphopeptide as AMpSTTSISSPQPGK (267–280) from HIP-55 protein. A series of fragment ions such as fragment ions of b and y series can be clearly identified. The detection of y series ions at m/z 1102.5695, 1269.5607, 1171.5980 and 1302.6601 indicated that phosphorylation occurred exclusively at Serine 269 in this peptide. Detection of b series ions also confirmed this identification. Figure 3B shows the spectrum of a doubly-charged molecular ion peak with m/z at 776.318, which is similar to Figure 3A. The corresponding peptide is identified as the same peptide sequence as in Figure 3A, but with an additional phosphorylation site at Threonine 271. As shown in Figure 3B, y series ions at m/z 900.4679, 1081.4956, 1182.5466 and 1349.5401 indicated that phosphorylation occurred exclusively at Serine 269 in this peptide. Detection of b series ions also confirmed this identification. As shown in both mass spectra, the base peak at m/z 301.19 was observed, which resulted from fragmentation at Proline 278. It was noticed that most of the b series ions were subject to neutral loss of 98 (a phosphate molecule).

Figure 3. (**A**) MS/MS spectrum of a doubly-charged peak at m/z 736.333. The corresponding peptide is identified as AMpSTTSISSPQPGK (267–280), of which Serine 269 is phosphorylated; (**B**) MS/MS spectrum of a doubly-charged peak at m/z 776.318. The corresponding peptide is identified as AMpSTpTSISSPQPGK (267–280), of which both Serine 269 and Threonine 271 are phosphorylated.



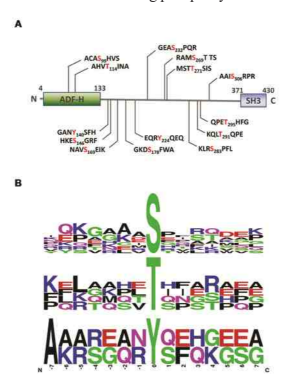
As another example, Figure 4A showed the MS/MS spectrum of a doubly-charged molecular ion peak with m/z at 697.321, from which a phosphopeptide as QLpTQPETHFGR(289–299) from HIP-55 protein was identified by database searching. The assignment of y series ions at m/z 971.4701, 1054.5074, 1152.4874 and 1167.5844 clearly indicated phosphorylation at Threonine 291, which was confirmed by detection of b series ions at m/z 242.1500, 325.1871, 423.1541 and 453.2197. The base peak at m/z 843.4137 was from the fragmentation at Proline 293. Figure 4B shows a similar spectrum of a doubly-charged molecular ion peak with m/z at 737.304, which was identified as the same peptide sequence (289–299) with two phosphorylation sites at Threonine 291 and 295. The detection of y series ions at m/z 697.3052, 826.3933, as well as the base peak ion at m/z 923.4073, confirmed the identification.

Figure 4. (**A**) MS/MS spectrum of a doubly-charged peak at m/z 697.321. The corresponding peptide is identified as QLpTQPETHFGR (289–299), in which Threonine 291 is phosphorylated; (**B**) MS/MS spectrum of a doubly-charged peak at m/z 737.304. The corresponding peptide is identified as QLpTQPEpTHFGR (289–299), in which both Threonine 291 and Threonine 295 are phosphorylated.



The phosphosite mapping of *HIP-55* protein were graphically displayed as Figure 5A. Furthermore, motif analyses of phosphosites were created and displayed using the Weblogo server (http://weblogo.berkeley.edu/). A modification site consists of the modified residue at the 0 position, plus the seven flanking amino acids *N*-terminal (positions –7 to –1) and *C*-terminal (positions +1 to +7) to the modification site. For the analysis of sequence features adjacent to the identified phosphosites of *HIP-55* protein, the 14 amino acids surrounding each phosphosite were extracted and aligned. The motif logo shows the difference in the frequency of amino acids surrounding the different phosphosites (serine, threonine, and tyrosine), suggesting the kinase recognition motif of these three kinds of phosphorylation sites are different in mammalian cells (Figure 5B). In addition, our results showed that serine and threonine residues undergo phosphorylation more often than tyrosine residues. The phospho-amino acid content ratio (pSer:pThr:pTyr) was 9:4:2 within *HIP-55* protein, which was consistent with previous reports of that in the whole-cell [17,18].

Figure 5. Graphical display and motif analysis of Phosphosites of *HIP-55* Protein. (A) Graphical display of phosphosite mapping of *HIP-55* Protein; (B) Frequency distribution of amino acid residues surrounding phosphorylation sites at positions –7 to +7.



Protein phosphorylation plays a significant role in a wide range of cellular processes. It is estimated that approximately one-third of all proteins in eukaryotic cells are phosphorylated at any given time [17]. Therefore, the mapping of phosphorylation sites of special interest proteins is the subject of a large body of research. Because of the prominent role of protein phosphorylation in signaling transduction, traditional studies focus on enzymes, receptors and ion channels which are switched "on" or "off" by phosphorylation and dephosphorylation in signaling transduction process [19–22]. However, one of the major goals of studying signal transduction is to determine the mechanisms that control cross-talk between signaling cascades and to determine how specificity in signaling is achieved. An emerging class of proteins that are major contributors to these processes are adaptor proteins [23]. By linking specific binding proteins together, adaptor proteins control cellular signals appropriately. Most adaptor proteins binding to partners function as signaling regulators in a phosphorylation-dependent manner. Therefore, identification of phosphorylation sites of adaptor proteins is of critical importance in the field of cellular signal transduction. HIP-55, as a signaling adaptor protein, has been shown as a common effector of antigen receptor-signaling pathways and regulates T-cell activation by bridging TCRs [7,9]. In our previous studies, we found Ser269/Thr291-phosphosites of HIP-55 mediated the interaction with 14-3-3τ (data not shown), which was confirmed by a recent research of 14-3-3-phosphoproteome [24]. Further, we also found pro-oncogenic function of HIP-55 through Ser269/Thr291-phospho-sensor motifs (data not shown).

3. Experimental Section

3.1. Chemicals and Materials

Sequencing-grade TPCK-modified trypsin was obtained from Promega (Madison, WI, USA). ACN, formic acid, TCEP, Supel-Tips Ti Pipette Tips were purchased from Sigma-Aldrich (St. Louis, MO, USA). EDTA-free protease inhibitor cocktail tablets were from Roche (Basel, Switzerland). Phosphatase inhibitor cocktail and Ni-NTA resin were from Pierce Biotechnology (Rockford, IL, USA). Anti-*HIP-55* antibody was purchased from BD Bioscience (San Diego, CA, USA). All other chemicals, as well as the Bradford protein assay kit, were from Bio-Rad (Hercules, CA, USA); the *HIP-55* plasmid was kindly provided by Prof. Haian Fu from Emory University. Ultra-pure water was prepared by a MilliQ water purification system (Millpore, Bedford, MA, USA).

3.2. Expression and Purification of HIP-55 Protein

The pDEST-His-*HIP-55* plasmid with full length human *HIP-55* gene was transfected transiently into HEK293 cells using Fugene HD (Roche) following the manufacturer's protocol. Cells were lysed 36 h post-transfection in His pull-down lysis buffer (1% Nonidet P-40, 137 mM NaCl, 1 mM MgCl₂, 40 mM Tris-Cl, 60 mM imidazole, 5 mM Na₄P₂O₇, 5 mM NaF, 2 mM Na₃VO₄, 1 mM phenylmethylsulfonyl fluoride, 10 mg/L aprotinin, 10 mg/L leupeptin). The lysate was cleared by centrifugation and a small aliquot of the supernatant was removed for Western Blot analysis. Then the supernatant was incubated with Ni-NTA resin at 4 °C for 2 h, which was pre-equilibrated with a buffer containing 100 mmol/L PBS, 6 mol/L guanidine HCl and 10 mmol/L imidazole; pH 7.4. After washes by the same buffer, the proteins bound to the Ni-NTA resin were eluted with a buffer containing 100 mmol/L PBS, 6 mol/L guanidine HCl and 250 mmol/L imidazole; pH 7.4.

3.3. Tryptic Digestion

The eluted proteins were precipitated by adding four volumes of -40 °C acetone and kept at -20 °C for at least 2 h. Then the protein pellet was redissolved with a buffer containing 0.1% SDS and 50 mmol/L ammonium bicarbonate, pH 8.5. The protein sample was reduced at 60 °C for 30 min by adding TCEP to a final concentration of 5 mmol/L, followed by alkylation with iodoacetamide at a final concentration of 4 mmol/L in the dark at room temperature for 60 min. Then the protein samples were cooled down to room temperature and mixed with freshly prepared sequencing-grade TPCK-modified trypsin buffer (15 μg/mL in 25 mmol/L ammonium bicarbonate, pH 8.5) at an enzyme/protein ratio of 1:100. Digestion was performed at 37 °C for at least 15 h and stopped by adding 10% formic acid to a final concentration of 1%. The tryptic digests were cleared by centrifugation and the supernatant was purified over C18 tips. For enrichment of phosphopeptides, the supernatant that had not been processed by C18 tips was loaded onto TiO₂ tips that were fully equilibrated with 3% formic acid. After washed with pure water three times, the bond peptides were eluted with 0.5% piperidine and then lyophilized.

3.4. Mass Spectrometry

The tryptic peptide samples were dissolved in buffer A (0.1% formic acid, 1% ACN), which were separated on a C18 reverse capillary column (150 mm \times 0.17 mm) with buffer B (0.1% formic acid, 99% ACN) over a 60 min gradient of 2%–40%. The eluted peptides were then delivered into an LTQ Orbitrap mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA) equipped with a nanospray source. The mass spectrometer was operated in a data dependent mode, in which five MS/MS scans on the most abundant ions detected in the MS scan (400–1500 m/z) were acquired. The dynamic exclusion time was set as 1 min. Singly-charged ions were excluded for MS/MS.

3.5. Peptide and Protein Identification

The collected spectra were analyzed by Thermo Proteome Discoverer 1.1 software (Thermo Fisher Scientific, Waltham, MA, USA), in which the built-in SEQUEST as used to search the data against IPI protein database (International Protein Index, ipi.HUMAN.v3.84, downloaded at ftp://ftp.ebi.ac.uk/pub/databases/IPI) with the following parameters: trypsin was selected with two maximum missed cleavage allowed; precursor mass tolerance was 15 ppm; fragment mass tolerance was 0.8 Da; carbamidomethylation of cysteine was set as static modification; phosphorylations of serine, threonine and tyrosine, as well as oxidation of methionine, were set as dynamic modifications. The results from SEQUEST searches were filtered with Peptide Confidence as High, in which XCorr threshold was set as 1.9, 2.3 for doubly, triply-charged ion peaks, respectively.

3.6. Motif Analysis of Phosphosites

Sequence features adjacent to the identified phosphosites (serine, threonine, and tyrosine) were created and displayed using the Weblogo server (http://weblogo.berkeley.edu/).

3.7. SDS-PAGE and Immunoblotting

Cell lysates (25 μg) were subjected to electrophoresis in 12% Tris-glycine-SDS polyacrylamide gel using a Mini-Cell system (Bio-Rad, Hercules, CA, USA). Gels were electroblotted to 0.2 μm pore size polyvinylidene fluoride (PVDF) membranes (Millipore, Bedford, MA, USA). Then the membranes were blocked with 1% nonfat dry milk in a buffer containing 25 mM Tris, pH 7.5, 150 mM NaCl, 0.05% Tween 20 for 1 h at room temperature. Membranes were then incubated with the primary antibody against *HIP-55* (BD Bioscience, San Diego, CA, USA) for 2 h at room temperature. After washing for 10 min in TBST solution, membranes were incubated with properly diluted secondary antibody conjugated with horseradish peroxidase (Sigma-Aldrich, St. Louis, MO, USA) for 1 h at room temperature. Western signals were developed with ECL chemiluminescent reagents (GE, Boston, MA, USA).

4. Conclusions

In the present study, HEK293 cells over-expressing human *HIP-55* protein were used to establish a profile of phosphorylation sites within *HIP-55*. Ni-NTA resin enrichment and gel electrophoresis

were utilized to collect the target protein and TiO₂ affinity chromatography and biological mass spectrometry were employed to investigate phosphorylation. Investigation of the mass spectra revealed 14 phosphorylation sites of *HIP-55* protein, several of which were identified for the first time. Our findings provide evidence for future investigations of the phosphorylation status of *HIP-55* protein of different subtypes and facilitate further studies on their biological functions in health and disease.

Acknowledgments

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Conflicts of Interest

The authors declare no conflict of interest.

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Mass Spectrometry Analysis Coupled with *de novo* Sequencing Reveals Amino Acid Substitutions in Nucleocapsid Protein from Influenza A Virus

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Abstract: Amino acid substitutions in influenza A virus are the main reasons for both antigenic shift and virulence change, which result from non-synonymous mutations in the viral genome. Nucleocapsid protein (NP), one of the major structural proteins of influenza virus, is responsible for regulation of viral RNA synthesis and replication. In this report we used LC-MS/MS to analyze tryptic digestion of nucleocapsid protein of influenza virus (A/Puerto Rico/8/1934 H1N1), which was isolated and purified by SDS poly-acrylamide gel electrophoresis. Thus, LC-MS/MS analyses, coupled with manual *de novo* sequencing, allowed the determination of three substituted amino acid residues *R452K*, *T423A* and *N430T* in two tryptic peptides. The obtained results provided experimental evidence that amino acid substitutions resulted from non-synonymous gene mutations could be directly characterized by mass spectrometry in proteins of RNA viruses such as influenza A virus.

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1. Introduction

Influenza virus has long been a global health threat since 1918 [1]. Although the annually circulating strains of influenza virus are not very virulent, there is still concern that the genome of these seasonal virus strains can mutate to acquire the ability to cause mortality in humans [2,3]. Additionally, for the avian influenza virus strains that usually show no adaptation to a human host, the virus genome can mutate to allow the virus to cross the species barrier to infect humans. For example, H5N1, H1N1 and recently reported H7N9 virus strains have shown their ability to cause severe infections in humans [4–6].

Mutations in viral genomes, some of which are non-synonymous mutations and thus result in amino acid substitutions, are often detected by gene sequencing [7–9]. With the introduction of soft ionization techniques such as ESI and MALDI, characterization of large biomolecules such as proteins has been achieved with high sensitivity and accuracy. Mass spectrometry has been used to analyze several mutations in hemoglobin variants [10,11]. Up to seven amino acid substitutions in HA of influenza A virus were revealed by mass spectrometry [12].

As influenza A virus has a relatively high mutation rate, there will always be an urgent need to detect variation in amino acid sequences resulting from non-synonymous SNPs that may have functional consequences. While both DNA and RNA have served as targets for most genotyping screen strategies, the other major functional molecule, protein, has recently been explored as a source for proteotyping, wherein a variety of protein forms from a single gene are characterized through

sophisticated mass spectrometric techniques [13]. Similar to DNA/RNA-based genotyping, proteotyping strategy can be applied on either a single protein [14] or on a proteome-wide scale [15]. Because influenza A virus continues to mutate to evolve, the previously established DNA/RNA-based PCR approaches often fail to detect the newly emerging strains due to sequence variation in primer and probe [16]. However, for the protein-based proteotyping strategy, the mutated peptides or modified peptides van be detected without the concerns in PCR approaches. Therefore, once the proteotyping strategy is optimized for any given strain, it should be effective to detect an array of isoforms of viral proteins, including the peptides upon modification and amino acid substitution. In this study, we report the characterization of the nucleocapsid protein (isolated and purified by SDS-PAGE) of influenza A virus by mass spectrometry. By manual interpretation of the MS/MS data, three amino acid substitutions were identified. The results indicated that mass spectrometry coupled with *de novo* peptide sequencing had the power to characterize the amino acid substitutions in proteins of RNA viruses such as influenza A virus.

2. Results and Discussion

2.1. Identification of NP Protein

Influenza virus was inoculated in chick embryos. Several serial passages were performed to enhance the rates of multigenic mutations. The virus particles were purified from the collected allantoic fluid and then lysed and separated on 12% SDS-PAGE. After staining with Colloidal Coomassie G250, two major bands were found at 15 and 56 kDa, respectively (Figure 1). The band at 56 kDa was cut off and subject to in-gel tryptic digestion. LC-MS/MS analysis of the obtained peptide mixture coupled with protein database searching identified a total of 18 unique peptides of nucleocapsid protein from influenza virus (A/Puerto Rico/8/1934 H1N1) (Table 1).

Figure 1. Identification of nucleocapsid protein from purified influenza virus (A/Puerto Rico/8/1934 H1N1). The purified virus was lysed and separated on 12% SDS-PAGE. The upper band around 56 kDa was cut off and subjected to in-gel digestion, followed by mass spectrometric analysis. Database searching identified 18 tryptic peptides (labeled with red) from nucleocapsid protein of influenza A virus. Manual interpretation of the obtained MS/MS data identified three amino acid substitutions (*R452K*, *T423A* and *N430T*, highlighted with yellow) within two tryptic peptides (labeled with green).

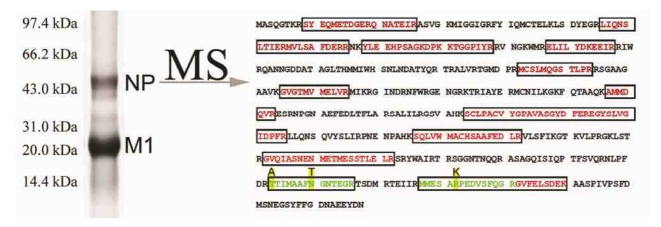


Table 1. Summary of tryptic peptides identified in nucleocapsid protein from influenza virus (A/Puerto Rico/8/1934 H1N1) by database searching.

Peptide No.	Peptide sequence	Charge status	Calculated m/z (monoisotopic)	Measured m/z (monoisotopic)	Residues
P1	TGGPIYR	2	382.21	382.21	92–98
P2	KTGGPIYR	2	446.26	446.29	91–98
Р3	A <u>MM</u> DQVR	2	441.69	441.70	237-243
P4	QNATEIR	2	416.22	416.22	20–26
P5	GVFELSDEK	2	512.25	512.26	462-470
P6	MVLSAFDER	2	534.26	534.27	66–74
P7	YLEEHPSAGK	2	565.78	565.78	78–87
P8	YLEEHPSAGKDPK	2	735.86	735.86	78–90
P9	LIQNSLTIER	2	593.84	593.85	56–65
P10	GVGT <u>M</u> V <u>M</u> ELVR	2	612.31	612.32	185–195

Table 1. Cont.

Peptide No.	Peptide sequence	Charge status	Calculated m/z (monoisotopic)	Measured m/z (monoisotopic)	Residues
P11	MVLSAFDERR	2	620.31	620.31	66–75
P12	EGYSLVGIDPFR	2	676.85	676.85	294-305
P13	MCSLMQGSTLPR	2	706.82	706.82	163-174
P14	SYEQMETDGER	2	680.77	680.77	9–19
P15	ELILYDKEEIR	2	710.89	710.89	107-117
P16	GVQIASNENMETMESSTLELR	2	1170.05	1170.07	362-382
P17	SCLPACVYGPAVASGYDFER	2	1110.00	1110.00	274-293
P18	SQLVWMACHSAAFEDLR	2	1010.97	1010.98	326-342

<u>M</u>: mono-oxidized methionine.

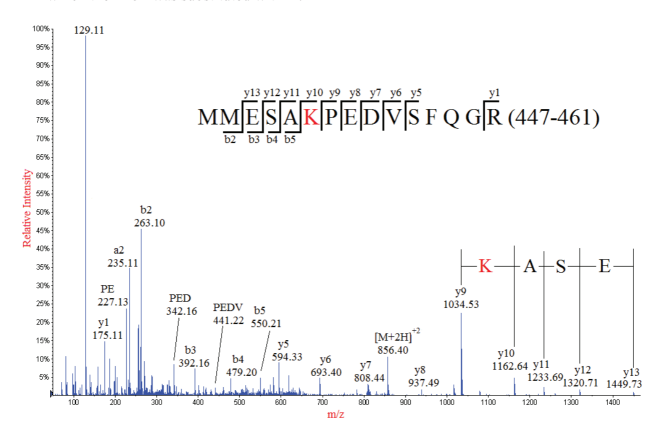
Besides the peptides identified by database searching, two additional mutated peptides were determined by manual interpretation of the available data, in which three amino acid substitutions were identified. Accordingly, both database searching and manual interpretation of the obtained LC-MS/MS data allowed the assignment of a total of 20 unique peptide sequences.

2.2. Identification of AA Substitution of R452K

Interpretation of the MS/MS spectrum of the doubly-charged ion peak MP1 at m/z 856.40 (Figure 2) allowed the identification of a partial sequence of ESA, considering the ion series of m/z1449.73, 1320.71, 1233.69, 1162.64 at the high mass end of the spectrum were y type fragment ions y13, y12, y11, y10, respectively. The sequence of ESA was readily to be located in one of the tryptic peptides of NP: MMESARPEDVSFQGR (447-461) with theoretical m/z value of 870.40 for its doubly-charged ion. Thus, a nominal mass shift of -28 Da was observed for the detected doubly-charged ion of MP1 in comparison with the molecular weight of the theoretical sequence of MMESARPEDVSFQGR (447–461) in NP, which might result from amino acid substitution of one of five residues in the theoretical sequence: $R \rightarrow Q/K$, $V \rightarrow A$, $M \rightarrow C$, $D \rightarrow S$ or $E \rightarrow T$. The possibility for amino acid substitution of M→C was readily eliminated because the Cys (C) residue would be chemically alkylated during sample preparation if Methionine (M447/448) was mutated into Cys (C). Noticeably, the fragment ion y9 at m/z 1034.53 adjacent to y10 ion (m/z 1162.64) in the high mass range of the MS/MS spectrum indicated that the residue next to the Alanine (A₄₅₁) should be either K or Q, considering that the calculated difference between 1162.64 and 1034.53 was identical to the nominal mass of 128 of either of these two amino acid residues. In addition, substitution of Arginine (R₄₅₂) with either Lysine (K) or Glutamine (Q) was also confirmed by the detection of the base peak at m/z 129, which was the immonium ion of either K or Q. Although both K and Q residues had identical nominal mass of 128, the exact masses of them were different (K with 128.095 and Q with 128.058). The precise mass difference between y9 and y10 was calculated as 128.11, suggesting that the R₄₅₂ was substituted by K, but not Q. This conclusion was well supported by the precise mass data of the immonium ion detected at m/z 129.11, which was much closer to the theoretic mass data of immonium ion of K (129.1022) than that of Q (129.0659). The assignments of most y series ions (from v5 to v13) clearly demonstrated the internal sequence of PEDV, eliminating the possibilities

of amino acid substitutions at E₄₅₄, D₄₅₅ and V₄₅₆. It should be noticed that there was a Proline (P) in the sequence, at which internal fragmentation could occur. Some internal sequences such as PE, PED, PEDV and PEDVS were detected and assigned, confirming that amino acid substitution should occur at R₄₅₂ but not E₄₅₄, D₄₅₅ and V₄₅₆. Additionally, detection of some of the a and b series ions such as a2, a3, b2 and b3 indicated that E₄₄₉ was not subject to amino acid substitution, confirming the substitution of R₄₅₂ \rightarrow K.

Figure 2. MS/MS spectrum of the doubly-charged ion at m/z 856.40 from the analysis of peak MP1.The mutated peptide (MMESAKPEDVSFQGR) of a normal sequence (residues 447–461) from tryptic digestion of nucleocapsid protein was identified, in which the R452 was substituted with K.



2.3. Identification of AA Substitution T423A and N430T

De novo sequencing of the MS/MS spectrum of a doubly-charged ion peak MP2 at m/z 720.36 identified a partial sequence of TIMAAFT with y series ions at m/z 1368.68, 1267.71, 1154.61, 1023.56, 952.53, 881.46, 734.42 and 633.35 (Figure 3), which was not found in the theoretical sequence of NP. However, investigation of the theoretical sequence of NP revealed a sequence of TIMAAFN (424–430), which was identical to the deduced sequence except for the N₄₃₀ residue. Therefore amino acid substitution of N₄₃₀ \rightarrow T was identified, which resulted in a mass shift of -13.01 Da. The identified sequence was contained in a tryptic peptide of NP: TTIMAAFNGNTEGR (423–436), of which the calculated m/z value of the doubly-charged ion was 741.86. However, a nominal mass shift of -43 but not -13.01 Da (N₄₃₀ \rightarrow T) was observed for MP2 when compared to the theoretical sequence of TTIMAAFNGNTEGR (423-436), suggesting that there might be at least

one additional amino acid substitution in the sequence, which resulted in an additional mass shift of -30.01 Da. Investigation of the rest of the residues of the tryptic peptide sequence (423–436) indicated that there were three amino acid residues that could result in a mass shift of -30.01 Da upon substitution: $T_{423}\rightarrow A$, $T_{433}\rightarrow A$, $T_{434}\rightarrow T$. The detection of the *y* series ion *y*13 at m/z 1368.66, as well as b_3 -H₂O, b_3 , b_2 -H₂O and b_2 ions, indicated that the first three residues in the peptide were ATI, thus confirming the identification of substitution of $T_{423}\rightarrow A$. Therefore, the peak MP2 was identified as the tryptic peptide in the residues from 423 to 436 with the two substitutions, namely $T_{423}\rightarrow A$ and $N_{430}\rightarrow T$.

Figure 3. MS/MS spectrum of the doubly-charged ion at m/z 720.36 from the analysis of peak MP2. The mutated peptide (ATIMAAFTGNTEGR) of a normal sequence (residues 423–436) from tryptic digestion of nucleocapsid protein was identified, in which the T423 and N430 were substituted with A and T, respectively.

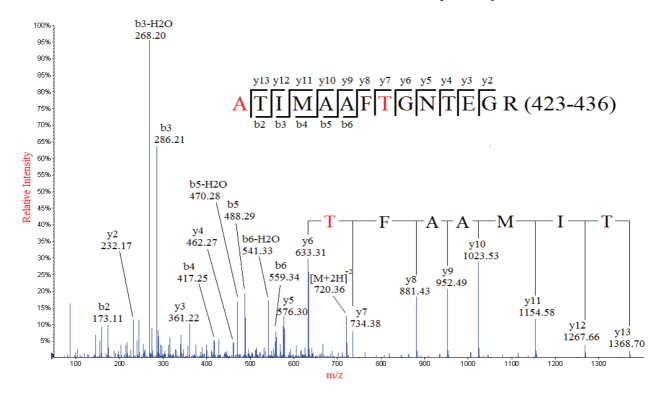
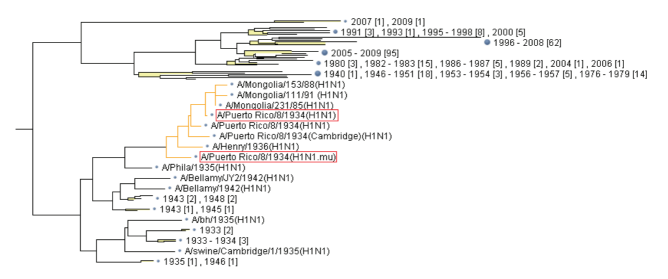


Figure 4. Part of detailed tree picture generated from linkage analysis of mutated nucleocapsid protein identified by MS/MS, by which the neighbor-joining method of clustering was used. The strain containing the mutated nucleocapsid protein (A/Puerto Rico/8/1934 (H1N1.mu)), as well as its original strain (A/Puerto Rico/8/1934 (H1N1)), is labeled with red rectangular box.



2.4. Bioinformatics Analysis

The sequences of nucleocapsid proteins were exclusively retrieved from "The FLU project" at GenBank. A protein sequence database containing the retrieved sequences and the mutated sequence was built and subject to multiple alignment and linkage tree analysis (Figure 4). The output file containing the whole tree data can be found in supplemental materials.

3. Experimental Section

3.1. Chemicals and Materials

Sequencing-grade TPCK-modified trypsin was purchased from Promega (Madison, WI, USA). Bradford protein assay kit, ammonium bicarbonate, dithiothreitol (DTT), iodoacetamide (IAA) were purchased from Bio-Rad (Hercules, CA, USA). All the other chemicals were purchased from Sigma-Aldrich (St. Louis, MO, USA). Influenza virus (A/Puerto Rico/8/1934 H1N1) was propagated in a biosafety level 2 (BL-2) containment facilities. Ultra-pure water was prepared by a MilliQ water purification system (Millpore, Bedford, MA, USA).

3.2. Virus Cultivation and Purification

Embryonated chicken eggs were inoculated with the influenza A virus (A/Puerto Rico/8/1934 H1N1) and incubated for 72 h at 37 °C. The allantoic fluid was harvested, followed by centrifugation at 5000 rpm for 15 min. The virus in the allantoic fluid was pelleted through a 4-step discontinuous gradient cushion consisting of 30%, 40%, 50% and 60% (w/v) sucrose, in a SW40 Ti rotor (Beckman-coulter, Fullerton, CA, USA) at 35,000 rpm at 4 °C for 60 min. The virus band between

40% and 50% sucrose was carefully collected, and suspended in 10 mM Tris-HCl pH 8.0, 150 mM NaCl. Aliquots of the purified virus sample were kept at 4 °C.

3.3. SDS-PAGE

The purified virus particles were lysed with 2× Laemmli sample buffer and kept at 95 °C for 5 min. The protein concentration was assayed with Micro BCA (bicinchoninic acid) protein assay kit (Pierce, Rockford, IL, USA). Electrophoretic separation was performed in a Mini-Cell system (Bio-Rad, Hercules, CA, USA), and run in 12% tris-glycine-SDS polyacrylamide gels with a 5% stacking gel. After electrophoresis, the gels were stained with colloidal Coomassie G250 and scanned with a calibrated densitometer (GS800, Bio-Rad).

3.4. In-Gel Digestion

Protein bands of interest were cut off from gels and washed with *Milli-Q* water three times. Then the gel pieces were destained with a solution of 50 mM NH₄HCO₃ in 50% ACN until the Coomassie blue in the gel became invisible. The destained gel pieces were reduced in 10 mM DTT, 50 mM NH₄HCO₃ aqueous solution at 60 °C for 60 min, followed by alkylation in 50 mM IAA, 50 mM NH₄HCO₃ aqueous solution at room temperature in dark for 30 min. The gel pieces were dehydrated with ACN, and then incubated in freshly prepared digestion solution containing 50 mM NH₄HCO₃ and 0.1 g/L TPCK-trypsin overnight at 37 °C. The resulting tryptic peptides were extracted with 5% trifluoroacetic acid (TFA) in 60% ACN and stored at -20 °C until LC-MS/MS analysis.

3.5. Capillary LC-MS/MS Analysis

The tryptic peptides were lyophilized and redissolved in high performance liquid chromatography (HPLC) buffer A (0.1% formic acid) and then separated on a C18 column (100 mm \times 180 μ m i.d.). The elution gradient was from 5% to 40% buffer B (0.1% formic acid, 99% ACN, flow rate: 0.2 μ L/min) for 90 min. The eluted peptides were then analyzed on an ABI QSTAR spectrometer using information dependent acquisition mode (IDA; Analyst QS, Applied Biosystems, Carlsbad, CA, USA) by selecting the three most intense ions for MS/MS analysis. A survey scan of 300–2000 Da was collected for 3 s followed by 5 s MS/MS scans of 40–1500 Da using the standard rolling collision energy settings. The dynamic exclusion time was set as 1.5 min.

MASCOT generic files were generated from the obtained MS data by using a script embedded in the Analyst QS 2.0 software (MDS Sciex, South San Francisco, CA, USA) and used to search against the Swiss-Prot protein database on a local MASCOT server (version 2.1, Matrix Science, London, UK). One missed cleavage was allowed. Carbamidomethylation of cysteines was specified as fixed modification, whereas oxidation of methionine was selected as variable modification. The mass tolerance was set to 0.3 and 0.6 Da for peptide and MS/MS ion masses, respectively. Manual *de novo* sequencing of peptide tandem mass spectra was performed with the aid of Pepsea (1.1) in Analyst QS 2.0 software (MDS Sciex).

3.6. Bioinformatics Analysis

The mutated nucleocapsid protein containing three amino acid substitutions was analyzed by using a suite of bioinformatics tools at NCBI (http://www.ncbi.nlm.nih.gov/genomes/FLU/FLU.html) [17].

4. Conclusions

We herein identified by LC-MS/MS analysis three amino acid substitutions in nucleocapsid protein from influenza virus (A/Puerto Rico/8/1934 H1N1). The three amino acid substitutions were located in two tryptic peptides of the nucleocapsid protein. One of identified amino acid substitutions, *R452K*, was located within the tryptic peptide MP1 (447–461), whereas the other two amino acid substitutions, *T423A* and *N430T*, were located within tryptic peptide MP2 (423–436). Both of the peptides were identified through manual interpretation of the relating MS/MS data, which included both calculation of high resolution MS data and assignment of fragment ions in MS/MS data. The outcome of this study indicated that the MS/MS analysis of amino acid substitutions might be useful in investigating the antigens from influenza viruses.

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Conflicts of Interest

The authors declare no conflict of interest.

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