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Quantification of pharmaceutical peptides in human plasma by LC-ICP-MS sulfur detection

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A method for quantification of a pharmaceutical peptide in human plasma was developed using gradient elution LC-ICP-MS. A membrane desolvation (MD) system was applied to remove organic solvents from the eluent prior to the detection as SO⁺ in the dynamic reaction cell (DRC) of the ICP-DRC-MS instrument and subsequent quantification by post-column isotope dilution (IDA). Plasma proteins were precipitated prior to analysis. Analytical figures of merit including linearity, precision, LOD, LOQ and accuracy were considered satisfactory for analysis of plasma samples. The selectivity of the developed method was demonstrated for five pharmaceutically relevant peptides: desmopressin, penetratin, substance P, PTH (1-34) and insulin. Preliminary experiments on an ICP-MS/MS system using oxygen to reduce the effect of organic solvents were also performed to compare sensitivity. The results of the study demonstrated that LC-ICP-MS post-column IDA may constitute a valuable additional tool in quantification of non-labelled peptides in the early drug development offering absolute quantification without need of species specific standards.

Introduction

The interest in peptides and proteins as drugs and excipients is continuously increasing in biomedical research. The number of peptide therapeutics reaching preclinical and clinical trials increases every year. Peptides are favorable as therapeutics as they offer good efficacy, safety and tolerability due to their composition of amino acids.1 During drug development, parameters such as stability, pharmacokinetics and metabolism must be monitored; therefore, sensitive and selective analytical methods are required. Quantitative analysis of peptides in biological matrices is challenging due to the similarity of pharmaceutical peptides to endogenous compounds. Peptide quantification has predominantly been performed by fluorescence detection (FLD) exploiting fluorophores naturally present in the peptide sequence or by adding a fluorescent label.2 Additionally, immunoassays, e.g. ELISA, are widely used for quantification of peptides and proteins.3 Nowadays, identification and quantification is predominantly performed by liquid chromatography mass spectrometry (LC-MS). Analysis of peptides by LC-MS in complex biological matrices such as plasma demands thorough sample preparation and species specific peptide standards for absolute quantification.4

The large potential of inductively coupled plasma mass spectrometry (ICP-MS) in quantitative analysis of biomolecules has already been highlighted in a number of reviews.5-7 Hyphenated to a separation technique, ICP-MS is a powerful technique for quantitative analysis of biomolecules offering low limits of detection, a wide dynamic range and in principle species independent sensitivity, thus, quantification using a single inorganic standard is possible. The premise of using ICP-MS for detection is the presence of an ICP-MS detectable element in the peptide structure such as sulfur (S), phosphorus (P) or selenium (Se), otherwise an ICP-MS detectable label must be inserted. Introduction of elemental labels as metal-chelating moieties or metal affinity tags has been widely reported.5,8 Labelling with metals ensure high sensitivity and selectivity, but the drawback of these approaches is alteration of the structure and physicochemical properties of the peptide. Furthermore, the stoichiometry of the label and the molecule must be controlled and known for quantitative purposes. Unequivocal stoichiometry is achieved using heteroelement tags, which are covalently bound in the peptide structure. Recently, we introduced Se as an elemental detection label for quantification of peptides.9 By exchange of the S-containing amino acid methionine (Met) with the Se-containing analogue selenomethionine (SeMet) during peptide synthesis, we introduced a label causing minimal alteration of the native peptide structure and physicochemical properties.10 The most attractive approach would therefore be to monitor a heteroelement in the native structure of the peptide. S is present in the natural occurring amino acids

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Met and cysteine (Cys) and thus constituting a versatile label in the majority of biomolecules, but measuring S in complex biological matrices by ICP-MS is not straightforward. The ionization efficiency of S in the argon plasma is limited due to the high first ionization energy (10.357 eV). Furthermore, detection of the main S isotope (32S, 94.99% (ref. 11)) is hampered by the formation of polyatomic interferences in the plasma, mainly dimers of oxygen (16O2+). Different instrumental setups and approaches to overcome these interferences have been developed. One approach is to use a quadrupole based collision/ reaction cell ICP-MS instrument. By using xenon as collision gas polyatomic interferences are removed by collision induced dissociation and allows for measurement of 32S.12 The main approach for collision/reaction cell instruments is to use oxygen as reaction gas for the formation of sulfur oxide (32S16O+) which is less interfered as compared to the S isotopes. 13 The application of high resolution sector field (SF) instruments facilitates the separation of the different masses of $^{32}S^+$ (m/z = 31.9898) and $^{16}O^{16}O^{+}$ (m/z = 31.9721) can be separated at a resolution of 4000 $(m/\Delta m)$. ^{14,15} Recently, ICP-MS/MS was introduced offering improved removal of polyatomic interferences and further reduction of the background level. The LOD of this instrument has thus been reported to be lowered to the ng $\rm L^{-1}$ range for S. ^{16,17}

Measurement of S in biological matrices demands hyphenation of a separation method to the ICP-MS in order to separate analytes from natural occurring S-species in the matrix. Peptide separation is most often performed by reversed-phase liquid chromatography (RP-LC) and gradient elution using either acetonitrile or methanol.18 The continuous change in organic solvent load from the eluent is not readily compatible with ICP-MS as it may cool the plasma, change ionization and thus potentially change sensitivity. Exposure to very high concentrations of organic solvent may even extinguish the plasma. Furthermore, the introduction of organic solvents may cause carbon build-up on the cones and decrease sensitivity over time. 19,20 By application of a membrane desolvation (MD) system the organic solvent is removed from the eluent prior to introduction to the ICP-MS and up to 100% methanol or acetonitrile at flow rates of 200 μL min⁻¹ are tolerated.21 Optimization of the MD is only possible for a single concentration level of organic solvent and the removal of eluent may therefore not be constant throughout the gradient, causing problems in species unspecific calibration. Isotope dilution analysis (IDA) can be applied to overcome this problem.22 The technique is based on measurement of the isotope ratios in a sample spiked with a solution enriched with a minor natural abundant isotope of the element. In species unspecific IDA, also known as post-column IDA, the enriched isotope is added to the chromatographic eluent after separation of the sample.²³ The technique and the equations used in calculations have been presented in a tutorial review by Rodriguez-Gonzáles.24 Sulfur post column IDA has been applied in several studies regarding protein quantification by ICP-MS. 22,25,26 The number of quantitative studies of peptides or proteins in complex biological matrices using ICP-MS detection of sulfur is very limited. Konz et al. developed a method for quantification of the peptide hepcidin in human urine by post column IDA27 but, the majority of studies does not report on using IDA for quantification.28-30

The aim of this study was to examine if it is possible to quantify therapeutically relevant levels of sulfur-containing pharmaceutical peptides in human plasma samples by LC-ICP-MS and post-column IDA.

Experimental

Instrumentation

ICP-DRC-MS. The ICP-MS instrument was an ELAN DRCe (Perkin Elmer SCIEX, Norwalk, CT, USA). Oxygen was applied as reaction gas with the following settings of the dynamic reaction cell: cell gas flow 0.35 L min^{-1} and R_{pq} 0.45. The sampler and skimmer cones were made of nickel. Lens voltage and RF power were optimized on a daily basis using a 100 μg S L⁻¹ sulfate solution in mobile phase (35% v/v acetonitrile (ACN), 0.1% v/v trifluoroacetic acid (TFA)). Data acquisition settings for speciation analysis were: dwell time 200 ms; sweeps per reading 1; and readings per replicate were adjusted to fit the time of the chromatographic run, 48SO+ and 50SO+ were monitored. A desolvation system (Aridus II, CETAC Technologies, Omaha, NE, USA) equipped with a 200 μL min⁻¹ C-flow PFA concentric nebulizer (CETAC Technologies) was employed to remove organic solvent from the eluent. Desolvation system settings: spray chamber temperature 110 °C, desolvator temperature 160 °C, sweep gas flow 7 L min⁻¹ and nitrogen gas flow 3 mL min⁻¹.

ICP-MS/MS. The analyses were performed using an Agilent 8800 ICP-MS/MS instrument equipped with a quartz injector torch with a 1.5 mm internal diameter (ID), platinum sampling and skimmer cones, a brass skimmer base and an x-lens (Agilent Technologies, Santa Clara, USA). An Ari Mist HP nebulizer from Burgener Research (Mississauga, ON, Canada) and a Peltier-cooled double-pass Scott type spray chamber at -5 °C (Agilent Technologies) were used for sample introduction. The ICP-MS/MS parameters were optimized using a sulfate solution containing 100 μg L⁻¹ S in mobile phase (35% v/v acetonitrile, 0.1% v/v TFA). Monitoring the m/z 48 and 50 for ⁴⁸SO⁺ and ⁵⁰SO⁺, respectively, the plasma conditions were optimized automatically on a daily basis. The custom tune function was applied for optimization of S in order to improve the specific sensitivity. The carrier gas (Ar) flow rate was set to 0.36 L min⁻¹ and the option gas, a blend of 20% oxygen in argon, was added at a ratio of 34% to the plasma in order to prevent a carbon build-up on the cones. The RF power was 1600 W and the sampling depth 7.5 mm. The ICP-MS/MS analyses were carried out applying the MS/MS mode and oxygen as reaction gas at a cell gas flow of 35% (corresponding to 0.35 mL min^{-1}).

LC-MD-ICP-DRC-MS. An Agilent 1100 series HPLC system equipped with a degasser, a quaternary pump, an auto sampler and a column oven was used for separation. The column was an Aeris PEPTIDE XB-C18, 3.6 μ m, 100 \times 2.1 mm ID column protected by a C18-Peptide SecurityGuard ULTRA cartridge (Phenomenex, Denmark). A multi-step gradient from 17-70% v/v of acetonitrile (with 0.1% v/v TFA added) was applied (0-1 min: 17-20%, 1-5 min: 20-35%, 5.1-5.5 min: 38%, 5.6-6 min: 70%). The flow rate was 200 μ L min⁻¹, the column temperature 60 °C, and the injection volume 5 µL.

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LC-ICP-MS/MS. A Dionex Ultimate 3000 UHPLC (Thermo Scientific) equipped with a degasser, a quaternary pump, an auto sampler and a thermostatted column compartment was used for separation. The column and mobile phase were as described above for LC-MD-ICP-DRC-MS, except the gradient was slightly modified in order to move the peptide peak further away from

the front peak of the chromatogram (0-2 min: 17%, 2-3 min:

17–20%, 3–7 min: 20–35%, 7.1–7.5 min: 38%, 7.6–8 min: 70%). Post column isotope dilution analysis. Quantification was performed by post column IDA using an enriched 34S sulfate standard solution 1000 µg S L⁻¹ in 35% v/v acetonitrile and 0.1% TFA. The ³⁴S solution was infused constantly at a flow rate of 10 µL min⁻¹ using a syringe pump, Harvard Apparatus 11 plus (Holliston, MA, USA) with a 5 mL syringe (Hamilton, Bellefonte, PA. USA). The flow rate was corrected from mass per minute to volume per minute using the relevant density. The syringe pump was connected to the eluent flow from the HPLC by a static mixing T-piece. The connection was made of PEEK tubing with an inner diameter of 0.125 mm before mixing and an inner diameter of 0.250 mm after mixing. The experimental isotope ratio 48SO/50SO was calculated on a daily basis by measurement of 100 µg L⁻¹ S (sulfate) standard solution in mobile phase (35% v/v ACN, 0.1% v/v TFA) and corrected for blank signal and mass bias by using an exponential mathematical model.24 Theoretical isotope ratios were calculated by using the natural sulfur abundances.11 The dead time was corrected automatically by the instruments. Mass flow chromatograms were obtained by application of the isotope dilution equation to each data point of the chromatograms. Peak areas were determined by integration of the chromatographic peaks using the OriginPro 2015 software (version 9.2, OriginLab, Northampton, MA, USA).

Chemicals. Penetratin (Pen) was purchased from CASLO (Lyngby, Denmark). Desmopressin European Pharmacopeia certified reference standard (CRS) and substance P was purchased from Sigma (St. Louis, MO, USA). PTH (1-34) was from Bachem (Bubendorf, Switzerland) and human insulin was kindly provided by Sanofi. ACN LC-MS grade was obtained from Th. Geyer (Renningen, Germany), TFA from VWR International (Lutterworth, UK), and trichloroacetic acid (TCA) from Fluka Chemie, (Buchs, Switzerland). Human plasma was obtained from the blood bank (Rigshospitalet, Copenhagen, Denmark). The 99.90% ³⁴S isotope was purchased from ChemGAS (Boulogne, France).

Standards. All peptide solutions were prepared in NoStick® hydrophobic microtubes from SSI (Lodi, CA, USA). An amount of 1 mg of Pen was dissolved in 1 mL of purified water. From this solution, samples for validation were prepared. Desmopressin CRS (0.94 mg as described in the certificate) was dissolved in 1 mL purified water. Substance P and PTH (1-34) was dissolved in purified water and human insulin was dissolved in 0.1% TFA all in concentrations of 1 mg mL⁻¹.

The ³⁴S enriched isotope standard was prepared by dissolving 20 mg elemental ³⁴S in 3 mL sub boiled nitric acid (65%) and digested in a microwave until a clear and colorless solution was obtained. The solution was diluted with purified water to

20 mL. The total S concentration of the solution was determined by reversed isotope dilution using a PlasmaCAL elemental standard of S.

Procedures

Sample preparation. Frozen human plasma was thawed and filtered through a Q-max RR syringe filter, 0.45 μ m cellulose acetate membrane. Defined amounts of Pen were added to human plasma reaching final concentrations of 2–50 μ M Pen in 50% plasma in purified water using NoStick® hydrophobic microtubes. Plasma proteins were precipitated by addition of 10% v/v TCA in purified water in volume ratio 1 : 5 of TCA and sample. The samples were centrifuged at 14 000g for 15 min at room temperature. The supernatant was diluted 1 + 1 in eluent (10% v/v ACN, 0.1% v/v TFA) prior to analysis.

Sample preparation by ultrafiltration was executed in 50% human plasma diluted in purified water by addition of TFA to a final concentration of 0.5% and ultrafiltration through 500 μ L centrifugal filter units, Amicon® Ultra with a semi-permeable membrane of molecular weight cut-off 30 kDa. The ultra filtrates were diluted 1 + 1 in eluent (10% v/v ACN, 0.1% v/v TFA) prior to analysis.

Sample preparation recovery. The recovery was determined in triplicates by spiking 10 μ M and 40 μ M Pen into 50% plasma before and after sample preparation. Recoveries were calculated as the ratio of peak areas of plasma spiked before and after sample preparation (mass flow chromatograms).

Linearity. Linearity was determined by addition of defined amounts of Pen to plasma prior to sample preparation reaching final concentrations of: 2, 5, 10, 20 and 50 μ M in 50% plasma. The samples were diluted 1 + 1 in eluent (10% v/v ACN, 0.1% v/v TFA) prior to analysis.

Precision. Precision was determined as the relative standard deviation (RSD%) of five injections of a 10 μ M sample after sample preparation and dilution as described above.

Limit of detection (LOD) and limit of quantification (LOQ). A sample of 2 μ M Pen spiked into plasma prior to sample preparation was analyzed (5 injections). The standard deviation of the mass flow chromatogram peak areas was calculated and LOD and LOQ determined as three times and ten times the standard deviation, respectively.

Accuracy. Accuracy of the method was determined by analysis of 5 μ M S (2.5 μ M desmopressin Ph. Eur. CRS) (three injections).

Selectivity. Selectivity of the chromatographic method was demonstrated for five S-containing, pharmaceutical peptides: desmopressin, Pen, substance P, PTH (1-34) and human insulin. The peptides were spiked individually in ultrafiltrated plasma to a final concentration of peptide corresponding to 10 μ M S.

Results and discussion

Method development

The aim of this study was to examine if it is possible to quantify pharmaceutical S-containing peptides in human plasma by JAAS Paper

LC-ICP-MS using the cell-penetrating peptide Pen as a model compound. Reversed phase gradient separation using methanol or acetonitrile is the preferred method for separation of peptides. In order to overcome the challenges arising from introduction of organic solvents to the ICP-MS, a membrane desolvation (MD) system (Aridus II, Cetac) was applied. The use of the MD system limits the flow rate of the separation method, as only nebulizers with a flow rate up to 200 μL min $^{-1}$ are available for sample introduction. Furthermore, the MD extends the distance between the HPLC column and the ICP-MS causing peak broadening. Narrow peaks and excellent resolution is therefore demanded for the separation when a MD system is applied.

Recently we developed an LC-ICP-MS method applying a linear methanol gradient (0.1% v/v acetic acid, 0.05% v/v TFA) for separation of a Se-labelled analogue of Pen and its degradation products during cell uptake studies. Separation of S-containing peptides in human plasma is more challenging due to the presence of large amounts of endogenous S compounds in the sample matrix. In order to separate low molecular weight S compounds and S-containing proteins in plasma, a multi-step acetonitrile (0.1% v/v TFA) gradient was developed for separation of Pen from S-containing plasma constituents. The column temperature was set to 60 °C to obtain narrow peaks and improve resolution. Satisfactory peak shape and separation of Pen from plasma components were obtained.

Quantification by post column isotope dilution analysis

The species independent sensitivity of LC-ICP-MS is lost when gradient elution is applied, as the shift in organic content in the eluent changes the sensitivity during elution of the species. Thus, the instrument sensitivity of early eluting species will differ from later eluting species. In order to compensate for these variations in sensitivity as well as drift of the instrument signal, post column IDA was employed for absolute quantification (details are described in the Experimental section). The mass bias factor (F) was estimated and applied in correction of the experimental isotope ratios of 32 S/ 34 S. Subsequent to correction, the isotope dilution equation²⁴ was applied in the entire chromatogram to obtain the mass flow data and the peak areas were integrated. An example of a mass flow chromatogram obtained from MD-ICP-DRC-MS analysis of blank plasma and plasma spiked with Pen is shown in Fig. 1.

Sample preparation

Plasma proteins should be removed prior to LC-ICP-MS analysis, but it is important that the analyte remains in the sample with a satisfactory and reproducible recovery of analyte. Working with peptide-containing samples can be a challenge due to unspecific adsorption of peptides to surfaces and other constituents of the matrix *e.g.* proteins. Adsorption is often dependent on the specific side chains on the amino acids of the peptide and may be reduced by choosing conditions which affects the equilibrium constant between the peptide and surfaces or between the peptide and proteins which it may adsorb to.³¹ Pen contains 16 amino acids (RQIKIWFQNRRMKWKK-NH₂) of which 7 are

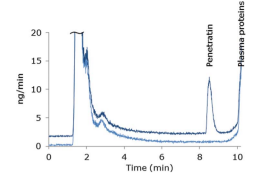


Fig. 1 Mass flow chromatograms from LC-MD-ICP-DRC-MS analysis of 50% plasma samples. Light blue: blank plasma, offset by -1.2 ng \min^{-1} . Dark blue: 50 μ M Pen spiked in 50% plasma prior to sample pretreatment.

basic residues, and thus have a net charge of +8 at physiological pH. In order to reduce ionic interactions of the peptide with the surroundings, negatively charged residues were reduced by acidifying samples during sample pretreatment. Two different approaches for removal of plasma proteins were pursued; precipitation and ultrafiltration. TCA (10% v/v, volume ratio 1:5) was chosen as precipitation reagent, ensuring acidic conditions. A protein precipitation efficiency above 99% has been reported for TCA.32 To determine the recovery of peptide after precipitation, spiked plasma samples were precipitated and the areas from the mass flow chromatograms compared to the areas of plasma samples spiked after protein precipitation. Recovery was determined at two concentration levels, 10 µM and 40 µM Pen; recoveries of 70.6 \pm 6.2% (n = 3) and 69.2 \pm 4.0% (n = 3) were obtained, respectively. Chromatograms of plasma samples showed that some S-containing protein still remained after precipitation with 10% TCA. For ultrafiltration, a semipermeable membrane with a molecular weight cut-off (MWCO) value of 30 kDa was used and the plasma acidified (0.5% v/v TFA) prior to filtration. S-containing plasma proteins were almost completely removed by ultrafiltration, but the recovery of 10 µM Pen was only 32.4 \pm 9.0%. Ultrafiltration is attractive due to the almost complete removal of proteins, but recovery and repeatability of Pen were not satisfactory. Therefore, protein precipitation with 10% v/v TCA was chosen for further experiments. The remaining protein content after precipitation is a drawback, but as Pen was clearly separated from the protein residues, this was accepted.

Selectivity

The selectivity of the chromatographic method was examined for a number of pharmaceutically relevant biomolecules: desmopressin, Pen, substance P, PTH (1-34) and insulin (Table 1). The species were all satisfactorily separated by extension of the 38% v/v ACN isocratic part of the gradient after elution of Pen. Chromatograms of the peptides are shown in Fig. 2. It must be noted that the developed method for Pen is not directly applicable to any peptide, as the properties of all peptides are different, the sample pre-treatment must be examined and optimized for each peptide of interest.

Table 1 Peptides used for demonstration of selectivity

Peptide	$M_{ m w}$	No. of amino acids	No. of S	
Desmopressin	1069.22	9	2	
Penetratin	2245.74	16	1	
Substance P	1347.63	11	1	
PTH (1-34)	4117.74	34	2	
Insulin	5807.6	110	6	

Analytical figures of merit

The chromatographic method was validated for Pen using the DRC instrument equipped with a MD system (MD-ICP-DRC-MS) and quantification by post column IDA. Oxygen was applied as reaction gas resulting in measurement of $^{32}S^{16}O^{+}$ at m/z 48. The analytical figures of merit are summarised in Table 2.

Linearity was examined in order to demonstrate a linear response in the applied concentration range. Pen was spiked into 50% v/v human plasma followed by protein precipitation reaching a final concentration of 0.5–15 μM Pen (15–450 μg S $L^{-1}).$ Linearity was established with correlation coefficients of 0.999. Precision was determined at two concentration levels (0.7 μM and 3.5 $\mu M).$ For both concentrations the precision were considered satisfactory.

LOD and LOQ were estimated and absolute LOD of 0.6 pmol S and LOQ of 1.9 pmol S were obtained. The LOD is considerably lower as compared to LODs reported by others using ICP-DRC-MS instruments and oxygen as reaction gas. For proteins, peptides or amino acids in standard solutions, LODs in the range of 2.3-3.1 pmols have been reported. 13,33,34 Only a limited number of studies, where hyphenated ICP-MS methods were used for absolute quantification of S in plasma, has been described. Koellensperger et al. obtained an LOD of 62 pmol S in albumin in human plasma using a two dimensional chromatographic system and ICP-DRC-MS.29 The improved LOD obtained in this study is explained by the application of the MD system. In the MD, the sample is heated and all volatile matrix constituents removed through a semipermeable membrane. As the sample is evaporated in the MD, the total amount of analyte is introduced into the ICP-MS. Using a regular ICP-DRC-MS setup with a spray-chamber and

a concentric nebulizer, only 2% of the sample spray reaches the detector. $^{\rm 35}$

The method was developed for Pen, which is not yet part of any approved drug formulation. The relevant plasma concentration is therefore not defined. For a well-known drug compound as human insulin, maximum plasma concentrations of 200 $\mu IU~mL^{-1}$ (1.2 mM insulin ≈ 7.3 mM S) have been reported after a single dose of 0.1 IU kg $^{-1}.^{36}$ The developed method would be applicable in a similar experiment as the LOD obtained in this study was 0.12 μM Pen (containing 1 S atom). The species dependent LOD will improve with the presence of multiple S atoms in the peptide. If peptides such as desmopressin (2 S atoms) or insulin (6 S atoms) were quantified, the LOD of the species would improve correspondingly.

As no certified standards are available for Pen, the accuracy of the IDA method was determined using desmopressin (Ph. Eur. CRS). Accurate quantification by IDA requires simultaneous interference free detection of two isotopes of S, in this case ³²S and ³⁴S.²⁵ Using oxygen as a reaction gas the interferences are limited by the formation of ³²S¹⁶O⁺ and ³⁴S¹⁶O⁺. Possible spectral interferences include ⁴⁸Ti⁺, ⁴⁸Ca⁺, ³⁶Ar¹²C⁺ and ³¹P¹⁶O¹H⁺ which may affect the ³²S¹⁶O⁺ signal, while ³⁴S¹⁶O⁺ can be interfered by ⁵⁰Ti⁺ if interferences are not removed in the DRC.¹⁵

Giner Martinez-Sierra *et al.* demonstrated applicability of using reaction cell technology and quantification of S (32 S 16 O $^{+}$) by post column isotope dilution in aqueous solutions of methionine with accuracies of 98% and 99%. 22 In our laboratory we have demonstrated accuracy of 98.2% for a certified sulfate standard using ion chromatography with isocratic elution hyphenated to the ICP-DRC-MS and quantification by post column IDA. Thus, the interferences are not considered devastating for the IDA quantification. The influence of matrix constituents from biological samples on IDA quantification is not known.

Application of the MD with ICP-DRC-MS may induce other challenges, as differences in sensitivity between species when using an MD system are well known.^{21,37} In order to obtain accurate quantification by IDA using a MD system, the isotope spike (³⁴S as sulfate) as well as the analyte (desmopressin or Pen) must be stable in the MD system and exhibit similar sensitivity. This was experimentally confirmed and has also

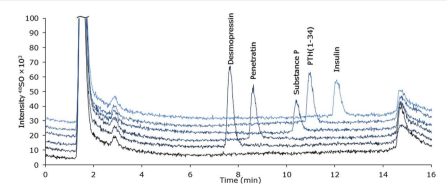


Fig. 2 MD-LC-ICP-DRC-MS chromatograms of S-containing peptides and protein (\sim 10 μ M S) spiked into ultra-filtrated human plasma.

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Analytical figures of merit of RP chromatography of Pen in human plasma using an MD-ICP-DRC-MS instrument equipped with an MD systems

RSD 0.5 μM level	RSD 3.5 μM level	Estimated LOD	Estimated LOQ	Linearity 0.5–15 μM	Accuracy ^a
4.0%	2.3%	0.6 pmol $3.6 \ \mu g \ L^{-1}$	1.9 pmol 12.0 $\mu { m g~L}^{-1}$	R^2 0.999	$102\pm2.7\%$
(n = 5)	(n = 5)	(n = 5)	(n=5)		(n=3)
^a Accuracy determi	ined for desmopressin (Ph	ı. Eur. CRS).			

been reported by Feldmann and colleagues who demonstrated equal and stable signals of sulfate and smaller peptides using

ICP-MS/MS detection

an MD-system.38

The chromatographic method was transferred to an ICP-MS/MS instrument in order to compare sensitivity. As this instrument was equipped with another LC-system, the ACN-gradient was slightly modified in order to move the Pen peak away from front peak tailing. One of the reasons for applying the ICP-MS/MS instrument was that this system can handle larger organic solvent amounts by addition of oxygen (20% in argon) to the plasma and therefore the more complicated MD set-up was avoided. A mass flow chromatogram is shown in Fig. 3. A comparison of the mass flow chromatograms from LC-MD-ICP-DRC-MS (Fig. 1) and LC-ICP-MS/MS (Fig. 3) reveal that the Pen peak is slightly broader using the ICP-DRC-MS, presumably because of the MD system and the different LC-equipment applied.

The main purpose of transferring the developed method to the ICP-MS/MS system was to compare sensitivity with the MD-ICP-DRC-MS set-up. The absolute LOD obtained by the ICP-MS/MS instrument was 0.5 pmol S and the absolute LOQ 1.6 pmol S and thus comparable to the MD-ICP-DRC-MS system.

In theory, interferences are even more efficiently removed in the ICP-MS/MS due to the dual selection of m/z in the first and third quadrupole (Q1 and Q3). When measuring S, m/z 32 and

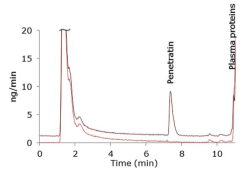


Fig. 3 Mass flow chromatogram from LC-ICP-MS/MS. 50 μM Pen spiked in 50% plasma prior to sample pre-treatment and 50% blank plasma. Dark red: Pen in plasma, light red: blank plasma, offset by -1.2 $ng min^{-1}$.

34 are exclusively selected in Q1 while m/z 48 and 50 interferences are rejected. Only the selected ions reach the reaction cell and react with O₂ forming ³²S¹⁶O⁺ and ³⁴S¹⁶O⁺. Finally, in the O3. m/z 48 and m/z 50 are selected while the interferences 16 O₂ and 16O18O+ are rejected improving the background level.39 Furthermore less noise was observed in the ICP-MS/MS chromatograms as compared to the MD-ICP-DRC-MS chromatograms. The ICP-MS/MS system was therefore expected to improve the LOD obtained by the ICP-DRC-MS. However, the results demonstrated, that the application of the MD system improved the LOD of the DRC-system to a level comparable to the ICP-MS/MS. Application of the MD system with the ICP-MS/ MS might also improve the LOD of this instrument as more sample is introduced and signal suppression from introduction of organic solvent is avoided. However, the MD introduces further complexity to the analytical system and increases the requirements for optimization which is not desirable.

The LOD obtained for the ICP-MS/MS is in accordance with the relatively few studies reported applying ICP-MS/MS hyphenated to a separation technique. Recently, LODs of 0.7 pmol S and 1.3 pmol S were reported for S-containing amino acids by LC-ICP-MS/MS analysis. 34 By capillary LC-ICP-MS/MS, an absolute LOD at fmol level has been reported.16,40 To the authors best knowledge, no studies on quantification of exogenous S in human plasma by LC-ICP-MS/MS have yet been published.

Precision of 8.1% and 5.6% were obtained with the ICP-MS/ MS system for 0.5 µM Pen and 3.5 µM Pen, respectively. This is slightly worse as compared to the MD-ICP-DRC-MS. As observed from the mass flow chromatograms in Fig. 1 and 3, the intensities of the Pen peaks are comparable, but the noise level was visibly improved using the ICP-MS/MS. The superior precision obtained by the MD-ICP-DRC-MS was therefore unexpected but may be explained by the settings of the data collection, as considerable fewer data points were collected for each chromatographic peak during the LC-ICP-MS/MS analysis as compared to the LC-MD-ICP-DRC-MS analysis.

The accuracy obtained by LC-ICP-MS/MS was, however, only 80.2%. The LC method was developed and thoroughly tested using MD-ICP-DRC-MS and subsequently transferred to the ICP-MS/MS system. Analyte adsorption to surfaces is an important issue in peptide analysis and could be a source of the poorer accuracy as the two systems were completely different. Thus, further experiments should be performed to examine this discrepancy between accuracies in the set-ups.

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Conclusion

A reversed phase gradient LC-ICP-DRC-MS method using a membrane desolvation system and post-column IDA was developed for quantification of an S-containing peptide in human plasma. The analytical figures of merit were considered satisfactory for analysis of human plasma samples. The obtained LOD was considerably improved as compared to LODs reported in literature using an ICP-DRC-MS instrument. By transferring the developed method to an ICP-MS/MS system, comparable LOD and improved noise level was achieved, but the accuracy was only 80.2%. Further optimization of the LC-ICP-MS/MS system is thus needed in order to determine the applicability of this method.

LC-ICP-MS analysis represents a promising tool for quantification of peptides and small proteins in complex biological matrices. The possibility of quantifying peptides without introduction of a detection label or need of species specific standards is obviously attractive. The selectivity obtained with this method emphasizes the potential as analytical technique in peptide drug development. Compared to UV-detection, ICP-MS offers improved selectivity as only sulfur-containing compounds are detected and may also be valuable for quantification of peptides with limited UV-absorbance. In addition, LC-ICP-MS may offer an alternative method in cases where ELISA is not easily executed or specific antibodies are unavailable.

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