Development of Mass Spectrometry-Based Methods for Quantitation and Characterization of Protein Drugs: Transferrin as a Model Drug Delivery Vehicle

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Development of Mass Spectrometry-Based Methods for Quantitation and Characterization of Protein Drugs: Transferrin as a Model Drug Delivery Vehicle

A Dissertation Presented
by
SHUNHAI WANG

Submitted to the graduate school of the University of Massachusetts Amherst in partial fulfillment of the requirement for the degree of

DOCTOR OF PHILOSOPHY

SEPTEMBER 2013

Department of Chemistry
DEVELOPMENT OF MASS SPECTROMETRY-BASED METHODS FOR 
QUANTITATION AND CHARACTERIZATION OF PROTEIN DRUGS: TRANSFERRIN AS 
A MODEL DRUG DELIVERY VEHICLE

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by

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DEDICATION

This dissertation is dedicated to my father, Changshou Wang and my mother Bingfang Yao. Their support, encouragement and constant love have sustained me throughout my life.
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At last, a special thank you to all my friends and family members. It is your support and encouragement that I could stay focused on this dissertation work.
ABSTRACT

DEVELOPMENT OF MASS SPECTROMETRY-BASED METHODS FOR QUANTITATION AND CHARACTERIZATION OF PROTEIN DRUGS: TRANSFERRIN AS A MODEL DRUG DELIVERY VEHICLE

SEPTEMBER 2013

SHUNHAI WANG, B.S., PEKING UNIVERSITY
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Directed by: Professor Igor A. Kaltashov

In the last two decades, protein drugs have enjoyed a rapid growth and achieved a tremendous success in treating human diseases. However, the presence of physiological barriers greatly impedes the efficient delivery of such unconventional large molecule drugs, and therefore limits their clinical utility. An elegant way to address this challenge takes advantage of certain endogenous transporter proteins, such as human transferrin (Tf), whose ability to traverse physiological barriers has been extensively exploited. However, methods to investigate Tf-based drug delivery remained insufficient and unsatisfactory until recent development of quantitative mass spectrometry (MS). Hereby, MS-based methods have been developed and validated for quantitation of exogenous Tf in biological fluids. Particularly, different O18-labeling based approaches have been evaluated, modified and developed in this work, in order to achieve the most reliable quantitation. Alternatively, a novel approach based on indium labeling and inductively coupled plasma mass spectrometry (ICP-MS) detection has been developed for sensitive
quantitation of Tf in biological fluids. The second aspect of this dissertation work focuses on the application of MS-based methods for characterization of protein drugs at different levels, ranging from protein identification, covalent structure, conformation, and interaction with physiological partners. Particularly, an O18-labeling assisted approach has been developed to identification of protein deamidation products. This new approach can readily distinguish between the two deamidated isomers. Also, an LC-MS based method has been developed for ranking the susceptibility of protein disulfide bonds to reduction, which could be applied to several disulfide bond-related analyses. Finally, a recently designed growth hormone transferrin fusion protein was studied using MS-based methods, and the molecular basis for its successful oral delivery was revealed.
# TABLE OF CONTENTS

ACKNOWLEDGEMENTS ..............................................................................................................      v

ABSTRACT .................................................................................................................................. vii

LIST OF TABLES ........................................................................................................................... xiii

LIST OF FIGURES ........................................................................................................................... xiv

CHAPTER

1. INTRODUCTION ......................................................................................................................... 1

   1.1 Protein drugs: development and challenges ........................................................................ 1
   1.2 Transferrin (Tf) as a drug delivery vehicle ........................................................................... 2
   1.3 Tf-based drug system under development ............................................................................ 3
   1.4 Challenges for Tf-based drug delivery ................................................................................... 5
   1.5 Methods to investigate Tf-based drug delivery ..................................................................... 6

2. DEVELOPMENT OF MS-BASED METHOD FOR QUANTITATION OF HUMAN TRANSFERRIN IN BIOLOGICAL FLUIDS ............................................................... 12

   2.1 Introduction ......................................................................................................................... 12
       2.1.1 Stable isotope labeled internal standard ......................................................................... 12
       2.1.2 Enrichment of human Tf ................................................................................................ 16
       2.1.3 LC-MS based protein quantitation .................................................................................. 17
   2.2 Results and discussion ......................................................................................................... 18
       2.2.1 MS-based method for quantitation of Tf* in bovine serum ........................................... 18
           2.2.1.1 Ni-NTA Affinity Purification .................................................................................. 18
           2.2.1.2 Preparation of internal standard by acid-catalyzed O$^{18}$ labeling ..................... 19
           2.2.1.3 Discovery of pitfalls in protein quantitation using acid-catalyzed O$^{18}$
labeling ................................................................. 21
  2.2.1.4 Quantitation of Tf* in Bovine Serum ......................... 24
  2.2.2 Cysteine O^{18}-labeling for accurate protein quantitation ..... 25
    2.2.2.1 The workflow of cysteine O^{18}-labeling for protein quantitation ..... 25
    2.2.2.2 Preparation of O^{18}-labeled iodoacetic acid (IAA) ......................... 26
    2.2.2.3 Cysteine O^{18}-labeling: a model protein study ......................... 26
    2.2.2.4 Labeling efficiency of cysteine O^{18}-labeling ............................ 27
    2.2.2.5 Stability of cysteine O^{18}-labeling .................................. 28
    2.2.2.6 The feasibility of using cysteine O^{18}-labeling for absolute protein quantitation ......................................................... 29
    2.2.2.6 Conclusion ........................................................................ 29
  2.2.3 Development of MS-based method for quantitation of Tf* in rat CSF ..... 30
    2.2.3.1 Ni-NTA affinity purification of Tf* from rat CSF ......................... 30
    2.2.3.2 Quantitation of Tf* using cysteine-containing peptides .................. 31
    2.2.3.3 Quantitation results by different reference peptides ..................... 32
    2.2.3.4 Calibration Curve and Method Validation ............................... 33
    1.2.3.5 Conclusion ......................................................................... 35
  2.2.4 Indium labeling and ICP-MS detection based method for Tf quantitation ..... 35
    2.2.4.1 Metal labeling and ICP-MS detection based methods for protein quantitation ............................................................. 35
    2.2.4.2 Indium as a tracer for Tf* quantitation .................................... 36
    2.2.4.3 ICP-MS analysis of serum sample spiked with In_{2}Tf .................... 37
    2.2.4.4 Quantitation methods for indium in serum and CSF .................... 38
  2.3 Conclusion ........................................................................ 39

3. O^{18}-LABELING BASED METHOD FOR IDENTIFICATION OF DEAMIDATION PRODUCTS . 67
  3.1 Introduction ..................................................................... 67
    3.1.1 Traditional methods for identification of deamidation products .......... 68
3.1.2 MS and LC-MS based methods for characterization of deamidation products 68
3.1.3 Tandem MS-based methods for differentiation of deamidation products ...... 70
3.1.4 An O^{18}-labeling approach to differentiation of deamidation products ........... 71
3.2. Results and discussion ........................................................................................................... 72
   3.2.1 O^{18}-labeling assisted approach to differentiation of deamidation products .... 72
   3.2.2 Preparation of O^{18}-labeled deamidation standards .................................................. 73
   3.2.3 A model peptide study ..................................................................................................... 74
   3.2.4 Method validation by ECD-based approach ................................................................. 75
   3.2.5 Characterization of deamidation products from human Tf ......................................... 76
   3.2.6 Elution order of aspartyl- and isoaspartyl-containing peptides in RP-HPLC ....... 78
3.3 Conclusion .................................................................................................................................. 79
4. RANKING THE SUSCEPTIBILITY OF PROTEIN DISULFIDE BONDS TO REDUCTION .... 85
4.1 Introduction ............................................................................................................................... 85
   4.1.1 Screening for possible functional disulfide bonds ......................................................... 86
   4.1.2 Disulfide bonds responsible for covalent aggregation of proteins ............................ 87
   4.1.3 Disulfide-based Protein Modification ............................................................................ 87
   4.1.4 Factors contribute to susceptibility of protein disulfide bonds to reduction ... 88
   4.1.5 Ranking the susceptibility of protein disulfide bonds to Reduction ............................. 89
4.2 Results and discussion ............................................................................................................. 91
   4.2.1 Experiment workflow ...................................................................................................... 91
   4.2.2 Determination of reduction percentage of protein disulfide bonds upon partial
       reduction ................................................................................................................................. 92
   4.2.3 Reduction percentage of disulfide-linked cysteine residues ........................................ 93
   4.2.4 Analysis of peptides containing two cysteine residues ............................................... 93
   4.2.5 Analysis of disulfide bonds in hTf .................................................................................. 95
   4.2.6 Disulfide bonds constraining the lobe movement in hTf .............................................. 97
   4.2.7 Possible disulfide-scrambling site in hTf .................................................................... 98
4.3 Conclusion ............................................................................................................. 100

5. GROWTH HORMONE-TF FUSION PROTEIN FOR SUCCESSFUL ORAL DELIVERY .... 109
   5.1 Introduction ............................................................................................................. 109
   5.2 Results and discussion ............................................................................................ 111
      5.2.1 Protein identification by native ESI MS and a bottom-up approach .......... 111
      5.2.2 Binding ability of GHT1 and GHTx to TfR receptor ........................................ 112
   5.3 Conclusion ............................................................................................................. 115

6. CONCLUSIONS AND FUTURE DIRECTIONS .......................................................... 119

BIBLIOGRAPHY ............................................................................................................. 122
# LIST OF TABLES

<table>
<thead>
<tr>
<th>Table</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Top five proteins identified by Mascot MS/MS ion search engine</td>
<td>40</td>
</tr>
<tr>
<td>2. $^{18}$O-incorporation efficiency in cysteine-containing tryptic peptides of Tf*</td>
<td>41</td>
</tr>
<tr>
<td>3. Validation of quantitation method of Tf* in rat CSF</td>
<td>42</td>
</tr>
<tr>
<td>4. Validation of quantitation method of indium in rat CSF</td>
<td>43</td>
</tr>
<tr>
<td>5. Analysis of disulfide bonds in hTf</td>
<td>101</td>
</tr>
<tr>
<td>6. Top five proteins identified in the first elution peak</td>
<td>116</td>
</tr>
</tbody>
</table>
# LIST OF FIGURES

<table>
<thead>
<tr>
<th>Figure</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-1</td>
<td>Mechanism of targeting tumor cells (left) and central nervous system (right) using TfR-mediated endocytosis/transcytosis</td>
<td>9</td>
</tr>
<tr>
<td>1-2</td>
<td>Tf-based strategies to deliver various therapeutic agents to tumor cells via TfR-mediated endocytosis</td>
<td>10</td>
</tr>
<tr>
<td>1-3</td>
<td>Structure of anticancer agent KP1019</td>
<td>11</td>
</tr>
<tr>
<td>2-1</td>
<td>A general workflow of MS-based method for protein quantitation</td>
<td>44</td>
</tr>
<tr>
<td>2-2</td>
<td>Strategies used for preparation of isotopically labeled internal standards</td>
<td>45</td>
</tr>
<tr>
<td>2-3</td>
<td>Investigate the recovery of Tf* after Ni-affinity purification by SEC.</td>
<td>46</td>
</tr>
<tr>
<td>2-4</td>
<td>Investigation of Ni-affinity purification efficiency on bovine serum sample</td>
<td>47</td>
</tr>
<tr>
<td>2-5</td>
<td>O$^{18}$-labeled peptide DGAGDVAFVK prepared by trypsin-catalyzed O$^{18}$ labeling (blue trace) and acid-catalyzed O$^{18}$ labeling (red trace)</td>
<td>48</td>
</tr>
<tr>
<td>2-6</td>
<td>Quantitation of Tf* in bovine serum using different reference peptides</td>
<td>49</td>
</tr>
<tr>
<td>2-7</td>
<td>Acid-driven deamidation demonstrated by peptide EFQLFSSPHGK from tryptic digestion of Tf*.</td>
<td>50</td>
</tr>
<tr>
<td>2-8</td>
<td>LC-MS/MS analysis of the structure of ionic species $I_0$, $I_8$, and $I_{13}$ related to tryptic fragment EFQLFSSPHGK</td>
<td>51</td>
</tr>
<tr>
<td>2-9</td>
<td>Quantitation of Tf* in bovine serum using different reference peptides</td>
<td>52</td>
</tr>
<tr>
<td>2-10</td>
<td>Calibration curves for quantitation of Tf* in bovine serum using reference peptides DSAHGFLK (left) and YLGEEYVK (right).</td>
<td>53</td>
</tr>
<tr>
<td>2-11</td>
<td>A schematic representation of applying Cys O$^{18}$-labeling for accurate protein quantitation.</td>
<td>54</td>
</tr>
</tbody>
</table>
2-12. ESI MS analysis of O\textsuperscript{18}-labeled IAA in the negative ion mode showing the isotopic distribution of IAA anion ................................................................. 55

2-13. Nano-LC/MS/MS analysis of tryptic peptides derived from a mixture of hTf capped with unlabeled and O\textsuperscript{18}-labeled IAA. ................................................................. 56

2-14. Stability of O\textsuperscript{18}-labeled peptide KPVEYANC*HLAR prepared by Cys O\textsuperscript{18}-labeling under acidic condition ................................................................. 57

2-15. Quantitation curves for Tf\* constructed using tryptic peptides WC*ALSHHER .... 58

2-16. Evaluation of enrichment or removal of proteins from Tf*/rat CSF sample after Ni-affinity purification ................................................................. 59

2-17. Quantitation method using peptide containing two cysteine residues .............. 60

2-18. Quantitation method using peptide containing a single cysteine residue ............. 61

2-19. Comparison of quantitation results by different reference peptides .................. 62

2-20. Calibration curve for quantitation of Tf\* in 50 \mu\textsuperscript{L} of rat CSF .................. 63

2-21. Structure of chelates DTPA (A) and DOTA (B) .............................................. 64

2-22. Full-scan spectrum of serum sample spiked with In\textsubscript{2}Tf acquired by ICP-MS .... 65

2-23. Calibration curves for quantitation of Indium in serum (left) and rat CSF (right) ... 66

3-1. Process of acid-catalyzed deamidation (at 50 °C with 1% TFA) of model peptide EWSVNSVGK in H\textsubscript{2}O\textsuperscript{18} ................................................................. 80

3-2. LC-MS analysis of the O\textsuperscript{18}-labeled acid-catalyzed (a) and base-catalyzed (b) deamidation standards generated using model peptide EWSVNSVGK ....................... 81

3-3. ECD MS/MS analysis of O\textsuperscript{18}-labeled deamidation standards (EWSVNSVGK) ........ 82

3-4. The aspartyl- and isoaspartyl-containing peptides assignment from tryptic digestion of hTf ................................................................. 83
3-5. Examples of peptides exhibited both normal and inverted elution order for their aspartyl- and isoaspartyl-forms ........................................................................................................... 84

4-1. Mass spectra of a doubly charged peptide ion FDEEFSEGC*APGSK alkylated with either IAA-16 (A) or IAA-18 (B) ........................................................................................................................................ 102

4-2. Reduction percentages of cysteine residues from the same disulfide bonds ........ 103

4-3. Mass spectra of three peptides that linked by two disulfide bonds .................... 104

4-4. (a) isotopic distribution of differentially alkylated peptide C*STSSLLEAC*TFR ... 105

4-5. A ribbon representation of the crystal structure of holo-hTf (3v83) ...................... 106

4-6. Reduction percentages of hTf disulfide bonds in both holo-form ....................... 107

4-7. A cartoon representation of a possible disulfide-scrambling site in hTf .............. 108

5-1. Peptide mass fingerprint analysis of high molecule weight fraction from SEC .......... 117

5-2. A. SEC analysis of GHTx, TfR, and their mixture ..................................................... 118
CHAPTER 1

INTRODUCTION

1.1 Protein drugs: development and challenges

High affinity and exquisite specificity towards the therapeutic targets are two of the most important features of an ideal drug. Indeed, those two features have led to a spectacular success of protein-based therapeutics in combating human diseases in the past decades. The first protein drug, biosynthetic “human” insulin was approved to therapeutic use in 1982. Twenty years later, protein-based therapeutics have already become one of the fastest growing segments among the new pharmaceutical products, which representing 44% of all drugs in the development phase and 27% of all drugs in both preclinical and clinical trials [1]. It is expected that protein-based therapeutics will continue to enjoy this growth, and gradually will outpace the traditional small molecule pharmaceuticals. In spite of the success, the clinical utility of many protein drugs is limited due to the inability to efficient and accurate delivery. A significant challenge faced by most protein drugs is the presence of various physiological barriers. For example, intracellular delivery of drugs is very difficult to achieve because the passive entry across the plasma membrane is restricted to molecules that are sufficiently non-polar and less than 500 Da in size [2]. However, none of the protein drugs fall into this category. In addition, when targeting central nervous system (CNS) is necessary, the existence of blood-brain barrier (BBB) amplifies the challenge. Consisting of tightly linked
cerebrovascular endothelial cells, the BBB presents a diffusional barrier between circulating blood and the interstitial space of the brain, making the brain inaccessible to large or hydrophilic molecules by diffusion. In fact, all the protein drugs and more than 98% of small-molecule drugs cannot cross the BBB passively [3]. Additionally, another physiological barrier that presents a formidable challenge in developing orally deliverable protein drugs is the intestinal epithelial barrier. Similar to the BBB, it tightly controls the passage of nutrients and chemicals, as well as protein drugs. In addition to these physiological barriers, the presence of intracellular and extracellular enzymes forms enzymatic barriers that further amplify the challenge of efficient delivery of protein-based therapeutics.

In spite of the difficulty, it has been long recognized that it is essential to develop a molecular mechanism to achieve efficient and accurate delivery of protein drugs [4]. An elegant strategy takes advantage of the ability of certain endogenous proteins to cross physiological barriers via receptor-mediated endocytosis/transcytosis. Those carrier proteins can be utilized as drug delivery vehicle, while the therapeutic proteins can be tethered to them as payload by either conjugation or as recombinant fusion proteins.

1.2 Transferrin as a drug delivery vehicle

Serum transferrin (Tf) is one such carrier protein whose ability to enter cells and cross the physiological barriers via transferrin receptor (TfR) mediated endocytosis/transcytosis has been extensively exploited. Several features have made Tf a
versatile drug delivery vehicle. First, Tf is one of the few proteins that can be internalized by cells via Tfr-mediated endocytosis. The pathway of endocytosis has been extensively studied. Fe₂Tf binds the receptor at the cell surface and both are internalized and transported to endosomes. ATP-dependent proton pumps then force hydrogen ions into the endosome reducing the pH to 5.5, which facilitates a conformational change in Tf, and its subsequent release of iron. Then the Apo-Tf/TfR complex returns to the cell surface where Apo-Tf is displaced by Fe₂Tf. Apo-Tf molecule then circulates until it again comes in contact with free iron, and the cycle is continued [5]. Importantly, overexpression of TfR is normally observed at the surface of rapidly growing malignant cells due to their elevated iron demands [6], providing the basis for targeted delivery of anti-cancer drugs to tumors (Figure 1-1, left panel). Furthermore, since TfR is one of the abundant proteins in the BBB, it is not surprising that Tf has been a focus of extensive efforts to target central nervous system via Tfr-mediated transcytosis (Figure 1-1, right panel). In addition, owing to the wide distribution of the Tfr on the inner lining of GI tract [7], Tf-based protein drugs have also been exploited for oral administration [8], which is often seen as the “holy grail” of protein therapy [9, 10].

1.3 Tf-based drug system under development

A variety of therapeutic agents (chemotherapeutic drugs, protein toxins, radionuclides, liposomes, modified viral particles, and nanoparticles) have been tethered to Tf, and studied for delivery to malignant cells by targeting Tfr on the cell
surface (Figure 1-2) [11]. For example, Hartinger et al. [12] recently reported a Ru-based anticancer agent termed KP1019 (Figure 1-3), which has already entered clinical trials. Taking advantage of its ability to form a covalent adduct with Tf, targeted delivery of this small molecule drug to tumor cells is enhanced. In another example, after conjugating to Tf, artemisinin can be delivered into tumor cells and consequently kill them, by producing free radicals from the reaction with iron [13]. Due to the elevated iron uptake by tumor cells, both toxicity and selectivity of this agent are enhanced. In addition to small molecule drugs, Tf has also been exploited as the delivery vehicle for various protein toxins, such as gelonin [14] and ricin [15]. Notably, CRM 107, a point mutant of diphtheria toxin (DT) with reduced non-specific binding to mammalian cells, has been conjugated to Tf and investigated as a treatment of malignant gliomas [16]. This is the only Tf-cytotoxin conjugate that has reached clinical trials in recent years. Furthermore, high molecular weight compounds, including liposome [17], viruses [18], and nanoparticles [19] have also been conjugated to Tf, and applied for targeted delivery to malignant cells. In addition to Tf, other proteins that bind to TfR have also been exploited as an alternative approach to target tumor cells, including anti-TfR monoclonal antibodies (mAbs) and Tf oligomers [20] (Figure 1-2).

As one of the very few proteins that can traverse the BBB, Tf has been a focus of extensive efforts to deliver therapeutic agents to CNS. In a recent study, human serum albumin nanoparticles which can be loaded with different drugs were conjugated to transferrin, and successful BBB-crossing was observed in an animal mode study [21].
Tf has also been exploited for the design of orally deliverable protein drugs. The wide distribution of TfR on the inner lining of the GI tract [7] provides a possibility of using TfR-mediated transcytosis to overcome the intestinal-epithelial barrier. Applying this strategy, Shen and co-workers [8] recently designed a human growth hormone/Tf fusion protein (GHT) that represented one of the very few examples of a protein drug capable of exhibiting therapeutic response after oral administration. Analysis of this fusion protein in our group [22] revealed the important role of protein aggregation to survive the harsh GI tract environment, and the key role of Tf-moiety to overcome the intestinal-epithelial barrier.

1.4 Challenges for Tf-based drug delivery

Despite the success of conjugating various therapeutic agents to Tf, their efficient delivery is challenging. It is well known that during the receptor-mediated endocytosis, Tf molecule always stays in the endosome while inside the cell, and is quickly recycled back to the cell surface (the entire cycle is only 4 to 5 min) [23]. In order to achieve efficient delivery, the drug molecule must dissociate from Tf and cross the endosomal membrane soon enough to avoid returning to the circulation. It is difficult to meet both requirements. For example, for every 10 million Tf/gelonin cytotoxin conjugates that are recycled, only one molecule of gelonin is actually delivered into the cell in a single cycle [14]. Apparently, the rapid recycling feature of Tf is undesirable for an efficient drug delivery vehicle. Thus, a lot of effort has been made to improve the delivery efficiency by
engineering Tf variants. In a recent study, Kamei et al [24] have applied a mathematical model of the Tf trafficking pathway in order to identify the key factors determining the cellular association of Tf, which has already been demonstrated to be correlated with the effectiveness of drug delivery [14]. In this study, they first identified that reduced iron release rate of holo-Tf can effectively increase its cellular association. By replacing the synergistic carbonate anion with oxalate [24], or using a genetically engineered Tf mutant (K296E/K534A) [23] that both led to higher affinity of Tf to iron and slower iron release, they observed a significant increase in Tf uptake by HeLa cells. In addition, application of Tf oligomers is another strategy to alter the Tf trafficking pathway in order to improve the drug delivery efficiency [20].

1.5 Methods to investigate Tf-based drug delivery

It thus is essential to investigate the ability of Tf and its variants as drug delivery vehicle for both cellular uptake and traversing the physiological barriers. To achieve this goal, accurate and sensitive quantitation of exogenous Tf in complex biological fluids is required. Currently, immunoaffinity-based assays are still the mainstay of detection and quantitation of proteins in complex matrices due to the high sensitivity and specificity. However, in the case of Tf quantitation, this well-established technique is less useful due to the presence of abundant endogenous Tf molecules in most biological fluids. For example, in a model study where quantitation of spiked human Tf (hTf) in bovine serum is required, the application of immunoaffinity-based method could be hampered due to
the cross-reactivity of anti-hTf antibody to abundant bovine Tf (bTf). Indeed, those two molecules share ca. 69% of sequence identity and even higher similarity, and therefore it will be an extremely challenging, time-consuming and resource-intensive effort to produce the antibody that can discriminate between those two molecules.

In fact, current methods used to study Tf-based drug delivery mostly rely on different labeling approaches. For instance, taking advantage of the binding ability of Tf to certain metal ions, radioactive labeling technique can be applied by incorporating radioactive metal ions such as $^{59}$Fe, $^{67}$Ga and $^{111}$In [25] into the Tf binding sites. The latter two are frequently used in medical and diagnostic applications [26-28]. In addition, radioactive-iodine labeling (usually $^{125}$I) is also frequently applied where the aromatic rings of tyrosine side chains are primarily labeled. Using this approach, detection and quantitation of Tf can be achieved with extremely high sensitivity by measuring the radioactivity [24]. However, due to the cost and safety concerns associated with handling radioactive compounds, the routine use of this technique is limited [29]. In addition, in the case of Tf labeling with radioactive metal ions, the quantitation results are less useful after the ion release during the endocytosis/transcytosis. Furthermore, this approach is not suitable for multiplexing so that two or more different forms of exogenous Tf cannot be investigated in a single experiment. Fluorophore labeling is another strategy to discriminate and quantitate exogenous Tf in the context of abundant endogenous Tf. A great advantage provided by this technique is the imaging capability, so that the bio-distribution of Tf can be studied [30]. In addition, applying this
strategy, different forms of exogenous Tf can be achieved simultaneously by using
different chromophores. However, for nearly all the labeling-based strategies,
quantitation of Tf is achieved indirectly by measuring the signal from reporter groups
(radioactive elements or fluorophores), and therefore labeling efficiency and stability
become great concerns.

In the 2nd chapter of this dissertation, novel methods based on mass spectrometry
and stable isotope labeling were developed and validated for quantitation of exogenous
Tf in biological fluids. Particularly, different O18-labeling based approaches have been
evaluated, modified and developed in order to achieve the most reliable quantitation.
Alternatively, a novel approach based on indium labeling and inductively coupled plasma
mass spectrometry (ICP-MS) detection has been developed for sensitive quantitation of
Tf in biological fluids. The 3rd chapter of this dissertation discussed the development of
an O18-labeling assisted approach to identification of protein deamidation products.
This new approach can readily distinguish between the two deamidated isomers. In the
4th chapter of this dissertation work, we discussed the development of an LC-MS based
method for ranking the susceptibility of protein disulfide bonds to reduction, which
could be applied to several disulfide bond-related analyses. Finally, in the 5th chapter, we
studied a recently designed growth hormone transferrin fusion protein, and the
molecular basis for its successful oral delivery was elucidated.
Figure 1-1. Mechanism of targeting tumor cells (left) and central nervous system (right) using TfR-mediated endocytosis/transcytosis. Figure from Kaltashov, I.A., et al., Adv Drug Deliver Rev, 2013, accepted.
Figure 1-2. Tf-based strategies to deliver various therapeutic agents to tumor cells via TfR-mediated endocytosis. Figure is from reference [11].
Figure 1-3. Structure of anticancer agent KP1019.
CHAPTER 2

DEVELOPMENT OF MS-BASED METHOD FOR QUANTITATION OF HUMAN TRANSFERRIN IN BIOLOGICAL FLUIDS

2.1 Introduction

Rapid growth of quantitative proteomics in which mass spectrometry (MS) has played a central role provides a new strategy for protein quantitation in a both sensitive and accurate fashion. By employing differential stable isotope labeling of proteins or peptides in two sample sets, mass differences can be created and recognized by MS analysis. Absolute quantitation of protein can be achieved by comparing the MS signal from the biological/clinical sample with the signal from the spiked internal standard present at a known concentration (Figure 2-1). More recently, a label-free quantitation method by MS has emerged as an alternative approach [31], and gained wide applications in quantitative proteomics. However, the most accurate quantitation is normally achieved using stable isotope labeled internal standards.

2.1.1 Stable isotope labeled internal standard

For accurate quantitation, it is essential to use an isotopically labeled internal standard to compensate for the matrix effects during LC-MS analysis. Based on when and how the isotopes are introduced, the internal standards can be divided into two categories: protein-based internal standard and peptide-based internal standard.
Apparently, for most accurate quantitation, protein-based internal standard is more ideal than peptide-based internal standard, since the former can be spiked into biological samples at a very early stage. As a result, all the downstream sources of quantitation error can be effectively removed (Figure 2-2) [32].

For example, isotopically labeled protein can be produced with relative ease in prokaryotic expression system during cell culture using \(^{15}\)N-enriched medium, which is the earliest possible point for introducing isotopes [33]. However, production of labeled protein in eukaryotic expression system by this approach is significantly more costly and labor-intensive. Instead, stable isotope labeling by amino acid in cell culture (SILAC) approach introduced by Mann and co-workers in 2002 has gained wider popularity [34]. Most frequently, \(^{13}\)C\(_6\)-arginine and \(^{13}\)C\(_6\)-lysine are used in the medium so that all the tryptic fragments of a protein carry at least one labeled amino acid, resulting in a constant mass increment that can be analyzed by MS. Alternatively, protein-based internal standard can also be produced by chemical labeling at the intact protein level. Amine-based labeling strategies (Lys, N-termini) are less favorable due to the frequent application of trypsin during protein proteolysis. As a result, thiol-based labeling strategies are the primary way to introduce isotopes at the intact protein level. For example, the ICAT approach has been widely applied for quantitative analysis of complex protein mixtures, where the thiol groups in protein were derivatized with isotope-coded reagents [35].

In comparison with isotopically labeled protein which is significantly more difficult
to produce, isotopically labeled peptides can be produced with relative ease via a variety of approaches. Indeed, the application of peptide-based internal standard in protein quantitation is much more frequent than the use of protein-based internal standard. Conveniently, isotopically labeled peptide can be synthesized based on a reference peptide from proteolysis of the protein of interest. Thus, absolute quantitation of the protein can be achieved by comparing the signal intensity from unlabeled (biological sample) and labeled peptide (internal standard). This approach was first introduced by Gygi and co-workers in 2003 termed AQUA [36], and it has been the most widely applied technique for protein quantitation. In addition, isotopes can also be introduced to peptides by chemical labeling. Particularly, amine groups (Lys, N-termini) are primarily targeted and can be derivatized with various labeling reagents, including iTRAQ [37], dimethyl labels [38] and tandem mass tag (TMT) reagents [39]. All those approaches have gained wide acceptance in the field of quantitative proteomics. Alternatively, enzyme-catalyzed $^{18}O$ labeling is another elegant way to introduce isotope labeling to proteolytic fragments. Yao and co-workers [40] first applied this technique in a comparative proteomics study. Despite the ease of use, application of this approach for accurate protein quantitation is hampered by the back exchange event during LC-MS analysis due to the residual protease activity in the system, although a lot of efforts have been devoted to reduce this effect [41-44]. In addition, an intrinsic disadvantage of enzyme-catalyzed $^{18}O$ labeling is the peptide-dependant $^{18}O$-incorporation efficiency, which further complicates the data analysis [45]. More recently, a modified approach
termed acid-catalyzed O\textsuperscript{18} labeling was introduced, which takes advantage of free exchange of carboxyl oxygens in peptides with solvent at low pH [46]. Despite the prolonged preparation time, this approach provides sufficient mass shift after labeling due to additional O\textsuperscript{18}-incorporation sites on the acidic residues. Furthermore, since no enzyme is employed during the labeling, back exchange rates could be significantly reduced. However, despite this variety of techniques, peptide-based internal standard is not ideal for protein quantitation, since it can only be spiked into biological samples after proteolysis, which inevitably introduces sources of quantitation error. In particular, digestion efficiency becomes a significant variable that can greatly affect the quantitation results. In fact, complete digestion of target protein is normally required for reliable quantitation, although it is frequently difficult to achieve.

Isotope effect is another important factor that needs to be taken into consideration during the selection of isotopes. Although it is generally valid that isotope labeling does not alter the physicochemical properties of a peptide, retention time shifts during RP-HPLC separation have been reported for deuterated peptides [47]. Obviously, identical retention time between unlabeled and isotope-labeled peptides is critically important for accurate quantitation, to ensure the identical solvent composition and ion suppression from co-eluting species, resulting in identical ionization efficiency of unlabeled and labeled peptides. Thus, instead of deuterium, isotope labels such as C\textsuperscript{13}, N\textsuperscript{15}, or O\textsuperscript{15} have been more widely applied, due to their negligible isotope effect.
2.1.2 Enrichment of human Tf

Since a protein drug in biological fluids typically represents a very small fraction of the total proteome, its detection by MS or LC-MS can be hampered by both insufficient signal intensity (ion suppression) and interference caused by co-eluting components of similar mass. The most straightforward way to overcome this difficulty is to reduce sample complexity by either removing the most abundant endogenous proteins or enriching the target protein via affinity-based purifications. For example, up to 21 high abundant proteins can be removed from plasma sample with relative ease using commercially available immunodepletion columns [48, 49]. Alternatively, if an appropriate antibody is available, the protein analyte can be simply enriched in one-step applying immunocapture/immunoprecipitation [50]. However, both those methods are less useful for enrichment of exogenous Tf (e.g., human Tf) from biological fluids due to cross-reactivity from abundant endogenous Tf (e.g., bovine Tf or rat Tf) or vice versa.

Instead, immobilized metal ion affinity chromatography [51] is employed in this work, which takes advantage of the coordinate binding of amino acids, particularly histidine, to metal ions, such as cobalt, nickel and copper [52]. Particularly, nitrilotriacetic acid (NTA) is applied as chelating ligand to retain the metal ions on the solid surface. Due to the strong binding ability of NTA to metal ions, a wide variety of binding/washing conditions (particularly stringent washing conditions) can be employed in order to enrich low abundance target protein from complex matrices. After
introducing a 6×His-tag to the N-terminus of hTf (referred to as Tf*), enrichment of Tf* can be achieved by applying Ni-NTA affinity purification.

2.1.3 LC-MS based protein quantitation

LC-MS based method using conventional separation conditions (4.6-mm i.d. column and 1.0 mL/min flow rate or 2.1-mm i.d. column and 0.2 mL/min flow rate) has been a mainstay for quantitation of small molecular compounds from biological fluids due to the high-throughput and robustness. However, for detection and quantitation of low levels of therapeutic proteins in complex matrices, LC-MS analysis with standard flow rate separation normally cannot meet the requirement of high sensitivity. Instead, capillary flow (1-10 μL/min) and nano flow (ca. 200 nL/min) are frequently applied in order to enhance the sensitivity by reducing dimensions of the system [53, 54]. In our study, a nanoLC-MS system interfaced by a nano ESI spray source is applied to provide necessary sensitivity for the quantitation of exogenous Tf* in biological fluids. To cope with the nano flow, digestion samples (ca. 50-200 μL) are first preconcentrated using a micro-trapping column at a relatively high flow rate (40 μL/min), and subsequently eluted to the analytical column at a low flow rate (200 nL/min). A control experiment showed that amount as low as 8 ng of Tf* can be confidently detected using this setup.
2.2 Results and discussion

2.2.1 MS-based method for quantitation of Tf* in bovine serum

In a model study, method consisting of Ni-affinity purification for protein enrichment, acid-catalyzed O\textsuperscript{18} labeling for preparation of internal standard and nanoLC-MS detection was developed for quantitation of low levels of Tf* in bovine serum.

2.2.1.1 Ni-NTA affinity purification

In order to improve the sensitivity of the method, Ni-NTA affinity purification was employed to extract low abundance Tf* from bovine serum samples. Briefly, 50 μL of Ni-NTA resin slurry was washed by 1 mL of binding buffer, and then transferred to the sample solution in a 1.5 mL Eppendorf tube. The binding was conducted at room temperature for 90 min with gentle shaking. Subsequently, the resin was washed by 500 μL of washing buffer (50 mM of potassium phosphate, 20 mM of imidazole, 6 M of guanidine hydrochloride, 1 M of sodium chloride and 10% v/v glycerol, pH 8.0) for two times in order to remove non-specific bound proteins. Finally, Tf* was recovered by adding 100 μL of elution buffer (50 mM of potassium phosphate, 300 mM of imidazole, pH 8.0) to the resin, and subsequently collecting the supernatant. The recovery of Tf* after Ni-affinity purification was investigated using size exclusion chromatography (SEC), and calculated to be ca. 80% based on the UV signal of Tf* peak (Figure 2-3).
After Ni-affinity purification, the complexity of the serum sample was significantly reduced, which can be clearly observed in SDS-PAGE analysis (Figure 2-4). By comparing the protein bands from the flow-through fraction and the purification fraction, it was clear that majority of serum proteins were efficiently removed after Ni-affinity purification.

The enrichment of Tf* from the bovine serum sample by Ni-affinity purification was further studied using a “bottom-up” approach. Briefly, bovine serum samples spiked with Tf* (1 µg of Tf* in 1 mL of bovine serum) were treated with or without Ni-affinity purification. After tryptic digestion, proteolytic peptides were subjected to LC-MS/MS analysis, and the results were searched against SwissProt protein database by Mascot MS/MS ions search engine [55] to identify the precursor protein(s). The top five protein hits in each sample were listed in Table 1. Without Ni-affinity purification, Tf* cannot be successfully identified due to the presence of abundant endogenous proteins (complete list was not shown in Table 1). In contrast, with the purification step Tf* can be confidently identified, emphasizing the necessity of applying Ni-affinity purification for detection of low levels of Tf* in bovine serum samples.

2.2.1.2 Preparation of internal standard by acid-catalyzed O\(^{18}\) labeling

O\(^{18}\)-labeling was selected for preparation of stable isotope labeled internal standard in our study due to the minimal isotope effect and the ease of use. Two classic approaches to introduce O\(^{18}\)-labeling have been reported, termed “enzyme-catalyzed
O$^{18}$ labeling” (Scheme 1) and “acid-catalyzed O$^{18}$ labeling” respectively (Scheme 2). Enzyme-catalyzed O$^{18}$-labeling is widely applied in quantitative proteomics. It takes advantage of the reversible binding between enzyme (mostly trypsin) and the proteolytic peptides in H$_2$O$^{18}$ so that both the C-terminal carboxylic oxygens can be exchanged to O$^{18}$, leading to a mass increment of 4 Da for each peptide. Despite the ease of use, this approach is limited by some intrinsic problems. First, complete labeling is difficult to achieve, resulting in overlap of isotopic peaks with unlabeled peptides. In addition, peptide-dependant labeling efficiency greatly complicates the data analysis [45]. Furthermore, back exchange of O$^{18}$-labeled peptides occurring in RP-HPLC separation can be significant due to the catalysis of residual trypsin. As a result, removal or inactivation of trypsin after labeling is normally required. Most of those problems can be effectively eliminated by increasing the mass shift after labeling. Indeed, the new approach, acid-catalyzed O$^{18}$ labeling, explored the possibility of labeling all the carboxylic acid groups (C-terminus, acidic residues) in order to increase the mass shift [46]. By incubating the tryptic fragments in H$_2$O$^{18}$ under acidic condition, all the carboxyl oxygens can be exchanged to O$^{18}$ upon equilibrium. Owing to the high frequency of acidic residues in the human proteome, nearly half of the tryptic fragments contain at least one acidic residue, providing a mass shift of 8 Da or more, which is sufficient to completely separate isotopic clusters from unlabeled and labeled peptides [46]. In addition, as no enzyme is required during the labeling, the back exchange rate of labeled peptides is also significantly reduced. However, this approach requires a considerably
longer preparation time of the internal standard (typically over 10 days), thus careful planning of the experiments is generally suggested. Both approaches were investigated in our study.

Scheme 1

Scheme 2

Figure 2-5 represents the labeling results of a tryptic fragment from Tf* by both trypsin-catalyzed O^{18} labeling (blue trace) and acid-catalyzed O^{18} labeling (red trace), in comparison with unlabeled peptide (black trace). Clearly, sufficient mass increment provided by acid-catalyzed O^{18}-labeling greatly simplifies the data analysis. Thus, this approach was selected to prepare an O^{18}-labeled internal standard for quantitation of Tf* in bovine serum.

2.2.1.3 Discovery of pitfalls in protein quantitation using acid-catalyzed O^{18} labeling

The quantitation method for Tf* in bovine serum was developed using acid-catalyzed O^{18} labeling. However, pitfalls of this approach were identified during the initial method development [56]. As shown in Figure 2-6, quantitation of Tf* in bovine serum was achieved using the identical amount of O^{18}-labeled internal standard, and
results were generated using several reference peptides (black bars). Surprisingly, a large
discrepancy in quantitation results was observed among different reference peptides.
Furthermore, even though the unlabeled to labeled ratio should not exceed 1.0
(preparation of internal standard does not expose Tf* to the affinity purification step,
thereby eliminating unavoidable protein loss), the apparent ratio for one of the peptides
shown in Figure 2-6 exceeded that level. Close examination of the data presented in
Figure 2-6 revealed a common structural feature among all peptides that yielded higher
quantitation levels of Tf * in bovine serum: each of these peptides has at least one
amino acid residue bearing an amide group on its side chain (Asn, Gln, and
carboxamidomethylated cysteine, Cys*). Because of the susceptibility of amides to acid
hydrolysis, acid-driven deamidation was suspected as a likely cause of the discrepancy in
quantitation results (Scheme 3). Indeed, if the fraction of deamidated peptides (from
O^{18}-labeled internal standard) was neglected during the ratio calculation, the apparent
level of Tf* in bovine serum would necessarily be inflated. In addition, acid-driven
deamidation in H_{2}O^{18} should add another 5 Da to the labeled peptides due to additional
incorporation of two O^{18} atoms (Scheme 3).

Scheme 3

An example of this phenomenon is shown in Figure 2-7. For peptide EFQLFSSPHGK,
a maximum mass increment of 8 Da was expected after acid-catalyzed O$^{18}$ labeling, due to O$^{18}$ incorporation at both the glutamic acid residue and the C-terminus. Simultaneously, acid-driven deamidation at glutamine residue led to additional incorporation of two O$^{18}$ atoms, resulting in a total mass increment of 13 Da. LC-MS analysis revealed the consistent results. In addition to unlabeled peptide ($I_0$) and O$^{18}$-labeled peptide ($I_8$) co-eluted at the first peak, another ionic species eluted slightly later with maximum mass increment of 13 Da ($I_{13}$) was also observed.

The deamidation site was further confirmed by tandem MS experiment (Figure 2-8). Collisional activation of the doubly charged ion corresponding to this peptide yielded a series of single charged y-ions. The fragment ion mass spectrum of the $I_{13}$ species did not show any additional mass shifts compared to that of $I_8$ for smaller fragments (up to y$_8$), suggesting that no modification was present in the last eight residues. However, an additional mass shift of 5 Da was observed for fragment ion y$_9$, clearly suggesting that deamidation of Glu followed by incorporation of two O$^{18}$ atoms into the newly formed carboxylic acid group. Similar analyses were carried out for other amide-containing peptides, and in all cases extensive deamidation was observed in internal standard prepared by acid-catalyzed O$^{18}$-labeling (data not shown). The extent of deamidation varied among different peptides, but on average it was within 15-30% of the total peptide population following 2 weeks of acid-catalyzed labeling (the typical amount of time needed to achieve a near-complete labeling of the acidic side chains).

Clearly, the loss of a significant fraction of an internal standard due to deamidation
led to an artificial increase of the apparent level of Tf* in bovine serum. An attempt to remediate this problem by including all deamidated species in the ratio calculation led to a significant decrease in the discrepancy of the quantitation results (Figure 2-9), but did not eliminate it completely. One important reason for the remaining discrepancy is the inadvertent change of the ionization efficiency of deamidated peptides compared to their intact counterparts (which is caused not only by the alteration of their covalent structure but also by a change in the solvent composition at different retention time and a different matrix/ion suppression effect from coeluting species). Another reason is the existence of a matrix effect during the tryptic digestion of serum sample and the internal standard, since the latter was prepared in the absence of bovine serum. However, this matrix effect on digestion efficiency is a universal problem for nearly all the quantitation methods using peptide-based internal standard, which can only be eliminated by switching to protein-based internal standard.

2.2.1.4 Quantitation of Tf* in bovine serum

As demonstrated in this study, selection of reference peptides is critically important in this approach. Apparently, amide-containing peptides should be excluded due to the susceptibility to acid-driven deamidation. In addition, for methods using peptide-based internal standard, complete digestion is required to minimize the matrix effect on digestion efficiency. Thus, only peptides without any missing cleavage can be used as reference peptides. And finally, the use of high abundance peptides can effectively
improve method sensitivity. Using two such peptides, calibration curves were generated for quantitation of Tf* in bovine serum (Figure 2-10). Excellent linearity was achieved for over two orders of magnitude in each case. Concentrations as low as 0.2 μg/mL of Tf* can be confidently quantitated using this approach, which is ca. 4,000 times lower than the concentration of endogenous bovine Tf.

2.2.2 Cysteine O\(^{18}\)-labeling for accurate protein quantitation

As discussed earlier and demonstrated in our first study, the application of peptide-based internal standard is not ideal for accurate protein quantitation. Particularly, the matrix effect on digestion efficiency can greatly compromise the quantitation results, since the digestion of biological sample and internal standard are performed separately. A logical solution for this problem is the use of a protein-based internal standard so that the addition of the internal standard to the biological sample can occur prior to digestion. However, classic approaches (enzyme-catalyzed O\(^{18}\)-labeling and acid-catalyzed O\(^{18}\)-labeling) only introduce the isotopes at the peptide level, thus are both suffering from this disadvantage. Hereby, we developed a new strategy of O\(^{18}\)-labeling at the intact protein level, termed cysteine O\(^{18}\)-labeling, in which the isotope incorporation occurs during the alkylation of cysteine residues. The feasibility of the new method for accurate protein quantitation has been demonstrated using Tf* as a model.

2.2.2.1 The workflow of cystein O\(^{18}\)-labeling for protein quantitation
The workflow of using cysteine $^{18}$O-labeling for protein quantitation is represented in Figure 2-11. Briefly, after reduction of disulfide bonds in both biological samples and internal standard, the cysteine residues were alkylated with either $^{16}$O-labeled or $^{18}$O-labeled iodoacetic acid. Subsequently, an internal standard of known quantity was spiked into the biological samples, followed by tryptic digestion of the mixture. Finally, the protein quantitation was achieved by comparing signal intensities of differentially alkylated products from a cysteine-containing peptide.

2.2.2.2 Preparation of $^{18}$O-labeled iodoacetic acid (IAA)

The $^{18}$O-labeled IAA was prepared by exchanging its oxygen atoms in carboxylic group with H$_2$O$^{18}$ at low pH. The labeling results are shown in Figure 2-12, where the isotopic distribution of IAA after a 24 hour incubation at 50 °C, in the presence of 1% TFA, shows > 85% exchange of both carboxylic oxygen atoms and > 98% exchange of at least one oxygen atom. These numbers yield the overall extent of $^{18}$O incorporation as 93%, whereas the theoretical value is calculated to be 93.9% after taking into account the presence of residual $^{16}$O in the reaction mixture. This consistent result suggested that the $^{18}$O exchange between IAA and water had reached the equilibrium.

2.2.2.3 Cysteine $^{18}$O-labeling: a model protein study

The alkylation of cysteine residues with $^{18}$O-labeled IAA was first studied using Tf*, which has 38 cysteine residues as a model protein. Figure 2-13 presents an LC-MS/MS analysis of the mixture of tryptic fragments from two batches of Tf* capped by
$O^{16}$-labeled and $O^{18}$-labeled IAA, respectively. The mass spectrum (Figure 2-13B) of a tryptic peptide KPVEYANC*HLAR indicates a maximum mass shift of 4 Da between unlabeled ($I_0$) and labeled ($I_4$) peptides. Extracted ion chromatograms plotted for their most abundant isotopic peaks, $I_0$ and $I_4$ (Figure 2-13A), clearly show identical elution times, which is critically important for reliable quantitation. The tandem MS data (Figure 2-13C and D) suggest that the presence of IAA moiety on a peptide ion does not interfere with its identification using CAD (contrary to ICAT labeling [57]). Furthermore, the isotopic distributions of $y^{4+}$ and $y^{5+}$ ions provide a clear indication as to the incorporation site of $O^{18}$ atoms within the peptide (cysteine residue).

2.2.2.4 Labeling efficiency of cysteine $O^{18}$-labeling

In this approach, $O^{18}$ labeling is introduced by alkylation reagent, thus the labeling efficiency of all the cysteine-containing tryptic fragments should be consistent. This assumption was demonstrated by calculating the labeling efficiency of all the detectable cysteine-containing peptides from tryptic digestion of Tf* (capped by $O^{18}$-labeled IAA). As shown in Table 2, for peptides containing one cysteine residue, the full labeling products (with both two $O^{18}$ atoms incorporated) made up to 87%–90%, consistent with the labeling results of IAA (vide supra). Unlike enzyme-catalyzed $O^{18}$ labeling which has peptide-specific $O^{18}$ incorporation rate, the use of $O^{18}$-labeled IAA generated almost identical labeling efficiency among all the cysteine-containing peptides. This feature greatly simplifies the data analysis, particularly in the case of insufficient mass shift.
(incomplete separation of isotopic distributions of labeled and unlabeled peptides). In fact, the ratio between unlabeled and labeled peptides can be easily estimated by the following equation:

$$\text{peptide ratio} = \frac{I_1}{I_1 \times (1 + \alpha)}$$

Where $\alpha$ ($\alpha=0.14$ in this work) is a fraction of partially labeled peptides (only one O$^{18}$ atom incorporated); and $I_1$ and $I_5$ represent the observed relative intensities of the second isotopic peaks for the unlabeled peptides and fully O$^{18}$-labeled peptides, respectively.

2.2.2.5 Stability of cysteine O$^{18}$-labeling

The back exchange in trypsin-catalyzed O$^{18}$ labeling is primarily contributed from the residual trypsin activity in the labeling solution. However, in this new approach, the O$^{18}$-incorporation sites (carboxymethylated Cys) cannot be recognized by trypsin, thus the back exchange rate is expected to be extremely low (without the catalysis of trypsin). To validate this assumption, stability of the cysteine O$^{18}$-labeled peptide KPVEEYANC*HLAR was investigated at both RP-HPLC conditions (0.1% FA) and near neutral pH (pH 8, for tryptic digestion) (Figure 2-14). The relative abundance of the isotopic peak $I_2$ as percentage of the total ionic signal was calculated and monitored over time. No noticeable back exchange was observed under RP-HPLC conditions for at least 8 hours, or under near neutral pH for at least 6 days. The excellent stability of O$^{18}$-labeling under near neutral pH allows overnight tryptic digestion and any other
sample processing step (as long as it does not require a long period of time in extreme pH conditions) after spiking the internal standard. In addition, the good stability of O\textsuperscript{18}-labeling under RP-HPLC conditions allows us to apply a long LC gradient separation for complex samples.

2.2.2.6 The feasibility of using cysteine O\textsuperscript{18}-labeling for absolute protein quantitation

To demonstrate the feasibility of using cysteine O\textsuperscript{18}-labeling for protein quantitation, we generated Tf\textsuperscript{*} samples capped with unlabeled and O\textsuperscript{18}-labeled IAA at various ratios covering three orders of magnitude, and performed nano-LC/MS analysis. Successful quantitation was achieved using two cysteine-containing tryptic peptides from Tf\textsuperscript{*} respectively (Figure 2-15). High correlation of theoretical and experimentally observed ratios was achieved ($R^2 > 0.99$). Two reference peptides both showed good linearity in up to three orders of magnitude. Furthermore, the quantitation results generated by these two peptides were highly consistent, highlighting the reliability of quantitation using protein-based internal standard.

2.2.2.6 Conclusion

Obviously this new method only targets cysteine-containing tryptic peptides and therefore it cannot be applied to quantitate proteins without cysteine residue. Nevertheless, even the most conservative estimates of the fraction of completely cysteine-free proteins in the proteomes of higher organisms put this number as low as 8%, while the SWISS-Prot database puts the number of human proteins that have at

29
least one cysteine residue in the sequence as high as 97%. As a result, this method can still be widely applied for tasks ranging from pharmacokinetics studies to quality control of biopharmaceutical products, where accurate protein quantitation is required.

2.2.3 Development of MS-based method for quantitation of Tf* in rat CSF

In spite of the success achieved by protein drugs in treating human diseases in the last decade, development of protein drugs targeting CNS has been notably slow, primarily due to the presence of blood-brain barrier (BBB). As one of the very few proteins that can penetrate the BBB via receptor-mediated transcytosis, Tf has been a focus of extensive efforts to achieve delivery of therapeutic agents to CNS [58]. In order to investigate the BBB-traversing ability of Tf, quantitation method for low levels of exogenous Tf in CSF samples is required. As discussed in section 1.1.5, immunoaffinity-based assays are less useful for this task due to the presence of abundant endogenous Tf in CSF. Although radioactive labeling based approaches provide extremely high sensitivity, protein quantitation is achieved indirectly by measuring the radioactivity, and therefore might be less accurate and reliable. Hereby, we developed an LC-MS based method for accurate and sensitive quantitation of Tf* in rat CSF, where the newly developed cysteine O^{18}-labeling approach was applied to produce internal standard.

2.2.3.1 Ni-NTA affinity purification of Tf* from rat CSF

Although CSF presents a significantly less complex system compared to plasma
(contains approximately 0.3% plasma proteins [59]), an enrichment step for exogenous Tf* is still required. Indeed, a similar “bottom-up” analysis (as discussed in section 1.2.1.1) of 1 µg of Tf* in 50 µL of rat CSF sample demonstrated the necessity of applying Ni-NTA affinity purification prior to LC-MS detection (data not shown). In addition, a comparative analysis of the flow-through fraction and purification fraction from Ni-affinity purification was also conducted to investigate the enrichment of Tf* and removal of endogenous proteins. Briefly, control sample with 1 µg of Tf* in 20 µL of rat CSF was treated with Ni-affinity purification. Cysteine residues from proteins in flow-through fraction and purification fraction were capped by O\(^{16}\)-labeled iodoacetic acid (IAA-16) and O\(^{18}\)-labeled iodoacetic acid (IAA-18) respectively. Subsequently, after combination of the two fractions, the sample was digested by trypsin, and subjected to LC-MS/MS analysis. By comparing the relative intensities of differentially alkylated peptides (ion \(I_0\) from flow-through fraction and ion \(I_4\) from purification fraction) from certain protein, the enrichment or removal of this protein can be evaluated. Clearly, after Ni-affinity purification, Tf* was greatly enriched to the elution fraction; in contrast, two of the most abundant endogenous proteins in CSF, rat Tf and rat albumin, were effectively removed (Figure 2-16).

2.2.3.2 Quantitation of Tf* using cysteine-containing peptides

\(O^{18}\)-labeled Tf* was used as internal standard for quantitation, which was prepared by alkylation of cysteine residues with IAA-18. Thus, only cysteine-containing peptides
could be used as reference peptides. Figure 2-18 represents an example of using a tryptic peptide with two cysteine residues for protein quantitation. Since the isotopic distributions from unlabeled and labeled peptides are completely separated due to sufficient mass shift (with a maximum of 8 Da), an m/z range (the grey area shown in Figure 2-17) covering all the isotopic peaks from either unlabeled or labeled peptide is extracted. Subsequently, quantitation can be achieved using a simple equation (Figure 2-17) with peak areas derived from extracted ion chromatograms (XICs).

In another case, protein quantitation can also be achieved using a reference peptide containing a single cysteine residue (Figure 2-18). Although isotopic overlapping between unlabeled and labeled peptides can be observed due to incomplete labeling, the interference can be simply corrected by introducing a constant (α) that represents the fraction of partially labeled peptide (with only one O\textsuperscript{18} atom incorporated). To further minimize the interference, the second isotopic peak of either unlabeled peptide (I\textsubscript{1}) or fully O\textsuperscript{18}-labeled peptide (I\textsubscript{3}) are extracted, and the peak areas under XICs are used for the ratio calculation (Figure 2-18).

2.2.3.3 Quantitation results by different reference peptides

A great advantage of using protein-based internal standard is to eliminate the discrepancies in digestion efficiency between biological sample and internal standard. This assumption was demonstrated by comparing quantitation results generated by different peptides, including those with missing cleavages. Triplicate samples containing
1.6 μg of Tf* in 50 μL of rat CSF were spiked with 0.4 μg of internal standard, and analyzed by this method. Ten cysteine-containing peptides were used for quantitation, and the results were normalized against the average value shown in percentage (Figure 2-19). Clearly, nearly all the quantitation results were well within ±15% range regardless of the peptide used. Importantly, even peptides with missed cleavages showed highly consistent quantitation results. Those observations indicated that the discrepancies in digestion efficiency were effectively reduced by using the protein-based internal standard. Furthermore, the feasibility of using peptides with missing cleavages for protein quantitation greatly expanded the selection of reference peptides. In fact, peptide DLLFRDDTVC*LAK was selected as a reference peptide in this study since it was unique to Tf* (in comparison with rat Tf) and could be consistently detected at low concentration, although it did contain a missed cleavage.

2.2.3.4 Calibration curve and method validation

Using peptide DLLFRDDTVC*LAK as the reference peptide, a linear calibration curve was plotted using measured concentration of Tf* versus the corresponding theoretical spiked concentration (20, 40, 100, 200, 800, 1600, and 3200 ng of Tf* in 50 μL of rat CSF). A 1/X² weighted regression was applied. Triplicate samples were measured for each data point, and the standard deviation was presented as the error bar. As shown in Figure 2-20, excellent linearity was achieved for more than two orders of magnitude.

**Accuracy and precision.** The accuracy and precision of this method were further
validated using QC samples prepared with concentrations in the low, medium and high parts of the calibration curve. As shown in Table 3, both precision (% CV) and accuracy at all quality control levels were well within 15%, indicating that the method is both precise and accurate.

**Selectivity and lower limit of quantitation (LLOQ).** The uniqueness of the reference peptide DLLFRDDTVC*LAK for Tf* was confirmed by a protein database search against UniProtKB/Swiss-Prot using the Protein BLAST search engine [60]. In addition, blank samples containing 50 µL of rat CSF were analyzed in triplicates by this method. No signal was observed at the retention time of the reference peptide. During the method development, the lower limit of quantitation of this method was estimated to be 20 ng of Tf* in 50 µL of rat CSF. Measurement of QC samples with this concentration demonstrated that both good precision (% CV) and accuracy (within 20%) can be achieved at LLOQ level (Table 3).

**Matrix effect.** It is well known that the total protein level in rat CSF can vary significantly from individual to individual [61]. Thus, in order to apply this method in animal studies, it is critically important to investigate the matrix effect. QC samples containing 500 ng of Tf* in 100 µL of rat CSF (the calibration standards were prepared using 50 µL of rat CSF) were analyzed in triplicates, and the results were compared with the value derived from the calibration curve. As shown in Table 3, both accuracy and precision of the measurements were well within 15% range, suggesting negligible matrix effect within two times of complexity.
1.2.3.5 Conclusion

Applying cysteine $^{18}$O-labeling, an LC-MS based method was developed and validated for quantitation of Tf* in rat CSF. The newly developed labeling strategy allows the preparation of a protein-based internal standard, which proved to be superior for MS-based quantitation. Excellent linearity, accuracy and precision were achieved using this method. In addition, minimal matrix effect was found after increasing the sample complexity by two times, suggesting the feasibility of applying this method in real animal studies. Finally, concentrations as low as 10 ng of Tf* in 50 μL of rat CSF can be confidently quantitated, providing necessary sensitivity for quantitation of low levels of Tf* in rat CSF after intravenous injection.

2.2.4 Indium labeling and ICP-MS detection based method for Tf quantitation

In addition to LC-MS based approach, we also explored the possibility of using inductively coupled plasma mass spectrometry (ICP-MS) for quantitation of Tf* in biological fluids. Using indium as a tracer, selective and sensitive quantitation of Tf* can be achieved by detection of indium by ICP-MS. This work was done in collaboration with Hanwei Zhao in our lab. Hanwei prepared the Tf bound with different metal ions, and investigated their stability and binding ability to TfR. I then acquired the ICP-MS spectra of $^{115}$In-Tf in serum samples, and developed and validated the quantitation method for indium in serum and rat CSF.

2.2.4.1 Metal labeling and ICP-MS detection based methods for protein quantitation
ICP-MS is an analytical tool widely used for elemental analysis, which consists of a high-temperature plasma for sample ionization and MS for ion separation and detection. During the last decade, ICP-MS has greatly expanded its applications into a wide variety of areas, particularly in the field of bioanalysis [62]. For example, it was reported that ICP-MS based methods have been applied for protein quantitation using a metal labeling strategy. Frequently, labeling of the protein with rare-earth metals is preferred, which allows sensitive quantitation due to their high ionization efficiency and extremely low background (absence of rare-earth elements in biological samples and no spectroscopic interferences). Figure 2-21 shows the structure of two metal chelates that are most frequently employed for protein derivatization. Both chelates DTPA [63] and DOTA [64] can form strong complexes with multiple lanthanides (e.g., log $K$ 22.5 for Gd[DTPA] and log $K$ 24.7 for Gd[DOTA] [65]), providing a possibility to achieve multiplexing. However, protein labeling efficiency with those metal chelates is generally low due to their relatively large size. In addition, extensive modification of a protein by metal chelates can significantly alter its biological activity or function, and therefore such studies have not explored the possibility of using metal-coded protein drugs for pharmacokinetic studies.

2.2.4.2 Indium as a tracer for Tf* quantitation

Instead of using metal chelates, we took advantage of the two metal binding sites in Tf. It is well known that Tf can bind to several other metal ions in addition to iron,
including aluminum, gallium, indium, and thallium [25]. The use of $^{67}\text{Ga}$- and $^{111}\text{In}$-bound 
Tf has even gained widespread medical and radiodiagnostic applications [26-28]. In our 
study, indium was selected as the tracer due to its high binding affinity to Tf. High 
resolution MS analysis by Hanwei Zhao in our lab could confidently discriminate apo-Tf 
and Tf bound with different metal ions based on accurate masses. MS analysis 
confirmed that indium binds to Tf with a constant ratio of 2:1, which provides an easy 
conversion between indium concentration and Tf concentration. In addition, 
indium-bound Tf was observed to be stable in the presence of ferric ions or other 
proteins (e.g. albumin), demonstrating the feasibility of using indium as a tracer for Tf in 
the circulation (data now shown). Furthermore, our study also revealed that 
indium-bound Tf had similar binding affinity to Tf receptor (low µM range) as diferric Tf, 
which is critically important to perform its biological functions (data not shown). At last, 
the absence of indium in all living systems provides an extremely low background during 
ICP-MS analysis, and therefore greatly improves the method sensitivity. 

2.2.4.3 ICP-MS analysis of serum sample spiked with In$_2$Tf 

Figure 2-22 shows a full-scan spectrum of serum sample spiked with In$_2$Tf acquired 
by ICP-MS. Signal from elements within 50 to 150 m/z range was recorded and shown in 
black trace. Colored bars represented the natural isotopic distributions of several 
elements. The excellent agreement between theoretical distribution and measured 
distribution further confirmed the identity of each element. As shown in the inset, low 
level of indium (0.5 ppb) can be readily detected without interferences from other
elements, suggesting good selectivity of this method.

2.2.4.4 Quantitation methods for indium in serum and CSF

Applying the new strategy, quantitation of Tf was converted to quantitation of indium in biological fluids, which can be easily achieved with high sensitivity even without purification step. Another great feature of ICP-MS analysis is the minimal matrix effect due to the complete ionization induced by high plasma temperature. In this study, 2.5% of nitric acid solution was used as surrogate matrix of serum and CSF to generate calibration curves (Figure 2-23). In both cases, excellent linearity was achieved for nearly three orders of magnitude.

Subsequently, the quantitation methods were validated for accuracy, precision and matrix effect. And all of them were well within 15% range (Table 4). Concentration as low as 0.02 ppb of indium can be confidently quantitated in CSF samples, providing necessary sensitivity for quantitation of low levels of In$_2$Tf in rat CSF after intravenous injection.
2.3 Conclusion

LC-MS based methods were developed and validated for quantitation of low levels of Tf* in biological fluids. Particularly, pitfalls in protein quantitation using a classic approach (acid-catalyzed O\textsuperscript{18} labeling) were discovered, and successful quantitation was achieved after circumventing those problems. In addition, a new strategy of O\textsuperscript{18}-labeling was developed to produce protein-based internal standard, which greatly improved the reliability of the quantitation results. Furthermore, an ICP-MS based method was also developed for quantitation of Tf in both serum and CSF samples using indium as a tracer. This novel approach took advantage of the metal-binding ability of Tf and the superb sensitivity of ICP-MS detection. Reliable quantitation of In\textsubscript{2}Tf can be easily achieved without protein enrichment, thus providing an attractive alternative to LC-MS based approach. Although MS-based method is not a mainstay for quantitation of protein drugs currently, development of novel methods as discussed in this work will definitely broaden the application of mass spectrometry for tasks ranging from pharmacokinetic studies to quality control of biopharmaceutical products.
Table 1. Top five proteins identified by MASCOT MS/MS ion search engine

<table>
<thead>
<tr>
<th>Protein List</th>
<th>Origin</th>
<th>Mascot Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Without Ni-affinity purification (1 µg of Tf*, 1 mL of bovine serum)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Albumin</td>
<td>Bos taurus</td>
<td>527</td>
</tr>
<tr>
<td>Transferrin</td>
<td>Bos taurus</td>
<td>472</td>
</tr>
<tr>
<td>alpha-2-HS-glycoprotein precursor</td>
<td>Bos taurus</td>
<td>251</td>
</tr>
<tr>
<td>Prothrombin</td>
<td>Bos taurus</td>
<td>101</td>
</tr>
<tr>
<td>apolipoprotein C-III precursor</td>
<td>Bos taurus</td>
<td>89</td>
</tr>
<tr>
<td>B. With Ni-affinity purification (1 µg of Tf*, 1 mL of bovine serum)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Albumin</td>
<td>Bos taurus</td>
<td>585</td>
</tr>
<tr>
<td>hemopexin precursor</td>
<td>Bos taurus</td>
<td>496</td>
</tr>
<tr>
<td>Transferrin</td>
<td>Human</td>
<td>356</td>
</tr>
<tr>
<td>complement factor H precursor</td>
<td>Bos taurus</td>
<td>332</td>
</tr>
<tr>
<td>apolipoprotein C-III precursor</td>
<td>Bos taurus</td>
<td>321</td>
</tr>
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</table>
Table 2. O^{18}-incorporation efficiency in cysteine-containing tryptic peptides of Tf*  

<table>
<thead>
<tr>
<th>Tryptic Peptides</th>
<th>Percentage of differentially O^{18}-incorporated peptides</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0xO^{18}</td>
</tr>
<tr>
<td>KPVEEYANC*HLAR</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>WC*ALSHHER</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>DYELLC*LDGTR</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>C*LKDAGDVAFVK</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>SVIPSDGPSVAC*VKK</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>DLLFRDTCV*LAK</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>FDEFSEGEC*APGSK</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>WC*AVSEHEATK</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>C<em>STSSLLEAC</em>TFR</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>EGTC<em>PEAPTDCE</em>KPVK</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>IEC<em>VSAETTEDC</em>IAK</td>
<td>&lt;1%</td>
</tr>
</tbody>
</table>
Table 3. Validation of quantitation method of Tf* in rat CSF

<table>
<thead>
<tr>
<th>low QC</th>
<th>theor conc (ng/50 μL)</th>
<th>number of replicates (n)</th>
<th>CV (%)</th>
<th>accuracy (%)</th>
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<tr>
<td></td>
<td>60</td>
<td>5</td>
<td>10.2</td>
<td>90.3</td>
</tr>
<tr>
<td>Medium QC</td>
<td>600</td>
<td>5</td>
<td>4.75</td>
<td>104</td>
</tr>
<tr>
<td>High QC</td>
<td>1600</td>
<td>5</td>
<td>9.92</td>
<td>101</td>
</tr>
<tr>
<td>LLOQ</td>
<td>10</td>
<td>5</td>
<td>10.3</td>
<td>81.2</td>
</tr>
<tr>
<td>Matrix effect</td>
<td>500 ng/100 μL</td>
<td>3</td>
<td>4.93</td>
<td>104</td>
</tr>
<tr>
<td>QC samples</td>
<td>theor conc (ppb)</td>
<td>number of replicates (n)</td>
<td>CV, %</td>
<td>accuracy, %</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>------------------</td>
<td>--------------------------</td>
<td>---------</td>
<td>-------------</td>
</tr>
<tr>
<td>Low</td>
<td>0.04</td>
<td>3</td>
<td>0.435</td>
<td>89.1</td>
</tr>
<tr>
<td>Medium</td>
<td>0.4</td>
<td>3</td>
<td>0.764</td>
<td>98.9</td>
</tr>
<tr>
<td>High</td>
<td>4</td>
<td>3</td>
<td>1.47</td>
<td>107</td>
</tr>
<tr>
<td>Low with 100 µL CSF</td>
<td>0.04</td>
<td>3</td>
<td>1.89</td>
<td>105</td>
</tr>
<tr>
<td>Medium with 100 µL CSF</td>
<td>0.4</td>
<td>3</td>
<td>0.975</td>
<td>106</td>
</tr>
<tr>
<td>High with 100 µL CSF</td>
<td>4</td>
<td>3</td>
<td>1.77</td>
<td>112</td>
</tr>
</tbody>
</table>
Figure 2-1. A general workflow of MS-based method for protein quantitation. Figure adapted from reference [77].
Figure 2-2. Strategies used for preparation of isotopically labeled internal standards. Blue: biological/clinical sample; yellow: internal standards. The horizontal line represents the stage where internal standards are spiked. Figure adapted from reference [31].
Figure 2-3. Investigate the recovery of Tf* after Ni-affinity purification by SEC. Black trace: control sample; blue trace: the first flow-through fraction; red trace: the elution fraction.
Figure 2-4. Investigation of Ni-affinity purification efficiency on bovine serum sample. Lanes: 1- NEB 2-212 kDa MW Ladder; 2- Tf*; 3- bovine serum; 4- Tf* + bovine serum; 5- flow through fraction; 6- purification fraction; 7- Tf*.
Figure 2.5. O18-labeled peptide DGAGDVAFVK prepared by trypsin-catalyzed O18 labeling (blue trace) and acid-catalyzed O18 labeling (red trace), in comparison with unlabeled peptide (black trace).
Figure 2-6. Quantitation of Tf* in bovine serum using different reference peptides. Identical amount of O18-labeled internal standard was added before LC-MS analysis.
Figure 2-7. Acid-driven deamidation demonstrated by peptide EFQLFSSPHGK from tryptic digestion of Tf*. Left column represents extracted ion chromatograms (XICs) of ionic species $I_0$, $I_8$ and $I_{13}$ respectively. Right column represents mass spectra at elution time 30’35"-30’45" and 30’00"-30’10” respectively.
Figure 2-8. LC-MS/MS analysis of the structure of ionic species $I_0$, $I_8$, and $I_{13}$ related to tryptic fragment EFQLFSSPHGK showing full views of fragment ions spectra (left column) and zoom views of m/z regions corresponding to $y^5^+$ and $y^6^+$ ions (right column). The retention times for the various parent ions are indicated in the total
Figure 2-9. Quantitation of Tf* in bovine serum using different reference peptides. Identical amount of O18-labeled internal standard was added before LC-MS analysis. Results were shown both with (grey bars) and without (black bars) correcting for internal standard deamidation effect. The inset shows the calibration curves for two select peptides before and after correcting for internal
Figure 2-10. Calibration curves for quantitation of Tf* in bovine serum using reference peptides DSAHGFLK (left) and YLGEEYVK (right).
Figure 2-11. A schematic representation of applying Cys O18-labeling for accurate protein quantitation.
Figure 2-12. ESI MS analysis of O18-labeled IAA in the negative ion mode showing the isotopic distribution of IAA anion.
Figure 2-13. Nano-LC/MS/MS analysis of tryptic peptides derived from a mixture of hTf capped with unlabeled and O18-labeled IAA. (A) Total ion chromatogram of the digest and extracted ion chromatograms of peak I₀ and I₄ representing unlabeled (black trace) and O18-labeled (gray trace) tryptic fragment KPVEYANC*HLAR. (B) Isotopic distribution of a triply charged peptide ion KPVEYANC*HLAR. (C) MS/MS (CID) of peptide ion KPVEYANC*HLAR shown in (B). (D) A zoom view of the MS/MS spectrum (c) showing y⁴⁺ and y⁵⁺ fragment ions.
Figure 2-14. Stability of O18-labeled peptide KPVEYANC*HLAR prepared by Cys O18-labeling under acidic condition (0.1% FA) (A) and near neutral pH (pH 8.0) (B). Number in the bracket represents the relative abundance of the isotopic peak $I_2$ as percentage of the total ionic signal.
Figure 2-15. Quantitation curves for Tf* constructed using tryptic peptides WC*ALSHHER (A and B) and WC*AVSEHEATK (C and D) plotted on logarithmic and linear scales.
Figure 2-16. Evaluation of enrichment or removal of proteins from Tf*/rat CSF sample after Ni-affinity purification. Blue: flow-through fraction; red: elution
Figure 2-17. Quantitation method using peptide containing two cysteine residues. The m/z range highlighted by grey area was extracted to generate XICs shown in the right column. The inset represents the control spectrum of this peptide alkylated by IAA-18.

\[ \text{ratio} = \frac{S_L}{S_H}, \quad \alpha (\alpha=0.14 \text{ in this work}) \text{ is the fraction of partially labeled peptides} \]
Figure 2-18. Quantitation method using peptide containing a single cysteine residue. The second isotopic peak of either unlabeled or fully labeled peptide highlighted by grey area was extracted to generate XICs shown in the right column. The inset represents the control spectrum of this peptide alkylated by IAA-18

\[
\text{ratio} = \frac{S_L}{S_H \times (1+\alpha)}, \quad \alpha (\alpha=0.14 \text{ in this work}) \text{ is the fraction of partially labeled peptides}
\]
Figure 2-19. Comparison of quantitation results by different reference peptides. Quantitation results from triplicate experiments were shown in different bars.
Figure 2-20. Calibration curve for quantitation of Tf* in 50 μL of rat CSF. A $1/X^2$ regression was applied. The inset shows an enlargement of the low end of the calibration curve.
Figure 2-21. Structure of chelates DTPA (A) and DOTA (B). Me³⁺: metal ion; FG: functional group
Figure 2-22. Full-scan spectrum of serum sample spiked with In$_2$Tf acquired by ICP-MS. Colored bars represent natural isotope distribution of several elements. The inset shows the signal from indium (both $^{113}$In and $^{115}$In) at a concentration of 0.5 ppb.
Figure 2-23. Calibration curves for quantitation of indium in serum (left) and rat CSF (right). The inset shows an enlargement of the low end of the calibration curve.
CHAPTER 3

\textbf{O^{18}-LABELING BASED METHOD FOR IDENTIFICATION OF DEAMIDATION PRODUCTS}

3.1 Introduction

As discussed in section 1, deamidation of asparagine has been observed to have a negative impact on protein quantitation. In fact, deamidation of asparagine and isomerization of aspartic acid in proteins are the most frequent non-enzymatic post-translational modifications (PTM) in vivo [66]. In a recent study, Liu et al. observed that endogenous IgG isolated from human serum possessed 23% deamidation on Asn384 [67]. Under physiological conditions, both deamidation and isomerization proceed through formation of a succinimide intermediate, and lead to a mixture of aspartyl (Asp) and isoaspartyl residues (isoAsp) [68]. Upon each deamidation event, an additional negative charge is introduced to the protein surface; furthermore, the formation of isoAsp residue elongates the protein backbone by inserting an extra methylene group, resulting in the β-peptide linkage. Thus, these modifications closely associate with protein conformation [69], function [70, 71], activity [72, 73], stability [74], aggregation [75] and even immune responses [76, 77]. \textit{In vivo}, the level of isoAsp is controlled by protein L-isoaspartyl O-methyltransferase (PIMT), which catalyzes the conversion of isoAsp back to Asp [78]. The lack of such a mechanism \textit{in vitro} leads to accumulation of isoAsp contents over time as Asn deamidation and Asp isomerization can also occur during protein production and long term storage. It is particularly
important for protein-based therapeutics, where the effect of isoAsp formation has been extensively examined [79]. Therefore, close monitoring of Asn and Asp degradation products in protein pharmaceuticals is highly desirable and necessary [80].

3.1.1 Traditional methods for identification of deamidation products

Traditional ways of detection and localizing isoAsp residues in stressed protein involve chemical hydrolysis and proteolysis. Edman sequencing is useful for identifying the formation of the isoAsp residue, because these reactions stop at the iso-peptide bond [81]. PIMT is also frequently applied, since it leads to the methylation of isoAsp residues as an intermediate step. Owing to the longer retention time of methylated isoaspartyl-containing peptide compared to its aspartyl counterpart on RP-HPLC analysis, valid assignment can be made [82]. Furthermore, the byproduct of the methylation reaction, S-Adenosyl-L-homocysteine (SAH), has been used to achieve global analysis of isoAsp residue content in protein samples, which has already been commercialized under the name IsoQuant [83, 84]. The utility of endoproteinase Asp-N for the differentiation of aspartyl- and isoaspartyl-containing peptides has also been explored, by taking advantage of the selective cleavage of aspartyl-containing peptides but not their isoaspartyl counterparts [85, 86].

3.1.2 MS and LC-MS based methods for characterization of deamidation products

Detection of deamidated peptides by MS is relatively straightforward, since a mass shift of +0.984 Da occurs upon each deamidation event, which can be readily detected.
by MS with high resolution. However, the differentiation between aspartyl- and isoaspartyl-containing peptides from Asn deamidation or Asp isomerization presents a significant challenge for MS due to their identical mass. Even when separated by HPLC, unambiguous identification of the two isomers can be difficult using LC-MS analysis alone. In fact, most of the assignments have been made based on either the ca. 3:1 relative peak intensity ratio or the elution order of the aspartyl- and isoaspartyl-containing peptides (isoaspartyl-containing peptides are believed to be more hydrophilic than their aspartyl counterparts due to the greater acidity at the side-chain, therefore have shorter retention time during RP-HPLC analysis). However, as has been reported by other researchers [86-88], the elution order is strongly dependant on the chromatographic conditions as well as the properties of the peptides. In fact, the inverted elution order (isoaspartyl-containing peptides have longer retention time than aspartyl-containing peptides) has been consistently observed in peptides with Asp/isoAsp residues located at the non-acetylated N-terminus [87, 89]. We have also observed examples with inverted elution order in this work (vide infra). Clearly, accurate assignment of aspartyl- and isoaspartyl-containing peptides cannot be achieved by elution order alone. Likewise, assignment of aspartyl- and isoaspartyl-containing peptides based on relative peak intensity ratio (ca. 1:3) is not always reliable. Since these peptides usually elute at different time, their relative ionization efficiencies can be modulated differently by mobile phase and/or co-eluting species, and may not reflect their fractional concentrations in solution. Furthermore, the 1:3 ratio had been
established based on the analysis of products of deamidation of short, unstructured peptides, while in proteins it may vary due to influence of the higher order structure [90, 91].

3.1.3 Tandem MS-based methods for differentiation of deamidation products

Other approaches to aspartyl- and isoaspartyl-containing peptides differentiation have been mainly focused on applying different tandem mass spectrometry techniques. Several successful examples have been reported using different collision-activated dissociation (CAD) techniques, in which the differentiation is mainly based on the specific reporter ions [92] or the relative fragment ion intensity ratios [93]. However, these approaches are highly dependent on the peptide sequence and very sensitive to experimental conditions. A much more successful strategy is the application of electron-based fragmentation techniques (electron capture dissociation, ECD, and electron transfer dissociation, ETD), which generate a pair of reporter ions (c+57 and z-57) that are unique to isoaspartyl-containing peptides [94], and another pair of reporter ions (z-44 and (M+nH)(n-1)−-60) that are unique to aspartyl-containing peptides [95, 96]. This approach has already been applied in a high-throughput manner to achieve proteome-scale identification and quantitation of isoaspartyl residues in biological samples [97, 98]. However, this technique is limited to ECD-capable instruments, and sometimes can be hampered due to the low abundance of the reporter ions [86].
3.1.4 An $^{18}$O-labeling approach to differentiation of deamidation products

Recently, a number of $^{18}$O-labeling based strategies were developed to facilitate the Asn deamidation or Asp isomerization studies [99, 100]. Most of them have taken advantage of the increased mass increment after deamidation due to $^{18}$O incorporation (from 1 Da to 3 Da). Particularly, by using $\text{H}_2\text{O}^{18}$ for sample storage and protein digestion, in vitro and in vivo occurring deamidation processes can be readily differentiated [101, 102]. However, none of these methods have explored the ability of using $^{18}$O-labeling for aspartyl- and isoaspartyl-containing peptides differentiation and assignment. In the current work, we demonstrate the possibility of using $^{18}$O-labeling and LC-MS method for unambiguous differentiation of aspartyl- and isoaspartyl-containing peptides. Taking advantages of different deamidation mechanisms in acidic and basic environments, isomer-specific mass tags have been introduced to $^{18}$O-labeled aspartyl- and isoaspartyl-containing peptides, which then can be easily distinguished by MS. Feasibility of this new method has first been demonstrated using a synthesized peptide EWSVNSVVGK. Subsequently, successful assignment was achieved for at least six Asn-containing peptides and two Asp-Gly motif-containing peptides, from the tryptic digestion of an 80 kDa non-glycosylated form of human serum transferrin (hTf), which is a part of a number of biopharmaceutical products that are currently under development [80, 103].
3.2. Results and discussion

3.2.1 O\textsuperscript{18} labeling assisted approach to differentiation of deamidation products

The mechanisms of Asn deamidation under both basic conditions and acidic conditions have been extensively studied [68]. Unlike deamidation under basic conditions, which generates both aspartyl- and isoaspartyl-containing peptides, acid-catalyzed deamidation only leads to formation of aspartyl-containing peptides. By preparing forced deamidation standards in H\textsubscript{2}O\textsuperscript{18} under different conditions, different levels of O\textsuperscript{18} incorporation in aspartyl- and isoaspartyl-containing peptides can be achieved, which can be exploited for unambiguous isomer differentiation (Scheme 4).

![Scheme 4](image)

As previously reported [56], acid-catalyzed labeling and deamidation result in O\textsuperscript{18} incorporation in all carboxylic groups, leading to a maximum mass increment of 4n + 9 Da (4 Da from each acidic residues, 4 Da from C-terminus, and 5 Da from deamidation-formed Asp residue), where n is the number of acidic residues and
carboxymethylated Cys residues in this peptide. On the other hand, base-catalyzed
deamidation only leads to a mass increment of 3 Da (1 Da from deamidation and 2 Da
from incorporation of one O\(^{18}\) atom), because only one O\(^{18}\) atom is incorporated during
the hydrolysis of succinimide intermediate. Indeed, due to the much lower rate of
Asp/isoAsp isomerization under basic conditions, the formation and subsequent
hydrolysis of succinimide intermediate for the second time is limited within the
experimental time-scale.

3.2.2 Preparation of O\(^{18}\)-labeled deamidation standards

Under basic conditions, Asn deamidation in proteins occurs at a much lower rate
compared to deamidation of short unstructured peptides due to the protection afforded
by the higher order structures. Therefore, both acid- and base-catalyzed deamidation
standards were prepared using peptides as starting material. Some variation in the
deamidation rate under basic conditions was still observed, since this process is
influenced by the sequence of the peptide, especially the neighboring residue on the
C-terminal side of Asn. In particular, deamidation occurs much faster in Asn followed by
a polar residue with a relatively small side chain (Gly, Ser, Thr and Asp), compared to that
followed by a hydrophobic residue with a bulky side chain (Val, Leu and Ile) [104]. In
order to accelerate the deamidation process, elevated pH (1% NH\(_4\)OH, pH ~ 10) and
temperature (50 °C) were applied. In contrast, the deamidation rate of Asn under acidic
conditions is barely affected by the peptide sequence. By incubating the model peptide
EWSVNSVGK in H$_2$O$^{18}$ at 50 °C and in the presence of 1% TFA, the deamidation process was monitored by LC/MS analysis overtime. After 24 hours’ incubation, nearly 15% of peptides were in aspartyl form, and the isotopic peak presenting maximum O$^{18}$ incorporation could be readily detected (Figure 3-1). Thus, both acid- and base-catalyzed deamidation standards were prepared by 24 hours’ incubation using the conditions described above.

3.2.3 A model peptide study

Forced deamidation of model peptide EWSVNSVGK was carried out in H$_2$O$^{18}$ under both acidic and basic conditions using the procedure described above. Subsequently, the O$^{18}$-labeled deamidation standards were analyzed by LC-MS. As previously reported [56, 105], oxygen atom exchange occurs between carboxylic groups and solvent at low pH, leading to a maximum mass increment of 8 Da ($I_6$) for peptide EWSVNSVGK. In addition, hydrolysis of the amide group in Asn residue under acidic conditions leads to an additional mass increase of up to 5 Da (the $I_{13}$ species). Indeed, only one deamidation product with a maximum mass increase of 13 Da (designated as peak 3 in Figure 3-2A) is observed under acidic conditions. In contrast, the base-catalyzed deamidation leads to the formation of two indistinguishable deamidation products (designated as peaks 1 and 3 in Figure 3-2B), with a maximum mass increase of 3 Da (the $I_3$ species) for each. Analysis of the mixture of acid- and base-catalyzed deamidation standards reveals a co-elution peak of species $I_3$ and $I_{13}$ (peak 3) which can be unambiguously assigned to
aspartyl-containing peptide. The other chromatographic peak for species $I_3$ (peak 1) can be assigned to isoaspartyl-containing peptide (Figure 3-2C). The corresponding mass spectra also allow the unambiguous differentiation between the two isomers to be made.

3.2.4 Method validation by ECD-based approach

To validate this approach, a mixture of acid- and base-catalyzed deamidation standards prepared from the model peptide EWSVNSVGK was further analyzed by LC-MS/MS using ECD to produce fragment ions. ECD MS/MS analysis allows a distinction to be made between aspartyl- and isoaspartyl-containing peptides, since it can generate a pair of reporter ions ($c_i$+58 and $z_{k-r}$-57) that are unique to isoaspartyl-containing peptides, and another pair of ions ($z_{k-r}$-44 and $(M+nH-60)^{(n-1)\text{+}}$) that are unique to aspartyl-containing peptides ($r$ is the position of Asp/isoAsp residue from the N terminus and $k$ is the total number of residues in the peptide). Both ion $I_3$ and ion $I_{33}$ (elution at $t_1$ and $t_3$) were selected for ECD MS/MS analysis (Figure 3-3). The reporter ion ($z_5$-57) was only observed from the fragmentation of ion $I_3$ at elution time $t_3$, which confirmed the identity of this species as base-catalyzed isoaspartyl-containing peptide. On the other hand, both ($z_5$-44) and $(M+2H-60)^{+}$ fragment ions were observed in the ECD spectrum of species $I_3$ acquired at elution time $t_3$, but not from the fragmentation of species $I_3$ at elution time $t_1$, which confirmed the identity of this species as base-catalyzed aspartyl-containing peptide. In addition, the corresponding ions ($z_5$+4)-44 and
(M+2H+8)-60 with additional incorporation of O$^{18}$ atoms were observed in the fragmentation of ion $I_{13}$, which confirmed the identity of this species as acid-catalyzed aspartyl-containing peptide. Despite the O$^{18}$ incorporation in the carboxylic group of Asp residue, the reporter ions did not carry any O$^{18}$ atom due to the consequent loss of O$^{18}$ atoms in the form of either CO$_2$ (44 Da) or C$_2$H$_4$O$_2$ (60 Da) during the fragmentation. These results are in excellent agreement with the proposed mechanism [94] for the neutral loss involving Asp residues during ECD fragmentation. The ECD MS/MS analysis clearly validates the O$^{18}$-labeling approach to identification of aspartyl- and isoaspartyl-containing peptides in deamidated peptides.

3.2.5 Characterization of deamidation products from human Tf

Following validation of the new methodology with ECD MS/MS, it was applied to detect and identify deamidation products of an 80 kDa plasma protein hTf, whose ability to traverse physiological barriers and be internalized by malignant cells is attracting significant attention in the biopharmaceutical sector [80]. Deamidation of protein pharmaceuticals is always a big concern during the protein production and storage. A general procedure to monitor the deamidation in proteins is by detecting the deamidated peptides in the protein digest (mostly by trypsin) mixtures. However, as has been reported before [88], the traditional overnight tryptic digestion can introduce false positives (artificial deamidation, especially in Asn-Gly motif). Therefore, in this study, a shortened digestion time (4 hours) was used to minimize the digestion-induced
deamidation. Accordingly, an extra step of removing the alkylating reagent as well as an increased enzyme to substrate ratio was applied in order to improve the digestion efficiency. The tryptic digests were further subjected to acid- and base-catalyzed deamidation in H₂O¹⁸, and followed by LC/MS analysis.

Successful assignment of aspartyl- and isoaspartyl-containing peptides was achieved for six Asn-containing peptides and two Asp-Gly motif containing peptides from hTf (Figure 3-4). Based on the number of acidic residues in the peptide (n), a specific ion (I₄n+9) was extracted together with ion I₃. As discussed earlier, the co-eluting species I₄n+9 and I₃ indicated the presence of aspartyl-containing peptide, while the other elution peak of ion I₃ indicated the presence of isoaspartyl-containing peptide. In the case of peptide DSGFQMNQLR, due to the presence of two Gln residues which are also targets of acid-catalyzed deamidation, three peaks of ion I₁₃ eluting at different times were observed. Nevertheless, even in this case the assignment of aspartyl-containing peptide can still be made based on the co-elution of one of such peaks with one of the elution peaks of ion I₃. By subtracting, the other two peaks of ion I₁₃ can be assigned to either one of the Gln-deamidated peptides. Furthermore, by extracting ion I₂, the isoaspartyl-containing peptide produced by Asp isomerization can be easily detected. Since Asp isomerization occurs much slower than Asn deamidation under basic conditions, it was only observed in peptides with an Asp-Gly motif which is the most susceptible isomerization site.
3.2.6 Elution order of aspartyl- and isoaspartyl-containing peptides in RP-HPLC

The elution order of aspartyl- and isoaspartyl-containing peptides from tryptic digestion of hTf was mostly in agreement with the general observation that isoaspartyl-containing peptides elute earlier than aspartyl-containing peptides due to the greater acidity at the side-chain. However, the inverted elution order did occur in the case of peptides NLREGTCPEAPTDECKPVK and DGAGDVAFKV (Figure 3-4), where both the Asn residue and Asp-Gly motif were localized at the N-terminus. It is likely that the more favorable interaction between the N-terminal amine group and the carboxyl group of isoAsp residue could increase the overall hydrophobicity of the peptide, leading to longer elution time (Figure 3-5). Indeed, another peptide CLKDGAGDVAFKV, as an extended version of peptide DGAGDVAFKV with three more residues at the N-terminal side of Asp, exhibited ‘normal’ elution order vis-a-vis aspartyl- and isoaspartyl-containing peptides (Figure 3-5). This could be explained by the unfavorable ionic interaction at N-terminus. Furthermore, peptide SVIPSDGPSVACVKK (Figure 3-5) also exhibited inverted elution order after Asp isomerization, probably caused by the hydrogen bonding between the side-chains of Ser and isoAsp residues. These results indicate the unreliability of assigning the aspartyl- and isoaspartyl-containing peptides based on elution order alone.
3.3 Conclusion

A new method based on O_{18} labeling and LC-MS detection was developed for unambiguous assignment of aspartyl- and isoaspartyl-containing peptides produced by Asn deamidation and Asp isomerization. By preparing the acid- and base-catalyzed deamidation standards in H_2O_{18}, specific mass tags were introduced to aspartyl- and isoaspartyl-containing peptides respectively, which could be easily distinguished by MS. Compared to the traditional method where the assignment of the isomers is based on the assumption of elution order, this new method is both more accurate and reliable. Furthermore, since the method can be applied to the entire digest of a protein, multiple assignments can be made simultaneously. The procedure is considerably more cost-effective and faster than the use of synthetic aspartyl- and isoaspartyl-containing peptide standards. The new method also offers an attractive alternative to the direct ECD MS/MS analysis of deamidation products, which relies on detection of reporter ions and, therefore, may not produce a conclusive answer if the fragment ion intensity is insufficient. Furthermore, isomer identification can be carried out with the new methodology using a generic and inexpensive LC-MS system without tandem capabilities, such as single quadrupole MS.
Figure 3-1. Process of acid-catalyzed deamidation (at 50 °C with 1% TFA) of model peptide EWSVNSVGK in H₂O. XICs of m/z range from 503 to 511 were generated to estimate the relative abundance of deamidation-produced aspartyl-containing peptide (left column). The mass spectra acquired across the deamidation peaks are shown in the right column.
Figure 3-2. LC-MS analysis of the O18-labeled acid-catalyzed (A) and base-catalyzed (B) deamidation standards generated using model peptide EWSVNSVGK, and their mixture (C). The total ion chromatogram (TIC) is shown in black trace, and the extracted ion chromatogram (XIC) of ionic species $I_0$, $I_3$, $I_6$, and $I_{13}$ are shown in grey trace, respectively. The elution peaks are labeled with numbers, and the corresponding mass spectra are shown in the insets. N: Asn; D: Asp; isoD: isoAsp.
Figure 3-3. ECD MS/MS analysis of O18-labeled deamidation standards (EWSVNSVGK). The inset in each panel represents the total ion chromatogram (TIC), and the ion selected for each ECD MS/MS analysis is highlighted by the grey area. The signature ions are also highlighted by the grey area. The number in brackets after the amino acid residues indicates the mass increment in this residue due to O18 incorporation.
Figure 3-4. The aspartyl- and isoaspartyl-containing peptides assignment from tryptic digestion of hTF. The number in brackets after the peptide sequence indicates the number of acidic residues and carboxymethylated Cys residues in this peptide. XICs of ionic species $I_0$, $I_3$ and $I_{4n+9}$ are shown in black, dark grey and light grey, respectively. N: Asn; D: Asp; isoD: isoAsp; E: Glu.
Figure 3-5. Examples of peptides exhibited both normal and inverted elution order for their aspartyl- and isoaspartyl-forms, and the possible explanations. The XICs of ionic species $I_0$ and $I_2$ are shown in black and grey, respectively.
CHAPTER 4

RANKING THE SUSCEPTIBILITY OF PROTEIN DISULFIDE BONDS TO REDUCTION

4.1 Introduction

Disulfide bonds are generally considered to play a significant role in maintaining and stabilizing structure of proteins, primarily by decreasing the entropy of the unfolded state [106]. They are frequently found in extracellular proteins where they can promote the resistance of proteins to proteolysis and denaturation in the harsh extracellular environment. In addition to the structural role, some disulfide bonds also possess roles in regulating molecular functions [107, 108]. For example, catalytic disulfide bonds present a special group of functional disulfides that can catalyze the formation, reduction, or isomerization of disulfide bonds in their protein substrate [109, 110]. More recently, it has been realized some other disulfide bonds can regulate protein function allosterically by triggering a conformational change upon their reduction [111, 112]. Those disulfides are termed ‘allosteric’ disulfides to distinguish them from the catalytic disulfides [108]. Indeed, disulfide bond structure is critically important to maintain protein structure, stability and biological functions, and therefore a lot of effort has been devoted to characterize this unique covalent bond in protein. A variety of analytical tools have been developed to identify the linkages of disulfide bonds, including NMR analysis [113] and Edman Degradation [114]. Particularly, MS-based methods have proved to be extremely powerful to elucidate very complicated disulfide connectivity [115-119]. In
contrast, the stability of those disulfide bonds, particularly under reductive conditions, has been less appreciated, although it can play a significant role in protein stability and function. For example, reduced binding affinity of rabbit IgG and human IgG to the first component of complement (C1q) has been observed upon partial reduction of their disulfide bonds. In order to identify those “weak spots” or active sites, it is important to develop a method for complete ranking of susceptibility of disulfide bonds upon reduction.

4.1.1 Screening for possible functional disulfide bonds

Despite the primary structural role of protein disulfide bonds, it has become clear that some disulfide bonds, particularly in cell-surface proteins, are targets of redox regulation in the immune system [120]. For example, during the priming and activation of T cells, an elevated thiol level on cell surface can be observed due to reduction of disulfide bonds in cell-surface proteins by thioredoxin [121]. Indeed, those functionally important disulfide bonds are frequently redox-labile, since they have to be sensitive to the changes in redox condition in order to switch between bonded and unbounded states. Taking advantage of this feature, a proteomic-based approach has been reported to systematically screen for membrane proteins containing labile disulfide bonds. The results suggested that reduction of certain disulfide bonds in a wide range of membrane proteins was common during immune activation [107].
4.1.2 Disulfide bonds responsible for covalent aggregation of proteins

Formation of covalent aggregates of proteins via intermolecular disulfide scrambling is a great concern for protein therapeutics, which can result in both diminished bioactivity and increased immunogenicity risk [122, 123]. For example, the dimerization of IgG2 molecule was reported to be a result of disulfide bond formation between cysteines located in the hinge region [124, 125]. Furthermore, intermolecular disulfide scrambling was also attributed to the particle formation of antibody under agitation stress [126]. Apparently, labile disulfide bonds in proteins are the primary targets for the formation of nonnative disulfide bonds, by providing active free cysteines, which emphasizes the importance of ranking the susceptibility of disulfide bonds in a protein to reduction.

4.1.3 Disulfide-based Protein Modification

Recently, a novel disulfide-based protein modification approach was reported in several studies, in which a disulfide bond is selectively reduced followed by bis-alkylation to insert the modification reagent (e.g., PEG) [127, 128]. Because the modification site can be well controlled by selective reduction under mild conditions, site-specific modification can be achieved; this is an attractive feature compared to traditional amine-based modification. In addition, since the reduced disulfide bond is re-bridged after bis-alkylation, the protein’s tertiary structure is normally maintained and the biological activity is preserved [129]. Frequently, crystal structure of a protein can be
used to estimate the relative surface accessibility of each disulfide bond. More recently, a computation approach has been developed for identification of solvent-accessible disulfide bonds using published structural information [130]. However, it is important to emphasize that the static image generated from X-ray crystallography cannot fully represent the dynamic properties of a protein in solution. Furthermore, in addition to solvent accessibility, other factors (as discussed below) have also been shown to contribute to the susceptibility of disulfide bonds upon reduction. Thus, a direct measurement of susceptibility of disulfide bonds in protein is favored in order to accurately predict the disulfide-based modification site.

4.1.4 Factors contribute to susceptibility of protein disulfide bonds to reduction

Obviously, disulfide bonds at the surface of a protein are probably more susceptible to reduction, since they are accessible to either small thiols or enzymatic reducing agents. However, there is evidence that, although rare, some buried disulfides can be readily reduced with the assistance of exposed disulfides via an intra-molecular thiol-disulfide exchange mechanism [131]. In addition, dynamic features of a protein in solution can also greatly affect the stability of disulfide bonds upon reduction, and these cannot be predicted based on the static image generated from X-ray crystallography [130]. A recent bioinformatics study also suggested that labile disulfide bonds are usually found to carry relatively high dihedral strain energy that make them more easy to cleave [108]. Furthermore, the presence of a positively charged neighboring residue near a
disulfide bond can also enhance its lability by providing a favorable electrostatic environment for increased local concentration of thiolated anion [132]. Finally, more recently, mechanical force was suggested as another important factor that affects the stability of disulfide bonds upon reduction, although those studies are currently limited to engineered disulfides [133, 134].

4.1.5 Ranking the susceptibility of protein disulfide bonds to Reduction

Indeed, the presence of multiple contributing factors greatly amplifies the challenge for prediction of susceptibility of disulfide bonds upon reduction. Although X-ray crystallography is extremely useful to probe the solvent accessibility and calculate the dihedral strain energy (DSE) of a disulfide bond based on protein crystal structure. For proteins without crystal structure or the resolution is insufficient, this approach is less useful. Indeed, crystallography has been proved to be particularly useful for fast- and large-scale screening of labile disulfides in the protein database [108]. However, direct measurement is still required to achieve more reliable and accurate ranking of the susceptibility of all disulfide bonds in a protein. Fortunately, the development of mass spectrometry provides a powerful tool to address this challenging task. Particularly, proteomic-based methods have been developed to determine the cysteine oxidation status using a differential alkylation approach [135]. More recently, a similar approach was applied to rank the susceptibility of disulfide bonds in human IgG1 antibodies [136].

In this study, we proposed a new LC-MS/MS based approach for the ranking of
susceptibility of disulfide bonds in human Tf using $^{16}$O/$^{18}$O labeled iodoacetic acid for differential alkylation. The preparation and quantitative application of $^{18}$-labeled iodoacetic acid (IAA-18) have previously been reported in our lab [137].
4.2 Results and discussion

4.2.1 Experiment workflow

15 µg of hTf was prepared in 20 µL of 50 mM potassium phosphate buffer at pH 7.4. Partial reduction was initiated by addition of DTT to a final concentration of 10 mM, and followed by incubation at 37 °C for 5 min. Reduction was quenched by a 25-fold dilution into the alkylation solution consisted of 10 mM of O\textsuperscript{16}-labeled iodoacetic acid (IAA-16) and 6 M of guanidine hydrochloride at pH 8.0. Alkylation of free cysteine residues was conducted at 37 °C for 30 min in the dark. Subsequently, the excess IAA-16 was removed by extensive buffer exchange into a denaturing buffer (50 mM of potassium phosphate, 6 M of guanidine hydrochloride, pH 8.0) using a Vivaspin concentrator with a 10 kDa molecular weight cut-off membrane. The complete reduction was accomplished by the addition of DTT to a concentration of 10 mM and incubation at 37 °C for 30 min. Finally, all the free cysteine residues in hTf were alkylated with 30 mM IAA-18 at 37 °C for 30 min in the dark. After tryptic digestion, peptide fragments were subjected to nanoLC-MS/MS analysis, and cysteine-containing peptides were identified and used for calculation of reduction percentage of each disulfide bond (Scheme 5).

Scheme 5
4.2.2 Determination of reduction percentage of protein disulfide bonds upon partial reduction

The IAA-18 solution can be easily prepared by exchanging carboxylic oxygen atoms of commercially available IAA in \( \text{H}_2\text{O}^{18} \) at low pH [137]. Compared to C13-labeled alkylating reagents that are more frequently used [136, 138], IAA-18 provides larger mass separation (4 Da) between the differentially alkylated peptides. Analysis of the control sample alkylated with IAA-18 solution revealed that less than 1% of alkylated peptides failed to incorporate any \( \text{O}^{18} \) atom (Figure 4-1B). In addition, it has been demonstrated that the ratio of fully labeled products (+4 Da) and partially labeled products (+2 Da) is constant among all cysteine-containing peptides [137]. This feature greatly simplifies the calculation of the reduction percentage, which can be estimated using the following equation:

\[
\text{Reduction} \quad (\%) = \frac{I_1}{I_1 + I_5(1+\alpha)}
\]

where \( \alpha \) (\( \alpha = 0.14 \) in this work) is the fraction of partially labeled peptides; and \( I_1 \) and \( I_5 \) represent the observed relative intensities of the second isotopic peaks for \( \text{O}^{16} \)-labeled peptides and fully \( \text{O}^{18} \)-labeled peptides, respectively (Figure 4-1C). Also, control experiments with 0 minute of reduction revealed the existence of after-quench reduction (data not shown), which occurred during the alkylation step (Scheme 5). The after-quench reduction would certainly contribute to the overall reduction percentage (from 3% to 8% among all the peptides), however it can be easily subtracted during data
4.2.3 Reduction percentage of disulfide-linked cysteine residues

An assumption was applied to the data analysis that two disulfide-linked cysteine residues should exhibit the same reduction percentage. Thus, the reduction percentage of a disulfide bond could be determined by either one of its cysteine residues. The validity of this assumption was demonstrated by six disulfides in hTf where both the cysteine residues could be measured (Figure 4-2). Indeed, identical reduction percentage (within experimental error) was observed between all the paired cysteine residues. This assumption was frequently applied in data analysis in order to improve the coverage of measurable disulfide bonds. In addition, this measurement could also be applied to assist the determination of unknown disulfide bond linkages by reducing possible combinations. For example, Cys-9 and Cys-118 in hTf were less likely to form the same disulfide bond, because they exhibited a significant difference in stability upon reduction (Figure 4-2).

4.2.4 Analysis of peptides containing two cysteine residues

In order to improve the coverage of measurable disulfide bonds, analyses of peptides with two cysteine residues were frequently required, although they produced convoluted mass spectra (Figure 4-3). To address this difficulty, the previously demonstrated assumption was applied (see Section 3.2.3). For example, the reduction percentage of two disulfide bonds within peptide EGTC*PEAPTDEC*KPVK could be easily
calculated using isotopic distribution of two of its disulfide-linked peptides SAGWNIPIGLLYC*DLPEPR and APNHAVVTRKDKEAC*VHK. Even in the absence of one disulfide-linked peptide (peptide APNHAVVTRKDKEAC*VHK in this case), reduction percentage of this disulfide bond could still be calculated using a mathematic method derived from two known isotopic distributions (Figure 4-3).

In the case of peptide C*STSSLEAC*TFR, none of the disulfide-linked peptides could be detected in LC-MS/MS analysis, and therefore tandem mass spectrum generated by CAD was applied for calculation of reduction percentage. As shown in Figure 4-4, the reduction percentage of the second disulfide bond could be easily calculated using the isotopic distribution of fragment ion $y_7^+$. Subsequently, the reduction percentage of the first disulfide bond could be calculated applying the same strategy discussed above. However, in order to acquire reliable and quantitative data from tandem mass spectra, it was critically important to ensure that ions generated from differentially alkylated peptides (ranging from 767 to 774 m/z) would be isolated to the first quadrupole and consequently get fragmented equally efficiently. The former was believed to be valid since isotope labeling should not affect the fragmentation efficiency of peptides. However, in order to isolate all the ions between the 767 to 774 m/z range for subsequent fragmentation without bias, the mass selection window was set to low. A control experiment using very low collision energy (5 eV) confirmed the equal efficiency of mass isolation (Figure 4-4b).
4.2.5 Analysis of disulfide bonds in hTf

Applying this method, the reduction percentage of 15 disulfide bonds in hTf (19 disulfide bonds in total) could be measured with good precision after 5 minutes of reduction by DTT (10 mM, 37 °C and pH 7.4), and therefore the susceptibility of those disulfide bonds to reduction could be readily ranked. As shown in Table 5, those 15 disulfide bonds exhibited a wide range of reduction percentage, ranging from 8% to 60%. In order to correlate the susceptibility with possible contributing factors, those disulfide bonds were also analyzed for solvent accessible surface areas (SASA), secondary structure feature and dihedral strain energy (DSE) [139]. All those structural analyses were based on a recently solved crystal structure of diferric-hTf (PDB ID 8v83) [140].

Not surprisingly, solvent accessibility was found to play one of the most important roles to modulate the susceptibility of a disulfide bond to reduction. For example, buried disulfide bonds such as Cys9-Cys48, Cys19-Cys39, Cys227-241, Cys355-368, Cys450-Cys523, Cys484-Cys498 and Cys563-577 all exhibited limited reduction with less than 20% reduced in 5 minutes. In contrast, solvent accessible disulfide bonds such as Cys137-Cys331 and Cys339-Cys596 were observed to be much more susceptible to reduction. In addition, flexibility of the protein segments where a disulfide bond is located was also observed to be important. For example, disulfide bonds connecting an α-Helix and a β-strand (Cys9-Cys48, Cys19-Cys39 and Cys355-Cys368) were found to be much more resistant to reduction comparing to those with one cysteine residues located
at a flexible loop region (Csy137-Cys331 and Cys339-Cys596). Although it is difficult to evaluate the effect of flexibility independently without the interferences from other factors (e.g., disulfide bonds located in flexible region are frequently also solvent accessible), it is reasonable to speculate that good flexibility can enhance the interaction of a disulfide bond with reducing agents, and therefore increase its reductive susceptibility. Furthermore, dihedral strain energy (DSE) has been recognized as an important factor for disulfide bond stability for decades. Applying five successive $\chi$ angles, strain energy of a disulfide bond can be estimated using the following empirical formula:

$$DSE(\text{kJ/mol}) = 8.37(1 + \cos 3\chi_1) + 8.37(1 + \cos 3\chi_1') + 4.18(1 + \cos 3\chi_2) +$$

$$14.64(1 + \cos 2\chi_3) + 2.51(1 + \cos 3\chi_3)$$

where $\chi_1$ and $\chi_1'$ are the dihedral angles of two $\text{C}_\alpha$-$\text{C}_\beta$ bonds, $\chi_2$ and $\chi_2'$ are the dihedral angles of two $\text{C}_\beta$-$\text{S}_\gamma$ bonds, and $\chi_3$ is the dihedral angle of $\text{S}_\gamma$-$\text{S}_\gamma$ bond [141]. For example, two disulfide bonds Cys339-Cys596 and Cys345-Cys377 in the C-lobe of hTf were calculated to carry exceptionally high strain energy of 39.5 KJ/mol and 22.7 KJ/mol respectively (a mean value of 14.8 KJ/mol was found in a recent study of 6874 unique disulfide bonds [108]). Interestingly, both the two disulfide bonds exhibited unusually high susceptibility to reduction, particularly when comparing to their homologous disulfide bonds Cys9-Cys48 and Cys137-Cys331 in the N-lobe (Table 5). Apparently, multiple factors could modulate the stability of a disulfide bond simultaneously, emphasizing the indispensability of applying a direct measurement rather than
prediction based on crystal structure alone.

4.2.6 Disulfide bonds constraining the lobe movement in hTf

As shown in Table 5, disulfide bond Cys345-Cys377 is nearly three times more susceptible to reduction than its homologous disulfide bond Cys9-Cys48 (reduction percentage of 29% and 11% respectively), despite the fact that those two disulfide bonds are both completely buried and share exactly the same nearby residues and secondary structure features. Similarly, disulfide bond Cys339-Cys596 is significantly more labile than disulfide bond Cys137-Cys331, although they are very similar in solvent accessibility and local environment. Indeed, this discrepancy in stability can be explained by the difference in dihedral strain energy of those disulfide bonds (Table 5). Crystal structure reveals that disulfide bonds Cys339-Cys596 and Cys345-Cys377 are both located within or near the linker region of hTf that connects the N-lobe and C-lobe (Figure 4-5). Interestingly, although the linker region is not structured in hTf, it adopts a helical conformation in its family member lactoferrin. In addition, the unstructured linker region in hTf possesses a unique disulfide bond Cys339-Cys596 that is missing in both lactoferrin and ovotransferrin. As a result, function of this disulfide bond was predicted to constrain the relative movement between the N- and C-lobes [142]. This assumption agrees very well with our observations. It was speculated that the tension in the linker region generated from constraining the lobe movements could contribute to the elevated strain energy of disulfide bond Cys339-Cys596, by distorting its thermally
stable configuration. Similarly, disulfide bond Cys345-Cys377 could also be destabilized by the same mechanism, since it is located right after the linker region.

In order to demonstrate this hypothesis, reduction percentages of those two disulfides were also measured in apo hTf. It was speculated that the tension in the linker region would be released after iron removal due to the opening of both lobes, and therefore the two stressed disulfide bonds could be stabilized. Indeed, upon iron removal the reduction percentages of both disulfide bonds Cys339-Cys596 and Cys345-Cys377 were reduced by ca. 30%. In contrast, most of the other disulfide bonds maintained the reductive susceptibility after the conformational change (Figure 4-6).

4.2.7 Possible disulfide-scrambling site in hTf

Disulfide bond Cys118-Cys194 was another labile disulfide bond that can be easily overlooked by structure-based screening methods. The crystal structure of hTf suggested that this disulfide bond was completely buried and the calculated dihedral strain energy was only slightly above the mean value (Table 5). In addition, data from hydrogen/deuterium exchange MS experiments also suggested this disulfide bond was located in a rigid region, thus lacking of flexibility. However, as shown in Table 5, this disulfide bond was observed to be extremely labile upon reduction. Close examination of the crystal structure revealed a disulfide-rich segment aligning closely to this disulfide bond, particularly disulfide bond Cys158-Cys174 (Figure 4-7). Although those nearby disulfide bonds could not be detected due to the large size of tryptic fragment, they
were believed to be highly susceptible to reduction because of good solvent accessibility and flexibility. Therefore, an intra-molecular thiol-disulfide exchange mechanism was proposed to explain the lability of disulfide bond Cys118-Cys194 (Scheme 6). As a result, this region in hTf was also speculated to be a highly possible disulfide-scrambling site.

Scheme 6

![Disulfide Scrambling Diagram]
4.3 Conclusion

A new LC-MS/MS based method has been developed for ranking of susceptibility of disulfide bonds in protein upon reduction using a differential alkylation strategy. Based on the isotopic distributions of cysteine-containing peptides from both MS1 and MS/MS spectra, nearly all the detectable disulfide bonds could be measured for reduction percentage upon partial reduction. Applying this method, 15 disulfide bonds in hTf were ranked based on susceptibility to reduction, and subsequently possible contributing factors such as solvent accessibility, flexibility and dihedral strain energy were discussed. Particularly, two disulfide bonds Cys339-Cys596 and Cys345-Cys377 in the linker region of hTf were demonstrated to be responsible for constraining the relative movement of N- and C-lobe. In addition, lability of another disulfide bond Cys118-Cys194 was believed to be a result of an intra-molecular thiol-disulfide exchange event, indicating a possible disulfide-scrambling site.
Table 5. Analysis of disulfide bonds in hTf

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<th>Cys-1 secondary structure</th>
<th>Cys-2 SASA (Å²)</th>
<th>Cys-1 secondary structure</th>
<th>Cys-2 SASA (Å²)</th>
<th>DSE (KJ/mol)</th>
<th>Reduction Percentage</th>
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<td>8.52</td>
<td>18.3±0.5%</td>
</tr>
<tr>
<td>loops or irregular</td>
<td>6</td>
<td>bend</td>
<td>3</td>
<td>21.0</td>
<td>15.1±1.1%</td>
</tr>
<tr>
<td>loops or irregular</td>
<td>53</td>
<td>loops or irregular</td>
<td>12</td>
<td>14.3</td>
<td>32.3±3.8%</td>
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Figure 4-1. Mass spectra of a doubly charged peptide ion FDEEFSEGC*APGSK alkylated with either IAA-16 (A) or IAA-18 (B). C: Mass spectrum of differentially alkylated peptide FDEEFSEGC*APGSK after partial reduction (10 mM of DTT, 37 °C, pH 7.4, and 5 min of incubation).
Figure 4-2. Reduction percentages of cysteine residues from the same disulfide bonds. The values are shown by different bars (dark grey and light grey) in pairs. The error bar represents the standard deviation of three replicate measurements.
Figure 4-3. Mass spectra of three peptides that linked by two disulfide bonds. Equations on the bottom represent the calculation method for reduction percentage of one disulfide bond using the other two isotopic distributions.
Figure 4-4. (a) Isotopic distribution of differentially alkylated peptide C*STSSLLEAC*TFR. (b) Mass isolation of the whole isotopic cluster to the first quadrupole using ‘low resolution mass selection’ setting and low collision energy (5 eV). (c) Isotopic distribution of fragment ion y₇⁺ from differentially alkylated peptide C*STSSLLEAC*TFR.
Figure 4-5. A ribbon representation of the crystal structure of holo-hTf (3v83). Two disulfide bonds with high strain energy are shown in red.
Figure 4-6. Reduction percentages of hTf disulfide bonds in both holo-form (light grey bars) and apo-form (dark grey bars). The error bar represents the standard deviation of three replicate measurements.
Figure 4-7. A cartoon representation of a possible disulfide-scrambling site in hTf. Disulfide bond Cys118-Cys194 is shown in red. The three nearby disulfide bonds are shown in purple.
CHAPTER 5

GROWTH HORMONE-TF FUSION PROTEIN FOR SUCCESSFUL ORAL DELIVERY

This work was done in collaboration with Dr Cedric E. Bobst in our lab. Cedric performed the SEC and native ESI MS analyses of GHT fusion protein and the binding with TfR. He also investigated the stability of GHT in a stomach-like environment. I performed all the bottom-up analyses for the identification of GHTx and the binding between GHTx and TfR.

5.1 Introduction

In spite of the great success achieved by protein drugs in treating human diseases in the last decade, the development of orally deliverable protein drugs has been notably slow [10]. Indeed, the normal fate of protein drugs after oral administration is degradation due to the harsh gastrointestinal (GI) tract environment (not only the extreme pH in stomach but also the presence of various proteases). Even the survival of such conditions cannot guarantee efficient delivery, which also requires the traversing of intestinal-epithelial barrier. One of the very few successful examples is a recently designed human growth hormone/transferrin fusion protein (GHT) that exhibited measurable therapeutic response after oral administration. In that study, significant weight gain was observed in growth hormone-deficient mice after oral administration with GHT fusion protein, but not with growth hormone alone [8]. However, molecular basis for this rare success was not elaborated, and elucidation might provide important
insights for the design of orally deliverable protein drugs in the future.
5.2 Results and discussion

Mass spectrometry has become a powerful tool that can assist all stages of protein drug development. Using MS-based approaches, protein drugs can be characterized at different levels ranging from protein identification, covalent structure, conformation, interaction with physiological partners and PK/PD studies. Thus, in this study the GHT fusion protein was analyzed primarily by MS-based methods.

5.2.1 Protein identification by native ESI MS and a bottom-up approach

Size exclusion chromatography analysis of GHT clearly suggested two distinct fractions, with one eluting very close to the column’s void volume and the second one eluted slightly earlier than the Tf control. Subsequently, the species in the second fraction was identified as the monomeric form of GHT (GHT1) based on the accurate mass provided by native ESI MS analysis. However, a similar strategy could not be applied for the first fraction due to the large size of the molecular species (light scattering measurements indicated that components in this fraction were submicron particles). Instead, a so-called “bottom-up” approach [143] was applied for protein identification, in which the proteins of interest are digested by endoproteases, followed by LC-MS/MS analysis of proteolytic peptides and finally peptide mass fingerprint analysis or protein-database search to identify precursor protein(s). Briefly, protein samples were denatured in 6 M of guanidine-HCl buffer, followed by reduction and alkylation of cysteine residues, and finally digestion by trypsin. Analysis of the LC-MS data using the amino acid sequence of
GHT identified fragments derived from both Tf and growth hormone moieties. In addition, peptides from the helical linker region of GHT were also identified (Figure 5-1). This experiment unequivocally confirmed the presence of GHT in the submicron particle fraction, in which this protein existed as large soluble aggregates (GHTx).

It is also worthy to mention that during the bottom-up experiments, GHTx exhibited significant resistance to tryptic digestion in comparison to the Tf control, suggesting a promoted ability of GHTx to resist hydrolysis. Thus, it was speculated that the aggregation state of GHT might play an essential role in its ability to survive the harsh stomach environment. To validate this hypothesis, the stability of GHT1 and GHTx in the stomach-like environment (pH 3.5, 37 °C and in the presence of pepsin) was investigated by Cedric. By monitoring absorbance (214 nm) at the intact protein peak, the extent of proteolysis of both GHT1 and GHTx was measured by SEC. Indeed, GHTx displayed considerable stability in stomach-like environment, with a negligible decrease in abundance after 5 hours of exposure to acid and pepsin. In stark contrast, only 15% of GHT1 protein survived after 1.5 hours, demonstrating the essential role of protein aggregation to withstand the harsh GI tract environment.

5.2.2 Binding ability of GHT1 and GHTx to TfR receptor

Nevertheless, survival of GHTx in GI tract environment cannot guarantee effective delivery, since it still needs to cross the intestinal-epithelial barrier. Due to the wide distribution of Tf receptor (TfR) on the inner lining of GI tract [7], the Tf moiety was
believed to play a critical role in this step by taking advantage of receptor-mediated
transcytosis. Thus, it was necessary to demonstrate the ability of the complex to bind to
tfR. Using both SEC and ESI MS, we investigated the binding ability of GHT1 to TfR.
Mixing purified GHT1 with an excess of TfR led to a new peak that eluted earlier than
free GHT1, indicating formation of a complex. The identity of this complex was
confirmed using a new ESI MS-based strategy developed in our lab [144]. Both
complexes GHT1-TfR and (GHT1)_2-TfR were observed, indicating the same binding
stoichiometry as Tf/TfR complexes. In addition, since excess of TfR led to complete
elimination of free GHT1, the binding affinity could be estimated to be submicromolar
range (all protein concentrations were in low- and submicromolar ranges).

Unfortunately, similar approaches could not be applied to investigate the binding
ability of GHTx to TfR. Owing to the large size, a retention time shift of GHTx upon
binding to TfR was simply not possible. However, we did note that addition of an excess
of TfR to GHTx led to a significant change in area of the early-eluting peak, indicating a
binding event between GHTx and TfR. In order to demonstrate the specific feature of the
binding, a control experiment was conducted where addition of an excess of albumin to
GHTx did not result in detectable change in area of the early-eluting peak. Although the
components in the early-eluting peak could not be analyzed by MS directly due to the
large size, isolation of this fraction followed by bottom-up analysis revealed the
presence of both GHT and TfR.

As shown in Table 6, the top 5 protein hits by MASCOT MS/MS ion search engine
was listed. Furthermore, identification of one such tryptic fragment (LTTDFGNAEK) from TfR was demonstrated in Figure 5-2. Both MS/MS pattern (Figure 5-2C) and retention time (Figure 5-2D) of this fragment derived from the early-elution peak closely matched that derived from the TfR control sample, indicating presence of TfR in the early-elution peak. Those results suggested that the aggregate state of GHT remained binding affinity to TfR, which was essential for successful oral delivery.
5.3 Conclusion

This study investigated the molecular basis responsible for a rare success of GHT as an orally deliverable protein drug. Once again, Tf molecule was demonstrated as a promising drug delivery vehicle to overcome the intestinal-epithelial barrier. In addition, this study emphasized the important role of soluble aggregates of proteins to achieve successful oral delivery, by promoting the stability of protein drugs in harsh GI tract environment. This example argued the traditional negative impression of protein aggregation phenomena, particularly in the biopharmaceutical field. Instead, it suggested a possibility of exploiting protein aggregation in the design of efficient oral deliverable protein drugs. Finally, this study further demonstrated that MS was a powerful tool for characterization of protein drugs at various levels, such as protein identification and monitoring interaction with physiological partners as shown in this example.
Table 6. Top five proteins identified in the first elution peak

<table>
<thead>
<tr>
<th>Protein</th>
<th>Protein Score</th>
<th>Peptide matches</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cationic trypsin OS=Bos taurus</td>
<td>836</td>
<td>10</td>
</tr>
<tr>
<td>Transferrin receptor protein 1 OS=Homo sapiens</td>
<td>94</td>
<td>5</td>
</tr>
<tr>
<td>Transferrin OS=Homo sapiens</td>
<td>89</td>
<td>7</td>
</tr>
<tr>
<td>Keratin, type I cytoskeletal 9 OS=Homo sapiens</td>
<td>85</td>
<td>4</td>
</tr>
<tr>
<td>Human growth hormone OS=Homo sapiens</td>
<td>81</td>
<td>2</td>
</tr>
<tr>
<td>Keratin, type II cytoskeletal 1 OS=Homo sapiens</td>
<td>81</td>
<td>5</td>
</tr>
</tbody>
</table>
Figure 5-1. Peptide mass fingerprint analysis of high molecule weight fraction from SEC. The sequence of growth hormone is in italics, the sequence of the linker is in bold, and the rest is the sequence of transferrin. The underlined peptides were identified within a mass tolerance of 50 ppm.
Figure 5-2. A. SEC analysis of GHTx, TfR, and their mixture. B. total ion chromatograms of tryptic digests of SEC fractions of TfR (blue trace) and GHTx/TfR mixture (red trace) highlighted in (A). C. reference MS/MS spectrum of a tryptic fragment T71 (LTTFDFAEK) of TfR derived from TfR fraction (blue) and GHTx/TfR fraction (red). D. extracted ion chromatogram of this tryptic peptide eluting at 25 min (marked with asterisk in (B). Figure is from reference [22].
CHAPTER 6

CONCLUSIONS AND FUTURE DIRECTIONS

Mass spectrometry has become a powerful tool that can assist the characterization of protein drugs at different levels ranging from protein identification, covalent structure, conformation, interaction with physiological partners and PK/PD studies. In this work, we focused on the development of MS-based methods for quantitation of transferrin, a promising drug delivery vehicle, in complex biological fluids. Particularly, pitfalls in a classic approach (acid-catalyzed O\textsuperscript{18} labeling) to preparation of O\textsuperscript{18}-labeled internal standard were discovered, and a novel approach (Cys O\textsuperscript{18}-labeling) was developed for reliable quantitation of exogenous transferrin in biological fluids. Alternatively, a new method based on indium labeling and ICP-MS detection was developed for ultrasensitive quantitation of Tf. Following method development and validation, those methods will be applied in an animal mode study (Rat), in order to investigate the biodistribution of human Tf after intravenous injection, as well as its BBB-traversing ability. In addition, MS-based methods also provide a possibility to achieve multiplexing so that different versions of protein drugs can be analyzed simultaneously. In proteomic-based approaches, multiplexing can be achieved with relative ease using protein-specific reference peptides. However, for ICP-MS based approach, the application of multiple metal ions or isotopes as tracers will be required in order to analyze more than one type of Tf in a single experiment. The selection of suitable tracers will require extensive
analysis of the metal ion including its binding affinity to transferrin, the binding affinity of metal-coded Tf to Tfr, and the ICP-MS detection of this metal.

In this work, a new method based on O\textsuperscript{18}-labeling and LC-MS detection was developed for unambiguous assignment of aspartyl- and isoaspartyl-containing peptides produced by Asn deamidation and Asp isomerization. This method provided a simple and reliable solution to a challenging analytical task, differentiation between deamidated isomeric peptides. In addition, this method can be easily applied to other protein drugs where deamidation is a great concern during protein production and long-term storage. Following the identification of deamidated species, accurate and sensitive quantitation of them presents another urgent and challenging task. We speculate that the application of MS-based methods in combination with stable isotope labeling will prove to be useful in this field, although elegant experimental design is required.

Furthermore, a method consisting of partial reduction, differential alkylation, and LC-MS analysis was developed in this work for ranking of susceptibility of protein disulfide bonds to reduction. Applying this method, 15 disulfide bonds in hTf was analyzed. Particularly, two disulfide bonds were found to play an important role in constraining lobe movement in hTf, which is in great agreement with other studies. In addition, a possible disulfide-scrambling site was found in hTf, although further study is required to confirm that. Furthermore, this approach will be applied to study covalent aggregates of proteins due to intermolecular disulfide scrambling. It is speculated that
the most labile disulfide bond in a protein is most likely responsible for disulfide-induced protein aggregation. This assumption will be validated by identifying the intermolecular disulfide linkages from protein aggregates induced by partial reduction.

Finally, our study also investigated the molecular basis responsible for a rare success of GHT as an orally deliverable protein drug. Tf molecule was demonstrated as a promising drug delivery vehicle to overcome the intestinal-epithelial barrier. In addition, the soluble aggregate state of GHT fusion protein was proved to be essential to achieve successful oral delivery, by promoting its stability in harsh GI tract environment. However, several questions were remained unanswered. For example, what’s the nature of GHT aggregates? Will the aggregates dissociate in the circulation after crossing the intestinal-epithelial barrier? What are the differences in pharmacokinetics properties between monomeric GHT and aggregate GHT? Answering those questions will greatly benefit the design of new protein drugs suitable for oral delivery.
BIBLIOGRAPHY


