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# DEVELOPMENT OF BIOANALYTICAL METHODS FOR CLINICAL APPLICATIONS AND DRUG SCREENING

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# DEVELOPMENT OF BIOANALYTICAL METHODS FOR CLINICAL APPLICATIONS AND DRUG SCREENING

### **XIAOHAN CAI**

### ABSTRACT

In the past decade, bioanalytical method development has become an integral part of clinical diagnosis, biomarker discovery, and drug discovery and development. The new and emerged bioanalytical technologies allow the quantitative and qualitative analysis of small molecules and biomolecules with high sensitivity and specificity. Specifically, the bioanalytical methods based on LC-MS and methylation-specific PCR are well suited for detecting low-abundance metabolites, proteins, and DNA in biofluids and tissues for biomarker investigation. They offer great clinical promises for early diseases diagnosis and therapeutic interventions. Besides, the LC-MS/MS quantitative method is essential for the estimation of pharmacokinetic and toxicological properties in drug screening.

In this work, modern bioanalytical technologies, together with their applications from biomarker discovery and validation in metabonomics, genomics and proteomics to drug discovery, were reviewed. Dependent on the type of molecules analyzed, different methods were established to achieve accurate and reliable detection. LC-MS/(MS) methods were developed and validated for quantitative analysis of bile acids and anti-cancer agent JCC76. The former has been successfully applied in a clinical study for the

diagnosis of inflammatory bowel diseases; and the latter has been utilized in a pharmacokinetics study for drug screening and optimization. In terms of proteomics profiling, a LC-MS/MS method was demonstrated for comparative analysis of serum peptides with the successful identification of a potential biomarker for ovarian cancer. Lastly, a comprehensive DNA methylation profiling for hepatocellular carcinoma was conducted through methylation-specific PCR methods. These methods enabled sensitive and specific detection of DNA hypermethylation on several tumor-associated genes.

In addition, this work discussed a major challenge of matrix effect in quantitative method development. Possible solutions were proposed for matrix effect prevention and troubleshooting. Moreover, standard addition coupled with internal standard method and optimizing sample extraction method was illustrated for compensating or minimizing matrix effect in chapter II and chapter III, respectively.

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### **CHAPTER I**

# INTRODUCTION OF BIOANALYTICAL METHODS FOR CLINICAL APPLICATIONS AND DRUG SCREENING

### 1.1. General introduction of bioanalytical methods and their applications

Bioanalytical science, with a focus on qualitative and quantitative measurements in biological materials, plays a key role in understanding diseases, clinical diagnosis, and drug discovery and development. The technologies in biomedical science have made significant progress over recent years. This facilitates bioanalytical method development of biomarker discovery, to become an integral component drug metabolism/pharmacokinetic (DMPK), and toxicological monitoring. Advanced technologies and enhancements of conventional platforms emerged from bioanalysis fulfill the requirements of clinical and pharmaceutical fields, including the improvement in mass spectrometry detection, fast chromatographic separation, high-throughput sample pretreatment, and melting curve analysis with high resolution for genomic assays.

Early diagnosis of diseases has great significance in improving survival rates and minimizing current invasive diagnostic procedures. This leads to another major clinical need in the accurate detection of molecular biomarkers for chronic illnesses and cancers. The biomarker study monitors different biological entities including nucleic acids, proteins, and metabolites to reflect the pathophysiology and progression of diseases. Ideal biomarkers need to be well-understood for their functions in the pathogenic processes and their values for clinical diagnostic, prognostic, and predictive outcomes. However, these molecular biomarkers often present in low abundance in the biological samples, bringing great challenges in reliable detection and validation. Although these challenges remain, there is a large number of biomarkers developed currently and to be assessed in clinical studies for their diagnostic and prognostic applications [1].

Besides the broad bioanalytical applications in biomarker discovery, the impressive growth of quantitative bioanalysis has been also well-documented in pharmaceutical drug discovery and development. In the past decade, more than 500 novel drugs were approved by U.S. Food and Drug Administration (FDA) to prevent and treat human diseases [2]. Each year, more than 3000 on-going clinical trials are carried out in the drug development phase [3]. Despite the enormous amount of lead compounds screened in the drug discovery phase, the drug development process is costly and risky with very low rate of clinical success. This drives the rational lead optimization in the earliest stage of drug discovery to improve the likelihood of drug approval and to prevent drug withdrawal on the market. Quantitative bioanalysis serves as a major tool for understanding pharmacological properties including absorption, distribution, metabolism, and elimination (ADME), as well as toxicity to guide the drug screening for lead candidates.

### 1.1.1. Clinical applications for biomarker studies

Biomarkers are defined as indicators of normal biological processes, pathogenic processes, or pharmacological responses to a therapeutic intervention according to the National Institutes of Health (NIH) working group [4]. Cancers are the most studied diseases for biomarker discovery since the early detection of cancers before metastasis is always desirable to greatly improve survival. With advanced technologies for molecular biomarker measurement, thousands of potential biomarker candidates have been discovered and linked with cancers. These disease-related molecules may involve in cell regulatory and post-translational modification processes as proteins, alter the expression of downstream target molecules as nucleic acids, and represent metabolic responses as endogenous small molecules.

Among different biomarker discovery tools, proteomics allows the identification, characterization, and quantification of differential protein expression involved in normal and pathological states. Blood is the most commonly used sampling source for proteome profiling since blood sampling is non-invasive compared to tissue biopsy. In cancers, the

intra- and inter-cellular events happen at the tumor tissue microenvironment, introducing the changes of accumulative protein expression in the circulating blood stream. The differentiated proteomic pattern may reflect the development of malignancy and can also provide diagnostic, prognostic and predictive value for cancers. Extra-cellular nucleic acids in biofluids is another popular source of biomarker investigation, since it can reflect the cancer cell transformation induced by gene mutation or hypermethylation [5]. The circulating free DNA and cancer-specific RNA have been profiled in numerous genomic biomarker studies for the diagnosis and staging of cancer diseases.

The epigenetic changes in DNA methylation are also commonly associated with tumorigenesis. Taking place in the promoter region of CpG islands, the methylation of cytosine residue in normal cells is well-maintained in a homeostasis through a feedback regulation of DNA methyltransferase mechanism [6]. The disturbances to normal cytosine methylation are fundamental contributors to the malignancy of cancer [7]: the hypermethylation of tumor suppressor genes results in the inactivation of gene transcription and thus the loss of their intrinsic functions; global change of hypomethylation in a wide area causes repeated sequences and transposable elements, ultimately resulting in the loss of genomic stability and increased mutations [8]. These aberration events are the rationales for discovering potential genomic biomarkers to aid the diagnosis, predict the clinical outcomes, and subsequently guide the therapeutic treatment. Therefore, it is of crucial importance to develop bioanalytical tools that support the profiling of aberrant methylation.

Besides the large biomolecules in cancer biomarker study, the endogenous metabolites in cells, tissues, and body fluids start to gain more interest for reflecting human health status from their quantitative information. The most common example in clinical application is the measurement of cholesterol to monitor the cardiovascular disease by physicians. The metabolic signatures of patients reflect the biochemical changes in diseases and the disturbed metabolic pathways. The analysis of glycerophospholipids, fatty acids, steroid metabolites, and bile salts have been conducted for examining abnormal lipid metabolism and interaction of intestinal microflora for liver and intestinal diseases. These research areas provide useful sources for metabolic biomarker profiling.

### 1.1.2. Bioanalysis applications in drug discovery and development

Drug PK and toxicity properties are key parameters in the screening and optimization of lead compounds in the drug discovery phase. An ideal drug candidate should demonstrate the ability be absorbed in the blood stream, reach desirable concentration for effective activity, and be eliminated without producing toxic metabolites.

High-throughput PK screening usually starts from *in-vitro* assays to study the drug-drug interaction and metabolism using liver microsomes and hepatocytes as experimental systems. However, the *in-vitro* results cannot truly represent the real physiological environment and may lead to mistaken conclusion about drug metabolism. Therefore, it is essential to assess the PK parameters *in vivo* to improve the candidate selection through animal models.

In order to accurately define the drug behaviors *in vitro* and *in vivo*, bioanalytical support has been a prerequisite in the pharmaceutical industry. The large amount of compounds involved in the lead optimization requires the quantitative method to be accurate, sensitive, and high-throughput to facilitate drug discovery. These requirements can be fulfilled by accurate sampling procedures, advanced chromatographic and mass spectrometric techniques, as well as automated sample preparation methods.

#### 1.2. Modern bioanalytical technologies

Modern bioanalytical technologies have been significantly broadened in the last decade, demonstrating its ability in accurate qualitative and quantitative determination for protein, nucleic acids, small molecular metabolites, and drugs in biological materials. The major methodologies used for proteomics investigation are based on mass spectrometry (MS). The dramatic progresses of MS instrumentation refine mass accuracy, resolution, and dynamic ranges, ensuring the successful detection of low abundance proteins in biofluids and structural confirmation with their characteristic precursor and fragment ions. In addition, the robust and reliable liquid chromatography (LC) system in low flow rate has greatly improved the sensitivity for the MS detection and confidence for structure illustration.

Besides its application in protein analysis, the hyphenation of LC and MS (LC-MS) is established as the state-of-the-art methodology for the quantitation of small molecular compounds due to its specificity and sensitivity. It is now widely accepted as the preferred method for the quantitative measurement of small molecule drugs and endogenous metabolites in various biological matrices including plasma, serum, blood, urine, intestinal fluid, and tissue.

With respect to epigenetic biomarker discovery, the majority of DNA methylation assays are based on bisulfite reaction, methylation-specific PCR (MSP), and melting curve analysis. Sodium bisulfite converts cytosine to uracil at unmethylated CpG site, leaving methylated one unchanged. The MSP methods with designed primers selectively amplify methylated DNA, bringing high analytical specificity and sensitivity.

### 1.2.1. Principles of LC-MS

### 1.2.1.1. LC separation

LC is the basic separation platform for bioanalysis. With this technique, target analyte can be separated with interfering protein, salts, and phospholipids content in complicated biological materials. The separation mechanisms of liquid chromatography are based on the distribution of analyte between the liquid mobile phase and a stationary phase. Depending on different type of stationary phases, different distribution mechanisms are applied.

Adsorption mechanism is applied for both normal-phase (NP) chromatography and reverse-phase (RP) chromatography. In NP chromatography, the stationary phase is a polar silica gel and the mobile phase is a non-polar solvent such as hexane, pentane, and chloroform. NP chromatography is preferable for non-polar analyte and the retention decreases as the non-polarity of analyte increases. Opposite to NP chromatography, the stationary phase of RP chromatography uses non-polar silica based packing materials after the surface modification with C8, C18, or phenyl. Accordingly, the retention decreases with increasing polarity of the compound and the amount of polar solvents. RP chromatography is suitable ideally for polar and ionic compounds, which makes it the most widely used LC application. The interaction of analyte with stationary phase and mobile phase solvent greatly depends on the hydrophobicity of the analyte.

Ion-exchange chromatography is based on the ion exchange equilibrium between the ionic or polar compounds with the stationary phase. With opposite charge with the ionic functional group of the stationary phase, the ionic compounds can be retained. The elution speed is related with the ionic strength of the counter-ions, pH environment, and the modifier contained in the mobile phase.

Size-exclusion chromatography is usually applied in the separation of macromolecules according to their ability to penetrate into the pores of stationary material. The elution time of analyte is merely based on their size, but not molecular weight. The retention decreases as the size increases.

Besides the aforementioned traditional chromatography, there are some modern approaches for improving chromatographic resolution and separation efficiency: ultraperformance liquid chromatography (UPLC), monolithic chromatography, and hydrophilic interaction chromatography (HILIC). Underlying the same basic principle with RP-LC, UPLC utilizes column with sub-2 µm particle size and a system that can handle elevated pressure. UPLC has great advantages in resolution, sensitivity, and speed over conventional HPLC, and thus is considered as a better tool for high throughput analysis. Monolith column is packed with highly porous material, which is designed to handle fast flow rate and ensure sufficient surface for separation at the mean time. Consequently, the separation speed and sample throughput are significantly increased. HILIC is a valuable alternative to NP chromatography for very polar compounds because polar compounds are hardly retained and experience bad reproducibility when using NP chromatography. In addition, the large portion of organic mobile phase used for the HILIC elution increases the sensitivity when coupling MS with LC for detection.

In general, a sample is separated and analyzed by LC in the following sequence: the sample solution is injected through an injection port, and then delivered by the mobile phase by high-pressure pumps, and finally flowed into the column for retaining and further elution (Fig. 1.1). The instrumentation design should consider the following issues: the high-pressure is generated when the solvents are pumped into the small particle filled stationary phase; the dead volume of connecting tubes, the injector, and the mixing valve should be minimized to prevent the reduction of analyte peak resolution; the sample

residue on the tubing and injector should be avoided for carry-over issue in quantitative analysis.

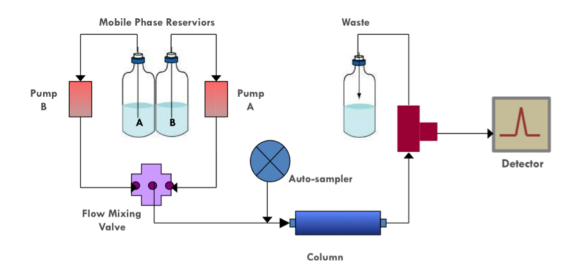


Figure 1.1, The instrumentation setup for LC separation

In this work, all small molecule compounds and peptides were separated by RP-LC with a C18 column and mass spectrometric detection. A guard column was used to prevent potential damages from the injected crude samples and proteins from the biological matrices. As the heart of LC system, the column needs to achieve adequate separation with a short time, maintain the precision for the retention, and have good stability in a broad pH range.

### 1.2.1.2. MS detection

MS has become a crucial part for pharmaceutical analysis and biomolecules research because of the improvements in ionization methods in the past decade. Taking advantages of powerful separation and sensitive detection, LC-MS analysis is well-suited for structural elucidation, accurate quantification, and metabolites prediction in complex biological matrices. The analyte in liquid flow eluted from the LC goes into three modules of ionization source, mass analyzer, and detector, undergoing ionization and evaporation, separation, and detection, respectively.

As the basic interfacing strategy, atmospheric-pressure ionization (API) enables the MS analysis by generating ions in a steam of liquid after LC separation. A number of API sources such as electrospray (ESI) and atmospheric-pressure chemical ionization (APCI) were developed to transfer the analyte from the liquid phase to the gas phase in MS with different ionization mechanisms.

In the ESI source, a highly positive or negative voltage is applied to the end of a steel capillary probe, where the sample solution is introduced (Fig. 1.2). When traveling along the capillary probe, the sample liquid is sprayed with excess charge. The nebulization process helps the formation of small droplets for ions in solvent vapor. During their flight from the electrical field to the MS proper, the droplets pass through an evaporation chamber, allowing the evaporation of solvents with the help of heating gas and nebulization gas. In the mean time, the quick evaporation process condenses the droplets and increases the surface charge density. At the end of the evaporation, the light solvent molecules diffuse away, leaving ions to enter into the MS analyzer.

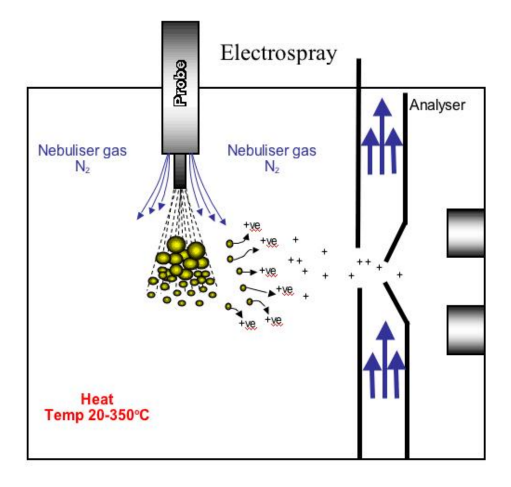


Figure 1.2, The schematic diagram of ESI source (http://www.ucl.ac.uk/ich/services/lab-services/mass\_spectrometry/metabolomics/hplc)

Unlike ESI, the ionization process of another liquid-based ionization source APCI occurs in gas phase. Without applying high voltage, the APCI capillary enables the volatile liquid sample to be heated and spayed first, and then a corona discharge needle with a high voltage generates ions from the aerosol cloud through interaction of reagent gas and electrons.

When ions are accelerated by the applied electric or magnetic field, the mass analyzer separates the ions according to their mass to charge ratio (m/z). Typical mass analyzers include quadrupole MS, ion-trap MS, and time-of-flight (TOF) MS with their own characteristics and applications.

The quadrupole instrument selects ions with a certain m/z to pass through the four parallel rods with certain direct-current potential and radiofrequency. Triple quadrupole MS allows the filtration of incoming ions by the first quadrupole, the fragmentation of selected ions by the second quadrupole, and the filtration of selected fragments by the third quadrupole, achieving high specificity. Similar to quadrupole MS, the ion-trap MS captures ions by a three dimensional manner in the electrical and magnetic combined fields (Fig. 1.3). The ion-trap MS takes advantages of its high sensitivity and resolution. The TOF MS accelerates ions and determines the flight time needed for ions moving from the ion source to the detector. TOF MS is characterized with advantages in high ion transmission and unlimited mass range, but with disadvantages in precursor ion selectivity. The modern MS instrument hybridizes different type of MS analyzers on one instrument, facilitating broader MS/MS applications.

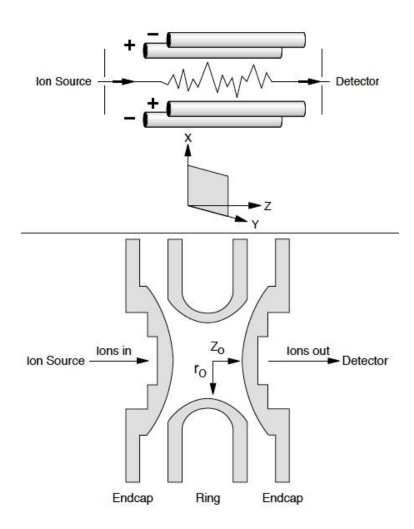


Figure 1.3, The operation of quadrupole MS and ion-trap MS (www.currentseparations.com/issues/16-3/cs16-3c.pdf)

### 1.2.2. Quantitative LC-MS(/MS) analysis for small molecules

The most prevalent method for small molecules quantification is based on quadrupole MS with ESI or APCI interface since 1990's. The detection of target analyte ions utilizes selective-ion-monitoring (SIM) and multiple-reaction-monitoring (MRM) approaches for LC-MS and LC-MS/MS methods, respectively.

The SIM mode detection is operated on single quadrupole instrument, with a particular m/z value selected for the target analyte ion. Because the impurities including proteins, phospholipids, and salts in the sample matrices may have the same m/z with target analyte, the LC-MS analysis requires more elaborate sample extraction and LC separation. Compared with SIM, MRM detection is more specific and sensitive since triple-quadrupole (QqQ) MS analyzer is applied for ion filtration and collision. Particular precursor ion and product ion are selected for detection based on their unique fragmentation pathway, resulting in a great improvement of signal to noise ratio (S/N) comparing to that of SIM.

The quantitative LC-MS(/MS) method development generally follows the workflow in Fig. 1.4. A successful bioanalytical method requires three interlinked methodologies in MS detection, chromatographic separation, and sample preparation. Some important aspects and challenges such as mobile phase choice, sample pretreatment, and matrix effect are discussed in the following sections.

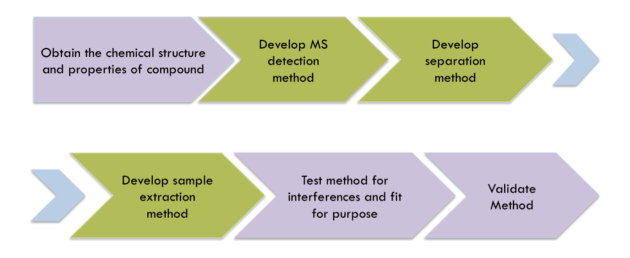


Figure 1.4, Method development workflow for small molecule bioanalysis

### 1.2.2.1. Mobile phase optimization

Mobile phase additives are often added in RP-LC for reproducible retention and the improvement of resolution and sensitivity when using MS as a detector. However, only volatile additives are compatible with LC-MS, because the non-volatile buffers such as phosphates may clog the ionization source and cause signal suppression. In addition, some volatile additives help the retention but deteriorate the MS ionization. For example, trifluoroacetic acid (TFA) is commonly used as an ion-pairing agent for increasing the retention of polar compounds. Nevertheless, it is also reported to induce significant signal suppression for some negatively and positively charged compounds [9-11].

Besides the LC modifiers, the pH of the mobile phase also has a large impact on both retention and ionization. By adding volatile acids such as formic acid, acetic acids, and their salts with ammonium in the mobile phase, the protonation of basic molecules under positive ionization mode is favored in acidic condition. Similarly, the deprotonation of some acidic molecules in negative-ion mode can increase the response by adding ammonium hydroxide as mobile phase additive. But these conditions may cause an adverse effect for retention if the hydrophobic interaction between analyte ions with the stationary phase is not sufficient [12]. In addition, the concentration of the additives is also critical since the MS response may be reduced under very high concentration, but concentration that is too low may lack buffer capacity. To solve the dilemma between retention and ionization, the selection of mobile phase composition needs to carefully consider all the characteristics of individual analyte.

### 1.2.2.2. Sample preparation

Although LC is a powerful tool for separation, the sample pretreatment for biological sample before injecting to LC-MS is essential for accurate and reproducible analysis. The biological sample matrices are very complicated with much higher content of proteins, salts, and endogenous lipids than target analyte. The large protein content in plasma sample is problematic due to its clogging of column and reducing analytical efficiency. Also, the endogenous interference and salts in most biological samples may suppress the ionization of analyte.

Conventionally, the sample cleanup has been performed by protein precipitation (PPT), liquid-liquid extraction (LLE), and solid phase extraction (SPE). PPT is popular when handling plasma sample because it is simple and fast. But the major disadvantage for PPT is that residues consisted of salts and endogenous material after the removal of proteins, which may greatly affect the MS detection [13]. Besides the use of organic solvents for denaturing proteins, other PPT additives such as acids, metal ions, and salts were reported to improve the efficiency of protein removal and disrupt the protein-drug binding [14].

LLE is an efficient technique to separate analyte from sample matrices based on the different distribution in the water-immiscible organic phase and aqueous phase. It is successful in giving excellent sample cleanup. But the disadvantages for LLE include the relatively large sample and solvent consumption, possible formation of emulsion, and unsuitability for hydrophilic compounds. Based on the conventional LLE, the salting-

out-assisted-LLE is developed as a more convenient alternative by adding concentrated ammonium salt solution into a mixture of biological sample and water-miscible solvents. In this way, high-throughput LLE can be applied through the automation of the handling process in 96-well plate.

The separation process of SPE method prior to sample analysis is similar to LC separation. The analyte is isolated relying on its affinity difference with the liquid sample solution and the solid SPE sorbents. Depending on the interaction of analyte and the selected solid phase, the SPE sorbents vary from polymer based ion-exchange materials to silica based materials. Typical SPE procedures start with the conditioning of the cartridge by a solvent or water (Fig. 1.5). Then the sample is added onto the cartridge and the analyte interacting with the sorbent is retained. While the interferences are removed after rinsing the cartridge with buffer or solvent, the analyte can be eluted with an organic solvent and further concentrated by evaporation and re-constitution.

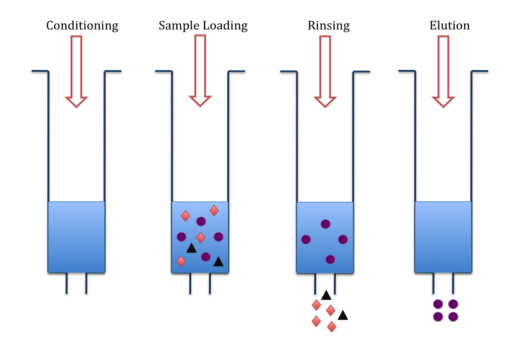


Figure 1.5, Illustration of four steps when performing SPE

Conventional SPE is performed on single cartridge, with the limitation for large volume of eluates and subsequent time-consuming evaporation process. The on-line SPE mode can be fully automated for directing injection of sample without any pretreatment. This advantage promotes the use of on-line SPE and 96-well plate, together with a column-switching system in high-throughput analysis to facilitate the extraction speed. The runtime for high-throughput analysis using SPE-LC-MS was reported to be within 5 min for many applications [15-17].

## 1.2.2.3. Matrix effect

Matrix effect is one of the major issues encountered during LC-MS method development and validation [18]. The phenomenon of matrix effect is observed when the ionization of analyte is suppressed or enhanced by the undetected co-eluting components from the biological matrix. The adverse results of matrix effect are reduced sensitivity for the detection and deteriorated precision and accuracy of the assay. According to the FDA guideline, it is required to assess the matrix effect when developing a reliable bioanalytical method.

In order to quantitatively determine the absolute matrix effect, an useful strategy was proposed by Matuszewski *et al.* [19]. The matrix effect is evaluated by comparing the signal response of analyte obtained from a neat solution with that from a post-extraction solution. In this way, two sets of samples are examined: one set is prepared by spiking standard analyte in neat solution and the other set is prepared by spiking standard analyte

at the same concentration in the extracted solution of blank biological sample, which is termed as post-extraction solution. The difference of response from these two sets determines whether signal is suppressed or enhanced. More importantly, the relative matrix effect should be evaluated by comparing the response of analyte in post-extraction solution from different blank matrix sources.

The post-column infusion of analyte is usually helpful to locate the co-eluting substances causing suppression in an LC run. A mixing tee is setup after the column elution and prior to MS ionization interface (Fig. 1.6). The post-extraction solution of a blank biological matrix is injected into the LC system, and then eluted by the mobile phase from the column. At the mixing tee, the matrix eluents mix with the analyte, which is infused constantly through an individual syringe pump. The MS monitors the signal change after the injection of post-extraction solution. The signal response of analyte should be expected to be steady in the absence of suppressing impurities. When the ion suppression or enhancement of analyte is present, the signal response will drop or increase at certain time points when interferences are eluted out, which can be easily observed on the chromatogram. In this way, the elution time of the ionization interferences and the extent of suppression or enhancement effect can be assessed through several continuous runs. The subsequent experimental design of analyte elution should avoid the co-elution with interferences.

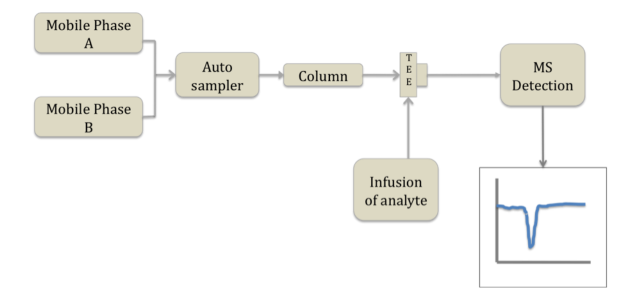


Figure 1.6, The post-column infusion experiment used for the assessment of matrix effect

Matrix suppression is induced by different reasons ranged from endogenous compounds from inadequate sample clean-up, ion-suppression mobile phase additives, choice of ionization method, to sample storage conditions. One of the most extensively used and efficient method to solve matrix effect issue is the utility of stable isotope labeled (SIL) internal standard. Since the SIL internal standard has very similar chemical structure and properties compared to the analyte, the ionization suppression or enhancement effect on both compounds is expected to be the same level. However, the SIL internal standard is costly and sometime hard to obtain. It is also problematic for tackling the "cross talk" problem if the purity of SIL internal standard is not adequate.

The matrix effect can also be minimized by improving the sample extraction method to remove the interferences. The endogenous compounds in biological samples have different polarity and thus are difficult to be completely removed by sample extraction methods. However, choosing the optimal sample preparation to reduce the amount of interferences is an efficient approach to ensure success in method development. Little *et al.* identified the phospholipids as a major contributor of matrix effect in blood and plasma by MS/MS using different extraction methods [20]. Their results suggested that the glycerophosphocholines caused matrix effect in both positive and negative ionization, with larger effect for isocratic elution than gradient elution. As the effect of different sample pretreatment methods on matrix effect, it is reported that LLE had lower signal suppression compared to SPE, followed by the PPT extract, which usually contains the most endogenous residues (Fig. 1.7) [21].

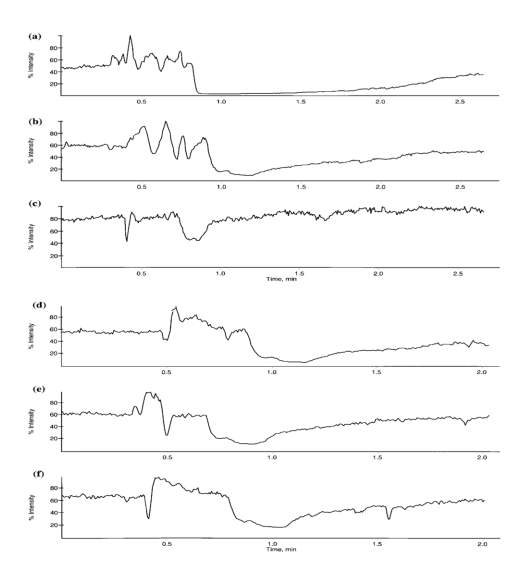


Figure 1.7, Infusion chromatograms (2.5 min) showing the ability of different sample preparation methods to remove endogenous sample components [21]. Panels (a) through (f) show the SRM XIC of a post-column infusion of phenacetin showing the effect of an on-column injection of 10 mL of a blank plasma sample prepared by each of the tested sample preparation methods. (a) Plasma protein precipitation blank. (b) Plasma Oasis SPE blank. (c) Plasma MTBE liquid-liquid extraction blank. (d) Plasma Empore C2 disk SPE blank. (e) Plasma Empore C8 disk SPE blank. (f) Plasma Empore C18 disk SPE blank.

Alternatively, adjusting the chromatographic conditions is another approach to reduce or eliminate the matrix suppression. In the RP-LC separation, the suppression effect is often found in an early time of isocratic elution program and at the end of gradient elution during the post-column infusion. Therefore, it is wise to alter the elution of target analyte at other regions of the chromatogram where the matrix effect is the lowest. Some mobile phase additives such as triethylamine (TEA) and TFA can also induce matrix effect in LC-MS analysis. The strong ion-pairing ability of these additives helps trap very polar compounds in the column and reduce peak tailing, but it also masks the detection by neutralizing the positive charge of analyte. The use of ammonium salts as a substitution or the choice of other columns with different retention mechanism can relieve this problem.

In bioanalytical method development, matrix effect is more frequently reported in ESI interface MS than APCI since the ionization mechanisms are different in these two sources. In ESI, the analyte is charged when traveling in the electrical probe, then nebulized to small droplets, and at last evaporated in the gas phase. When the interfering compounds compete with the analyte for the surface charge, the charge transfer occurs if the interferences have higher proton affinity, causing the lost of charge for the analyte and the decrease of MS intensity. Compared to ESI, the APCI of analyte in liquid undergoes opposite sequence for evaporation and ionization. The evaporation of liquid solvent takes place in the capillary before the ionization by charge transfer from the corona probe.

In addition to the endogenous compounds causing matrix effect problems, other often neglected sources from the dosing vehicles and blood anticoagulant can also result in ion suppression. Dosing vehicles including propylene glycol, Tween 80, and hydroxypropyl- $\beta$ -cyclodextrin are often used in the pre-clinical PK studies. Undetected matrix effect in the post-dose samples would give underestimated drug concentration and generate PK results with large errors. It also has been suggested that heparin should be avoided for separating plasma from blood during sample handling [22]. Sodium EDTA usually is preferred for anticoagulation in the PK and toxicokinetics studies for the prevention of matrix effect.

### 1.2.3. Mass spectrometric analysis for proteins

In the scientific process of biomarker discovery and evaluation, a lot of assays including ELISA, functional assays, flow cytometry, immunohistochemistry, and MS have been developed for biomarker analysis [23]. The interfacing of LC with MS, with the ability in qualification and quantitation, is the principal technique to define proteomes. Direct sequencing can be obtained by generating protein or peptide signature spectra and then imputing the spectra fingerprint into proteomics database.

#### 1.2.3.1. Identification of differentially expressed protein candidates

Body fluids are the major sources to characterize proteome. However, the analysis of proteins in human blood or urine presents a lot of challenges owing to the dynamic range

of protein concentration and complexity of sample components. The pre-fractionation of proteins is necessary to deplete the high-abundance proteins before MS analysis. With this purpose, different separation methods have been well developed prior to protein characterization.

Two-dimensional polyacrylamide gel electrophoresis (2D-PAGE) tandem with MS analysis is the first tool used to separate and identify differentiated proteins [24]. 2D-PAGE separates thousands of proteins by two steps. Firstly, samples are isolated by their different isoelectric points and by their molecular sizes. In this step, the protein pattern can be visualized by staining the gel. Secondly, the spots representing differentially expressed proteins are excised and digested into peptides by enzymes prior to MS analysis. However, when both abundant and less abundant proteins are presented on the same gel, it is necessary to determine the relative intensity of protein spots. Matrixassisted desorption/ionization time of flight mass spectrometry (MALDI-TOF MS) is commonly applied in spot identification to generate protein or peptide fingerprint. The MALDI-TOF MS analysis takes advantages in unlimited mass range and fast analysis speed. Bioinformatics can subsequently be utilized to search the proteomic database to identity the targeted proteins or directly analyze the results for discriminating protein patterns from control to patient samples.

Although 2D-PAGE with MALDI-TOF MS analysis is a simple and widespread tool for the analysis of complex protein mixtures, it have several limitations such as the relative low resolution of 2D-PAGE separation and the reproducibility of detection, and it is restricted to protein less than 20k Da [25]. Micro-scale or nano-scale chromatographic separation enhances the speed and efficiency comparing to conventional 2D-PAGE, facilitating the incorporation of high resolution LC in tandem MS as the primary protein identification platform.

The LC-MS/MS protein analysis can be performed on various mass spectrometers, such as quadurpole and linear ion traps, Orbitrap, and quadrupole-TOF. The bottom-up method for protein primary sequence determination involves the enzymatic digestion of protein into small peptides (Fig. 1.8). Retaining the digested peptide in acidic condition, the C18 reverse-phase microcapillary or nano-LC columns are the most commonly used ones to fractionate peptides with high resolution.

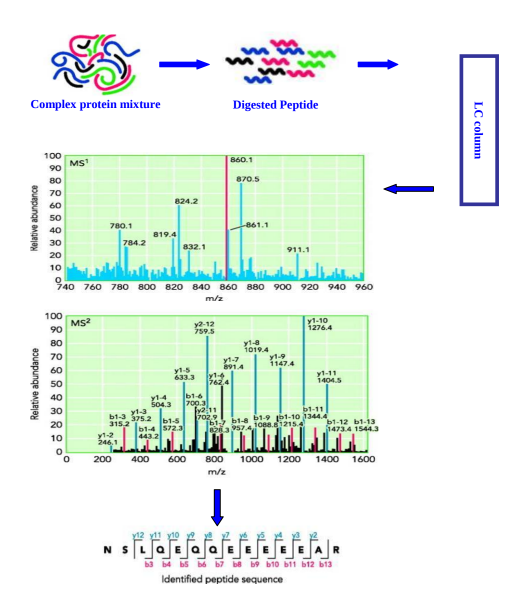


Figure 1.8, Bottom-up proteomics analysis using LC-MS/MS. Trypsin digestion is the first step for the analysis of a complex protein mixture. Afterward, the sample containing tryptic peptide is injected into the HPLC column of the LC-MS/MS system. The mass spectrometer generates parent ( $MS^1$ ) spectrum and fragmentation ( $MS^2$ ) spectrum created by CID. The computer will use the MS2 spectrum together with MS1 spectrum to compare the database containing theoretical peptide spectra to identify the protein of origin.

The gradient program of the LC system helps the elution of the peptides. When delivered to the mass spectrometer in acidic environment, the peptides are converted into cationic form with single or multiple charges [26]. The first stage is to produce MS<sup>1</sup> spectra for the m/z of various peptides. To generate more information for peptide identification, the fragments of selected peptide ions (precursor ions) are initiated. To produce fragment ions, tandem mass spectrometric analysis (MS/MS) uses collision-induced dissociation (CID) to generate MS<sup>2</sup> data for peptide sequence information. By searching the protein database with the acquired MS/MS spectra, the software program can identify protein with matching peptide mass fingerprinting. Clearly, the higher mass accuracy obtained from precursor ions and product ions, the more confidence for protein identification.

## 1.2.3.2. Quantitative protein analysis

With the speed-up of generating biomarkers, the center of proteomics has shifted from biomarker discovery to biomarker evaluation and quantification. It is necessary to compare the amount of differentially expressed proteins among healthy and disease status. Therefore, the development of accurate methods to quantify biomarkers of interest is a reliable theme for biomarker evaluation. Currently, the established techniques for protein and peptides quantification include two general strategies: 1) non-labeling methods by correlating MS signal with relative protein quantity and 2) the use of stable isotope incorporation prior to MS analysis. In non-labeling quantification, the quantitative information for specific peptide is derived from according ion intensity or peak areas in complex peptides mixture. The accurate quantification needs a careful calibration of instrument system and normalization of MS spectra. For LC-MS/MS quantification, peptide ion intensity counting of MS mode and spectral counting in MS/MS mode are extensively used for ion detection. With respect to peptide ion counting, the number and intensity of precursor ion or peak area at selected m/z can be obtained. The spectral counting approach refers to the number of fragment ions generated from selected peptide. The peptide levels are yielded by comparing referring the abundance of peptide between samples in two or more separate LC-MS/MS runs. Label-free approach for quantitative proteomics is preferable due to its low cost and no limitation for sample numbers [27].

However, there are still controversies for the reliability of label-free quantitative analysis. The accurate quantification of this approach requires the minimization for variations between different runs, high resolution for the chromatography to finding correlated peptide, and high MS accuracy to prevent interfering signals with similar m/z. It also assumes that the linearity of response is the same for every peptide, but in fact the spectrum count response varies from different peptides. Because of the dynamic range of peptides in a sample, the existence of high abundance peptide in the complex mixture will affect the accuracy of low abundance peptide quantization.

The major stable isotope labeling methods include using isotope coded affinity tags (ICAT) and isobaric tags for relative and absolute quantification (iTRAQ) [28]. The

basic principle of ICAT relies on a special affinity tag to react with cysteine residues and allows differentially labeled samples to be resolved with MS analysis. Two different tags, which are identical except for one has hydrogen and the other one has deuterium atoms in the linker, react with samples before protein digestion [28]. Their mass difference allows the relative quantitative measurement by MS. The disadvantages of ICAT are its limitation for proteins containing cysteine residues only and high cost due to the isotope reagent.

The iTRAQ approach allows multiplexed quantification by targeting all free amines at N terminus of all peptides and the epsilon-amino group of lysine residues. This technique enables the analysis of up to 8 samples in one run. The iTRAQ tags contain reporter group with mass from 114 to 117 Da, balance group from 28 to 31Da, and a protein-reactive group. During each MS scan, each labeled peptide displays the same mass to charge ratio. However, the dissociation of the reporter groups displays different m/z after the fragmentation under MS/MS mode. Signals from peptides after isobaric labeling are acquired for both MS and MS/MS scanning mode, thus improving the sensitivity and specificity of detection [29]. This property has great potential in the quantification of low abundance proteins. iTRAQ coupled with LC-MS/MS has been used as in the serum biomarker in several studies and it shows promise in determining differential expression profiles for cancer diagnosis, prognosis or monitoring of treatment [30].

When the quantitative information is derived for cataloging the protein files in a sample, the biomarker development requires careful follow-up validation, which includes the application of targeted proteomics methods. Targeted proteomics focused on individual proteins or a panel of proteins. When the use of isotope labeled methods only can provide relative quantitative information, the use of a standardized reference in the sample provides the absolute quantification information (AQUA). It is performed by spiking an isotopically-labeled internal standard into a biological sample prior to MS analysis. The ratio of labeled to unlabeled peptide determined by MS analysis can be calculated and the abundance of specific protein can be derived. Combining AQUA with MRM method in MS analysis, the specificity and accuracy can be improved for absolute quantification. MRM involves the selection of parent ions and then monitoring the fragmentation ions from the selected parent ions, thus enabling this technique highly specific and sensitive. Because the MRM detection limit allows at low as ng/ml level, this method offers the most promise for biomarker validation.

To develop proteomics biomarkers with high specificity and sensitivity for clinical application, several phases including biomarker discovery, evaluation, determination of biological relevance, and development of clinical assay are need to follow. Huge challenges were presented such the complexity, variation and dynamic range of proteins in biological samples. The new MS technologies play the most critical role in the improvement of resolution and sensitivity, bringing promises for more clinical successes.

### 1.2.4. PCR based methods for methylation profiling

DNA methylation is the most studied epigenetic changes related to normal biological processes and many diseases, especially in cancer development. This covalent modification of cytosine mostly happens at CpG dinucleotides rich sites, which are associated with gene promoters. While the methylation in normal cells is regulated by the DNA methyltransferases in a steady status for a stable genome, the aberrant gene methylation represses the transcription of downstream genes. In the process of tumorigenesis, a large number of tumor suppressor genes were found to be hypermethylated in the promoter regions [31]. Based on these findings, the investigation on DNA methylation has become one of the most popular areas in molecular oncology. Technologies for the genomic DNA methylation profiling are designed with enormous improvements with regarding to sensitivity, the elimination false-positive results, and sample throughput.

In currently used methodologies for methylation profiling, bisulfite conversion based method is the fundamental one to investigate the gene-specific methylation. With standard sequencing method, the similar base-pairing sequence of methylated and unmethylated cytosine cannot be distinguished. Sodium bisulfite treatment with genomic DNA can solve this problem by chemical reaction. Under certain conditions, sodium bisulfite specifically deaminates unmethylated cytosine to uracil but leaves the methylated cytosine unchanged (Fig. 1.9.). The uracil is replaced with thymine in the followed PCR amplification after bisulfite conversion. After this reaction, standard

methods such as sequencing, pyrosequencing, PCR, or mass spectrometry can be used to analyze the bisulfite-converted DNA product [32-34].

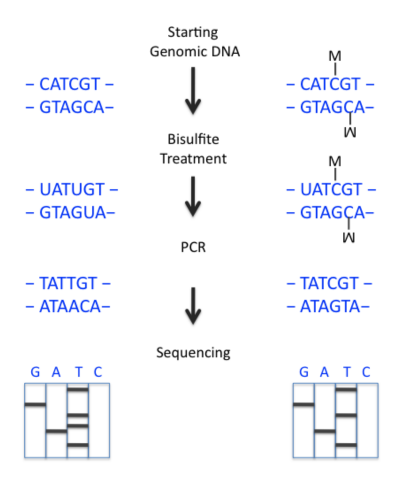


Figure 1.9, Strategy to distinguish unmethylated cytosine and methylated cytosine by bisulfite reaction

However, bisulfite treatment may give inaccurate results when the incomplete conversion of unmethylated cytosine is misinterpreted as methylated cytosine. It is critical to ensure the reaction is complete for unmethylated DNA by spiking known unmethylated DNA with the reaction as control. Another issue is the partial degradation as a result of DNA depurinization at acidic pH, which limits the sensitivity of the PCR reaction [35]. This problem can be overcame by adjusting the proper bisulfite reaction conditions of pH, temperature and the time of reaction [35].

Direct sequencing of bisulfite treated DNA allows the detection of methylation for each CpG dinuclotide within the analyzed area, thus is considered as the golden standard for methylation profiling. Nevertheless, the cost and labor intenseness of this approach is extremely high for large-scale sample analysis.

Alternatively, the differences between unmethylated and methylated DNA sequence can be characterized by melting curve analysis. Unmethylated and methylated DNA has different GC content after bisulfite conversion, presenting varied resistant levels to melting. To detect the signal of PCR product, fluorescence dyes are used for specific binding with double-stranded DNA (ds DNA). The fluorescence signal is monitored as the temperature increases, producing a melting curve to depict the relationship between fluorescence intensity with the increase of temperature. The characteristics of PCR product is indicated by the fluorescence peak with a certain melting temperature on the derivative melting curve. The fully methylated PCR product and fully unmethylated DNA shows distinct melting peaks, while a mixture of both may show a complex pattern with both melting peak characters [36] (Fig. 1.10).

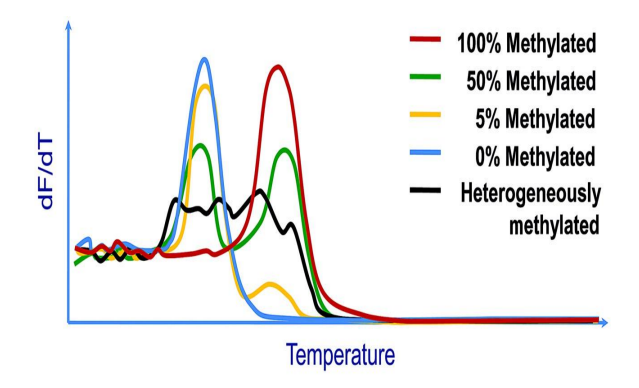


Figure 1.10, Melting curve analysis on fully methylated, fully unmethylated, and mixtures of both PCR products for methylation detection [34].

These sequencing and MCA methods such as combined bisulfite restriction analysis (COBRA), melting curve analysis methods including methylation-sensitive singlenucleotide primer extension (MS-Snupe), and methylation-sensitive melting curve analysis (MS-MCA) all reply on the PCR amplification prior to methylation detection. Methylation-independent PCR primers utilized in these methods allow proportional amplification of methylated and unmethylated templates. However, the templates of methylated DNA have higher GC content than unmethylated templates, leading to different amplification efficiencies and PCR bias for unmethylated product. Although many attempts are reported to overcome this problem such as increasing the annealing temperature during amplification and proper primer design [37], the sensitivity of these methods remains to be improved.

The development of methylation-specific PCR (MSP) in the mid-90s permits the simple and fast analysis of the DNA methylation status after bisulfite conversion. MSP method is highly sensitive and specific with designed PCR primer for the amplification of methylated sequence only. It was reported that MSP was able to detect 0.1% methylated template in a pool of unmethylated DNA [38]. MSP is also related with high falsepositive rates caused by the incomplete bisulfite conversion and possible contamination during analysis, which can be alleviated by increasing melting temperature and more stringent amplification conditions.

The development of quantitative MSP (qMSP) resolves the limitation of MSP based on real-time PCR (RT-PCR). The amount of initial DNA product can be determined with

high precision and a wide range. In addition to unknown DNA products, standard methylated DNA after bisulfite treatment is serially diluted and amplified for standard curves in separated reactions. Since the quantitative results are relative to the standards, the absolute quantity of DNA only can be obtained when the absolute copies of the standards are known.

As an alternative mechanism for cancer development, aberrant gene methylation has been found in the patterns of hypomethylation for global genome change and hypermethylation for specific tumor suppressor genes. The DNA methylation profiling holds promises for the biomarker discovery of cancers and other diseases for early detection. Nevertheless, these potential biomarkers are relative low abundant in biofluids or tissues, requiring the developed detection method for gene methylation to be highly sensitive and specific for potential clinical applications. A large number of target genes have been identified for colorectal cancer, breast cancer, and hepatocellular carcinoma. Moreover, the DNA methylation targets are more frequently to be observed as a panel of multiple genes rather than a single gene, suggesting a direction for improving the specificity of cancer screening.

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# **CHAPTER II**

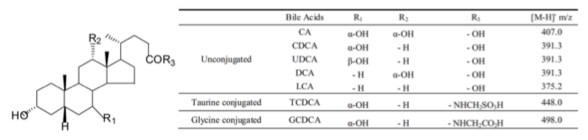
# QUANTITATIVE ANALYSIS OF BILE ACIDS FOR INFLAMMATORY BOWEL DISEASES

### 2.1. Introduction

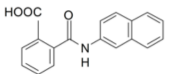
## 2.1.1. Bile acids

Bile acids are produced in the hepatocytes through the oxidation of cholesterol. They are composed of a steroid structure with side chain terminating in a carboxylic acid and hydroxyl groups (Fig. 2.1). After conjugating with taurine or glycine in the liver, bile acids are excreted into gallbladder, released into intestinal tract, reabsorbed in the terminal ileum, and then back to enterohepatic circulation, leaving a small part entering into the colon [1,2] (Fig. 2.2). In the terminal ileum, cecum and colon, primary bile acids

including cholic acid (CA) and chenodeoxycholic acid (CDCA) are metabolized by bacterial flora through two major reactions: one is deconjugation to produce free bile acids and taurine or glycine from conjugated bile acids; and the second one is dehydroxylation to form secondary bile acids deoxycholic acid (DCA), ursodeoxycholic acid (UDCA) and lithocholic acid (LCA) [3]. The numerous metabolic conversions of bile acids bring the complexicity of bile acids composition in biological fluids.



1a Structure of bile acids



1b Structure of the internal standard NPA [M-H]<sup>-</sup> m/z: 290.1

Figure 2.1, Chemical structures of bile acids (a) and internal standard NPA (b)

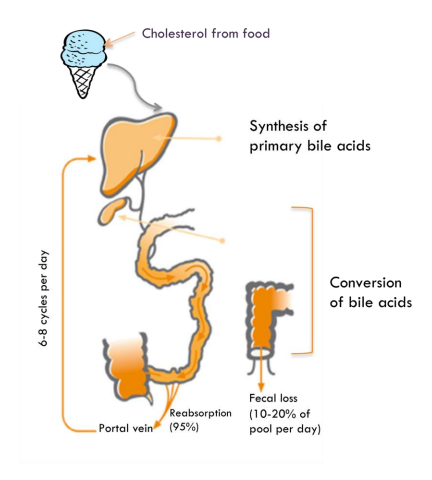


Figure 2.2, Bile acids metabolism

Under the physiological conditions, bile acids are only present at low concentrations in the peripheral circulation due to the hepatic extraction and intestinal absorption. The excretion of bile acids from feces stands for 10% -15% of total daily bile acids production in human. In hepatobiliary and intestinal diseases, the cholesterol metabolism is disturbed, affecting the synthesis of bile acids and conversion of bile acids in different biological fluid (serum, bile, urine, and feces). The bile acids quantitative and qualitative changes in feces provide possible correlation with the development and prevention of liver diseases and intestinal diseases.

## 2.1.2. Inflammatory Bowel Diseases (IBD)

Inflammatory bowel disease (IBD), including Crohn's disease (CD) and ulcerative colitis (UC), significantly affects the quality of life of many patients in western world [4]. A standard surgical procedure called ileal pouch anal anastomosis (IPAA) is usually used for the treatment of UC. However, patients often develop inflammatory complications of ileal pouch after the surgery, among which, pouchitis is the most common and non-specific one [5](Fig. 2.3). The non-specific symptoms of pouchitis bring great challenges in the diseases diagnosis and management.

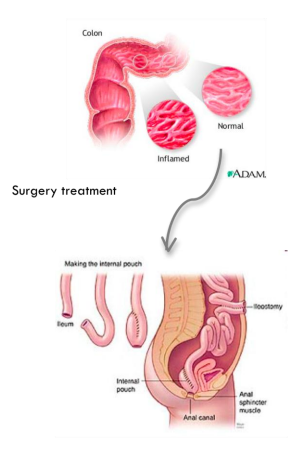


Figure 2.3, IBD and its treatment by IPAA surgery (http://www.adam.com/)

The pathophysiology of pouchitis is not completely understood yet, while one of the important factors could be the bacterial overgrowth in the ileal pouch reservoir [6]. In hepatobiliary and intestinal disorders, the abnormal cholesterol metabolism affects the bile acids synthesis, enterohepatic circulation, and biotransformation process, thus changing its composition in biofluids. Kruis *et al.* and Natori *et al.* reported that altered bacterial conversion of primary to secondary bile acids was related to abnormal intestinal microflora in UC patients [7, 8]. Therefore, monitoring the change of bile acids profile through bile acids quantification may serve as a useful diagnostic test for the ileal pouch diseases developed for the treatment of IBD.

# 2.2. Development of a quantitative bioanalytical method for fecal bile acids

### 2.2.1. Challenges in method development

Due to the complex composition and minor difference between bile acids components, the quantification of fecal bile acids is challenging. Most of the current methods use gas chromatography–mass spectrometry (GC-MS) for the analysis of bile acids in feces [9, 10]. Because of the comprehensive bile acids analysis requires the separation of free bile acids and the conjugated ones, the major disadvantage of the GC-MS approach is the tedious and time-consuming derivatization and pre-fractionation steps prior to analysis.

LC-ESI-MS is a powerful technique for the simultaneous analysis of multiple bile acids in biological materials including plasma, serum and urine with high specificity and selectivity [11,12]. One of the challenges is the separation for the dihydroxy  $C_{24}$  bile acids presenting isomeric structures. The detection of bile acids is often operated in SIM mode for free bile acids and MRM mode for glycine and taurine conjugates. There was also a report that utilized LC-ESI-MS to analyze fecal bile acids, but its accuracy and precision were not reported [13].

Matrix effect is another common problem associated with analysis of a complex biological sample with mass spectrometry. This phenomenon in ESI was observed when the target analyte is co-eluted with other bio-fluid components [14]. The endogenous phospholipids, proteins and fatty acids in the biological samples are the main source of the matrix effect. The ionization of analyte in mass spectrometry can vary greatly, which in turn significantly affects accuracy and precision of measurements [15,16]. The affected signal intensity can be explained as the competition among ions for the limited droplet surface charge [17]. For example, Scherer *et al.* reported the matrix effect during the analysis of bile acids in serum [18]. We also encountered serious matrix suppression problem during the course of our own analysis of fecal materials.

In this work, we have developed a simple and effective method termed as standard addition with internal standard (SA-IS) method to overcome the large quantitative errors brought by matrix effect. Combining standard addition with internal standard, the SA-IS method takes advantages of both, which compensates matrix effect and variations in sample preparation and MS detection. In this study, we studied seven bile acids CA, CDCA, DCA, UDCA, LCA, taurochenodeoxycholic acid (TCDCA), and glycochenodeoxycholic acid (GCDCA) using the SA-IS method along with a simple sample pretreatment procedure. The sensitivity, accuracy, reproducibility of this method were evaluated and improved to the acceptable range for the quantitative determination of bile acids in fecal materials such as pouch aspiration. We expect that the SA-IS approach developed in this work can be a general method for the quantitative analysis of other complex samples by LC-ESI-MS, in which the matrix effect exists.

## 2.2.2. Chemicals and methods

2.2.2.1. Chemicals

CA, CDCA, DCA, UDCA, LCA, GCDCA, TCDCA, N-1-napthylphthalamic acid (NPA), ammonium hydroxide and ammonium acetate were purchased from Sigma-Aldrich Chemical (St Louis, MO, USA). HPLC-grade methanol and acetonitrile were also purchased from Sigma-Aldrich. Deionized water was generated by a Millipore water purification system (Billerica, MA, USA). Syringes and syringe filters were obtained from VWR international (Wester Chester, PA, USA).

#### 2.2.2.2. Sample collection

The pouch aspiration used in this study is one kind of homogenous fecal materials. Diagnostic or surveillance pouch endoscopies, a subspecialty of the Pouthitis Clinic at Cleveland Clinic (Cleveland, OH, USA), were performed as a part of routine clinical care. The project was approved by Institutional Review Board at Cleveland Clinic. Informed consent was obtained from all patients. Before pouch endoscopy, each enrolled patient was given one Fleets®enema, and in 5–10 min the patient went to empty the pouch. Pouch aspiration samples were collected during pouch endoscopy and stored at  $-20^{\circ}$ C.

#### 2.2.2.3. Instrumentation and LC-MS conditions

The LC-MS system consisted of an Agilent 1100 series HPLC (Santa Clara, CA, USA) and a Waters Micromass Quattro II triple quadrupole mass spectrometer (Manchester, UK). A Luna C18 column (150 mm, 2mm i.d., 5  $\mu$ m) with a guard column (40 mm, 2mm i.d.) from Phenomenex (Torrance, CA, USA) was used for the chromatographic separation. The mobile phase was a gradient mixture of 10 mM ammonium acetate– ammonium hydroxide buffer at pH 8.0 (A) and 10mM ammonium acetate in acetonitrile-methanol 3:1 (B). The gradient elution program is illustrated in Figure 2.4. The HPLC column was first equilibrated with an initial mobile phase of 30% B. The mobile phase was then increased to 65% B within 6 min and increased to 72% B from 6.1 to 14 min. Afterward the mobile phase was changed to 90% B in 1 min and maintained for 5 min. At 20 min, the proportion was adjusted back to the initial ratio condition of 30% B and maintained for 10 min to re-equilibrate the column. During the entire analysis the flow rate was 200  $\mu$ L/min. The injection volume was 10  $\mu$ L and the total run time was 30 min for each sample including the re-equilibration.

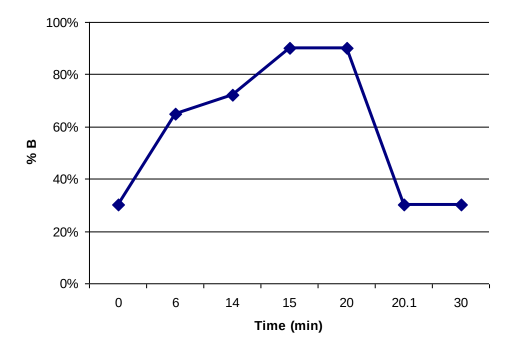


Figure 2.4, The gradient elution for the separation of bile acids

Negative ESI mode was adopted for the mass spectrometer operation. A small portion of flow (50  $\mu$ L/min) was introduced to the mass spectrometer with a post-column split ratio of 1:4. Direct infusion of each bile acid at 1  $\mu$ mol/L and the internal standard (IS) at 10  $\mu$ mol/L was used to fine tune the mass spectrometry conditions. The ion source temperature was maintained at 95 °C and the capillary voltage was set at 3.0 kV. Nitrogen nebulization and drying gas were held at 12 and 300 L/h, respectively. Cone voltage was at 50 V. Detection of bile acids was performed using selective-ion monitoring (SIM) mode. The deprotonated molecules of the free and conjugated bile acids were recorded at m/z 375.2 (LCA), 391.3 (UDCA, CDCA and DCA), 407 (CA), 448 (GCDCA), and 498 (TCDCA). Micromass Masslynx (Version 3.3) was utilized for system operation, data acquisition, and data processing.

#### 2.2.2.4. Stock and working solutions

Stock solutions of each bile acid and IS were prepared at a concentration of 14 mmol/L by carefully weighting each compound and dissolving them in methanol. Stock solutions were stored under  $-20^{\circ}$ C. The dilution buffer, 70% mobile phase B, was used for the preparation of bile acid working solutions at 20, 40, 80, 100, 200, 400, 800, 1000, 1600, and 2000 µmol/L per compound. The one with the highest concentration was prepared by mixing equal volumes of seven bile acid stock solutions together; other lower concentrations were obtained by serial dilution. The IS working solution at 4000 µmol/L was prepared by diluting the IS stock solution.

2.2.2.5. Pouch aspiration calibrators and Quality Control (QC) samples

The collected pouch aspiration samples were homogenous and thawed before use. The blank pouch aspiration lots, which had no detectable ion signals for given bile acids, were obtained from different IBD patients. The calibrators and QC samples used different two lots of blank pouch aspiration. Bile acids calibrators of 0.05, 0.1, 0.25, 0.5, 1, 2.5 and 5  $\mu$ mol/L were obtained by spiking 15  $\mu$ L bile acid working solutions of 20, 40, 100, 200, 400, 1000 and 2000  $\mu$ mol/L into 30  $\mu$ L blank pouch aspiration, respectively. In the same way, bile acids QC samples of 0.2, 2 and 4  $\mu$ mol/L were prepared by spiking 15  $\mu$ L bile acids working solutions of 80, 800 and 1600  $\mu$ mol/L into 30  $\mu$ L blank pouch aspiration, accordingly. There is a 400 times' concentration difference between the stock solution and wanted final concentration of bile acids considering the sample treatment procedures described in the following section.

## 2.2.2.6. Sample treatment of calibrators and QC samples for internal calibration

After thawing the pouch aspiration calibrators and QC samples to room temperature from -20 C, they were homogenized using a mixer. Prior to sample extraction, each aliquot of 45 µL calibrators and QC samples was mixed with 15 µL IS working solution for the internal standard calibration. For the sample preparation, each of the mixed samples was added with ethanol to obtain a total volume of 600 µL. The whole mixture was sonicated by an ultrasonic bath for 15 min. After a centrifugation at 10,000 rpm for 5 min, 100 µL supernatant was removed and diluted 10 times using the dilution buffer. The solution was

then filtered using a syringe filter with a pore size of 0.45  $\mu$ m. Finally, the filtrated solution was injected into LC-MS system for analysis.

2.2.2.7. Sample treatment of QC and patient samples for SA-IS

Two portions of QC sample and patient sample were needed for each analysis when applying the SA-IS method. The QC sample analysis required each portion as 45  $\mu$ L and the patient samples analysis required each portion as 30  $\mu$ L. Before the sample extraction, one portion was spiked with 15  $\mu$ L IS working solution, and the other was spiked with both the 15  $\mu$ L IS working solution and 15  $\mu$ L bile acid working solution at 800  $\mu$ mol/L. The following sample preparation procedure was the same as that described in the section "Sample Pretreatment of Calibrators and QC Samples for Internal Calibration".

2.2.2.8. Matrix effect and recovery

For the evaluation of matrix effect in pouch aspiration, the chromatographic peak area ratios of bile acids to IS in matrix-contained solution were compared with those for the non-matrix contained solution. The matrix-contained samples were prepared as follows: a 30  $\mu$ L blank pouch aspiration sample was extracted, diluted, and filtrated by the procedure described above. Further, bile acids and IS working solution were spiked into the post-extraction (SPE) solution. The non-matrix contained solutions of bile acids were the reference samples and were prepared by diluting bile acids work solution using

dilution buffer. The matrix effect was determined in two different lots of blank pouch aspiration under three concentration levels: 0.2, 2 and 4 µmol/L.

For the determination of extraction recovery, the chromatographic peak area ratio of bile acids to IS in the spiking-before-extraction (SBE) solution was compared with that in the SPE solution. The preparation method of bile acids and IS in the SBE solution was the same as preparation of QC standards. The recovery was also determined under three concentration levels: 0.2, 2 and 4  $\mu$ mol/L.

#### 2.2.2.9. Method validation

After the development of this SA-IS method, we validated it by assessing the linearity, limit of detection (LOD), lower limit of quantification (LLOQ), and the intra- and interassay precision and accuracy. Bile acids calibration curves were constructed over the range of 0.05-5  $\mu$ mol/L. The peak area ratios of bile acids to the IS in the pouch aspiration sample (y) were regressed against the concentration of spiked standard bile acids (x).

The LOD and LLOQ samples were prepared by spiking 15  $\mu$ L bile acid working solutions with 30  $\mu$ L blank pouch aspiration. The LOD for bile acids in pouch aspiration was determined when the signal-to-baseline noise ratio was above 3. The LLOQ was the lowest concentration of bile acids in pouch aspiration that could be determined within a precision and an accuracy of 20%.

Intra-assay precision and accuracy were determined in five replicates of QC samples at 0.2, 2 and 4  $\mu$ mol/L prepared in the same run. Inter-assay precision and accuracy were measured in three different runs. The precision was expressed in terms of percentage coefficient of variation (CV%) and accuracy was expressed in terms of percent relative errors (RE%).

#### 2.2.2.10. Method application

To test the feasibility of this SA-IS method in fecal material analysis, we determined the bile acids profiles for a preliminary clinical application. The clinical pouch aspiration samples were collected from the patients diagnosed as having pouchitis and normal pouch after IPAA surgery.

#### 2.2.3. Results and discussion

#### 2.2.3.1. Separation of bile acids

A broad range of the bile acids physicochemical properties including PKa and hydrophobicity, along with the minor difference between some isomeric bile acids, bring great challenges in bile acids separation. The free and conjugated bile acids have different PKa range at  $3.9 \pm 0.1$  and  $5.0 \pm 0.1$ , respectively [19]. The Log D values of bile acids distribute from 0 to 4 for all bile acids [20]. Among these seven compounds, UDCA and

CA are the most hydrophilic bile acids, while DCA and LCA are the most hydrophobic ones.

The separation of bile acids CDCA, UDCA, DCA, LCA, CA, TCDCA, and GCDCA was achieved with a Luna C18 column from Phenomenex. The isomeric bile acids CDCA and DCA were the most difficult to separate. The mobile phase was optimized to achieve complete separation of these two isomers by using different ratio of methanol, acetonitrile, and mobile phase additives. The addition of methanol to the organic mobile phase could sharpen bile acids peaks [21]. The retention times and sensitivity of bile acids were improved by adding ammonium salts into the mobile phase is basic, the variation of log P on the change of pH is the lowest, and consequently the deprotonation of bile acids is promoted. Three different concentrations of ammonium acetate: 2, 10, and 20 mM were tested in the mobile phase. We found that 10 mM ammonium acetate yielded the highest ionization intensity and best separate the analytes within 20 min and to sufficiently isolate three isomers of UDCA, CDCA, and DCA, as well as other bile acids (Fig. 2.5).

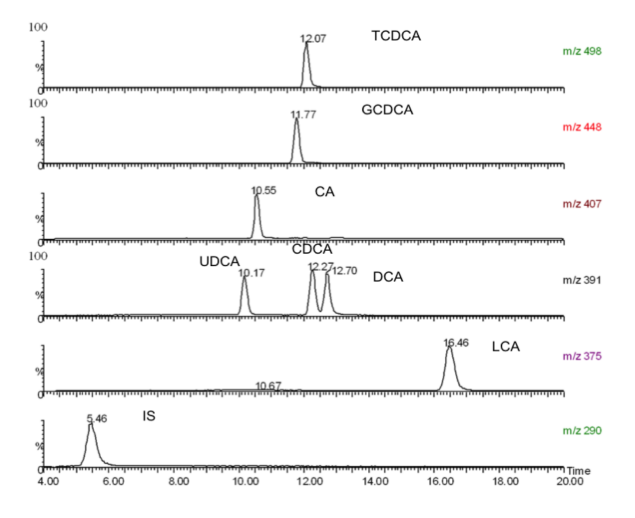


Figure 2.5, Representative mass chromatograms of bile acids at 0.1  $\mu mol/L$  and IS NPA at 10  $\mu mol/L$ 

## 2.2.3.2. Matrix effect

The variation of analyte signal after the extraction process for the biological sample is often contributed by extraction recovery and matrix effect. Recovery variation is mainly caused by the extraction efficiency, while matrix effect is introduced by co-eluting interference existing in the extracted solution. When the matrix interference shows higher proton affinity than targeted analyte, proton is transferred from the ionized analyte to the interference, causing the lost ion intensity of partial analyte [14]. It has been reported that, when LC-MS was used to analyze biological samples such as urine and plasma, matrix effect often occurred and caused deterioration of the precision and accuracy of the analysis [22-24].

The evaluation of matrix effect during LC-MS analysis usually employed the following methods: post-column infusion, post-extraction spike, and the comparison of standard calibration slopes. In this study, the post-extraction spike method and the comparison of standard calibration curves were employed to assess the matrix effect on bile acids in pouch aspiration samples.

NPA was selected as IS to normalize MS signal for all bile acids because of their similar structures and chemical properties [25](Fig. 2.1). After the normalization of MS variation by the IS, the matrix effect (ME %) was measured by comparing the peak area ratios of

bile acids to IS spkied in post-extraction solution and with those in dilution buffer according to [26]:

$$ME\% = (1 - \frac{R_{SPE}}{R_{DB}}) \times 100\%$$

where  $R_{SPE}$  represents the peak area ratio determined in the SPE solution of blank pouch aspiration and  $R_{DB}$  represents the peak area ratio determined in the dilution buffer. The positive ME% indicates matrix suppression of target analytes; the negative ME% indicates an enhancement of ionization by sample matrix; and zero of ME% indicates no matrix effect is present. In addition, we compared the ME% from two different batches (No.398 and No. 957) of blank pouch aspiration at three concentrations: 0.2 µmol/L, 2 µmol/L, and 4 µmol/L.

In addition, three calibration curves of bile acids were established in the non-matrix contained solution, matrix contained solution after extracting blank plasma sample lot 1, and matrix contained solution after extracting blank plasma sample lot 2 for matrix effect evaluation also. The slopes were compared between three calibration curves.

As shown in Table 2.1, the ME% in sample 398 (except for 4 µmol/L CDCA and DCA) ranged from 7.6 to 38%, indicating severe ion signal suppression by the pouch aspiration sample matrix. In sample No. 957 the ME% was even larger. These results were confirmed in the results from the calibration curves for bile acids in different matrix solutions (Fig. 2.6). The slopes of calibration curves established in non-matrix contained solution were higher than that in two different matrix-contained solutions. In addition, the level of ion suppression resulting from the pouch aspiration matrix varied greatly

between different samples. This is not surprising due to the highly complex nature of fecal materials such as pouch aspiration.

Bile Acids	Concentration (µmol/L)	No. 398 ME (%)	No. 957 ME (%)
	0.2	30.6	45.1
CA	2	27.2	41.8
	4	11.8	29.9
	0.2	23.5	37.2
CDCA	2	23.9	37.3
	4	-1.2	18.9
	0.2	29.2	30.9
UDCA	2	19.2	28.3
	4	7.6	21.7
	0.2	28.9	37.3
DCA	2	21.1	31.7
	4	-0.3	21.6
	0.2	37.1	47.3
LCA	2	24.2	18.4
	4	9.7	13.3
	0.2	38.0	49.3
TCDCA	2	24.7	36.4
	4	8.3	26.7
	0.2	35.0	38.1
GCDCA	2	25.1	29.0
	4	10.7	22.4

Table 2.1 Matrix effect determined in blank pouch aspiration sample No. 398 and No. 957 for bile acids at three concentrations (n=3)

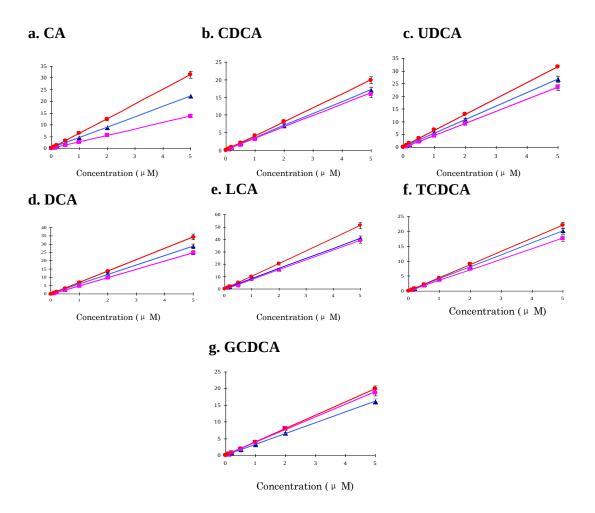


Figure 2.6, Calibration curves of bile acids: CA (a), CDCA (b), UDCA (c), DCA (d), LCA (e), TCDCA (f), and GCDCA (g) established in different solution sets: non-matrix contained solution (• in orange), extracted solution of blank pouch aspiration lot #1 ( $\blacktriangle$  in blue), and extracted solution of blank pouch aspiration lot #2 ( $\blacksquare$  in purple)

#### 2.2.3.3. Internal standard calibration

To evaluate the availability of internal calibration for compensating matrix effect and determining bile acids in pouch aspiration, calibration curves were established from pouch aspiration calibrators. The calibrator and QC standards were prepared from two different batches of blank pouch aspiration samples. The QC standards of three spiked concentrations were measured based on the calibration curves, and the results were listed in Table 2.2.

For most of the bile acids, the analytical accuracy with the internal standard method was low considering RE% ranges from -50.4% to 64.6%. These large quantitative errors resulted from the sample-to-sample variations in the matrix components. More specifically, such great errors were caused by significant differences between the level of matrix effect of QC standards and that of calibrators. Therefore, the internal standard calibration was not suitable for bile acids analysis when the matrix varied among samples.

Bile Acids	Spiked Concentration (µmol/L)	Accuracy RE(%)
	0.2	64.6
CA	2	22.7
	4	21.3
	0.2	-17.1
CDCA	2	-50.4
	4	-23.5
	0.2	39.8
UDCA	2	5.5
	4	-13.6
	0.2	36.3
DCA	2	7.4
	4	22
	0.2	15.6
LCA	2	-6.4
	4	5.2
	0.2	7.4
TCDCA	2	-17.3
	4	15.4
	0.2	-19.5
GCDCA	2	-45.3
	4	20.6

Table 2.2 The accuracy results determined by internal standard method (n=5)

#### 2.2.3.4. SA-IS method

Several approaches were reported to reduce matrix suppression [15,16,27]. One of most effective approaches is the use of isotope-labeled analogues as internal standards. However, the presence of multiple analytes in pouch aspiration requires multiple isotope-labeled internal standards, which are costly and difficult to obtain.

Using appropriate sample extraction procedure is another approach to reduce the matrix effect. We tried to clean up the co-eluting substance by LLE and SPE. However, since the hydrophobicity of bile acids has a broad range, it is difficult to develop an LLE method ensuring low matrix interference and good recovery. Also the large number of lipids and lysophospholipids, which are similar with bile acids with regarding to structure and polarity, presents in pouch aspiration, causing the co-elution of bile acids and the interference when using SPE methods. In addition, the matrix interference, with significant sample-to-sample variation, could not be removed completely by the same sample cleanup method. As a result, all of these purification attempts failed to reduce the matrix effect within  $\pm 15\%$  for all the bile acids.

Standard addition was another option reported to compensate the matrix effect in LC-MS analysis [28,29]. It measures the concentration of analyte in an unknown sample by comparing the response difference between before and after the addition of a known amount of analyte to that of the original sample. However, standard addition alone cannot correct the loss of analytes associated with sample preparation. As a result, we

investigated the SA-IS method that combined standard addition with internal standard to measure the concentrations of bile acid components in pouch aspiration. The use of standard addition will eliminate adverse matrix effect on accuracy and precision, while the use of internal standard will compensate for incomplete extraction and variability in sample preparation and MS detection.

The SA-IS method is illustrated in Fig.2.7.  $C_{unk}$  represents the unknown concentration of a bile acid in a pouch aspiration sample;  $C_{SA}$  represents the spiked concentration of the bile acid in the same pouch aspiration after standard addition. The peak area ratios of the bile acid to IS in the original sample and in standard addition samples are  $R_{unk}$  and  $R_{SA}$ , respectively. The peak area ratios are regressed against the concentration of bile acid spiked in.  $C_{unk}$  is calculated according to:

$$C_{unk} = \frac{C_{SA} \times R_{unk}}{(R_{SA} - R_{unk})}$$

With the SA-IS method, each pouch aspiration sample was divided into two aliquots: the first one was spiked with only IS, and the other was spiked with the bile acids mixture and IS. The validation results showed that the SA-IS is effective in correcting the matrix effect and more accurate than internal standard method.

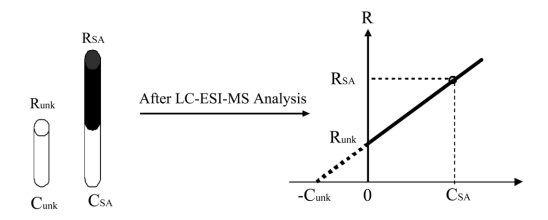


Figure 2.7, Illustration of SA-IS method

# 2.2.3.5. Method validation

**Linearity:** Two requirements are needed when applying the SA-IS: 1) a good linear relationship between the analyte concentration and according signal response; 2) zero signal response in the absence of analyte. The linearity were studied by analyzing the calibrators, which were blank pouch aspiration samples spiked with bile acids and IS. We observed that calibration curves established by plotting the peak area ratio of analyte to IS vs. the concentrations of each bile acid were linear within the range of 0.05  $\mu$ mol/L-5  $\mu$ mol/L with correlation coefficient of 0.9964 - 0.9999 (Table 2.3).

Bile Acids	Range	Calibration curve of standard bile acid in blank pouch aspiration sample			
	(µmol/L)	slope	y-intercept	correlation coefficient	
CA	0.05-5	4.331	0.063	0.9998	
CDCA	0.05-5	3.543	0.016	0.9964	
UDCA	0.05-5	5.163	0.012	0.9991	
DCA	0.05-5	6.129	0.018	0.9998	
LCA	0.05-5	7.782	0.023	0.9988	
TCDCA	0.05-5	3.869	0.076	0.9999	
GCDCA	0.05-5	3.129	0.048	0.9996	

Table 2.3 Calibration curve results for bile acids spiked in pouch aspiration

**Sensitivity:** The sensitivity of this method was assessed in term of the limit of detection (LOD) and the lower limit of quantification (LLOQ). For all the bile acids, the LOD was 1 nmol/L using a signal-to-noise ratio of 3. LOD was determined by spiking the bile acid standards into the blank pouch aspiration samples. The LLOQ was 0.05 µmol/L as the lowest end of the calibration in this study.

**Recovery:** We performed the recovery study by determining the peak area ratios of bile acids to IS in SPE solution and SBE solution. The percent recovery was calculated as:

Recovery% = 
$$\binom{R_{SBE}}{R_{SPE}}$$
 × 100%

Independent of the LC-ESI-MS interface and ionization method, the recovery measures the efficiency of bile acids extraction process during sample pre-treatment. The recovery of different bile acids varied from 91% to 109.8%, 86.7% to 99.8% and 91.3% to 111.4% under 0.2µmol/L, 2µmol/L and 4µmol/L levels, respectively (Table 2.4). This showed that our sample preparation method was adequate to recover bile acids from pouch aspiration.

Bile Acids	Concentration (µmol/L)	Recovery (%)
CA	0.2	109.8
	2	98.9
	4	108.5
CDCA	0.2	91.1
	2	86.7
	4	97.0
UDCA	0.2	105.7
	2	99.8
	4	91.3
DCA	0.2	91.0
	2	91.5
	4	99.4
LCA	0.2	102.7
	2	92.1
	4	107.6
TCDCA	0.2	102.5
	2	96.0
	4	109.6
GCDCA	0.2	105.8
	2	95.7
	4	111.4

Table 2.4 Recovery of bile acids at three concentrations (n=3)

Accuracy and Precision: Accuracy in terms of percent relative error was determined five replicates per concentration for three different QC samples of 0.2, 2 and 4  $\mu$ mol/L. The relative error of the SA-IS method was -7.0% to 12.1%, which was within the acceptable values for FDA guideline (± 20% for LLOQ and ± 15% for all QC samples) (Table 2.5). The intra- and inter-assay precision of this SA-IS method were investigated by five replicating analysis of the QC samples in triplicate run. The CV % ranged from 0.8% to 11.4%. The results have shown that this analytical method is accurate and precise for the quantification of bile acids in pouch aspiration within the range of 0.05 µmol/L-5 µmol/L.

Bile Acids	Spiked Concentration (µmol/L)	Accuracy RE(%) (n=5)	Intra-assay CV(%) (n=5)	Inter-assay CV(%) (n=3)
	0.2	11.1	3.9	4.2
CA	2	12.1	2.7	7.4
	4	2.7	5.6	6.2
	0.2	9.1	3.8	7.1
CDCA	2	9.7	2.2	5.8
	4	0.4	11.4	11
	0.2	-3.1	3.8	4.3
UDCA	2	7.8	5.8	9.5
	4	-1.5	7.6	5.6
	0.2	1.3	4.7	7.6
DCA	2	8.4	3.8	0.5
	4	-2	3.5	5.9
	0.2	2.1	4.3	7.5
LCA	2	1.9	1.5	2.9
	4	-7	4.6	4.6
	0.2	7.3	4.4	4.5
TCDCA	2	1.5	3.7	9.4
	4	-2.4	4.1	2.8
	0.2	8.8	5.7	4.2
GCDCA	2	7.6	3.1	2.8
	4	0.7	5.3	4.3

Table 2.5 Accuracy and intra- and inter-precision in the analysis of bile acids in pouch aspiration by the SA-IS method

### 2.2.3.6. Method application

During the treatment of UC, pouchitis is the most common and non-specific inflammatory disease developed in the built ileal pouch after IPAA surgery. Currently the diagnosis criteria of pouchitis are mainly based on clinical symptoms and endoscopic and histologic value inflammation [5]. To study of the bile acids profile for diseases developed after IPAA, we applied the SA-IS method to measure the concentration of bile acids in the pouch aspiration samples from the patients diagnosed as pouchitis (No. 570 and No. 588) and normal pouch (No. 839 and No. 232). The SIM chromatograms demonstrated that all bile acids peaks were clearly detected and sufficiently separated (Fig. 2.8). Comparing to normal pouch patients, pouchitis patients showed a relatively higher concentration of total bile acids and increased ratio of conjugated bile acids (Table 2.6). However, a broader investigation with more pouch aspiration samples will be required in future studies to determine whether the bile acids profile will become a new index for the diagnosis of pouchitis.

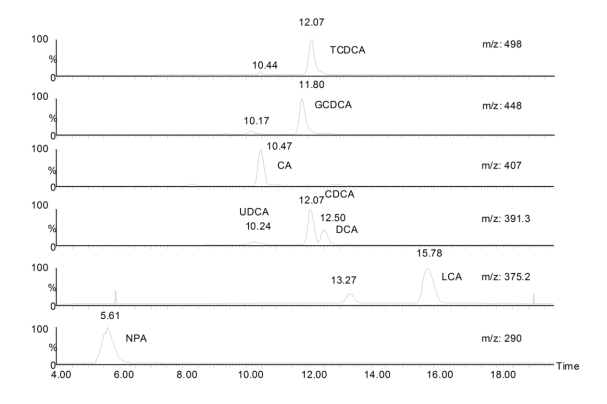


Figure 2.8, Chromatograms of bile acids in pouch aspiration sample No. 588

Bile Acids Class		Found concentrations (µmol/L)			
	Bile Acids	Pouchitis		Normal Pouch	
		No. 570	No. 588	No. 839	No. 232
	CA	125.5	119.1	114.8	143.5
Drimon	CDCA	84.7	124	82.2	120.2
Primary	TCDCA	26.9	48.8	15.3	26.4
	GCDCA	82.5	69.4	18.7	12.3
Secondary	UDCA	20.5	41.5	ND <sup>a</sup>	9.6
	DCA	78.2	58.2	35.6	17.6
	LCA	13.6	35.9	23.8	37.9
Total Bile Acids (µmol/L)		431.9	496.9	290.4	367.5
Secondary Bile Acids (%)		26	27.2	20.5	17.7
Conjugated Bile Acids (%)		25.3	23.7	11.7	10.5

Table 2.6 Bile acids concentrations in four pouch aspiration samples of patients diagnosed as pouchitis and normal pouch after IPAA

<sup>a</sup> Not detected

The SA-IS method developed in this work can potentially be used to study bile acids profile in patients with other types of IBD, from whom fecal materials can be collected during the routine colonscopic surveillance. In addition, the SA-IS method is accurate and robust, thus it can serve as a reference for researchers to develop other technologies to profile bile acids in fecal materials.

#### 2.3. Conclusion

Severe matrix effect was observed when quantifying bile acids in pouch aspiration by LC-ESI-MS. Ionization suppression caused by co-eluting matrix components varied greatly among samples, and it introduced large errors for the measurement using the internal standard method. In this study, a standard addition coupled with internal standard method has been developed and validated to solve the matrix effect problem. We demonstrated that this SA-IS method could effectively correct the ionization suppression effect caused by matrix molecules, thus significantly improve the accuracy of measurements. To the best of our acknowledge, this is the first study utilizing the combination of standard addition and internal standard to correct matrix effect during the LC-MS analysis. Compared with existing methods, the SA-IS method involves internal standard and simple sample preparation. The method was validated to ensure high sensitivity, accuracy, and precision. We have successfully demonstrated its potential use in analysis of bile acids in pouch aspiration. The SA-IS method can be a general method for bioanalysis in the presence of matrix effect.

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# **CHAPTER III**

# THE DEVELOPMENT OF A ROBUST AND SENSITIVE LC-MS/MS METHOD FOR THE QUANTIFICATION OF AN ANTI-CANCER AGENT IN RAT PLASMA

# 3.1. Introduction of anti-cancer agent JCC76

In United States, more than 40,000 women die each year from metastatic breast cancer. Overexpression of human epidermal growth factor receptor 2 (Her2) occurs in about 25%-30% metastatic breast cancer. As a member of ErbB family of receptor tyrosine kinases [1], HER2 is preferred for ligand binding and the receptor dimerization activated multiple downstream signaling cascades, which promote cellular proliferation, survival, migration, invasion, and differentiation. HER2 overexpressed breast cancer has an increased tendency for metastasis and leads to a relative resistance to same cytotoxic and hormone therapy [2,3]. Therefore, HER2 tumors are considered to be more aggressive and often have poor prognosis [4]. Previous research showed that the level of HER2 expression was closely related to the tumor growth, indicating HER2 as on of the most valuable targets for breast cancer therapy [5,6].

Nimesulide is a nonsteroidal anti-inflammatory drug (NSAIDs) that inhibits cyclooxygenase-2 (COX-2) activity. Besides its anti-inflammatory activity, nimesulide has been reported to inhibit proliferation and induce apoptosis of a variety of human cancer cell lines including lung, ovarian, and breast cancer [7-9]. Other research demonstrated that the novel sulfonanilide derivatives of nimesulide significantly increased the activity in inhibiting breast cancer cell growth in comparison to that of nimesulide [10-11]. JCC76 {Cyclohexanecarboxylic acid [3-(2,5-dimethyl-benzyloxy)-4-(methanesulfonylmethyl-amino)-phenyl]-amide} is a novel compound deviated from nimesulide without COX-2 inhibitory activity [12]. Recently, this lead compound JCC76 was found to have potent activity against human epidermal growth factor receptor 2 (HER2) overexpressing breast cancer [10].

In the *in vitro* study, JCC76 was found to dramatically inhibit HER2 overexpression cell proliferation (i.e., SKBR-3, BT474 and MDA-MB-453 cell lines) [13]. In addition, this compound induced apoptosis in HER2 overexpressing cells (i.e., SKBR-3), but it was less active in HER2 negative cells (i.e., MCF-7 and MDA-MB-231), which suggested the selective inhibition of HER2 overexpressing breast cancer cells [13]. In the *in vivo* study, JCC76 significantly decreased the size of breast tumor in the mice xenograft experiments [14]. Based on these results of pharmacological study, JCC76 demonstrated high potential to be a target therapeutic agent acting on HER2 positive breast cancer.

In order to provide insights into the drug developability for targeted breast cancer therapy, a quantitative method for JCC76 is needed for the further pharmacological and toxicological studies. To date, no analytical assay for JCC76 has been reported. The low concentration of JCC76 in blood requires the measurement to be sensitive and specific. Because of the multiple blood sampling from a single animal model during a short time, only a low volume of sample can be used for each analysis. In previous studies, the extraction methods for the class of COX-2 inhibitors, which have similar chemical structure with JCC76, employed large amount of sample volume (200 – 500  $\mu$ L) and solvent consumption (4-8 mL), and complex sample pre-treatment procedures of solid phase extraction for each analysis [15-17].

In this work, we have developed a single-step LLE method to clean up the sample matrix and to ensure high extraction recovery. The extraction procedure is fast and simple, and it consumes less sample volume and solvent. Furthermore, the present study provided a short time for each LC-MS/MS analysis and high sensitivity with a lower limit of quantification (LLOQ) of 0.3 ng/mL. Finally, we successfully illustrated the preclinical application of this method with a pharmacokinetics study of JCC76 in rats.

#### **3.2.** Material and methods

#### 3.2.1. Reagents and chemicals

JCC76 and IS (compound 75) were synthesized according to previously published procedures (Fig. 3.1) [18]. Their purities were higher than 97%, confirmed by NMR and HPLC analysis. Methanol, acetonitrile, ethyl acetate, and methyl tert-butyl ether (MTBE) of HPLC grade and ammonium formate were purchased from Sigma-Aldrich (St. Louis, MO, USA). N-hexane of HPLC grade was obtained from Pharmco-AAPER (Shelbyville, KY, USA). Deionized water was purified by Barnstead NANOpure® water purification system from ThermoScientific (Waltham, MA, USA). Pooled blank rat plasma was purchased from Lampire Biological Laboratories (Pipersville, PA, USA).

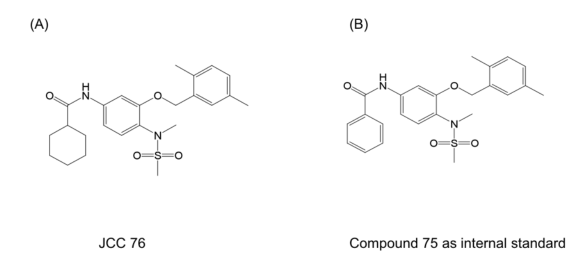


Figure 3.1, Chemical structures of JCC76 and the internal standard Compound 75

#### 3.2.2. LC-MS/MS instrumentation

The HPLC system consists of two LC-20AD pumps, a DGU-20A<sub>3</sub> degasser, a SIL-20AC autosampler, and a CBM-20A module (Shimazu, Tokyo, Japan). The chromatographic separation was performed on a Luna C18 column (2.0 mm × 150 mm, 5  $\mu$ m) with a guard column (2 mm × 40 mm, 5  $\mu$ m) from Phenomenex (Torrance, CA, USA). The mobile phase was a mixture of aqueous ammonium formate (pH 3.7; 5 mM)-methanol (1:9, v/v). Isocratic elution at a flow rate of 0.2 mL/min was employed. The injection volume was 5  $\mu$ L and each run time was 5 min.

The mass spectrometric detection was performed on an AB Sciex QTrap 5500 system (AB Sciex, Toronto, Canada) with positive electrospray ionization (ESI<sup>+</sup>). The multiple reaction monitoring (MRM) function was used to measure the transition of m/z 445 to 366.3 and m/z 439.3 to 360.3 for JCC76 and IS, respectively. The optimized parameters for detecting JCC76 and IS were set as following: the ion spray voltage was 5500 eV; the temperature was at 450 °C; the heating gas (GS1), nebulization gas (GS2), and curtain gas (CUR) were 30, 40, and 45 psi, respectively. Compound parameters, including declustering potential (DP), entrance potential (EP), collision energy (CE), and collision exit potential (CXP) for both JCC76 and IS, were set at 60, 10, 21, and 10 V, respectively. Data acquisition and quantitation were performed using Analyst software version 1.4.2.

#### 3.2.3. Preparation of calibration standards and QC samples

The stock standard solutions of JCC76 and IS were prepared by dissolving each compound in acetonitrile at 1 mg/mL and stored at -20°C. One set of JCC76 working solutions at 3, 10, 20, 100, 200 and 1000 ng/mL, was prepared by serial dilution from the stock solution with water-acetonitrile (1:1, v/v), and then used for preparing the calibration standards. Another set of JCC76 working solutions at 3, 9, 90, and 900 ng/mL was made in the similar way, and used for preparing QC samples. The working solution of IS was prepared by diluting the IS stock solution to 50 ng/mL. All of the working solutions were freshly prepared before use.

The calibration standards were prepared by spiking 5  $\mu$ L of JCC76 working solutions into 45  $\mu$ L blank rat plasma to give the final concentration of JCC76 at 0.3, 1, 2, 10, 2 and 100 ng/mL. The QC samples were prepared in a similar way at 0.3, 0.9, 9 and 90 ng/mL, representing lower limit of quantification (LLOQ), low QC (LQC), middle QC (MQC) and high QC (HQC) of JCC76 in plasma, respectively. All of the calibration standards and QC samples were further treated in the same sample preparation procedure described below.

#### 3.2.4. Sample preparation

Aliquots of 50  $\mu$ L rat plasma sample, from the calibration standards, QC samples and pharmacokinetics study samples, were mixed with 5  $\mu$ L IS working solution (50 ng/mL). After vortexing for 10 seconds, the samples went through a single step liquid-liquid extraction with 500  $\mu$ L of an MTBE-hexane mixture (1:2, v/v). After vortexing for 60 s,

the mixture was centrifuged at 10,000 rpm for 5 min. Then 420  $\mu$ L supernatant was separated, and evaporated using a centri-vap vacuum evaporation system (Labconco, MO, USA). The dry residues were reconstituted in 42  $\mu$ L 50% acetonitirle for LC-MS/MS analysis.

#### 3.2.5. Method validation

**Calibration, sensitivity, and selectivity:** Calibration curves were constructed by using the peak area ratios of JCC76 to IS (y) versus concentrations of JCC76 (x) in the calibration standards. The weighted linear regression was generated by using 1/x as weighting factor. The LLOQ was determined as the lowest concentration in calibration curve that can be quantified with the accuracy and precision within 20%. The selectivity of this method was evaluated by testing the presence of the interfering peak in blank plasma samples from six different sources.

**Matrix effect and recovery:** The absolute/relative matrix effect and recovery were studied at three QC levels: 0.9, 9 and 90 ng/mL. The absolute matrix effect was determined by comparing the peak area of analyte spiked in the post-extraction solution of blank plasma with those of standard solution at equivalent concentration. The relative matrix effect was studied by comparing the peak area ratio of analyte and IS spiked in the blank plasma post-extraction solution with that in standard solution. The post-extraction solution solution was prepared by extracting blank plasma using procedures in section 2.4.

The absolute recovery was determined by comparing the peak area of JCC76 spiked in plasma before extraction with that in post-extraction spiked sample. The relative recovery was determined by comparing the peak area ratio of JCC76 to IS spiked in plasma before extraction with that in post-extraction spiked sample.

Accuracy and precision: Intra- and inter-assay precision and accuracy were assessed using QC samples at four different concentrations: LLOQ, LQC, MQC and HQC. Intraassays were carried with five replicates (n=5) for each concentration in the same day, while inter-assays were performed with five replicates (n=5) in different days. The precision results were expressed as percent relative standard deviation (% RSD) and the accuracy results were expressed as percent relative error (% RE).

The dilution QC was also prepared to study the accuracy and precision in cases when samples concentration was above the highest concentration of the calibration curve. The dilution QC samples (n=5) contained were prepared by spiking JCC76 into blank rat plasma at 900 ng/mL, and then were 10 times diluted with blank rat plasma before extraction. Following the same sample preparation procedures, the dilution QC samples were analyzed and their concentrations were compared with the nominal concentration.

**Stability:** The storage stability was investigated with blank plasma spiked with JCC76 at LQC level (0.9 ng/mL) and HQC level (90 ng/mL) in triplicates going through the following conditions: sitting in room temperature for 4, 6 and 24 hr, three freeze-thaw cycles, -20°C for 30 days, and post-extraction storage at 4°C for 24 hr. For the freeze-

thaw stability study, the spiked samples were subjected to three freeze (-20°C)-thaw (room temperature) cycles and each cycle was 24 hr.

3.2.6. Pharmacokinetics study

The feasibility of this quantitative method was tested through a pharmacokinetics study of JCC76 in rats. Male Sprague–Dawley rats (each weight 300 – 350 g) were purchased from Charles River Laboratories International (Spencerville, OH, USA). The animals were housed in a 12 h light/dark cycle room with free access to food and water for at least 7 days to adapt to the environment. All the animal experiment procedures were performed under the guideline approved by Institutional Animal Care and Use Committee at Cleveland State University.

Before the intraperitoneal(i.p) administration of JCC76 at a single dose of 5 mg/kg, animals were fasted overnight but with free access to water. Blood samples of 150  $\mu$ L each time point were collected from the saphenous veins and femoral veins into heparinized tubes at 0 hr (before drug administration) and at 0.25, 0.5, 1, 1.5, 2, 3, 4, 6, and 8 h after dosing. The blood samples were centrifuged immediately at 10,000 rpm for 5 min in room temperature. The plasma samples were separated and store at – 20 °C until analysis.

The concentration of JCC76 in rat plasma versus time profiles were analyzed to estimate pharmacokinetics parameters using WinNonlin® software version 5.2 (Pharsight Corporation, Mountain View, CA, USA).

#### **3.3. Results and Discussion**

#### 3.3.1. Mass spectrometric and chromatographic conditions

In order to optimize the MS parameters, we introduced standard JCC76 and IS in 90% methanol at 200 ng/mL through infusion into the mass spectrometer at 10  $\mu$ L/min. The mass spectra from full positive scan of JCC76 and IS showed the protonated molecules [M+H]<sup>+</sup> with m/z 445.3 and 439.3, respectively. The most abundant product ion after fragmentation was at m/z 366.3 for JCC76 and m/z 360.3 for IS (Fig. 3.2). The fragmentation reactions for both compounds were proposed as the loss of  $-SO_2CH_3$  by 79 u. As a result, the ion detection employed multiple reaction monitoring (MRM) mode with selecting the transition of m/z 445.3  $\rightarrow$  366.3 for JCC76 and m/z 439.3  $\rightarrow$  360.3 for IS. The collision energy, spray voltage, and ion-spray voltage were fine-tuned to obtain the highest MS response.

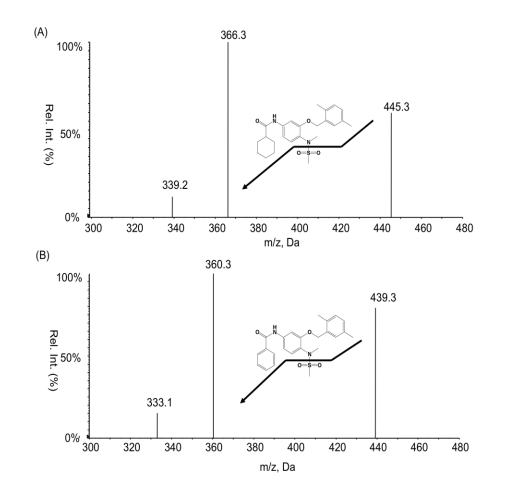


Figure 3.2, The precursor/product ion spectra and proposed fragmentation pathways for (A) JCC76 and (B) the internal standard compound 75

The solubility of JCC76 is poor in water with a predicted high Log D value of 4.86. Therefore, we chose to use a large portion of methanol in the mobile phase to elute the compound from a C18 column. Since the addition of organic acids promotes the protonation of the analyte under positive ESI, several mobile phase modifiers including formic acid, acetic acid, ammonium formate, and ammonium acetate with different concentrations were tested to optimize the chromatographic results. We observed that mobile phase consisting of 5mM ammonium formate-methanol (1:9, v/v) yielded the highest MS response of JCC76, which increased by 2 folds comparing water-methanol (1:9, v/v) without any modifier. The retention times were 3.04 min for IS and 3.5 min for JCC76 by the isocratic elution. Using the above mobile phase, the peaks of analyte and IS were symmetrical, the sensitivity was improved, and the total run time was controlled within 5 min.

### 3.3.2. Sample extraction

PPT has been used extensively for the preparation of plasma samples because of its simplicity [19]. To simplify our sample extraction technique, the PPT method for plasma cleaning-up was tested, but matrix suppression of 30%-33% was observed. Adding acetic acid or formic acid into rat plasma prior to the PPT pre-treatment could relieve the ion suppression of the analyte by 6-8%. However, it decreased the extraction recovery by 13-16%. LLE was reported to be more efficient and to provide cleaner extracts than PPT and the SPE for sample preparation in most cases [20]. Much effort was put into finding an

optimal extraction solvent with high extraction efficiencies and minimal matrix effect for JCC76.

We tested extraction solvents ranged from polar to non-polar, which included ethyl acetate, chloroform, methyl chloride, diethyl ether, MTBE, and hexane. Among theses solvents, non-polar ones including MTBE and hexane have much higher extraction recovery than other solvents for JCC76 (Fig. 3.3), which implies that the non-polar solvents favored the extraction for hydrophobic compounds. Regarding the matrix effect, the extraction with MTBE yielded the lowest ion suppression, which agreed with other reports that suggested MTBE is especially efficient in reducing matrix effect by removing the phospholipids in plasma [21]. Based on these observations, parallel extraction experiments based on different ratios of MTBE and hexane mixtures were performed. Finally, we observed that the optimized solvents mixture consisted of MTBE-hexane (1:2, v/v) (Fig. 3.4). This LLE method produced reproducible results of matrix effect less than 10% and high recovery above 90% for the extraction of JCC76 from rat plasma. The extraction procedure is simple, and it uses small amounts of solvent and sample volume.

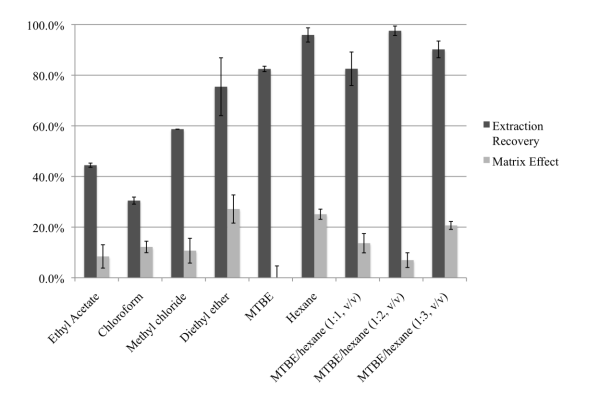


Figure 3.3, The comparison of matrix effect and recovery of JCC76 in rat plasma among different LLE solvents. Each column represents the mean  $\pm$  S.D. (n = 3)

#### 3.3.3. Method validation

#### 3.3.3.1. Linearity, sensitivity, and selectivity

JCC76 calibration curves were established using double blank (blank plasma sample with neither JCC76 or IS), zero blank (blank plasma with IS only), and six non-blank calibration standards at the concentrations of 0.3, 1, 2, 10, 20, and 100 ng/mL. The IS concentration in zero blank and the calibrations standards was 5 ng/mL. The peak area ratio of JCC76 to IS (*y*) versus JCC76 concentration (*x*) was plotted using 1/x as weighting factor. The linear regression equation (the slope and intercept in the mean  $\pm$  SD) obtained in five different days was  $y = (0.192 \pm 0.002)x + (0.00054 \pm 0.0006)$ . The linearity was excellent over the range of 0.3-100 ng/mL with the correlation coefficients above 0.9993 for all calibration curves built in different days. Accuracy and precision of all calibrators were within 15% (Table 3.1). As the lowest concentration on the calibration curve, the LLOQ was 0.3 ng/mL. It was sufficient to determine the concentration of JCC76 in rat plasma for pharmacokinetic study.

Nominal	Determined	A courses 0/	Precision %	
Concentration	Concentration	Accuracy %		
0.3	$0.29\pm0.007$	-3.3%	3.9%	
1	$0.98\pm0.017$	-2.3%	1.7%	
2	$1.89\pm0.138$	-5.8%	7.3%	
10	$10.15\pm0.084$	1.5%	0.8%	
20	$19.70\pm0.310$	-1.5%	1.6%	
100	$100.28\pm3.581$	0.3%	3.6%	

Table 3.1 Accuracy and Precision of JCC76 calibration standards over 0.3 - 100 ng/mL

The selectivity was investigated from blank plasma from six different sources. LC-MS/MS chromatograms of blank plasma, blank plasma spiked with JCC76 and IS, and a rat plasma sample after i.p. administration of JCC76 were compared (Fig. 3.4). No endogenous inferences at the retention times at 3.5 min for JCC76 and at 3.04 min for IS were found in six different blank plasma samples, indicating high selectivity and specificity of this method for the analysis of JCC76 in rat plasma matrix.

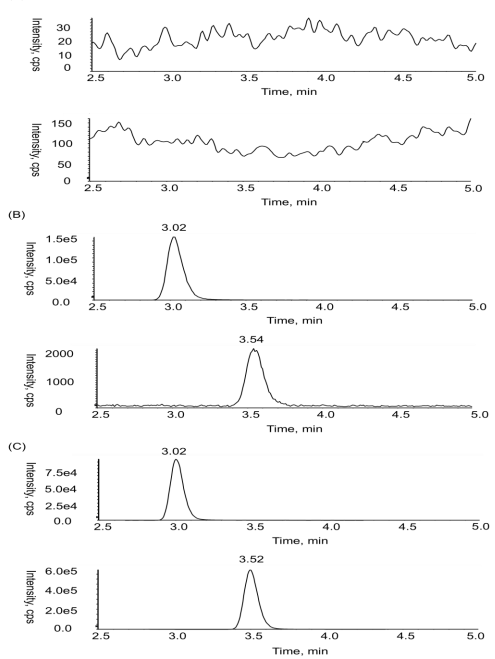


Figure 3.4, The MRM chromatograms of (A) blank rat plasma, (B) blank plasma spiked with JCC76 at LLOQ level (0.3 ng/mL 3.54 min) and IS (5 ng/mL, 3.02 min), and (C) a real rat plasma sample collected 8 hr after an i.p. administration of 5 mg/kg JCC76 spiked with IS

#### 3.3.3.2. Matrix effect and recovery

The matrix effect and extraction recovery was further evaluated at three levels: 0.9, 9, and 90 ng/mL. As shown in Table 3.2, the absolute matrix effect at three concentrations ranged from 3.1% to 9.2%, and the relative matrix effect ranged from -6.1% to 4.3%, indicating no obvious signal suppression or enhancement for the ionization of JCC76 in rat plasma matrix. The extraction of JCC76 at three levels showed absolute recovery of 89.5% to 97.3% and relative recovery of 105.1% to 106.1% after the IS normalization. The results indicated that the extraction procedure was not only sufficient to remove the interference impurities from the sample matrix.

	Matrix Effect		Recovery		
Concentration(ng/mL)	Absolute	Relative ME	Absolute	Relative	
(lig/lill2)	ME	Relative WIL	Recovery	Recovery	
0.9	3.1%	-6.1%	96.9%	106.1%	
9	8.8%	3.7%	89.5%	105.1%	
90	9.2%	4.3%	97.3%	105.1%	

Table 3.2, Absolute and relative matrix effect and recovery of JCC76 in rat plasma

3.3.3.3. Accuracy, precision, and dilution integrity

As the results shown in Table 3.3, the inter- and intra-assay accuracy and precision were within  $\pm$  10%, indicating the this method is accurate, precise, and reproducible. Since some samples containing JCC76 above the highest concentration of calibration curve, we also investigated the accuracy and precision after ten times dilution of dilution QC by blank sample matrix. The results showed that the intra- and inter-accuracy of the dilution samples were -0.1% and 2.1%, respectively. The intra- and inter-precision results were 0.8% and 1.6%. This result suggests that the plasma sample can be diluted and then analyzed when the concentration of JCC76 is above the upper limit of calibration.

	Intra-assay			Inter-assay				
Nominal	Determined	Accuracy	SD	Precision	Determined	Accuracy	SD	Precision
(ng/mL)	(ng/mL)	%RE	50	%RSD	(ng/mL)	%RE	50	%RSD
0.3	0.297	-1.1%	0.004	1.4%	0.301	0.3%	0.010	3.3%
0.9	0.870	-3.3%	0.010	1.1%	0.915	1.7%	0.072	7.9%
9	8.63	-4.1%	0.095	1.1%	9.04	0.5%	0.395	4.4%
90	84.2	-6.5%	1.272	1.5%	87.3	-3.1%	2.481	2.8%
900	899	-0.1%	7.483	0.8%	919	2.1%	14.618	1.6%

Table 3.3, Intra- and Inter- assay accuracy and precision of JCC76 in rat plasma

The results of the stability tests were summarized in Table 3.4. At room temperature, JCC76 was found to be stable for at least 24 h. The post-extraction stability study of JCC76 indicated its stability in the reconstitution solvent for at least 24 h. After three freeze-thaw cycles, the recovery of JCC76 was 78.2 % at LQC and 85.5 % at HQC levels. The long-term storage stability for JCC76 at -20 °C for 30 days was 89.2 % and 79.3% at LQC and HQC levels, respectively.

Concentration	Recovery	
(ng/mL)	%	
0.9	99.3%	
90	95.3%	
0.9	92.3%	
90	90.4%	
0.9	99.9%	
90	99.2%	
0.9	78.2%	
90	85.5%	
0.9	97.6%	
90	101.2%	
0.9	89.2%	
90	78.3%	
	(ng/mL) 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90 0.9 90	

# Table 3.4, Stability test of JCC76 in rat plasma

#### 3.3.3.5. Pharmacokinetic study

The feasibility of this method was tested by the application of a preclinical pharmacokinetics study in rats. The mean JCC76 concentration in plasma versus time profile was presented in Fig. 3.5. The maximum concentration of JCC76 ( $C_{max}$ ) in plasma was 528 ng/mL, which was reached at 0.5 h ( $T_{max}$ ) after dosing. The pharmacokinetics parameters were estimated through compartmental analysis and the concentration – time profile was found to fit a two-compartment model. The estimated AUC<sub>0-x</sub>, total body clearance, and volume of distribution were 1962 (ng\*h)/mL, 2.5 L/(h\*kg), and 7.0 L/kg, respectively.

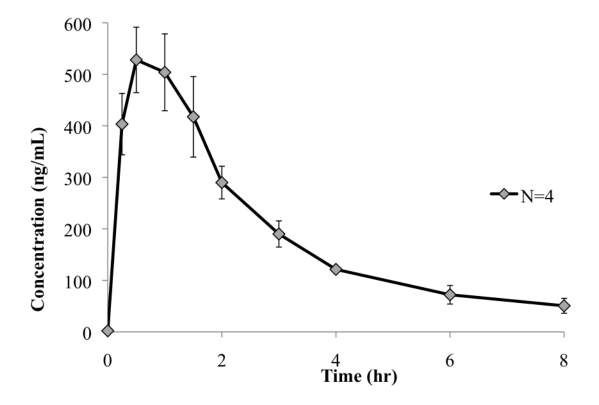


Figure 3.5, Mean plasma concentration-time profile of JCC 76 after the i.p. injection of JCC 76 at a single dose of 5 mg/kg. Each point represents the mean  $\pm$ S.D. (n = 4)

#### 3.4. Conclusion

In the presented work we have developed and validated a LC-MS/MS method for the quantification of JCC76 in a biological matrix for the first time. This method is simple, sensitive, and specific for the analysis of JCC76 in rat plasma. It used a one-step liquid-liquid extraction for sample preparation and a short time for LC-MS/MS analysis. The LLOQ of this method was as low as 0.3 ng/mL and the accuracy and precision were lower than 10%. The stability tests were performed under different storage and handling conditions and the results showed the suitability of this method for high throughput routine analysis. We have successfully applied this method in the determination of JCC76 in rat plasma for the pharmacokinetics study. This method will be further used in pharmacokinetics and toxicokinetics studies in animals in the future. It could be valuable for supporting the new drug investigation and application.

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#### **CHAPTER IV**

# PROTEOMICS STUDY FOR POTENTIAL BIOMARKER ANALYSIS OF OVARIAN CANCER

#### 4.1. Introduction

Ovarian cancer is the leading cause of death from gynecological malignancies worldwide. The lifetime risk of ovarian cancer for women is about 1.6% and the risk increases with age and deceases with pregnancy. The majority of ovarian cancers tend to present as advanced stage, resulting in as high mortality rate as 56% [1,2].

The International Federation of Obstetrics and Gynecology (FIGO) staging system describes the early stage (stage I and II) and advanced stage (stage III and IV) of ovarian cancer as follows: stage I disease involves one or both ovaries; stage II disease is defined as the spread tumor limited to pelvis; stage III disease involves the spread tumor with

peritoneal implants; and stage IV tumor is the distant spread tumor of metastasis [3, 4]. The early diagnosis of ovarian cancer can increase survival significantly. When the tumor is still confined to the ovary in early stage, detection and surgical removal of cancerous tissues result in cure for over 90% patients [3]. However, the ovary is not symptomatic in the stage I and II. Compared to breast, prostate, and colon cancer, ovarian cancer is anatomically more difficult to be assessed during physical examination due to the location of the ovary. Therefore, a very promising approach to improve the mortality rate for ovarian cancer is to discover reliable biomarkers and to develop adequate sensitive and specific screening test for early detection.

The biomarkers associated with the development of ovarian malignancy have been investigated in blood, tissue, and other biofluids using DNA microarrays and proteomics. With advantages in characterizing post-translation modifications of proteins, proteomics is believed to be one of the most attractive approaches for biomarker discovery. The technological obstacles of proteomics profiling for ovarian cancer include the lack of sensitivity of mass spectrometric detection, the mask of low-molecular weight proteins by the abundant proteins, and the discrimination of ovarian carcinoma from benign tumors.

The only approved serum biomarker CA-125 has been used for remission monitoring of ovarian cancer, but not for screening. It fails to reach the sensitivity and specificity for a screening test of early stage ovarian cancer detection. The elevated level of CA-125 was found in approximately 80%-85% of patients with advanced stage ovarian cancer, but in only 50% of patients in early stage [5]. In addition, elevated levels of CA-125 are also

associated with a variety of other conditions including other cancers (pancreatic, breast, bladder, liver and lung), benign and malignant breast and colon diseases, peritoneal irritants, and benign gynecologic diseases [6-8]. Nevertheless, new technologies in mass spectrometry for discovering other novel proteomics markers are emerging, aimed to improve the sensitivity and specificity of current screening tests.

Due to the dynamic range of proteins present in the serum, high abundance proteins were removed to increase the possibility of low molecular weight protein detection. Matrix-assisted desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS) has been used as an fundamental tool for proteomic study for its efficiency in detecting peptides as low as 1 femtomole [9]. In addition, there is no mass range limitation for the MALDI-TOF analysis and it is preferable for fast analysis, allowing 100 samples to be finished in less than 10 min [10]. For these major advantages, MALDI-TOF-MS was used for the quick scan of large biomolecules existed in serum samples from ovarian cancer patients. Our group reported differentiated mass spectra derived from cancer patient and normal sera by MALDI-TOF-MS analysis for three major peptide peaks of 1260, 1465, and 1545 Da [11] (Fig. 4.1).

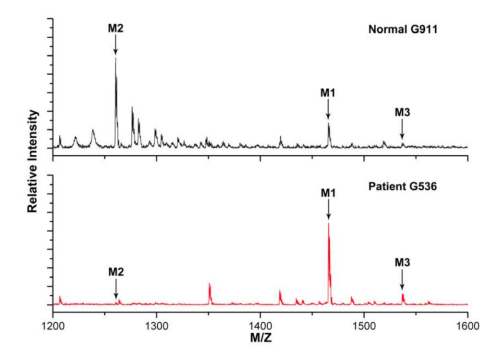


Figure 4.1, Previous reported analysis on MALDI-TOF-MS pattern for ovarian cancer patient and normal control samples [11].

Based on these previous findings, comparative analysis of low molecular weight protein profiling was further preformed by LC-MS/MS in this work. Several sets of samples from patients with ovarian carcinoma, benign tumors, and healthy normal controls were examined to identify carcinoma-associated proteins or peptides as valuable biomarkers for ovarian cancer progression. One of the major MS peaks, which discriminated normal control with cancer patients in MALDI-TOF scanning, was defined by LC-MS/MS analysis. The developed LC-MS/MS method greatly improved the reproducibility in peptides profiling. In addition, the protein fractionation method was refined in terms of higher extraction efficiency and less impurities. At last, the discovered biomarker was validated in benign and carcinoma samples for the improvement of early detection.

#### 4.2. Materials and Methods

#### 4.2.1. Materials

Acetonitrile and chloroform of HPLC grade, TFA, and formic acid were purchased from Sigma-Aldrich Chemical (St Louis, MO, USA). Deionized water was generated by a Millipore water purification system (Billerica, MA, USA). Phosphate-buffer saline (PBS) solution was purchase from BIO-RAD (Hercules, CA, USA).

#### 4.2.2. Patient samples

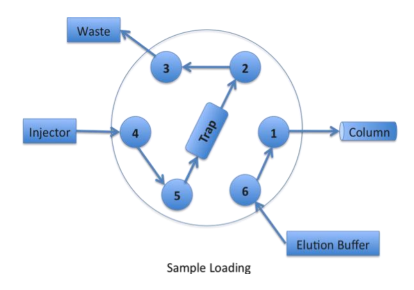
Human serum samples from healthy individuals were purchased from Sigma-Aldrich (St Louis, MO, USA). The serum samples from patients diagnosed with advanced stage ovarian cancer of carcinoma (N=40) and benign diseases (N=20) were obtained from Gynecologic Oncology Group (GOG) tissue bank (Buffalo, NY, USA). Among the forty carcinomas samples, there were twenty papillary serous carcinomas, ten mucinous carcinomas, five endometrioid carcinomas, and five clear cell carcinomas; As the twenty benign samples, there were ten serous benign, five mucinous benign, and five other benign. This project was approved by the Institutional Review Board at Cleveland State University.

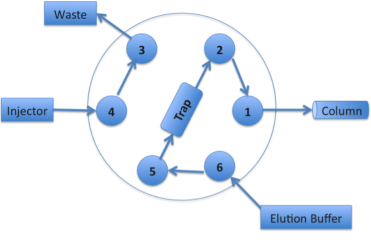
#### 4.2.3. Serum peptide fractionation

The low molecular weight peptides were separated from the abundant proteins in serum by protein precipitation. An aliquot of 40  $\mu$ L serum was mixed with 1 mL 90% methanol in water. After vortexing for 1 min, the mixture was centrifuged at 10,000 rpm by a Beckman Coulter Allegra<sup>TM</sup> X-22R Centrifuge for 5 min to precipitate the proteins. The clear supernatants were removed and then evaporated to dryness by centri-vap vacuum evaporation system (Labconco, MO, USA). The dry residues were re-dissolved in 400  $\mu$ L water. Afterward, the solution was cleaned by adding 200  $\mu$ L chloroform for liquid-liquid extraction. The chloroform phase and aqueous phase were well mixed by vortexing for 2 min. The mixture was centrifuged at 10,000 rpm for 5 min and the upper layer was carefully removed. After evaporating and concentrating the sample to 60  $\mu$ L, an aliquot of 10  $\mu$ L solution was injected to the LC-MS/MS system for analysis.

## 4.2.4. Online peptide trap setup

Prior to the micro-flow LC separation, the injected peptide sample was cleaned and desalted by an online peptide trap setup. As illustrated in Fig. 4.2, at the sample loading position, the trap peptide cartridge (CapTrap, Michrom) was connected with sample injector; at sample elution position, the peptide trap was connected with elution buffer pump and capillary HPLC column. After the sample was loaded into the peptide trap in 10 min, the column switch was changed to sample elution position. The switched elution buffer eluted the sample from the trap to the HPLC column.





Sample Elution

Figure 4.2, Online peptide trap setup

#### 4.2.5. Serum protein analysis by LC-MS/MS

The micro-LC-MS/MS system consisted of an Aglient 1100 HPLC, an online-extraction system, and a Bruker HCT 3000 plus ESI IonTrap Mass Spectrometer. The chromatographic separation of peptides was carried out on a Vydac C18 capillary reverse phase HPLC column (100 mm  $\times$  300 µm ID, 5 µm) at room temperature (23 °C). The mobile phase used 0.1 % formic acid in water: acetonitrile (99:1, v/v) as aqueous phase A and 0.1% formic acid in water: acetonitrile (1: 99, v/v) as organic phase B. The following gradient elution program was applied for all the peptide analysis: at the beginning of LC, 2% B was hold for 10 min, then increased to 90 % B in 70 min, following by maintaining 90% B for 20 min, and at last returned to 2% B for 10 min for re-equilibration. The flow was maintained at 5 µL/min through the analysis.

The mass spectrometer was operated in positive ESI mode. MS/MS data were acquired at a scanning mode of standard-enhanced at a range of 350- 1500 m/z. The nebulization gas, drying gas, and dry temperature were set as: 11 psi, 5 L/min, and 300 °C, respectively. The tandem MS data were used to search matched peptide fingerprints against NCBInr protein database by Mascot Program (http://www.matrixscience.com). The following searching parameters were applied: no trypsin, no taxonomy or modification, peptide mass tolerance is  $\pm 1.0$  Da, MS/MS tolerance is  $\pm 0.5$  Da, all possible charge states (i.e. + 1, +2, and +3), and a mass window between 0 and 100 kDa. To reduce the number of false-positive signals, only significant hits with at least 4 matching peptide masses were considered as final results.

# 4.3. Results

4.3.1. Identification of differentially expressed low molecular weight peptides

Based on the results of MALDI-TOF-MS pattern analysis reported previously [11], the clinical serum samples of 20 ovarian caner patient and 18 normal healthy controls generated a panel of biomarkers. By comparing the typical MS pattern, significant differences were observed regarding three peaks with the following m/z: 1465 Da, 1260 Da, and 1535 Da [11]. However, this method is poor in reproducibility and is not able to identify the peptide with corresponding MS spectra.

To compensate for this disadvantage, ESI-MS was used for improving the reproducibility in this study since no crystallization process during ionization is involved in ESI compared to MALDI. The mass spectrometer generated the spectra of both precursor ions and fragmentation ions, which were further analyzed by the computer to search for matching protein information in the database. Since HPLC was often necessary to separate the complex proteomic sample prior to the MS analysis, we conducted capillary LC to analyze pre-fractionated serum samples from ovarian cancer patient group and normal control group.

Initially, the mixture of six patient samples and the mixture of six ovarian cancer patient samples were analyzed. From the resulting chromatograms, we noted that the major difference between these two sample mixtures was a peak appearing at 42 min in cancer group but not in control group (Fig. 4.3). In the averaged MS spectra generated by this characteristic peak, the double charged MS peak of m/z 733 had the highest signal intensity. This finding was consistent with the result from previous MALDI-TOF-MS analysis, which showed the most abundance peptide peak at 1465 Da.

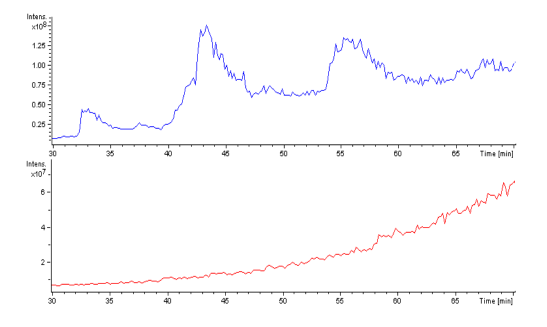


Figure 4.3, Differentiated LC-MS/MS chromatograms from ovarian cancer group (top) and control group (bottom)

Further, the mascot database searching was applied using the MS/MS peptide data acquired from the ovarian cancer sera sample. The matched peptide information was identified as des-alanine fibrinopeptide A (des-alanine-FPA) with the sequence of DSGEGDFLAEGGGVR. Des-alanine-FPA is derived from FPA by losing an amino acid and its molecule weight is 1465 Da. Interestingly, we did not find the intact FPA of 1535 Da in neither the cancer patient samples nor in the normal healthy individuals. By carefully looking at the results of Mascot searching, we found that the des-alanine-FPA fragment fingerprints matched up with the MS spectra of peak eluting at 42 min in ovarian cancer serum sample (Fig. 4.4).

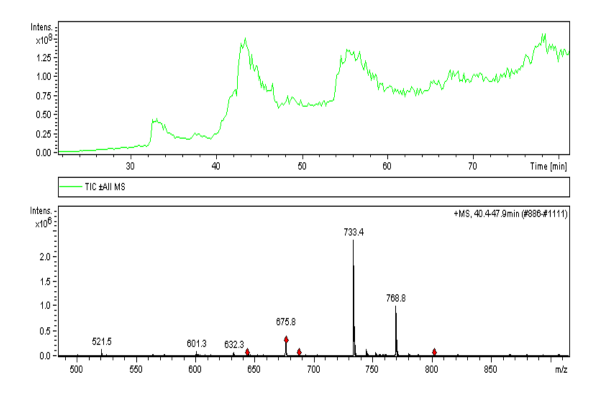


Figure 4.4, The MS spectrum of chromatographic peak eluting at 42 min, with matched MS information with des-alanine-FPA

#### 4.3.2. Protein fractionation method by organic solvent precipitation

Previously our group has successfully removed the abundant proteins by utilizing 80% acetonitrile solution and obtained better prefractionation results compared to centrifugal ultrafiltration. To further optimize this sample preparation method, we compared the signal intensity of des-alanine-FPA in the supernatants after 80% acetonitrile precipitation and 90% methanol precipitation. The results indicated that 90% methanol precipitation yielded higher intensity of des-alanine-FPA and its fragments in the MS spectrum, suggesting a better recovery of this biomarker.

The MS and MS/MS spectra were examined from the whole gradient program to find out when des-alanine-FPA was mainly eluted. This potential biomarker was separated from other hydrophobic components, which were mainly eluted out at the end of elution program with high percentage of organic phase. However, the full scan chromatogram indicated that the intensity of des-alanine-FPA was much lower than these components. These hydrophobic components may overload the reverse phase column and decrease the efficiency of chromatographic separation after a few continuous injections. To prevent the reduction of column robustness, we added a liquid-liquid extraction procedure to remove hydrophobic components after organic solvent precipitation.

The supernatants after 90% methanol precipitation were subjected to a liquid-liquid extraction method using chloroform as the extracting solvent. By comparing the chromatograms with and without chloroform extraction, the MS total ion chromatograms of ovarian cancer serum sample showed much lower intensity of hydrophobic

components, but without much effect on des-alanine-FPA level (Fig. 4.5). In addition, we did not observe any decrease in the resolution and intensity of the differentiated chromatographic peaks in five continuous injections. It proved that the liquid-liquid extraction method was effective to reduce the complexity of serum by removing partial hydrophobic components.

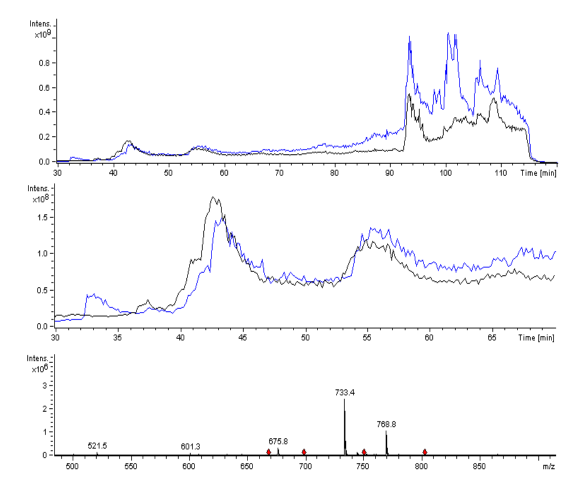


Fig. 4.5, Improvement of chromatograms by adding a chloroform-water extraction procedure into the sample preparation. **Blue line** indicate sample without chloroform-water extraction and **black line** indicate sample with chloroform-water extraction.

#### 4.3.3. LC-MS/MS analysis of different sample sets

Given the finding of des-alanine-FPA as a potential biomarker by LC-MS/MS analysis, we further evaluated this biomarker in each serum sample of the patients with carcinoma, patients with benign conditions, and healthy control. The tandem MS information for each sample was processed and used for the searching of the des-alanine-FPA fingerprints.

The criteria to confirm whether des-alanine-FPA exists in different samples include the MS/MS database search and the elution time at 42 min. Des-alanine-FPA peak was predominantly found in both the benign and carcinoma groups. Analysis of 20 benign samples yielded detectable des-alanine-FPA peak in 16 samples with the percentage of 80%. Among the 40 carcinoma samples subjected to analysis, 37 samples were observed with des-alanine-FPA peak, indicating the sensitivity of this biomarker as 93%. With regarding to the control group, we observed that three samples (N=20) showed desalanine-FPA signal and all other samples had negative results, suggesting a good specificity result of 85%.

Considering half of the malignant tumors are transformed from the pre-existing benign cysts, the up-regulated des-alanine-FPA appearing in a high percentage of benign set is not surprising. However, when comparing the peak area or signal intensity of desalanine-FPA peak in the benign and carcinoma sets, we did not observe any significant difference on its expression levels to discriminate these two sets. This is possible if desalanine-FPA is not only involved in the slow growth of pre-malignant lesions before carcinogenesis, but also is associated with the fast transformation of normal ovary to carcinoma.

#### 4.4. Discussion

In this study, we demonstrated a LC-MS/MS method for the low molecular serum protein profiling in the biomarker discovery of ovarian cancer. One of the obstacles for protein biomarker discovery in serum is the dynamic range of protein components. To overcome this problem, extensive pre-fractionation steps were usually needed to increase the likelihood of low abundance biomarker detection. It has been reported that organic solvent precipitation was used to extract low molecular weight protein in human serum samples [12]. Compared to other methods, organic solvent precipitation method is simple, convenient, and processes high recovery for low abundance peptides.

The sample pretreatment method utilized in this work was modified based on previous reports. It improved reproducibility and sensitivity by reducing the hydrophobic residues in the serum sample solution after the depletion of high abundance proteins. The developed LC-MS/MS method showed that the chromatogram of serum low molecular weight peptides differentiates from ovarian cancer patients to healthy control in a major marker peak. Further, this peak was characterized as des-alanine FPA, which is derived from fibrinopeptide by losing an amino acid of alanine.

Fibrinopeptide A (FPA) is generated from fibrinogen by thrombin during blood coagulation. Thrombosis and disseminated intravascular coagulation are common complications of cancer, and such a procoagulant state in cancer arises mainly from the capacity of tumor cells to express and release procoagulant factors. It has been reported that FPA related peptides were up-regulated in different type of cancers including ovarian cancer [13-16]. In addition, there was finding of elevated phosphor-FPA as post-transcription modification in serum of advance staged ovarian cancer samples [13]. These findings revealed a high degree of biological relevance between cancer and fibrinogen fragments. However, the mechanism of high level des-alanine-FPA and its correlation with ovarian cancer has not been reported yet. It is essential to carry out a careful follow-up validation using rational and rigorous methodology for ovarian cancer biomarker discovery.

In the validation of des-alanine-FPA level in the sample with different sets of ovarian cancer diseases, we found that both benign and carcinoma sets had up-regulated desalanine-FPA expression compared to normal control. The challenge of differentiating benign tumors with malignant carcinoma for ovarian cancer screening remained in this work. We have not found any relation between the level of this biomarker to define different disease states such as benign and carcinoma.

Serum was used as the sampling source in this study because it contains the circulating proteins and peptides shed from cancerous cells and tissues into blood. Compared to plasma, it is less complex due to the coagulation process and the removal of clotting factors during specimen collection. Serum is generally used as an acceptable starting material for many diagnostic assays and proteomics investigations. However, the serum sampling process usually requires the coagulation process of whole blood for 30-45 min, which initiates a cascade of enzymes reactions such as the formation of solid fibrin by serine proteases. During this clotting process, FPA is removed from the n-terminal segment of the alpha chains of fibrinogen by the thrombin. We were concerned that the FPA level and especially des-alanine-FPA level could be affected by clotting process of serum sampling. Variability introduced through sample collecting and storage can be misinterpreted as des-alanine-FPA level changes resulting from ovarian tumorigenesis.

Consequently, further study should be carried out in examining whether the specimen sampling and storage as well as our current sample preparation method affect the desalanine-FPA level. More importantly, the verification of the des-alanine-FPA level in the plasma is essential to move this work forward. Compared to serum, the sample source of plasma avoids the specimen-to-specimen variation of clotting extent and duration and could better represent real blood proteomes.

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#### **CHAPTER V**

# COMPREHENSIVE ANALYSIS OF TUMOR-ASSOCIATED DNA METHYLATION IN HEPATOCELLULAR CARCINOMA

#### 5.1. Introduction

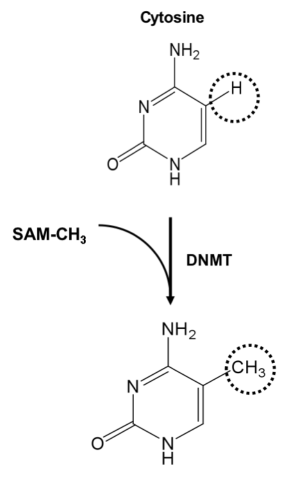
Hepatocellular carcinoma (HCC) is the one of the most aggressive malignancies with the third mortality among cancer worldwide [1]. Highly prevalent in Asia and Africa, HCC is recently reported to be on the rise in many developed countries including United States, Japan, and Western Europe [2]. The clinical management of HCC, depending on its stage, includes curative approaches such as resection, orthotopic liver transplantation, local ablation, radioembolization, and sorafenib treatment [3]. Despite significant advances in HCC management, the survival rate of HCC is low due to its poor diagnosis and prognosis, high recurrence, and resistance to chemotherapy and radiotherapy [4].

The major etiological risk factors of HCC for cancer development have been well understood. These factors include hepatitis virus infection, chronic alcohol consumption, liver cirrhosis, and aflatoxin intake [5,6]. However, the majority of HCC were detected in non-resectable advanced stages, which prevents potential curative treatments.

The early diagnosis of HCC presents a lot of challenges since the imaging techniques such as CT and MRI scan are not adequate to distinguish HCC with long-term liver cirrhosis [7]. The most commonly used tumor biomarker  $\alpha$ -fetoprotein (AFP) is helpful in the surveillance tests, but its use is also controversial because of lacking diagnostic accuracy and sensitivity [8]. Other serum biomarkers such as lectin-bound AFP, or descarboxy-prothrombin (DCP) have large limitations because they are not consistent or particularly precise for the early diagnosis of HCC [9]. Therefore, it is urgent to develop novel biomarkers for the early detection of malignancy of hepatocytes for better screening tests.

The carcinogenesis for HCC development involves complicated genetic and epigenetic changes. Genomic alterations such as gene mutations, chromosomal amplifications and deletions, and unstable genomics were closely associated with HCC [10,11]. In addition, epigenetic alterations, including DNA methylation, histone modification, and altered expression of chromatin-modifying enzymes were also frequently observed in malignant transformation of hepatocytes as an alternative pathway [12]. Aberrant gene methylation has been well documented as the best-known epigenetic event in different cancers including HCC [13,14]. When DNA methylation occurs in CpG dinucleotides, a methyl

group from S-adenosyl methionine (SAM-CH<sub>3</sub>) is added at the carbon five position of the cytosine ring through covalent bond (Fig. 5.1). This reaction is catalyzed by DNA methyltransferases 1, 3a, or 3b (DNMT).



5-methylcytosine

Figure 5.1, Cytosine methylation catalyzed by DNMT

In cancers, global hypomethylation causes the instability of chromosomes and local changes of promoter methylation (hypermethylation) of CpG islands in tumor suppressor genes lead to transcriptional silencing [15,16]. Besides the direct inactivation of tumor suppressor genes, DNA hypermethylation can also block transcription factors and silence DNA repair genes, resulting in the loss of downstream gene functions and the accumulation of genetic lesions [17].

In HCCs, a growing number of genes with aberrant DNA hypermethylation have been identified. Epigenetic silencing of the tumor suppressors RASSF1A and p16 (CDKN2A) were frequently reported in previous studies. Hu L *et al.* detected the promoter hypermethylation of RASSF1A in both serum and tissue DNA in HCC [18]. Similarly, Formeister *et al.* reported the increased methylation of RASSF1A, p16 (CDKN2A), APC, GSTP1, and RIZI in tumorous tissues comparing with adjacent non-tumorous HCC tissues [19]. In addition, this phenomenon has been observed in the non-cancerous cirrhotic tissues of HCC patients, supporting that the hypothesis that epigenetic inactivation is an early event of HCC [20]. These findings help the understanding of hepatocarcinogenesis in early stage and support the potential biomarker discovery for early diagnosis through methylation alterations.

The RASSF1A gene is one major isoform of Ras-Association Domain Family 1 (RASSF1). It maintains the genomic stability and modulates a broad range of cellular functions including apoptosis, cell motility and invasion for normal cell functions [21]. The expression of RASSF1A is ubiquitous in non-cancerous tissues, but is reduced in

cancer cell lines and tissues. Mice with knocked out RASSF1A gene has the tendency to develop tumors, correlating impaired RASSF1A expression with tumorigenesis [22]. The lost of RASSF1A function has been found in 37 tumor types due to the promoter methylation [23].

P16 is a cyclin-dependent kinase inhibitor (cdki) with the biochemical ability to form a complex with cdk 4-6. This binding process inhibits cell cycle progression and tumor progression. P16 gene mutation, promoter methylation, and deletion are counted as the causes of its frequent inactivation [24]. Loss of p16 expression is not only reported in tumor development, but also for the prognosis of tumor progression [5,26].

In this study, we compared the DNA methylation status in paired HCC and noncancerous liver tissues for a panel of 21 tumor-related genes. Among these selected genes, p16, RASSF1A, E-Cadherin, MSH2, CCND2, SEMA3B, SPINT2, SFN, MYC, MAGEA3B, FHIT, and MGMT were reported to have methylation rates higher than 40% for HCC [19, 27-30]. The other 9 genes: p14, p15, GATA4, NDRG4, OPCML, SEPT9, SFRP2, TFPI2, and ALX4 were documented to have elevated methylation in colon cancer tissues [31-34]. Using bisulfite deamination treatment and methylation-specific polymerase chain reaction (MSP) method, we obtained the distinct HCC methylation profiles and evaluated their combinational use for the detection of HCC.

# 5.2. Materials and methods

# 5.2.1. Collection of clinical tissue specimen

We analyzed 80 tissue samples, consisting HCC and paired non-HCC liver tissues as control, from 40 patients who underwent curative resection surgery between 2001 and 2007. All patients were subjected to pathological diagnosis and classification for different stages. Informed consent was obtained before the study from each patient. A summary of the clinicopathological features is listed in Table 5.1. For all patients, liver tissue samples were collected from the cancerous and the adjacent non-cancerous surgical margin. The adjacent tissues included 16 cirrhotic samples and 24 normal samples. All of these tissues were stored as formalin-fixed paraffin-embedded (FFPE) samples.

		HCC patients
NO. of Patients		
	Male	32
	Female	8
Age (average ±SD)		50.5±8.2
Etiology		
	HBV	8
Tumor size		
	>5cm	16
	<5cm	24
Cirrhosis		
	Yes	24
	No	16
Nodules		
	Yes	25
	No	15
Stage		
	Ι	2
	II	17
	II-III	8
	III	13

Table 5.1, Clinical and pathological characters of 40 HCC patients

#### 5.2.2. DNA isolation from liver tissues

Since all the tissue samples were fixed by formalin and embedded by paraffin, they were cut into 5  $\mu$ m sections prior to xylene extraction. Every 6-7 sections of FFPE samples were incubated with 1.5 mL of xylene and vortexed for 30 min. Followed by the centrifugation at 10,000 rpm for 5 min, the supernatant was removed to a clean Eppendorf tube. These steps were repeated for 3 times by adding ethanol into the pellet to remove the residual xylene. At the end, the resulting tissue pellet was collected after the evaporation of ethanol residue.

Genomic DNA automated extraction was performed on the Maxwell® 16 Instrument (Promega, Madison, WI) with Maxwell® 16 DNA Purification Kits. Tissue pellet was lysed, purified, and washed by cell lysis solution, MagneSil PMPs, and wash buffer, respectively.

# 5.2.3. Sodium bisulfite conversion

Prior to sodium bisulfite conversion, DNA concentration was quantified by RT-PCR amplification. Standard unmethylated and methylated genomic DNA was purchased with CpGenome DNA modification Kit (Millipore, Billerica, MA) for establishing standard curve. Different concentration of standard DNA were obtained by serial dilution and then amplified with the  $\beta$ -actin (ACTB) primer: 5' GGCGGCACCACCATGTACCCT 3' and 5' AGGGGCCGGACTCGTCATACT 3'. Sample DNA concentration was calculated

utilizing the standard calibration curve of cycle numbers and log concentration. Based on these results, the final genomic DNA concentrations of all samples were normalized within the same concentration through dilution adjustment.

Sodium bisulfite reaction was performed on cancerous, adjacent cirrhosis, and normal control DNA samples as well as standard unmethylated and methylated DNA samples using the EZ DNA Methylation-Gold Kit (Zymo Research, Irvine, CA). Briefly, the bisulfite reagent was prepared as 5.5 mol/L sodium bisulfite in the mixture of M-Dilution buffer and M-Dissolving buffer (6:1, v/v). Every 100 ng DNA samples were mixed with 50  $\mu$ L bisulfite reagent. The reaction took 150 min in the following program: 0-10 min at 95 °C, 10-150 min at 64 °C, and then cool down to 4 °C at the end of the reaction. The modified DNA sample was then desalted using KingFisher Flex Magnetic Particle Processors (Thermo Scientific, Asheville, NC). The whole purification consisted of sample binding process utilized 200  $\mu$ L EZ beads for each sample. The desulfonation step used 0.2 mmol/L NaOH in PEG binding buffer, and the ethanol wash used 80% ethanol and 100% ethanol with or without salts. At the end, the converted DNA sample was eluted in 10 mM Tris buffer and store at -20 °C for the further use.

MSP amplifications of 21 genes including p16, RASSF1A, MSH2, CCND2, SEMA3B, SPINT2, SFN, MYC, MAGEA3B, FHIT, E-Cad, MGMT, p14, p15, GATA4, NDRG4, OPCML, SEPT9, SFRP2, TFPI2, and ALX4 were performed on MJ Research Chromo4 Real Time PCR Instrument (Bio-Rad, Hercules, CA) and LightCycler 480 (Roche Diagnostics; Switzerland). The primer sequences of each locus for MSP were described in Table 5.2. The reaction mixture included 1  $\mu$ L bisulfite-modified genomic DNA, 4  $\mu$ L primer solution at 5  $\mu$ mol/L, and 5  $\mu$ L Type-it® HRM master mix (contains DNA polymerase, EvaGreen dye, optimized concentration of Q-solution, dNTPs, and MgCl<sub>2</sub>) (Qiagen,Valencia, CA) for a total volume of 10  $\mu$ L. The following PCR program was applied: 95 °C denaturing for 10 min and 50 cycles of 95 °C for 45 s, annealing temperature for 30 s, and extension at 72 °C for 45 s.

# Table 5.2, The MSP primer sequences of 21 genes

Gene	MSP primer 5'-3'	Annealing Temperatu re ( °C)
	F: TTATTAGAGGGTGGGGGGGGGATCGC	
p16 (INK4A)	R: CCACCTAAATCGACCTCCGACCG	66
	F: GTGTTAACGCGTTGCGTATC	
RASSF1A	R: AACCCCGCGAACTAAAAACGA	57
	F: GGCGGATTT TAT CGTAGTCG	
CCND2	R: CTCCACGCTCGA TCCTTCG	62
	F: TGGTTAGGCGGGGGTATTTTC	
SEMA3B	R: TCAACAATAAAAACGAAAACG	58
	F: CGGGCGTTTTTATATTGAAGGTTC	
SPINT2	R: ACGCCACCAACCGTTAAAATCTCG	60
МҮС	F: TAGAATTGGATCGGGGGTAAA	
	R: CGACCGAAAATCAACGCGAAT	57
SFN	F: TGGTAGTTTTTATGAAAGGCGTC	
(has-mir-219-2)	R: CCTCTAACCGCCCACCACG	58
	F: TTTGTTCGGAATTTAGGGTAGTATC	
MAGEA3	R: GTCGCTCGTTACTCAAAACG	60
	F: TCG TGG TCG GAC GTC GTT C	
MSH2	R: CAA CGT CTC CTT CGA CTA CAC CG	60
	F: GAAGGTAGGGGGGGGGGGGGGGAGGTAAGTT	
FHIT	R: CGTAAACGACGCCGACCCCACTA	68

Gene	MSP primer 5'-3'	Annealing Temperature (°C)	
	F: TAATTAGCGGTACGGGGGGC		
E-Cadherin		66	
	R: CGAAAACAAACGCCGAATACG		
MGMT	F: ATTTGGTGAGTGTTTGGGTCGTTTC	57	
	R: AAAACGCACCTAAAACTCGCCC	GCCC	
OPCML	F: CGTTTAGTTTTTCGTGCGTTC	62	
	R: CGAAAACGCGCAACCGACG		
P15	F: GCGTTCGTATTTTGCGGTT	58	
	R: CGTACAATAACCGAACGACCGA		
SFRP2	F: GGGTCGGAGTTTTTCGGAGTTGCGC	60	
	R: CCGCTCTCTTCGCTAAATACGACTCG		
TFPI2	F: TTTCGTATAAAGCGGGTATTC	57	
	R: ACGACCCGCTAAACAAACG		
ALX4	F: CGTTCGCGTTTTCGTTCGTCGTTTGC	58	
	R: ACGACGAACCCTCCCGACTCTACG		
GATA4	F: AGGTTAGTTAGCGTTTTAGGGTC	60	
	R: ACGACGACGAAACCTCTCG		
NDRG4	F: TTTAGGTTCGGTATCGTTTCGC	60	
	R: CGAACTAAAAACGATACGCCG		

#### 5.2.5. Specificity and sensitivity of MSP

The specificity of the MSP assay was evaluated by detecting the bisulfite-modified standard unmethylated DNA and methylated DNA through high-resolution-melting (HRM) analysis repeatedly. The melting temperatures for each gene were recorded as reference for the patient sample analysis. Sensitivity of MSP was determined by mixing bisulfite-modified standard methylated DNA (1%) with standard unmethylated DNA (99%) together.

#### 5.2.6. Quantitative methylation analysis

The quantitative methylation analysis was carried out on MJ Research Chromo4 Real Time PCR Instrument. Bisulfite-converted standard DNA sample of known concentration was serial diluted and amplified using bisulfite-converted ACTB primer. The standard curve prepared from this step was used for determining reference quantity of total DNA( $Q_{total}$ ) amount for each patient sample. The amplification condition was as following: 95 °C denaturing for 10 min, 50 cycles of 95 °C for 45 s and 60 °C annealing for 30 s, and extension at 72 °C for 45 s. Patient samples that were negative for ACTB gene amplification were exclude from the study.

The methylated DNA reference( $Q_M$ ) was determined by constructing the standard curve of bisulfite-converted standard methylated DNA in serial concentrations. The conditions

for the amplification were using the MSP primer and specific annealing temperature. In this case, only the converted methylated DNA can be amplified.

# 5.3. Results

# 5.3.1. Specificity and sensitivity of MSP method

The specificity of the MSP methods on 21 genes was evaluated using standard unmethylated DNA as negative control and methylated DNA as positive control to ensure the completion of bisulfite conversion. In addition, two blank samples, one from the blank control added before the bisulfite conversion and the other added before the PCR reaction, were used to account for false positive. As shown in Fig. 5.2, this highly specific test on the SFN gene was illustrated in the PCR amplification and the subsequent melting curve analysis for standard methylated DNA and unmethylated DNA as well as a blank control. The selected primer and optimized PCR conditions ensured that only methylated DNA after bisulfite conversion could be amplified for all genes. The specificity of these MSP methods is high for all 21 genes.

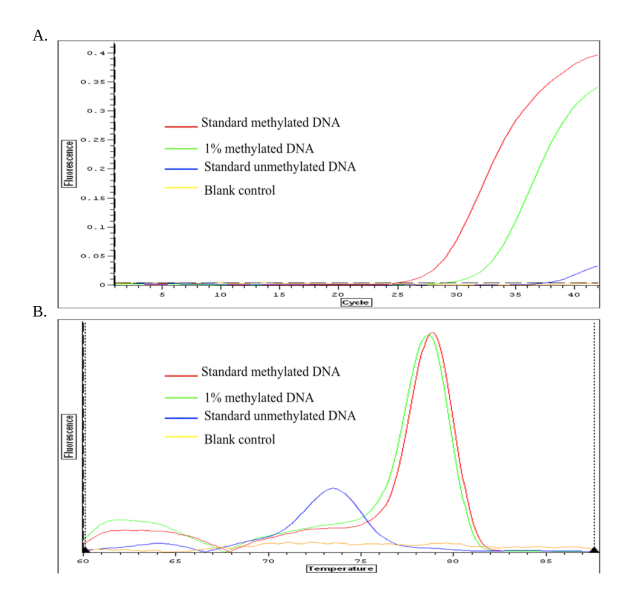


Figure 5.2, The amplification results of standard methylated DNA, 1% methylated DNA, standard unmethylated DNA as negative control, and blank control are shown in RT-PCR (A) and melting curve analysis (B).

The sensitivity of the MSP methods was assessed by mixing 1% standard methylated DNA and 99% standard unmethylated DNA together. Repeated amplifications of the mixed sample were performed by RT-PCR. For most genes, the MSP methods could detect 1% methylation to ensure enough sensitivity of tests (Fig. 5.2).

During the patient sample MSP analysis, control samples including negative and positive control, and two blank samples, as well as 1% standard methylated DNA were amplified on the same plate with patient samples. These controls ensured the completion of bisulfite reaction and the reliability of the results.

# 5.3.2. Gene-specific promoter methylation analysis

In epigenetic studies of HCC, the aberrant promoter methylation of p16, RASSF1A, E-Cadherin, MSH2, CCND2, SEMA3B, SPINT2, SFN, MYC, MAGEA3B, FHIT, and MGMT was frequently reported in previous studies [19, 27-30]. We selected these genes for methylation screening. In addition, another nine genes including p14, p15, GATA4, NDRG4, OPCML, SEPT9, SFRP2, TFPI2, and ALX4, which were found to be highly hypermethylated in colon cancer and related to the tumorigenesis [31-34], were also examined in this study. For all the 40 pairs of matched cancerous and noncancerous liver tissue samples, 4 pairs of patient samples were found to have very low amount of DNA and they were excluded from the methylation profiling.

Among all the patient samples, 14 pairs of HCC cancerous and non-cancerous adjacent tissues were first randomly selected for the screening of all genes in the training phase, and then another 22 pairs of matched liver tissue samples were used for validating the screened genes. The reported highly methylated genes on colon cancer were found to have negative results for all HCC samples. In addition, the methylation rates on E-Cadherin, MGMT, CCND2, MYC, and FHIT were very low on both cancerous and non-cancerous tissues. For all other genes, the methylation rates were listed in Table 5. 3.

	Cancerous li	Cancerous liver tissue		Adjacent	
Gene				non-cancerous tissue	
	No. of samples methylated	Methylation rates	No. of samples methylated	Methylation rates	
P16	15(36)	41%	2(36)	6%	
RASSF1A	32(36)	88%	10(36)	28%	
SPINT2	23(36)	57%	2(36)	6%	
SFN	33(36)	92%	32(36)	89%	
SEMA3B	24(36)	67%	20(36)	56%	
MAGEA3	22(36)	61%	24(36)	67%	
MSH2	22(36)	61%	19(36)	53%	
CCND2	2(14)	14%	1(14)	7%	
МҮС	0(14)	0%	1(14)	7%	
FHIT	1(14)	7%	3(14)	21%	

Table 5.3 The methylation profiles of ten tumor suppressor genes in the screening test

Based on these results from the screening test, p16, RASSF1A, SPINT2, SFN, SEMA3B, MAGEA3, and MSH2 showed relatively high methylation frequencies on HCC cancerous tissues. As a result, these seven genes were further selected for the validation analysis on the other 22 pairs of samples (Table 5. 3). For genes p16, RASSF1A, and SPINT2, they showed significant higher methylation rates in cancerous samples over noncancerous samples. However, we found that the methylation frequencies of SFN, SEMA3B, MAGEA3, and MSH2 were also high in adjacent noncancerous sample and their methylation profiles were not adequate to distinguish these two groups.

The high methylation frequencies in non-cancerous tissues on SFN, SEMA3B, MAGEA3, and MSH2 may be related to the field cancerization of HCC, which initiates the malignancy with the accumulation of epigenetic and genetic damages on several tumor suppressor genes [35, 36]. To test this explanation, we examined the methylation status of the aforementioned seven genes in the validation study from 36 pairs of HCC and their corresponding non-cancerous tissues. Four possibilities of methylation status in paired samples from the same patient were proposed: 1) positive methylation in both cancerous and non-cancerous tissues as  $C_{(+)}/NC_{(+)}$ ; 2) positive methylation in cancerous tissue but negative in non-cancerous tissue as  $C_{(-)}/NC_{(-)}$ ; 4) negative methylation in cancerous tissue but positive in non-cancerous tissue as  $C_{(-)}/NC_{(+)}$ . The results after the comparison of each gene showed that most tissue pairs had accordant alterations with the methylation status of  $C_{(+)}/NC_{(+)}$  at 40%,  $C_{(+)}/NC_{(-)}$  at 25%, and  $C_{(-)}/NC_{(-)}$  at 24%. The discordant methylation appeared in 3% tissue pairs as  $C_{(-)}/NC_{(+)}$  for genes p16, RASSF1A, SEMA3B, MAGEA3, and MSH2 (Table 5.4).

Genes	C(-)/NC(-)	C(-)/NC(+)	C(+)/NC(-)	C(+)/NC(+)
P16	20(56%)	1(3%)	14(39%)	1(3%)
RASSF1A	3(8%)	1(3%)	23(64%)	9(25%)
SPINT2	13(4%)	0(0%)	21(58%)	2(6%)
SFN	3(8%)	0(0%)	1(3%)	32(89%)
SEMA3B	10(28%)	2(6%)	6(17%)	18(50%)
MAGEA3	11(31%)	3(8%)	1(3%)	21(58%)
MSH2	13(36%)	1(3%)	4(11%)	18(50%)
Average%	24%	3%	25%	40%

Table 5.4. Methylation status in paired cancerous and non-cancerous samples (n=36) from the same patient

## 5.3.3. Quantitative methylation analysis

Although the promoter methylation on SFN, SEMA3B, MAGEA3, and MSH2 were high in HCC patient samples, the MSP analysis cannot differentiate the non-cancerous and cancerous tissue with the assistance of the melting curves. To obtain the quantitative information, we further performed RT-PCR methylation analysis of these three genes on 36 pairs of matched samples. The quantitative results were expressed as methylation percentage, which was determined by the following equation:

$$Methylation\% = \frac{Q_M}{Q_{Total}} \times 100\%$$

 $Q_M$  and  $Q_{Total}$  were the reference methylated and total DNA quantity, respectively. As shown in Figure 5.3, Standard curves were established to determine  $Q_M$  and  $Q_{Total}$ .

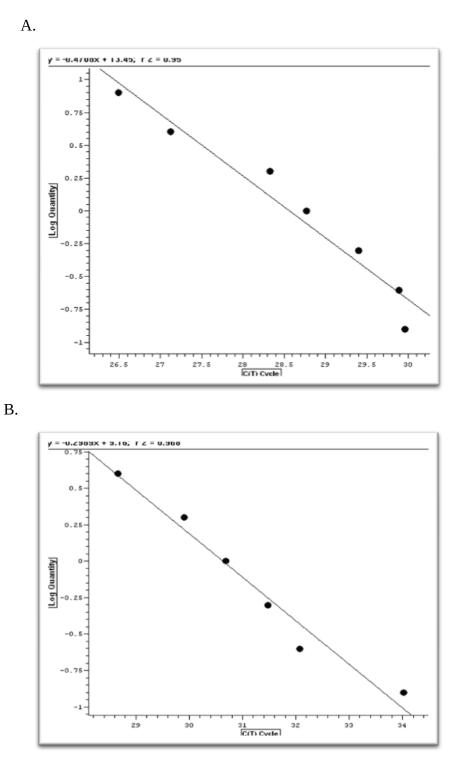


Figure 5.3, Standard curves constructed for quantifying methylated DNA for gene SFN (A) and total DNA using primer ACTB (B)

As summarized in Table 5.5, the quantitative results in 36 pairs of matched HCC tissues showed higher methylation percentage on gene SFN and MSH2 for cancerous samples comparing with the non-cancerous ones. The methylation percentage was low in both tissues for gene SEMA3B, and it appeared to be higher in non-cancerous tissue than in cancerous tissue for gene MAGEA3. Currently, the cut-off value cannot be set to define the cancerous and non-cancerous tissues based on their methylation percentage.

Table 5.5 The methylation percentage of 4 genes on 36 pairs of HCC cancerous/non-cancerous samples

Genes	HCC Cancerous Samples	Non-Cancerous Samples	Ν	
	(Average ±S.D)	(Average ±S.D)		
SFN	76% ±39%	37% ±25%	32	
SEMA3B	14% ±8%	7% ±5%	18	
MAGEA3	59% ±17%	94% ±26%	21	
MSH2	$57\% \pm 28\%$ ,	21% ±19%	18	

## 5.4. Discussion

Epigenetic analysis of promoter methylation plays an important role in the process of tumorigenesis and biomarker discovery for disease diagnosis. Recent studies have demonstrated that HCC had a high frequency of promoter methylation of multiple genes [18-19, 27-30]. However, many issues regarding the comprehensive methylation profile of a large pool of tumor-associated genes and the mechanisms of epigenetic phenomena in cancer progress remain elusive. In this study, we decided to investigate the promoter methylation status of multiple genes including p16, RASSF1A, E-Cadherin, MSH2, CCND2, SEMA3B, SPINT2, SFN, MYC, MAGEA3B, FHIT, MGMT, p14, p15, GATA4, NDRG4, OPCML, SEPT9, SFRP2, TFPI2, and ALX4. These genes are involved in different molecular pathways of carcinogenesis such cell-cycle regulatory (p16, p14, and p15), apoptosis (RASSF1A, SEMA3B and FHIT), cell adhesion (E-Cadherin), and DNA repair (MGMT). Understanding their functions in the advance tumor stages of HCC provides the ability to predict the premalignant conditions for early diagnosis.

The current study has successfully identified altered methylation status on several genes using the developed quantitative methylation profiling method. Sodium bisulfite reacted with the methylated CpG site specifically, enabling the discrimination of methylated DNA with unmethylated DNA by methylation-specific PCR. Among the 21 tested genes, p16, RASSF1A, SPINT2, SFN, SEMA3B, MAGEA3, and MSH2 had higher frequencies of promoter methylation in the training phase, in which 14 pairs of cancerous versus noncancerous adjacent tissues were used. Although hypermethylation of E-Cadherin, CCND2, MYC, FHIT, and MGMT in HCC was reported in previous studies for HCC [28-30], it was not observed in the present study. Interestingly, the methylation status of all colon cancer related genes including p14, p15, GATA4, NDRG4, OPCML, SEPT9, SFRP2, TFPI2, and ALX4 showed negative results for all HCC samples. This may be explained by that the molecular functions of these genes in carcinogenesis are specific to colon cancer, rather than HCC.

Based on the results of training phase, all the genes with low methylation rates in HCC samples were eliminated, leaving seven genes for the validation and further screening on another 22 pairs of matched samples. The methylation status of these seven genes were profiled and shown that high methylation frequency of both cancerous tissue and adjacent non-cancerous tissue for the majority of sample pairs. This phenomenon suggested field cancerization in the surgical margin. For gene SPINT2 and SFN, their methylation status was accordant in all the sample pairs, revealing the monoclonal expansion model in the early cancer event. The discordant methylation status of p16, RASSF1A, SEMA3B, MAGEA3, and MSH2 may result from the polycolonal origins of HCC. These results indicated the complicated origins in hepatocarcinogenesis, involving both monoclonal and polyclonal expansion of preneoplastic cells.

Due to the silenced gene expression in cancerous and non-cancerous pairs for most hypermethylated genes, the methylation frequencies were not adequate to predict the early event of HCC. We developed the comparative quantification assay for SFN, SEMA3B, MAGEA3, and MSH2 to separate HCC cancerous and non-cancerous samples. SFN and MSH2 demonstrated higher methylation percentage compared to other genes. However, adequate cut-off value to define cancerous from non-cancerous tissues was not available for these two genes. These results may be due to the fact that HCC sample pairs used in the study were mainly from advanced stage, and the field cancerization caused the accumulative epigenetic changes in the surgical margins.

In conclusion, we examined the gene-specific alterations through promoter methylation profiling on 21 genes using bisulfite conversion and MSP. With the comparison of HCC cancerous and non-cancerous tissues, frequent CpG island hypermethylation was found for p16, RASSF1A, and SPINT2. In addition, we observed the consistent methylation status in cancerous and adjacent non-cancerous tissue for other hypermethylated genes SFN and MSH2, resulted from field cancerization of HCC. In the future, the quantitative methylation assays will be performed on early stage of HCC samples to better assess the ability of these genes for its early screening. The results from the present study supported that the DNA methylation could be an important event during carcinogenesis and a potential biomarker for HCC diagnosis.

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