

**β -CYCLODEXTRIN FUNCTIONALIZED IONIC LIQUID AS
HIGH PERFORMANCE LIQUID CHROMATOGRAPHY
CHIRAL STATIONARY PHASE FOR THE
ENANTIOSEPARATION OF NATURAL PRODUCTS AND
PHARMACEUTICALS**

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**FACULTY OF SCIENCE
UNIVERSITY OF MALAYA
KUALA LUMPUR**

2017

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**THESIS SUBMITTED IN FULFILMENT OF THE
REQUIREMENTS FOR THE DEGREE OF DOCTOR OF
PHILOSOPHY**

**FACULTY OF SCIENCE
UNIVERSITY OF MALAYA
KUALA LUMPUR**

2017

UNIVERSITY OF MALAYA
ORIGINAL LITERARY WORK DECLARATION

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Name of Degree: Degree of Doctor of Philosophy

Title of Project Paper/Research Report/Dissertation/Thesis (“this Work”):

β -Cyclodextrin functionalized ionic liquid as high performance liquid chromatography chiral stationary phase for the enantioseparation of natural products and pharmaceuticals

Field of Study: Analytical Chemistry

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ABSTRACT

The demanding for enantiomerically pure (enantiopure) compounds, especially for pharmaceutical field has been attracting great attention during last decades. Direct enantioseparation by chiral stationary phases (CSPs) using high performance liquid chromatography (HPLC) remains as the most important technique for enantioseparation. The development of novel stable and powerful CSPs is therefore important. The first part of this study involved a facile and reliable preparation of CSPs. Thus, β -cyclodextrin was functionalized with ionic liquids (ILs) namely 1-benzylimidazole (1-BzIm) and 1-decyl-2-methylimidazole (C_{10} MIm) with tosylate as anion produced β -CD-BIMOTs and β -CD-DIMOTs respectively. β -CD-BIMOTs and β -CD-DIMOTs were attached to the modified silica to obtain the CSPs. The performances of the synthesized CSPs were determined by examining the capability of enantioseparation of selected analytes: flavonoids (flavanone, hesperetin, naringenin and eriodictyol), β -blockers (atenolol, metoprolol, pindolol and propranolol) and Non-steroidal anti-inflammatory drug (NSAIDs) (ibuprofen, fenoprofen, ketoprofen and indoprofen). The performance of β -CD-BIMOTs and β -CD-DIMOTs stationary phases was also compared with native β -CD stationary phase. The results indicated that β -CD-BIMOTs stationary phase afforded more favorable enantioseparations than β -CD-DIMOTs and native β -CD based stationary phases. Therefore, the optimization for enantioseparation of selected analytes (flavonoids, β -blockers and NSAIDs) and evaluation of interactions was further investigated on β -CD-BIMOTs stationary phase. The selected flavonoids, flavanone and hesperetin obtained high resolution factor in reverse phase mode. Meanwhile naringenin and eriodictyol attained partial enantioseparation in polar organic mode. In order to understand the mechanism of separation, the interaction of selected flavonoids and β -CD-BIMOTs was studied using spectroscopic methods which are 1H NMR, NOESY and UV/Vis spectrophotometry. The result for enantioseparation of selected β -blockers, propranolol and metoprolol showed good enantioresolution compared to atenolol and pindolol. The results suggested that the lipophilic property and the structure of propranolol and metoprolol that enable the formation of inclusion complex contribute to better enantioseparation. This observation was proven by 1H NMR and NOESY of β -CD-BIMOTs/ β -blockers. The effect of the types and variation of mobile phase composition on enantioseparation of NSAIDs was also studied on β -CD-BIMOTs CSP. From the result of enantioseparation, ibuprofen and indoprofen achieved the better resolution than ketoprofen and fenoprofen due to their favorable orientation to fit into the β -CD-BIMOTs cavity. This orientation was depending on the structure of NSAIDs.

ABSTRAK

Permintaan yang tinggi terhadap sebatian enantio yang asli, terutamanya dalam bidang farmaseutikal telah menjadi perhatian sejak berdekad yang lalu. Pemisahan enantio secara langsung oleh fasa pegun kiral (CSP) menggunakan kromatografi cecair prestasi tinggi (HPLC) adalah teknik yang penting untuk pemisahan enantio. Oleh itu, perkembangan penghasilan CSP yang terbaru perlu diambil kira. Bahagian pertama kajian ini adalah melibatkan penyediaan CSP yang sangat mudah. Untuk itu, β -cyclodextrin telah difungsikan dengan cecair ionik (ILs) iaitu 1-benzylimidazole (1 BzIm) dan 1-Decyl-2-methylimidazole ($C_{10}MIm$) dengan tosylate sebagai anion masing-masing menghasilkan β -CD-BIMOTs dan β -CD-DIMOTs. β -CD-BIMOTs dan β -CD-DIMOTs dilekatkan pada silika terubahsuai untuk menghasilkan fasa pegun kiral. Prestasi fasa pegun kiral ini diukur dengan keupayaan pemisahan enantio terhadap analit yang terpilih: flavonoid (flavanone, hesperetin, naringenin dan eriodictyol), β -blockers (atenolol, metoprolol, pindolol dan propranolol) dan ubat anti-radang bukan steroid (NSAIDs) (ibuprofen, fenoprofen, ketoprofen dan indoprofen). Prestasi fasa pegun β -CD-BIMOTs dan β -CD-DIMOTs juga telah dibandingkan dengan fasa pegun β -CD asli. Keputusan menunjukkan bahawa fasa pegun β -CD-BIMOTs mencapai pemisahan enantio yang lebih baik daripada fasa pegun β -CD-DIMOTs dan fasa pegun β -CD asli. Oleh itu, pengoptimuman pemisahan enantio terhadap analit yang terpilih (flavonoid, β -blockers dan NSAIDs) dan penilaian interaksi yang terlibat disiasat dengan menggunakan fasa pegun β -CD-BIMOTs. Flavonoid seperti flavanone dan hesperetin memperolehi faktor resolusi yang tinggi dalam mod fasa terbalik. Sementara itu, naringenin dan eriodictyol mencapai separa pemisahan enantio dalam mod organik berkutub. Untuk memahami mekanisma pemisahan, interaksi flavonoid dan β -CD-BIMOTs dikaji menggunakan kaedah spektroskopi iaitu 1H NMR, NOESY dan spektrofotometri UV-Vis. Keputusan pemisahan enantio β -blockers menunjukkan resolusi enantio propranolol dan metoprolol adalah lebih baik berbanding atenolol dan pindolol. Ini kerana sifat lipofilik serta struktur propranolol dan metoprolol yang membolehkan pembentukan kompleks kemasukan berlaku dan seterusnya menyumbang kepada pemisahan enantio yang lebih baik. Interaksi ini dibuktikan dengan 1H NMR dan NOESY β -CD-BIMOTs/ β -blockers. Pemisahan enantio NSAIDs dengan β -CD-BIMOTs turut dikaji berdasarkan jenis dan kepelbagaian komposisi fasa bergerak. Berdasarkan keputusan pemisahan enantio, ibuprofen dan indoprofen mencapai resolusi yang lebih baik berbanding ketoprofen dan fenoprofen kerana orientasi yang sesuai untuk mereka dimuatkan ke dalam rongga β -CD-BIMOTs. Orientasi ini bergantung kepada struktur NSAIDs itu sendiri.

ACKNOWLEDGEMENTS

It is with great pleasure I convey my sincere appreciation to all who made my doctoral degree a success. First and foremost, I would like to express my gratitude to my research advisor, Dr. Sharifah Mohamad and Dr Tay Kheng Soo, for their support, encouragement, patience and guidance during my graduate studies. I will never forget what I have learned from them and I look forward to our future endeavors. I consider myself extremely fortunate to have the opportunity to work under their supervision.

I greatly appreciate the assistance of all faculty and staff in the Department of Chemistry at the University of Malaya, especially Miss Norzalida Zakaria for assisting me in using NMR, and other lab assistants for helping me to maintain HPLC instruments and training me in the use of other instrumentation. I also thank my lab members (FD-L5-4) and other colleagues (Dr. Muggundha, Dr. Nur Nadhirah, Dr Saliza, Dr. Mazidatul, Siti Farhana, Khalijah, Shabnam, Fairuz Liyana, Naqiyah Farhan, Syed Fariq, Ahmad Razali, Nur Faizah and Nur Atiqah) for their help and friendship.

Last but not least, I would especially like to thank my parents, mother Siti Omar, father Rahim Yussof, sisters Nur Syuhada and Nuratikah for their unconditional love and support during my life.

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LIST OF SYMBOLS AND ABBREVIATIONS

1-BzIm	:	1-Benzylimidazole
1D	:	1 dimension
2D	:	2 dimension
ACN	:	Acetonitrile
C ₁₀ MIm	:	1-decyl-2-methylimidazole
CD	:	Cyclodextrin
CIP	:	Cahn-Ingold-Prelog priority
CoA	:	Coenzyme A
CSP	:	Chiral stationary phases
DCM	:	Dichloromethane
DMF	:	N, N-Dimethylformamide anhydrous
DMSO-D ₆	:	Dimethyl Sulfoxide
EMIMLpro	:	1-ethyl-3-methyl-imidazolium L-proline
FDA	:	Food and Drug Administration
FT-IR	:	Fourier transforms infrared
HILIC	:	Hydrophilic Interaction Liquid Chromatography
HOAc	:	Acetic acid
HPLC	:	High performance liquid chromatography
HAS	:	Human serum albumin
ILs	:	Ionic liquids
LE-CE	:	Ligand-exchange capillary electrophoresis
LE-MEKC	:	Ligand-exchange micellar electrokinetic capillary chromatography
MD	:	Molecular dynamics

MDPCCD	: Mono-6-(3-methylimidazolium)-6-deoxyper (3,5-dimethylphenylcarbamoyl)- β -cyclodextrin chloride
MeOH	: Methanol
MPCCD	: Mono-6-(3-methylimidazolium)-6-deoxy-perphenylcarbamoyl- β -cyclodextrin chloride
NaOH	: Sodium hydroxide
NMR	: Nuclear Magnetic Resonance
NOESY	: Nuclear Overhauser Effect Spectroscopy
NSAID	: Non-steroidal anti-inflammatory drugs
ODPCCD	: Mono-6-(3-octylimidazolium)-6-deoxyper (3,5-dimethylphenylcarbamoyl)- β -cyclodextrin chloride
OH	: Hydroxyl
OPCCD	: Mono-6-(3-octylimidazolium)-6-deoxyperphenylcarbamoyl- β -cyclodextrin chloride
PG	: Prostaglandin
T3	: Triiodothyronin
T4	: Thyroxin
TDI	: Toluene 2,4-diisocyanate
TEA	: Triethylamine
TEAA	: Triethylamine acetate
TGA	: Thermo gravimetric analyses
Ts ₂ O	: <i>p</i> -Toluene sulfonic anhydride
VAMPCCD-POLY	: 6 ^A -(N,N-allylmethylammonium)-6-deoxyperphenylcarbamoyl- β -cyclodextrin chloride
VIMPCCD-POLY	: 6 ^A -(3-vinylimidazolium)-6-deoxyperphenylcarbamate- β -cyclodextrin chloride
β -CD	: β -Cyclodextrin
β -CD-BIMOTs	: Mono-6-deoxy-6-(3-benzylimidazolium tosylate)- β -CD

β -CD-DIMOTs : Mono-6-deoxy-6-(3-decyl-2-methylimidazolium tosylate)- β -CD

β -CDOTs : 6-O-Monotosyl-6-deoxy- β -CD

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CHAPTER 1: INTRODUCTION

1.1 Background of study

In chemistry, chirality refers to a molecule that containing asymmetric center (chiral atom or chiral center) and thus it can occur in a pair of isomer which is two mirror images of each other. This pair of isomer is called enantiomers or optical isomers (Figure 1.1). Chirality is important because the biological properties of enantiomers may differ significantly. Using ethambutol and thalidomide as examples, one enantiomer of ethambutol is used to treat tuberculosis while the other isomer causes blindness. *R*-thalidomide is a sedative and effective against morning sickness, whereas *S*-thalidomide is causing the birth defect (Sekhon, 2013; Blaschke *et al.*, 1978). A guideline was issued in 1992 by US Food and Drug Administration (FDA) that each drug enantiomer must be studied separately for its pharmacological pathways, and only therapeutically active isomer is allowed to be marketed (Stinson, 2000).

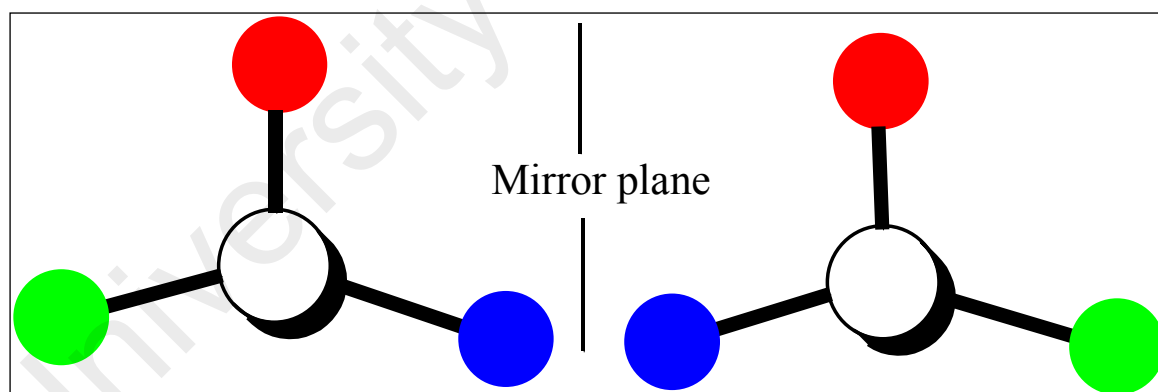


Figure 1.1: Chiral molecule

In laboratory, most compounds are produced as racemic mixture that containing equal amount of enantiomers. Ideally, the desired pure enantiomer could be obtained by direct asymmetry synthesis without further treatment (Pazos *et al.*, 2009; Svang-Ariyaskul *et al.*, 2009; Karnik & Kamath, 2008; Kaluzna *et al.*, 2005; Missio & Comasseto, 2003). However, this approach is not always efficient or cost effective. By using chiral catalysts for asymmetric reaction, catalyst efficiency, reaction conditions and kinetics should be considered. Furthermore, there are no general chiral catalysts for all asymmetric reactions. In order to obtain the pure enantiomer, the separation of an enantiomeric mixture or so called enantioseparation is often necessary (Schurig, 2002; Szejtli, 1998). The enantioseparation method includes enzymatic resolution, the diastereomers crystallization or direct chromatographic separation (Lorenz & Seidel-Morgenstern, 2014; Allenmark, 1989).

Recently, high performance liquid chromatography (HPLC) is becoming more widely used instrument for the direct separation of chiral compounds. An advantage of HPLC is that it can be used to separate enantiomers which are non-volatile, polar, or ionic. There are several approaches that have been used to achieve enantioseparation using HPLC. The simplest way to achieve the enantioseparation is to add chiral additives directly into the mobile phase of HPLC (Zhang *et al.*, 2005). This approach affords satisfactory separation with simpler operation. However, the used of chiral additives could not be regenerated after separations. In addition, the preparation of the chiral additives can be laborious and expensive. Consequently, another more practical approach is to use chiral HPLC column that containing chiral stationary phases (CSPs). In this method, the chiral selector is physically adsorbed or covalently bonded to the solid support for the preparation of CSPs. There are several types of CSPs applied in HPLC such as pirkle-type CSPs, polysaccharide-based CSPs, cyclodextrin-based CSPs, macrocyclic antibiotics-based CSPs, chiral crown ether-based CSPs, protein-based

CSPs and molecular imprinting-based CSPs. Herein, this dissertation focuses on cyclodextrin (CD) based CSPs.

CDs are natural cyclic oligosaccharides consisted of six or more glucose units joined through α -1, 4 linkage (Figure 1.2a). CDs contain hydrophobic center and hydrophilic outer surface (Figure 1.2b). Due to the chair conformation of the glucose units, the CDs are shaped like a truncated cone rather than perfect cylinders as illustrated in Figure 1.2b. CDs are classified by the number of glucose unit. α -CD, β -CD, γ -CD containing six, seven and eight glucose unit, respectively. β -CD based CSPs are among the most widely used CD in HPLC due its special sizes of its hydrophobic cavity (cavity size: α -CD < β -CD < γ -CD) (Stalcup *et al.*, 1990; Armstrong *et al.*, 1986; Armstrong *et al.*, 1985; Armstrong & DeMond, 1984).

When β -CD is used as CSP, chiral recognition can be achieved via the interaction between chiral β -CD and enantiomers (Gubitz & Schmid, 2009). The example of interaction is illustrated in Figure 1.3. The β -CD molecule contains 35 chiral centers. Enantiomers can interact via van der Waals dispersion forces with the hydrophobic cavity which is due to methylene hydrogen. β -CD also has a C_7 symmetry axis and 14 hydroxyl groups situated at the exterior of the cavity. Thus, a number of potential interactions might be present between these hydroxyl groups and enantiomers. If the enantiomer has suitable polar substituents group such as hydroxyl, carbonyl, carboxyl, amino and phosphate, one or more favorable hydrogen bonds can be formed with the β -CD CSP. Additionally, repulsive interaction due to steric hindrance around the chiral atoms of CD provides conformational control that can advocate the chiral separation (Hinze *et al.*, 1985; Daffe & Fastrez, 1983). These properties of β -CD has led to its widely used as stationary phase, particularly in HPLC for the separation of chiral compounds (Juvancz & Szejtli, 2002).

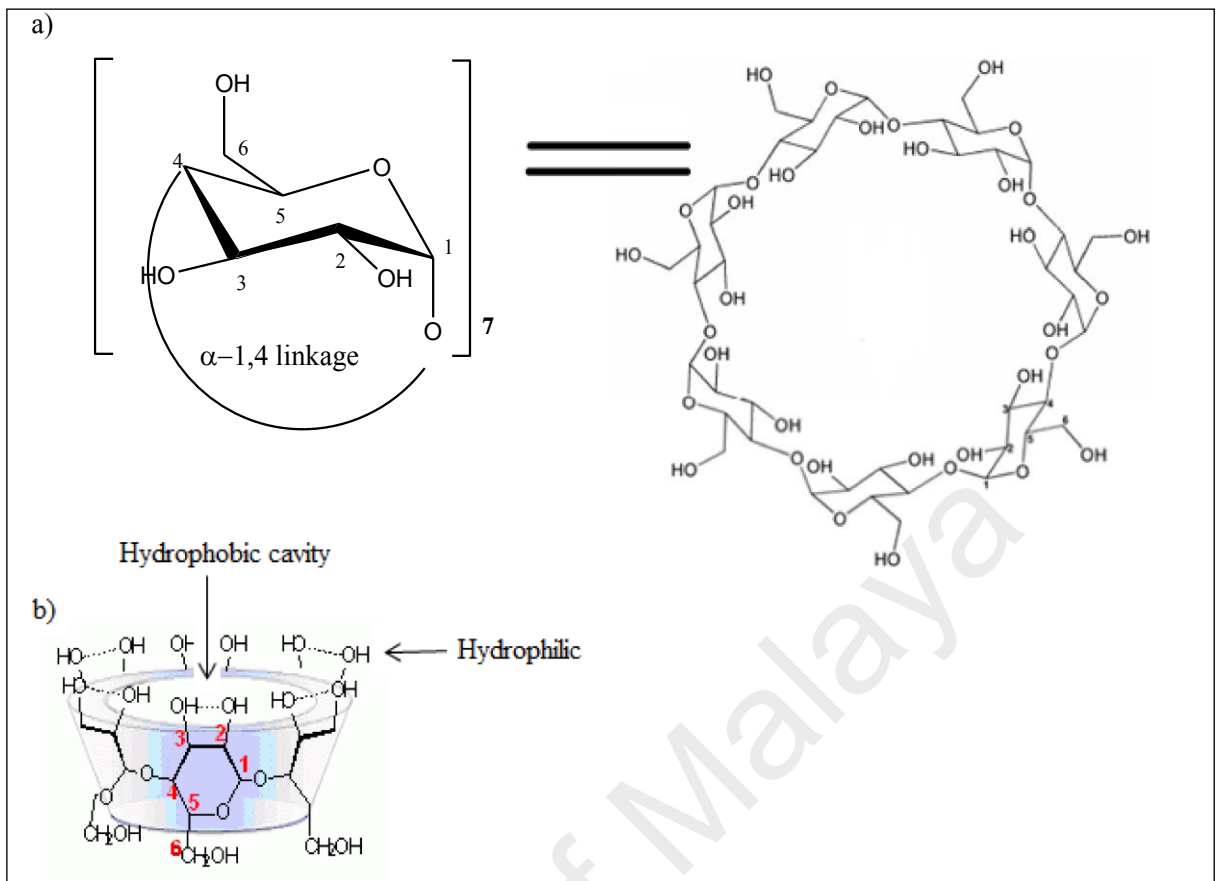


Figure 1.2 : a) Chemical structure of CD b) Molecular shape of CD

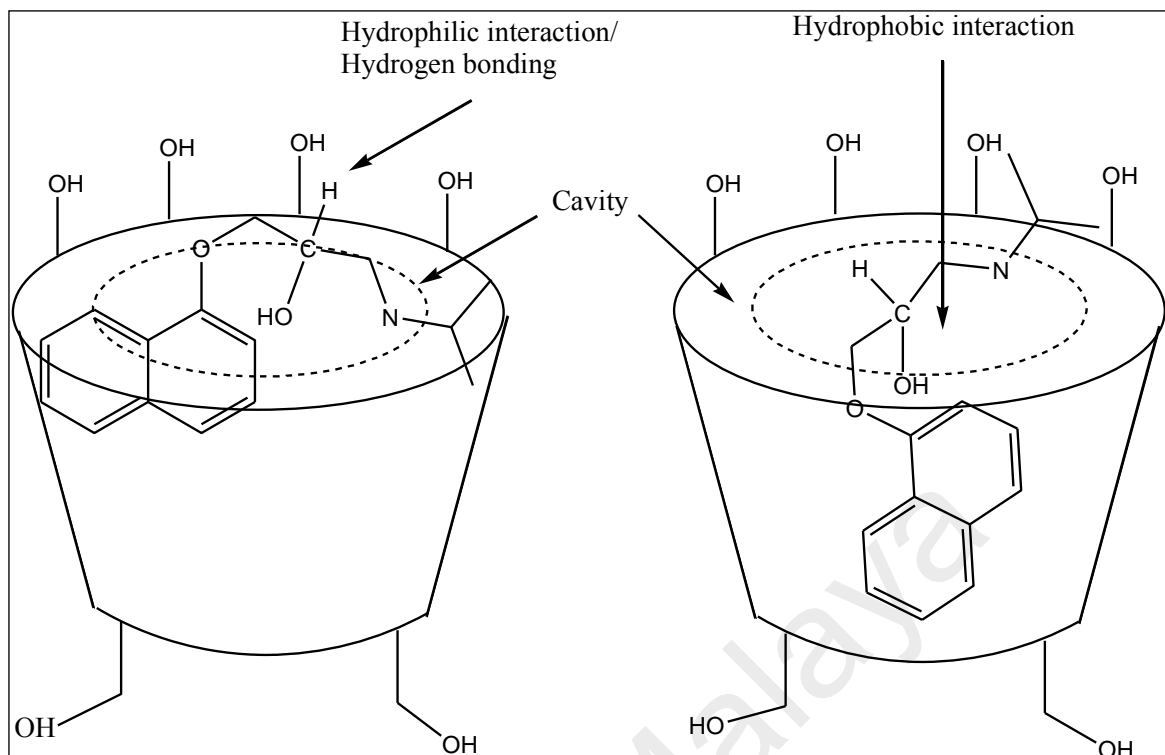


Figure 1.3: Illustration of the interaction between β -CD and enantiomer

In most cases, the cylindrical binding cavity of native β -CD is found to be too symmetrical to induce large enantioselectivities (Szejtli, 1994). Due to the native β -CD based CSP is unable to achieve satisfactory separation of enantiomers (Stalcup *et al.*, 1990), additional substituents are often introduced in order to achieve better chiral recognition. Therefore, various efforts have been directed toward developing new β -CD derivative-based CSPs to enhance the chiral separation (Wang *et al.*, 2010; Ciucanu, 1996; Ciucanu & Konig, 1994). Some common substitution groups that have been used to modify β -CD were alkyl, acetyl, benzoyl, hydroxypropyl, phenylcarbamoyl (naphthylethyl carbamoylated or 3,5-dimethylphenyl carbamoylated), *p*-toluoyl, carboxymethyl, pyridylethylene diamine and nitropyridylethylene diamine (Xiao *et al.*, 2009; Han *et al.*, 2005; Tang *et al.*, 2005a; Tang *et al.*, 2005b; Lipka *et al.*, 2003; Armstrong *et al.*, 1998; Chang *et al.*, 1992). Among various substitution groups, the aromatic ring substituted β -CD-based CSPs have been labeled as a multi-modal CSPs due to its ability to interact with enantiomers at various bonding sites. The aromatic

substituted β -CD-based CSPs not only afford hydrogen bonding effects and dipole-dipole interactions, but also hydrophobic and π - π interactions during enantioseparation. The different substitution groups on the aromatic ring can further alter the nature of π - π interaction to make them more suitable for the separation of various enantiomers. Recently, the 6-hydroxyl group of CD was bonded with ionic liquids (ILs) such as imidazole or pyridine in order to introduce additional π - π interaction and ionic interaction (Xiao *et al.*, 2009; Tang *et al.*, 2005a; Tang *et al.*, 2005b).

Ionic liquids (ILs) are a class of salt, in which the ions are poorly coordinated. Consequently, these compounds are in liquid form at the temperature of below 100 °C (Subramaniam *et al.*, 2010; Fontanals *et al.*, 2009). ILs has unique properties, such as non-volatility, non-flammability, low viscosity, and has chemical and electrochemical stability (McEwen *et al.*, 1999), and also can remain in the liquid state over a wide range of temperature. ILs could be hydrophobic and hydrophilic depending on the cationic and anionic characteristic. This dual nature role of ILs indicated their usefulness as stationary phase in chromatography (Anderson & Armstrong, 2003). On the other hand, ILs molecules also consist of high charge region and low charge region (Canongia Lopes & Padua, 2006). This property of ILs contributes to the electrostatic and dispersive interaction which useful for mechanism of enantioseparation (Anderson & Armstrong, 2003).

In this study, β -CD was first functionalized with ILs. The selected ILs were 1-benzylimidazole and 1-decyl-2-methylimidazole with tosylate as anion named β -CD-BIMOTs and β -CD-DIMOTs respectively. Then, β -CD functionalized ILs were then bonded onto modified silica gel to obtain CSPs. The performance of both CSPs for the enantioseparation was evaluated using flavonoids (flavanone, hesperetin, naringenin and eriodictyol), β -blockers (propranolol, metoprolol, pindolol and atenolol) and non-

steroidal anti-inflammatory drugs (NSAIDs) (ibuprofen, fenoprofen, indoprofen and ketoprofen). In addition, the mechanisms of enantioseparation were investigated experimentally through the inclusion complexes formation study. This inclusion complexes study gave an insight into the interaction between CSP and the selected analytes during HPLC separation.

1.2 Objectives of the research

The objectives of this study were:

1. To synthesis β -cyclodextrin functionalized ionic liquid (1-benzylimidazole and 1-decyl-2-methylimidazole) based CSPs.
2. To examine the performance of the synthesized CSPs for the separation of flavonoids, β -blockers and NSAIDs group with optimization of mobile phase.
3. To investigate the mechanism of separation of flavonoids, β -blockers and NSAIDs.

1.3 Outline of thesis

The present thesis is organized into five chapters. Chapter 1 gives a brief introduction on research background, research objectives, and scope of study. A review of related literature is presented in Chapter 2. Chapter 3 presents the experimental procedure for the synthesis of β -CD based-CSP and the preparation of inclusion complex. Chapter 4 discussed the characterization of the synthesized β -CD based-CSP, and the evaluation of synthesized CSPs performance and the mechanism of enantioseparation of flavonoids, β -blockers and NSAIDs. Finally, the overall conclusions, together with recommendations of future works are provided in Chapter 5.

CHAPTER 2: LITERATURE REVIEW

2.1 Chirality

The word “chiral” derives from the greek word “*cheir*” which means hand. In chemistry, chirality was first discovered by Louis Pasteur in 1848. Pasteur conducted an experiment in which he produced crystals salt known as racemic acid. The crystals were of divided into two forms, known as "+" and "-" forms, which is mirror images of one another. Pasteur shone polarized light through each solution of these salts, and found that the two solutions had equal but opposite optical activity. Thus, Pasteur identified, for the first time, the two enantiomers of a chiral substance, and recognized the existence of molecular chirality (Arjomandi-Behzad *et al.*, 2013). Chirality was later defined by Lord Kelvin in 1906 as the non-superimpose ability of a molecule on its mirror image (Evans & Kasprzyk-Hordern, 2014). Chiral molecules are also called optical isomers because the solutions of different enantiomer rotate plane-polarized light in different direction. The optical isomer or enantiomer which rotates plane-polarized light in the clockwise direction is designated as dextrorotatory (*D*) or (+)-enantiomer. In contrast, its antipode (e.g., opposite enantiomer) which rotates plane-polarized light in the counter clockwise direction is designated as levorotatory (*L*) or (–)-enantiomer (Agustian *et al.*, 2016). An equal mixture of each of the enantiomer is known as a racemic mixture (Zhang *et al.*, 2014).

Generally, molecular chirality is mainly due to the stereogenic centers of sp^3 hybridized carbon atoms that bear four different substituents. Apart from carbon, boron, nitrogen, phosphorus and sulphur also have stable chiral centers. The most important nomenclature system for denoting enantiomers is the *R/S* system. Absolute configuration of the isomer are performed by labeling each chiral center *R* or *S* according to a system by which each substituents are assigned a priority, according to

the Cahn-Ingold-Prelog priority rules (CIP), based on atomic number (Zhang *et al.*, 2014).



Figure 2.1: Examples of how to design configuration using Cahn-Ingold-Prelog priority rules

On a molecular level, chirality represents an intrinsic property of the “building blocks of life”, such as amino acids and sugars, and therefore, of peptides, proteins and polysaccharides (Zhang *et al.*, 2014). For example, amino acids are all presence in *L*-configuration rather than *D*-configuration. Meanwhile, natural sugars are presence in *D*-configuration. Consequently, metabolic and regulatory processes mediated by biological systems are sensitive to stereochemistry and different responses can be often observed when comparing the activities of a pair of enantiomers in biological system. Therefore, stereochemistry is an important consideration when studying xenobiotics, such as drugs, agrochemicals, food additives, flavors or fragrances. Drug action is the result of pharmacological and pharmacokinetic processes, by which it enters, interacts and leaves the body. Thus, straight regulations have been demanded by US Food and Drug Administration (FDA) towards marketing the single-enantiomer of drugs (Zhang *et al.*, 2014). FDA demands full documentation of pharmacological and pharmacokinetic (activity and toxicity) profiles of each individual enantiomer, as well as the racemic

mixture of drugs from the manufacturer. Therefore, it is necessary to have reliable analytical methods for the separation of each individual enantiomer and isolate the pure enantiomers. Chirality is also important in the agrochemical and food industry. In the food industry, a significant number of additives, flavors, fragrances and fumigants, preservatives, growth regulators, pesticides and herbicides are chiral molecules (Sekhon, 2013). Enantiomers in agrochemicals can have diverse effects on plants and insects, and cause negative effects to the environment and human health (Zsila, 2013). For examples, several European governments only allow the application of pesticide mecoprop and dichlorprop in the form of *R*-enantiomers (Author, 2004). All metalaxyl fungicidal activity is resided with the active *R*-enantiomer. The degradation of metalaxyl was shown to be enantioselective with the fungicidally active *R*-enantiomer being degraded faster than the inactive *S*-enantiomer, resulting in residues enriched with *S*-metalaxyl when the racemic compound was applied (Sekhon, 2013). In addition, *R*-enantiomer of fipronil, a phenylpyrazole insecticide, was more toxic to *Ceriodaphnia dubia* (water flea) than the *S*-enantiomer but in other studies the *S*-enantiomer was shown to have significantly more androgen and progesterone activity than the *R*-enantiomer (Negru *et al.*, 2015).

2.2 Enantiomeric separation technology

2.2.1 Development of chiral separation technologies

During the past decades, the requirement of enantiomeric separation emerges rapidly in the area of food safety, environmental analyses, agrochemical and drug industries (Bubalo *et al.*, 2014). In the preparation of single enantiomer, enantioseparation at analytical scale is important for determining enantiomeric purity (Dai *et al.*, 2013). Since enantiomers have identical physical and chemical properties except for the rotation of the plane of polarized light, chiral separation has been

considered as one of the most challenging tasks in chemistry. The enantioseparation can be divided in two classes: non-chromatography and chromatography.

For non-chromatography methods, Louis Pasteur discovered the spontaneous enantiomeric resolution by crystallizing separately each isomers of salt crystal as mentioned previously at section 2.1. After that, a considerable number of optical compounds were resolved mainly by fractional crystallization of the diastereomeric salts (Ismail *et al.*, 2016). Generally, reaction of a racemic acid or base with an optically active base or acid gives a pair of diastereomeric salts. Members of this pair exhibit different physicochemical properties (e.g., solubility, melting point, boiling point, adsorption, phase distribution) and can be separated owing to these differences by crystallization.

For chromatography methods, the earliest report of chiral separation was carried out by Gil-Av and his coworkers in 1966. They found that optically active stationary phase consisting of N-trifluoroacetyl-L-phenylalanine cyclohexyl ester was successfully applied to separate the enantiomers of trifluoroacetyl derivatives of some amino acids (Arjomandi-Behzad *et al.*, 2013). Since then, chromatography approaches are rapidly becoming the most commonly used enantioseparation approach in both analytical and preparative scale.

The publication for HPLC in the area of enantioseparation has been growing rapidly in recent years due to its easy-handling (Lin *et al.*, 2014). Separation of chiral compounds can be carried out using HPLC through direct and indirect methods. Indirect methods are based on the addition of chiral additive to the mobile phase. Direct methods separate the isomers on chiral stationary phases (CSPs). Generally, CSPs is prepared by adsorbing or covalently bonding the chiral selector onto solid support. Chiral selector is the chiral component of the separation system that is able to interact

enantioselectively with the enantiomers to be separated (Saleem *et al.*, 2013). Figure 2.2 illustrates the structures of the various chiral selectors. However, research findings have found that there are no universal CSP or chromatographic conditions which enabling the enantioseparation for all compounds. For most of the CSPs, small changes in the analytes' structures and/or chromatographic conditions would exert a strong impact on the efficiency of enantioseparation. Therefore, many parameters of chromatographic conditions in HPLC need to be optimized to resolve the enantiomers (Ismail *et al.*, 2016).

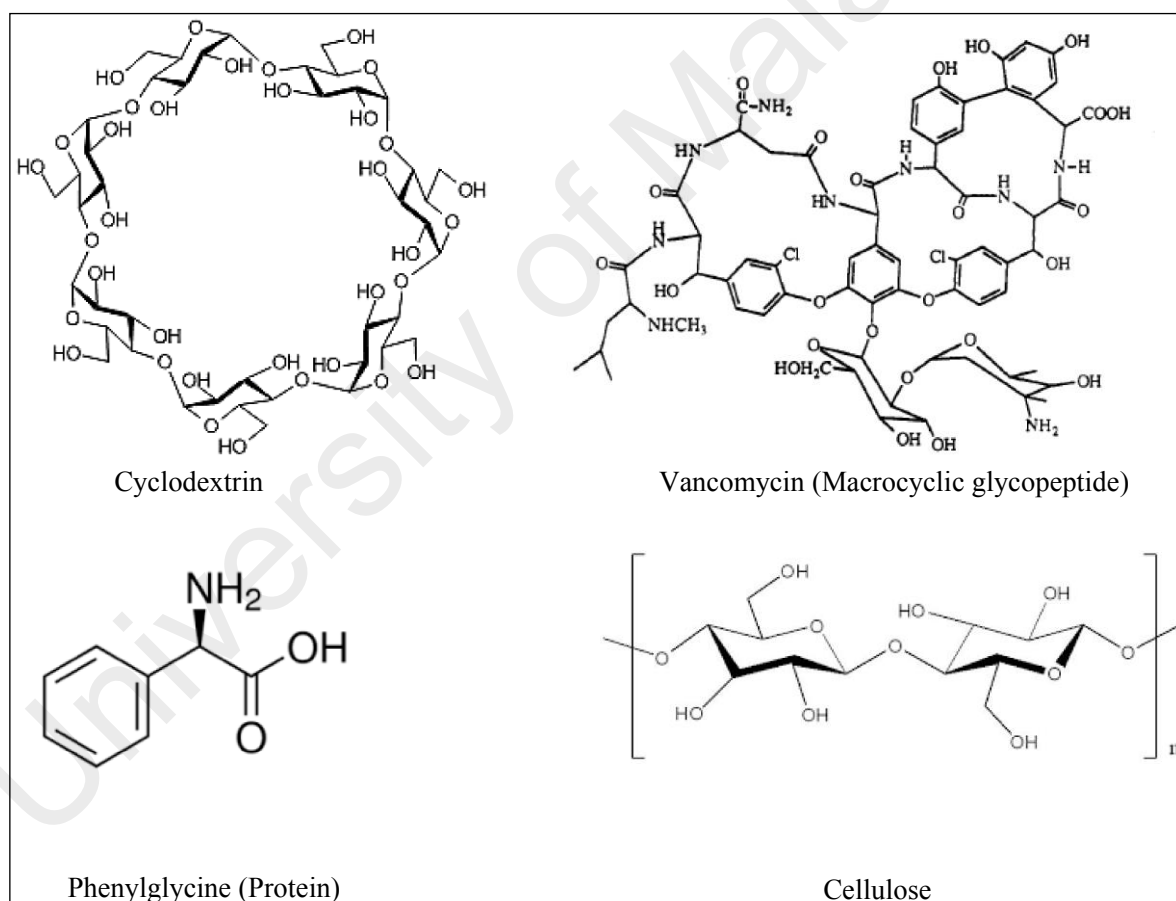


Figure 2.2: Common structures of chiral selectors

2.2.2 Development of chiral stationary phase

CSPs have been studied extensively since Davankov's review on the application of natural sorbents (proteins, carbohydrates, and optically active quartz) and also artificial dissymmetric sorbents (based on silica gel and activated carbon) as stationary phase for the ion exchange chromatography in the early 1970s (Arjomandi-Behzad *et al.*, 2013). Driven by the growth of asymmetric organic synthesis leading to chiral drugs, food additives, fragrances, agricultural chemicals and many other important chiral intermediates, the development of CSPs has grown rapidly. Various CSPs were developed and applied in various chiral resolution technologies. Firstly, Davankov *et al.* developed metal ion complexes for enantioseparations (Arjomandi-Behzad *et al.*, 2013). After that, by linking small chiral molecules onto stationary phase, brush type chiral stationary phases were prepared (Valente & Soderman, 2014). Pirkle *et al.* developed the first commercial column with brush type chiral stationary phase (Figure 2.3) for HPLC in 1981 (Valente & Soderman, 2014). Most recently, naturally occurred chiral macromolecules such as cyclodextrins, celluloses, macrocyclic glycopeptides and proteins were modified for the application of enantioselective processes (Wang *et al.*, 2011b).

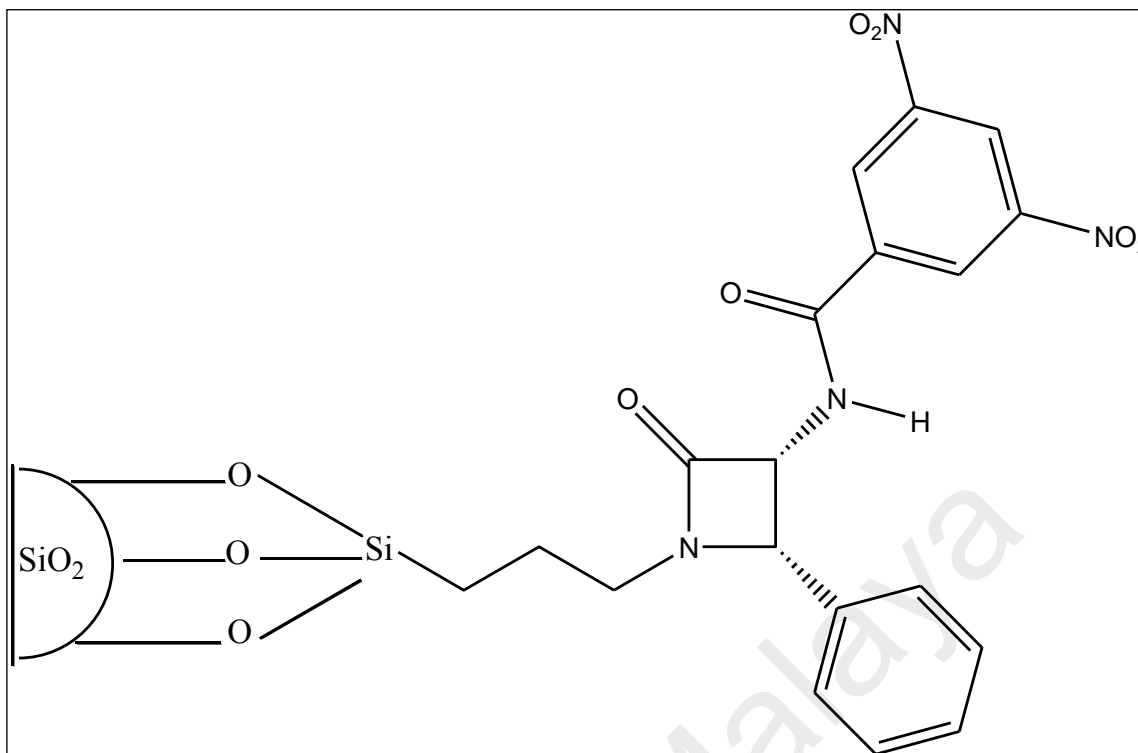


Figure 2.3: Molecular structure of the first commercial chiral column (Pirkle 1-J-column)-Brush type CSP

2.3 Cyclodextrin and its applications in enantioseparation

Cyclodextrins (CDs) are toroidal structural molecules. The α -, β -, γ - CD consist of six, seven and eight α -(1, 4)-linked D-(+)-glucose units, respectively (Figure 2.4). CDs are presence as chiral molecule due to the presence of chiral center of glucose units. The special properties of CDs originate from their unique truncated cone shape structures. The interior cavity of the cone is highly hydrophobic and the exterior is hydrophilic owing to hydroxyl (OH) group (Tang & Tang, 2013). The truncated cone of CDs consists of secondary OH groups at C2 and C3 and primary OH at C6 (Figure 2.4). The hydrogen at C1, C2, and C4 are located at the outside surface of the torus. The OH groups combined with the hydrogen atoms outside surface of CD build up a polar exterior to compatible with polar environments. The cavity interior is lined with the glucose ring oxygen atoms, as well as with the hydrogen atoms at C3 and C5 thus gives

the cavity some Lewis-base character (Zhang *et al.*, 2005). These characteristics endow CDs with a special capacity which can accommodate large variety of organic and inorganic compounds through inclusion complexation (Schurig & Juza, 2014).

As shown in Table 2.1, three types of CDs have different sizes of cavity. A general consideration is that small size hydrophobic organic molecules form the most stable complex with α -CD but the weakest with γ -CD. Secondly, neutral molecules generally bind more tightly with native CDs than their charged species. Compared with the α - and γ -CDs, β -CD is more widely investigated in separation science due to their high chemical stability and low cost. In addition, β -CD also has the special size of its hydrophobic cavity (cavity size: α -CD < β -CD < γ -CD) which affords to form inclusion complexes with numbers of organic and inorganic compounds (Valente & Soderman, 2014).

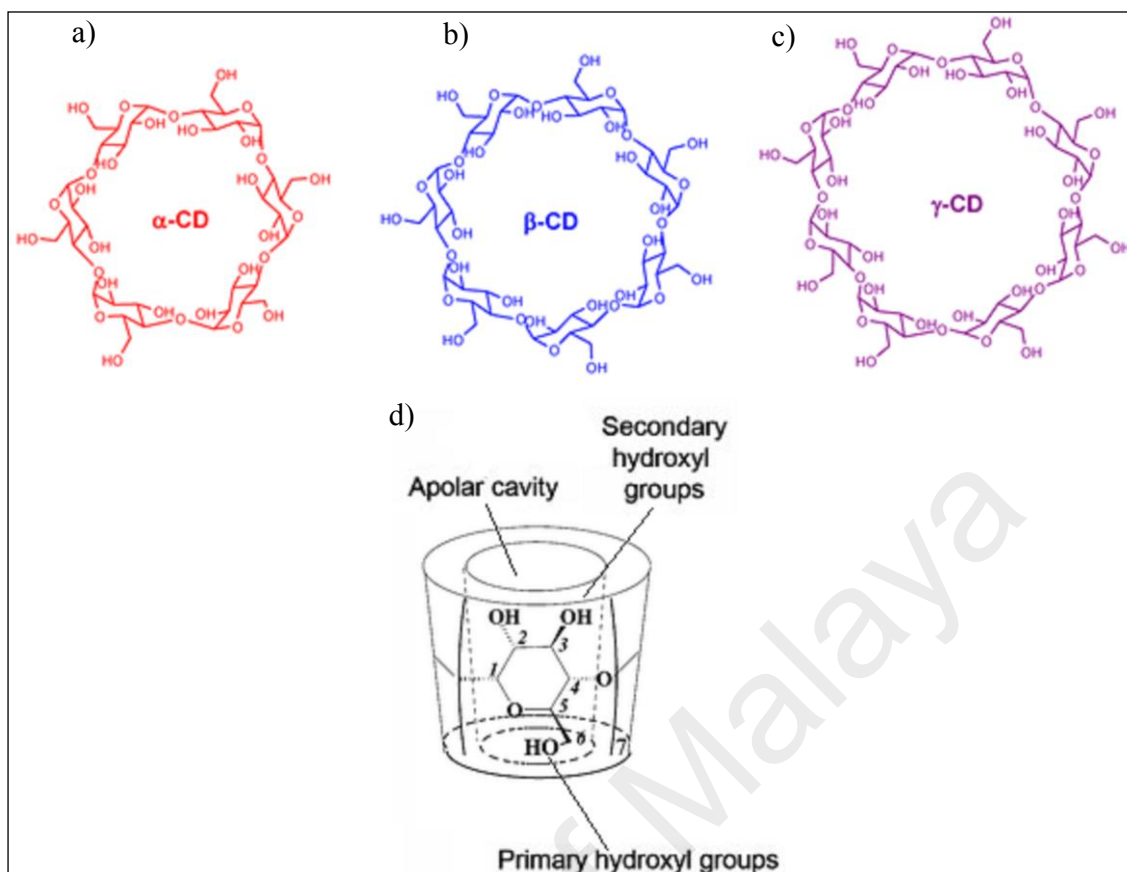


Figure 2.4: Illustration of a) α -CD, b) β -CD, c) γ -CD and d) side view of CD represent the position

Table 2.1: Physical and chemical properties of CD molecules (Bender & Komiyama, 2012)

Cyclodextrin	No of glucose units	Molecular mass (g/mol)	Cavity diameter (nm)	No. of stereogenic center	Water solubility (g/100 mL)
α	6	972	0.49	30	14.5
β	7	1135	0.62	35	18.5
γ	8	1297	0.79	40	23.3

For the mechanism of enantioseparation, according to Armstrong *et al.* (1986), there are a number of requirements for chiral recognition by CD. For example, an inclusion complex must be formed, and there must be relatively tight fit between the complexed moiety and the CD (Wang *et al.*, 2011b). The chiral center and one substituent of the chiral center of an analyte must be near and interacts with the mouth of the CD cavity. The unidirectional OH groups at C2 and C3 located at the mouth of CD cavity are particularly important in chiral recognition in order to satisfy the requirement of the “three-point” model. The “three-point” model was introduced by Pirkle at 1989 to elaborate the enantioseparation on CSPs (Valente & Soderman, 2014). According to Pirkle’s model, chiral recognition requires three interactions with at least one of them has to be stereoselective. Pirkle’s model can be illustrated by a representative enantioseparation in Figure 2.5.

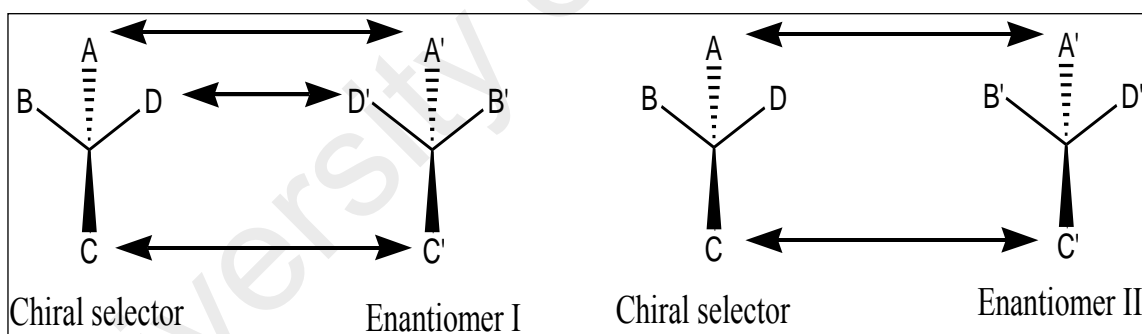


Figure 2.5: The “three point” model

As illustrated in Figure 2.5, three interactions of A—A', C—C' and D—D' between the chiral selector and enantiomer (I) whereas, only two interactions A—A' and C—C' are formed between chiral selector and enantiomer (II). The discrimination effect of the two enantiomers falls on the interaction of D—D' and resulting in the different of elution order of the two enantiomers.

The first application of CDs for enantioseparation was reported in 1959 in which CDs were employed as a selective precipitation or crystallization agent for occlusion compounds (Szente & Szemaan, 2013). From then on, CDs were studied either as mobile phase additives or stationary phases in chromatographic separation (Zhang *et al.*, 2015b). CDs derived stationary phases were originally designed for enantiomeric separation, structural and geometrical isomers separation. Early studies of CDs based stationary phases for enantioseparation focused on the polymerized CDs which were not robust in chiral discrimination and often overloaded with distorted peaks (Bender & Komiyama, 2012). Thereafter, researchers investigated the development of covalently bonded CD based CSPs. In 1984, the first stable CD CSP (Cyclobond I) with high coverage of the CD was developed by Armstrong & DeMond (1984). Subsequently, the CD derived CSPs were also commercialized by their group and hundreds of chiral compounds have been resolved on these CSPs using HPLC (Dai *et al.*, 2013).

The properties of the CD can be modified by replacing one or more primary or secondary OH groups with different moieties (Ong *et al.*, 2008). For CD, the three OH groups at the glucose units are different in reactivity due to the different acidities and sterical hindrance. Of the three types of OH groups present in CD rim, the most nucleophilic are primary OH at C6, the least nucleophilic are secondary OH at C2 and the most inaccessible are secondary OH at C3. This forms the basis for a broad spectrum of regioselective alkylations and acylations which have been applied to modify the CDs for CSPs (Schurig & Juza, 2014).

The modified CDs with certain functional moieties can provide potentially additional useful interaction sites and accommodate a variety of spatial requirements to produce highly selective separations for a versatile array of analytes. The substitution groups that have been incorporated onto CDs were alkyl, acetyl, hydroxypropyl,

phenylcarbamoyl groups (naphthylethyl carbamoyl or 3,5-dimethylphenyl carbamoyl) (Figure 2.6) (Dai *et al.*, 2013).

Generally, the OH groups, especially the secondary OH groups allow CD to interact with analytes via hydrogen bonding or dipole-dipole interaction. Although methylation of the OH groups reduced the hydrogen bonding sites but it enlarges the hydrophobic cavity and thus, enhances the steric interactions. These CSPs exhibit good enantioselectivities to some specific solutes such as furan derivatives, tetralins and melatonin ligand. The chiral recognition of these CSPs is implemented through hydrophobic and steric interactions between the analytes and the methoxy groups on the CD rim after inclusion complex formation (Han *et al.*, 2005; Lipka *et al.*, 2003). Since methylation could not introduce diverse effective interaction sites (like hydrogen bonding and π - π interaction sites), these CSPs are less effective towards a wide range of chiral compounds.

Hydroxypropylated CD-based CSPs (Figure 2.6 (iii)) have been considered as a very successful CSP. The OH groups of this CD derivative increase the flexibility of hydrogen bonding and provide additional hydrogen bonding sites with analyte. Many chiral compounds that are partially resolved on unmodified CD-based CSP could undergo baseline resolution using similar separation conditions on these hydroxypropylated CSPs. Enhanced enantioseparation of some important drugs like conazoles, methadone, sertraline, Jacobsen's Catalyst and strigol can be achieved using 2-hydroxypropyl- β -CD (Liu *et al.*, 2015). However, the preparation process for these CSPs is relatively tedious and costly.

Substituted phenyl or naphthylethyl carbamoylated CD CSPs (Figure 2.6 (iv)) have been labeled as multi-modal CSPs due to their various bonding sites. It is not only afford hydrogen bonding effects and dipole-dipole interactions but also hydrophobic

and π - π interactions. In addition, the different substitution groups on the aromatic rings can enhance the nature of π - π interaction to make them more suitable for the separation of various racemates. Besides, an ionic interaction site was introduced by incorporating ionic liquid (IL) moiety such as imidazole or pyridine groups into the structure of CD and make them suitable for the enantioseparation of charged and polar analytes (Wang *et al.*, 2012b, 2012a; Wang *et al.*, 2012c; Wang *et al.*, 2008).

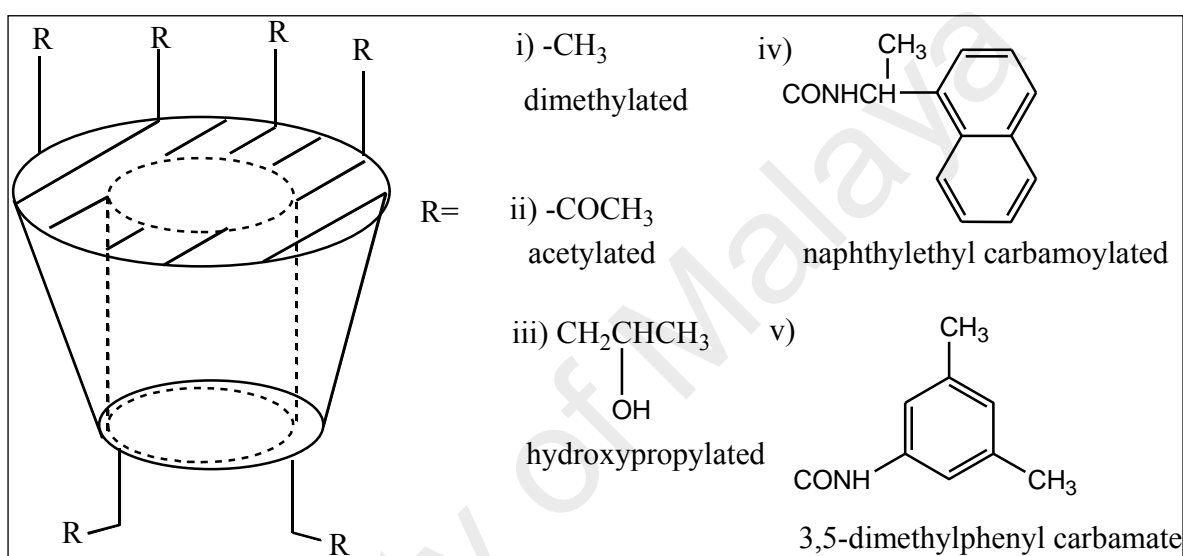


Figure 2.6: Common derivatives group of CD

2.4 Ionic liquid in enantioseparation

Ionic liquids (ILs) belong to salt-like materials which are liquid below 100 °C and even below room temperature (Yao *et al.*, 2014b). As salts they are by essence made of cation and anion. The term ILs covers inorganic as well as organic molten salts. ILs are usually composed of bulky, nonsymmetrical organic cation such as alkyl-imidazolium, pyridinium or pyrrolidinium, ammonium or phosphonium. Anions could be inorganic, including chloride, tetrafluoroborate, or hexafluorophosphate (Figure 2.7) (Bubalo *et al.*, 2014). The anion is not necessarily to be inorganic; ILs possessing

organic anions such as tosylate and methanesulfonate are also commercially available (Figure 2.7).

Owing to tunable properties which can be selected by choosing appropriate cationic or anionic constituents, they can be applied as mobile phase additive or stationary phase in chromatographic analysis. Compared with ILs used as mobile phase additives in HPLC, the application of ILs as stationary phases is fewer. Armstrong *et al.* (1999) and Anderson and Armstrong (2003) applied the ILs (1-Butyl-3-methylimidazolium hexafluorophosphate [BMIM][PF₆] and chloride [BMIM][Cl]) as stationary phases for gas chromatography (Zhang *et al.*, 2015a). They claimed that the dual nature of ILs is the main factor that contributed to the effective separation of polar and nonpolar compounds. Afterward, the applications of ILs in chromatography have been increased significantly.

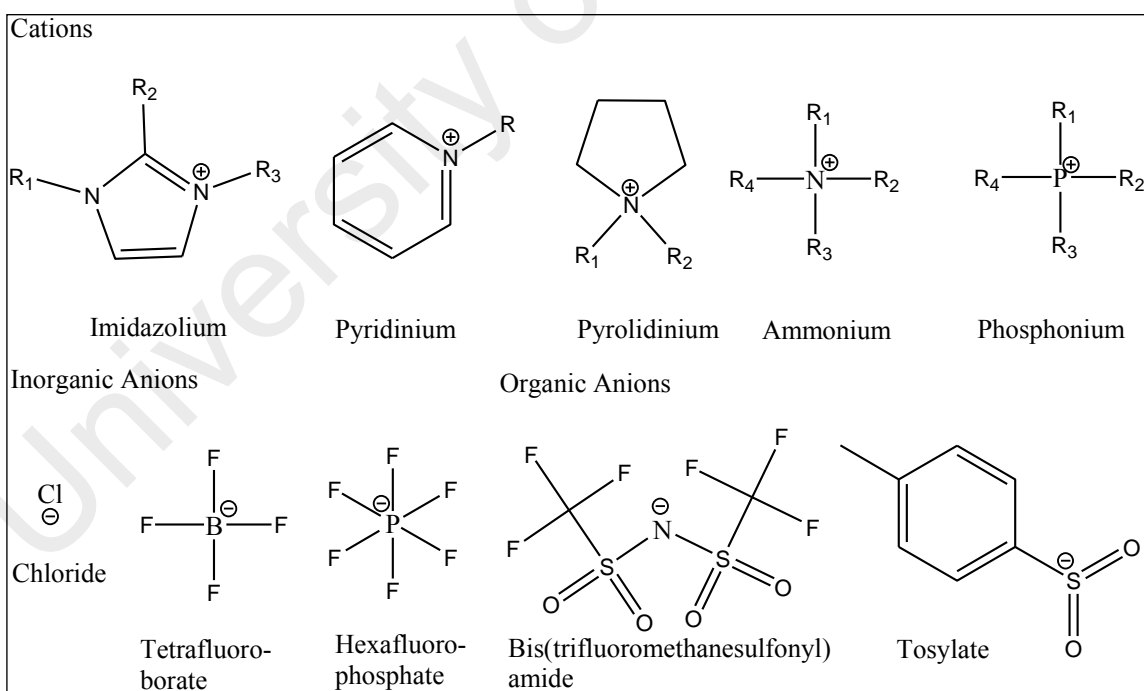


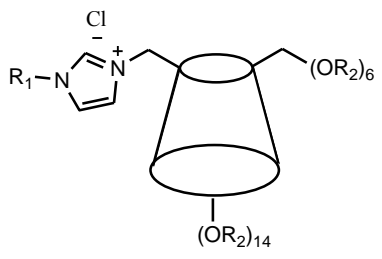
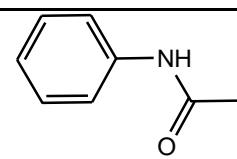
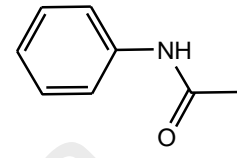
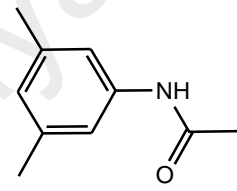
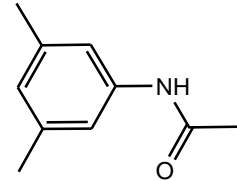
Figure 2.7: Common structures of cation and anion of ILs

Extending ILs to the realm of chiral separations has been done in two ways: (1) the ILs itself can be chiral or (2) a chiral selector can be dissolved in an achiral ILs. The first approach is not popular since the synthesis of chiral ILs is tedious and required expensive reagents. Thus, the second approach is the most preferred method. Modification of chiral selector with ILs yielded the CSPs with ion exchange properties. Consequently, the chiral separation mechanism involving ILs relies on multi modal interaction such as donor-acceptor interactions (hydrogen bonding, π - π interaction) and ionic interactions.

Lately, Wang *et al.* (2008) have physically coated a series of alkyimidazolium modified β -CD onto porous spherical silica gel to develop a series of β -CD-IL based CSPs namely mono-6-(3-methylimidazolium)-6-deoxy-perphenylcarbamoyl- β -CD chloride (MPCCD), mono-6-(3-methylimidazolium)-6-deoxyper (3,5-dimethylphenylcarbamoyl)- β -CD chloride (MDPCCD), mono-6-(3-octylimidazolium)-6-deoxyperphenylcarbamoyl- β -CD chloride (OPCCD) and mono-6-(3-octylimidazolium)-6-deoxyper (3,5-dimethylphenylcarbamoyl)- β -CD chloride (ODPCCD) (Table 2.2). These CSPs were used for the chiral separation of 18 aryl alcohols using HPLC and supercritical fluid chromatography (SFC). Among these CSPs, OPCCD, consisting of an *n*-octyl group on the imidazolium moiety and phenylcarbamoyl groups, exhibited the best separation ability for the aryl alcohols. Chromatographic studies revealed that the CSPs consisting of long alkyl group on the imidazolium moiety on the CD ring can provide enhancement of analyte-chiral substrate interactions over CSPs bearing the short alkyl group on the imidazolium moiety on the CD ring.

Later, Wang prepared another two β -CD-ILs CSP by graft polymerization of 6^A-(3-vinylimidazolium)-6-deoxyperphenylcarbamate- β -CD chloride or 6^A-(N,N-allylmethylammonium)-6-deoxyperphenylcarbamoyl- β -CD chloride onto silica to obtain VIMPCCD-POLY and VAMPCCD-POLY CSPs, respectively (Wang *et al.*, 2012b; Wang *et al.*, 2012c). These CSPs were used to separate the enantiomers of 12 pharmaceuticals and six carboxylic acids under reverse phase and normal phase mode. VIMPCCD-POLY exhibited higher enantioselectivities towards most of the selected analytes than VAMPCCD-POLY in normal-phase HPLC (Wang *et al.*, 2012c). The higher enantioselectivity was attributed to the additional π - π conjugation and electrostatic interactions formed with the aromatic imidazolium moiety. Meanwhile, the planar imidazolium moiety was found to make the CSP more accessible to the analytes than the tetrahedral ammonium moiety. The chiral separation abilities of VAMPCCD-POLY and VIMPCCDPOLY were also compared in SFC (Wang *et al.*, 2012a). The electrostatic force generated from the cationic imidazolium moiety was found to be important in the retention and chiral separation of 14 racemates, encompassing flavanones, thiazides and amino-acid derivatives.

Table 2.2: Chemical structures of the cationic functionalized β -CDs (Wang *et al.*, 2008)

Chemical structure	CSPs	R ₁	R ₂
	MPCCD	-CH ₃	
	OPCCD	-C ₈ H ₁₇	
	MDPCCD	-CH ₃	
	ODPCCD	-C ₈ H ₁₇	

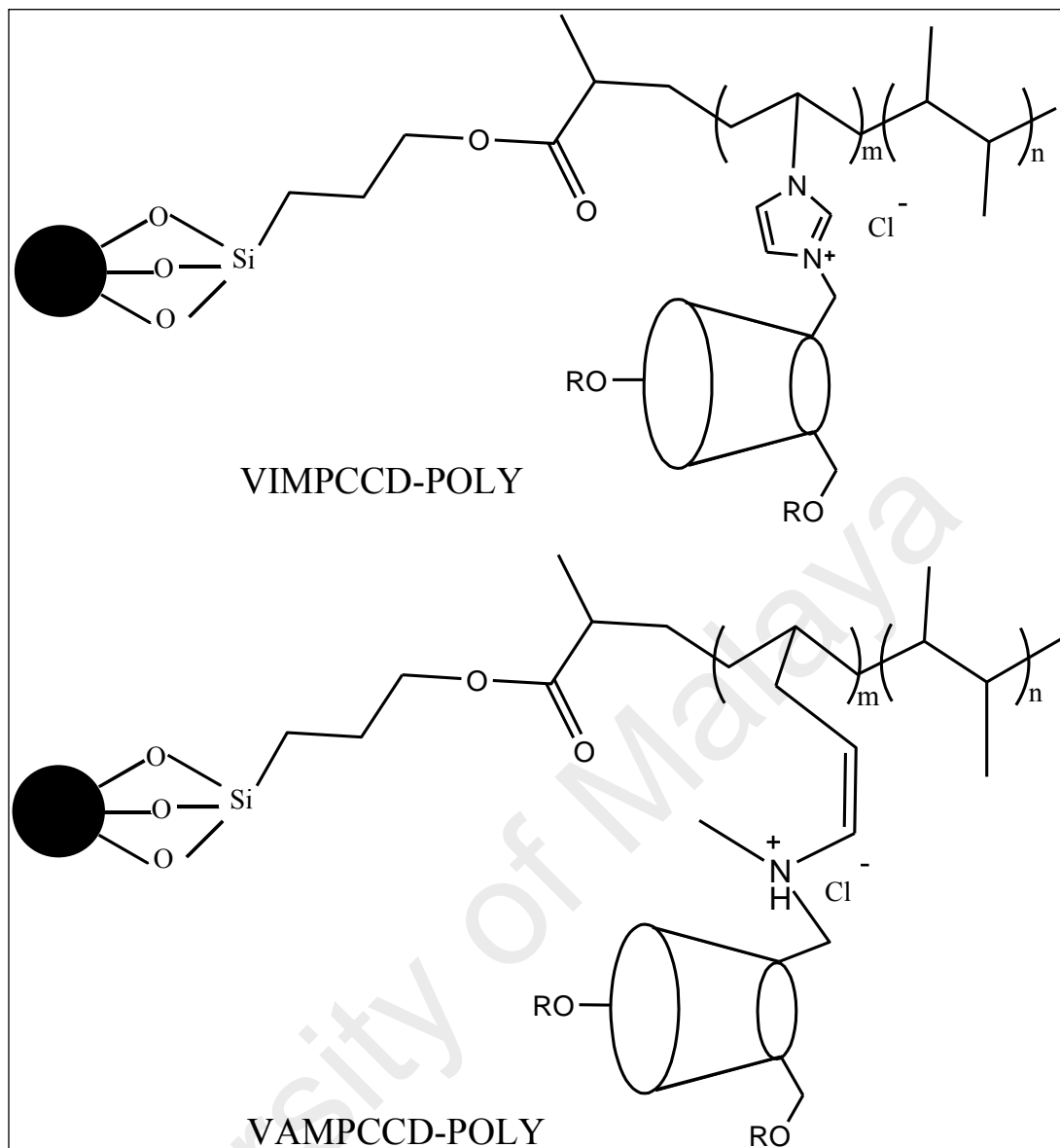


Figure 2.8: Structures of VIMPCCD-POLY and VAMPCCD-POLY CSPs (Wang *et al.*, 2012c)

Cooperative effects of β -CD and ILs as CSPs have been studied by Zhou *et al.* (2010) who functionalized β -CDs with ILs. Zhou *et al.* (2010) substituted the 6-tosyl- β -CD with 1,2-dimethylimidazole (Figure 2.9 (i)) or 1-amino-1,2,3-triazole (Figure 2.9 (ii)). Then, the functionalized β -CDs-ILs was bonded to silica gel to obtain CSPs. The presence of ILs was found to enhance the enantioselectivity of the synthesized CSPs towards α -nitro alcohol, α -hydroxylamine and aromatic alcohol. Zhou *et al.* (2010) stated that the π -conjugation through lone pair electron of NH_2 in 1-amino-

1,2,3-triazole was electronically stronger than the π -conjugation through the two CH_3 groups in 1,2-dimethylimidazole. Therefore, 1-amino-1,2,3-triazole cation was much more electronically stabilized. Consequently, 1-amino-1,2,3-triazole cation forming a loose ion pair with its counter ion (OTs^- or NO_3^-) and it was more readily participates anionic exchange with analytes. Whereas 1,2-dimethylimidazole cation has a higher affinity to anion and could form a tight ion pair (Zhang & Lv, 2006) with its counter ion (OTs^- or NO_3^-). CSPs containing 1-amino-1,2,3-triazole was found to lead to the higher resolution factors for the acidic analytes. Moreover, the CSPs consist of NO_3^- anion paired with either 1,2-dimethylimidazole or 1-amino-1,2,3-triazole cation always provided higher resolutions than the CSPs consist of OTs^- anion. It was suggested that NO_3^- anion has more hydrogen bonding sites and less sterically hindered to easier the interaction with the analytes.

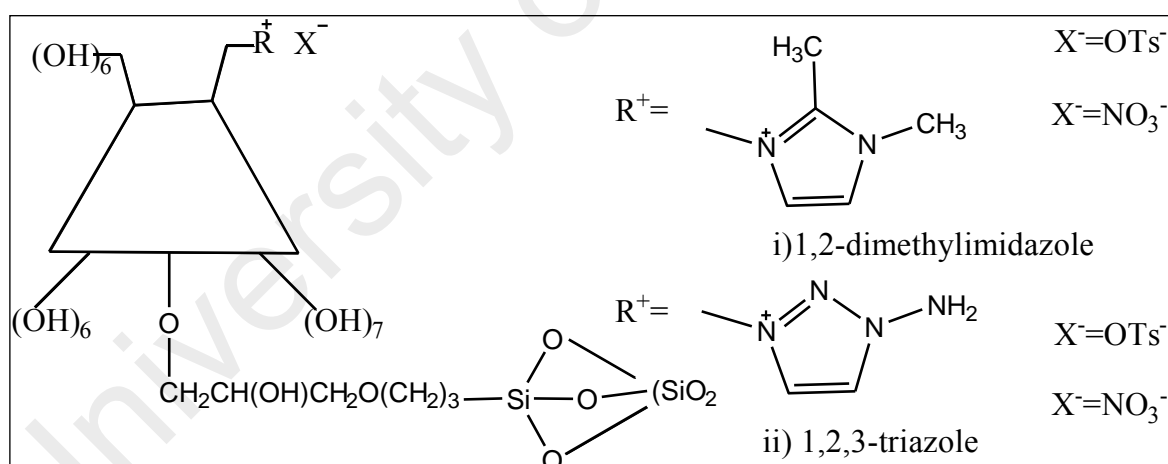


Figure 2.9: Structure of functionalized IL-bonded CSPs (Zhou *et al.*, 2010)

Recently, Yao *et al.* (2014b) has applied the simple thiol-ene click chemistry to anchor vinyl imidazolium β -CD onto thiol silica to form a novel β -CD-based CSP with ionic property (Figure 2.10 (i)). This new CSP enhanced chiral separation towards dansyl (Dns) amino acids, carboxylic aryl compounds and flavonoids by HPLC as compared with CSP that prepared through azide/alkynyl click reaction (Yao *et al.*,

2014b) . At the same year, Yao *et al.* (2014a) has synthesized triazole-bridged β -CD CSP. The performance of triazole-bridged β -CD CSP (Figure 2.10 (ii)) was compared with the previous thioether-bridged β -CD CSP (Figure 2.10 (i)) for enantioseparation of 26 isoxazoline derivatives. Most of the selected analytes was well resolved ($R_s > 1.5$) under reversed phase mode for both CSPs.

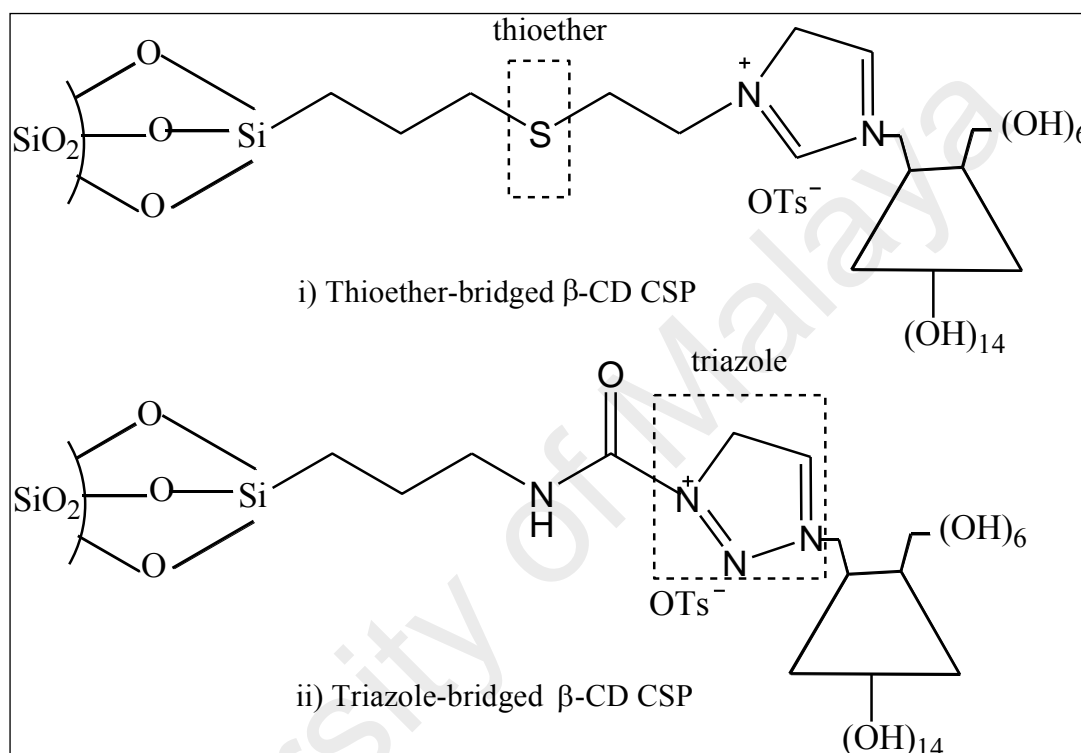


Figure 2.10: Structure of Thioether-bridged β -CD and Triazole-bridged β -CD CSPs (Yao *et al.*, 2014a)

Li *et al.* (2014) prepared four β -CD derivatives functionalized by ILs, in which the substituents and β -CD cavity are linked by a CH₂-N=C bonding and the corresponding CSPs based on silica were namely (a) mono-6-deoxy-6-(p-N,N,N-trimethylaminobenzimide)- β -CD nitrate CSP, (b) mono-6-deoxy-6-(p-N,N,N-trimethylamino-benzimide)- β -CD tosylate CSP, (c) mono-6-deoxy-6-(p-N-methylimidazolemethyl-benzimide)- β -CD nitrate CSP and (d) mono-6-deoxy-6-(p-N-methylimidazolemethylbenzimidide)- β -CD tosylate CSP. The excellent enantioseparation

was obtained for most of 1-phenyl-2-nitroethanol derivatives, aromatic alcohol and ferrocene derivatives. The analytes with small volume was found to achieve better enantioseparation on CSP (b) with smaller volume of cation and anion. Thus, they summarized that not only the structure matching between β -CD derivatives and the analytes that contributed to the enantioseparation, but the cooperation of cationic and anionic substituents also play a significant role in the enantioseparation.

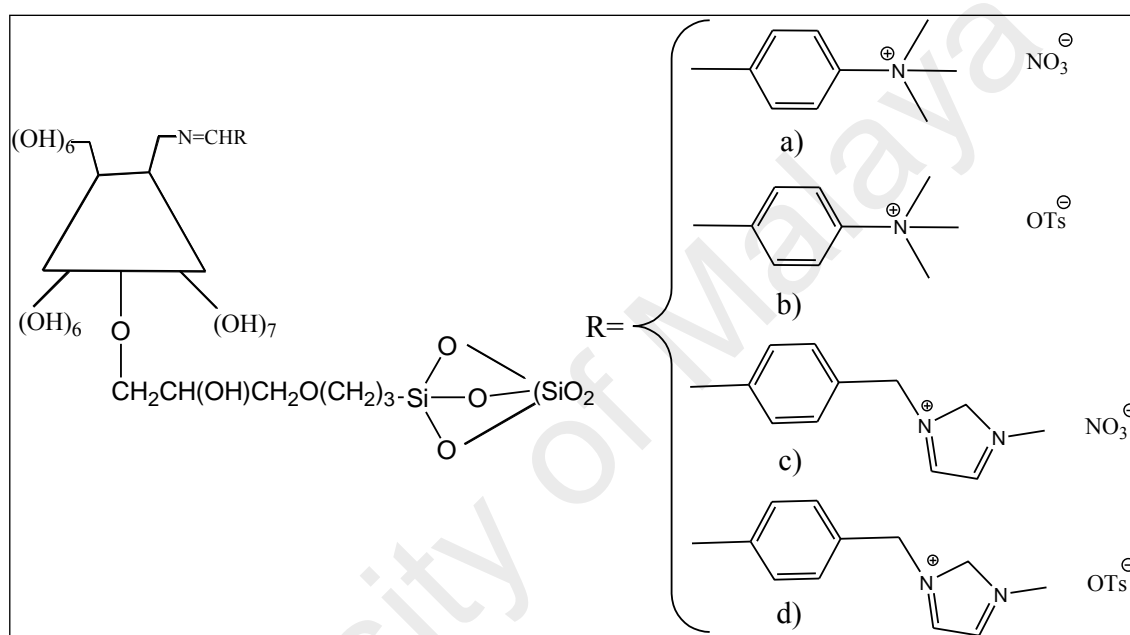


Figure 2.11: Structure of β -CD derivatives functionalized by ILs (Li & Zhou, 2014)

Liu *et al.* (2015) successfully fabricated the IL, 1-ethyl-3-methyl-imidazolium L-proline (EMIMLpro) onto the surface of $\text{Fe}_3\text{O}_4@\text{SiO}_2$ nanospheres. Complete resolution for separation of tryptophan racemate via the $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{HMIDI-EMIMLpro}$ nanospheres (Figure 2.12) was eventually achieved by centrifugal chiral chromatography using a spiral tube assembly mounted on a type-J coil planet centrifuge. The newly synthesized nanosphere are promising materials for chiral separation of racemates, because they can provide a huge surface area to accommodate

chiral selectors and are easy to be recycled through an external magnetic field (Liu *et al.*, 2015b).

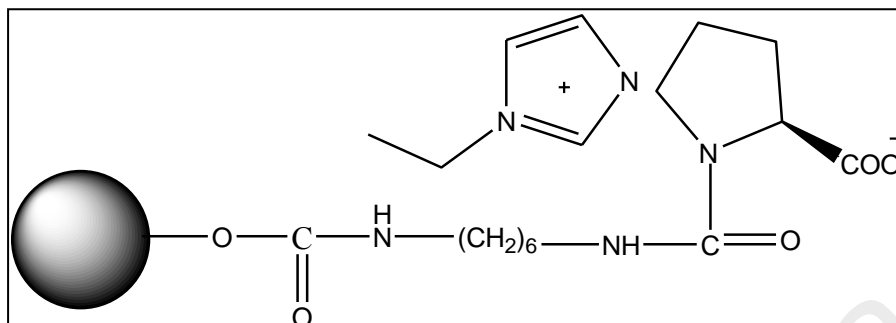


Figure 2.12: Structure of $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{HMDI-EMIMLpro}$ (Liu *et al.*, 2015b)

A novel amino acid IL, tetramethylammonium L-hydroxyproline (Figure 2.13), was first applied as a chiral ligand to evaluate its enantioselectivity towards several aromatic amino acids in ligand-exchange capillary electrophoresis (LE-CE) and ligand-exchange micellar electrokinetic capillary chromatography (LE-MEKC) (Liu *et al.*, 2015a). In the LE-CE system, excellent separations were achieved for tryptophan and 3, 4-dihydroxyphenylalanine. Meanwhile, the separations of the enantiomers of tryptophan, phenylalanine, and histidine were all improved in LE-MEKC system.

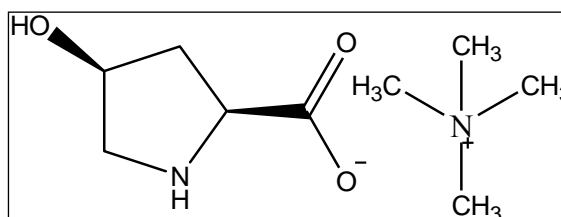


Figure 2.13: Structure of tetramethylammonium L-hydroxyproline (Liu *et al.*, 2015a)

The latest research based on CD functionalized IL was reported by Li *et al.* (2016). Li and co-workers were prepared and evaluated four single thioether bridged cationic CD CSPs with different spacer length, selector concentration and rim functionalities (Figure 2.14). The enantioseparation ability of prepared CSPs were evaluated by separating over forty enantiomers including isoxazolines, dansyl amino acids, flavonoids, tröger's base, 4-chromanol, bendroflumethiazide and styrene oxide. Most of the enantiomers were well resolved (Li *et al.*, 2016).

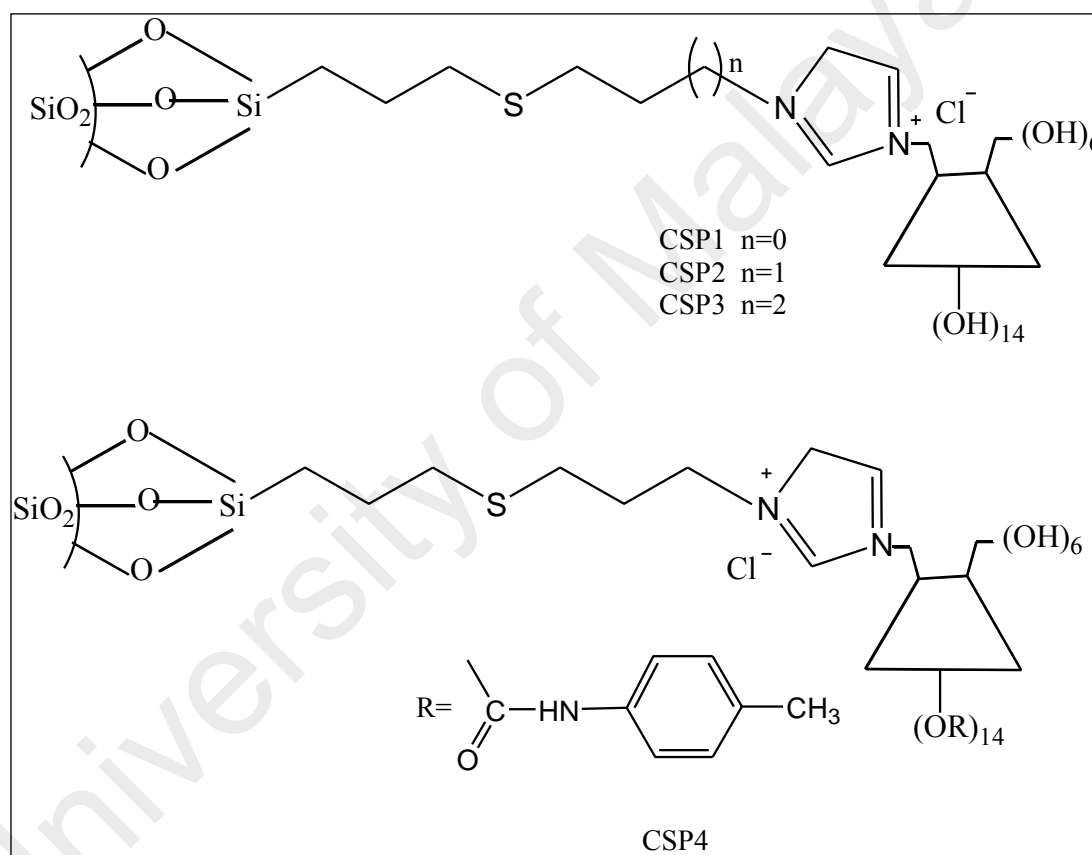


Figure 2.14: Novel cationic CSP (Li *et al.*, 2016)

2.5 Selected chiral compounds

2.5.1 Flavonoids

Flavonoids are a class of secondary metabolites of the plant and fungus. Chemically, they have the general structure of a 15-skeleton (15 carbon atoms), which consists of two phenyl rings (A and B) and a heterocyclic ring (C) (Figure 2.15). Flavonoids are divided into subclasses as showed in Table 2.4.

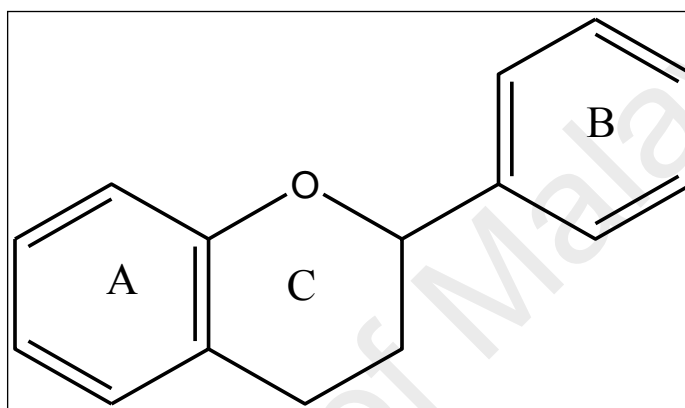


Figure 2.15: Basic chemical structure of flavonoid

Within the large family of flavonoids, flavanones possess a unique chiral structural which distinguishes them from all other classes of flavonoids. All the flavanones have a chemical structure based on a $C_6-C_3-C_6$ (Figure 2.16) configuration consisting of two aromatic rings joined by a three-carbon link (Tiwari *et al.*, 2013). Flavanones present a single stereogenic center at C (2) of chromanone core (Figure 2.16).

Among various flavanones, hesperetin, naringenin and eriodictyol (Figure 2.17) are the most abundant flavonoids that widely distributed in plants. Traditionally, researchers are attracted with the organoleptic properties of flavanones, such as bitterness or taste (Zid *et al.*, 2015). In recent decades, flavanones are increasingly being recognized for their nutritional value since they may reduce the risk of chronic diseases and in general it gives a positive effect to the health (Tucker & Robards, 2008;

Scalbert *et al.*, 2005). Recent studies have shown that naringenin possesses activities such as anti-inflammatory (Park *et al.*, 2012), anticancer (Sabarinathan *et al.*, 2011, 2010), antimetastasis (Qin *et al.*, 2011), normalizing lipids (Cho *et al.*, 2011; Goldwasser *et al.*, 2010), anti-hyperglycemia (Annadurai *et al.*, 2012), and anti-hypercholesterolemia (Chanet *et al.*, 2012). Eriodictyol can provide a cytoprotective effect in ultraviolet (UV)-irradiated keratinocytes (Lee *et al.*, 2011), induce long-term protection in ARPE-19 cells (Johnson *et al.*, 2009), and prevent early retinal and plasma abnormalities in streptozotocin induced diabetic rats (Bucolo *et al.*, 2012).

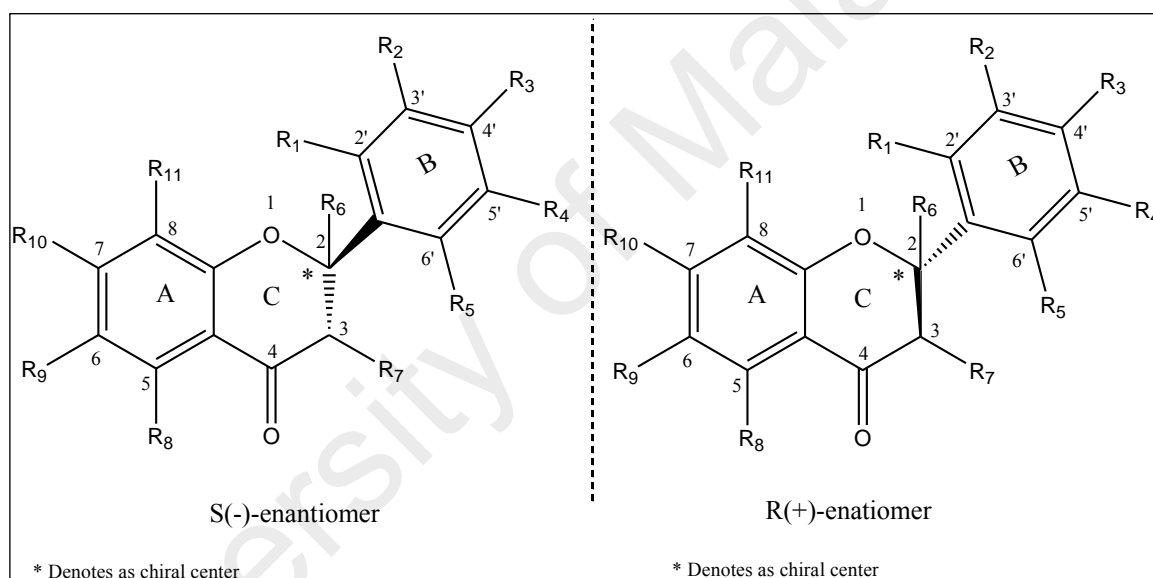


Figure 2.16: Spatial dispositions of the enantiomers of chiral flavanones

Table 2.3: Common dietary flavonoids

Flavonoids subclass	Dietary flavonoids	Common food source
Anthocyanidins	Cyanidin, Delphinidin, Malvidin, Pelargonidin, Peonidin, Petunidin	Red, blue, and purple berries; red and purple grapes; red wine
Flavonols	Monomers (Catechins): Catechin, Epicatechin, Epigallocatechin Epicatechin gallate, Epigallocatechin gallate Dimers and Polymers: Theaflavins, Thearubigins, Proanthocyanidins	Catechins: Teas (particularly green and white), chocolate, grapes, berries, apples Theaflavins, Thearubigins: Teas (particularly black and oolong) Proanthocyanidins: Chocolate, apples, berries, red grapes, red wine
Flavanones	Hesperetin, Naringenin, Eriodictyol	Citrus fruit and juices, e.g., oranges, grapefruit, lemons
Flavonols	Quercetin, Kaempferol, Myricetin, Isorhamnetin	Widely distributed: yellow onions, scallions, kale, broccoli, apples, berries, teas
Flavones	Apigenin, Luteolin	Parsley, thyme, celery, hot peppers
Isoflavones	Daidzein, Genistein, Glycitein	Soybeans, soy foods, legumes

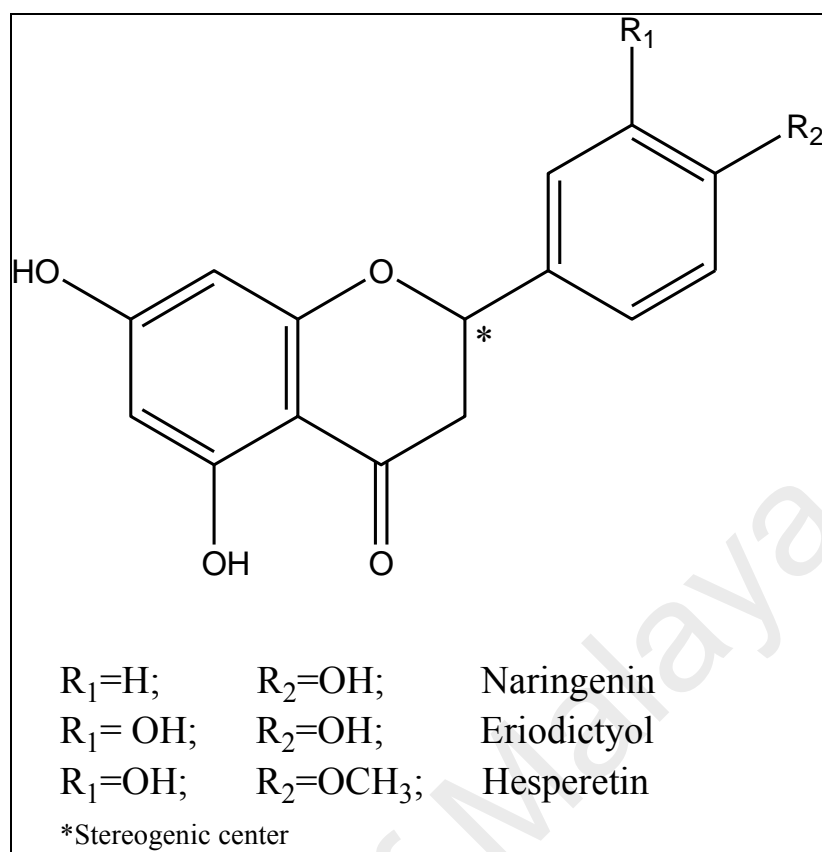


Figure 2.17: Chemical structures of some flavanones

The vast majority of flavanones can be purchased from chemical companies, but they are mainly available as racemates. Until now, there are only three stereochemically pure flavanones that are currently marketed internationally. Eriodictyol is marketed as the pure *S*-enantiomer by Fluka (Buchs, Switzerland). Homoeriodictyol is marketed as the pure *S*-enantiomer by Indofine Chemical Company (Hillsborough, NJ), Extrasynthese (Genay, France), and ITI International Inc. (Miami, FL). Finally, taxifolin is marketed as the pure *2R, 3R*-enantiomer by Alexis Biochemicals (San Diego, CA), Fluka (Buchs, Switzerland), and Extrasynthese (Genay, France) (Yanez *et al.*, 2007). As pharmaceutical related compounds, biological activity of flavonoids may result from a single enantiomer. Therefore, there is a need for stereospecific assay methods for the quantitation and effectively isolate the pure flavonoid enantiomers for their pharmacometric study in *in vivo* and *in vitro* models.

2.5.2 β -blocker drugs

β -adrenergic blocking agents (β -blockers) are basic drug that are frequently used for the treatment of angina pectoris and cardiovascular (Saleem *et al.*, 2013). β -blockers competitively binds to β -adrenergic receptor located at the heart and /or nonvascular smooth muscle. β -blockers inhibit the action of adrenergic agents (stimulants) by reducing the force of the heart muscle contraction and tend to reduce the heart rate. These drugs do not seem to produce vasodilation (widening of blood vessels resulting relaxation of the muscular walls of the vessels) as in the case of α -adrenergic blocking agents (Arjomandi-Behzad *et al.*, 2013). It is well known that β -blockers are chiral and their enantiomers have different potential of pharmacological and therapeutic effects (Evans & Kasprzyk-Hordern, 2014). *L*-isomer of all β -blockers is more potent in blocking β -adrenoceptors than their *D*-isomer. For example, *S*(-)-propranolol is 100 times more active than its *R*(+)-propranolol (Evans & Kasprzyk-Hordern, 2014). It has been demonstrated that *R*-propranolol can inhibit the conversion of thyroxin (T4) to triiodothyronin (T3) (Stoschitzky *et al.*, 1992; Harrower *et al.*, 1977; Wiersinga & Touber, 1977). Therefore, *R*-propranolol might be used as a specific drug without β -blocking effects to reduce plasma concentrations of T3 particularly for patients who suffering from hyperthyroidism. Meanwhile, racemic propranolol cannot be administered because of contraindications for β -blocking drugs (Stoschitzky *et al.*, 1998). Therefore, it is important to isolate and separate the enantiomer of β -blockers for further application in pharmaceutical field since each isomer give the different effect to the body metabolism. Figure 2.18 showed the studied β -blockers.

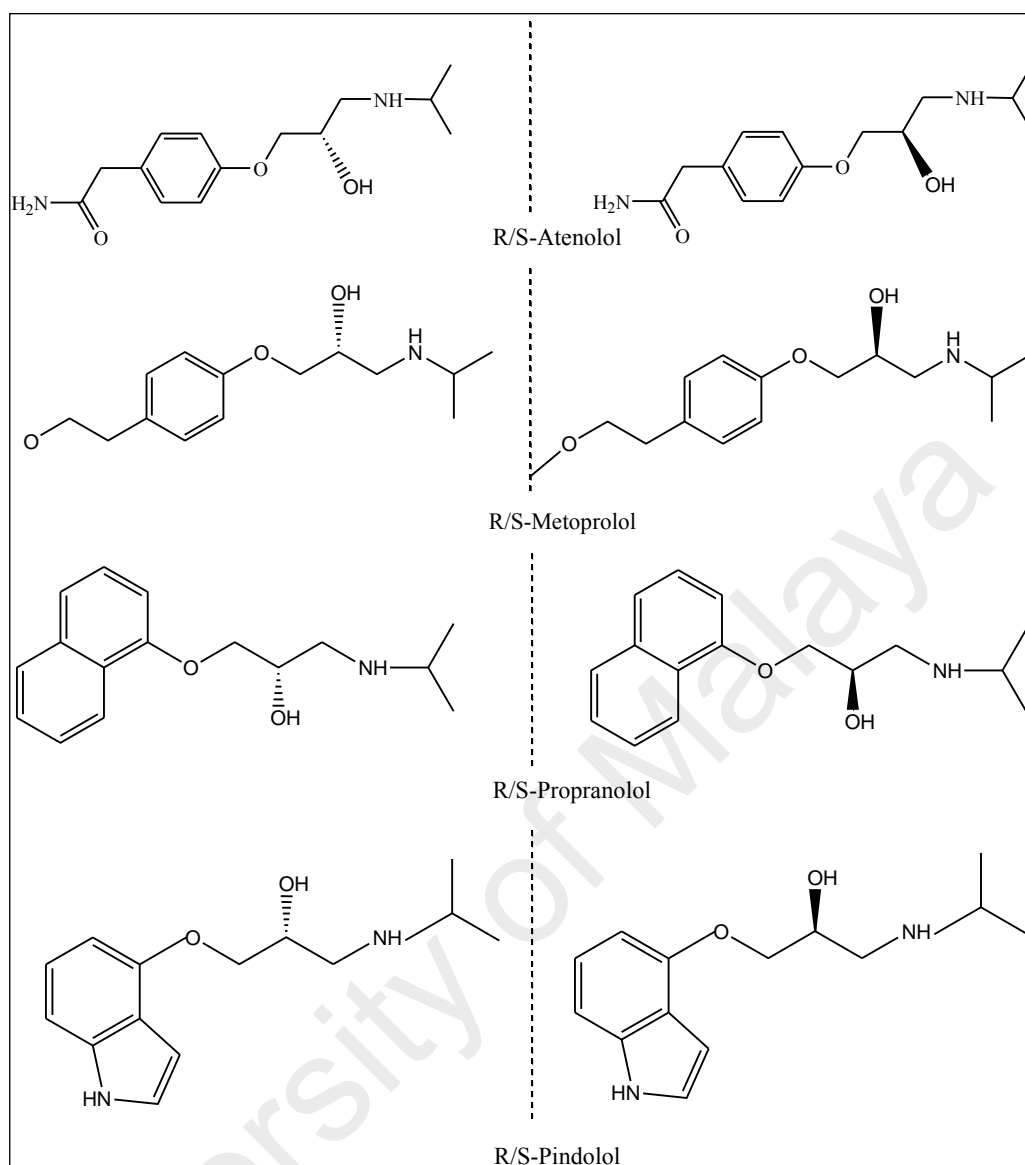


Figure 2.18: Structure of studied β -blockers

2.5.3 Non-steroidal anti-inflammatory drugs (NSAIDs)

Profen (2-arylpropionic acids) is an important group of non-steroidal anti-inflammatory drugs (NSAIDs), characterized by a chiral carbon atom next to the carboxylic acid group (Figure 2.19). The common anti-inflammatory mechanism of NSAIDs are inhibiting cyclooxygenase or 5-lipoxygenase and reducing the biosynthesis of prostaglandin (PG) to achieve the anti-inflammatory effect. It is well known that the pharmacological activities of the S-enantiomers of many NSAIDs are higher than that

of their *R*-enantiomers (Sekhon, 2013). Some reports have shown that the protein binding to NSAIDs have stereoselectivity (Zsila, 2013).

For ibuprofen, it is mainly the *R*-enantiomer that binds with human serum albumin (HSA) and the two enantiomers can be mutually replaced. In *in vivo* study, the *R*-enantiomer of ibuprofen undergoes unidirectional chiral inversion to *S*-enantiomer. This occurs to the extent about 65%, whereas there is no bio-inversion of *S*- to *R*-ibuprofen (Zhang *et al.*, 2014). Although this would favor the used of racemic ibuprofen, since most of its inactive enantiomer is converted to active form, conversion of racemic ibuprofen to *S*-ibuprofen results in variability of clinical response, including delayed onset of activity, and difficulty in achieving an optimal dose, also the formation of coenzyme A (CoA) thioester during bio-inversion of *R*- to *S*- ibuprofen may resulting toxic effects (e.g. interference of lipid anabolism/catabolism) (Podar *et al.*, 2016). In addition, *R*-ibuprofen bio-activation is susceptible to biological factors and certain drugs.

Most or all cyclooxygenase inhibitory activity of ketoprofen is attributed to the *S*-enantiomer (Podar *et al.*, 2016). The *R*-enantiomer is 30 to 5000 times less potent as an inhibitor of cyclooxygenase-1 and about 100 times less potent as an inhibitor of cyclooxygenase-2 (Negru *et al.*, 2015; Cooper *et al.*, 1998). In addition, *S*-ketoprofen has been found to be significantly less ulcerogenic in the rat gastrointestinal tract as compared to the racemic ketoprofen and that *R*-enantiomer may contribute to the pathogenesis of ulcers (Hardikar, 2008). In order of the different pharmacokinetic effect between each isomer of NSAIDs, they are raising the method to isolate and separate the individual isomers of the NSAIDs via chromatography.

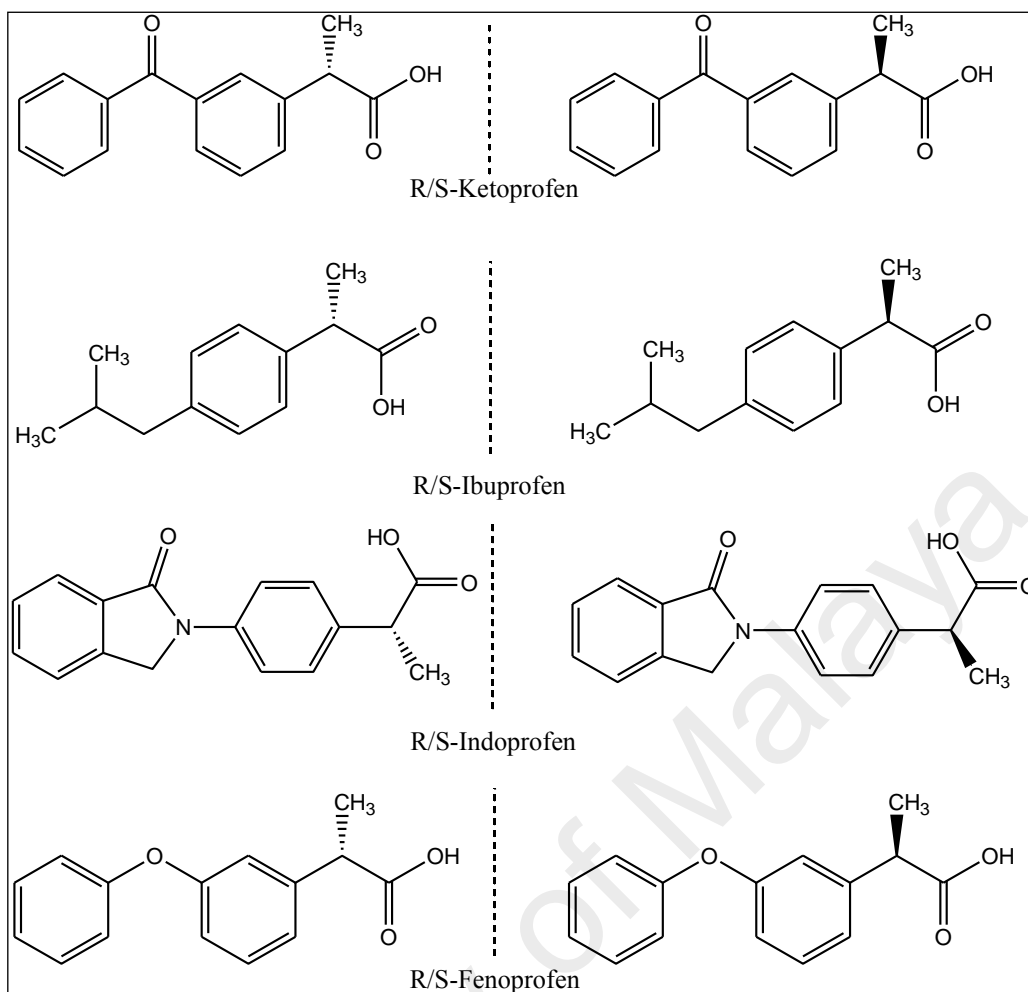


Figure 2.19: Structure of selected NSAIDs

CHAPTER 3: EXPERIMENTAL

3.1 Chemicals, materials and reagents

β -CD was purchased from Acros (Geel, Belgium) (99%). 1-Benzylimidazole (1-BzIm) (99%), 1-decyl-2-methylimidazole (C₁₀MIm) (97%) and toluene 2,4-diisocyanate (TDI) (95%) were supplied by Sigma Aldrich (Buches SG, Switzerland). Anhydrous N,N-Dimethylformamide (DMF), anhydrous hexane, HPLC grade of acetonitrile (ACN) and methanol (MeOH), *p*-toluene sulfonic acid, *p*-toluene sulfonyl chloride and Kromasil spherical silica gel (100Å pore size and 5µm particle size) were purchased from Merck (New York, NY, USA).

Flavonoids group consisting of hesperetin, naringenin and eriodictyol were purchased from Roth Karlsruhe (Germany) while flavanone was purchased from Sigma Aldrich (Buches SG, Switzerland). Propranolol, metoprolol, atenolol and pindolol were supplied from Sigma Aldrich (Buches SG, Switzerland). Ketoprofen, ibuprofen, indoprofen and fenoprofen were also purchased from Sigma Aldrich (Buches SG, Switzerland). The standard stock solutions of flavonoids, β -blockers and NSAIDs (500 mg/L) were prepared separately by dissolving them in MeOH and were stored in a dark amber glass at 4 °C.

3.2 Instruments

Fourier transform infrared (FT-IR) spectra were recorded using Perkin–Elmer RX1 FT-IR (Perkin Elmer, Waltham, MA, USA) in the ranged 4000 to 400 (cm⁻¹). ¹H NMR, ¹³C NMR, and NOESY spectra were recorded on AVN 600 MHz (Bruker, Fällanden, Switzerland), and Dimethyl Sulfoxide (DMSO-D₆) was used as solvent. Thermogravimetric analyzers were examined using TGA 4000 (Perkin Elmer, USA). A linear heating rate was set at 20 °C per min within the temperature ranged from 50 °C to 900 °C in a stream of nitrogen atmosphere. The chromatographic data was performed

using a HPLC system consisted of a LC-20AT pump, a SPD-M20 detector, a SIL-20AHT auto sampler, a CTO-20AC column oven and CBM-20A communication bus module (Shimadzu, Japan).

3.3 Preparation of β -CD based chiral stationary phase

The preparation of β -CD based CSP was carried out by synthesizing β -CD functionalized IL and then immobilized onto modified silica.

3.3.1 Synthesis of β -CD functionalized ionic liquid

β -CD functionalized IL was prepared according to the previous report (Raovv *et al.*, 2013), as shown in Figure 3.1. First, 6-O-monotosyl-6-deoxy- β -cyclodextrin (β -CDOTs) was prepared as describe by Zhong (Raovv *et al.*, 2013). Then, the reaction was carried out by reacting β -CDOTs with IL (1-BzIm/C₁₀MIm). Since tosylate is a good leaving group, imidazole can easily undergo the nucleophilic substitution.

The reaction was performed as follows: A suspension of β -CD (11.5 g, 10 mmol) and *p*-toluenesulfonic anhydride (Ts₂O) (4.9 g, 15 mmol) in 250 mL of water was stirred at room temperature for 2 h. Then, solution of NaOH (5.0 g in 50 mL of H₂O) was added, and after 10 min, the reaction mixture was filtered through the celite on the sintered glass funnel to separate the excess tosylate. The filtrate was brought to pH 8 by the addition of ammonium chloride (13.4 g). The precipitate of β -CDOTs was obtained and cooled at 4 °C overnight. Then, the dried β -CDOTs (1.00 g, 0.78 mmol) and 1-BzIm (10 mole equivalent) were dissolved in anhydrous DMF (40 ml) and the solution was stirred at 90 °C under N₂ atmosphere. After two days, the resultant solution was cooled to room temperature and acetone slowly was added. The mixture was stirred for 30 minutes, and thereafter, filtered and washed the obtained β -CD-

BIMOTs (mono-6-deoxy-6-(3-benzylimidazolium tosylate)- β -CD) in excess amount of acetone.

The same procedure was applied for synthesizing β -CD-DIMOTs (mono-6-deoxy-6-(3-decyl-2-methylimidazolium tosylate)- β -CD) using $C_{10}MIm$ replacing 1-BzIm. The characterized results showed that β -CD-BIMOTs and β -CD-DIMOTs had been successfully prepared. From 1H NMR result, the chemical shifts of imidazole ring (Hf, He, and Hd) appeared in the downfield region since the protons were deshielded upon functionalization. A new peak was observed in proton (H6*, 3.9 ppm) and carbon signal (C6*, 45 ppm), which belonged to the substituted CD. All the protons of β -CD still appeared after the reaction because the functionalization process occurred at only one of the primary hydroxyl groups of β -CD. The obtained product was successfully characterized using several analytical techniques. Both structures of β -CD-BIMOTs and β -CD-DIMOTs are illustrated in Figure 3.2 and Figure 3.3.

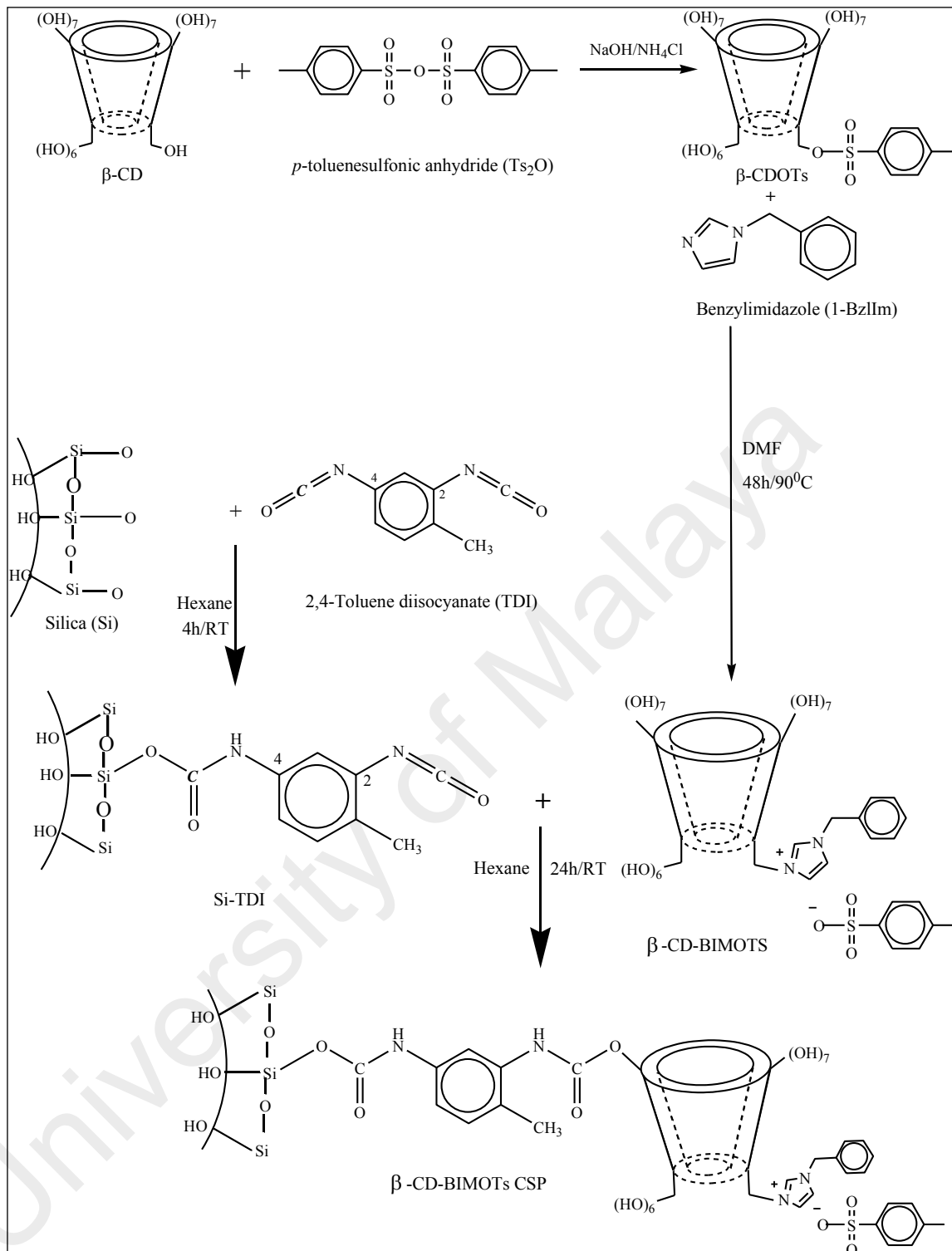


Figure 3.1: Synthesis pathways of $\beta\text{-CD-BIMOTs CSP}$

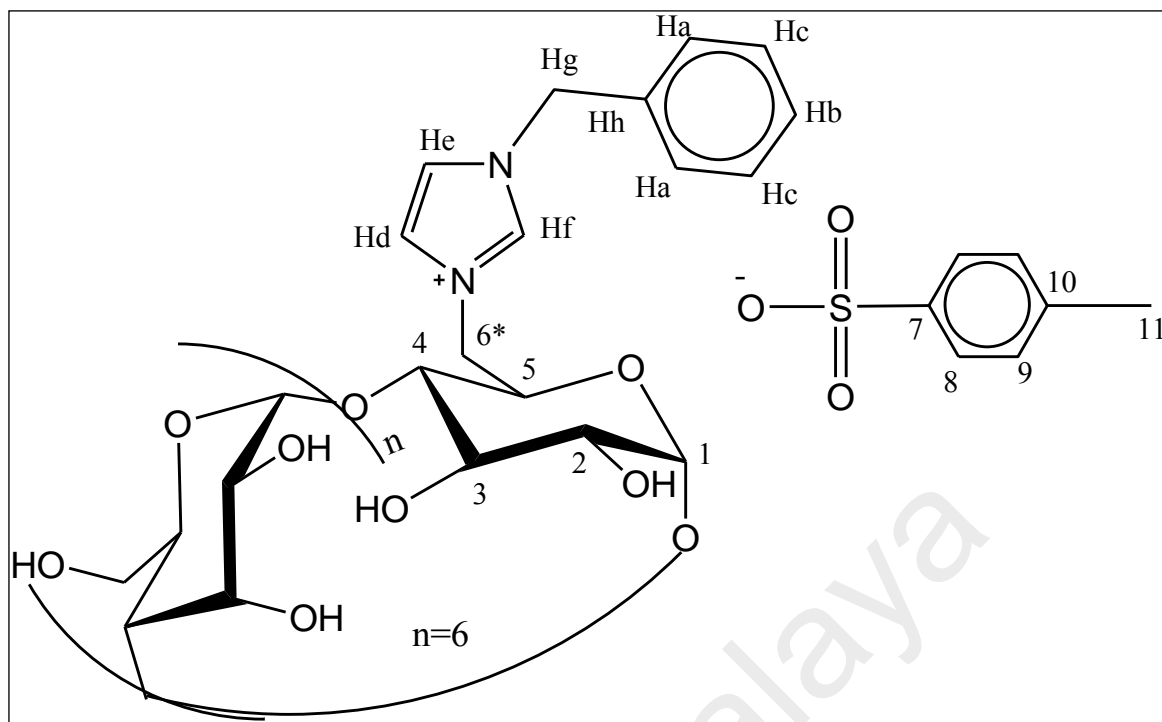


Figure 3.2: Structure of β -CD-BIMOTs

FT-IR/KBr, cm^{-1} : 3297 (OH), 2922 (C-H), 1652 (C=C), 1152 (C-N).

^1H NMR, DMSO- D_6 : Hf (9.28, s), He (7.94, s), Hd (8.20, s), Hc (7.75, s), Hb (7.80, t), Ha (7.46, s), Hg (5.18, s), H8 (7.41, d), H9 (7.10, d), OH-2-OH-3 (5.50–5.80, m), H1 (4.83, s), OH-6 (4.47–4.6, m), H6* (3.91), H3, H5, H6 (3.40–3.63), H2-H4 (3.20–3.40, m), H11 (2.08, s).

^{13}C NMR, DMSO- D_6 : Ca (127), Cb (123.4), Cc (128.3), Cd (128), Ce (119), Cf (136.9), Cg (52), Ch (137.8), C7 (145.26), C10 (137.3), C9 (128.7), C8 (125.6), C1 (101.8), C4 (81.16), C2 (73.27), C3 (71.6), C5 (69.37), C6 (60.03), C6* (45.2), C11 (21.97).

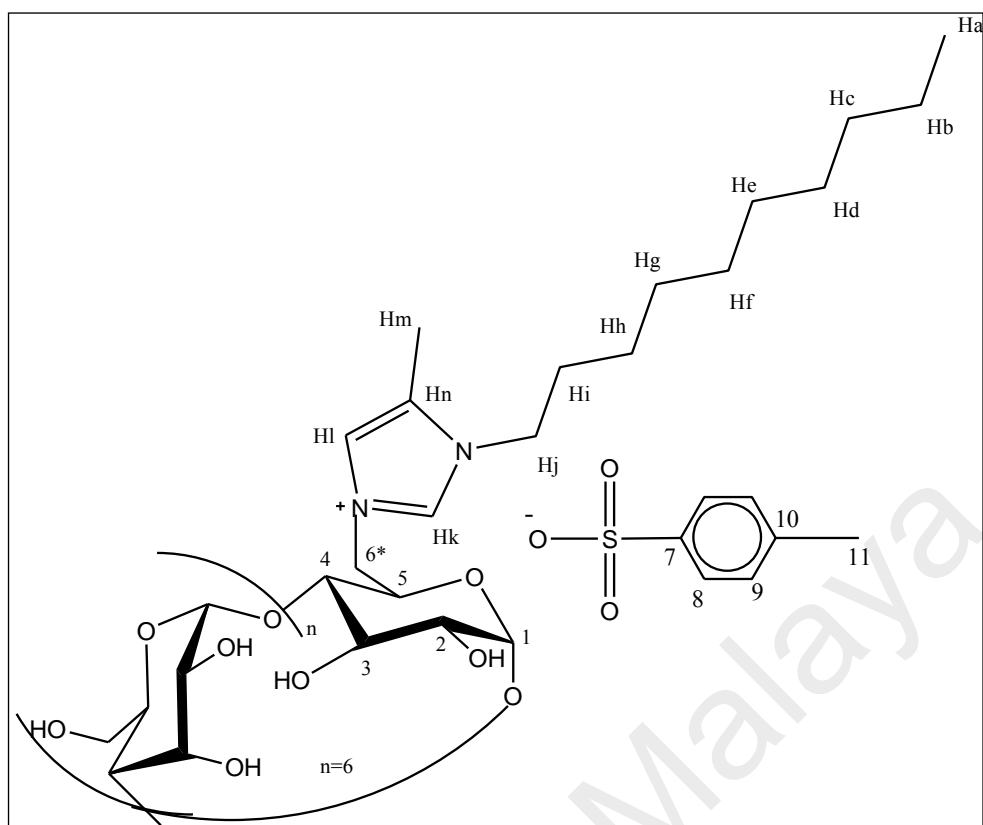


Figure 3.3: Structure of β -CD-DIMOTs

FT-IR/KBr, cm^{-1} : 3297 (OH), 2922 (C-H), 1652 (C=C), 1152 (C-N).

^1H NMR, DMSO- D_6 : Hl (7.68, s), Hk (7.61, s), Hb-Hj (1.23-1.28, t), Ha (0.85, t), H8 (7.46, d), H9 (7.11, d), OH-2-OH-3 (5.64–5.79, m), H1 (4.83, s), OH-6 (4.44–4.54, m), H6* (3.91), H3, H5, H6 (3.54–3.63), H2-H4 (3.20–3.34, m), H11 (2.28, s).

^{13}C NMR, DMSO- D_6 : Ca (16.13), Cb (19.79), Cc (28.62), Cd (22.48), Cg (22.48), Ch (21.38), Ci(22.48), Cj (31.37), Ck (126.42), Cl (128.75), Cm (14.40), Cn (129.84), C9 (128.17), C8 (126.06), C1 (102.38), C4 (81.95), C2 (73.49), C3 (72.43), C5 (70.74), C6 (60.36), C6* (45.66).

3.3.2 Immobilization of β -CD-BIMOTs and β -CD-DIMOTs onto modified silica to obtain the CSP

Silica is the most suitable inert support for stationary phase, because of its high physical strength, chemical inertness and high thermal resistance (Arakaki *et al.*, 2000; Alimarin *et al.*, 1987; Cassim & Yang, 1969). The immobilization was performed by reacting the β -CD functionalized IL with modified silica gel that bearing carbamate group as linker (Zhang *et al.*, 1999).

First, the modified silica gel was prepared as reported (Yatabe & Kageyama, 1994). The modified silica gel was prepared by reacting TDI with silica gel in dry hexane for 4 h at room temperature to obtain Si-TDI. Upon completion of the reaction, the product was filtered, rinsed thoroughly by hexane and dried under reduced pressure. Later, the Si-TDI (5g) was stirred in anhydrous hexane (200 mL) through continuous stream of nitrogen at room temperature. After 30 min, a solution of β -CD functionalized IL (β -CD-BIMOTs or β -CD-DIMOTs) (1.8 g) was added. Stirring was continued for 24 h. The obtained solid was filtered and wash with toluene, acetone and distilled water to afford purified product. The obtained product was characterized using FT-IR and TGA.

3.3.3 Synthesis of native β -CD (n- β -CD) as chiral stationary phase

Native β -CD as CSP was prepared by immobilizing the native β -CD onto Si-TDI. The procedure was similar as the immobilization of the β -CD-BIMOTs and β -CD-DIMOTs onto Si-TDI.

3.4 Column packing approach

The synthesized CSPs were packed with hexane into empty stainless steel column (250 mm \times 4.6 mm I.D.). First, the CSPs (2.5 g) was suspended in approximately 15 ml of HPLC grade hexane and then poured into the column. The

CSPs were packed into the stainless steel column with a 1525 binary HPLC pump. The flow rate and pressure was first settled at 24.00 ml/min and 4000 Psi respectively. After that, the pressure was increased stepwise until the back pressure reached 8000 Psi. The pressure and flow rate was keep constantly for 1 h.

3.5 HPLC analysis instrumentation and conditions

The newly packed column was flushed with 100 % hexane at a flow rate of 0.2 ml/min for 24 hours. The flow rate was increased to 0.5 ml/min for getting the stable baseline. All analyses were performed at ambient temperature at 25 °C. The analytes solutions at concentration of 500 mg/L were prepared by dissolving flavonoids, β -blockers and NSAIDs separately in MeOH. The injection volume was 20 μ l. The flow rate was fixed at 0.5 ml/min for all analytes. The buffer of triethylamine acetate (TEAA) was prepared by adding triethylamine (TEA) with acetic acid (HOAc) to adjust the pH of mobile phase. The amount of additives in the buffer was recorded as the total weight of both acetic acid and TEA in buffer (w/v).

3.6 Calculations of chromatographic data

Figure 3.4 illustrated the example of chromatogram of two well resolved enantiomers and its chromatographic data. Three important terms used in this regard are k' (capacity factor or retention factor), α (selectivity factor or separation factor) and R_s (resolution factor). k' is a measurement of time of a solute is retained on the column. Retention is a function of affinity of the solute to the stationary phase. The stronger the attraction between the solute and the column material, the longer is the retention. α is a measurement of selectivity of the column for any pair of solutes. R_s is a measurement of how well the enantiomers have been separated. The baseline resolution is achieved when $R_s \geq 1.5$. The k' , α and R_s were calculated using the following equations:

$$k' = \frac{(t_R - t_0)}{t_0} \quad 3-2$$

$$\alpha = \frac{k_2'}{k_1'} = \frac{(t_{R2} - t_0)}{(t_{R1} - t_0)} \quad 3-3$$

$$R_s = \frac{2 \times (t_{R2} - t_{R1})}{(W_1 + W_2)} \quad 3-4$$

The dead time (t_0) is the time for the mobile phase to pass through the column, which relates to the efficiency of the column. The retention time (t_R) is the retention time corresponding to each isomer in the chromatographic separation. t_{R2} and t_{R1} represents the retention times of the second and first isomers respectively, and W_1 and W_2 are the corresponding base peak width.

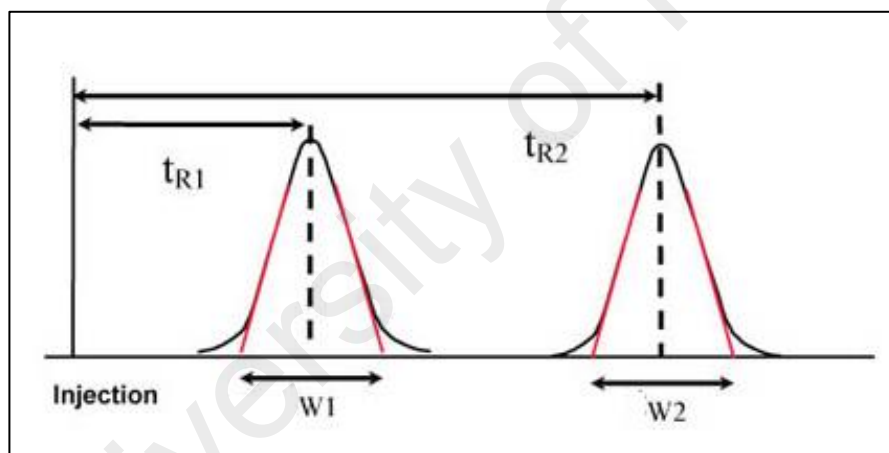


Figure 3.4: Two enantiomerically related peaks and the measurements required to calculate k_1' , k_2' , α and R_s

3.7 Preparation of inclusion complex

3.7.1 Preparation of kneaded complex

The inclusion complex of β -CD-BIMOTs with analytes was prepared using conventional kneading method (Cwiertnia *et al.*, 1999). Equimolar amount of β -CD-BIMOTs and analytes were kneaded with mortar and pestle in minimal ethanol to form homogenous paste (Figure 3.5). The complex was kneaded for 30 min and dried to constant mass. After drying, a white powder was obtained. The final product was characterized in the liquid state by one dimensional (1D) ^1H NMR and two dimensional (2D) ^1H NMR NOESY. For ^1H NMR and NOESY, the spectra were obtained from the samples that prepared using β -CD-BIMOTs and analytes with the ratio of 1:1. The samples were dissolved in DMSO-d_6 . Seven hundred microliter of solutions were introduced into standard 5 mm NMR tubes and the spectra were recorded at 300.15 K. For NOESY experiments, the spectra were recorded with a mixing time of 700 ms with 256 increments and 40 scans.

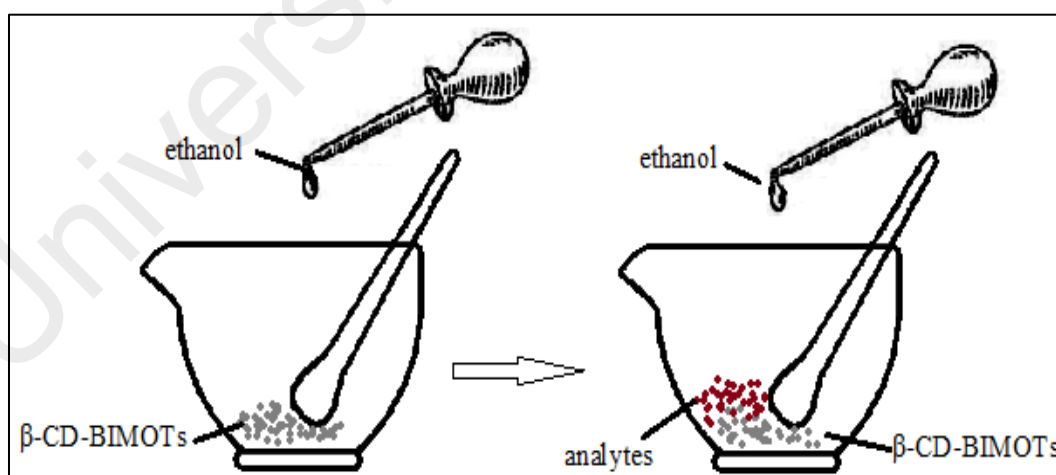


Figure 3.5: Schematic of kneading method

3.7.2 Determination of formation constant

UV-Visible spectrophotometer with 1 cm quartz cuvette was used for this experiment. The absorption spectrum of β -CD-BIMOTs and analytes complex was recorded against blank reagent. Blank reagent was prepared with the same concentration without the addition of analytes. In addition, absorption spectra of each analyte and β -CD functionalized ionic liquid were also recorded. For the formation constant curve, the concentration of analytes was held constant at 0.01 mM, meanwhile the concentration of β -CD functionalized ionic liquid was varied (0.001, 0.002, 0.003 and 0.005 M). The formation constant and stoichiometry of the β -CD functionalized ionic liquid inclusion complex was obtained from the Benesi-Hildebrand equation (Equation 3-5) (Qian *et al.*, 2008).

$$\frac{1}{(A-A_0)} = \left[\frac{1}{(A'-A_0)} \right] + \left[\frac{1}{K(A'-A_0)[\beta\text{-CD-BIMOTs}]} \right] \quad 3-5$$

In the above equations, A_0 is the intensity of absorption of the guest without β -CD functionalized ionic liquid, A is the absorbance with a particular concentration of β -CD functionalized ionic liquid, A' is the absorbance at the maximum concentration of β -CD functionalized ionic liquid used and K is the formation constant. Linearity is obtained in the plot of $1/(A - A_0)$ versus $1/K(A' - A_0)[\beta - \text{CD} - \text{BIMOTS}]$ for 1:1 complexes (Equation 3-5). The formation constant (K) was calculated from the slope of Benesi-Hildebrand plot using the Equation 3-6.

$$K = \left[\frac{1}{\text{slope } (A'-A_0)} \right] \quad 3-6$$

CHAPTER 4: RESULTS AND DISCUSSION

4.1 Characterization of β -CD Based Chiral Stationary Phase

4.1.1 FT-IR analysis

The spectra of β -CD, β -CD-BIMOTs and β -CD-DIMOTs are shown in Figure 4.1. Meanwhile, the main frequencies of β -CD, β -CD-BIMOTs and β -CD-DIMOTs are shown in Table 4.1. The broad O-H stretching band around 3200-3300 cm^{-1} (Figure 4.1) for β -CD, β -CD-BIMOTs and β -CD-DIMOTs are corresponded to the multiple -OH functional groups in β -CD molecules. O-H stretching, C-H stretching, and C-N bending (refer Table 4.1 for assignment) were observed as the most obvious band in the IR spectra of both β -CD-BIMOTs and β -CD-DIMOTs. The intense band at 1657 cm^{-1} referred to C=C aromatic ring of 1-BzIm moieties was observed at β -CD-BIMOTs spectra (Figure 4.1 (b)). The weak bands known as overtones at 1665-2000 cm^{-1} were correlated to aromatic ring of benzene was also observed at β -CD-BIMOTs spectra. Moreover, the band of C-H of β -CD-BIMOTs and β -CD-DIMOTs spectra (Figure 4.1(b and c)) that occurred at 2925 cm^{-1} are more intense than the band of C-H of β -CD spectra (Figure 4.1(a)). These prove that β -CD was successful functionalized with 1-BzIm or C₁₀MIm and β -CD-BIMOTs and β -CD-DIMOTs were obtained.

The spectra and assignment peak of Si-TDI (modified silica), native β -CD CSP, β -CD-BIMOTs CSP and β -CD-DIMOTs CSP are shown in Figure 4.2 and Table 4.2, respectively. Spectra of Si-TDI (a) shows the presence of the isocyanate (O=C=N-) group at 2280 cm^{-1} . TDI has two isocyanate groups with different activities towards OH groups that located at the para-position and ortho-position, respectively. The two isocyanate groups in TDI react at different rates with the para-position (approximately four times more reactive than the ortho-position) (Arnold *et al.*, 1957; Simons & Arnold, 1956). Hence, the isocyanate functional groups in TDI (para position) reacted

with OH groups on the surface of silica and formed Si-TDI. The remaining isocyanate group at ortho-position would react with secondary OH group of β -CD or β -CD functionalized ionic liquid. Therefore, the isocyanate peak was disappeared after immobilization of native β -CD, β -CD-BIMOTs and β -CD-DIMOTs onto Si-TDI to obtain CSP as shown in Figure 4.2 (b), (c) and (d).

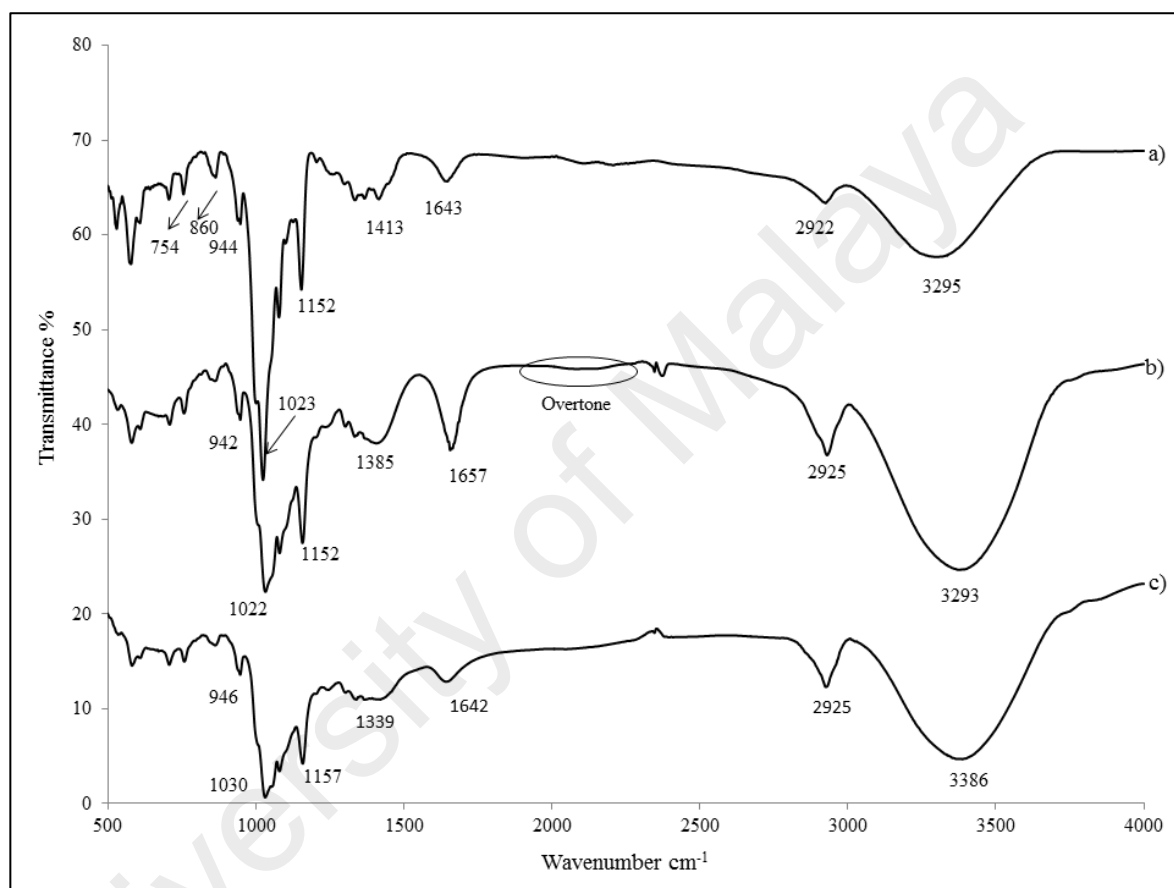


Figure 4.1: FT-IR spectrum of a) β -CD b) β -CD-BIMOTs c) β -CD-DIMOTs

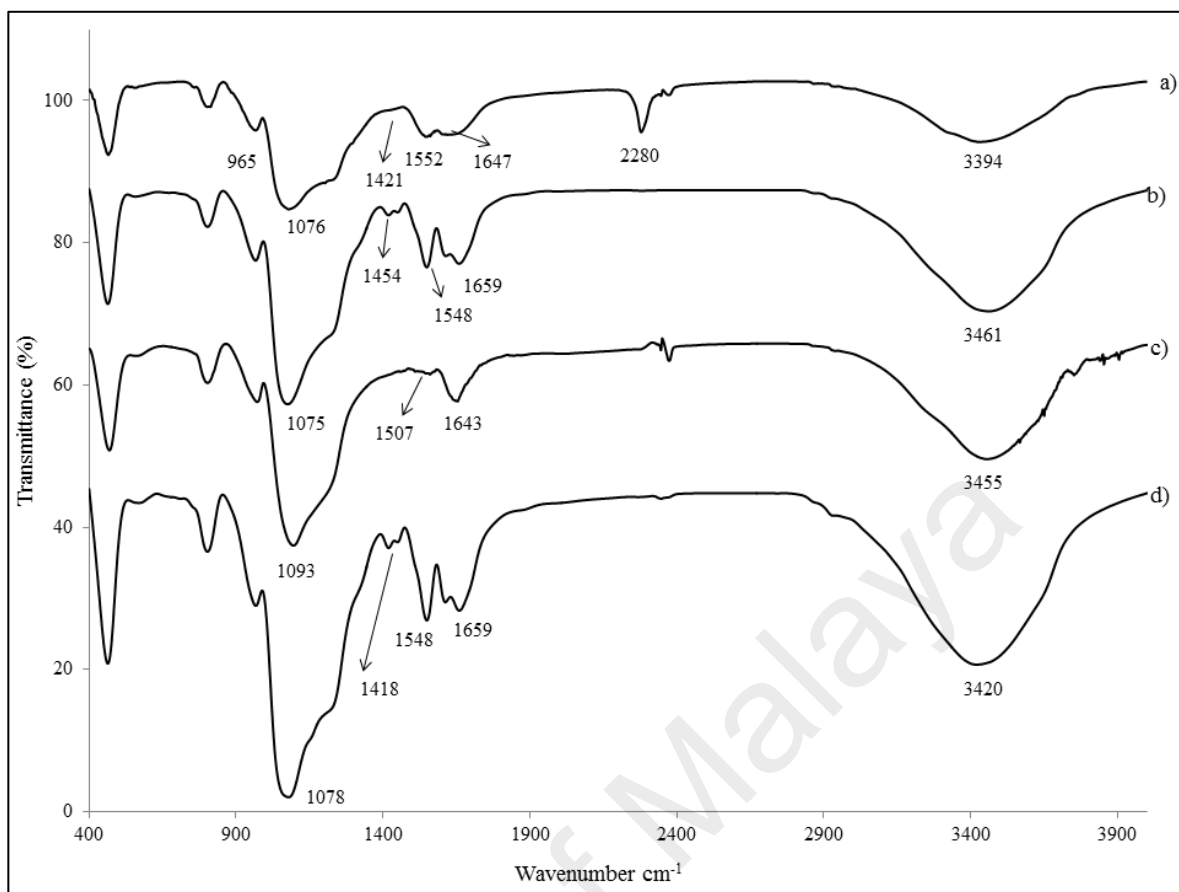


Figure 4.2: FT-IR spectrums of a) Si-TDI b) native β -CD CSP c) β -CD-BIMOTs CSP
d) β -CD-DIMOTs CSP

Table 4.1: Main IR frequencies for β -CD, β -CD-BIMOTs and β -CD-DIMOTs with assignments

Wavelength cm^{-1}	Assignments	β -CD	β -CD-BIMOTs	β -CD-DIMOTs
3295	O-H stretch	√		
3293	N-H, O-H stretch		√	
3386	N-H, O-H stretch			√
2922	C-H stretch	√		
2925, 1385	C-H stretch, bend		√	
2925, 1339	C-H stretch, bend			√
1643, 1023	C-O stretch	√	√	
1657	C=C aromatic (1-BzIIIm)		√	
1642, 1030	C-O stretch			√
1413	O-H, CH ₂	√		
1152	C-C-C	√		
1157	C-N		√	
1157	C-N			√
944, 860, 754	-CH, =CH ₂ , CH	√	√	√

Table 4.2: Main IR frequencies for Si-TDI, native β -CD CSP, β -CD-BIMOTs CSP and β -CD-DIMOTs CSP with assignments

Samples	Wavelength cm^{-1}	Assignments
Si-TDI	3297	O-H stretch
	2280	N=C=O stretch
	1647, 1552	NHCO carbamate linkage
	1421	Aromatic group in TDI
Native β -CD CSP	3461	O-H stretch
	2280	Absence of N=C=O
	1659, 1548	NHCO carbamate linkage
	1548, 1454	Aromatic group in TDI
β -CD-BIMOTs CSP	3455	N-H, O-H stretch and imidazole ring
	2280	Absence of N=C=O
	1653	C=C aromatic (1-BzIIIm)
	1507	C-C stretch in aromatic
	1093	C-O stretch
β -CD-DIMOTs CSP	3420	N-H, O-H stretch and imidazole ring
	2280	Absence of N=C=O
	1659, 1078	C-O stretch
	1548, 1418	Aromatic group in TDI

4.1.2 Thermalgravimetric analysis

TGA was performed on the Si-TDI, native β -CD CSP, β -CD-BIMOTs CSP and β -CD-DIMOTs CSP in the temperature range of 50 to 900 °C. Based on the thermograms shown in Figure 4.3, it can be seen that there is an initial loss of weight at temperature below 100 °C for all samples. This was attributed to the removal of physically adsorbed water and/or remaining solvent residues. Physically adsorbed water was removed completely by further heating to around 200 °C. TDI attached to the silica surface decomposed in the region between 125 and 250 °C (Guo *et al.*, 2005). Moreover, Si-TDI revealed a smaller, but noticeable, weight loss in the region from 250-600 °C. This can be attributed to the dehydration of the silica surface, in which silanol groups condense to siloxanes, a process known to occur in this thermal region (Poole, 2003). The thermogram of β -CD-BIMOTs CSP and β -CD-DIMOTs CSP showed two very distinct weight loss that occurred at the range of 210-357 °C and 400-600 °C. The first of these two weight loss was attributed to the decomposition of organic moieties at the surface. The second weight loss was associated with the decomposition of the residual methoxy groups on silica (Antochshuk & Jaroniec, 2000). In addition, the thermogram of native β -CD CSP, β -CD-BIMOTs CSP and β -CD-DIMOTs CSP attributed to the weight loss at 600-900 °C due to decomposition of the β -CD. By comparing Figure (c) and (d), it is clear that β -CD-BIMOTs-CSP shows more pronounced weight loss than β -CD-DIMOTs-CSP at all isothermal temperatures. This may be due to the long alkyl chain of β -CD-DIMOTs-CSP prevent it to be very volatile at high temperatures (Lu *et al.*, 2002). The temperature of weight loss with detail assignment is shown in Table 4.3.

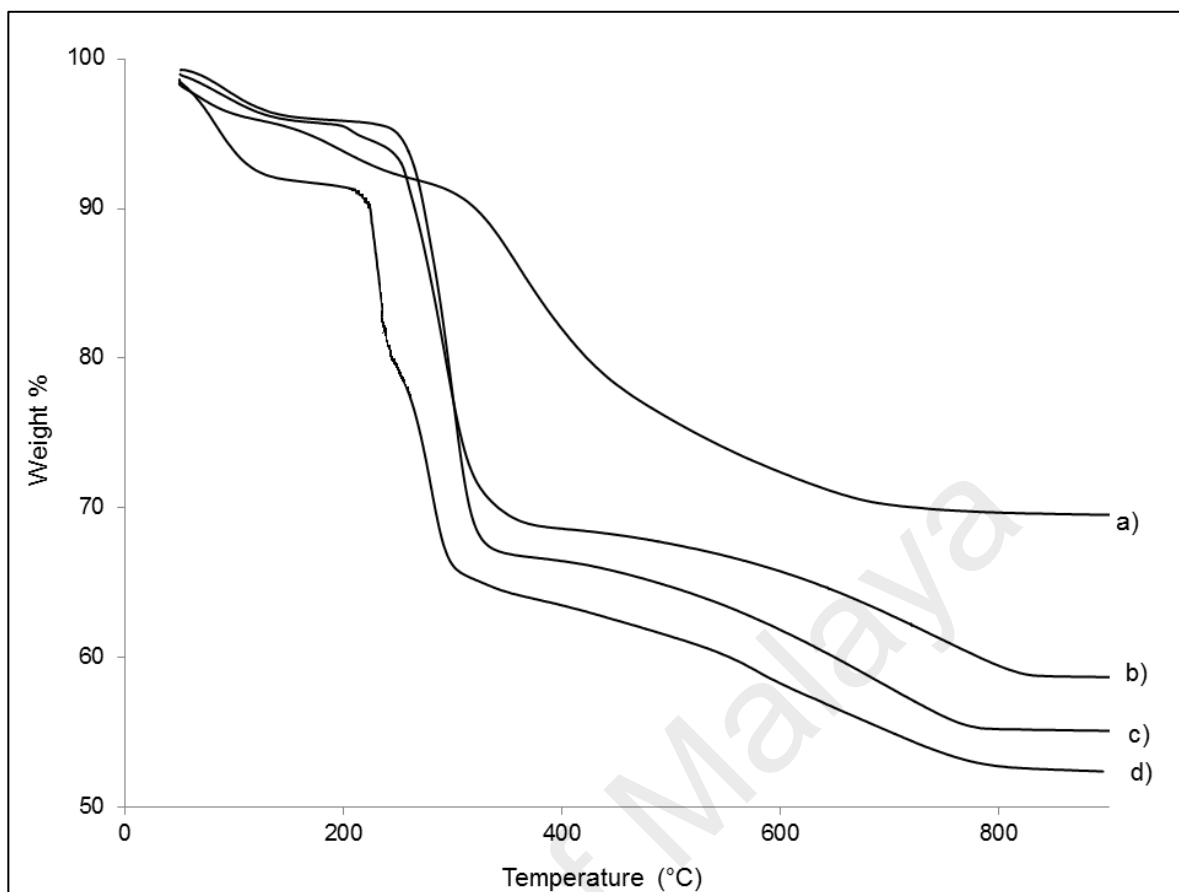


Figure 4.3: Thermogram of a) Si-TDI b) native β -CD CSP c) β -CD-BIMOTs CSP d) β -CD-DIMOTs CSP

Table 4.3: The assignment for temperature of weight loss

Samples	Region (°C)	Weight loss (%)	Assignment
Si-TDI	50-100	4	Water loss
	125-250	2	TDI
	250-600	28	Silanol condensation
Native β -CD CSP	50-100	3	Water loss
	125-250	3	TDI
	250-600	24	Silanol condensation
	600-900	10	β -CD
β -CD-BIMOTs CSP	50-100	3	Water loss
	125-250	3	TDI
	215-357	26	1-BzIm, OTs
	357-900	11	Silanol condensation, β -CD
β -CD-DIMOTs CSP	50-100	7	Water loss
	125-250	2	TDI
	211-357	15	C ₁₀ Mim, OTs
	357-900	12	Silanol condensation, β -CD

4.2 Screening performance of CSPs

Different moieties that functionalized on β -CD possess different effects to the separation of chiral compounds. Herein, the effect of different group at the side chain of imidazolium cation of IL was studied. The performance of β -CD-BIMOTs CSP and β -CD-DIMOTs CSP were compared with native β -CD based CSP for the enantioseparation of flavonoids, β -blockers and NSAIDs. As shown in Table 4.4, the chromatograms showed that most of the flavonoids, β -blockers and NSAIDs were enantioseparated using β -CD-BIMOTs CSP as compared to β -CD-DIMOTs CSP and native β -CD based CSP. This result might due to the β -CD-BIMOTs CSP that displayed additional interaction with analytes which enhanced the enantioseparations. β -CD-BIMOTs CSP is prefer to be approached by planar analytes due to the planar aromatic

of 1-BzIIm (Wang *et al.*, 2012c). This might attributed to the π - π interaction between analytes and β -CD-BIMOTs CSP that enhanced the enantioseparation. In addition, the long alkyl chain is preferably covered the partial cavity (Meier-Augenstein *et al.*, 1992) resulting decreased the chiral selectivity of β -CD-DIMOTs CSP. Thus, the optimization of mobile phase for the enantioseparation of flavonoids, β -blockers and NSAIDs on β -CD-BIMOTs CSP was studied. Furthermore, the mechanism of the enantioseparation was also evaluated.

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Table 4.4: The chromatogram for the enantioseparation of selected flavonoids, β -blockers and NSAIDs on β -CD, β -CD-BIMOTs and β -CD-DIMOTs CSPs

Analytes	CSPs		
	β -CD	β -CD-BIMOTs	β -CD-DIMOTs
Flavonoids			
β -blockers			

Table 4.4, continued

Analytes	CSPs		
	β -CD	β -CD-BIMOTs	β -CD-DIMOTs
NSAIDs			

Flavonoids: a) flavanone b) hesperetin c) naringenin d) eriodictyol

β -blockers: a) propranolol b) metoprolol c) pindolol d) atenolol

NSAIDs : a) fenoprofen b) ibuprofen c) indoprofen d) ketoprofen

Condition: i) 90/10 ACN/water ii) 50/50 ACN/water iii) 30/70 ACN/water

4.3 Enantioseparation performance of Flavonoids

The type and composition of organic modifier as mobile phase are important factors that affect the enantioseparations. Adjusting the pH of mobile phase for reverse phase mode would also influence the forms of analytes and thus affect the enantioseparation. As presented in Table 4.5, high R_s values indicated the good enantioseparation for flavanone ($R_s=1.63$) and hesperetin ($R_s=1.06$) with the mobile phase of MeOH/water:50/50 and ACN/water:50/50, respectively. In addition, flavanone also obtained good enantioseparation ($R_s=1.86$) in ACN/buffer at pH 4. However, a low R_s value was obtained for flavanone when ACN/buffer pH 9 was selected as mobile phase. Meanwhile, the enantiomers of naringenin and eriodictyol were not resolve at all using all selected mobile phases. Moreover, it can be seen that the k_1' values of flavonoids decreased with increasing content of organic solvent. This was a common rule in reverse phase mode due to the increasing content of organic solvent that led to the increased of elution strength of mobile phase. Thus, flavonoids easily can be displaced from the stationary phase.

Flavanone obtained good enantioseparation in most of the mobile phase conditions which might due to its hydrophobic properties that facilitated the inclusion complex formation with hydrophobic cavity of β -CD-BIMOTs CSP. Moreover, flavanone with aromatic rings without any substituent may experience less steric hindrance for inclusion complex formation with cavity of β -CD-BIMOTs CSP. In addition, the carbonyl group and aromatic ring of flavanone can form hydrogen bonding and π - π interaction, respectively, with β -CD-BIMOTs CSP which can further enhance the enantio-recognition. Flavanone is classified as neutral compound as compared with hesperetin, naringenin and eriodictyol which are weakly acidic in nature (Ng *et al.*, 2002). Thus, at pH 4 and 7, flavanone is remained neutral and preferable to form

inclusion complex with cavity of β -CD (Raov *et al.*, 2013). Meanwhile, flavanone is known to undergo ring opening under basic condition to the corresponding unstable 2'-hydroxyl substituted chalcones (Figure 4.4) (Wistuba *et al.*, 2006) which might be a reason in the decreasing R_s value at pH 9.

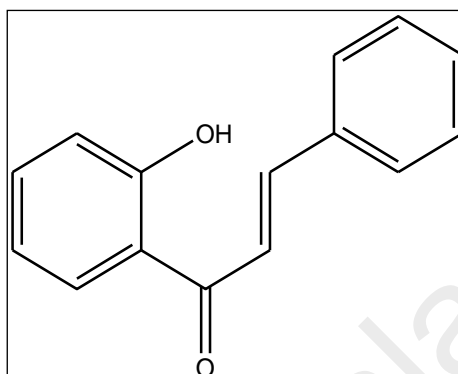


Figure 4.4: Structure of 2'-hydroxyl substituted chalcones

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Table 4.5: Chiral separation data for the flavonoids on β -CD-BIMOTs CSP in the reverse mobile phase

Flavonoids	Conditions	pH 4			pH 7			pH 9		
		k_1'	k_2'	R_s	k_1'	k_2'	R_s	k_1'	k_2'	R_s
Flavanone	a	0.34	0.48	0.64	0.33	0.49	0.45	0.38	0.85	0.79
	b	2.09	5.24	1.86	0.47	0.71	0.81	0.33	0.46	0.46
	c	2.77	2.77	0	2.61	2.61	0	2.51	2.51	0
	d	7.23	7.23	0	1.44	2.05	0.76	1.92	3.34	0.93
	e	2.27	3.58	0.85	2.58	4.31	1.63	6.84	6.84	0
Hesperetin	a	1.18	1.18	0	0.47	0.76	0.45	0.79	0.79	0
	b	1.49	1.49	0	0.37	1.36	1.06	1.61	1.61	0
	c	9.75	9.75	0	4.43	7.14	0.92	4.31	4.31	0
	d	1.35	1.35	0	1.29	1.29	0	1.80	1.80	0
	e	-	-	-	16.19	16.19	0	4.18	4.18	0
Naringenin	a	0.27	0.27	0	0.28	0.28	0	0.28	0.28	0
	b	0.62	0.62	0	0.84	0.84	0	0.97	0.97	0
	c	1.54	1.54	0	4.16	4.16	0	5.29	5.29	0
	d	0.68	0.68	0	0.12	0.12	0	0.83	0.83	0
	e	-	-	-	0.18	0.18	0	3.61	3.61	0
Eriodictyol	a	0.22	0.22	0	0.32	0.32	0	0.34	0.34	0
	b	0.34	0.34	0	0.34	0.34	0	0.34	0.34	0
	c	0.35	0.61	0.26	0.36	0.36	0	0.37	0.37	0
	d	-	-	-	0.19	0.19	0	0.82	0.82	0
	e	-	-	-	0.34	0.34	0	4.09	4.09	0

Conditions pH 7: a) ACN/water-90/10 b) ACN/water-50/50 c) ACN/water-30/70 d) MeOH/water-90/10 e) MeOH/water-50/50

Conditions pH 4 or 9: a) ACN/buffer-90/10 b) ACN/buffer-50/50 c) ACN/buffer-30/70 d) MeOH/buffer-90/10 e) MeOH/buffer-50/50

According Li *et al.* (1992), the formation of inclusion complex is an important interaction to achieve better enantioseparation (Li & Purdy, 1992). In order to study the interaction for the enantioseparation, ^1H NMR and NOESY of β -CD-BIMOTs/flavonoids complexes were studied. The deduced structures of the β -CD-BIMOTs and β -CD-BIMOTs/flavonoids complexes are shown in Figure 4.5 and Figure 4.6, respectively. Chemical shift (δ) variations can provide evidence for the formation of inclusion complexes in solution. The values of the δ for different protons in β -CD-BIMOTs and β -CD-BIMOTs/flavonoids complexes are listed in Table 4.6. The induced shift ($\Delta\delta$) is defined as the difference in chemical shift in the presence or absence of analytes. In this study, the induced shift was calculated using Eq. 4-1:

$$\Delta\delta = \delta(\text{complex}) - \delta(\text{free}) \quad 4-1$$

Normally, the inclusion of an apolar region of an analyte into the hydrophobic cavity would affect the inner protons of the glucose units of β -CD, namely, H3 and H5 (Zhang *et al.*, 1990), whereas the protons on the exterior torus of β -CD (H1, H2 and H4) would also be affected if there are any hydrogen bonding involved. As the result, the chemical shifts of β -CD-BIMOTs protons (H1, H2, H3, H4 and H5) would change as the presence of analytes.

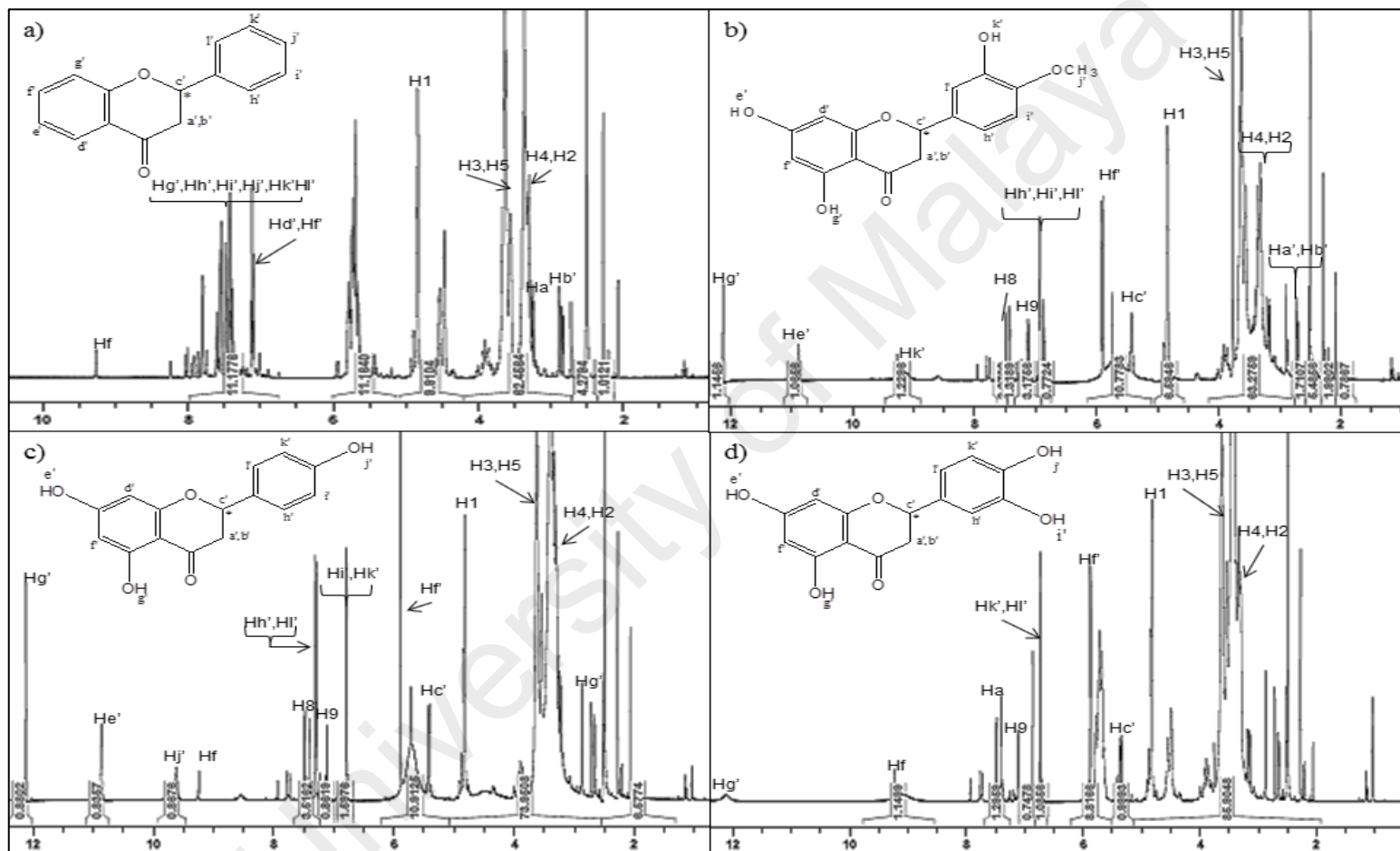


Figure 4.6: The deduced structure of a) β -CD-BIMOTs/flavanone complex, b) β -CD-BIMOTs/hesperetin complex, c) β -CD-BIMOTs/naringenin complex d) β -CD-BIMOTs/eriodictyol complex

For β -CD-BIMOTs/flavanone complex (Table 4.6), the significant changes were observed on $\Delta\delta$ at H5 proton located at the cavity of β -CD-BIMOTs due to inclusion complex formation. In addition, there is large shift at H2 proton located at the exterior torus of β -CD-BIMOTs caused by hydrogen bonding. The NOESY spectra in Figure 4.7 shows the cross-peak between H1, H2 and H5 protons of β -CD-BIMOTs with Hg' and Hj' protons of flavanone proved that the inclusion complex and hydrogen bonding were formed between flavanone and β -CD-BIMOTs.

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Table 4.6: Chemical shifts (δ) and induced shifts ($\Delta\delta$) of β -CD-BIMOTs and β -CD-BIMOTs/flavonoids

	β -CD- BIMOTs	β -CD- BIMOTs/Flavanone	β -CD- BIMOTs/Hesperetin	β -CD- BIMOTs/Naringenin	β -CD- BIMOTs/Eriodictyol				
	δ	δ	$\Delta\delta$	δ	$\Delta\delta$	δ	$\Delta\delta$	δ	$\Delta\delta$
H1	4.8405	4.8872	0.0467	4.8381	-0.0024	4.8241	-0.0164	4.8365	-0.004
H2	3.3312	3.2568	-0.0744	3.3214	-0.0138	3.2406	-0.0946	3.34	0.0048
H3	3.6394	3.6392	-0.0002	3.6401	0.0007	3.6253	-0.0141	3.6235	-0.0159
H4	3.3716	3.3797	0.0081	3.3552	-0.0164	3.3989	0.0273	3.4438	0.0722
H5	3.5777	3.5572	-0.0205	3.5586	-0.0191	3.5443	-0.0334	3.5428	-0.0349
H6	3.9225	3.9110	-0.0115	3.9185	-0.004	3.9053	-0.0172	3.8979	-0.0246
H8	7.4215	7.4374	0.0159	7.4276	0.0061	7.4128	-0.0087	7.4105	-0.011
H9	7.1112	7.1142	0.0030	7.1281	0.0169	7.1174	0.0062	7.1199	0.0087
H11	2.0847	2.0821	-0.0026	2.0844	-0.0003	2.0706	-0.0141	2.0698	-0.0149
Ha	7.4314	7.4827	0.0513	7.4995	0.0681	7.4873	0.0559	7.4756	0.0442
Hb	7.7957	7.8025	0.0068	7.8019	0.0062	7.7771	-0.0186	7.765	-0.0307
Hc	7.7542	7.7892	0.035	7.7552	0.001	7.738	-0.0162	7.7274	-0.0268
Hd	-	-	-	-	-	-	-	-	-
He	7.9563	7.9472	-0.0091	7.9456	-0.0107	7.9333	-0.023	7.9312	-0.0251
Hf	9.234	9.2696	0.0302	9.2744	0.035	9.2419	0.0025	9.2252	-0.0142
Hg	5.4371	5.4471	0.0100	5.4191	-0.018	5.4067	-0.0304	5.4000	-0.0371

-: overlap peak

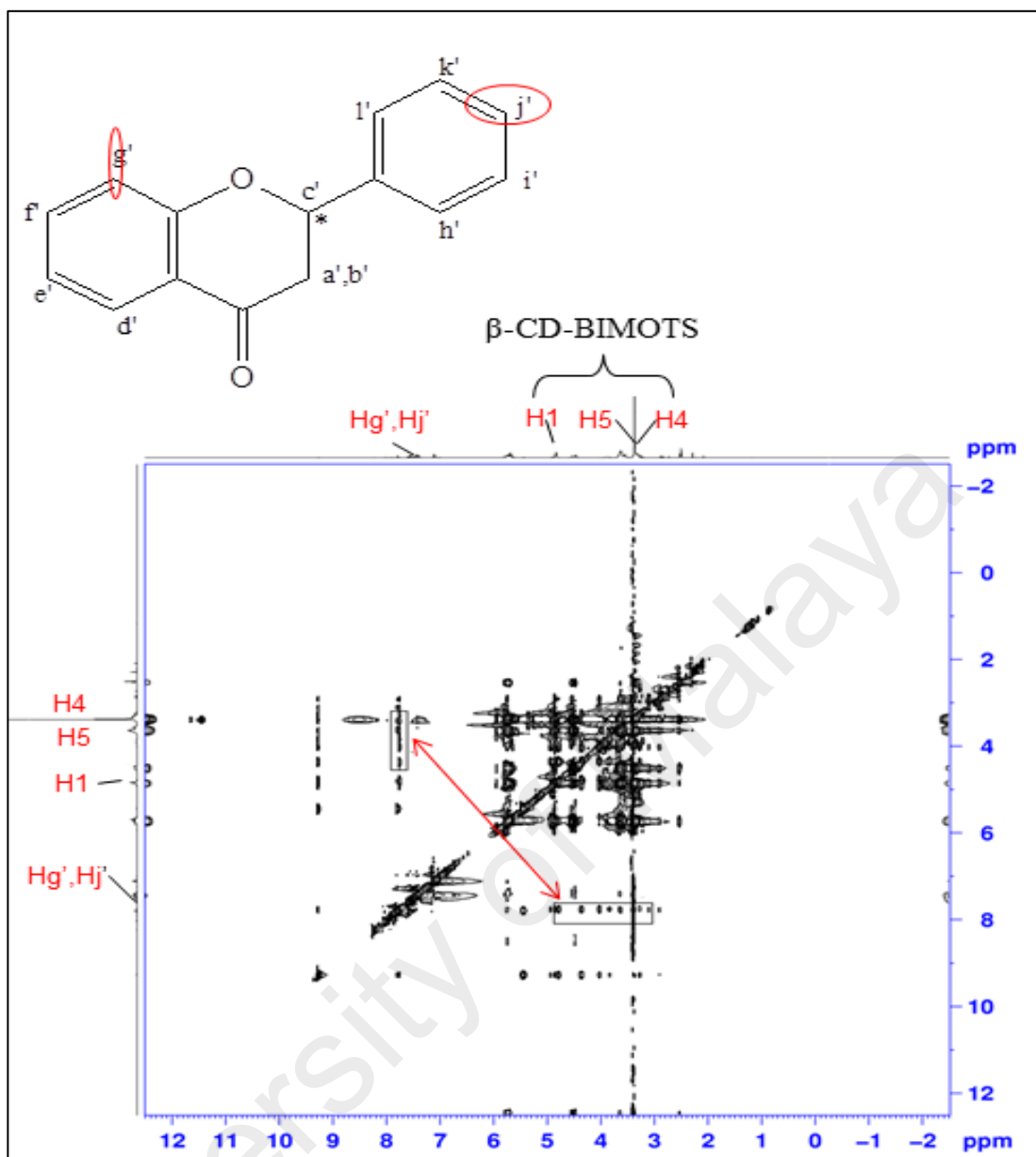


Figure 4.7: NOESY spectra of β -CD-BIMOTs/flavanone

Meanwhile, for hesperetin which is weakly acidic (pK_a 7.9) also formed neutral species at pH 7 and able to form inclusion complex with the cavity of β -CD-BIMOTs CSP. Thus, it was effectively enantioseparated using β -CD-BIMOTs based CSP (Table 4.5). Hesperetin bearing methoxy group is more hydrophobic than naringenin and eriodictyol. Therefore, hesperetin has greater affinity towards the cavity of β -CD-BIMOTs CSP as compared to naringenin and eriodictyol. Hesperetin was not enantioseparated at pH 4 and 9. At acidic pH, hesperetin is in neutral form (Ficarra *et al.*, 2002) but the TEAA species in the mobile phase compete with it for the inclusion formation (Kavalirova *et al.*, 2004). Meanwhile, the protonated hesperetin at pH 9 was not favored to form inclusion complex with β -CD (Raovv *et al.*, 2013). This finding further support the role of inclusion complex formation in enantioseparation of β -CD based CSPs. Moreover, OH groups and aromatic rings of hesperetin can form hydrogen bonding and π - π interaction with β -CD-BIMOTs CSP and thus enhanced the enantioseparation. These interactions were further proven using ^1H NMR and NOESY of β -CD-BIMOTs/hesperetin complex. The β -CD-BIMOTs/hesperetin complex shows appreciable shift at H4 proton at exterior torus of β -CD-BIMOTs because of hydrogen bonding. There are also large shift at H5 proton located in cavity of β -CD-BIMOTs (Table 4.6) which related to the formation of inclusion complex. In addition, the NOESY spectra (Figure 4.8) shows the cross-peaks between H3, H4 and H5 protons of β -CD-BIMOTs with He', Hg', and Hk' protons of hesperetin also proved that the inclusion complex and hydrogen bonding were formed with β -CD-BIMOTs which enhanced the enantioseparation.

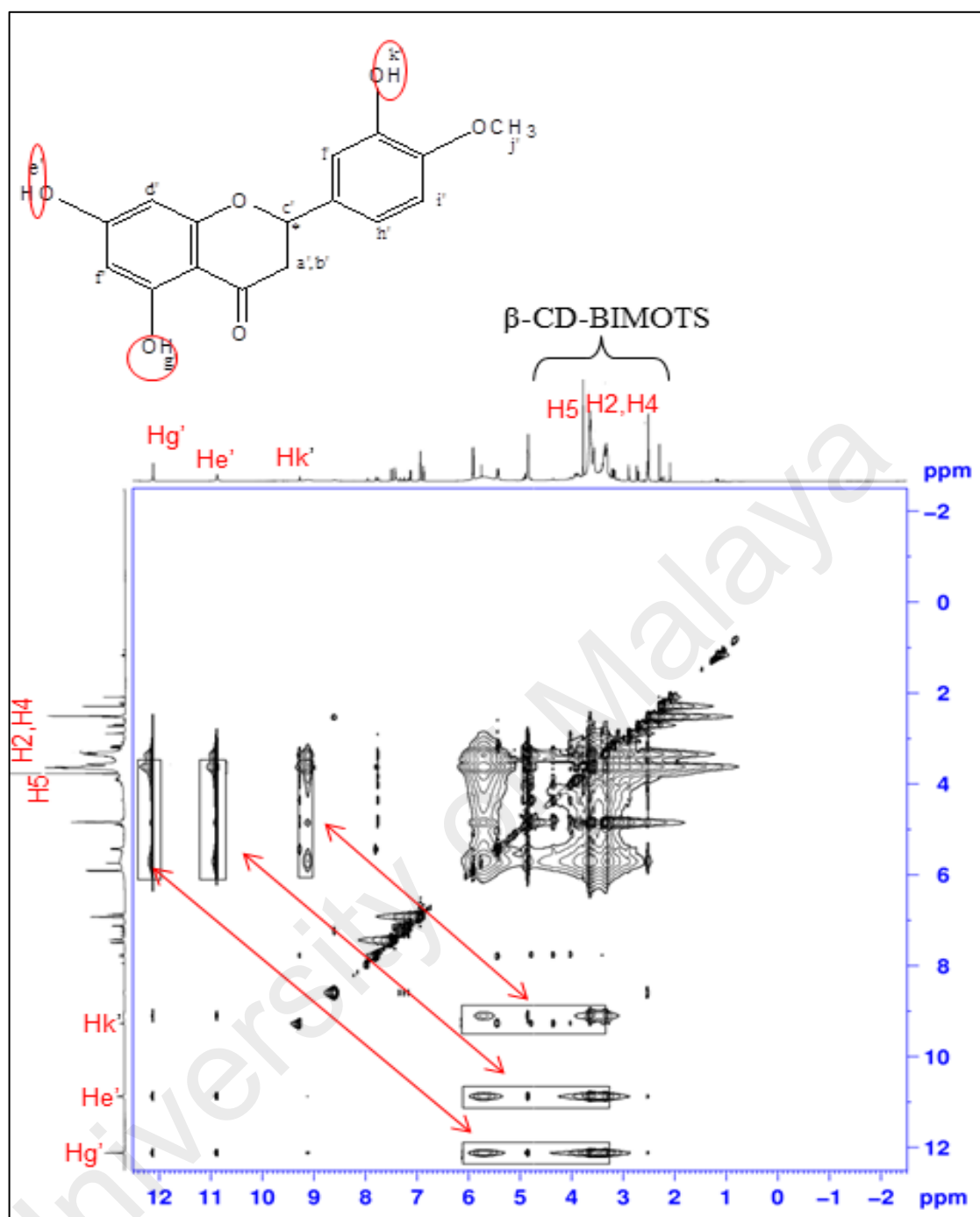


Figure 4.8: NOESY spectra of β -CD-BIMOTs/hesperetin

As shown in Table 4.5, naringenin and eriodictyol are not resolved in the reverse phase mode. Naringenin and eriodictyol contains highly polar moieties (OH) which might weaken the hydrophobic interaction with β -CD-BIMOTs cavity and retard the formation of inclusion complexes. Naringenin and eriodictyol might prefer to form hydrogen bonding at exterior torus instead of interior cavity of β -CD-BIMOTs CSP. Moreover, the presence of OH functionality as electron donating group could increase the electron density of aromatic ring of naringenin and eriodictyol and facilitate the π - π repulsion which weaken the π - π interaction (Hunter *et al.*, 2001). It can be deduced that hydrogen bonding is not sufficient to produce enantio-recognition. ^1H NMR of β -CD-BIMOTs/naringenin and β -CD-BIMOTs/eriodictyol complexes were studied to get detail information of the interaction. Large $\Delta\delta$ of H2 and H4 protons of β -CD-BIMOTs with the presence of naringenin and eriodictyol was observed, respectively (Table 4.6). In addition, NOESY spectra for β -CD-BIMOTs/naringenin complex (Figure 4.9) showed the cross-peak between He', Hg' and Hj' protons of naringenin with H2 proton of β -CD-BIMOTs. In NOESY spectra of β -CD-BIMOTs/eriodictyol complex (Figure 4.10), there are cross-peak between Hc', Hg' and Hf' protons of eriodictyol with H4 proton of β -CD-BIMOTs. These results suggest that there are hydrogen bonding between naringenin and eriodictyol at exterior torus of β -CD-BIMOTs.

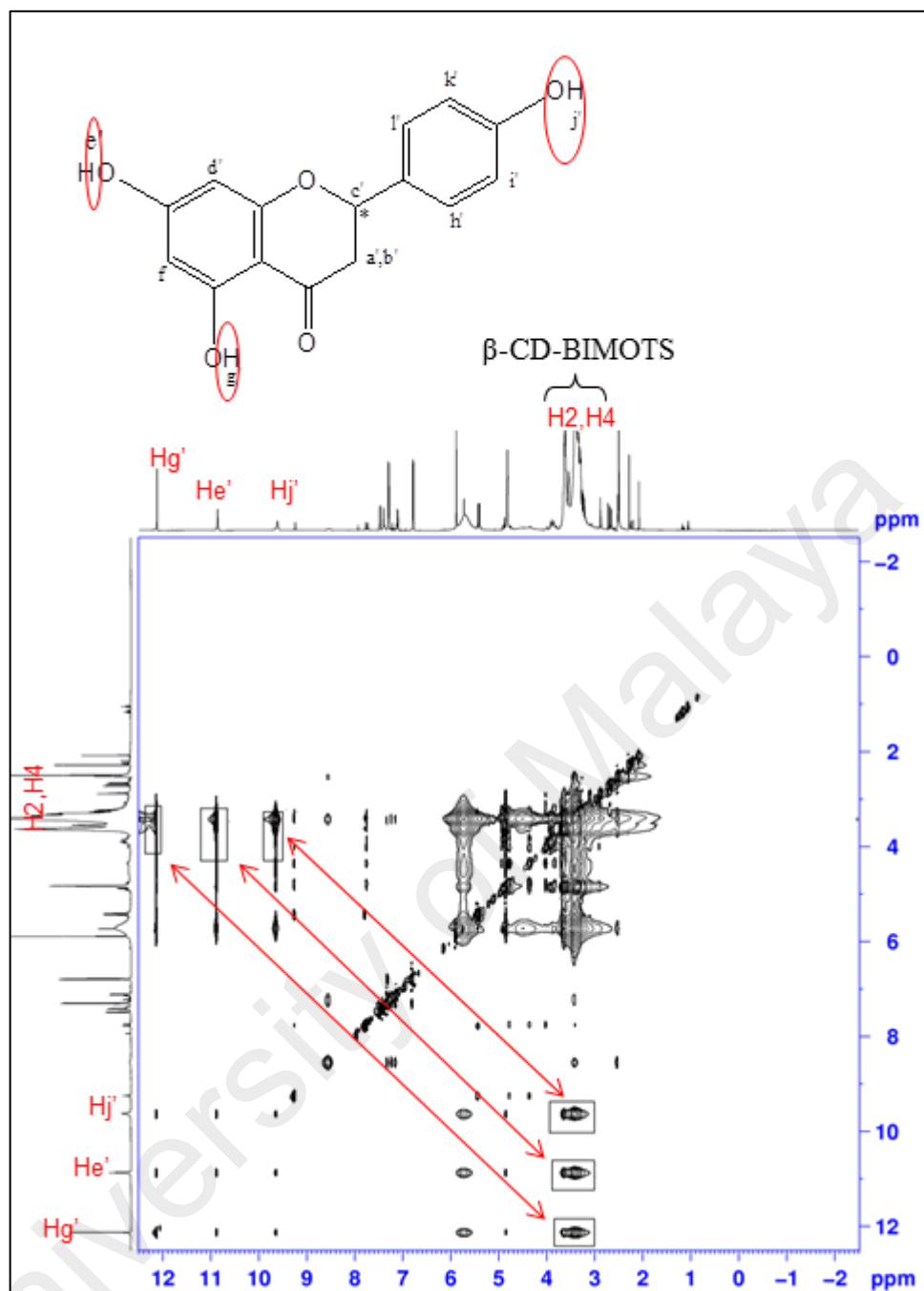


Figure 4.9: NOESY spectra of β -CD-BIMOTs/naringenin

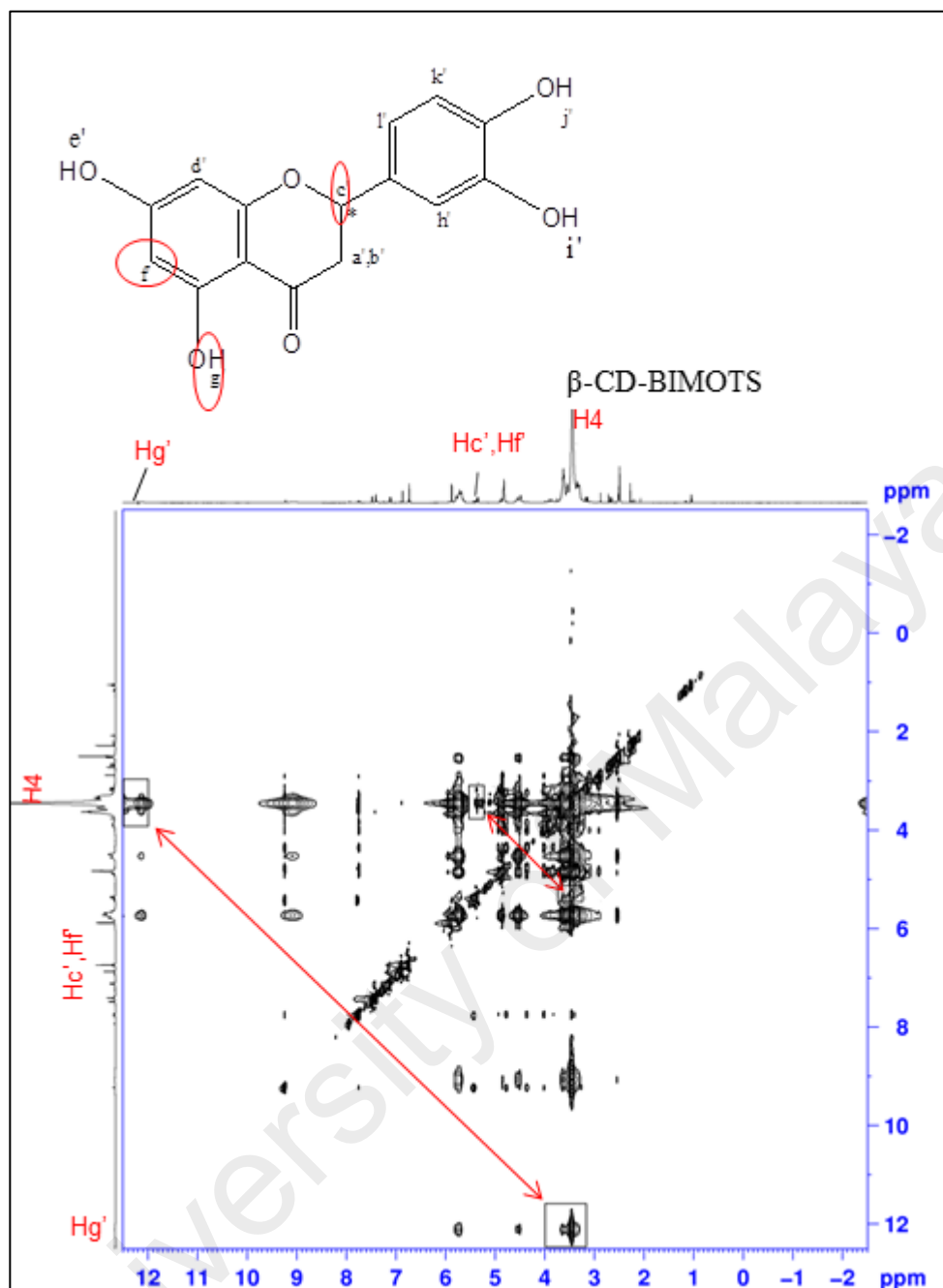


Figure 4.10: NOESY spectra of β -CD-BIMOTs/eriodictyol

As a part of the optimization, the polar organic mode with different additives was used to improve the enantioseparation of naringenin and eriodictyol. This system can be used to resolve compounds that cannot be separated in the reverse phase mode. In this study, the mobile phase of polar organic mode was the mixture of ACN and MeOH. The selected additives were TEA and HOAc (Kafkova *et al.*, 2005). In the polar organic mode, the relative high concentration of organic solvents occupies the relatively hydrophobic cavity of β -CD. Armstrong *et al.* (1993) proposed that the analytes may form a “lid” over the “mouth” of the cavity. Moreover, the retention and selectivity are mainly due to the polar OH groups at the rims of β -CD forming hydrogen bonding with analytes (Chang *et al.*, 1993). Thus, the total number of OH moiety at flavonoids would affect the enantioseparation. The HPLC chromatograms shown naringenin achieved better enantioseparation at higher amount of TEA (Figure 4.11) meanwhile eriodictyol was resolved at higher amount of HOAc (Figure 4.12). At higher amount of TEA, naringenin which has less OH groups than eriodictyol tends to carry less number of deprotonated OH. Thus, naringenin prefer to form electrostatic interaction associated with hydrogen bonding which facilitated the enantioseparation. Meanwhile, eriodictyol which has highest number of deprotonated OH led to the stronger electrostatic interaction with β -CD-BIMOTs and thus inhibit the enantioseparation.

At higher ratio of HOAc, both of naringenin and eriodictyol are in neutral form. Under this condition, enantioseparation of eriodictyol was achieved better than naringenin. This might due to the structure of eriodictyol with 4 OH groups that have high capability to form hydrogen bonding at the exterior torus of β -CD-BIMOTs. It can be deduced that the better enantioseparation in the polar organic mode shows the importance of the hydrogen bonding and/or electrostatic interaction for the chiral recognition of naringenin and eriodictyol.

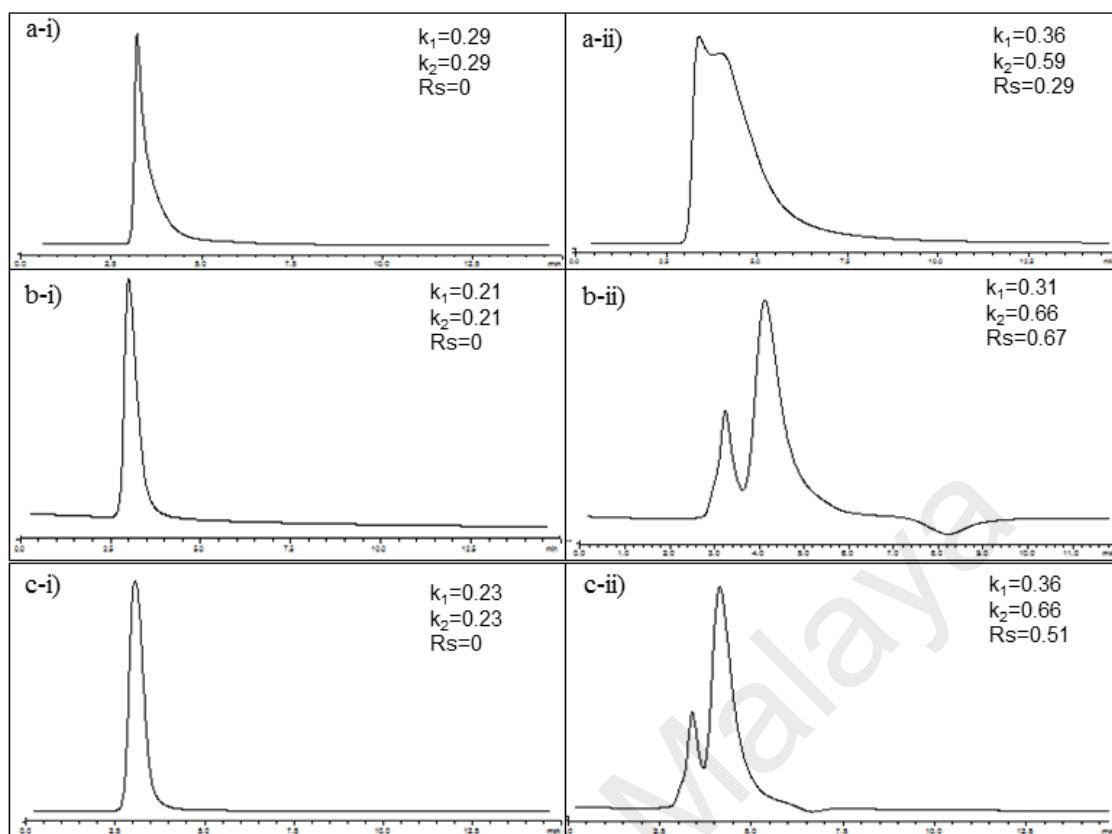


Figure 4.11: HPLC chromatograms of naringenin in polar organic mode. Mobile phase composition, ACN/MeOH/TEA/HOAc (v/v/v/v): a-i) 90/10/1/3, a-ii) 90/10/3/1, b-i) 50/50/1/3, b-ii) 50/50/3/1, c-i) 30/70/1/3 and c-ii) 30/70/3/1

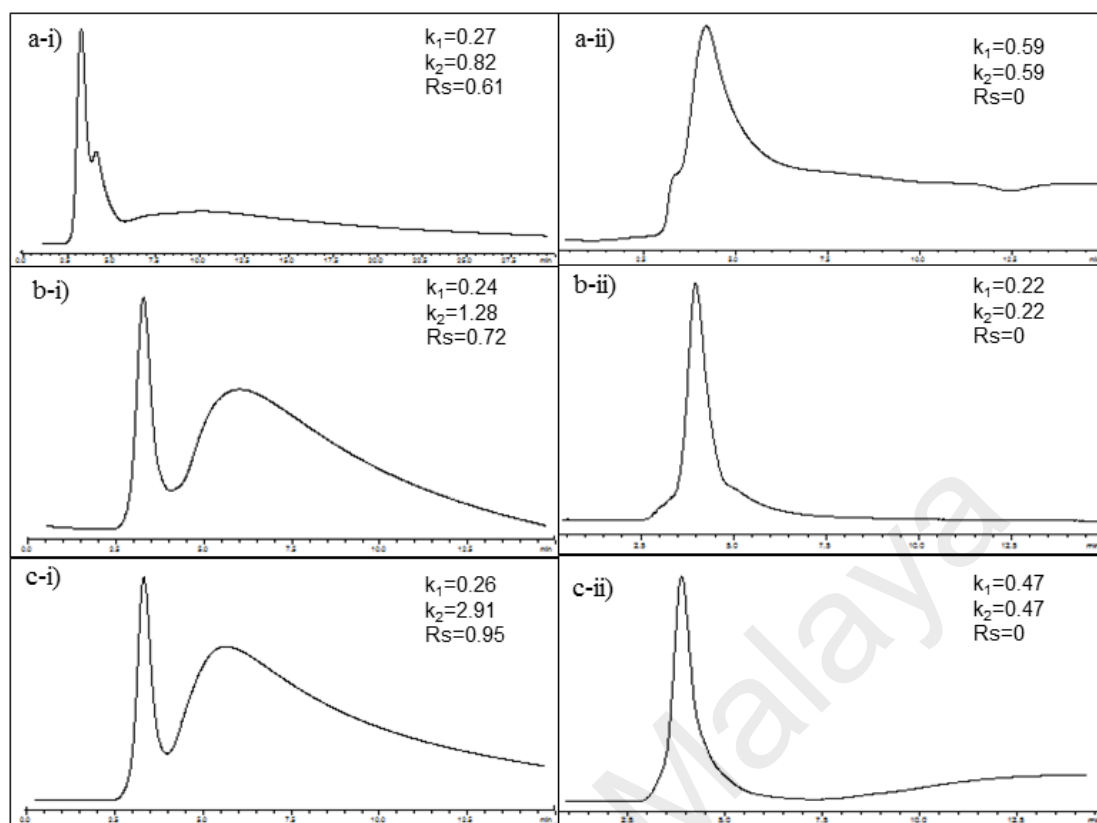


Figure 4.12: HPLC chromatograms of eriodictyol in polar organic mode. Mobile phase composition, ACN/MeOH/TEA/HOAc (v/v/v/v): a-i) 90/10/1/3, a-ii) 90/10/3/1, b-i) 50/50/1/3, b-ii) 50/50/3/1, c-i) 30/70/1/3 c-ii) 30/70/3/1

The chromatogram of eriodictyol (Figure 4.12(c-i)) with the broad and tailing peak was caused by the formation of strong hydrogen bonding with β -CD-BIMOTs CSP. Thus, it can be deduced that the higher number of OH groups leads to the stronger interaction with β -CD-BIMOTs CSP and thus, inhibit the enantioseparation. Consequently, the formation constant (K) was determined to study the strength of the interaction between flavonoids and β -CD-BIMOTs. In the experiment, the plots of absorption for β -CD-BIMOTs, flavonoids and β -CD-BIMOTs/flavonoids complexes were first measured (Figure 4.13) by monitoring the UV spectra. The results showed that β -CD-BIMOTs had a λ_{\max} in the range of 230-260 nm. The absorption spectra of flavanone displayed two well-defined λ_{\max} at 250 and 320 nm meanwhile naringenin, hesperetin and eriodictyol displayed one λ_{\max} at 320 nm. The λ_{\max} of β -CD-

BIMOTs/flavonoids complex was observed at 230-260 nm referred to β -CD-BIMOTs. Meanwhile, the λ_{max} at 320 nm of β -CD-BIMOTs/flavonoids complex was referred to flavonoids. It was observed that the absorption spectra of all β -CD-BIMOTs/flavonoids complexes showed both hyperchromic and hypochromic effect. Increase in absorption at λ_{max} is defined as hyperchromic effect and decrease in the absorption at λ_{max} is defined as hypochromic effect (Hu *et al.*, 2012; Ventura *et al.*, 2006). Hyperchromic effect that observed in the UV spectra of β -CD-BIMOTs-flavonoids at 320 nm was due to the electron perturbation at the chromophore of flavonoids (Ventura *et al.*, 2006). Meanwhile the hypochromic effect is due to the intercalative mode involving the stacking interaction (Hu *et al.*, 2012) which was mainly referred to π - π interaction between aromatic ring of flavonoids and β -CD-BIMOTs. The hypochromic effect for β -CD-BIMOTs-flavanone was not observed due to the overlapping of absorption band at 250 nm (Figure 4.13(a)). Both hyperchromic and hypochromic effects observed in the absorption spectra of β -CD-BIMOTs-flavonoids proved that there were multiple interactions between β -CD-BIMOTs and flavonoids.

The K values were then calculated (using Equation 3-6) from the slope of $\frac{1}{(A-A_0)}$ versus $\frac{1}{[\beta\text{-CD-BIMOTs}]}$ of β -CD-BIMOTs/flavonoids as shown in Figure 4.14. In Table 4.7, the K values obtained are in the following order: β -CD-BIMOTs/hesperetin < β -CD-BIMOTs/flavanone < β -CD-BIMOTs/naringenin < β -CD-BIMOTs/eriodictyol. This deduced that the strength of interaction is correlated with the substituted OH group at flavonoids. Previous study reported that hydrogen bond is the strongest non-covalent interactions with 2-10 kcal/mol stabilization energy (Frieden, 1975). Naringenin and eriodictyol that possess 3 and 4 OH groups experienced highest K values indicating the stronger hydrogen bond formation. Indeed, these results clarified that naringenin and eriodictyol interacted at the external torus of β -CD-BIMOT. Meanwhile, the small K

values for flavanone and hesperetin proven that the inclusion complex was formed due to hydrophobic interaction and facilitated the enantioseparation.

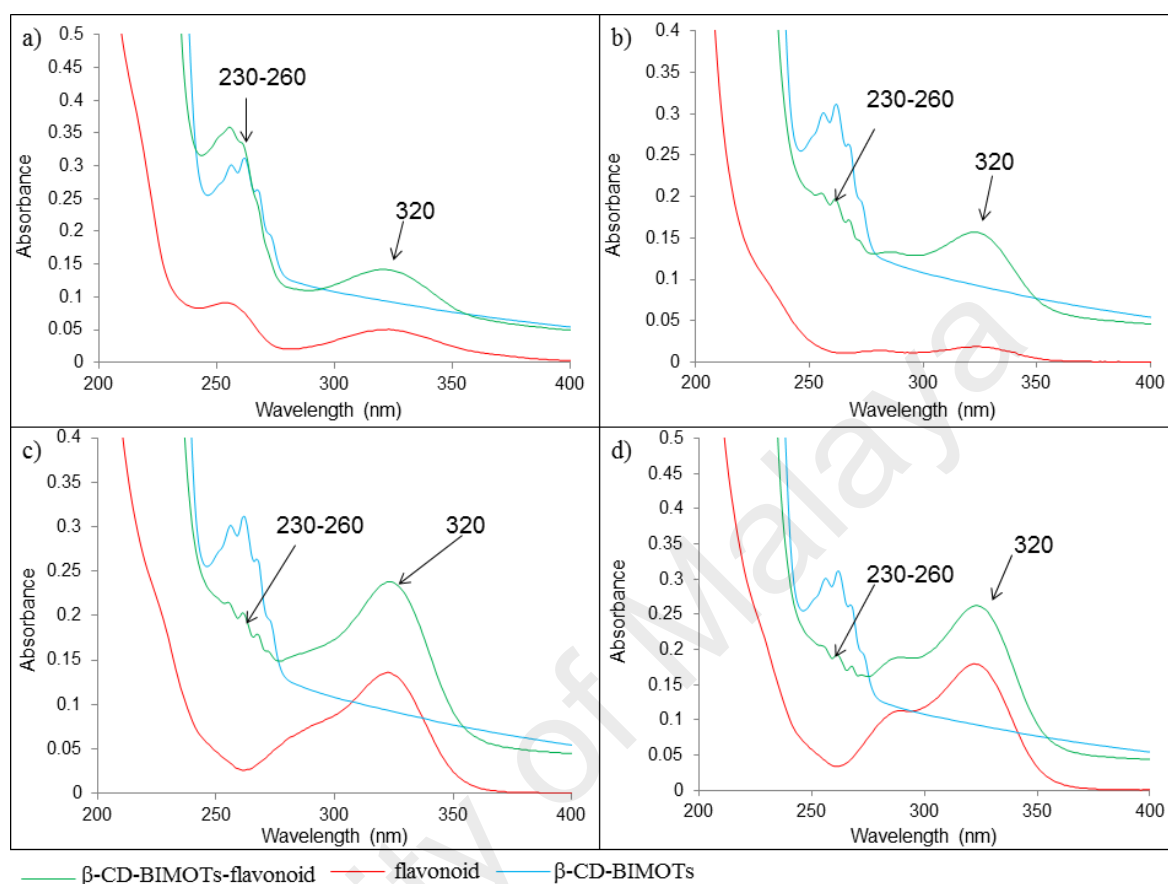


Figure 4.13: Absorption spectra of a) β -CD-BIMOTs/flavanone b) β -CD-BIMOTs/hesperetin c) β -CD-BIMOTs/naringenin d) β -CD-BIMOTs/eriodictyol with $[\beta$ -CD-BIMOTs]: 0.032mM [Flavonoids]: 0.01mM; T = 25 °C

Table 4.7: *K* values for β -CD-BIMOTs/flavonoids

Flavonoids	<i>K</i>
Flavanone	722
Hesperetin	572
Naringenin	1077
Eriodictyol	6032

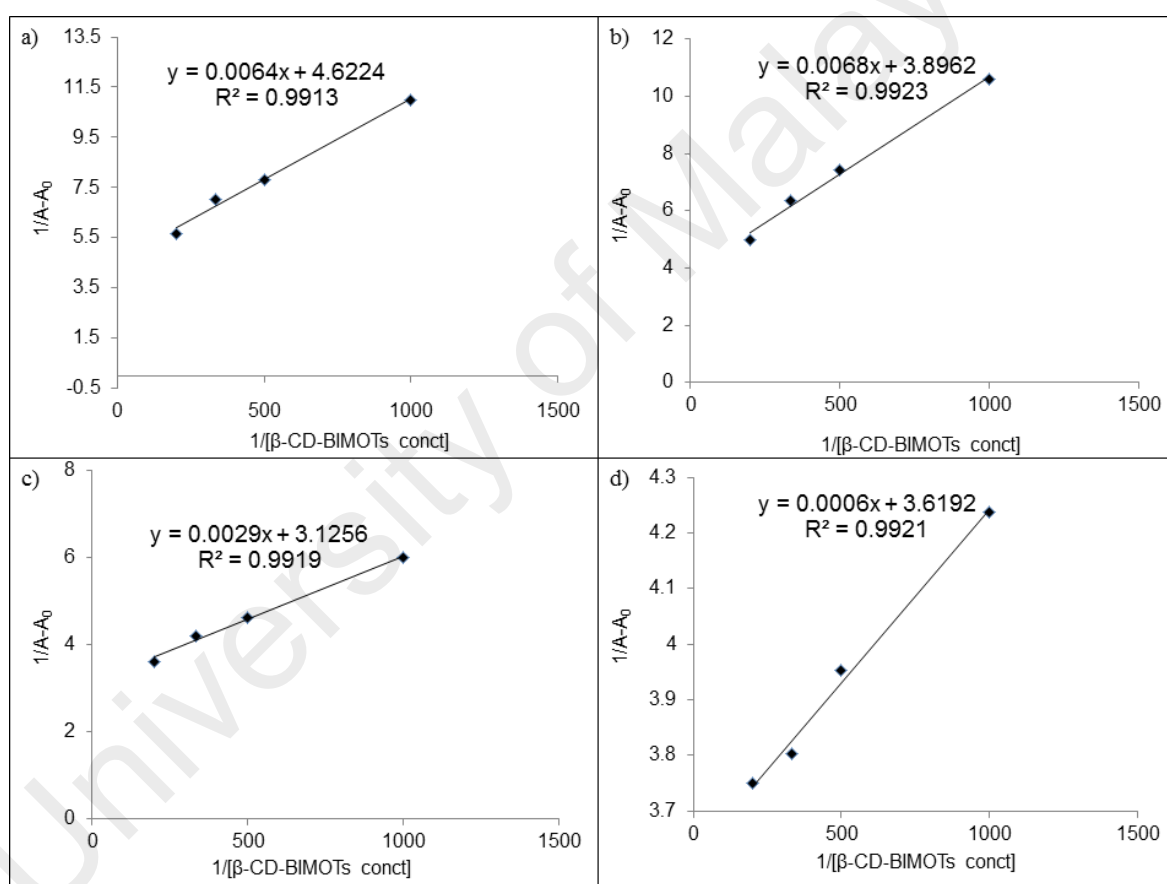


Figure 4.14: Benesi-Hildebrand plot of $1/A-A_0$ versus $1/[\beta\text{-CD-BIMOTs}]$ for a) β -CD-BIMOTs/flavanone, b) β -CD-BIMOTs/hesperetin, c) β -CD-BIMOTs/naringenin d) β -CD-BIMOTs/eriodictyol

4.4 Enantioseparation performance of β -blockers

The enantioselective ability of β -CD-BIMOTs CSP was also examined for chiral compounds with basic properties, β -blockers to study the enantiomeric behavior and the mechanism of enantioseparation. The baseline separation was achieved for the enantiomers of propranolol and metoprolol as shown in Table 4.8. Among the selected β -blockers, propranolol and metoprolol achieved the R_s values of 3.10 and 2.38, respectively. Complete enantioseparation of propranolol and metoprolol was achieved in 30 min. However, for pindolol and atenolol, no peak was observed even after 120 min due to the high retention of these compounds onto β -CD-BIMOTs CSP. β -Blockers can be divided according to its lipophilic (propranolol and metoprolol) and hydrophilic (pindolol and atenolol) nature (Borchard, 1998). The result indicated hydrophilic atenolol and pindolol with polar amide and indole moiety showed stronger interaction with CSP that contribute to high retention. On the other hand, it is proven that the β -blockers with lipophilicity properties were well enantioseparated than the hydrophilic β -blockers.

The enantioseparation of propranolol and metoprolol were separated excellently using β -CD-BIMOTs CSP and this might due to the formation of inclusion complex between the analytes and β -CD through the stereogenic center of β -CD located at the interior cavity. In order to verified this interaction, the inclusion complexes of β -CD-BIMOTs and selected β -blockers were prepared. ^1H NMR and NOESY were used to study the interaction between β -CD-BIMOTs and β -blockers in the complexes. The values of the chemical shifts (δ) and induced shifts ($\Delta\delta$) for different protons in β -CD-BIMOTs, β -blockers and β -CD-BIMOTs/ β -blockers complexes are listed in Table 4.9 and Table 4.10.

Table 4.8: Chiral separation data for the β -blockers on β -CD-BIMOTs CSP in neutral pH mobile phase

β -blockers	Conditions	β -CD-BIMOTs CSP			
		k_1'	k_2'	α	R_s
Atenolol	ACN/water-90/10	n.a	n.a	n.a	n.a
	ACN/water-50/50	n.a	n.a	n.a	n.a
	ACN/water-30/70	n.a	n.a	n.a	n.a
Metoprolol	ACN/water-90/10	2.04	3.64	1.78	2.38
	ACN/water-50/50	0.58	0.58	1.00	0
	ACN/water-30/70	0.65	0.65	1.00	0
Propranolol	ACN/water-90/10	2.83	4.88	1.72	3.10
	ACN/water-50/50	0.79	1.01	1.27	0.46
	ACN/water-30/70	0.84	1.10	1.30	0.43
Pindolol	ACN/water-90/10	n.a	n.a	n.a	n.a
	ACN/water-50/50	n.a	n.a	n.a	n.a
	ACN/water-30/70	n.a	n.a	n.a	n.a

n.a: not available

Table 4.9: Chemical shifts (δ) corresponding to β -CD-BIMOTs in presence of β -blockers

	β -CD-BIMOTs	β -CD-BIMOTs/ atenolol		β -CD-BIMOTs/ metoprolol		β -CD-BIMOTs/ propranolol		β -CD-BIMOTs/ pindolol	
	δ	δ	$\Delta\delta$	δ	$\Delta\delta$	δ	$\Delta\delta$	δ	$\Delta\delta$
H1	4.8405	4.8301	-0.0104	4.8249	-0.0156	4.8285	-0.012	4.8329	-0.0076
H2	3.3312	3.3483	0.0171	3.3425	0.0113	3.3042	-0.027	3.3476	0.0155
H3	3.6394	3.6311	-0.0083	3.6274	-0.0120	3.6309	-0.0085	3.6335	-0.0059
H4	3.3716	3.4304	0.0588	3.4660	0.0944	3.3762	0.0046	3.4391	0.0675
H5	3.5777	3.5488	-0.0289	3.5464	-0.0313	3.5531	-0.0246	3.5580	-0.0197
H6	3.9225	3.9473	0.0248	3.9272	0.0047	3.9041	-0.0184	3.9041	-0.0184
H8	7.4215	7.4212	-0.0003	7.4202	-0.0013	overlap	-	7.4361	0.0146
H9	7.1112	7.1227	0.0115	-	-	7.1192	0.0008	7.1259	0.0147
H11	2.0847	2.0797	-0.0050	-	-	-	-	-	-
Ha	7.4314	7.4798	0.0484	7.4752	0.0438	7.4832	0.0518	7.4896	0.0582
Hb	7.7957	7.7903	-0.0054	7.7892	0.0350	7.8063	0.0106	7.8081	0.0124
Hc	7.7542	7.7402	-0.014	7.7391	-0.0151	7.7490	-0.0052	7.7473	-0.0069
Hd	-	-	-	-	-	-	-	-	-
He	7.9563	7.9440	-0.0123	-	-	-	-	-	-
Hf	9.2394	9.2606	0.0212	9.2807	0.0413	9.3132	0.0738	9.3379	0.0985
Hg	5.4371	5.4400	0.0029	5.4460	0.0089	5.4369	-0.0002	5.4482	0.0111

 $\Delta\delta$: induced shifts

-: overlap peak

Table 4.10: Induced shifts ($\Delta\delta$) corresponding to β -blockers in presence of β -CD-BIMOTs

	β -CD- BIMOTs/atenolol	β -CD- BIMOTs/metoprolol	β -CD- BIMOTs/propranolol	β -CD- BIMOTs/pindolol
	$\Delta\delta$	$\Delta\delta$	$\Delta\delta$	$\Delta\delta$
Ha'	0.1059	0.0995	-0.0018	0.1140
Hb'	0.1345	0.0055	-	0.2063
Hc'	0.1060	0.0995	-0.0018	0.1140
Hd'	-	-0.0075	-0.0160	-0.0067
He'	0.1596	-0.0057	-0.0044	0.1766
Hf'	0.0545	-	-0.0063	-
Hg'	-	-0.0075	-0.0246	-0.0067
Hh'	-0.0008	-0.0269	0.0794	0.0248
Hi'	0.0118	0.0048	-0.0012	0.0041
Hj'	0.0096	0.0064	-0.0027	0.0162
Hk'	0.0096	-0.0009	-0.0025	0.0162
Hi'	0.0132	0.0003	-0.0131	0.0836
Hm'	0.0054	-	-0.0037	-0.0006
Hn'	-	-	-0.0011	0.0056
Ho'	-	-	-0.0055	-

-: overlap peak

The deduced structures of β -CD-BIMOTs/ β -blockers complexes are shown in Figure 4.15. For β -CD-BIMOTs/ β -blockers complexes, the presence of propranolol and metoprolol showed appreciable shift of H5 proton of β -CD-BIMOTs (Table 4.9). The upfield shifts for this proton proved the existence of an interaction between the analytes and the interior proton of β -CD-BIMOTs. Additionally, the larger $\Delta\delta$ value of H1' proton was observed for propranolol (Table 4.10). This indicated the perturbation at the aromatic ring of propranolol which might due to π - π interaction with IL at β -CD-BIMOTs. In contrast, the $\Delta\delta$ values of aromatic protons (Hi', Hj', Hk', Hl') of metoprolol were relatively small (Table 4.10). This result suggested that propranolol achieved better enantioseparation than metoprolol because of the additional π - π interaction that contributed by IL at β -CD-BIMOTs. Moreover, the greater shift of H4 proton of β -CD-BIMOT-metoprolol was observed as compared to other complexes. Higher electronegativity of oxygen atom at the methoxy group of metoprolol caused the lower electron density around the H4 proton. As a result, the proton was deshielded and experienced higher chemical shift. In Figure 4.16, the cross peak between Hm' and Hn' protons of propranolol with H5 proton β -CD-BIMOTs complex was observed in NOESY spectra. Meanwhile, in Figure 4.17, the cross peak between Hi' and Hj' protons of metoprolol with H5 proton of β -CD-BIMOTs complex was also observed. This indicated the interaction of propranolol and metoprolol at the interior protons of β -CD-BIMOTs.

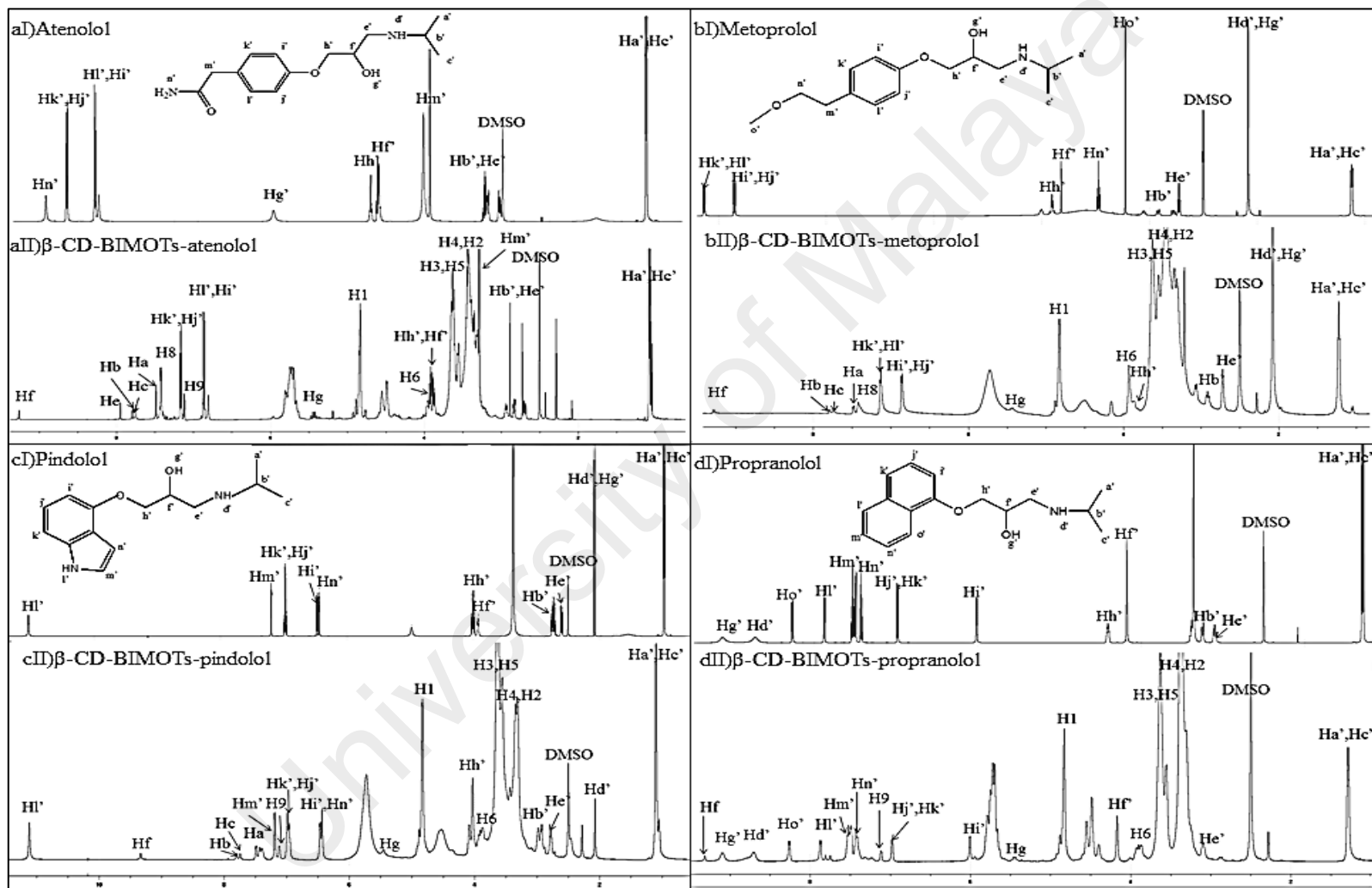


Figure 4.15: The deduced structure of β -CD-BIMOTs/ β -blockers complexes: a) atenolol, b) metoprolol, c) Pindolol, d) Propranolol

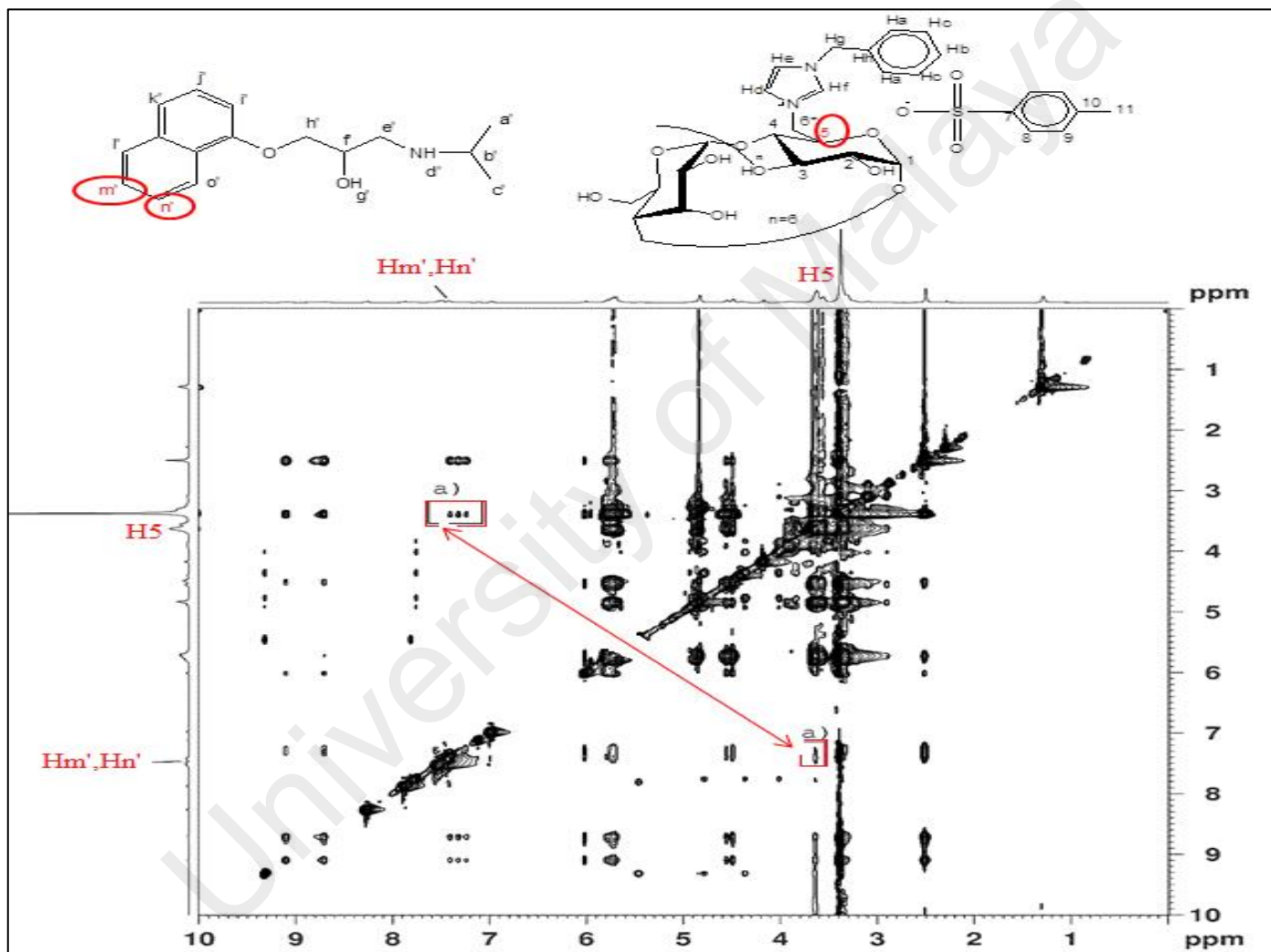


Figure 4.16: 2D NOESY spectra of β -CD-BIMOTs/propranolol complex

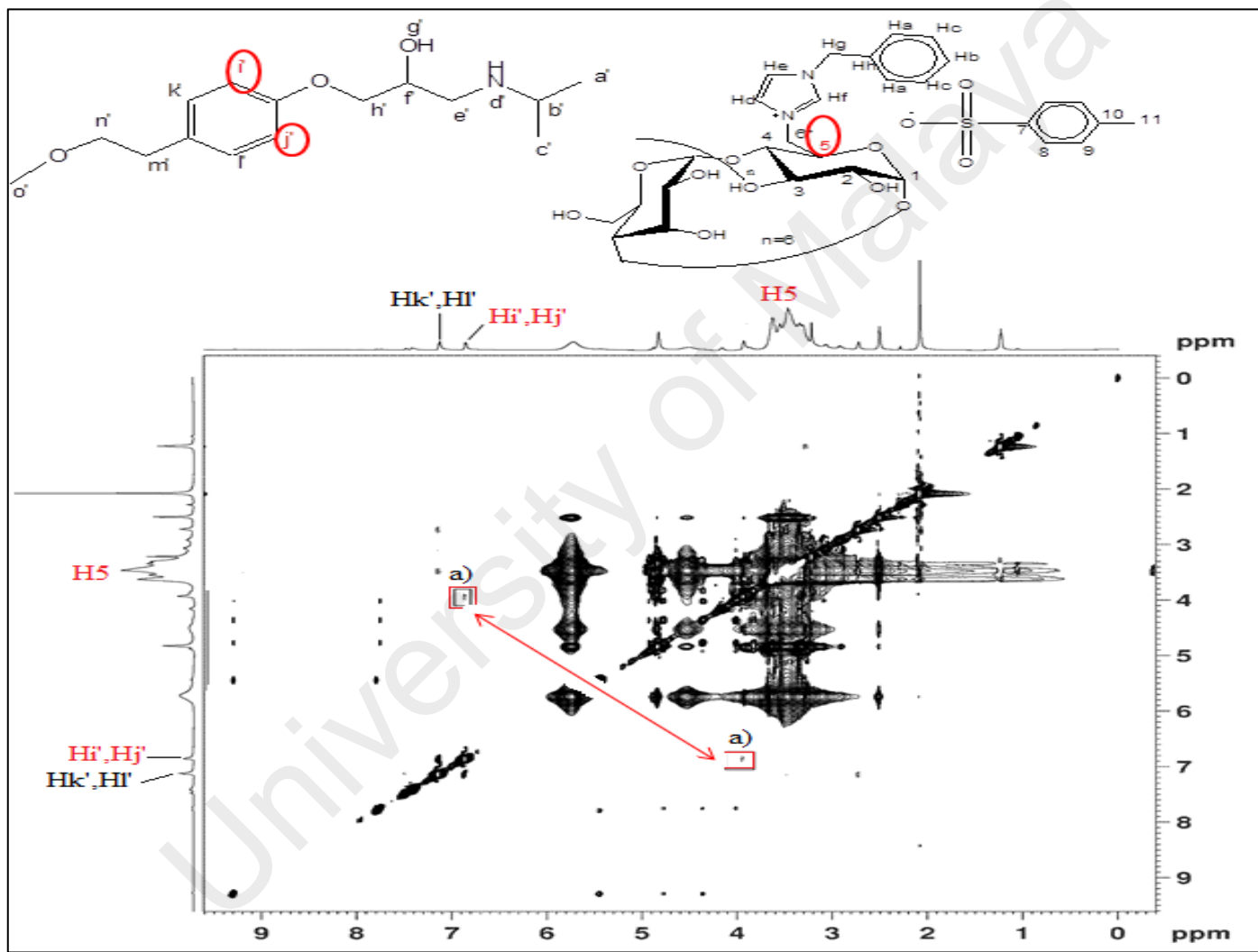


Figure 4.17: 2D NOESY spectra of β -CD-BIMOTs/metoprolol complex

From the ^1H NMR studied (Table 4.9), H4 (exterior proton) at β -CD-BIMOTs was experienced appreciably shifted downfield after forming complexes with pindolol or atenolol. This result suggested that pindolol and atenolol were not forming inclusion complex but it formed hydrogen bonding with exterior torus of β -CD-BIMOTs. Moreover, the large $\Delta\delta$ values were observed for Ha', Hb' and Hc' of pindolol and atenolol (Table 4.10). For β -CD-BIMOTs/pindolol complex, the NOESY spectra showed the cross-peak between H1' proton of pindolol with H1 and H4 protons of β -CD-BIMOTs (Figure 4.18). Meanwhile, β -CD-BIMOTs/atenolol complex showed the cross-peak between Hj' and Hk' protons of atenolol and H4 protons of β -CD-BIMOTs (Figure 4.19). This result indicated the close interaction of pindolol and atenolol at the exterior protons of β -CD-BIMOTs

The composition of the mobile phase also plays an important role in enantioseparation. The effect of ACN contents on enantioseparation of selected β -blockers can be seen from Table 4.8. The high k_1' and k_2' of propranolol and metoprolol at high organic content (90 % ACN) showed the normal phase behavior of the β -CD-BIMOTs CSP. On the other hand, when organic content is low (30 % ACN), the high k_1' and k_2' of propranolol and metoprolol showed typical reverse phase behavior of β -CD-BIMOTs CSP. Therefore, the retention behavior of β -blockers can be considered as the mixed reverse-normal separation mode (Guo *et al.*, 2009). In this separation mode, the retention mechanism is based on the distribution of the analytes between the ACN-rich mobile phase and water enriched layer adsorbed onto the polar stationary phase (Buszewski & Noga, 2012). Thus, for more hydrophilic analytes (pindolol and atenolol), partitioning equilibrium is shifted towards the immobilized water layer on the stationary phase, causing the analytes retained longer in column.

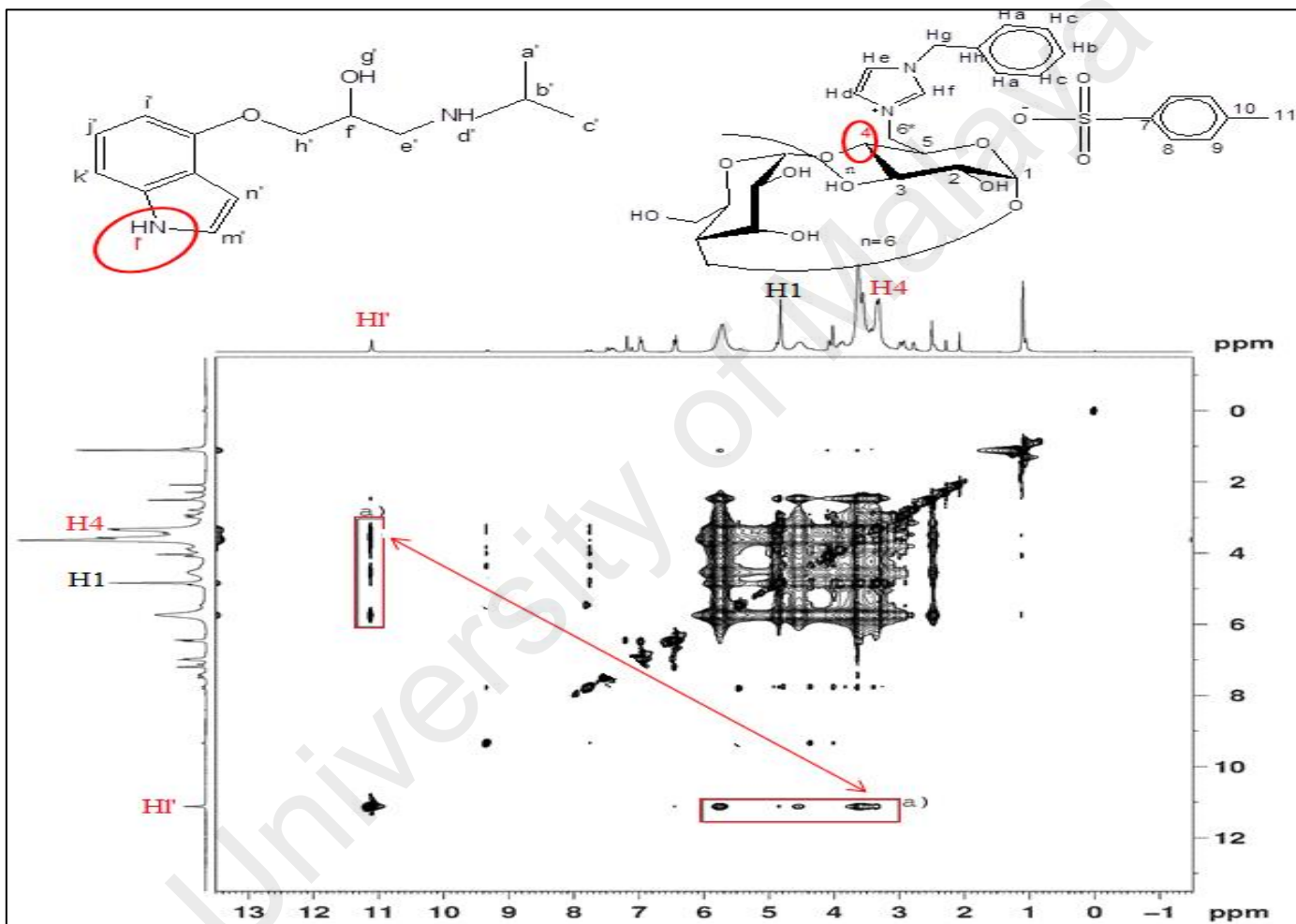


Figure 4.18: 2D NOESY spectra of β -CD-BIMOTs/pindolol complex

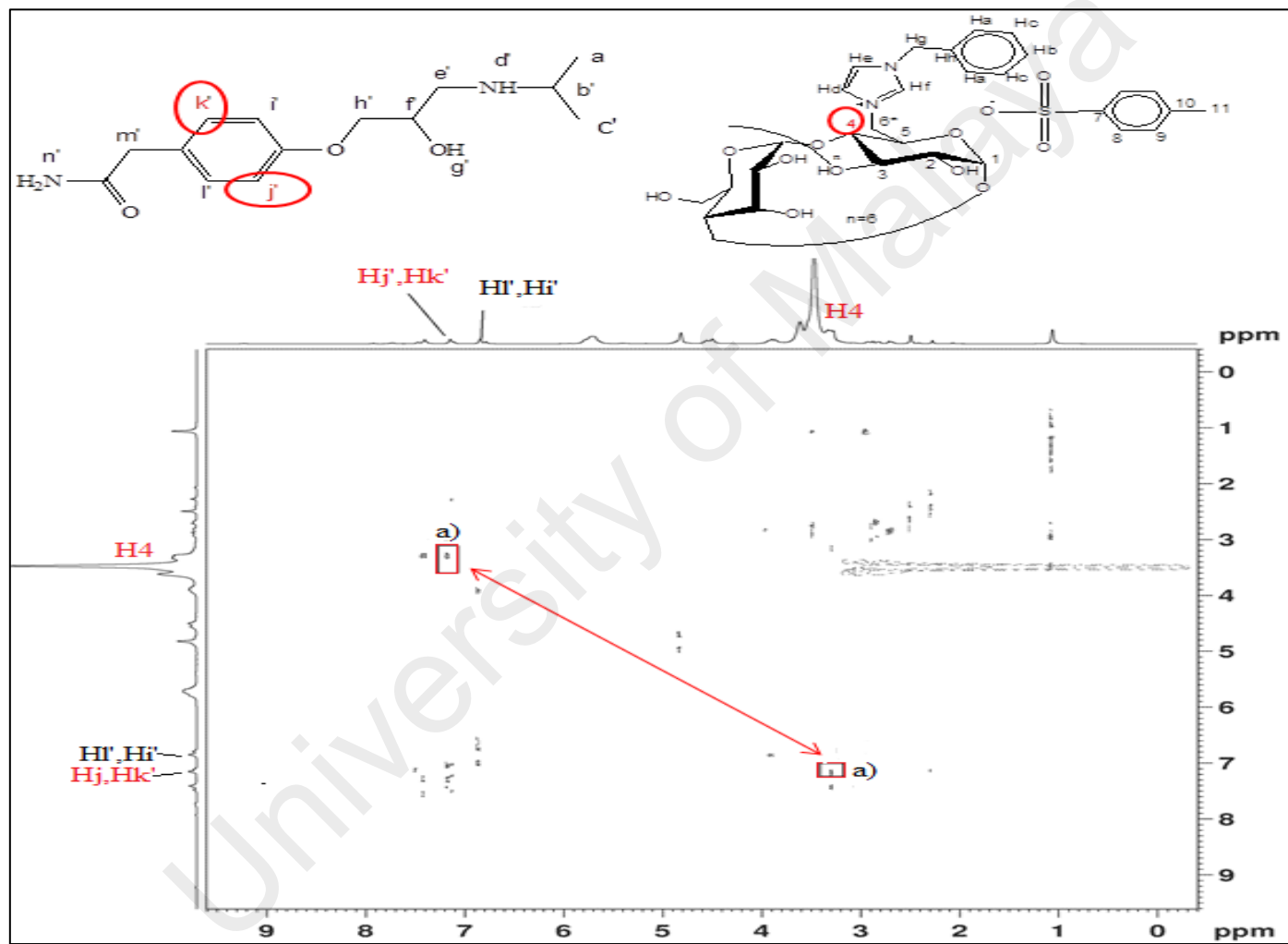


Figure 4.19: 2D NOESY spectra of β -CD-BIMOTs/atenolol complex

TEAA buffer was used to control the pH of mobile phase and ionic strength. Buffer can influence the degree of ionization of analytes and resulting in different retention behavior. The chromatograms in Figure 4.20 show the effect of pH towards the enantioseparation of β -blockers. Propranolol and metoprolol were not enantioseparated at pH 4 and 9. Meanwhile, they are well enantioseparated at pH 7. This is due to the deprotonation and protonation of β -blockers at pH 4 and 9, respectively. Protonated and deprotonated analytes were not favorable for the formation of inclusion complex with β -CD (Raovv *et al.*, 2013). This finding further support the role of inclusion complex formation in enantioseparation of β -CD based CSPs. Meanwhile, the retention time of pindolol and atenolol was reduced at pH 4 and 9 as compared to pH 7. Due to both of analytes and β -CD-BIMOTs CSP acquiring positive charges at pH 4, the electrostatic repulsion occurred and it reduced the retention time of analytes. At basic pH, the abundance of TEAA species reduces the retention time due to the competition between TEAA and protonated analytes.

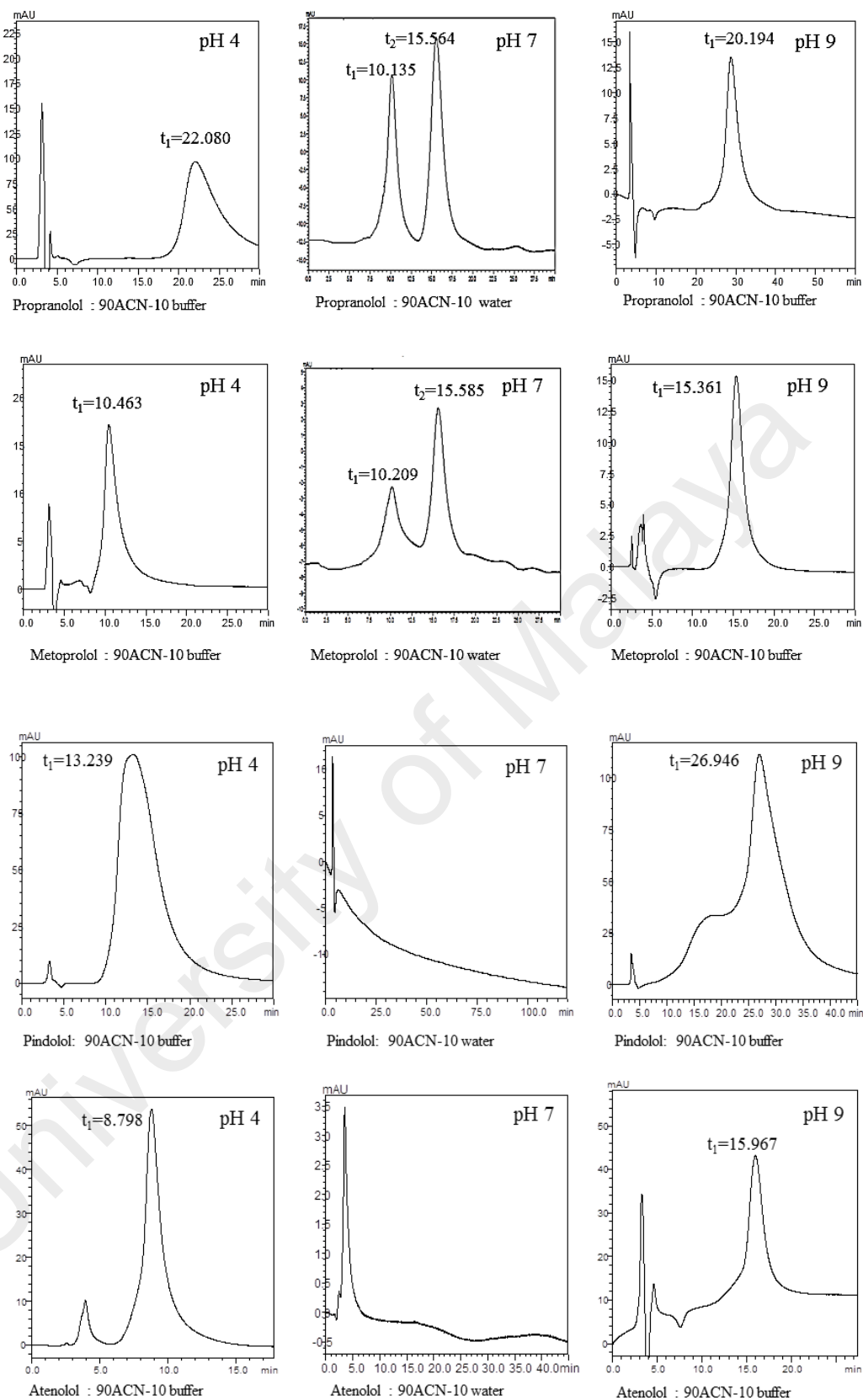


Figure 4.20: The chromatograms of propranolol, metoprolol, pindolol and atenolol responding to different pH of mobile phase

4.5 Enantioseparation performance of NSAIDs

In the final part of this work, the enantioselective ability of β -CD-BIMOTs CSP was examined using chiral compounds with acidic properties, NSAIDs. The influence of mobile phase on the separation of the NSAIDs enantiomers was investigated. The effect of organic solvents (ACN and MeOH) on retention time and resolution was also evaluated (Table 4.11). The R_s values for all selected NSAIDs were higher in ACN mobile phase. Compared to MeOH, ACN has greater solvent strength, therefore less retention were found at equivalent volume of mobile phase (50 %).

The effect of the amount of ACN on enantioseparation of selected NSAIDs was evaluated by varying the percentage of ACN in mobile phase (Table 4.11). The high k_1' and k_2' of NSAIDs at 90 % of ACN showed the normal phase behavior of the β -CD-BIMOTs-CSP. On the other hand, when at 30 % of ACN, the high k_1' and k_2' of NSAIDs showed the typical reverse phase behavior of β -CD-BIMOTs CSP. Therefore, the retention behavior of NSAIDs can be considered as the mixed reverse-normal separation mode (Guo *et al.*, 2009) similar with the retention behavior of β -blockers.

As given in Table 4.11, ibuprofen was completely resolved with R_s value of 2.51. Indoprofen showed partial separation with R_s value of 1.09. Ketoprofen and fenoprofen also partially enantioseparated with fenoprofen attained the lowest R_s value of 0.54. The high R_s values of ibuprofen and indoprofen are probably due to the *para* position of the substituent (containing the chiral center) on the aromatic ring. Previous study revealed that *para*-substituted aromatic rings can fit properly into the CD cavity (Fanali & Aturki, 1995) forming inclusion complex. However, the extent of the penetration mode is also depending on the polarity and feature structure of analytes (Nunez-Aguero *et al.*, 2006). Thus, this result showed that the hydrophobic ibuprofen achieved better enantioseparation than more polar indoprofen (Velkov *et al.*, 2007).

Meanwhile, the relatively low R_s values of ketoprofen and fenoprofen were because its substituent that located at *meta* position (Fanali & Aturki, 1995) that make their orientation in an unfavorable way to fit into the β -CD-BIMOTs cavity.

Table 4.11: Chiral separation data for the NSAIDs on β -CD-BIMOTs CSP

NSAID	Condition	k_1'	k_2'	α	R_s
Ibuprofen	ACN/water-90/10	0.29	1.17	4.04	2.51
	ACN/water-50/50	0.43	0.43	1.00	0
	ACN/water-30/70	1.23	1.23	1.00	0
	MeOH/water-90/10	0.16	0.16	1.00	0
	MeOH/water-50/50	0.77	0.77	1.00	0
Indoprofen	ACN/water-90/10	3.35	3.35	1.00	0
	ACN/water-50/50	0.15	0.51	3.39	1.09
	ACN/water-30/70	0.16	0.48	3.02	0.68
	MeOH/water-90/10	0.26	0.26	1.00	0
	MeOH/water-50/50	3.23	3.23	1.00	0
Ketoprofen	ACN/water-90/10	0.76	1.01	1.33	0.43
	ACN/water-50/50	0.46	0.94	2.06	0.72
	ACN/water-30/70	0.52	1.14	2.20	0.88
	MeOH/water-90/10	2.54	2.54	1.00	0
	MeOH/water-50/50	5.12	5.12	1.00	0
Fenoprofen	ACN/water-90/10	1.04	1.04	1.00	0
	ACN/water-50/50	0.07	0.07	1.00	0
	ACN/water-30/70	0.11	0.50	4.55	0.54
	MeOH/water-90/10	0.06	0.06	1.00	0
	MeOH/water-50/50	1.05	1.05	1.00	0

Even though the polarity of fenoprofen and ibuprofen are close to each other ($\log P_{\text{fenoprofen}}=3.8$, $\log P_{\text{ibuprofen}}=3.7$) (Velkov *et al.*, 2007), ibuprofen achieved higher R_s value at high organic solvent content (90 % ACN) mobile phase. This result suggested that ibuprofen can be fitted into β -CD-BIMOTs cavity whereas fenoprofen with two aromatic rings was less favorable to be fitted into β -CD-BIMOTs cavity due to steric hindrance effect. Previous simulation study (Nunez-Aguero *et al.*, 2006) showed the formation of moderate and weak hydrogen bonding between the carboxyl group of ibuprofen and hydroxyl groups of β -CD during complexation. Therefore, a part of inclusion complex formation, hydrogen bonding also plays a role to enhance the enantioseparation of NSAIDs. Additionally, ketoprofen which composed of almost similar structure (two aromatic rings) as fenoprofen achieved better enantioseparation than fenoprofen. This might due to the presence of carbonyl group in ketoprofen which enhanced the formation of hydrogen bonding with β -CD-BIMOTs rather than ether linkage in fenoprofen (Lommerse *et al.*, 1997).

In order to verify the interactions of enantioseparation, ^1H NMR and NOESY of β -CD-BIMOTs/NSAIDs complexes were studied. The values of chemical shifts (δ) obtained from ^1H NMR for different protons in β -CD-BIMOTs, NSAIDs and β -CD-BIMOTs/NSAIDs complexes are listed in Table 4.12 and 4.13. The deduced structures β -CD-BIMOTs/NSAID complexes are shown in Figure 4.21, respectively.

Table 4.12: Chemical shifts (δ) corresponding to β -CD-BIMOTs in the presence of NSAIDs

	β-CD-BIMOTs	β-CD-BIMOTs/ Ibuprofen	β-CD-BIMOTs/ Indoprofen	β-CD-BIMOTs/ Ketoprofen	β-CD-BIMOTs/ Fenoprofen				
	δ	δ	$\Delta\delta$	δ	$\Delta\delta$	δ	$\Delta\delta$	δ	$\Delta\delta$
H1	4.8405	4.8369	-0.0036	4.8316	-0.0089	4.8337	-0.0068	4.8280	-0.0125
H2	3.3312	3.3200	-0.0112	3.3474	0.0162	3.3015	-0.0297	3.3118	-0.0194
H3	3.6394	3.6387	-0.0007	3.6323	-0.0071	3.6284	-0.011	3.6326	-0.0068
H4	3.3716	3.4056	0.0340	3.4292	0.0576	3.3985	0.0269	3.4132	0.0416
H5	3.5777	3.5597	-0.018	3.5536	-0.0241	3.5458	-0.0319	3.5530	-0.0247
H6	3.9225	3.9091	-0.0134	3.9045	-0.018	3.9048	-0.0177	3.8803	-0.0422
H8	7.4215	7.4422	0.0207	7.4318	0.0103	7.4182	-0.0033	7.4209	-0.0006
H9	7.1112	7.1189	-0.0077	7.1268	0.0156	7.1196	-0.0084	-	-
H11	2.0847	-	-	-	-	-	-	-	-
Ha	7.4314	7.4877	0.0563	7.4835	0.0521	7.4737	0.0423	7.4834	0.052
Hb	7.7957	7.8149	0.0192	-	-	-	-	7.7896	-0.0061
Hc	7.7542	7.7516	-0.0026	-	-	-	-	7.7410	-0.0132
Hd	-	-	-	-	-	-	-	-	-
He	7.9563	7.9921	0.0358	-	-	7.9378	-0.0185	7.9399	-0.0164
Hf	9.2394	9.3362	0.0968	9.3202	0.0808	9.2240	-0.0154	9.3217	0.0823
Hg	5.4371	5.4514	0.0143	5.4146	-0.0225	5.4036	-0.0335	5.4459	-0.0088

 $\Delta\delta$: induced shifts

-: overlap peak

Table 4.13: Induced shifts ($\Delta\delta$) corresponding to NSAIDs in the presence of β -CD-BIMOTs

	β -CD-BIMOTs/ Ibuprofen	β -CD-BIMOTs/ Indoprofen	β -CD-BIMOTs/ Ketoprofen	β -CD-BIMOTs/ Fenoprofen
	$\Delta\delta$	$\Delta\delta$	$\Delta\delta$	$\Delta\delta$
Ha'	-0.0022	-0.0044	-0.0183	0.0132
Hb'	-0.0041	-0.0022	-0.0048	0.0133
Hc'	0.0072	-0.0044	-0.0083	0.0132
Hd'	-0.0030	-0.0141	-0.0070	0.0677
He'	-0.0033	-0.0051	-0.0119	0.0677
Hf'	-0.0023	-0.0081	-0.0083	0.0237
Hg'	-0.0011	-0.0081	-0.0052	0.0238
Hh'	-0.0020	-0.0086	-0.0046	0.0373
Hi'	-	-0.0086	-0.0042	0.0099
Hj'	-0.0029	-	0.0155	-
Hk'	-	-0.0235	-0.0098	0.0258

-: overlap peak

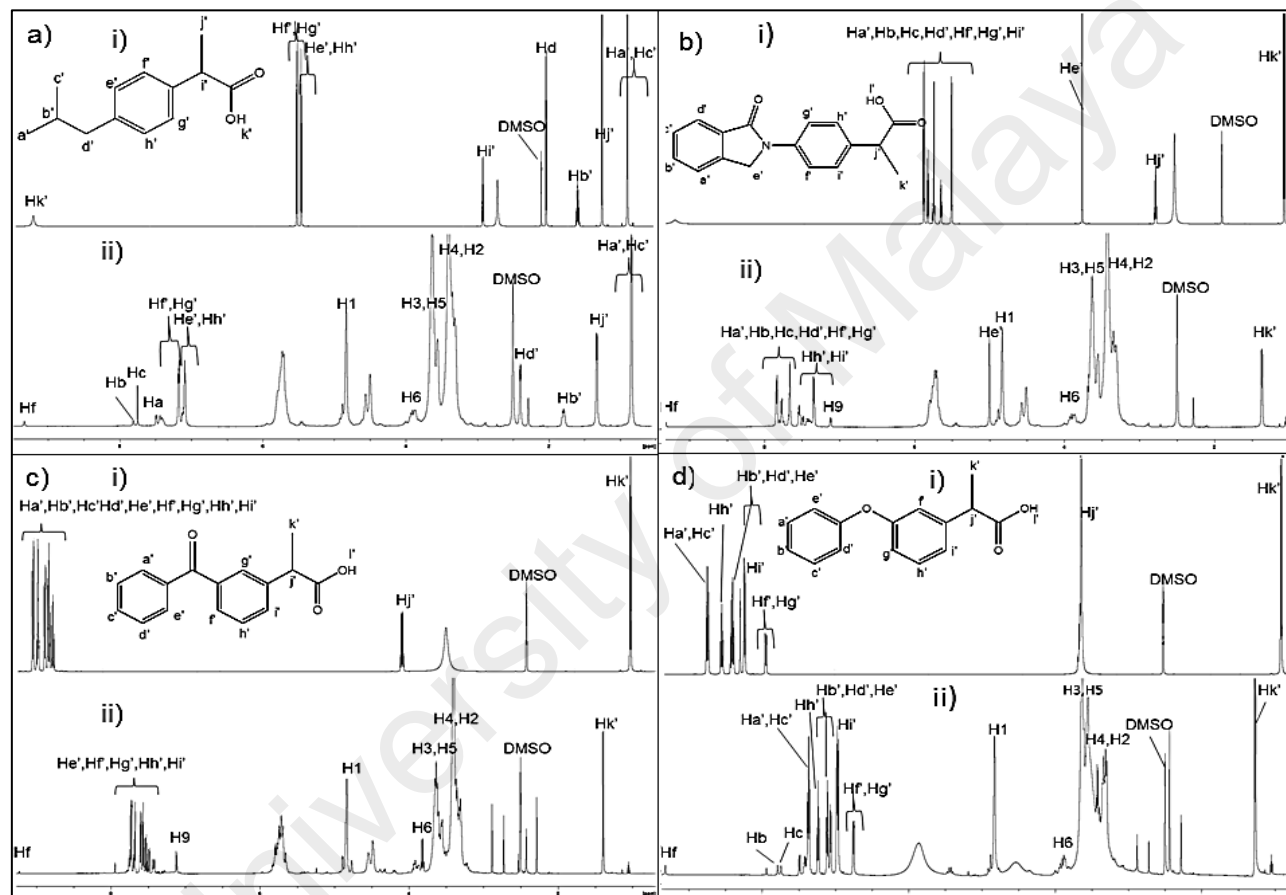


Figure 4.21: The deduced structure of NSAID/ β -CD-BIMOTs complexes: (a) i) ibuprofen ii) β -CD-BIMOTs/ibuprofen, (b) i) indoprofen ii) β -CD-BIMOTs/indoprofen (c) i) ketoprofen ii) β -CD-BIMOTs/ketoprofen, (d) i) fenoprofen ii) β -CD-BIMOTs/fenoprofen

The presence of ibuprofen, indoprofen, ketoprofen and fenoprofen was found to cause appreciable shift at H4 and H5 protons of β -CD-BIMOTs (Table 4.12) due to the formation of hydrogen bonding and inclusion complex, respectively. Significant change at Hc' proton of ibuprofen (Table 4.13) was observed. This result indicated that isobutyl moiety of ibuprofen was included into the cavity of β -CD-BIMOTs. However, the cross peak between proton of isobutyl ibuprofen with H5 proton of β -CD is absent in the NOESY spectra of β -CD-BIMOTs/ibuprofen (Figure 4.22). Perhaps, the great difference between isobutyl size and the internal β -CD diameter, (≈ 4.3 and 7.8 Å, respectively) is responsible for this weak interaction (Nunez-Aguero *et al.*, 2006). But, there were cross peak between Hf', Hg' and Hj' protons of ibuprofen with H5 proton of β -CD-BIMOTs confirmed the penetration aromatic moiety into the β -CD-BIMOTs cavity. The appreciable shift was also observed for the aromatic proton of indoprofen (Hd', Hh', Hi'), ketoprofen (Ha', He') and fenoprofen (Hd', He') (Table 4.13) as evidenced of inclusion complexes. This result was further strengthen with the NOESY spectra of β -CD-BIMOTs/indoprofen, β -CD-BIMOTs/ketoprofen and β -CD-BIMOTs/fenoprofen (Figure 4.23-4.25) showed the cross-peak between Hh', Hi' (proton indoprofen), He' (proton ketoprofen) and Ha', Hc', Hi' (proton fenoprofen) with H5 proton of β -CD-BIMOTs.

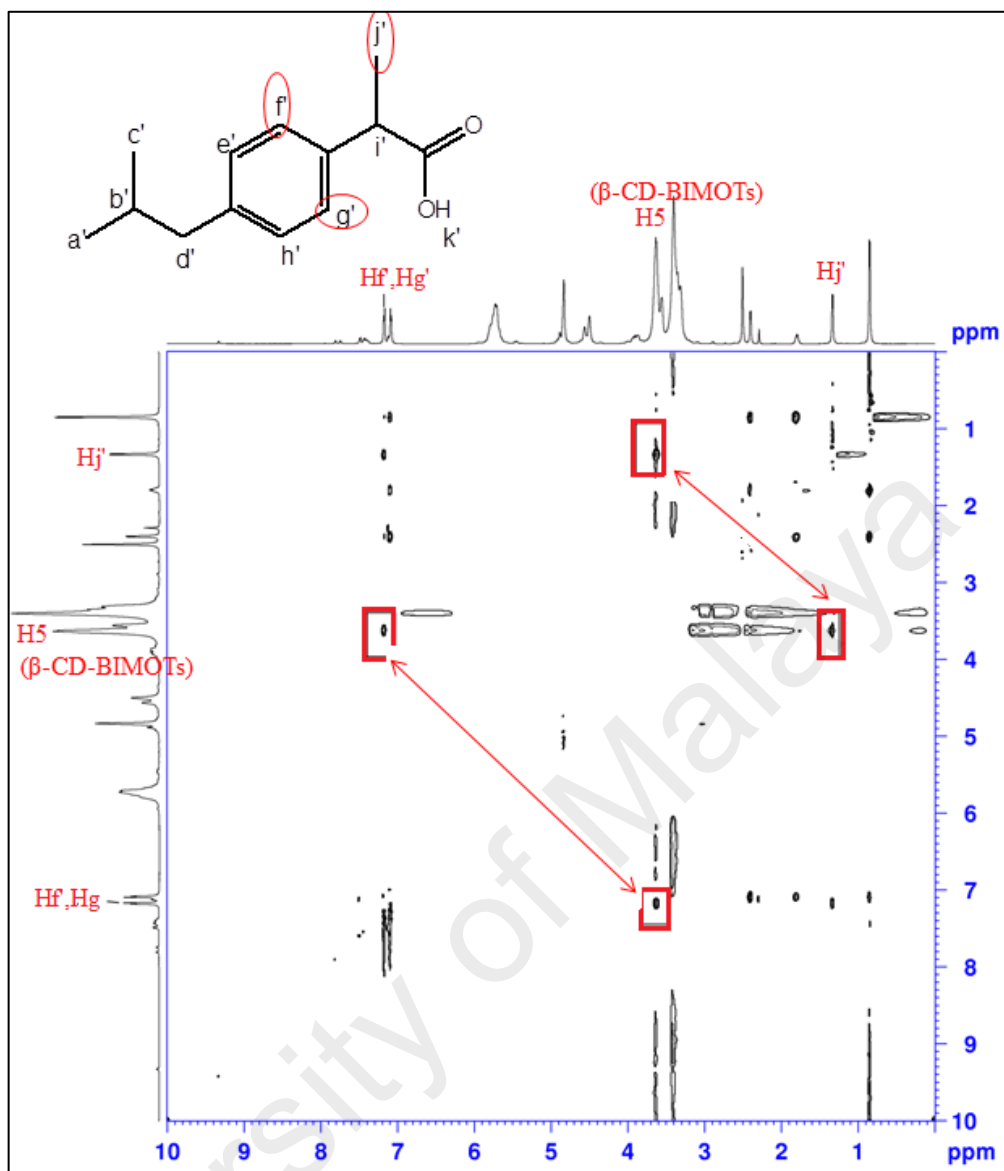


Figure 4.22: NOESY spectra of β -CD-BIMOTs/ibuprofen

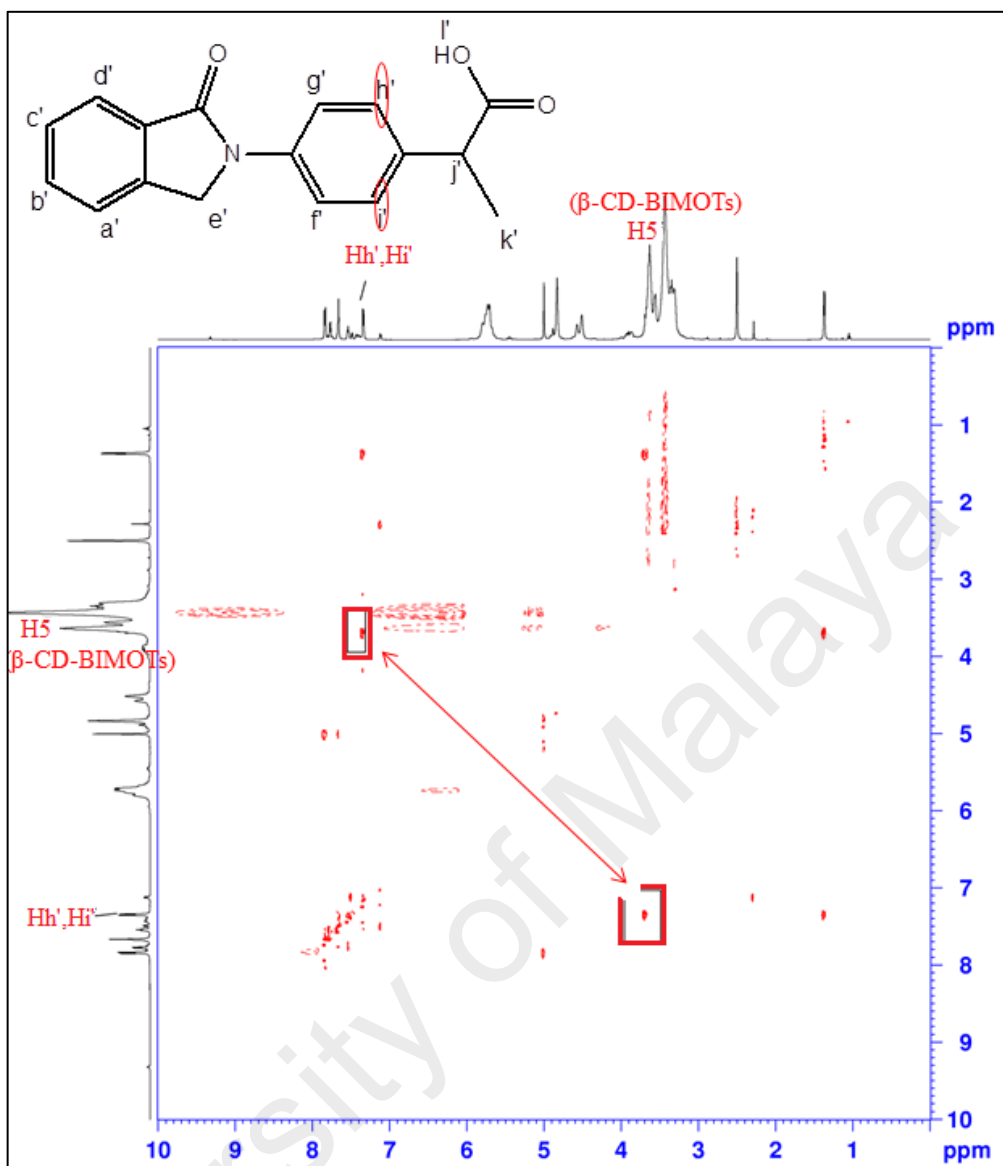


Figure 4.23: NOESY spectra of β -CD-BIMOTs/indoprofen

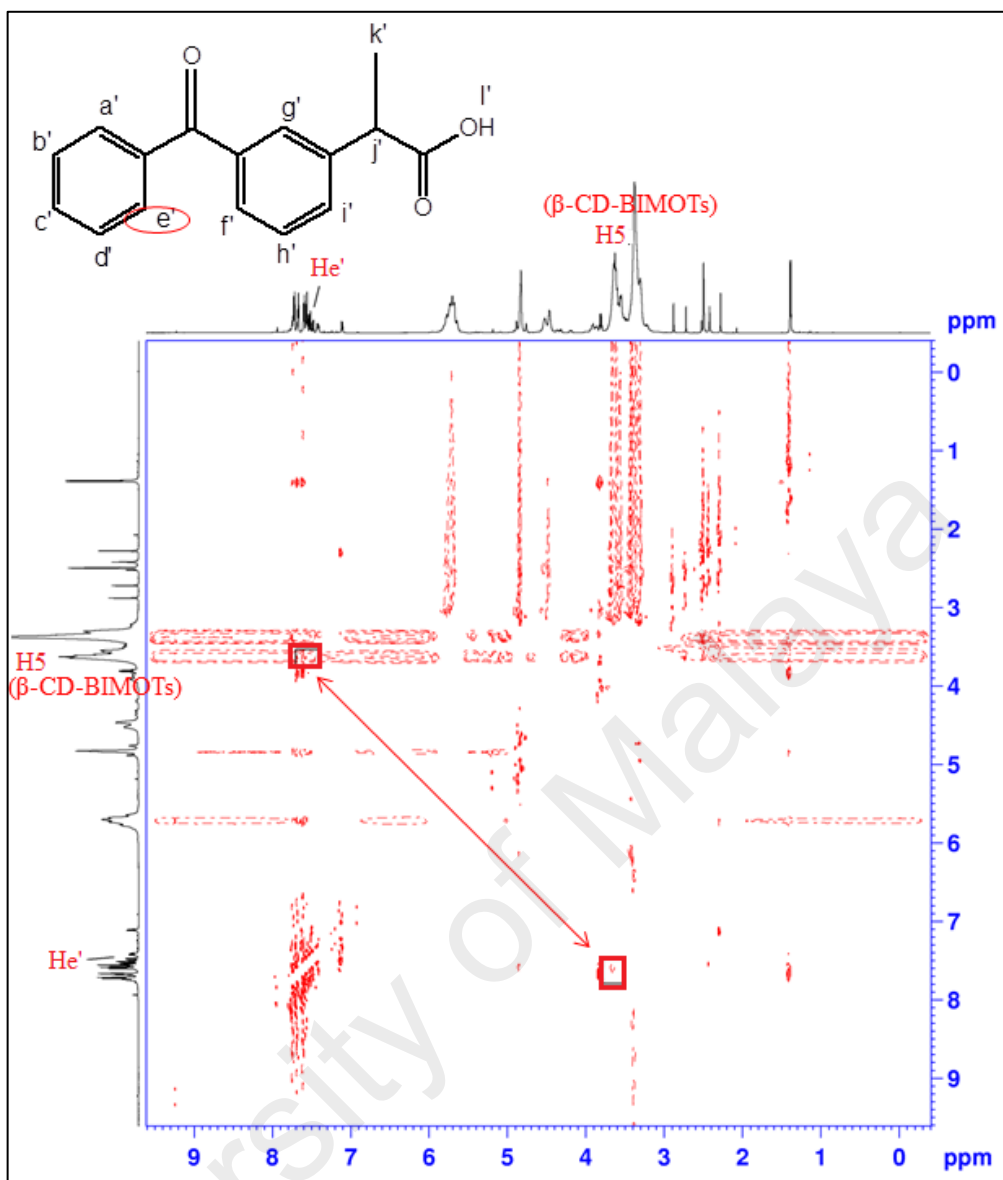


Figure 4.24: NOESY spectra of β -CD-BIMOTs/ketoprofen

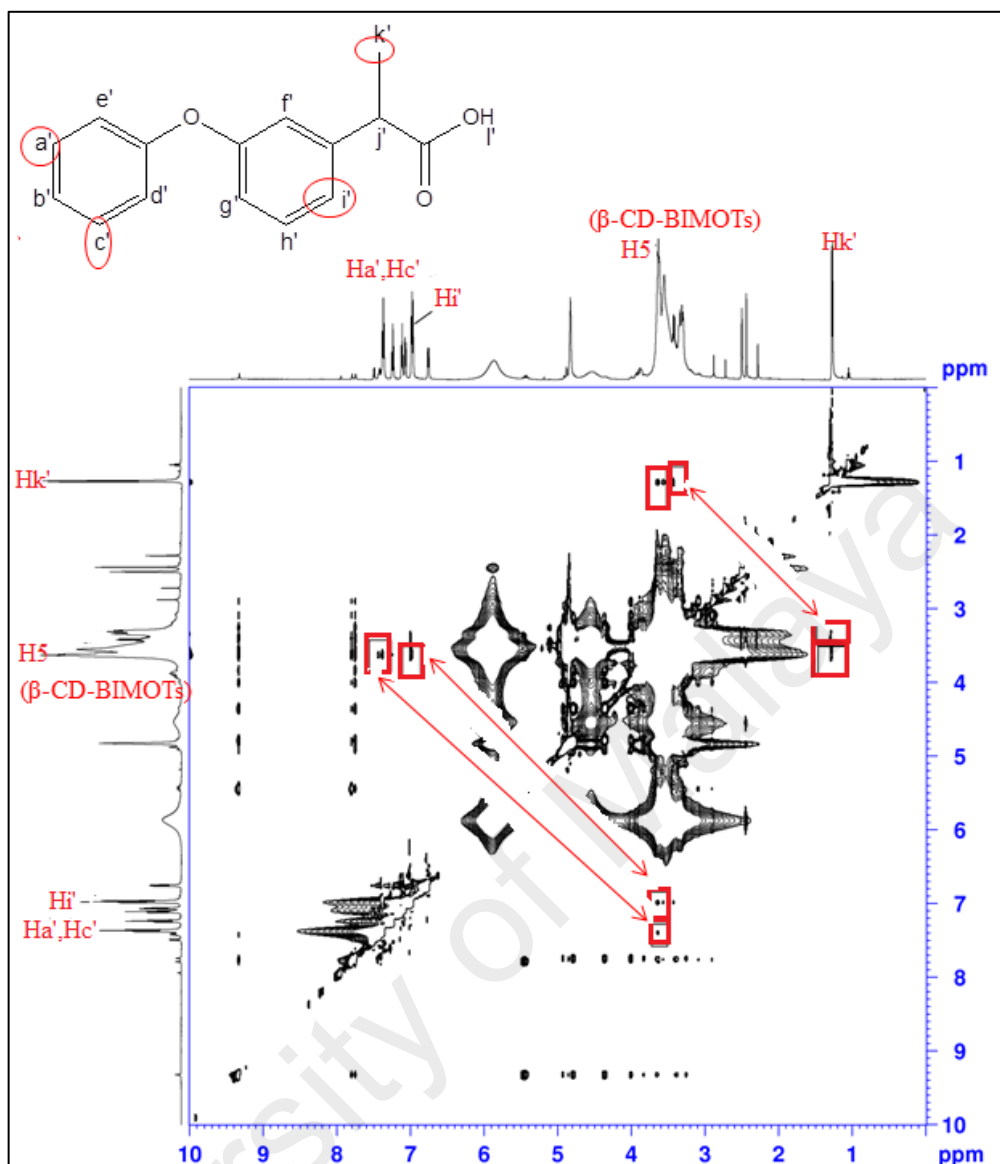


Figure 4.25: NOESY spectra of β -CD-BIMOTs/fenopropfen

The UV/Vis absorption spectra of β -CD-BIMOTs/NSAIDs complexes were further investigated to acquire more information on the interaction between NSAIDs and β -CD-BIMOTs. The plots of UV/Vis absorption for β -CD-BIMOTs, NSAIDs and β -CD-BIMOTs/NSAIDs complexes are presented in Figure 4.26. The results showed that β -CD-BIMOTs showed a λ_{\max} in the range of 230-260 nm. The λ_{\max} of β -CD-BIMOTs/ibuprofen, β -CD-BIMOTs/indoprofen and β -CD-BIMOTs/fenoprofen complexes appeared at 262, 256 and 256 nm, respectively referring to β -CD-BIMOTs. This absorbance undergoes the hyperchromic effect (increased of absorbance) and shifted bathochromically (change of absorbance to a lower frequency). Meanwhile, the absorbance of β -CD-BIMOTs/ketoprofen experienced the hypochromic effect (decreased of absorbance). The bathochromical shift is because of partial shielding of the chromophore electrons (Wang *et al.*, 2011a) in the β -CD-BIMOTs cavity. Both of hyperchromic and hypochromic effects was due to the π - π^* transition of dipole moments of aromatic ring. The transition dipole moment of this chromophore will interact with the induced dipoles of the neighboring chromophores, depending on their relative orientation. If the dipoles are along the same axis and one behind the other, then the intensity of the absorption band will be increased, and hyperchromic effect is observed. Conversely, if the dipoles are parallel and adjacent, a decrease in intensity of the absorption band occurs, and hypochromic effect is observed (Peral & Gallego, 2000). Moreover, hypochromic effect on β -CD-BIMOTs/ketoprofen also attribute by the limitation for π - π^* transition because of hydrogen bonding (Peral & Gallego, 2000) at carbonyl group between aromatic rings of ketoprofen. The variations that occur in the UV/Vis spectra are consequence of complexation of NSAIDs with β -CD-BIMOTs accompanied by π - π interaction and hydrogen bonding. These results proved the role of IL which provides π - π interaction which is the superposition of inclusion complex and hydrogen bond for the enantioseparation of NSAIDs.

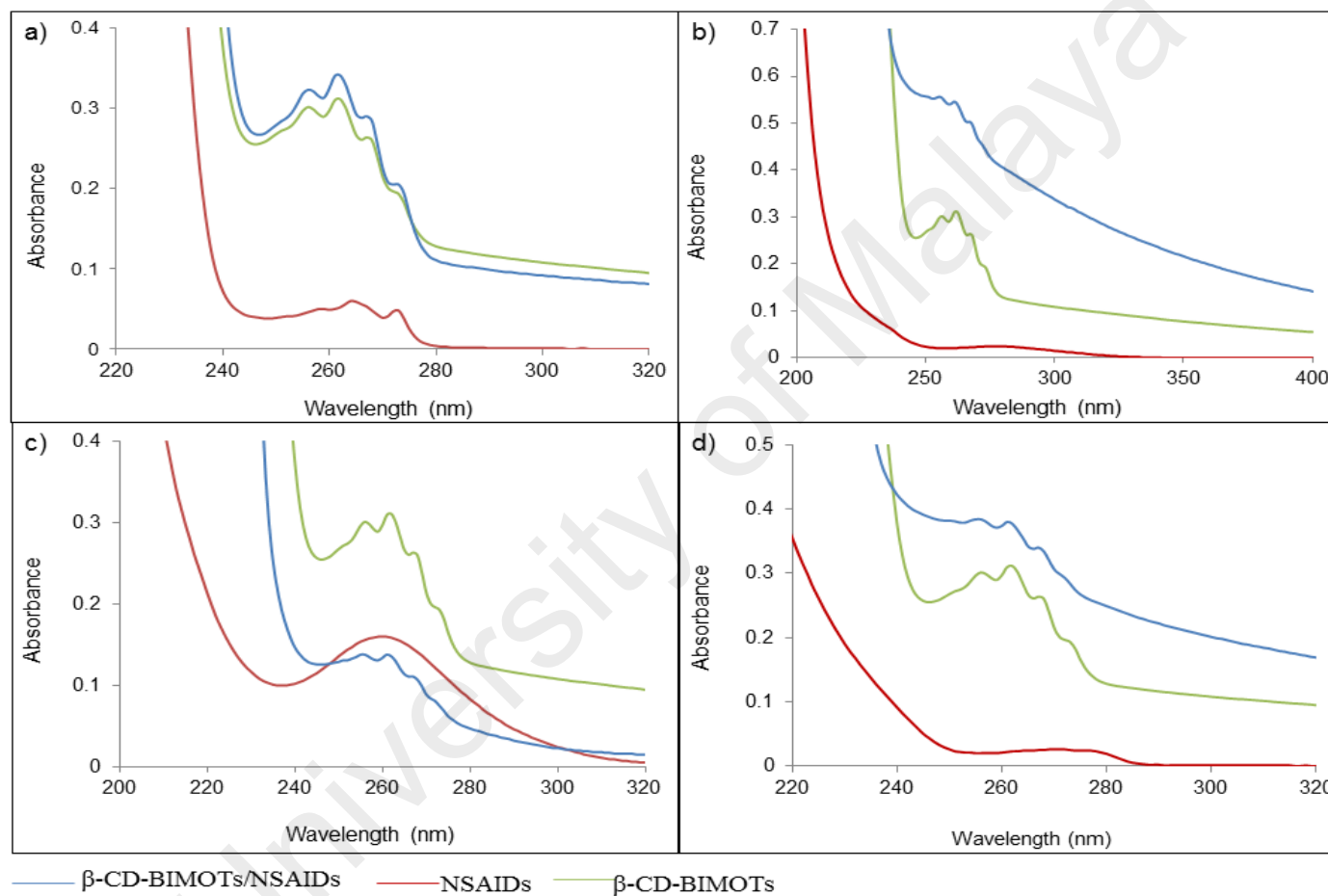


Figure 4.26: Absorption spectra of a) β -CD-BIMOTs/ibuprofen b) β -CD-BIMOTs/indoprofen c) β -CD-BIMOTs/ketoprofen d) β -CD-BIMOTs/fenoprofen with $[\beta$ -CD-BIMOTs]: 0.032mM [NSAIDs]: 0.01mM; T = 25 °C

CHAPTER 5: CONCLUSIONS AND FUTURE RECOMMENDATIONS

5.1 Conclusions

In this study, two new β -CD functionalized IL based CSPs (β -CD-BIMOTs and β -CD-DIMOTs) were successfully synthesized, characterized and compared their performance with native β -CD CSP. The β -CD-BIMOTs and β -CD-DIMOTs CSPs were characterized using various tools and the result obtained was compared with native β -CD CSP.

The performance evaluation of β -CD-BIMOTs, β -CD-DIMOTs and native β -CD as CSPs for the enantioseparation of neutral flavonoids, basic β -blockers and acidic NSAIDs groups was investigated. Although native β -CD has been reported as versatile and efficient for enantioseparation, however it is limited to certain class of analytes. The β -CD-BIMOTs herein have shown even greater chiral resolution capabilities. The result showed that the IL moieties substituted on the β -CD enhanced the enantioseparation. In contrast to the native β -CD CSP, the β -CD functionalized IL based CSP presents the variety interactions with the analytes. β -CD-BIMOTs CSP was more accessible and able to provide more interaction sites compare to β -CD-DIMOTs CSP.

Applying β -CD-BIMOTs as CSP, the influences of organic modifier and analytes's structure was investigated in detail. The following points can be summarized from the series of elaborate investigations of the CSP in reverse phase and polar organic mode HPLC.

- a) The number of OH group substituted at flavonoids strongly affected the choice of mobile phase mode and further affected the enantiomeric separation. In this dissertation, β -CD-BIMOTs CSP was well resolved the enantiomer of flavanone and partially resolved for hesperetin, naringenin and eriodictyol. The broader enantio-recognition abilities of β -CD-BIMOTs CSP

towards flavanone and hesperetin were attributable to the hydrophobic interaction, hydrogen bonding and π - π interaction. Meanwhile, the chiral recognition for naringenin and eriodictyol were attributed to the exterior interaction with β -CD-BIMOTs CSP such as hydrogen bonding and π - π interaction. Different interactions have been proposed to explain these diversities of inclusion complex for different types of flavonoids.

- b) The enantioseparation that attained for the basic β -blockers group is different from the neutral flavonoids group since the mixed mode reverse-normal mobile phase was observed rather than reverse phase. High polarity of atenolol and pindolol retaining them onto the stationary phase and inhibit the chiral recognition. Even though ion pairing reagent such as TEAA was used to accelerate the elution of polar analytes, but the chiral recognition was not improved. Propranolol and metoprolol obtained good enantioresolution as compared to atenolol and pindolol. This result suggested that the lipophilic property and the structure of propranolol and metoprolol enabled the formation of inclusion complex which contributed to better enantioseparation. This observation was proven by ^1H NMR and NOESY of β -CD-BIMOTs- β -blockers inclusion complexes. According to ^1H NMR and NOESY, propranolol and metoprolol showed the interaction at the interior torus of β -CD-BIMOTs which indicates the formation of inclusion complex. However, atenolol and pindolol showed the strong hydrogen bonding at exterior torus of β -CD-BIMOTs and causing the poor enantioseparation.
- c) The β -CD-BIMOTs CSP depicted good enantioseparation for most of NSAIDs. It was proven through ^1H NMR, NOESY and UV/Vis studied that all selected NSAIDs were enantioseparated due to the superposition of hydrogen bonding, inclusion complex and π - π interactions with β -CD-

BIMOTs CSP. Moreover, the extent of the inclusion mode was affected the enantioseparation. The inclusion mode depends on the polarity and feature structure of analytes. Ibuprofen and indoprofen achieved the good resolution because of the *para* position of the substituent (containing the chiral center) on the aromatic ring can fit properly into the β -CD cavity forming inclusion complex. Meanwhile, the relatively low R_s values of ketoprofen and fenoprofen was because of its substituent in the *meta* position that make their orientation in an unfavorable way to fit into the β -CD-BIMOTs cavity.

As a whole, the combine effect of hydrophobic inclusion complex, hydrogen bonding and π - π interaction resulted in improved the chiral selectivity. β -CD-BIMOTs which provide the additional interaction which is π - π interaction showed the important role of IL to enhance the enantioseparation of analytes.

5.2 Future work suggestions

In this study, β -CD-BIMOTs and β -CD-DIMOTs CSP have been applied in reverse phase and polar organic mobile phase. Chromatographic conditions have been optimized. The possible chiral recognition mechanisms have been investigated using qualitative tools such as NMR and UV/Visible. However, the influences of π - π interaction, hydrogen bonding and hydrophobic inclusion complexation on chiral separation are not quantitatively calculated. Molecular modeling may be useful addition information for theoretical understanding and prediction of the chiral separation mechanism. Only tosylate ion was chosen as the counterion in β -CD-BIMOTs and β -CD-DIMOTs CSPs. Investigations on chiral ionic liquid had revealed that anions may also affect enantioseparation processes. It will be interesting to change the counterions in the CSPs to investigate their influence on chiral resolution as well.

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LIST OF PUBLICATIONS AND PAPERS PRESENTED

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3. Rahim, Nurul Yani, Tay Kheng Soo, Sharifah Mohamad.” Chromatographic and spectroscopic studies on the β -cyclodextrin functionalized ionic liquid as chiral stationary phase: Enantioseparation of NSAIDs”. *Adsorption and Separation Technology*, DOI: 10.1177/0263617416686798.
4. Nurul Yani Rahim, Sharifah Mohamad, Tay Kheng Soo. 2013. Ionic cyclodextrins chemically-bonded chiral stationary phases for high-performance liquid chromatography. International Conference on Ionic Liquids 2013 (ICIL 13). 11-13 December 2013, Langkawi Island, Kedah.
5. Nurul Yani Rahim, Sharifah Mohamad, Tay Kheng Soo. 2014. Preparation of ionic liquid β -cyclodextrin immobilization on functionalized silica gel as chiral stationary phase for High Performance Liquid Chromatography. 6th International Conference on Postgraduate Education (ICPE-6). 17-18 December 2014, University Teknikal Malaysia Melaka.
6. Nurul Yani Rahim, Sharifah Mohamad, Tay Kheng Soo. 2015. 28th Regional Symposium of Malaysian Analytical Sciences. 17-20 August 2015, Weil Hotel, Ipoh, Perak.