Interchim manufacturing capabilities

Interchim : Core manufacturing competance and family values

Interchim has over 40 years of manufacturing experience.

Our competence has been built upon an unique technological understanding established through the product supply of many of the recognized manufacturers in the chromatographic field for two generations. The family ethos that runs through Interchim forms the cornerstone for the retention of core understanding of market requirement both within the company organization and through long standing partnerships.

Interchims proprietary range of chromatographic solutions are recognized as providing this sector with cutting edge technological advances relative to the current and progessive requirements of the analyst.

InterFine Chemicals, Chemistry Division :

- · Custom synthesis from mg to Kgs. Innovative and commercially known compounds
- · Building blocks & Intermediates, heterocycles, boronic acids, screening compounds
- Process optimization
- Scale up
- Flash Chromatography purification
- · Traditional techniques for purification
- · Custom made packaging for combi-chem

InterBiotec, BioScience Innovations Division :

- Antibody production & labelling
- Fluorescent & enzymatic labelling
- · Micro-plate custom coating service
- Specialty development kits
- Specialty biochemical reagents

InterBiochrom, Biochromatography Group :

- Antibody immobilization & purification
- Affinity & LC purification under client protocols
- HPLC & MPLC analysis

Interchrom, Chromatography Division :

- · HPLC & GC custom column Mfg.
- HPLC & GC development method & optimization
- Contract laboratory
- Buffer Mfg.
- Custom SPE column & plate Mfg.

Historical development of Interchim's chromatography technologies :

- 2007 Strategy[™] 1.7 µm and 2.2 µm Stationary phase ultra fast analysis & high throughput Modulo-cart[™] XS HPLC hardware technology for ultra high pressure
- 2006 Rodeo™ HPLC method optimization through column coupling technology
- 2005 Uptisphere[®] Strategy™ Latest generation of Silica 100 Å – 425 m²/g
- 2003 UptiBond[™] Premium Multi GC applications capillary columns Atoll[™] Polymeric sorbent for Solid Phase Extraction (SPE)
- 2002 Upti-Select kit HPLC method development tool (in collaboration with LETIAM-IUT laboratory in Orsay) Upti-Clean™ 1st ultra pure spherical based silica for SPE PuriFlash™ Flash Chromatography columns & cartridges
- 2000 Modulo-cart[™] Quick Seal State-of-the art HPLC hardware
- 1997 Upti-prep[®] Interchim's ultra pure based silica Uptisphere[®] Third generation HPLC columns & stationary phases
- **1995** Modulo-cart[™] (version 1)
- 1994 0.5 mm i.d micro columns (in collaboration with Rodhia's MAN service)
- 1992 Chirachrom[™] chiral columns (in collaboration with CNAM's general chemistry lab)
- 1985 HPLC Interchrom columns
- 1975 Glass & Stainless Steel GC packed columns

Biochromatogra









HPLC method development

Carotenoid isomers subcritical chromatographic test for C18 HPLC column classification

A test has been developed at LETIAM (IUT Orsav), within the analytical chemistry laboratory. Paris Sud, that discriminates more than ten groups of reverse stationary phases. from an analysis of carotenoid pigments. Stationary phase properties can then be broadly differentiated between each other. Major features can then be compared for over 160 commercial C18 silica based HPLC columns. From mainstay generation phases such as Licrosorb. Kromasil & Partisil to more current developments, such as Uptisphere and Zorbax.

The test demonstrates, in a single analysis, three primary features of the stationary phase.

- Hydrophobicity, overall phase retention capacity, reliant on bonding rate and specific surface area.
- Steric selectivity, the ability to separate molecules which carry differing 3D structures (isomers).
- Accessibility to residual silanols, relevant for basic compounds.

Contrary to chemometric approaches, this test presents, simply & concisely, through two successive diagrams (highlighted on the following page), the classification of stationary phases that exhibit similar chromatographic behavior relative to phases with different chemical properties.

This test is performed in subcritical fluid chromatography, a rapid and effective technique offering an inherent value. The important eluting force of the modified subcritical fluid (carbon dioxide & methanol mixture) and its weak viscosity, lead to a reduced molecular kinetic dispersion (narrow peaks) and rapid analysis from high mobile phase flow rate capabilities. Results obtained with this methodology are correlated with those obtained using hydro-organic mobile phases.

Test conditions :

1. Injection of a15 µl diluted mixture of partially isomerizated beta-carotene, containing all-trans, 9-cis, 13-cis (main isomer cis), 15-cis, and zeaxanthin (di-hydroxylated betacarotene)



2. The selected mobile phase is a mixture of methanol-carbon dioxide (15:85, v/v), at 25°C, outlet pressure 15MPa, flow rate: 3ml /min & detection wavelength at 440 nm.



HPLC method development

Results

Stationary phase classification is achieved in a two step process. Step one provides a primary classification through shape recognition and accessibility to polar sites, defining groups in which the stationary phase are placed. Step two examines the impact of hydrophobic characteristics, of a given phase, for a specified group.

Step 1:

Primary classification is established from the comparison of the selectivity value of shape recognition & polar accessibility, for each stationary phase.

Chart placement

a. X-axis

The selectivity, between the 13-cis isomer (bent form) and the all-trans (linear form) of the beta-carotene, is plotted on X-axis.

The cis/trans isomer separation of the beta-carotene is achieved through a 3D discrimination or shape recognition, by the stationary phases of the linear molecule vs the bent molecule. The variation in the tri-dimensional geometry of the two molecules directly results in differing interactions within the stationary phase.

A comparison of the results obtained through these studies, with other recognized tests of known stationary phases (such as NIST 869), provides a comprehensive organization of phases analyzed :

- Monomeric phases (mono-layer bonded silica) with low bonding density for cis/trans selectivity from 1 to 1.1
- Monomeric phases with high bonding density or partially polymeric (polymolecular layers bonded silica) from 1.1 to 1.2
- Polymeric phases above 1.2

Monomeric phases can be manufactured through bonding mono, di or tri-functional silanes upon a silica without water content. Polymeric phases are obtained from tri-functional silanes containing trace water. Typically, polymeric phases show a capacity for the separation of PAH (Polycyclic Aromatic Hydrocarbons) and rigid isomers of different stereochemistries.

b. Y-axis

Selectivity plotted on the Y-axis is given by the selectivity of all trans beta-carotene and zeaxanthin. Polar interactions are exhibited between hydroxyls of the zeaxanthin and the residual silanols of the stationary phase and /or additional accessible polar sites eg. (silane for polyfunctional phases, amide, urea, ether-oxide, quaternary ammonium or carbamates for polar embedded phases also called shielded). The higher the selectivity value, the lower this type of interaction is exhibited within the stationary phase. Therefore the less polar groups contained within the phase, the weaker the retention of zeaxanthin.

This selectivity ranges from 0.3 for phases exhibiting polar linked groups, to almost 20 for the most protected phases.

Polar site accessibility is significant from 0.3 to 1, moderate from 1 to 5, low from 5 to 10, and very low above 10. Increased selectivity is, in the majority of instances, relative to multiple end-capping treatments (single or combined), providing an indication of residual silanol coverage. Other non end-capped phases, with high bonding density, display high selectivity suggesting a reduced accessibility to the polar sites available on the stationary phase.







HPLC method development



Step 2 :

The first step groups columns that will achieve similar separations. Further classification for each group is achieved (4 5,7 & 8 are displayed) from further examination of silanol accessibility relative to hydrophobicity. For each group a separation will be similar but analysis times will differ. This provides a further indication for the optimization of a method, for a given separation.

Group 4





HPLC method development

Group 5



Group 7



Group 8





HPLC method development

Test consideration for polar linked alkyl & polar end-capped phases

Polar linked alkyl and polar end-capped phases are designed to analyze basic compounds. The polar linked alkyl phase achieves sharp peak definition whilst the polar endcapped phase can be used with a 100% aqueous mobile phase. A major difference between polar linked alkyl and hydrophilic end-capped phases is a weaker hydrophobicity of the polar embedded phases. The hydrophobicity of the hydrophilic end-capped or sulfonamide phases compares to classical C18.

In supercritical chromatography, the lack of water within the mobile phase allows for the polar linked groups and the hydrophilic end-capping to display strong interaction with hydroxyl groups of the zeaxanthin analyte. The strength of these interactions can therefore be evaluated.

Selectivity less than 1 on the X axis indicates an inversion of retention between zeaxanthin and the β -carotene, this is a direct result of strong interaction of the two hydroxyls contained within the zeaxanthin molecule. Amide chemistry display a stronger interaction and subsequent peak inversion than the interaction shown by carbamate functionality. These are subsequently stronger than phases carrying ether chemisty.

Polar linked phases interact with water of a hydro-organic mobile phase forming a hydrophilic barrier, stopping basic compounds accessing the residual silanols.

Step 2 representation of hydrophobicity /silanol accessibility places phases in distinct groups. Amide phases form a cluster (group A). Group B displays Carbamate phases. Group C classifies two polar end-capped and the sulfonamide phase. Finally, group D covers ether phase technology.

A few phases exhibit distinctive characteristics. Uptisphere® Strategy™ RP is one of these unique phases.





HPLC method development

Upti-select[™] Kit A tool to achieve consistent, reliable method development through a simple 2-step process

Interchim's C18 stationary phases have been developed to reflect the criteria outlined by the E. Lesellier et al. Tchapla test. Uptisphere[®] and Uptisphere[®] StrategyTM are the brand names for Interchim's proprietary HPLC technology, our stationary phases fall into strategic areas of the primary groups within the classification.

Place	Stationary Phase	Group	19.5				1
Place 20 27 51 64 116 138 155 156 157 158 159	Stationary Phase Uptisphere HDO Uptisphere NEC Uptisphere ODB Uptisphere HSC Uptisphere HSC Uptisphere FLP IUptisphere BioP II Uptisphere BioP I Uptisphere WOD Uptisphere WRP Strateov Pro	Group G10 G3 G4 G5 G13 G1 G4 G4 G4 G4 G4 G3	19,5 -		64 + + 162	1 G01	013
160 161 162	Strategy RP Strategy C18-2 Strategy C18-3	G10 G8 G5	9,5 -	019 20 4 50	64 51 • 157 • 156 158 • 155		
			4,5 -		103 +159 +27 21 		• ¹¹⁶
			-0.5	s 0.36	1,05	1.15	1,25

The Interchim Upti-selectTM kit contains a range of 5 proprietary columns, across the classification, for the initiation of column selection for analytical method development.

Upti-select[™] kit :

- Provides a method to predict and select the optimal column for analysis
- Eliminates column selection uncertainty
- Enhances the process toward column optimization
- Reduces development costs



HPLC method development

Step 1 :

Inject sample for analysis and the proprietary kit reference mixture onto the Uptisphere® 5 µm ODB packed column.

Determine :

- The retention time of a Reference Peak (RP) from the analysis sample.
- The retention time of the five probes from the Reference Mixture (RM) [calibrate the system]

Choose a retention factor for the reference peak (RP) : K'_RP

An Excel spread sheet indicates the percentage of methanol and /or acetonitrile required, within the mobile phase, to reach the expected retention factor K'_{RP} on each of the five columns.

Step 2 :

Prepare the mobile phase according to step 1. Inject the sample of interest onto the five columns (including ODB column for verification). Choose the column highlighting the most suitable separation. Start column optimization.

Upti-select[™] Kit and Upti-select[™] Kit Pack components

Upti-select[™] kit consists of 5 columns featuring identical dimensions and including all necessary hardware. i.e.

- A Reference column packed with the Uptisphere[®] 5 μm ODB
 Four additional columns with complementary selectivity based upon classification chart. e.g Uptisphere[®] 5 μm HDO Uptisphere[®] 5 μm NEC
 - Uptisphere® 5 µm HSC
 - Uptisphere[®] 5 µm TF
- Column connectors, (# CH953820)

Upti-select[™] kit Pack also includes:

- Proprietary Reference Mixture (RM) containing five probes for test calibration
- CD-ROM (includes : excel spread sheet for automated organic percentage calculation).
- Instruction booklet

Description		Sizes	P/N
Upti-select Kit [™] Upti-select Kit [™]		50 x 1.0 mm 50 x 2.0 mm 150 x 2.0 mm 250 x 2.0 mm 50 x 3.0 mm 150 x 3.0 mm 50 x 4.6 mm 150 x 4.6 mm 150 x 4.6 mm 250 x 4.6 mm	S64060 S57950 T07670 T80410 S64050 T77680 S57930 S64100 S57920 S57900
Upti-select Kit [™] Upti-select Kit [™]	Pack Pack Pack Pack Pack Pack Pack Pack	50 x 1.0 mm 50 x 2.0 mm 150 x 2.0 mm 250 x 2.0 mm 50 x 3.0 mm 150 x 3.0 mm 50 x 4.6 mm 100 x 4.6 mm 150 x 4.6 mm 250 x 4.6 mm	BB2280 BB2270 BB2260 BB2250 BB2240 BB2230 BB2210 BB2200 BB2200 BB2140
Reference Mixture (RM)	5 ml		BB2290
Upti-select Kit™ (includes : Reference Mix	ture (PM) + CD	Upgrade	BB2310



HPLC method development

Upti-select[™] Kit methodology example : Anti-UV agents

Step 1 :

a) The anti-UV agent sample contained the following components :

- A: 2-Ethylhexyl-2-Cyano-3.3-Diphenylacrylate
- B: 2-Ethylhexyl-3-[4-Methoxyphenyl]-2-Propenoate
- C: 4-t-Butyl-4'-Methoxy-Dibenzoyl-Methane
- D: 2-Ethylhexyl Salicylate

The sample was injected onto the Uptisphere® 5 μm ODB, 250 x 4.6 mm. Conditions : MeOH-H_2O (80/20) - 1ml/mn - 25°C - UV : 238 nm.

Peaks B and C co-eluted and the analysis time was more than 60 mn Peak D was thus chosen as the reference peak (RP)

b) The Reference Mixture (RM) was injected under the same conditions onto the same column.





The retention of peak D and the retention time of the five probes (RM) were introduced into the excel spread sheet.

The retention factor of peak D was fixed at 8.

The calculation table provided the percentage values of methanol and acetonitrile for each mobile phase for each column to achieve $K'_{RP'}$

Step 2 :

The mobile phase were prepared according to Step 1. The sample of interest was injected onto the 5 kit columns according to the fixed conditions to obtain K'_{RP} = 8 for peak D.



HPLC method development

Chromatograms w/Methanol



Chromatograms w/Acetonitrile



Observations

Results show K'_{RP} very close to 8 for each of the five Uptisphere[®] columns. [This verifies our log P calculation and column transposition equations are relevant]

- The A, B, C, D elution order is preserved on the HSC, ODB, HDO, and NEC columns.
- C and D are inverted on the TF column.
- A better separation is observed on Uptisphere® NEC and TF.
- The B,C pair achieve a better separation when the column features higher silanol accessibility.
- Co-elution occurs on the ODB, HDO, and NEC, columns.

Conclusion

Uptisphere® 5 μm TF and a methanol phase was identified as the most appropriate column for this application.

Final validated method Separation was optimized using Inter-

chim's Rodeo[™] concept. A running time of 6.88 mm was established for the last peak and a resolution

of 3.02 for the critical pair (B. D).





HPLC method development

Stationary Phase characterization for basic compound analysis

LGVP

The Laboratory of Pharmaceutical Analytical Chemistry – University of Geneva, Switzerland (LCAP) has established a global reputation for separative method development. Their mandate is to evaluate the fundamental aspects that influence the current drives in HPLC.

'Hot topics' such as monolithic sorbents, sub-2 micron particles and high temperature HPLC (T°> 60°C) are some of their recent considerations and further information on these studies are available upon request.

LCAP has developed and established chromatographic tests since 1993 to classify silica based HPLC columns for basic compound separation. Such separations have historically been a complicated process using traditional bonded silica based HPLC sorbent generally due to potential ion exchange interactions that can occur within the conventional pH range. For nearly two decades, manufacturers have attempted to overcome problems associateed with basic compound separation by offering new, chemically modified, chromatographic phases. LCAP's current evaluations revolve around the analysis of a mixture containing 7 basic complementary compounds, that allow the rapid discrimination of phase behavior in working conditions (hydro organic buffered mobile phases at pH 3 and 7).

LCAP basic compound test

Chemometric tools such as Principal Components Analysis (PCA) or Hierarchical Classification Ascendant (HCA), are used to visualize and subsequently classify column performance relative to similar behavior patterns. Column function is classified, within the LCAP test, by the hydrophobic power and the residual interaction with basic compounds. The classification obtained with the LCAP test is correlated with recognized tests published in the scientific literature. LCAP utilizes compounds similar in structure to those found in the pharmaceutical industry and applies mobile phase conditions used in this environment. This establishes the LCAP test as providing relevant information for column suitability for basic compound analysis and discriminates chromatographic sorbents with similar properties.

A number of stationary phases from different manufacturers were tested with the 7 compound test mix at pH 3 & 7.

Retention, efficiency, asymmetry values were measured and the phases studied were represented according to their similarity with PCA and HCA chemometric tools.

Phases were plotted, within a 2 dimensional setting, relative to residual silanol activity at pH: 7 for the principal components within the mix.

HCA computation subsequently groups stationary phases displaying similar chromatographic behavior whilst retaining the "non visible" information within the PCA representation.

Hierarchical Classification Ascendant (HCA)

Column	Bonding	Chemistry	Manufacturer
Zorbax	C18	Bidentate	Agilent
Discovery RP Amide	C16	Polar	Supelco
Nautilus	C18	Polar	Macherey Nagel
Supelcosil ABZ+	C16	Polar	Supelco
Stability BS C23	C16	Polar	CIL-Cluzeau
Performance	C18	Monolithic	Merck
SpeedRod	C18	Monolithic	Merck
Xterra	C18	Polar (Hybrid)	Waters
Nucleosil HD	C18	High density	Macherey Nagel
Eclipse XDB	C18	High density	Agilent
Luna	C18	High density	Phenomenex
Nucleodur	C18	C18 (ultra pure)	Macherey Nagel
Acclaim	C18	C18	Dionex
Nucléosil AB	C18	C18	Macherey Nagel
Nucléodur Pyramid	C18	C18 endcapped Hydr	Macherey Nagel
Symmetry Shield	C18	Polar	Waters
Uptisphere ODB	C18	C18 endcapped	Interchim
Uptisphere TF	C18	C18 endcapped polyf.	Interchim
Uptisphere HDO	C18	C18 mixed-endcapped	Interchim
Uptisphere HSC	C18	C18 endcapped	Interchim
Uptisphere NEC	C18	C18 non-endcapped	Interchim





HPLC method development



6 main components





Hydrophobic properties



6 main components



ACP 2D - AHC	
95 % variability	

B.13



Silanol activity

HPLC Method Optimization- RODEO™

Method Development - to - Method Optimization

Interchim's general guide to selecting the most appropriate stationary phase can be represented as follows :

- For samples primarily containing basic compounds : initiate evaluation using the 3 column Upti-select basic compound development kit.
- Sample containing a mixture of basic, neutral and acidic compounds.
 Intitiate testing with the broad application Uptisphere Strategy column (5 μm C18-2, 250 x 4.6 mm), conditions : ACN H₂O (70/30) & (30/70). If unsuccessful, apply Upti-select Kit[™] methodology with standardized phase diversity and/ or apply other selectivities such as cyano, amino, ion exchange.

The appropriate column can then be optimized. Interchim have established the RODEOTM concept highlighted below to address column optimization.

The RODEO[™] concept

The RODEO[™] concept provides a novel approach for HPLC method optimization. It takes advantage of the ability to effectively couple multiple Uptisphere HPLC columns together with couplers from the Uptisphere hardware range (see Uptisphere Modulo-Cart hardware details later in this HPLC section).

The chromatograms on the following page demonstrate, and additional studies have shown, that when working under the same analytical conditions and evaluating at the same stage of analysis, pressure (P), efficiency (N), asymmetry (As), resolution (Rs) and retention factor (k') are extremely similar between a Modulo-Cart Quick Seal 250mm column and five Modulo-Cart Quick Seal 50 mm columns coupled together irrespective of the column internal diameter.

RODEO[™] therefore provides a very simple procedure for the optimization of required column dimension, mobile phase, and appropriate temperature & is discussed further on the following pages.





[Modulo-Cart Quick seal technology exhibits 15% increased efficiency compared to other HPLC hardware. It is a real hand-tight fitting and the carbon doped PEEK on stainless steel guarantees long column lifetime].





HPLC Method Optimization - RODEO[™]

Exploring the difference between a 250 mm Modulo-cart column and five 50 mm columns coupled together with Modulo-cart couplers.



Dimension	to	k'	Р	As	N	Rs
250 mm	2.35	6.88	61	1.17	21148	8.58
5 x 50 mm	2.4	6.8	63	0.99	22574	8.56



