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Chemometric Curve Resolution for Quantitative Liquid Chromatographic Analysis

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Chemometric Curve Resolution for Quantitative Liquid Chromatographic Analysis

A dissertation submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy in Chemistry at Virginia Commonwealth University

By

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# List of Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>1D</td>
<td>first dimension</td>
</tr>
<tr>
<td>1D</td>
<td>one-dimensional</td>
</tr>
<tr>
<td>2D</td>
<td>second dimension</td>
</tr>
<tr>
<td>2D</td>
<td>two dimensional</td>
</tr>
<tr>
<td>2D-LC</td>
<td>two-dimensional liquid chromatography</td>
</tr>
<tr>
<td>2D-LC</td>
<td>two-dimensional assisted liquid chromatography</td>
</tr>
<tr>
<td>8-MOP</td>
<td>8-methoxypsoralen</td>
</tr>
<tr>
<td>ALS</td>
<td>alternating least squares</td>
</tr>
<tr>
<td>°C</td>
<td>degree Celsius</td>
</tr>
<tr>
<td>C18</td>
<td>octadecyl carbon chain</td>
</tr>
<tr>
<td>c2DALC</td>
<td>combined two-dimensional assisted liquid chromatography</td>
</tr>
<tr>
<td>CBP</td>
<td>4-chlorobenzophenone</td>
</tr>
<tr>
<td>COW</td>
<td>correlation optimized warping</td>
</tr>
<tr>
<td>DAD</td>
<td>diode array detector</td>
</tr>
<tr>
<td>EFA</td>
<td>evolving factor analysis</td>
</tr>
<tr>
<td>EMG</td>
<td>exponentially modified Gaussian</td>
</tr>
<tr>
<td>GRAM</td>
<td>generalized rank annihilation method</td>
</tr>
<tr>
<td>HILIC</td>
<td>hydrophilic interaction liquid chromatography</td>
</tr>
<tr>
<td>IEC</td>
<td>ion exchange chromatography</td>
</tr>
<tr>
<td>IKSFA</td>
<td>iterative key set factor analysis</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>------------------------------------------------------------</td>
</tr>
<tr>
<td>IOPA</td>
<td>iterative orthogonal projection approach</td>
</tr>
<tr>
<td>HTLC</td>
<td>high temperature liquid chromatography</td>
</tr>
<tr>
<td>HPLC</td>
<td>high performance liquid chromatography</td>
</tr>
<tr>
<td>HSM</td>
<td>hydrophobic subtraction model</td>
</tr>
<tr>
<td>LC</td>
<td>liquid chromatography (used interchangeably with HPLC)</td>
</tr>
<tr>
<td>LC x LC</td>
<td>comprehensive two-dimensional liquid chromatography</td>
</tr>
<tr>
<td>LCCC</td>
<td>liquid chromatography under critical conditions</td>
</tr>
<tr>
<td>Mamp</td>
<td>methamphetamine</td>
</tr>
<tr>
<td>MCR</td>
<td>multivariate curve resolution</td>
</tr>
<tr>
<td>MDMA</td>
<td>methylenedioxymethamphetamine</td>
</tr>
<tr>
<td>MHC</td>
<td>multiple heartcutting two-dimensional liquid chromatography</td>
</tr>
<tr>
<td>MoxyAmp</td>
<td>methoxyamphetamine</td>
</tr>
<tr>
<td>MS</td>
<td>mass spectrometer or mass spectrometry</td>
</tr>
<tr>
<td>MS/MS</td>
<td>tandem mass spectrometry</td>
</tr>
<tr>
<td>OBGC</td>
<td>orthogonal background correction</td>
</tr>
<tr>
<td>OPA</td>
<td>orthogonal projection approach</td>
</tr>
<tr>
<td>PARAFAC</td>
<td>parallel factor analysis</td>
</tr>
<tr>
<td>PARAFAC2</td>
<td>parallel factor analysis version 2</td>
</tr>
<tr>
<td>PCA</td>
<td>principal components analysis</td>
</tr>
<tr>
<td>PLS</td>
<td>partial least squares</td>
</tr>
<tr>
<td>QuEChERS</td>
<td>quick, easy, cheap, effective, rugged, safe (acronym for extraction method)</td>
</tr>
<tr>
<td>RMSPE</td>
<td>root mean square percent error</td>
</tr>
<tr>
<td>RP</td>
<td>reversed phase liquid chromatography</td>
</tr>
<tr>
<td>%RSD</td>
<td>percent relative standard deviation</td>
</tr>
</tbody>
</table>
SE  standard error
SEC  size exclusion chromatography
SIMPLISMA  simple-to-use interactive self-modeling mixture analysis
sLC x LC  selective comprehensive two-dimensional liquid chromatography
UHPLC  ultra-high performance liquid chromatography
UV  ultraviolet light
Vis  visible light
List of Variables

\( \alpha \)  
chromatographic selectivity

\(<\beta>\)  
broadening correction factor

\( \lambda \)  
wavelength

\( \sigma \)  
standard deviation

\( \theta \)  
angle

\( A \)  
coefficient representing eddy diffusion in the van Deemter equation

\( A \)  
matrix representing sample profiles in PARAFAC

\( B \)  
coefficient representing longitudinal diffusion in the van Deemter equation

\( C \)  
coefficient representing resistance to mass transfer in the van Deemter equation
OR concentration

\( C \)  
matrix representing chromatographic profiles in MCR-ALS and PARAFAC

\( E \)  
residual error after MCR-ALS or PARAFAC fitting

\( f_{\text{coverage}} \)  
fractional coverage of the two dimensional separation space

\( H \)  
height equivalent of a theoretical plate

\( I \)  
number of data points in the chromatographic dimension

\( J \)  
number of data points in the spectral dimension

\( K \)  
number of samples represented in a data array

\( k \)  
retention factor

\( L \)  
column length
$m$  
model coefficients

$N$  
chromatographic efficiency

$n_c$  
peak capacity

$n_{mob}$  
number of moles in the mobile phase

$n_{stat}$  
number of moles in the stationary phase

$r^2$  
coefficient of determination

$^{1}R_S$  
first dimension chromatographic resolution

$^{2}R_S$  
second dimensional chromatographic resolution

$R_S$  
chromatographic resolution

$R_s$  
threshold chromatographic resolution needed for sufficient calibration with MCR-ALS

$S$  
matrix representing spectral profiles in MCR-ALS and PARAFAC

$S/N$  
signal-to-noise ratio

$t_M$  
dead time

$t_R$  
retention time

$t_S$  
sampling time

$w$  
peak width at 4σ

$X$  
raw data array used in MCR-ALS and PARAFAC

$x$  
factors used for building a model

$y$  
true concentration

$\hat{y}$  
predicted concentration
Abstract

CHEMOMETRIC CURVE RESOLUTION FOR QUANTITATIVE LIQUID CHROMATOGRAPHIC ANALYSIS

By Daniel Wesley Cook

A dissertation submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy at Virginia Commonwealth University

Virginia Commonwealth University, 2016

Director: Sarah C. Rutan, Professor, Department of Chemistry

In chemical analyses, it is crucial to distinguish between chemical species. This is often accomplished via chromatographic separations. These separations are often pushed to their limits in terms of the number of analytes that can be sufficiently resolved from one another, particularly when a quantitative analysis of these compounds is needed. Very often, complicated methods or new technology is required to provide adequate separation of samples arising from a variety of fields such as metabolomics, environmental science, food analysis, etc.

An often overlooked means for improving analysis is the use of chemometric data analysis techniques. Particularly, the use of chemometric curve resolution techniques can mathematically resolve analyte signals that may be overlapped in the instrumental data. The use
of chemometric techniques facilitates quantitation, pattern recognition, or any other desired analyses. Unfortunately, these methods have seen little use outside of traditionally chemometrics focused research groups. In this dissertation, we attempt to show the utility of one of these methods, multivariate curve resolution-alternating least squares (MCR-ALS), to liquid chromatography as well as its application to more advanced separation techniques.

First, a general characterization of the performance of MCR-ALS for the analysis of liquid chromatography-diode array detection (LC-DAD) data is accomplished. It is shown that under a wide range of conditions (low chromatographic resolution, low signal-to-noise, and high similarity between analyte spectra), MCR-ALS is able to increase the number of quantitatively analyzable peaks. This increase is up to five-fold in many cases.

Second, a novel methodology for MCR-ALS analysis of comprehensive two-dimensional liquid chromatography (LC x LC) is described. This method, called two dimensional assisted liquid chromatography (2DALC), aims to improve quantitation in LC x LC by combining the advantages of both one-dimensional and two dimensional chromatographic data. We show that 2DALC can provide superior quantitation to both LC x LC and one dimensional LC under certain conditions.

Finally, we apply MCR-ALS to an LC x LC analysis of fourteen furanocoumarins in three apiaceous vegetables. The optimal implementation of MCR-ALS and subsequent integration was determined. For these data, simply performing MCR-ALS on the two dimensional chromatogram and manually integrating the results proved to be the superior method. These results demonstrate the usefulness of these curve resolution techniques as a compliment to advanced chromatographic techniques.
Chapter 1: Overview of Objectives

With the ever increasing need to analyze complex chemical samples, analysis methods must constantly evolve. These complex analyte mixtures arise from a wide range of fields such as food science, environmental science, metabolomics [1], proteomics [2,3], and many others. These fields can produce samples with well over 1,000 analytes with multiple classes of analytes present [4–6]. Commonly, innovations in instrumental technology and instrumental methods drive the field of analytical separations; however, innovations in data analysis techniques can also offer powerful tools to complement existing instrumental methods.

Liquid chromatography (LC) has advanced greatly in the few decades since its inception. Both advances in the theoretical understanding and practical innovations have enabled its widespread adoption and today it is one of the most widely used techniques for chemical analysis. Chapter 2 presents a brief overview of the fundamentals of liquid chromatography and describes comprehensive two-dimensional liquid chromatography (LC x LC) [7,8], a particularly promising innovation in the field of separation science. LC x LC allows for a much greater number of analytes to be separated due its use of two coupled chromatographic separations with different selectivities. Even with advanced LC instrumentation and methods, the number of analytes able to be separated in a single analysis is finite and overlap of analyte signals is still common, particularly when short analysis times are desired and/or complex samples are being
analyzed. These peak overlaps along with other instrumental effects can degrade the quantitative performance of these methods [9].

The objective of the research described in the following chapters was to utilize chemometric techniques to improve quantitative liquid chromatographic analyses by extracting underlying quantitative information from data that may be corrupted by background, noise, interfering species, and other instrumental effects. The chemometric curve resolution techniques discussed in the following chapters mathematically resolve analyte signals from one another and from background and noise by analyzing the data holistically. Most traditional data analysis techniques rely on single-channel detection (i.e., a single wavelength or mass-to-charge value), even if multiple channels of data are collected from the instrument. Data from multichannel detectors are often visualized and analyzed at a single wavelength in the case of ultraviolet-visible (UV-Vis) detection or an extracted ion chromatogram in the case of mass spectrometric (MS) detection. Chemometric curve resolution techniques use the complete data by treating them as higher order data arrays, making use of the full spectral dimension in the data (e.g., ultraviolet visible or mass spectra) [10]. Descriptions of these curve resolution methods and other chemometric treatments of chromatographic data are presented in Chapter 3.

Three major goals guide the work described in the following chapters. First, many curve resolution techniques have been used in the literature without a detailed study on the abilities and limitations of these methods. We aim to characterize one such technique called multivariate curve resolution-alternating least squares (MCR-ALS) [11,12]. While used extensively in the literature, MCR-ALS has yet to find widespread use in routine analyses outside of traditionally chemometric research laboratories. This is possibly due to the misperception that it is difficult to implement and does not provide a significant advantage for chromatographic analyses. In
Chapter 4 it is demonstrated that MCR-ALS does provide a clear advantage and can produce up to a five-fold increase in effective peak capacity, a measure of the maximum number of analyzable peaks in a given separation. This is demonstrated over a range of conditions using a design of experiments approach allowing for the generation of an approximate model of the quantitative performance of MCR-ALS.

The second goal of this work was to investigate the use of MCR-ALS to improve the quantitative abilities of LC x LC. LC x LC aims to resolve analyte signals by adding a second dimension of chromatography. While this can provide significantly higher peak capacities, it can come at the cost of quantitative performance. Thus far in the literature the quantitative performance of LC x LC has typically been inferior to that of traditional one-dimensional (1D) chromatography. This is attributed to effects introduced during the transfer of the first dimension (1D) effluent to the second dimension (2D) of separation. Therefore, quantitative information is preserved in the 1D separation. The work described in Chapter 5 aims to extract the quantitative information of the 1D separation with the assistance of the 2D separation. This is done by utilizing the greater separation of peaks in the 2D to improve MCR-ALS analysis of the 1D separation, containing severely overlapped chromatographic peaks. This approach is named two-dimensional assisted liquid chromatography (2DALC).

Finally, our third goal was to demonstrate the use of MCR-ALS in a relevant, real-world LC x LC analysis. Chapter 6 describes the analysis of furanocoumarins from apiaceous vegetables with LC x LC and MCR-ALS. Furanocoumarins are a class of compounds of great interest due to their high bioactivity including interactions with the liver enzymes responsible for the metabolism of many pharmaceuticals[13,14]. In order to investigate the physiological effects from the consumption of these vegetables, it is crucial to determine the levels at which the
compounds are present within certain vegetables. To determine the best method for obtaining the concentrations of these compounds in vegetable samples, three implementations of MCR-ALS were investigated, as well as two strategies for the subsequent peak integration step. It was found that for this data set, LC x LC quantitation was competitive with that of one-dimensional LC when MCR-ALS was used. It was also found that, while tedious, manual integration of the resolved chromatographic peaks yielded superior quantitative results.

Through the work described in the following chapters, it is clear that MCR-ALS has great potential for improving quantitative liquid chromatographic performance. Through the ability to handle peak overlap, MCR-ALS can enable the use of shorter analysis times of more complex samples; however, rather than MCR-ALS being considered an alternative to improved separations, MCR-ALS should be thought of as a complementary technique that allows good separations to be made even better. This is shown by its applicability to LC x LC where MCR-ALS can assist in the quantitation of analytes while simultaneously increasing peak capacity even further than with LC x LC alone. Chapter 7 draws conclusions from the work presented here and provides future directions for this work.
Chapter 2: Liquid Chromatography

2.1. Fundamentals of Liquid Chromatography

Liquid chromatography (LC) is one of the most widely used methods of chemical analysis. LC separates analytes based on their interactions with a mobile phase and a stationary phase, either through partitioning or adsorption. These interactions differentially retard analytes giving rise to separation. These interactions are dictated by three main properties of the molecules: electrical charge, molecular size, or polarity [15]. The discussion below will focus on polarity; however, many of the concepts apply equally well to electrical charge. Also in this discussion and the following chapters, high performance or high pressure liquid chromatography (HPLC) will be used interchangeably with LC as it will be the focus of the work presented here. HPLC, rather than gravity-fed chromatography, forces the mobile phase through the stationary phase at higher pressure [16].

Separation in LC is driven by the differential retention of each analyte on the stationary phase. The extent of this retention can be quantified by a metric called the retention factor, $k$. This value is equal to the ratio of moles of analyte in the stationary phase, $n_{\text{stat}}$, to the moles of analyte in the mobile phase, $n_{\text{mob}}$, as shown in Eq. 2.1. Experimentally, it can be calculated based on the time it takes for the analyte to elute from the column, also known as retention time ($t_R$),
and the time it takes for an unretained compound to move through the column, also known as dead time ($t_M$).

$$k = \frac{n_{\text{stat}}}{n_{\text{mob}}} = \frac{t_R - t_M}{t_M}$$ (2.1)

For two analytes to be separated in a given analysis, they must have sufficiently different values of $k$. This difference is captured in a metric known as selectivity ($\alpha$), defined as the ratio of the $k$ values of two analytes as shown in Eq. 2.2. Selectivity is a key term in the calculation of chromatographic resolution ($R_s$), which is the most common measure of the separation of two compounds. It incorporates the efficiency ($N$) of the separation and the selectivity, both assuming a Gaussian peak shape [17]. Importantly, $R_s$ can be measured simply from a chromatogram using the retention times ($t_R$) and peak widths ($w$), where $w$ is considered the peak width at four times the standard deviation of the peak ($\sigma$).

$$\alpha = \frac{k_2}{k_1}$$ (2.2)

$$N = 16 \left( \frac{t_R}{w} \right)^2 = \left( \frac{t_R}{\sigma} \right)^2$$ (2.3)

$$R_s = \frac{1}{4} \sqrt{N} \left( \frac{\alpha - 1}{\alpha} \right) \frac{k}{k + 1} = \frac{2(t_{r,2} - t_{r,1})}{w_1 + w_2}$$ (2.4)

The relative importance of each factor in the value of $R_s$ is shown in Fig. 2.1. It can be seen that $\alpha$ is has a significant effect on the peak resolution, particularly at greater than $R_s = 0.5$; however, it should be noted that for a resolution greater than 1.5, the peaks are resolved to the baseline, meaning no further quantitative advantage is gained by an increased resolution.
When trying to increase the $R_s$ between two peaks, that is, to decrease the overlap of the peaks, the peaks can either be made to elute further apart, increasing $\alpha$, or the width of the peaks can be decreased, achieved by increasing $N$. In real-world chromatography, analysis time matters. Increasing $\alpha$ can be achieved by increasing retention, thereby increasing the time it takes for analytes to move through the column; however, at a certain point this becomes impractical due to longer separation times. The more popular approach is to increase $N$. From Eq. 2.3, it can be seem that the practical implication of increasing $N$ is to decrease the peak width ($w$). Decreasing $w$ allows for a greater number of peaks to be resolved in a shorter amount of time. This concept is known as peak capacity ($n_c$) and is defined as the maximum number of analyzable peaks (usually defined at $R_s = 1$) in a separation [20]. At an $R_s$ of less than one, it becomes difficult to differentiate peaks and quantitation greatly degrades due to the inability to integrate the individual peaks. Eq. 2.5 shows the formula for $n_c$ when using LC with a mobile phase gradient [20] where $t_{\text{gradient}}$ is defined as the time from the beginning to the end of the mobile phase
gradient. This equation assumes a random distribution of peaks from the beginning to the end of the mobile phase gradient.

\[ n_c = \frac{t_{\text{gradient}}}{w} = \frac{t_{\text{gradient}}}{4\sigma} \]  

(2.5)

The factors affecting \( N \) are explained using the common van Deemter equation. This simple equation (Eq. 2.6) equates the height equivalent of a theoretical plate (\( H \)), defined as the length of the column (\( L \)) divided by \( N \), to three phenomena in chromatography. First, the \( A \) term describes the eddy diffusion through the column; in other words, the effect of many different paths an analyte can take through the column. The more different paths that the analyte can travel, the higher the value of \( H \), which leads to a broader chromatographic peak. Second, the \( B \) term describes the longitudinal diffusion occurring in the column. This is inversely proportional to the linear velocity (\( u \)), which is related to flow rate; the slower the analyte moves, the more time it has to diffuse into a wider analyte band, leading to a broader peak. Finally, the \( C \) term describes the resistance to mass transfer of the analyte into the stationary phase. Figure 2.2 graphically depicts the van Deemter equation along with the influence of each term on the separation. The minimum value of \( H \) on the black curve in Fig. 2.2 is considered the optimum linear velocity. Often, however, this optimum velocity is slower than desired for quick analyses times and thus a tradeoff is made between linear velocity and minimal plate height. The penalty for using a linear velocity greater than the optimum velocity is determined by the magnitude of the \( C \) term, as shown by the blue line in Fig. 2.2.
Figure 2.2. Van Deemter plot and individual term contributions to the van Deemter curve (black). $H_{\text{min}}$ and $u_{\text{opt}}$ represent the minimum plate height at the optimum linear velocity.

$$H = A + \frac{B}{u} + Cu$$  \hspace{1cm} (2.6)$$

$$H = \frac{L}{N}$$  \hspace{1cm} (2.7)

The major contributing factor to all three terms in the van Deemter equation is the column. Column manufacturers are constantly innovating to create new packing materials in order to increase the efficiency of the columns. Traditionally, the silica particles used for packing are fully porous particles; recently, however, superficially porous, or core shell, particles are becoming very popular [21]. These particles consist of a solid core with a porous outer shell. The main implications of this are a decrease in the $C$ term (i.e., increasing the speed of mass transfer) and a decrease in the $A$ term due to improved column packing [22]. A decrease in the $C$ term lowers the slope of the blue curve in Fig. 2.2, which allows for an increase in linear velocity with lesser effects on $H$. Monolithic columns, consisting of a solid rod of porous silica or other material, also have this same advantage [23]. Decreasing particle size has been found to significantly increase efficiency as well. Not only do these small particles increase the speed of
mass transfer, they also pack more uniformly, decreasing the $A$ term, eddy diffusion. However, this comes at the cost of higher backpressure, leading to a need for ultra-high performance liquid chromatography (UHPLC), with pressures up to 1000 bar (14,500 psi) or more.

While chromatographic technologies are constantly improving, increasing column efficiency and leading to higher peak capacities, other approaches must be considered. In addition, increased peak capacities may not solve the problem of co-elution, which is common between analytes with similar chemical properties, such as isomers. In these cases, improved selectivity is required. Harnessing the different selectivities of two columns in one analysis presents a powerful separation method. Mixed mode columns, which consist of two functionalities, such as anion exchange and octadecyl carbon chains (C18), on a single column provide separation based on both ion exchange and polarity [24]. This type of column provides different $\alpha$ values for many compounds, but does not provide an increased peak capacity and necessitates the purchase (or synthesis) of a new column whenever a different selectivity is desired. Recently, we published a method for the synthesis of stationary phase gradients which allow for the fine tuning of chromatographic selectivity [25]. These stationary phase gradients were created on in-house synthesized monolithic columns by infusing an aminosilane functionalizing reagent through a bare-silica column creating a column with both amine and silica surface functionalities. The surface coverage and gradient steepness of the functional groups on the column support can be easily controlled by varying the time of infusion and concentration of reagent. This approach can be also extended to multiple functionalities, such as phenyl and C18 [26]. Although this approach makes tuning the selectivity of these columns simpler, the in-house synthesis of columns does not reach the same efficiencies as commercialized columns.
Another approach is the coupling of two or more separate columns. This can be accomplished in two main ways: serial connections or multidimensional coupling. Serially connected columns consist of two or more columns connected end-to-end, providing added selectivity. These columns can simply been connected with a piece of tubing; however, this potentially adds significant dead volume, leading to broadening of the chromatographic peaks. Commercialized versions of serially connected column, such as the POPLC® system (Bischoff Analysentechnik, DE), use specialized column “segments” which connect to one another with no additional tubing, thus eliminating the majority of dead volume between the columns. Still, the additional connections can prove to be problematic [27] and the column choices are limited by the offerings of a single manufacturer. Another approach, and the one focused on in Chapters 5 and 6, is two-dimensional liquid chromatography (2D-LC), in which two columns are coupled in an orthogonal manner.

2.2. Basics of Two-Dimensional Liquid Chromatography (2D-LC)

The 2D-LC technique combines two individual separations into a single analysis in order to provide greater separation power for a greater number of compounds due to its increased selectivity and increased peak capacities. The coupling of two separations is accomplished via a sampling valve placed between the two columns. This sampling valve collects a pre-defined volume of effluent from the first dimension (1D) column and then injects it onto the second dimension (2D) column. This is described further in the section 2.2.1. This sampling and 2D separation can occur in two main ways. In heart-cutting methods, a single section of the 1D chromatogram, often fully encompassing the peak(s) of interest, is collected in a single aliquot and transferred to the 2D column [28,29]. This creates a 1D chromatogram and a single 2D chromatogram. Multiple heart-cutting (MHC) methods do this for two (or more) sections of the
1D chromatogram, with each 1D section resulting in a single 2D chromatogram [30–32]. One potential disadvantage to this method is that any slight resolution obtained in the 1D separation will be lost due to mixing of the aliquot in the sample loop. Comprehensive 2D-LC (LC x LC) collects the entire 1D effluent in small aliquots (> 3 aliquots per peak [33]) and sequentially injects each aliquot onto the 2D column. This creates several 2D chromatograms, which when plotted together creates a two-dimensional (2D) chromatogram, with the 1D time on one axis and the 2D time on the second axis. In LC x LC the 2D separation time dictates the 1D sampling rate. Even at fast 2D separation times (< 1 min), the 1D separation is severely undersampled. A fairly recent variation of LC x LC, termed selective LC x LC (sLC x LC), can be viewed as a hybrid method of heart-cutting and comprehensive [34–37]. sLC x LC selects a single window in the 1D chromatogram similar to heart-cutting, but rather than collecting the entire window in a single aliquot, it collects several smaller aliquots, more similar to LC x LC. This produces multiple 2D chromatograms, each with a 1D axis for only the window selected. This method has the advantage of allowing for longer 2D separation times, for reasons that will be discussed further in the instrumentation section below. A graphical comparison of MHC, LC x LC, and sLC x LC is shown in Fig. 2.3.

In order to gain the most advantage from 2D methods, it is desired that the selectivities of the two separations be as different, or orthogonal, as possible [29]. This means that the retention of the compounds on the 1D column should be uncorrelated with the retention of the 2D column. This suggests that the most powerful combinations of separations would involve two different phases of separation and, indeed, these combinations are often seen in the literature. These include ion exchange (IEC) with reversed phase (RP) [38], size-exclusion (SEC) with RP [39], hydrophilic interaction liquid chromatography (HILIC) with RP [40], and others. Normal phase
Figure 2.3. Three modes of 2D-LC. (A) This diagram shows the 1D chromatogram and the collection of the 1D effluent samples across the 1D chromatogram. Each box represents a single aliquot collected in a single sample loop. Here, two peaks are selected for further analysis using MHC or sLC x LC. (B) These plots show the resulting chromatograms for each of the methods. MHC and sLC x LC result in two separate chromatograms for each of the two 1D windows collected. LC x LC results in a single comprehensive 2D chromatogram.

has also been used with RP [41]; however, solvent compatibility between the two separations must be considered. For example, normal phase LC uses highly non-polar organic solvents with increasing polarity to elute compounds. RP LC uses highly aqueous solvents with decreasing solvent polarity to elute compounds. Ignoring potential solvent immiscibility issues, if compounds elute from the 1D normal phase column in non-polar solvents and are injected onto
the 2D RP column, no retention (or minimal retention) will be observed. To circumvent these issues, complex methods are required. On the other hand, a very popular choice is the use of two RP separations [7,42–44]. While an increase in retention correlation is seen, the plethora of different RP column chemistries commercially available often allows for sufficient orthogonality between the two dimensions. Online selection tools are available to assist in the selection of orthogonal columns. HPLCcolumns.org [45], for example, makes use of the hydrophobic subtraction model (HSM) of selectivity [46] to compare over 600 commercially available RP columns from over 30 different manufacturers. By incorporating empirical values for hydrophobicity, steric effects, hydrogen bond acidity and basicity, and cation-exchange activity, this website calculates a similarity value between two columns. In their primer to LC x LC, Carr and Stoll attempt to compare different mode combinations according to several factors [42]. A portion of their table is reproduced here in Table 2.1. While the scoring is somewhat subjective, it presents a good overall picture of the different combinations. From this table, it can be seen that while the combination of two RP separations suffers slightly from lack of orthogonality, it is superior to all other combinations, particularly in terms of the wide range of compounds able to be analyzed by RP separations as well as the peak capacity per unit time of RP separations.

For the work in the following chapters, only comprehensive LC x LC and sLC x LC with RP conditions in both dimensions will be considered. LC x LC provides a significant increase in separation space allowing more peaks to be detected in a given analysis time. In terms of peak capacity, LC x LC provides an ideal 2D peak capacity \( (n_{c,2D}) \) equal to the product of the two individual separations’ peak capacities \( (n_{c} \text{ and } n_{c}) \); however, this 2D peak capacity is impacted by two major factors: peak broadening due to the undersampling of the 1D chromatographic peak and correlated retention between the two columns.
First, undersampling in LC x LC is due to the reality that the sampling rate of the 1D chromatogram is dictated by the speed of the 2D separation. This indicates a need for rapid separation in the 2D. To accomplish this, a combination of short column lengths and high temperature LC (HTLC) is often used in the 2D separation due to fact that higher temperatures allow for higher linear velocities owing to reduced solvent viscosities and therefore reduced backpressure [7]. Column efficiency is also increased using HTLC, particularly at high linear velocities [47]. This can be explained using the van Deemter equation (Fig. 2.2). At high temperatures, resistance to mass transfer (C term) is decreased and thus an increase in linear velocity has a lesser effect on H. These factors allow for a fast 2D separation; however, even with a 2D separation time of 12 seconds (to our knowledge, the fastest gradient separation achieved thus far in LC x LC [48,44]), the sampling rate of the 1D chromatogram is equal to 0.08 Hz, leading to the 1D chromatographic peak being represented by only a few data points, leading to a broadening of the peak [49]. A correction factor for broadening, $<\beta>$, can be incorporated in the

### Table 2.1. Comparison of different combinations for LC x LC

<table>
<thead>
<tr>
<th>1D Mode</th>
<th>IEC</th>
<th>SEC</th>
<th>NP</th>
<th>RP</th>
<th>HILIC</th>
<th>HILIC</th>
<th>SEC</th>
<th>SEC</th>
<th>LCCC</th>
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<td>RP</td>
<td>RP</td>
<td>RP</td>
<td>RP</td>
<td>RP</td>
<td>HILIC</td>
<td>RP</td>
<td>IEC</td>
<td>RP</td>
</tr>
<tr>
<td>Orthogonality</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>Peak capacity</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>++</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>--</td>
<td>+</td>
</tr>
<tr>
<td>Peak capacity/time</td>
<td>-</td>
<td>--</td>
<td>+</td>
<td>++</td>
<td>+</td>
<td>+</td>
<td>--</td>
<td>--</td>
<td>+</td>
</tr>
<tr>
<td>Solvent compatibility</td>
<td>+</td>
<td>+</td>
<td>--</td>
<td>++</td>
<td>+</td>
<td>++</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Applicability</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>++</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Score</td>
<td>4</td>
<td>3</td>
<td>1</td>
<td>9</td>
<td>5</td>
<td>2</td>
<td>-2</td>
<td>-3</td>
<td>2</td>
</tr>
</tbody>
</table>

*IEC – ion-exchange; RP – reversed phase; SEC – size-exclusion; NP – normal phase; HILIC – hydrophilic interaction; LCCC – liquid chromatography under critical conditions; Adapted from ref. [42].
definition of \( n_{c,2D} \). Davis, Stoll, and Carr define \(<\beta>\) as shown in Eq. 2.8 [49], where \( t_s \) is the sampling time and \( \sigma \) is the 1D peak width before sampling.

\[
\langle \beta \rangle = \sqrt{1 + 0.21 \left( \frac{t_s}{\sigma} \right)^2}
\]  

(2.8)

Correlated retention is also a major factor when estimating peak capacity. When retention in both dimensions of separation is strongly correlated, the chromatographic peaks elute along a diagonal line across the 2D separation space. If the two separations are completely uncorrelated, or orthogonal, the peaks appear evenly spread across the separation space. As stated previously, the combination of two RP separations will never be completely orthogonal and thus the separation space will always be under-utilized. The extent of this under-utilization of the separation space is captured in a metric called fractional coverage (\( f_{\text{coverage}} \)) and can be estimated via many different methods [50,51]. Figure 2.4 shows the cases of high and low \( f_{\text{coverage}} \). Typically, low fractional coverage is a result of the peaks falling along a diagonal line across the 2D separation space, however it can also be caused by weak retention in one dimension of the separation, leading to elution along a horizontal or vertical line in the 2D chromatogram. Eq. 2.9 incorporates both \( f_{\text{coverage}} \) and \( <\beta>\) into the calculation of \( n_{c,2D} \) [52].

\[
n_{c,2D} = \frac{n_c \times n_c \times f_{\text{coverage}} \times \frac{1}{\langle \beta \rangle}}{n_c}
\]  

(2.9)
As stated previously, mobile phase issues are of great concern in LC x LC. Even with RP in both dimensions, the compatibility of the solvents must be considered. When using gradient elution, compounds that elute towards the end of the 1D gradient are contained in a highly organic mobile phase composition. This sample is then injected onto the 2D column which is at a low organic composition. This solvent mismatch can cause broadening or even splitting of analyte peaks [53–55]. An area of interest in LC x LC literature is the promotion of on-column focusing in order to give better peak shapes. On-column focusing occurs when a compound is highly retained at the front of the column, leading to a narrower analyte band [56–58]. In LC x LC this can be achieved by diluting the 1D effluent with a weaker (i.e., aqueous) solvent in order to increase retention at the head of the column. While this decreases the analyte concentration and greatly increases the injection volume, Stoll et al. showed that the effect of on-column focusing was great enough to allow for narrower peaks and thus higher signal-to-noise ratios (S/N) [59] in comparison to the case with no dilution. Another approach is to modulate the temperature at the front of the column such that at injection the head of the column is at a lower

\[\text{Figure 2.4.} \text{ Diagram comparing cases of low fractional coverage (A) and high fractional coverage (B). Each dot represents an analyte peak. Low fractional coverage is often a consequence of correlated retention between the two separations causing peaks to elute along a diagonal line across the separation space.}\]
temperature, which increases retention and focuses the analytes [60]. This has been applied mostly to capillary LC due to the low thermal mass of capillary columns.

2.2.1. Instrumentation

High performance liquid chromatography requires the use of a high pressure pump to deliver the analyte mixture and mobile phase to the column where the analytes are separated. After separation, the analytes are detected using a detector suitable for the target analytes, often a diode array detector (DAD) or a mass spectrometer (MS). The choice of detector is crucial to the success of the analysis. Mass spectrometers are available with varying levels of mass resolution and have become a powerful detector in terms of both selectivity and sensitivity. When tandem mass spectrometry (MS/MS) is utilized, sensitivity and selectivity are enhanced even further along with the ability to obtain structural information about the compounds being analyzed. Mass spectrometers, however, are costly compared to alternatives such as DADs and require much more upkeep and maintenance. Issues with ion suppression are also commonly present in MS when multiple compounds coelute [61,62]. This occurs when two compounds enter the ionization source (e.g., electrospray ionization) and one compound negatively affects the ionization of the second compound. The exact mechanism of this process is not fully understood. The most common cause of ion suppression is interferences in the sample; however, ion suppression can also be caused by compounds introduced during sample preparation or even from tubing on the instrument [62]. These effects can lead to a drastic decrease in instrument sensitivity towards certain compounds in the sample. DADs, which measure the ultraviolet-visible absorption of analytes, are much more inexpensive and robust, but can suffer from lower sensitivity and lower selectivity due to the broad absorption bands of most organic compounds. Stoev and Stoyanov have shown, however, that the reliability of compound identification is
similar between DAD and low-resolution tandem MS [63]. Coelution in DAD is also less problematic as the analyte signals are additive when coelution occurs (assuming the total amount of analytes eluting is within the linear range of the detector). Combined with the use of curve resolution methods (described in the next chapter), LC-DAD detection can be a powerful method for analysis.

The instrumentation used for LC x LC is similar to that used for 1D LC with a few notable differences. Figure 2.5 shows a basic LC x LC set up. Two pumps are required for the two dimensions of separation. Figure 2.5 also shows the instrument set up with dual DADs. While the 1D DAD is not strictly required, it is useful in optimizing the instrumental conditions as it allows for the visualization of the 1D chromatogram prior to sampling. It can also serve to improve the quantitative abilities of LC x LC as described in Chapter 5. The sampling of the 1D effluent is accomplished via a sampling valve with two sampling loops. A 10-port/2-position valve is shown in Fig. 2.5, but concentric 8-port/2-position valves are becoming common as well [44,64,30]. Both valve types accomplish the same task of collecting the 1D effluent in the two sample loops. An optional dilution pump is shown in Fig. 2.6. This pump can be used to dilute the 1D effluent with a weak solvent in order to promote on-column focusing in the 2D, as discussed in the previous section. Splitting of the 1D effluent can also be employed to facilitate method development by de-coupling the flow rate of the two dimensions [65].
Figure 2.5. Diagram representing a typical setup for LC x LC, with a 10-port/2-position valve. The dashed outlines on the dilution pump and 1D DAD indicate these components are optional.

In LC x LC only two sampling loops are required due to the fact that as one loop is being filled with 1D effluent, the contents of the second loop are being delivered to the 2D column. The valve switches and these loops switch roles. sLC x LC requires a more complex sampling valve setup. sLC x LC collects many aliquots of the 1D effluent and stores them in sample loops rather than immediately injecting them onto the 2D column. This allows for fast sampling of the 1D separation and longer separations on the 2D column. Because of this, ten or more sampling loops are required. These are typically configured in what is sometimes called a “sampling deck [66]” or a “parking deck [30].” The instruments typically contain two of these sampling decks, where each one is used for a single peak window. This allows for a selected peak window to be sampled 10 or more times, which allows for a much faster 1D sampling rate than allowed in comprehensive LC x LC.

Data are collected at the 2D DAD as a sequence of 2D chromatograms, corresponding to the 2D separation of each sampled volume of the 1D effluent. In order to visualize these data as a 2D chromatogram, the sequence of 2D chromatograms must be folded into a 2D chromatogram.
Figure 2.6. Folding of instrumental data into a 2D chromatogram. Data is collected as a string of 2D chromatograms as shown in (A) separated by the dashed lines. These 2D chromatograms are aligned perpendicular to the 1D time axis as shown in (B). Typically this 2D chromatogram is visualized as a contour plot as shown in (C).

As shown in Fig. 2.6. Then, the data can be visualized as either a contour plot or a 3D plot. Note that these plots can only represent a single wavelength at a time so the choice of wavelength at which to visualize the data is important.

Despite its complexity both in instrumentation and method development, LC x LC provides a powerful analysis method for complex samples that are difficult to analyze with 1D-LC alone. Instrument manufacturers such as Agilent [67] and Shimadzu [68] are starting to recognize the strength of LC x LC and are starting to market LC x LC systems. This will likely lead to more widespread adoption of the technique due to the increased support and technical assistance that comes with the purchase of a commercial instrument.

One potential drawback of LC x LC, however, is its quantitative abilities. Typically, LC x LC has been seen to have poorer quantitative abilities compared with 1D-LC. This can be attributed to several factors which are described further in Chapter 5. The work described in
Chapters 5 and 6 aims to apply chemometric curve resolution to the analysis of LC x LC data in order to improve the quantitative abilities of LC x LC. These curve resolution methods are described in the following chapter.
Chapter 3: Chemometric Techniques for Liquid Chromatography


While LC is a powerful analysis technique, the data obtained from such analyses are often complex and necessitate the use of advanced data analysis techniques. Chemometrics provides many useful data analysis tools by utilizing mathematical concepts to solve chemical problems. Svante Wold, widely considered one of the fathers of chemometrics, defined chemometrics as "How to get chemically relevant information out of measured chemical data, how to represent and display this information, and how to get such information into data [69]." This definition encompasses both post-acquisition analysis of the data as well as pre-acquisition design of experiments in order to collect data that contains the most information pertinent to the goal of analysis.

Post-acquisition data analysis can be divided further into preprocessing and pattern recognition. Pattern recognition aims to find underlying trends in the data for easier visualization of the data or for more targeted purposes such as discriminating between two sample groups (e.g., healthy versus diseased individuals). This includes methods such as principal components analysis (PCA), partial least squares (PLS) [70], and cluster analysis, to name a few [71].
3.1. Traditional Preprocessing Methods

Preprocessing methods are often employed prior to pattern recognition in order to remove unwanted contributions to the data to reveal relevant signals in those data. Chromatographic data consist of three main contributions to the instrumental signal as shown in Fig. 3.1. Figure 3.1.A. depicts the chromatogram obtained instrumentally. This is an combination of the analytical signal, background, and noise (Figs 3.1.B,C,D, respectively). Background and noise reduction methods are among the most common preprocessing techniques. These aim to eliminate the background and noise signals from the raw data leaving only the analytical signal. This analytical signal contains the information about the compounds analyzed and thus the information that is relevant to the analysis.

To remove the background, several methods of baseline correction have been developed. The most commonly used method in currently available software packages is curve fitting [10].

Figure 3.1. The instrumental signal (A) consists of the analytical signal (B), the background (C), and the noise (D). Adapted from Matos et al. [72] and Amigo et al. [10].
This approach attempts to estimate the baseline by fitting a curve under each peak based on the area around the base of each peak. These fits can be global fits, or can be localized, which may provide a better fit but may not be continuous between segments [10]. When analyzing complex samples this approach can be problematic; if the chromatogram does not have baseline resolution around many of the peaks, it can be difficult to fit a suitable curve to estimate the baseline. Background correction methods have also been proposed for LC x LC. Often, these background methods extend 1D methods to 2D chromatograms. Filgueira, et al. extended traditional 1D background correction methods (median filtering and polynomial fitting) to LC x LC chromatograms [73]. Their method, named orthogonal background correction (OBGC), applies the selected method at each extracted 1D chromatogram (at each 2D time point). This implementation is orthogonal to how the data is collected (individual 2D chromatograms). The authors found this to be a very effective and easy to implement method.

Most noise reduction techniques make use of the fact that noise in data is typically high-frequency with low peak widths. The most well-known form of noise reduction is smoothing. This is often performed with Savitzky-Golay smoothing in which a few data points are captured in a window and those points are multiplied by a set of coefficients and summed [74]. The new value replaces the original center point of the window. The window is moved through the data, multiplying the data points within each window by the coefficients at each step. A popular variation on this is matched filtering in which the coefficients used in each window are set to match the expected peak shape [75,76]. For example, in chromatography, the analytical signal is expected to be a Gaussian peak shape so the matched filter is set to be a Gaussian peak with a similar peak width. Danielsson et al. proposed using a second-derivative Gaussian peak as the
filter [77]. By using the second-derivative, linear background contributions are reduced to zero, essentially performing background subtraction and noise reduction in the same step.

Once background and noise contributions are removed from the data, peaks must be detected to allow for further analyses. It is most often desired to detect peaks in an automated fashion particularly in complex samples and/or with a large number of samples. Two major approaches to peak detection are the use of derivatives and peak fitting. When the first-derivative of a chromatographic peak is taken, the zero-crossing point corresponds to the peak maximum. The peak width can be estimated using the second-derivative, where two zero-crossing points exist near the edges of a chromatographic peak [9]. Peak fitting can also be used for peak detection. In this approach the user or software uses a fixed peak model and optimizes the fit of the model to the data. Gaussian peak models are often used [10]; however, most chromatographic peaks exhibit non-Gaussian characteristics due to tailing and sometimes fronting. Very often, an exponentially modified Gaussian (EMG) model is employed due to its ability to model a tailing chromatographic peak [78]. A multitude of other peak models are also available which may fit peaks better depending on the data being fit. Almost 90 of these models have been compiled by Di Marco and Bombi [79]. Determination of the peak shape can be a tedious “guess-and-check” process, particularly if the peaks do not conform to a Gaussian or EMG shape. In cases where peak overlap is severe it is particularly difficult, if not impossible to choose determine the best peak shape. Amigo et al. demonstrated that in some cases, two peak shape models seem to fit the data equally well, but it is impossible to know which model better explains the true underlying peak shape [10]. Peak fitting methods are known as hard models in that the model is set and the parameters are estimated to best fit this model. A more flexible method is known as soft modeling, or self-modeling. These methods are explained in section 3.2.
3.2. Multiway Curve Resolution Methods

Most traditional methods treat the data using a single or a few detector channel(s) (i.e. a single wavelength or mass-to-charge value); however, this approach can exclude important information that may be contained in other channels such as other analytes or information about the background. Multiway curve resolution analyses treat the full spectrum and the chromatographic dimension as a single array of data from which components can be extracted. These components ideally correspond to each compound present in the sample. These methods can accomplish background and noise reduction by treating them as one or more extra component(s) in the data. In theory these can greatly simplify analysis by eliminating the need for multiple algorithms for each preprocessing step, which may affect the results of one another. Often, these methods eliminate the need for separate peak picking algorithms as well because each compound is ideally contained in a separate component. These components can then be utilized for the further pattern recognition steps. Two of the most widely used curve resolution methods are parallel factor analysis (PARAFAC) and multivariate curve resolution-alternating least squares (MCR-ALS).

3.2.1. Data Structure

In order to apply multiway methods of analysis to the entire dataset, the data must be organized into a single data array. These arrays are classified by the number of dimensions they contain, resulting from the instrumental measurement. Figure 3.2 depicts the types of data
structures typically encountered. The data produced by LC-DAD analysis, for example, are classified as a second-order structure (*i.e.*, a matrix) with both a temporal and a spectral dimension. When multiple samples are included, the data becomes a three-way data array (*i.e.*, a cube). LC x LC-DAD adds a second temporal dimension creating a third-order array for a single sample and a four-way array for a multi-sample dataset. LC-MS/MS adds a second spectral dimension; however, the spectra collected in the second dimension of MS are not independent of the selected precursor mass, and therefore cannot be represented as an additional multilinear dimension in the data.

These data can be analyzed with curve resolution algorithms (*i.e.*, PARAFAC or MCR-ALS) in order to resolve pure analyte signals from one another and from background signals. These pure analyte signals can then be used for quantitation or pattern recognition.

**Figure 3.2.** Data structures resulting from instrumental techniques of increasing complexity.

Adapted from Olivieri [80].
3.2.2. Parallel Factor Analysis (PARAFAC)

Parallel factor analysis is one technique capable of handling higher order data. PARAFAC is based on the assumption that the instrumental response, $X$, is a trilinear combination of pure compound responses in each dimension: the concentrations of the compounds in each sample, the chromatographic profiles, and the spectra (for LC-DAD data), and has dimensions of $I \times J \times K$ [81]. Here, $I$ is the number of chromatographic time points, $J$ is the number of m/z or wavelength points and $K$ is the number of samples. The PARAFAC model can be represented below:

$$\text{vec}(X) = \sum_{n=1}^{N} a_n \otimes c_n \otimes s_n + \text{vec}(E)$$  \hspace{1cm} (3.1)

where $a_n$, $c_n$ and $s_n$ are the sample, chromatographic and spectral profiles for the $n^{th}$ component and which form matrices $A$ ($I \times N$), $C$ ($J \times N$) and $S$ ($K \times N$), respectively. $E$ represents the error contribution and has the same dimensions as $X$ [72]. The $\otimes$ symbol represents the Kronecker product, and the vec operator rearranges the multidimensional array into a vector [81]. This is depicted graphically in Fig. 3.3. If multi-dimensional chromatographic data are being analyzed, a fourth (or more) dimension(s) can be added to the term within the summation.

![Graphical depiction of the PARAFAC model (Eq. 3.1) with two components. Adapted from Bro [82].](image)
The PARAFAC model is trilinear (when analyzing third-order data such as LC-DAD) which restricts its use to data whose retention times do not shift between runs. This is a strict requirement in LC where retention time shifts can occur due to slight inconsistencies in flow rate and mobile phase gradient as well as column aging. If shifts in retention occur, an alignment step, such as correlation optimized warping (COW) [83], is required, or an algorithm such as PARAFAC2 can be used, which attempts to handle retention time shifts within the model. Bortolato and Oliveri showed, however, that PARAFAC2 did not perform well when interferents were present [84]. Several algorithms exist for fitting the PARAFAC model to obtain the sample, chromatographic, and spectral profiles. These include generalized rank annihilation method (GRAM) and alternating least squares (ALS) [81,82].

Khakimov et al. [85] employed PARAFAC2 for the analysis of LC-MS data in a study of a plant’s resistance to insects. With the exception of splitting the chromatograms into smaller segments to decrease data complexity, no preprocessing steps were performed. Combining the results of PARAFAC with pattern recognition, these authors were able to correlate four previously reported compounds and five previously unknown compounds to the plants’ resistance to insects.

Synovec and colleagues have used PARAFAC extensively for the analysis of two dimensional gas chromatography (GC)-MS data [86–89]. After processing raw chromatograms in LECO ChromaTOF software [90], PARAFAC was used to resolve the chromatograms for the purpose of quantification. Their approach consists of a pattern recognition step to find the peaks that change the most between samples, then using ChromaTOF software to match mass spectra.
to a library. Finally, the relative concentrations of the analytes of interest were found using peak areas determined with PARAFAC [86]. They were able to identify 26 metabolites that differed between yeast metabolizing glucose and yeast metabolizing ethanol, with ratios ranging from 0.02 to 67 for the compounds [88], as well as identifying 63 metabolites between 5 strains of yeast and correlating them with RNA data to create pathways which could be compared across strains [86].

Due to the trilinearity requirement, PARAFAC is difficult to implement in LC x LC. Retention time shifts are very common in both separation dimensions. In the second dimension, retention time shifts mainly occur due to misalignments of the 2D chromatograms. This is very often due to slight variations in the valve timing. In the first dimension, slight retention time shifts are amplified by the undersampling of the 1D separation. Due to these requirements, it is often very difficult to implement PARAFAC in LC x LC, and often LC, analyses. PARAFAC can also be time consuming do to the computational demands compared to other techniques. For datasets with many samples, this can be another potential disadvantage.

3.2.3. Multivariate Curve Resolution (MCR)

Another method of curve resolution which is much more flexible is called multivariate curve resolution (MCR). MCR is a based on a bilinear model, rather than the trilinear model of PARAFAC, which makes MCR impervious to retention time shifting. The only requirement of MCR is that the spectral profiles of each analyte do not vary over the dataset. DAD is a very reproducible method which almost always meets this requirement, whereas other method such as mass spectrometry can have slight variations in spectra between each chromatographic time points.
The MCR model is shown in Eq. 3.2 and Fig. 3.4, where $X$ is the raw data matrix (as described in the previous section) and $C$ and $S$ are matrices consisting of the pure analyte (and background) chromatographic and spectral profiles, respectively.

$$X = CS^T + E$$  \hspace{1cm} (3.2)

**Figure 3.4.** (A) This graphic depicts the MCR model for a LC-DAD dataset with $I$ time points, $J$ wavelengths, and $N$ components representing the data. The yellow areas represent how each point in the raw data ($X$) is represented by the chromatographic ($C$) and spectral ($S$) profiles and error ($E$). Figure inspired by Rutan et al. [11]. (B) This graphic shows an example of a 2-compound spectrochromatogram resolved into two pure analyte chromatographic and spectral profiles.

Unlike PARAFAC, which can accept higher order data structures, MCR requires a matrix as input. For a single LC-DAD run, the data are contained in a matrix which fits the MCR model; however, when multiple samples are used, the data must be rearranged from a three-way array to a matrix. As shown in Fig. 3.5, this is accomplished by concatenating the chromatograms from each sample end-to-end along the time dimension to create a single augmented time dimension...
while retaining the spectral dimension. This matrix can then be used for MCR-ALS analysis. After analysis the data is then re-folded to determine the component profiles in each sample.

Similar to a multi-sample dataset, LC x LC-DAD adds another dimension to the data, creating a third-order data array, or a four-way array when multiple samples are analyzed. This requires two rearrangement steps. First, the two separation dimensions are unfolded to a single time dimension consisting of the $^2$D chromatograms appended to one another similar to how the data are collected (see Fig. 2.6). Then, the time and sample dimensions are unfolded to a single dimension as depicted in Fig. 3.5. This results in a single spectral dimension and a single augmented time dimension which can be analyzed via MCR-ALS.

![Diagram of data rearrangement](image)

**Figure 3.5.** Rearrangement of a multisample LC-DAD dataset into a single matrix for MCR analysis. $\lambda$ represents the spectral dimension, while $t$ represents the time dimension. (A) This graphic depicts the process graphically while (B) shows the process with realistic chromatograms with the wavelength axis going into the page.

### 3.2.3.1. Alternating Least Squares (ALS)

Data collected from the instrument are represented by $X$ in Eq. 3.2 but the goal of MCR is to obtain the chromatographic and spectral profiles for each analyte and background.
contribution, contained in the $C$ and $S$ matrices, respectively. To accomplish this, an alternating least squares (ALS) methodology is most often used. ALS works by solving Eq 3.2 for $C$ and $S$ alternatively using least squares solutions as shown in Eqs. 3.3 and 3.4.

$$C = XS(S^T S)^{-1}$$ (3.3)

$$S^T = (C^T C)^{-1} CX$$ (3.4)

Before the ALS process can begin, a number of components (the number of columns in $C$ and $S$) must be chosen and an initial guess, or estimate, must be obtained for either $C$ or $S$ as a starting point for ALS. The selection of the correct number of components is data dependent and often multiple numbers of components are tested for the best MCR-ALS performance. Scree plots, shown in Fig. 3.6., can offer a starting point for this determination. These plots are obtained from the singular values (from a singular value decomposition) of the data plotted versus the number of components [71]. A break or a shoulder in the trend can indicate a reasonable starting point for determining the number of components.

**Figure 3.6.** Scree plot for estimating the number of components for MCR-ALS. The possible break point occurs at 4 components indicating a starting point of 4 components for MCR-ALS.
The initial guess is typically derived from the raw data. Several methods have been developed for this purpose. If an initial guess of the chromatographic profiles, \( \mathbf{C} \), is desired, evolving factor analysis is a popular choice [91,92]. Most often, however, an initial guess of the spectral profiles, \( \mathbf{S} \), is preferred. All of the methods for obtaining an initial guess of \( \mathbf{S} \) involve extracting individual spectra from the raw data. The methods differ in how this is accomplished. Methods include SIMPLISMA [93,94], iterative key set factor analysis (IKSFA) [95,96], and iterative orthogonal projection approach (IOPA) [9,97], which is used throughout the work presented in the following chapters. IOPA, developed previously in our lab [9,97], builds off the previously developed orthogonal projection approach (OPA) [98,99] which extracts a set of spectra that represent the most different, or orthogonal, spectra in the dataset by defining a dissimilarity metric. This dissimilarity metric is defined as the determinant of a matrix containing the set of spectra for the initial guess. IOPA adds an iterative step in which each spectrum in the initial guess is replaced by each spectrum in the raw data to maximize the orthogonality between the spectra set used for the initial guess. If some or all of the analyte spectra are known \textit{a priori}, they can be used in the initial guess; however, OPA is still performed to obtain initial estimates of the background and potential interferent spectra [9].

Once the initial guess is obtained, it is used in either Eq. 3.3 or 3.4 to initiate ALS. ALS alternatively solves for \( \mathbf{C} \) and \( \mathbf{S} \) until either a predefined number of iterations is met or a solution is converged upon as defined by a minimum change in fit error (\( E \) in Eq. 3.2). A defining factor of ALS is that every time an updated estimate of \( \mathbf{C} \) or \( \mathbf{S} \) is calculated, the column vectors contained in these matrices are subjected to constraints. Mathematically, a wide range of possible solutions are possible for Eqs. 3.3 and 3.4, a problem often referred to as rotational ambiguity. A much smaller range of solutions are physically reasonable, however. For instance, analyte
chromatographic and spectral profiles cannot be negative. To drive the ALS solutions to physically reasonable solutions, constraints are imposed. These constraints, which are typically only applied to components (columns in C and S) that correspond to true analyte signals, most commonly include non-negativity, unimodality, normalization, and spectral selectivity. Non-negativity replaces any negative values with zeros, recognizing that concentration values and spectral intensities of chemical species cannot be negative. Unimodality recognizes that well-behaved, spectrally distinct analytes should only produce a single maximum in their chromatographic profiles. Unimodality eliminates any secondary maxima. Selectivity allows for any known values to be input into the C and/or S matrices. This is most often used in the spectral profiles to incorporate the knowledge that most analytes will not absorb at higher wavelengths and therefore the absorbance values at those wavelengths should be zero. It has been observed that spectral selectivity greatly helps in resolving analyte signals from background signals [100,101]. Selectivity can also be used in the chromatographic dimension or the sample dimension. Sample selectivity is especially useful when blanks are present in the dataset, and it is desired to set an analyte signal to zero in that sample. Normalization of the spectra solves the problem of intensity ambiguity. That is, normalization ensures that any intensity information is contained in the chromatographic profiles rather than the spectral profiles, allowing for quantitation based on the chromatographic profiles.

In the next chapter, we will explore the advantages of applying MCR-ALS to LC-DAD data, particularly in terms of its ability to increase the number of quantitatively analyzable peaks in a chromatogram.
4.1. Introduction

When developing chromatographic methods, the primary goal is often to increase the separation between analytes in order to facilitate further analysis. Sufficient separation is particularly important when quantitation is the final goal of the analysis in order to enable the precise and accurate recovery of peak areas (or peak height). Typically, the extent of separation is defined by chromatographic resolution ($R_S$) as discussed in Chapter 2 and the ideal separation ability is estimated by peak capacity ($n_c$). By definition, peak capacity is defined as the number of peaks that can be separated at a $R_S$ of 1.0 in an analysis [20]. As discussed in Chapter 3, curve resolution methods such as multivariate curve resolution (MCR-ALS) can resolve overlapping chromatographic peaks, allowing for quantitative analysis at low $R_S$.

Optimization of the chromatographic conditions is a crucial step in developing a chromatographic method; however, method development can be time consuming and may still not lead to satisfactory results. Similar attention to the application of chemometric approaches to enhance the analysis is often not considered as carefully. These methods can assist in both qualitative and quantitative analysis. MCR-ALS has been reported in the literature as early as 26 years ago [102,103] and has been utilized in a multitude of papers in many fields since then. Specifically, MCR-ALS has been applied to liquid chromatographic (LC) data for various
applications including, but not limited to, environmental analyses [104–106], omics studies [107–109], detection of biomarkers [110,111], food analyses [92,112,113], and pharmaceutical analyses [114–116]. Several software packages for MCR-ALS are available including a program by Tauler and colleagues [117,118] and the ALS package for the R programming environment by Mullen [119]. Commercial chemometric programs are also available which include MCR-ALS such as the PLS toolbox (Eigenvector Research, Inc., Manson, WA), which works through the MATLAB programming environment.

Despite the demonstrated utility of MCR-ALS, it has yet to find widespread use in chromatography outside of literature reports. It is our belief that this is due to an assumption that MCR-ALS (and other chemometric techniques) are overly complex and difficult to implement. There have also been very few papers characterizing the performance of MCR-ALS and other curve resolution techniques and demonstrating their potential in everyday analyses. Davis, Rutan, and Carr [120] investigated the use of chemometrics to analyze LC x LC data in a theoretical study involving statistical overlap theory [121,122] and multivariate selectivity [123]. Multivariate selectivity is related to the quantitative precision that can be obtained when using PARAFAC [82]. In that work, the authors found that in a separation space with a peak capacity of 100 with 200 randomly positioned peaks actually present, 120 peaks would be able to be analyzed quantitatively if PARAFAC were to be used. In comparison, only 20 peak maxima were able to be identified in the chromatograms without chemometric assistance. That work provided a detailed theoretical demonstration of the potential advantages of chemometrics, but no curve resolution was actually performed and no experimental deviations from ideality (background, retention time shifts, etc.) were considered. Fraga, Bruckner, and Synovec also investigated the use of chemometric curve resolution in two dimensional separations, but with
generalized rank annihilation method (GRAM) [124]. They found that, on average, the number of analyzable peaks was increased by a factor of two; however, no spectral information was utilized in that work, relying solely on the additional information provided by a second dimension of separation. This work also neglected the influence of background on the performance of the curve resolution techniques, which may be significant in many cases.

There is a need for a general characterization of MCR-ALS performance over a range of conditions in order to provide general guidelines of when it can provide an advantage in an analysis. In the work in this chapter, LC-DAD data have been simulated under a range of conditions with real instrumental background signals and analyzed with MCR-ALS. Then, quantitation was performed and a model of quantitative MCR-ALS performance was created.

4.2. Methods

All simulated data creation and MCR-ALS analyses were performed in MATLAB (R2013a, Mathworks, Inc., Natick, MA, USA). MCR-ALS was performed using an in-house program. Modeling was performed using JMP statistical software (version 12.0.1, SAS Institute, Inc. Cary, NC, USA).

4.2.1. Design of Experiments

To investigate the effects of different factors on the quantitative performance of MCR-ALS, simulated LC-DAD data were created varying the signal-to-noise ratio (S/N), the chromatographic resolution ($R_S$) and the spectral similarity as measured by the coefficient of determination ($r_{spectra}^2$). A three factor, three level full-factorial experimental designs was created
to investigate the effect of each of these three factors on the quantitative ability of MCR-ALS. The factor levels are listed in Table 4.1.

<table>
<thead>
<tr>
<th>S/N Peak B</th>
<th>Chromatographic Resolution (R_s)</th>
<th>Spectral Similarity (r_spectra)²</th>
</tr>
</thead>
<tbody>
<tr>
<td>26</td>
<td>0.25</td>
<td>0.5</td>
</tr>
<tr>
<td>256</td>
<td>0.5</td>
<td>0.95</td>
</tr>
<tr>
<td>513</td>
<td>1.0</td>
<td>0.98</td>
</tr>
</tbody>
</table>

*S/N of reference peak A = 1026; S/N calculated at middle calibration point

Simulated chromatograms were created using two Gaussian peak shapes with a set peak width (σ = 0.05 min). One peak, designated peak A, was held at a roughly constant intensity and at the same position to be used as the reference peak. The intensity and position of the second peak (peak B) were varied to give different S/N and R_s values. To create a calibration set, five samples were created. If peak A were held at exactly constant intensity across these five samples, any “cross-talk” between the resolved chromatographic profiles of the two peaks may influence the calibration quality of peak B. To account for this, a slight variation in the intensity of peak A (uncorrelated with peak B) across the samples was introduced in the data. A spectral dimension was added by first creating four artificial spectra. These were created by combining Gaussian peaks of varying positions (along the wavelength dimension) and peak widths to create realistic spectra. One spectrum was set as a reference, while the other three were used to create different
spectral correlations with the reference spectrum (listed in Table 4.1). These spectra are shown in Fig. 4.1.

Background data were added to each sample using real instrument backgrounds taken from 15 blank, gradient chromatographic runs. For each sample within a dataset, two backgrounds were chosen at random from the set of 15 backgrounds and a weighted average was calculated using random weights. These were then added to the simulated data. By taking randomly weighted averages, an infinite number of different backgrounds were able to be created within a reasonable range. Examples of the simulated chromatograms are shown in Fig. 4.2.
Figure 4.2. Examples of simulated chromatograms at varying $S/N$ and $R_S$ values. Each color represents one of the five different samples in each dataset.

Twenty-seven of these datasets were created with different combinations of the factors shown in Table 4.1, each with the same set of five backgrounds. Each of these datasets was then analyzed individually with MCR-ALS.

4.2.2. Multivariate Curve Resolution- Alternating Least Squares

For each of the 27 datasets, MCR-ALS was performed to resolve peak A and peak B from each other and from background in order to build a calibration curve based on the area of peak B. IOPA was used as an initial guess. Non-negativity and selectivity were used in both spectral and chromatographic dimensions. Unimodality was used in only the chromatographic dimension and normalization was used in the spectral dimension. A background smoothness constraint was used in the chromatographic dimension on the background component(s). This constraint ensures the background chromatographic profiles are smooth, which is expected for backgrounds caused by a changing mobile phase composition. This constraint is based on Eilers’ perfect smoother
algorithm [125] which requires a single value be input to determine the extent of smoothing. In order to remove any peak-like features in the background, this value was set high to effectively “over-smooth” the background chromatographic profiles. For all analyses, except in cases where $S/N$ was 26, three components were used. For the cases $S/N$ was 26, five components were needed to sufficiently resolve peak from the background.

After MCR-ALS resolution, the area of peak B was calculated for each sample by summing all intensities in the resolved chromatogram of peak B and a calibration curve was built. From this calibration the quality of fit was calculated using the coefficient of determination of the calibration curve ($r^2_{cal}$).

4.2.3. Monte-Carlo Simulations

After the MCR-ALS analyses of the 27 datasets, the set of background chromatograms was changed to a new, randomly generated set of backgrounds produced in the same manner as the previous set. The entire process was repeated including creation of datasets and MCR-ALS. This process was repeated a total of 50 times to give 50 replicates for each of the 27 combinations of factors, each with a slightly different background.

4.2.4. Modeling

From the calibration curves, the coefficient of determination ($r^2_{cal}$) was calculated. A model was built using the average $r^2_{cal}$ from each combination of factors ($n = 50$) as the response variable and $S/N$, $R_s$, and $r^2_{spectra}$ as the predictors. Before building the model, both the spectral and calibration $r^2$ variables were converted to angles using Eq. 4.1 [126]. Using angles ($\theta$) rather than $r^2$ as the model variables allows for greater sensitivity particularly at higher values of $r^2$.
where a difference between 0.9 and 0.999 is mathematically small, but very meaningful in terms of fit quality. In terms of angles the difference between these two values is 10-fold. Of note is the inverse relationship between $r^2$ and $\theta$.

$$r^2 = \cos^2 \theta$$  \hspace{1cm} (4.1)

A model was built including independent terms and interaction terms, to predict the quality of the calibration using the three predictors. The generic model is shown in Eq. 4.2, where $b$ represents the intercept, the $m$ values represent the coefficients, and the $x$ terms are the predictors, defined in Table 4.2. Only terms whose coefficients were statistically significant ($p<0.05$) were included in the final model.

$$\log \theta_{cal} = b + \sum_{i=1}^{3} m_i x_i + \sum_{i=1}^{3} \sum_{j=1}^{3} m_{ij} x_i x_j + m_{1,2,3} x_1 x_2 x_3$$  \hspace{1cm} (4.2)

**Table 4.2. Predictors used in building prediction model**

<table>
<thead>
<tr>
<th>$i$ or $j$</th>
<th>$x$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>log($\theta_{\text{spectra}}$)</td>
</tr>
<tr>
<td>2</td>
<td>log(S/N)</td>
</tr>
<tr>
<td>3</td>
<td>log($R_s$)</td>
</tr>
</tbody>
</table>

4.3. Results and Discussion

A model was built with $S/N$, $R_s$, and $\theta_{\text{spectra}}$ as predictors and $\theta_{cal}$ as the response variable as shown in Eq. 4.2 and Table 4.2. All terms were log-scaled to ensure no negative predicted values. Each independent term and cross-term was evaluated for its significance in affecting the
model using significance testing. Terms with \( p \)-values < 0.05 were included in the final model. The terms included are listed in Table 4.3 along with their corresponding coefficient (\( m \)), standard error (\( SE \)), and \( p \)-value. To interpret this model, it is important to remember the inverse relationship between \( r^2 \) and \( \theta \). A larger \( \theta_{cal} \) indicates a worse \( r_{cal}^2 \) for the calibration curve and a larger \( \theta_{spectra} \) indicates more dissimilar spectra. As shown in Table 4.3, \( \theta_{spectra} \) has an inverse relationship with \( \theta_{cal} \); as the spectra corresponding to peaks A and B become more dissimilar, the calibration fit quality improves, as expected. Similarly, \( S/N \) and \( R_S \) have inverse relationships with \( \theta_{cal} \); as the peak becomes less intense and the peaks become more overlapped, the calibration fit quality worsens. Several cross-terms were also found to be statistically significant including a quadratic term for \( \theta_{spectra} \) as well as all of the interaction terms between the predictors.

Table 4.3. Coefficients and errors for predictive model of calibration quality

<table>
<thead>
<tr>
<th>Term*</th>
<th>( m )</th>
<th>( SE )</th>
<th>( p )-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>1.15</td>
<td>0.11</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( \theta_{spectra} )</td>
<td>-1.670</td>
<td>0.075</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( S/N )</td>
<td>-0.646</td>
<td>0.030</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( R_S )</td>
<td>-0.35</td>
<td>0.15</td>
<td>0.0331</td>
</tr>
<tr>
<td>( (\theta_{spectra})^2 )</td>
<td>-1.70</td>
<td>0.34</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( \theta_{spectra} * S/N )</td>
<td>0.47</td>
<td>0.10</td>
<td>0.0001</td>
</tr>
<tr>
<td>( \theta_{spectra} * R_S )</td>
<td>-2.00</td>
<td>0.30</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( S/N * R_S )</td>
<td>0.50</td>
<td>0.12</td>
<td>0.0007</td>
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<tr>
<td>( (\theta_{spectra})^2 * R_S )</td>
<td>-4.3</td>
<td>1.4</td>
<td>0.0061</td>
</tr>
<tr>
<td>( \theta_{spectra} * S/N * R_S )</td>
<td>1.46</td>
<td>0.39</td>
<td>0.0016</td>
</tr>
</tbody>
</table>

*All variables are log-scaled; cross-terms are mean centered
Using this model, quantitative performance can be predicted with different combinations of $S/N$, $R_S$, and $\theta_{\text{spectra}}$ (or $r_{\text{spectra}}^2$). Of particular interest in this work is the prediction of the resolution required to obtain satisfactory calibration curves given a particular $S/N$ and spectral similarity. Figure 4.3 shows the effect of $R_S$ on $r_{\text{cal}}^2$ at three different $S/N$ values. A threshold value of $r_{\text{cal}}^2$ was defined at 0.999 for defining a satisfactory calibration, which is shown by the dotted line in Fig. 4.3. The point at which the calibration fit quality rises above 0.999 is considered the minimum resolution for quantitative analysis, $R_s$. As is shown in the figure, this value greatly depends on $r_{\text{spectra}}^2$, represented by the colored lines.

**Figure 4.3.** Predicted calibration quality ($r_{\text{cal}}^2$) versus $R_S$ at $S/N = 100$ (A), 50 (B), and 26 (C). The dotted line represents a threshold $r_{\text{cal}}^2$ of 0.999, above which represents a satisfactory calibration curve.
Figure 4.4 shows the relationship between $R'$ and $S/N$ at various values of $r_{spectra}^2$. As expected, as the correlation between the spectra decreases, the ability to perform quantitative analysis at lower $R_S$ and $S/N$ improves. For example, at $r_{spectra}^2 = 0.75$, satisfactory quantitation at $S/N = 50$ is possible at a resolution as low as 0.25. With more similar spectra (i.e., higher correlation), MCR-ALS does not provide satisfactory results for peak resolution because MCR-ALS works by separating peaks based on the difference between the spectra of the different compounds. This explains the results in Fig. 4.4 that at higher values of $r_{spectra}^2$ the $R'$ increases and is greater than one at low $S/N$; however, $R'$ is a function of the threshold $r_{cal}^2$. In Fig. 4.3 it can be seen that the $r_{cal}^2$ is still greater than 0.99 in all cases, showing that MCR-ALS is able to resolve the analyte signals with similar spectra and low $S/N$, just with a worse calibration fit quality.

![Figure 4.4](image)

**Figure 4.4.** Calculated minimum resolution required for quantitation ($R'$; defined at $r_{cal}^2 = 0.999$) as a function of $S/N$. Each line represents a different spectral correlation ($r_{spectra}^2$) as shown in the legend. Values in the gray area are below the range of the model and thus are extrapolated values.
4.3.1. Effective peak capacity in MCR-ALS

As shown in the previous section, MCR-ALS is able to resolve severely overlapped signals permitting quantitative analysis at low chromatographic resolution. In order to determine the enhancement in peak capacity, an effective peak capacity in MCR-ALS ($n_{c,MCR}$) is defined by Eq. 4.4 which includes the parameter, $R_s'$, the minimum required resolution for quantitation. Traditionally $n_c$ for a gradient LC separation is defined as the analysis time ($t_{analysis}$) divided by the peak width at four times the standard deviation ($\sigma$) of the peak. By incorporating this traditional peak capacity, $n_c$, into Eq. 4.3, Eq. 4.4 is obtained. From this equation, it can be shown that the peak capacity enhancement is proportional to the inverse of $R_s'$.

$$n_{c,MCR} = \frac{t_{analysis}}{4\sigma R_s}$$  \hspace{1cm} (4.3)

$$n_{c,MCR} = \frac{n_c}{R_s'}$$  \hspace{1cm} (4.4)

Conventionally, peak capacity is defined at $R_s = 1$, giving the traditional peak capacity equation. When using MCR-ALS, quantitation is possible at resolutions less than one, as shown in Fig. 4.3. Using $R_s'$ values from Fig. 4.4 in Eq. 4.3, the effective peak capacity can be estimated and the peak capacity enhancement can be calculated. For example, in a chromatogram where the target peak has a S/N of 100 ($S/N_{reference} = 1026$) and the $r_{spectra}^2$ of the two compounds is 0.9, then $R_s' = 0.45$, meaning there will be a 2.2-fold enhancement in peak capacity. At higher S/N values, the peak capacity is enhanced even more. Quantitative analysis can be performed at $R_s = 0.20$, corresponding to a 5-fold enhancement, at an S/N as low as 52, depending on the correlation between the compound spectra.
4.4. Conclusions

While MCR-ALS has been well documented in the literature, it has not yet become a standard tool in the analytical chemist’s arsenal. In the work presented here, three crucial factors impacting the performance of MCR-ALS – S/N, $R_s$, and $r_{spectra}^2$ – were varied and the potential for peak capacity enhancement in quantitative LC-DAD analysis was evaluated. It was found that MCR-ALS significantly increased the effective peak capacity. In many cases, the number of quantitatively analyzable peaks was increased by a factor of five or more compared to the traditional peak capacity at a resolution of one. This has the potential to significantly impact analyses by reducing the need for long analysis times and/or complicated methods to completely separate analyte peaks for subsequent quantification. The data simulated in this work included no pure standards. Previous work in our lab found that including pure standards in the data structure submitted to MCR-ALS improves the performance by providing a better initial guess to initiate the ALS algorithm [9].

In this work MCR-ALS was applied to LC-DAD data; however, the technique is broadly applicable to other instrumental methods including LC x LC, LC-MS, various spectroscopies, etc. Based on the results here, MCR-ALS can be expected to perform even better with increased peak separation (as in LC x LC) or with decreased spectral correlation (as in high resolution mass spectrometry). We believe that the ability of MCR-ALS to quantify significantly overlapped analyte peaks along with its relative ease-of-use lends itself well to being utilized much more often in the analytical laboratory for both routine and complex analyses.
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Chapter 5: Two-Dimensional Assisted Liquid Chromatography


Section 5.4.3 was adapted from D.W. Cook, S.C. Rutan, D.R. Stoll. 2016 *in preparation for submission to Chemom. Intell. Lab. Syst.*

5.1. Introduction

The increasing need for the analysis of complex samples necessitates the development of new analytical techniques and data analysis strategies. Particularly in “-omic” type applications, these samples can contain several hundred to several thousands of compounds [4]. For these applications traditional 1D chromatography is being pushed to its limits in regards to peak capacity, particularly when analysis time is limited. Many of these applications require analyte quantitation not merely detection. To do this the analytes must be adequately resolved in order to accurately determine how much is present in the sample [101].

Comprehensive two-dimensional liquid chromatography (LC x LC) can provide a significant advantage over one-dimensional liquid chromatography (1D-LC) in terms of resolving power. Ideally, the theoretical peak capacity for a two-dimensional (2D) separation is
equal to the product of the peak capacities of each separation dimension; however, this peak capacity is significantly smaller than ideal due to undersampling and lack of orthogonality between the two dimensions as discussed in Chapter 2. Despite these limitations Stoll, Wang, and Carr determined through both theoretical and experimental studies that when separation times are greater than 10 minutes and the 2D separation is conducted sufficiently rapidly, LC x LC has superior effective peak capacity as compared to 1D-LC [48,52,65,127]. While multi-dimensional chromatography has definite advantages in terms of peak capacity and peak capacity per unit time, the precision and accuracy of these methods compared to 1D chromatography often are not considered. Indeed, multi-dimensional liquid chromatography methods often suffer in terms of quantitative performance in comparison to their traditional 1D counterparts [101]. Stoll et al. reported percent relative standard deviations (%RSD) for two dimensional (2D) peak areas ranging from 0.7% to upwards of 15% with most falling between 1.5% and 7% when manual peak integration was employed [59]. In another paper, Stoll et al. compared the precision of 1D-LC and LC x LC and found that the 2D peak areas were on average seven-fold less precise based on the %RSD [52]. Kivilompolo et al. reported %RSDs for peak volumes ranging from 3%-13% for antioxidants in wines and juices [128]. Dugo et al. were able to quantify more polyphenols in red wines due to the increased resolving power of LC x LC; however, for the compounds detected in both LC x LC and 1D-LC, the %RSD in LC x LC was 12-fold higher than in 1D-LC [129]. A summary of these results is presented in Table 5.1. While the quantitative ability of LC x LC is improving with improved instrumentation and methods, poor quantitation still remains an issue and is unlikely to completely disappear without even more complex instrumentation and methods.
Table 5.1. Literature reports of quantitation in LC x LC

<table>
<thead>
<tr>
<th>Work</th>
<th>LC x LC Result</th>
<th>1D-LC Result</th>
<th>LC x LC Integration Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stoll, et al 2014 [59]</td>
<td>%RSD: 0.7 – 15%; most between 1.5 – 7%</td>
<td>Not reported</td>
<td>Manual integration</td>
</tr>
<tr>
<td>Stoll, et al 2008 [52]</td>
<td>%RSD: 1.5-8%</td>
<td>%RSD: 0.2-2.2%</td>
<td>Manual integration</td>
</tr>
<tr>
<td>Kivilompolo, et al. [128]</td>
<td>%RSD: 3-13%</td>
<td>Not reported</td>
<td>Manual integration</td>
</tr>
<tr>
<td>Dugo, et al. [129]</td>
<td>%RSD: 1.1-4.3%</td>
<td>%RSD: 0.1-2.2%</td>
<td>Manual integration</td>
</tr>
<tr>
<td></td>
<td>*LOQ ~6 x higher for LC x LC</td>
<td></td>
<td>Chrom square software (Chromaleont)</td>
</tr>
<tr>
<td>Place, et al. [130]</td>
<td>%RSD: 1.8-5.4%</td>
<td>%RSD: 2.7-4.7%</td>
<td>Automated integration of individual 2D slices</td>
</tr>
</tbody>
</table>

Factors that can contribute to this poorer precision include the use of multiple sample delivery valves and loops, which require precise control; the sampling of the first dimension; and higher background signals at the 2D detector. These high background signals result from the use of fast second dimension gradients. When short overall analysis times are required, and a gradient is used in the second dimension, the speed of the gradient causes a substantial increase in the 2D baseline due to dynamic refractive index effects in the UV-visible detector cell [131]. Dilution of the analytes is also of concern. Dilution can occur in two ways. First, the nature of LC x LC causes the analytes to be diluted when being delivered to the second dimension [57]. One strategy to counteract the dilution issue is to increase the volume of 1D effluent injected onto the 2D column, thereby increasing the number of moles of analyte delivered to the second dimension; however, this can lead to volume overload of the 2D column. This problem can be somewhat ameliorated by on-column focusing in the second dimension as described in Chapter 2. This on-column focusing can be enhanced by intentionally diluting the 1D effluent with a
weak solvent before delivery to the second dimension. This is useful for avoiding injection broadening on the $^2$D column caused by the delivery of the $^1$D effluent in a stronger solvent [59,35]. The amount of dilution needed is dependent on the analytes and should be optimized to avoid unnecessary loss of the signal-to-background caused by the loss of signal intensity. Sampling of the $^1$D separation can also cause a loss of S/N due to the division of a single chromatographic peak into three or more peaks.

Even given the superior peak capacity of LC x LC, many analytes may still be poorly resolved in complex samples, making quantification difficult. Another approach for improving peak resolution is to use a curve resolution technique, such as MCR-ALS. Using MCR-ALS, Bailey et al. were able to detect 18 peaks in a separation space which had a calculated chromatographic peak capacity of only 5 using a DAD detector [101]. Curve resolution can essentially improve resolution without increasing the complexity of instrumentation, while maintaining the inherent quantitative abilities of the instrument. This is demonstrated in Chapter 4 where MCR-ALS was shown to resolve chromatographic peaks with a resolution of less than 0.25 depending on the spectra similarity and S/N. While curve resolution methods work well for moderate to significantly overlapped peaks, they can fail when peaks are severely or completely overlapped and have low S/N. This is shown in Fig. 4.3, where the performance of MCR-ALS began to degrade at resolutions as high as 0.5 if the spectral similarity between the analytes was very high ($r_{spectra}^2 > 0.9$). Having the additional chromatographic resolution provided by LC x LC can potentially create a powerful technique when both LC x LC and MCR-ALS are combined.
In this chapter, a new strategy is presented, termed 2D assisted liquid chromatography (2DALC), for improving quantitation in LC x LC. This strategy is outlined graphically in Fig. 5.1. In this approach the higher resolving power of LC x LC is combined with the superior precision available from 1D chromatography by using a diode array detector (DAD) at the end of both the first and second dimension columns. By quantifying the 1D peaks, this strategy partially overcomes the inevitable resolution loss caused by the undersampling of the first dimension chromatogram, as well as the decrease in the signal to background ratio resulting from dilution encountered in LC x LC. This is accomplished by first using MCR-ALS to get an improved estimate of the pure component spectra from the 2D DAD data and then using those resolved spectra to initiate MCR-ALS analysis of the 1D DAD data. Calibration and quantification can then be performed using the resolved 1D chromatograms.

![Diagram](image)

**Figure 5.1.** Overall strategy for analyzing LC x LC-DAD data using 2DALC
5.2. Strategy

Figure 5.1 shows the overall strategy developed here. First, the 2D chromatogram is analyzed with MCR-ALS to obtain resolved spectra of the real chemical components. Then, these spectra are used to initialize MCR-ALS on the 1D chromatographic data to obtain the resolved 1D chromatograms. Calibration is performed with these resolved 1D chromatograms in order to assess the quantitative performance of this strategy.

5.2.1 Instrumental Setup

As discussed in Chapter 2, LC x LC commonly uses one detector at the end of the 2D column, which, after rearranging the data gives the 2D chromatogram. For this work, a multichannel detector is also needed at the end of the 1D column as shown in Fig. 5.2. In this case, both detectors were diode array detectors (DAD); however, this strategy should work with other multichannel detectors as well as long as the two detectors are identical. With this setup, both a 1D chromatogram from the 1D DAD and a 2D chromatogram from the 2D DAD can be produced. The addition of a detector after the first dimension will induce peak broadening due to added extra column volume; however, under the conditions of this experiment we estimate that the 1D detector will add at most a few percent to the 1D peak width prior to sampling (assuming a flow cell contribution of 1 µL²). When this is compared to the broadening of 1D peaks due to its undersampling [49], it is evident that the contribution of the 1D detector to the effective 1D peak width is very minor. As discussed in the previous section, pre-dilution with a weak solvent was also employed in this setup as shown in Fig. 5.2.
Figure 5.2. Instrumental setup for LC x LC with dilution and dual DAD detectors. Reproduced from Stoll et al. [59] with permission.

5.3. Experimental

All calculations and programs were written in MATLAB version R2013a (Mathworks, Inc. Natick, MA). ACD/Labs ChromProcessor 9.0 (Advanced Chemical Development, Inc. Toronto, Canada) was used to translate experimental data from Chemstation software files (Agilent Technologies Santa Clara, CA, USA, rev. C.01.05) to MATLAB format.

5.3.1. Simulated Datasets

The data discussed in this chapter consists of both simulated and experimental results, with each chromatographic analysis consisting of a 2D chromatogram and a 1D chromatogram, as shown in Fig. 5.3.

The simulated chromatograms consisted of two peaks, created by adding Gaussian peaks onto real, independent instrumental backgrounds. Backgrounds were collected on an Agilent 1290 Infinity 2D-LC system (Agilent Technologies, Waldbronn, Germany), which included a prototype Agilent pump to control the flow transferred to the second dimension. A more detailed
description of the instrumentation can be found in Allen et al. [44]. Mobile phases in both dimensions were water as the A solvent and acetonitrile as the B solvent. The first dimension consisted of an SB-C18 column (30 mm x 4.6 mm; 5 µm; Agilent Technologies, Little Falls, DE) maintained at 40 °C and a flow rate of 0.500 mL/min. A gradient from 5%-95% B over 10 minutes was used for a 15 minute time of analysis. The 1D effluent was split to 0.05 mL/min (see reference [65] for more details about splitting) and directed to a pair of 40 µL sample loops connected to an 8-port valve. The sample loops were alternatively filled with 1D effluent and then the effluent was delivered to the 2D column by switching the valve position. The cycle time was 12 sec and the gradient was 0%-100% B over 9 s. The second dimension consisted of a Poroshell 120 C18 column (30 mm x 2.1 mm; 2.7 µm; Agilent Technologies, Little Falls, DE) maintained at 80 °C with a flow rate of 3 mL/min.

For 2D chromatograms, Gaussian peaks were used in the second dimension (sampled at 0.0250 sec intervals) and sampled Gaussian peaks were used in the first dimension (sampled at 12 second intervals), using an in-house MATLAB program [132]. Two separate sets of simulated data, differing in only the amount of dilution of the 1D effluent before delivery to the 2D column, were created. The peak heights and widths were chosen to approximately match experimental data obtained for an amphetamine mixture. For low dilution conditions, the peak height ratio between 1D and 2D chromatograms was 2:1. For high dilution, the ratio was 10:1. The chromatographic resolution of the peaks was varied from 0.01 to 1 in both dimensions. At each resolution, 15 “samples” were created at varying concentrations including 9 calibration samples, 4 test samples, and 2 blanks. These concentrations are listed in Table 5.2. The spectra for the two analytes were taken from pure samples of 3-methoxyamphetamine (MoxyAmp) and 3,4-methylenedioxy-N-methylamphetamine (MDMA). The similarity of each compound’s spectra to
the rest of the data, including background, was measured using the first order selectivity metric developed by Lorber [123,133] and calculated as described in Cantwell et al. [134]. The selectivity of the analysis of MoxyAmp with respect to the background spectra and the MDMA was 0.714, and the selectivity of MDMA with respect to the background spectra and MoxyAmp was 0.311. Another measure of spectral selectivity, and that used in Chapter 4, is $r_{spectra}^2$, which equals 0.49 for MDMA and MoxyAmp. From this point on, MoxyAmp will be referred to as peak A and MDMA will be referred to as peak B.

Table 5.2. Relative concentrations of samples in the simulated dataset

<table>
<thead>
<tr>
<th></th>
<th>Concentrations</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Peak A</td>
<td>Peak B</td>
</tr>
<tr>
<td>Blank</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Calibration 1</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>Calibration 2</td>
<td>1.00</td>
<td>0.25</td>
</tr>
<tr>
<td>Calibration 3</td>
<td>1.00</td>
<td>0.50</td>
</tr>
<tr>
<td>Calibration 4</td>
<td>0.25</td>
<td>0.60</td>
</tr>
<tr>
<td>Calibration 5</td>
<td>0.50</td>
<td>1.00</td>
</tr>
<tr>
<td>Calibration 6</td>
<td>0.80</td>
<td>1.00</td>
</tr>
<tr>
<td>Calibration 7</td>
<td>0.60</td>
<td>0.80</td>
</tr>
<tr>
<td>Calibration 8</td>
<td>0.10</td>
<td>0.70</td>
</tr>
<tr>
<td>Calibration 9</td>
<td>0.90</td>
<td>0.10</td>
</tr>
<tr>
<td>Blank</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Test 1</td>
<td>0.50</td>
<td>0.50</td>
</tr>
<tr>
<td>Test 2</td>
<td>0.66</td>
<td>0.48</td>
</tr>
<tr>
<td>Test 3</td>
<td>0.39</td>
<td>0.77</td>
</tr>
<tr>
<td>Test 4</td>
<td>0.88</td>
<td>0.84</td>
</tr>
</tbody>
</table>
5.3.2. Experimental Datasets

The experimental data consisted of selective LC x LC (denoted sLC x LC) analyses of three furanocoumarins – psoralen, angelicin, and 8-methoxypsoralen (8-MOP). The multivariate selectivities of these compounds’ spectra were calculated to be 0.160, 0.204, and 0.156, respectively. The pairwise $r^2_{spectra}$ values for the furanocoumarin spectra were: psoralen/8-MOP 0.91; psoralen/angelicin, 0.95; and 8-MOP/angelicin, 0.89. The data consisted of three mixtures and three pure calibration standards for each compound. The concentrations of the components in each sample are listed in Table 5.3. In this work, sLC x LC separations of the furanocoumarins were performed using an Agilent 1290 Infinity 2D-LC system (Agilent Technologies, Waldbronn, Germany). Mobile phases in both dimensions were 10 mM phosphoric acid as the A solvent and acetonitrile as the B solvent. The 1D separations were carried out on a Poroshell 120 PFP (100 mm x 2.1 mm i.d.; Agilent Technologies, Little Falls, DE) column maintained at 40 °C with a flow rate of 0.15 mL/min and a 5 µL injection. A gradient was used with the following conditions: 40%-55% B from 0-6 min; 55%-95% from 6-7 min; 95%-40% from 7-7.01 min; and held at 40% until 9 min. The 1D effluent was sampled six times at 5-sec intervals starting at 3.75 min into 40 µL loops following dilution with water at 0.25 mL/min. Each 1D effluent sample was then delivered sequentially to the 2D column, which was a Zorbax SB-C18 (30 mm x 2.1 mm i.d.; 3.5 µm; Agilent Technologies, Little Falls, DE) column maintained at 60 °C with a flow rate of 2.5 mL/min under isocratic conditions (17.5% B) over 45 sec. For a more detailed description of the implementation of sLC x LC, the reader is referred to Stoll et al. [37].
Table 5.3. Concentrations of samples in the experimental dataset

<table>
<thead>
<tr>
<th></th>
<th>Concentrations (mg L(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Angelicin</td>
</tr>
<tr>
<td>Blank</td>
<td>-</td>
</tr>
<tr>
<td>Blank</td>
<td>-</td>
</tr>
<tr>
<td>Mix 1</td>
<td>4.0</td>
</tr>
<tr>
<td>Mix 2</td>
<td>6.0</td>
</tr>
<tr>
<td>Mix 3</td>
<td>1.5</td>
</tr>
<tr>
<td>Psoralen Standard</td>
<td>-</td>
</tr>
<tr>
<td>Psoralen Standard</td>
<td>-</td>
</tr>
<tr>
<td>Psoralen Standard</td>
<td>-</td>
</tr>
<tr>
<td>8-MOP Standard</td>
<td>-</td>
</tr>
<tr>
<td>8-MOP Standard</td>
<td>-</td>
</tr>
<tr>
<td>8-MOP Standard</td>
<td>-</td>
</tr>
<tr>
<td>Angelicin Standard</td>
<td>1.0</td>
</tr>
<tr>
<td>Angelicin Standard</td>
<td>2.5</td>
</tr>
<tr>
<td>Angelicin Standard</td>
<td>7.0</td>
</tr>
</tbody>
</table>
Figure 5.3. Representative chromatograms and spectra from the data analyzed in this work. The left column is the simulated data under high dilution conditions and experimental data is shown in the right column. A and B show the 2D chromatograms (at 200 nm) with a box around the actual section analyzed (the colorbars indicate the intensities in mAU). C and D show the 1D chromatograms (at 200 nm) and E and F show the pure spectra of the compounds. The peak resolution in both chromatographic dimensions is 0.5 (before undersampling) in A and C. See section 5.3 for chromatographic conditions.

To decrease the complexity of the data analysis and speed up analysis, regions of interest were selected from the raw data for analysis in both the 1D and 2D chromatograms (see the black
boxes in Figs. 5.3.A and 5.3.B). Before MCR-ALS analysis, the data must be augmented as described in Chapter 3.

5.3.3. MCR-ALS

MCR-ALS was performed using an in-house algorithm adapted from a previously described algorithm [135]. The code is reproduced in Appendix A. Non-negativity was applied as a constraint for the 1D and 2D chromatograms, unimodality was used for the 1D chromatograms, and spectral selectivity was used to set absorbances at the longer wavelengths to zero in the 2D chromatograms to assist in the differentiation of the signal from the background [101]. These constraints were applied only to the components which represented real chemical compounds; background components were not constrained.

5.3.4. Calibration

The resolved 1D chromatograms were integrated over the full retention time range to obtain the peak area. This gave acceptable results because of the complete resolution of the signals from each other and from the background. For the resolved 2D chromatograms, Savitzky-Golay second-derivative peak detection was employed to deal with the incomplete resolution of the analyte signals from the background signal. This was implemented using a Savitzky-Golay second-derivative smoothing [74,75] and then finding the peak start and end points based on a noise threshold in the second derivative. A window size of 19 points for the simulated chromatograms and 115 points for the experimental chromatograms for the second derivative smoothing was used, and the noise threshold was set at 0.0025 mAU. The peak area was then determined by summing the signal above a linear baseline created between these points in the MCR-resolved chromatogram. The areas of the calibration samples were then used to create a
calibration curve. The peak areas of the test samples were then used to estimate the predictive ability of this method, measured by root mean squared percentage error (RMSPE) as defined by Eq. 5.1 [136], where \( y_i \) is the true concentration, \( \hat{y} \) is the calculated concentration from the calibration curve, and \( n \) is the number of samples.

\[
RMSPE = \sqrt{\frac{\sum(y_i - \hat{y})^2}{n}} \times 100\%
\]  

(5.1)

5.4. Results

5.4.1 Simulated Data

Chromatograms were simulated to investigate the effect of peak resolution on the quantitative capabilities of 2DALC for both low dilution and high dilution conditions. RMSPE was used to compare the quantitative ability of 2DALC to the four alternative methods listed in Table 5.4. Examples of the \( ^1D \) and 2D data are shown in Figs. 5.3A and 5.3C, and the results are summarized in Figs. 5.4 and 5.5. These figures show the RMSPE as a function of both the first and second dimension resolutions. The bottom left corner of each panel in Figs. 5.4 and 5.5 shows the RMSPE for the case where no resolution is achieved in either the first or second dimension, and the top right corner of each panel shows the RMSPE for the case of high resolution in both dimensions. As expected, there is a clear trend where the performances of all of the methods improve as the resolution in both dimensions improves.
Table 5.4. Quantitation methods studied in this work

<table>
<thead>
<tr>
<th>Method</th>
<th>Chromatogram used for quantitation</th>
<th>Method of obtaining initial guess</th>
</tr>
</thead>
<tbody>
<tr>
<td>1D-IOPA</td>
<td>1D</td>
<td>IOPA</td>
</tr>
<tr>
<td>1D-Pure</td>
<td>1D</td>
<td>Pure analyte spectra with OPA estimated backgrounds</td>
</tr>
<tr>
<td>2DALC</td>
<td>1D</td>
<td>2D resolved spectra with OPA estimated backgrounds</td>
</tr>
<tr>
<td>2D-IOPA</td>
<td>2D</td>
<td>IOPA</td>
</tr>
<tr>
<td>2D-Pure</td>
<td>2D</td>
<td>Pure analyte spectra with OPA estimated backgrounds</td>
</tr>
</tbody>
</table>

When the two methods of resolving 1D chromatograms are compared (rows 1 and 2 in Fig. 5.4), it can be seen that IOPA begins to degrade at resolutions less than 0.3 versus using the pure spectra, where quantitative results can be achieved even when peaks are almost completely overlapped, (represented as a resolution of 0.01 in Fig. 5.4 and 5.5). A similar pattern is observed when the two methods of resolving 2D chromatograms are compared (rows 4 and 5 in Fig. 5.4 and rows 2 and 3 in Fig. 5.5). For peak A, as long as one dimension has a resolution greater than 0.4, the RMSPE is similar to that obtained using pure spectra as initial guesses. For peak B, the results are similar at low dilution; however, at high dilution 2D-IOPA fails to give acceptable quantitative results at any 1D resolution if the 2D resolution is low. In low dilution conditions (Fig. 5.4) the RMSPE is almost equal between peak A and peak B. In the high dilution conditions (Fig. 5.5) peak B has higher RMSPE values, which is consistent with the higher relative residual background in the resolved chromatographic profile due to the lower multivariate selectivity of this peak relative to the background and peak A (0.311 vs. 0.714 for peak A). In addition, some of the points in the 2D graphs do not follow the trends seen for peak A under the high dilution conditions. We attribute this to the difficulty in obtaining accurate peak
areas due to the low S/N, despite using peak detection to assist in the differentiation of peaks from residual noise in the resolved components.

2DALC is superior to 2D-IOPA (row 3 vs. row 4 in Fig. 5.4 and row 1 vs row 2 in Fig. 5.5). Under the low dilution conditions, the difference is small, but for higher dilution conditions, the advantage of quantitation in 2DALC is clear. Both peaks have poorer quantitation in 2D-IOPA when the resolution is less than 0.5 in both dimensions. Peak B has poorer quantitation at all resolutions when 2D-IOPA is used. There are also more inconsistencies in the patterns of both peaks, owing to the lower S/N in the 2D chromatogram. When 2DALC is used, quantitative performance is slightly better under low dilution conditions (compared to high dilution, compare row 3 of Fig. 5.4 to row 1 of Fig. 5.5), whereas the extent of dilution has a much bigger impact on the quantitative performance of 2D-IOPA (compare row 4 of Fig. 5.4 to row 2 of Fig. 5.5). Based on this we conclude that 2DALC is less sensitive to variations in S/N at the 2D detector.

2DALC also has advantages over 1D-IOPA (row 2 in Fig. 5.4). At 1D resolution of 0.25 1D-IOPA begins to fail. When the 2D resolution is greater than that of the 1D resolution, the spectrum obtained from the 2D chromatogram provides a better initial guess than IOPA analysis of the 1D data, and therefore provides better quantitation under both low and high dilution conditions. The exception is when 1D resolution is greater than 0.25; in this case IOPA is able to obtain a sufficiently accurate initial guess from the 1D chromatographic data leading to results that are equivalent to those obtained using 2DALC.
Figure 5.4. Results for simulated data at low dilution conditions (peak height ratio of 2:1). Colorbars indicate RMSPE. Scales for the graphs showing pure spectra initial guesses have different scales than the other graphs to better show the results. Resolution is defined as the chromatographic resolution before undersampling.
Figure 5.5. Results for the simulated data at high dilution conditions (peak height ratio of 10:1). For comparison to 1D results see rows 1 and 2 of Fig. 5. Colorbars indicate RMSPE values. Scales for the graphs showing pure spectra initial guesses have different scales than the other graphs to better show the results.

5.4.2 Experimental Data

Three furanocoumarins – angelicin, psoralen, and 8-MOP – were analyzed via sLC x LC. More information on these compounds is provided in Chapter 6. The dataset consisted of chromatograms for two mixtures analyzed in duplicate and three separate samples containing pure standards for each compound, as outlined in Table 5.3. All samples were analyzed,
representing a targeted analysis, in which the compounds present in the mixture are known ahead of time. The corresponding 2D and 1D chromatograms for one of the mixtures are shown in Figs. 5.3B and 5.3D, respectively. The results of this analysis are shown in Fig. 5.6. We observe that the RMSPE values for the 1D-IOPA and 1D-pure methods are almost equivalent, and superior to all other methods shown. This is because IOPA easily finds almost pure spectra from the standards to use as an initial guess. This indicates that when all of the compounds being analyzed are known, 2D chromatography does not provide an advantage over 1D chromatography for quantitation, when the spectra of the overlapped components are distinguishable and no spectrally similar interferents are present. In other words, spectral resolution along with the improved precision of 1D chromatography provides the necessary quantitative information without the need for the additional chromatographic resolution provided by the 2D column, as long as pure standards of the target compounds are available.

![Graph showing quantitation error for five methods applied to targeted sLC x LC analysis of furanocoumarins](image)

**Figure 5.6.** Quantitation error for the five quantitation methods applied to the targeted sLC x LC analysis of furanocoumarins (based on experimental data).

Often, such as in the case of “discovery” type metabolomics investigations, the analytes are not known prior to analysis; this is known as an untargeted analysis. To investigate this type of problem, the analysis of the furanocoumarin sLC x LC data was repeated, but without using
the data for the pure standard samples; this meant that there were no calibration standards included in the analysis. Therefore, the quantitative performance of the different methods was estimated by dividing the peak area by the concentration of the analyte in the mixture. The percent relative standard deviation (% RSD) was then calculated. As seen in Fig. 5.7, \(^1\)D-IOPA now gives the worst results due to the inability to obtain a sufficiently accurate initial guess, due to the severe overlap of compounds. Both 2D methods give similar results, with angelicin having a very good % RSD, but psoralen and 8-MOP still having a very poor % RSDs. The best results in the “untargeted” analysis are given by the \(^1\)D-pure method; however, this method is only used for comparison because pure spectra are not available prior to analysis in the untargeted case. 2DALC has the next best % RSD, being much better than both 2D-IOPA and \(^1\)D-IOPA. In this case, \(^1\)D-IOPA gives the worst results.

![Figure 5.7. Quantitation error for the five quantitation methods applied to the untargeted sLC x LC analysis of furanocoumarins (based on experimental data).](image)

**5.4.3. Combined 2DALC (c2DALC)**

An alternative approach to 2DALC was also investigated. In this variant, 2DALC was modified to perform the MCR-ALS analyses on the \(^1\)D and the 2D chromatograms simultaneously. This approach was named combined 2DALC (c2DALC). This method makes
use of a data augmentation step similar to that shown in Fig. 3.5. After the rearrangement of the 2D chromatogram, the 1D chromatogram is concatenated with the reshaped LC x LC chromatogram along the same augmented axis. This strategy is depicted in Fig. 5.8. This strategy is possible because both the 1D and 2D detectors collect spectral data with the same wavelength range and the same spectral resolution, meaning that spectra from both detectors have identical wavelength axes [137,138]. By concatenating the 1D and 2D chromatograms and performing a single MCR-ALS analysis, as opposed to a two-step process in 2DALC, the spectral information found in the 2D chromatogram can more directly assist in the resolution of 1D chromatogram, assuming the spectra are the same between the two detectors. As in 2DALC, the resolved 1D chromatogram is used for peak integration and subsequent quantitation.

**Figure 5.8.** Overview of the strategy utilized in c2DALC. First, the 2D chromatogram is unfolded and then concatenated with the 1D chromatogram. This concatenated chromatogram is then resolved with MCR-ALS and the 1D portion (represented by the gray shading) is integrated. Multiple samples can also be used by adding another concatenation as shown in Fig. 3.5.
The assumption that the spectra of the compounds will be identical between the two detectors is often incorrect. Because a molecule’s spectral properties are often influenced by its solvent environment [139], it can be expected that spectral shifts may occur between the 1D and the 2D detectors. As the molecules elute off of each column, they will be contained in a certain solvent composition and therefore a certain solvent polarity. Most often the mobile phase composition during the elution of a compound from the 1D column will be different than that from the 2D column. Even if the polarity difference is slight, spectral shifts may occur. To investigate the effect of these spectral shifts, data were simulated as described in section 5.3.1. One dataset was created with furanocoumarin spectra and the other was created with amphetamine spectra, which show less pronounced solvent effects on their spectra. The amphetamine spectra are also more dissimilar to each other compared to the furanocoumarin spectra. As demonstrated in Chapter 3, this will have an effect on all of the implementations of MCR-ALS tested here. The spectra used for the 1D and the 2D chromatograms were taken from MCR-ALS resolved experimental data from the 1D and 2D detector, respectively. This allowed the use of realistic spectral shifts in the data. As shown in Fig. 5.9, the furanocoumarin spectra have more pronounced spectral shifts than the amphetamine spectra. The chromatographic resolution in the second dimension (2RS) was held constant at 0.4 and the first dimension resolution (1RS) was varied to determine what effect resolution had on the performance of the methods tested.

Both datasets were analyzed with c2DALC, 2DALC, 1D-IOPA, and 2D-IOPA. The results are shown in Fig. 5.10 with 1RS values of 0.2, 0.3, and 0.5. In addition to integrating the 1D chromatogram in c2DALC as discussed above, the 2D chromatogram was also integrated.
Figure 5.9. Spectra used for data simulation. (A) shows the furanocoumarin spectra and (B) represent the amphetamine spectra: methylenedioxyamphetamine (MDMA), methoxymethamphetamine (MoxyMeth), and methamphetamine (Mamp). Spectral correlations ($r^2$) are a measure of the spectral shifts between the $^1$D spectra (solid lines) and the $^2$D spectra (dashed lines). These correlation values were calculated over the spectral range from 200-400 nm.

These results are labeled c2DALC (2D) in Fig. 5.10. In general, as $^1R_S$ decreases, the performance of $^1$D-IOPA improved, as expected. This trend generally held for 2DALC and c2DALC as well; however, the %RSD was generally higher for c2DALC versus 2DALC. For these data, 2D-IOPA most often performed the best at low $^1R_S$, with a few exceptions; this is most likely due to sufficient resolution of the compounds in the second dimension ($^2R_S = 0.4$) for MCR-ALS analysis. The trends seen in the angelicin plot are mostly different than the other compounds. This is most likely due to the fact that the angelicin elutes in the middle of psoralen and 8-MOP in the $^2$D separation causing peak overlap on both sides of the peak. In the $^1$D
Figure 5.10. Comparison of c2DALC, 2DALC, and other methods at varying 1D resolution. 2D resolution was held constant at 0.4. c2DALC (2D) represents the results when the 2D chromatogram was integrated after c2DLAC as opposed to the 1D chromatogram. Results from both furanocoumarins and amphetamines are shown, where the amphetamine spectra are less affected by the solvent difference between the 1D and 2D detector.

Another interesting trend shown in Fig. 5.10 is that of c2DALC when the 2D chromatogram is integrated (labeled as c2DALC (2D)), rather than integrating the 1D chromatogram. For many of the compounds, the %RSD values are lowest for c2DALC (2D) at low 1R_S values and increase as 1R_S increases. This is the opposite trend as c2DALC (where the 1D chromatogram is integrated). The goal of c2DALC is the simultaneous optimization of the 1D and 2D chromatograms. Ideally, the optimal solution would match the true peak profiles for both the 1D and 2D chromatograms.
(i.e., %RSD close to zero for both chromatograms); however, the opposing trends suggest that at low values of $R_S$, c2DALC is optimizing the fit based on the 2D spectra whereas at higher values of $R_S$ c2DALC is optimizing based on the 1D spectra. Due to this finding, and the observation that c2DALC did not provide a significant advantage over 2DALC, we decided not to pursue c2DALC for further analyses, such as those in Chapter 6.

5.5. Conclusions

A new method of quantitation in LC x LC based on the use of multivariate detection with a DAD in both dimensions has been developed. This method, 2DALC, uses MCR-ALS analysis of the 2D chromatogram to obtain a better initial guess for the component spectra for subsequent MCR-ALS analysis of the 1D chromatogram than would be obtained using the 1D detector data alone. For targeted analyses, where the spectra of all detected compounds are known prior to analysis, use of the 1D chromatography with pure spectra to initiate MCR-ALS is superior to all other methods tested, including all 2D methods. However, in contrast to untargeted analyses, where an initial guess of the analyte spectra must be extracted from the chromatograms for the unknown sample, 2DALC provides the best quantitation. Further, although not investigated in this work, we believe that 2DALC will provide quantitative advantages in targeted analysis in which unknown interferences overlap the target analytes. Based on simulated chromatograms, when the 1D peak resolution is less than 0.3, 2DALC gives better quantitation than the other, more conventional methods of data analysis tested (i.e., improved the accuracy by up to 14-fold compared to 1D-LC and up to 3.8-fold compared to LC x LC with a single multivariate detector). As chromatographic resolution improves, the quantitative performance of 2DALC and the conventional methods converge. The superiority of 2DALC over traditional IOPA-based methods increases as chromatographic resolution decreases. Additionally, 2DALC performs
increasingly better compared to other methods as sample dilution between the two separation dimensions of 2D-LC increases.

The performance of 2DALC depends on the spectral dissimilarity between the compounds themselves, and between the compounds and the background. The more similar the spectra, the harder it is for MCR-ALS to resolve severely overlapped peaks, even with the assistance of a second dimension of separation. This changes the threshold chromatographic resolution, below which point an advantage is gained by using 2DALC. This observation also depends on the degree of dilution between the first and second dimension separations. The use of 2DALC for high dilution cases provides better quantitation compared to quantitation based on the 2D detector signal because of the lower S/N in the 2D chromatogram. Even in low dilution cases, the background in the 2D detector signal is still worse than the 1D signal. The experimental data shown here involved a relatively high level of dilution. For other, more hydrophobic analytes, less dilution might be required, which should lead to improved performance from the MCR-ALS analysis of the 2D chromatogram [35]. It was shown that 2DALC is less sensitive to the amount of dilution, and therefore is recommended whenever dilution of any magnitude is applied. The results shown for simulated 2DALC also represent an optimistic case in which the spectra are identical between the two detectors. While this is often not the case, the experimental data confirmed that 2DALC can still provide a quantitative advantage.

A combined 2DALC approach was also investigated in which the 1D and 2D chromatograms were simultaneously resolved with the goal of improving 2DALC due to a more direct application of the 2D spectra to the 1D MCR-ALS analysis. It was found, however, that the effects of different solvent compositions on the analyte spectra at the two separate diode array detectors degraded the performance of c2DALC. While in cases of minimal spectral shifts,
c2DALC may provide a slight advantage over 2DALC; however, since it was found that even slight shifts greatly affect c2DALC, it was not tested in further analyses.

2DALC is a simple way to improve the accuracy and precision in the LC x LC analysis of compounds present in complex samples by combining LC x LC equipped with dual DADs and chemometrics, where neither method alone provides the desired quality of quantitation.
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Chapter 6: Comparison of Curve Resolution Strategies in LC x LC: Application to the Analysis of Furanocoumarins in Apiaceous Vegetables


6.1. Furanocoumarins

Many naturally occurring compounds in foods we eat are considered bioactive, with either positive or negative effects on our bodies. The complexity of the interactions of these compounds with our bodies very often means that these compounds can affect multiple biological pathways simultaneously, sometimes with both positive and negative effects. Many times the difference between beneficial and harmful effects is dependent on the concentration of the compound. In pharmaceuticals, medications must be dosed as to maximize the positive effects while minimizing the negative effects (i.e., side-effects). In foods it is also important to understand the effect of different compounds and their concentration-dependent interactions within the body.

One class of compounds known to have significant biological activity is furanocoumarins, found in significant levels in citrus fruits and apiaceous vegetables. As their
name suggests, furanocoumarins are defined by their structure of coumarin with a fused furan ring as shown in Fig. 6.1. The two common furanocoumarin isomers, psoralen and angelicin, are also shown in Fig. 6.1. Most furanocoumarins can be classified as a derivative of one of these two base structures.

Figure 6.1. Two basic furanocoumarin isomers, psoralen and angelicin, showing the furan and coumarin subunits in gray boxes.

Furanocoumarins are of great interest because of their interaction with the P450 enzymes found in the liver. P450 enzymes play a crucial role in the metabolism of many medicines and endogenous compounds. The interaction of furanocoumarins with these enzymes can have major implications on a medication’s effectiveness. One well-known effect, known colloquially as the “grapefruit juice effect,” is caused by furanocoumarins, namely bergamottin, interacting with the CYP3A4 enzyme. This enzyme is one of the most common P450s involved in the metabolism of many widely used medications such as statins, some blood-pressure drugs, and some anti-anxiety medications, to name a few [140]. Traditionally, grapefruit juice has been contraindicated with statin use, due to the increased bioavailability of the statins resulting from the decreased metabolism by the CYP3A4 enzyme [14]. Furanocoumarins are also known to cause photosensitivity, a condition sometimes called bartender or margarita dermatitis due to the rash that outdoor bartenders can develop from handling citrus fruits.
While often associated with their negative effects, furanocoumarins’ biological activity can often be harnessed for powerful medical treatments. Using grapefruit juice to inhibit CYP3A4 has been shown to increase bioavailability of a cancer therapeutic [141]. The photosensitizing effect of psoralen has been incorporated with ultra-violet therapy to increase its efficacy for diseases such as eczema, psoriasis, cutaneous T-cell lymphoma and others [142]. The inhibition of procarcinogen activation has also been studied. Procarcinogens are compounds that have the potential to become carcinogenic after activation by enzymes such as P450s. By inhibiting these enzymes by consumption of furanocoumarin rich foods, procarcinogen activation may be suppressed [143]. Finally, bergamottin has been proposed for treating overdoses of acetaminophen by inhibiting the conversion of excess acetaminophen to a toxic metabolite via the P450 enzymes [144].

Due to the important bioactivity of furanocoumarins, it is crucial to develop methods that are able to quantitatively analyze their presence in foods. Because of the complex sample matrix that is present in food analysis, it is necessary to employ more advanced analysis techniques. In this case, LC x LC is used to provide separation of the target analytes from the matrix, interfering compounds, and one another. Due to the complexity and relevance of furanocoumarin analysis, it was chosen for testing quantitation using different implementations of MCR-ALS for LC x LC-DAD data. Fourteen furanocoumarins were targeted in three apiaceous vegetables – parsley, parsnips, and celery. The structures and abbreviations of the fourteen targeted furanocoumarins are shown in Fig. 6.2.
The goal of the work in this chapter was to compare the quantitative performance of three methods of implementing MCR-ALS for LC x LC-DAD data. 2DALC as well as 1D and 2D analysis (both with IOPA initial guesses) were compared in terms of their quantitative performance for this dataset. In addition two methods of integrating the resolved chromatographic peaks are compared. The first method was manual integration of the resolved chromatographic peaks by visually choosing peak start and end points and drawing a linear baseline between the two points. The area between this baseline and the peak was then calculated. The second method, which we call summation, was performed by simply adding the intensities across the entire resolved component. Ideally, MCR-ALS removes most of the background contributions and interfering compounds signals (except spectrally identical compounds). This means that any points with non-zero intensity should correspond to the analyte

Figure 6.2. Structures and names of 14 furanocoumarins and the internal standard (IS) comprising the target analyte set

1. Psoralen
2. Angelicin
3. 8-Methoxypsoralen (8-MOP)
4. Sphondin
5. Isopimpinellin
6. 5-Methoxypsoralen (5-MOP)
7. Oxypeucudanin
8. Imperatorin
9. Dihydroxybergamottin
10. Trioxsalen
11. Phellopterin
12. Isoimperatorin
13. Epoxybergamottin
14. Bergamottin
IS 4-Chlorobenzophenone
signal. By summing all intensities, this should provide a simple alternative to the tedious manual integration approach.

6.2. Experimental

LC x LC analyses, including sample preparation, were carried out in the lab of Dwight Stoll at Gustavus-Adolphus College. Fourteen furanocoumarin compounds, listed in Fig. 6.2, along with one internal standard, 4-chlorobenzophenone (CBP) were analyzed. Samples of three apiaceous vegetable types – parsley, parsnip, and celery – were then analyzed for the presence and concentration of these 14 target compounds.

6.2.1. Plant Extracts

Samples of parsley, parsnips, and celery were purchased from a local grocery store and were provided by Sabrina Trudo and colleagues at the University of Arkansas. Sample preparation was accomplished using a QuEChERS methodology [145,146]. In general, the QuEChERS method entails the following steps: homogenization of sample with acetonitrile and internal standard, buffering and phase separation with salts, centrifugation and extraction of organic phase, and sample clean-up with dispersive solid phase extraction. For the vegetables in this work, samples were prepared combining 5.0 g of wet vegetable matter with 10 mL of water and 10 mL of acetonitrile (ACN) with 0.1% acetic acid. This mixture was then homogenized in an Ultra Turrax T25 Homogenizer (IKA Laboratories, Wilmington, NC) for 3 min. A 1 g portion of sodium acetate and 4 g of magnesium sulfate were added to the homogenate and hand-mixed by inversion for 2 min. Samples were then centrifuged at 3,000 rpm for 5 min at room temperature which resulted in three distinct layers: ACN, vegetable matter, and aqueous. Following centrifugation, 1.8 mL of the supernatant (i.e. the ACN layer) was transferred to a 2
mL dispersive solid extraction (dSPE) tube (United Science 20 CarbonX QuEChERS; Center City, MN; part number 05040114) to remove highly non-polar compounds such as pigments and lipids from the extract. Water was added to the dSPE tube to 10% of total volume to improve retention of undesirable non-polar compounds and tubes were vortexed with a Mini Vortex (VWR Part Number 12620-852, Radnor, PA) at the maximum speed setting. Samples were again centrifuged at 6,000 rpm for 30 sec and the supernatant was transferred to a HPLC vial and diluted 1:2 with water prior to LC analysis.

6.2.2. Chromatographic Conditions

The LC x LC analyses were performed on modules from the Agilent Technologies 1290 Infinity line (Agilent Technologies, Waldbronn, Germany). Samples were injected using an autosampler (Model G4226A). First and second dimension flows were generated using binary pumps (Model G4220A); a quaternary pump (Model DEQAT00023) was used to dilute 1D effluent. Thermostated column compartments were used in both dimensions (Model G1316C) and nominally identical diode-array UV detectors (DADs) (Model G4212A) were used at the outlets of the 1D and 2D columns. Low dispersion flow cells (800 nL volume, Model G4212-60038) were used in both the 1D and 2D DAD detectors. At the outlet of the 1D flow cell a pressure relief valve (Model G4212-60022) was used to protect the cell from breaking due to downstream pressure. A 2-position 8-port Duo-valve (Model 5067-4214) fitted with two 80 µL stainless-steel loops mounted on an external valve drive (Model G1170A) was used to transfer fractions of 1D effluent to the 2D column. The 2D-LC instrument was controlled by OpenLab Chromatography Data System (Agilent Technologies. Rev. C.01.07).
The $^1$D separation was performed using an Agilent Poroshell 120 PFP (100 mm x 2.1 mm i.d.) column with 5 mF sodium phosphate at pH 6 (A) and methanol (B). The unit milliformal (mF) used here indicates the amount of sodium phosphate added to solution rather than the amount of phosphate species that exists in solution after dissociation. A gradient was used with the following conditions: 45-45-75-100-100-45-45 % B from 0-12.5-29.5-31.5-33-33.01-36 min; this is shown in Fig. 6.3. The temperature was held at 20°C and the flow rate was 0.25 mL/min from 0-7.5 min and 0.125 mL/min from 7.51 to 36 min. Beginning at 7.0 min, the $^1$D effluent was diluted with 20 mM phosphoric acid in water at 0.15 mL/min to promote focusing of the analyte bands at the inlet of the $^2$D column [19]. The diluted effluent was collected in 80 µL sample loops before injection on the $^2$D column.

The $^2$D separation was performed from 7.5-36 min using an Agilent Zorbax SB-C18 rapid resolution high definition (RRHD) (50 mm x 3.0 mm i.d., 1.8 µm) column using 20 mM phosphoric acid in water (A) and acetonitrile (B) as mobile phase components. The temperature was held at 50°C and the flow rate was 2.5 mL/min. The $^2$D mobile phase was held isocratic at 42% B until 12.5 min and then a shifted gradient was employed as shown in Fig. 6.3.

6.2.3. Data Analysis

All computation was performed on a Dell Precision T3600 with an Intel Xeon E5-1620 CPU at 3.60 GHz and 32.0 GB of RAM. Data files were converted from Agilent .D format to MATLAB .mat format files using ACD/Lab Spectrus Processor (Advanced Chemical Development, Inc., Toronto, Canada). All other computational analysis was performed in MATLAB version R2013a (Mathworks, Inc., Natick, MA).
MCR-ALS was performed using an in-house algorithm which accepts LC x LC data in the original four-way data format (2D time x 1D time x spectra x sample). In the current work, non-negativity and selectivity were used in both the chromatographic and spectral dimensions. Unimodality was used in some cases; however, in many cases, particularly when analyzing the 1D chromatograms, interferents were present in the resolved component chromatograms, meaning that the true, underlying chromatographic profile contained more than one peak. Implementing unimodality in these cases would incorrectly constrain the chromatographic profiles, leading to incorrect results.

Prior to MCR-ALS analysis, the data were divided into sections to minimize chromatographic complexity and data size. The sections chosen for analysis are shown in Fig. 6.4. For MCR-ALS analysis, the full dataset was subdivided into three smaller datasets. Each dataset contained all samples from a single vegetable type \((n_{\text{parsley}} = 56; n_{\text{parsnip}} = 56; n_{\text{celery}} = 54)\). Each dataset also contained two blanks and three calibration sets, each consisting of five standard mixtures with analyte concentrations ranging from 1 – 50 µg/mL. Internal standard
calibration curves were created by integrating the resolved chromatograms, with each of the methods described in the following section. Because the concentrations of many of the compounds were found at the low end of this concentration range, calibration standards that were much higher than the predicted concentration in the extract were eliminated from the calibration curves, and analyte concentrations were predicted with this new calibration curve. The narrower range of concentrations allowed for increased accuracy of prediction.

Figure 6.4. Representative chromatograms of a standard (blue) where all compound concentrations are 10 µg/mL and a parsley extract (red; offset by 50 mAu in (A)). (A) and (B) show the 1D and 2D chromatograms, respectively. The boxes indicate the windows chosen for analysis and the numbers labeling the peaks correspond to the compounds in Fig. 6.2. IS = internal standard.
MCR-ALS was performed on the data using three strategies: 1D, 2D, and 2DALC. The 1D analysis corresponds to MCR-ALS being applied to only the 1D chromatogram, collected at the 1D detector. Likewise, the 2D analysis corresponds to MCR-ALS being applied to only the 2D chromatogram, collected at the 2D detector. Both of these methods used IOPA as the initial guess. Finally, 2DALC is the strategy described in Chapter 5, utilizing both the 1D and 2D chromatograms.

6.3. Results

Extracts from the three vegetable types – parsley, parsnips, and celery – were analyzed via LC x LC. Using diode array detectors (DAD) placed at the end of both the 1D and 2D columns, both 1D and 2D chromatograms were collected. This allowed for the comparison of quantitative performance of 1D and 2D chromatography. Representative chromatograms from both a standard and a parsley extract are shown in Fig. 6.4. From this, it can be seen that, as expected, the peaks are separated much better in the 2D chromatogram; however, interferents are still present. It is also important to note that lower level interferents in the 2D may not be seen in the 2D chromatogram due to the contour plot style in which minor peaks may not be seen.

6.3.1. Comparison of Integration Methods

Ideally, MCR-ALS completely resolves the pure analyte signals from the background and noise. This would allow for the integration over the entire pure analyte component. This is simply a summation of the total signal intensity contained in the resolved chromatographic profile of each component as shown in Fig. 6.5B. Unfortunately, MCR-ALS with DAD data rarely eliminates background and noise completely. This complicates the summation process by including residual background producing an overestimate of peak
Figure 6.5. Graphical representation of integration methods compared. (A) shows a MCR-ALS resolved LC x LC chromatogram. (B) and (C) show the rearranged LC x LC chromatogram with summation and manual integration, respectively. The shaded area represents the area calculated for quantitation. The percent difference in calculated peak area for these two chromatograms is 9.9%. The results from the two integration methods are identical for 1D LC.

area. In contrast, manual integration is performed by visually estimating the peak start and end points, drawing a linear baseline between those points, and integrating the peak area between that baseline and the peak as shown in Fig. 6.5C.

To compare manual integration to simple summation, resolved chromatographic profiles for both the calibration set and the unknown samples were integrated using both methods. Calibration curves were constructed and the concentrations in the samples were predicted. Since the samples were run in duplicate, the differences between the duplicates can be taken as a measure of the precision of each integration method. Figure 6.6 displays the sum of these differences across all
samples for five of the furanocoumarin compounds. By comparing Fig. 6.6A and 6.6B (showing the two integration methods), it can be seen that four of the five compounds showed better agreement between duplicates for all methods when manually integrated, as expected. The fifth compound, isoimperatorin, showed similar results between summation and manual integration across the three methods. This indicates that there is a significant amount of residual background and/or noise in four of the components which can be excluded with manual integration. While manual integration is considerably more tedious, the results are superior in terms of quantitation.

![Figure 6.6.](image)

**Figure 6.6.** Comparison of quantitative performance between different implementations of MCR-ALS. The difference in predicted concentration between instrumental duplicates is used to estimate the quantitative performance. (A) shows the results when the peaks are manually integrated. (B) shows when the resolved chromatograms are integrated over the entire component.
6.3.2. Comparison of Curve Resolution Strategies

In addition to manual versus summation approaches, it is important to understand the performance of the three methods tested – \( ^1 \)D, 2D, and 2DALC. Figure 6.6 compares the agreement between duplicates for the three methods tested. Figure 6.6A shows the comparison when employing manual integration. When comparing the \( ^1 \)D and 2D data analysis, the results are similar, with the 2D analysis having better agreement between the duplicates for 8-MOP and isoimperatorin, while the duplicates for imperatorin agree better with \( ^1 \)D analysis. However, note that these differences are not very large. Overall, \( ^1 \)D and 2D are similar due to the ability of manual integration to exclude interferents and background. Figure 6.6B shows the results for the summation approach. Overall the agreement is worse than for manual integration due to the residual background that occurs in all methods. For most of the compounds the \( ^1 \)D analysis has better agreement than the 2D analysis, with the exception of psoralen, which eluted early and had more interferents that may not have been completely resolved with MCR-ALS. Interestingly, when manual integration was applied, 2DALC had the best agreement only for imperatorin; however, for the summation approach, 2DALC performed the best or was equal to the best method for all but psoralen, most likely due to less residual background. For summation, any slight errors in MCR-ALS analysis would be overshadowed by residual noise, but when manual integration is employed, the small differences between resolution results obtained using the different methods become more evident.

For the best quantitative results, it is shown that 2D with manual integration is the best method for this data set; however, this process can be very time consuming. If the summation approach is to be utilized the \( ^1 \)D data provides superior results due to the fact that the signal-to-background ratio is higher in the \( ^1 \)D chromatograms. This is, however, limited to cases in which
there are no interfering compounds, meaning a compound with the same spectra occurs in the same resolved component as the target compound. In the presence of interfering compounds, the additional chromatographic resolution provided by LC x LC is key.

6.3.3. Vegetable Results (from 2D Manual)

Concentrations for the furanocoumarins in all three vegetable types were calculated using the 2D chromatograms with manual integration, which was determined to be the most precise method tested here as discussed in sections 6.3.1 and 6.3.2. The results are shown graphically in Fig. 6.7. Out of the fourteen compounds targeted in this analysis, only ten compounds were detected and the concentrations of these ten compounds are represented in the figure. Four compounds – trioxsalen, phellopterin, epoxybergamottin, and bergamottin – were not detected in any vegetable type in this study. It is interesting to note the wide range of concentrations for many of the compounds even within the same vegetable type. The sources of this large variation are unknown at this time; however, they may be related to the freshness, purchase data, geographical origin, or species of the vegetables. Correlations between sample information and analyte concentrations are currently being investigated by Sabrina Trudo (University of Arkansas, Fayetteville, AR) whose goal is to study the physiological effects of the consumption of these vegetables on rats, particularly in terms of anti-cancer properties [147]. Overall, the parsley samples had the highest concentrations of furanocoumarins, followed by parsnips, and then celery, on a per gram basis, but since the typical serving of parsley is much smaller than parsnips or celery, this does not reflect an actual dietary load.
Figure 6.7. Concentration (C) of detected furanocoumarins in the three vegetables as calculated with manual integration of the 2D chromatogram. The box represents the median and the 1st and 3rd quartiles. Each point represents the average concentration of instrumental duplicates in each sample. Only compounds which were found to be present in at least one vegetable type are shown. The total numbers of samples analyzed were: \( n_{\text{parsley}} = 56; n_{\text{parsnip}} = 56; n_{\text{celery}} = 54 \).

6.4. Conclusions

When quantitative analysis is performed it is important to consider the impact of all steps on the analysis. This is especially true when chemometric techniques are performed on the data. Here, several strategies for MCR-ALS analysis for LC x LC were compared along with the subsequent integration methods. In most cases 2DALC performed similarly to resolving the 1D
chromatogram by itself. Even though spectral shifts were present, the two-step strategy of 2DALC allowed for the 2D spectra to be further optimized for the resolution of the 1D chromatogram.

As expected, manual integration was superior to summation approaches; however, this approach is time consuming for studies involving many samples and for 2D chromatograms in which several 2D peaks must be integrated for a single chromatogram. If summation is to be used, the 1D chromatogram gave better agreement between replicates, but caution must be used so that interfering peaks are not included in the component window being summed. It is possible that the resolved chromatograms could be submitted to peak detection for a more automated method of excluding residual background [130]. Depending on the performance of the peak detection algorithm, results from this approach may approach those obtained from manual integration. This would facilitate the use of the 2D chromatographic data to obtain superior quantitative results.
Chapter 7: Conclusions and Future Work

As defined in Chapter 1, three main goals were set for the work in this dissertation: 1) to characterize the abilities and limitations of MCR-ALS for LC-DAD data; 2) to improve quantitative abilities of LC x LC through the use of a novel implementation of MCR-ALS; and 3) to demonstrate the utility of MCR-ALS for real-world LC x LC analyses. Work was performed towards these goals in Chapters 4-6.

7.1. Reflections on Chapter 4

Goal 1 was addressed in Chapter 4 where MCR-ALS was implemented on data over a wide range of conditions such as varying $S/N$, $R_s$, and spectral similarity. An effective peak capacity was defined where the number of analyzable peaks within a separation was defined by a calculated threshold $R_5$ rather than a $R_S$ of one. It was found that even when the target peak was of low intensity (i.e. low $S/N$) and was overwhelmed by a coeluting peak, excellent quantitation was achieved even at $R_S$ values less than 0.25. This was particularly true when analyte spectra were sufficiently dissimilar. These values corresponded to an up to five-fold increase in effective peak capacity. This finding clearly demonstrates the potential of MCR-ALS to improve separations. This work is particularly meaningful due to the inclusion of background contributions to the simulated data, which is often neglected in similar literature reports.
While we believe these results to be representative of expected experimental results, future work may include the validation of this work with the use of experimental data. By collecting experimental data under a range of conditions such as varying mobile phase composition and different analyte concentrations, the chromatographic resolution and $S/N$ can be controlled. Spectral similarity can be controlled through the careful selection of different analyte pairs. While significantly more involved than the simulated data sets used in Chapter 4, a smaller range of data (within the ranges used in the simulations) would be able to be used in order to validate the simulation results, rather than creating a full model.

The approach used in Chapter 4 can also be used as the basis for a comprehensive comparison between MCR-ALS and other curve resolution techniques such as PARAFAC and PARAFAC2. For this comparison, retention time shifting should be added as an additional parameter because these shifts are known to affect the performance of PARAFAC [4].

7.2. Reflections on Chapter 5

The second goal was addressed in Chapter 5. A new implementation of MCR-ALS was developed for LC x LC data in which data from two detectors were utilized to take advantage of the resolution of the 2D separation and the quantitative superiority of the $1^D$ separation. It was found that a clear advantage was present for the simulated data, particularly when dilution was significant. At $1^D$ peak resolutions of less 0.3, 2DALC provided a clear advantage. While no spectral shifts were incorporated in this data, the experimental data clearly showed that for the untargeted analyses, 2DALC did provide better quantitation for two of the three compounds. c2DALC was also investigated; however, it was found that even minor spectral shifts greatly affected the results and thus, we decided not to pursue c2DALC further. It is important to note
that while 2DALC was proven effective in this data, it may not always be superior to 2D or 1D chromatography alone, as seen in Chapter 5; however, the ability to obtain \( ^1 \text{D} \) and 2D chromatograms from an LC x LC analysis gives the analyst the ability to choose the best method for each analyte. Each of the three methods – \(^1 \text{D}-\text{IOPA}, \text{2D-IOPA}, \text{and 2DALC} \) – are relatively simple to run, giving the analyst the opportunity to evaluate which method is best for the given data. It is very possible that certain compounds that are not strongly overlapped in the \(^1 \text{D} \) may work better with \(^1 \text{D}-\text{IOPA} \) while other overlapped compounds in the same analysis may work better with 2DALC or 2D-IOPA, depending on the resolution in the \(^2 \text{D} \) separation.

The work in this chapter also showed two types of analyses: targeted and untargeted. The targeted analysis showed that when standards are available for all compounds, MCR-ALS was able to provide excellent calibration and prediction errors. Future work should investigate the intermediate case in which only some of the compounds are known prior to analysis. This would represent the case in which unknown compound spectra would need to be estimated from the raw data, while known compound spectra are also included in the analysis.

7.3. Reflections on Chapter 6

In Chapters 4 and 5, mostly simulated data were created in order to control chromatographic parameters such as \( R_5, S/N \), and spectral similarity. This allowed us to investigate the effect of each of these parameters individually, which would have been difficult to completely control with experimental data. In order to demonstrate the applicability to real-world analyses, we tested methods of MCR-ALS analysis on a targeted analysis of furanocoumarins from three apiaceous vegetables: parsley, parsnips, and celery. The LC x LC data were analyzed using the three methods outlined in Chapter 5: \(^1 \text{D}-\text{IOPA}, \text{2DALC}, \text{and 2D-}

IOPA. It was found that for this data set all three methods tested performed approximately equally in terms of the precision of quantitation. 2D-IOPA had the advantage that it was able to separate chromatographically overlapped, spectrally identical interferents from the target compounds, which MCR-ALS is unable to do. As discussed in Chapter 4, MCR-ALS requires a minimal level of spectral dissimilarity and chromatographic resolution in order to resolve overlapped analyte signals. Manual integration was also found to provide greater quantitative precision over the summation approach despite MCR-ALS removing most of the background. This suggests that the quantitative abilities in LC x LC are comparable to 1D-LC for these data when MCR-ALS is utilized. These findings are impressive when compared to many of the previous literature reports of quantitation in LC x LC. Because of this, we used 2D-IOPA with manual integration to quantify all 14 furanocoumarins. We were able to detect 10 of the 14 furanocoumarins at varying concentrations. Overall, the concentrations were low and often lower than the lowest calibration point; however, an estimate of concentration was able to be calculated. The concentrations of each furanocoumarin were found to vary greatly even within a single vegetable type. This may be due to vegetable freshness, specific plant species, geographical origin, or a plethora of other factors. It may also be due to natural, random variations, which would be a significant finding itself. Studies of correlations of the concentrations with exact sample information are ongoing. These samples will also be used to study the physiological effects of these compounds on rats. Although not performed in this work, peak detection may provide an alternative to manual integration that performs better than the summation approach, but still in a semi-automated fashion. The peak detection approach in Chapter 5 may work; however, selection of parameters for this approach (noise threshold,
window size, etc.) can still be tedious. Future work may provide a more automated peak detection method that can complement our work presented here.

7.4. Outlook and Future Work

It is clear from the work presented in this dissertation, along with the body of work in the literature, that MCR-ALS and other curve resolution techniques have the potential to greatly impact the field of separation science. While widespread adoption of these techniques has yet to occur, it is undeniable that these methods provide valuable tools for the analytical chemist.

While the identification power of diode array detection has been shown previously [63] and has been shown in the previous chapters to be particularly powerful in terms of quantitation using MCR-ALS, curve resolution with mass spectrometric detection must be explored due to the increasing adoption of both low and high resolution mass spectrometry. Currently in our laboratory and in a select few other labs, work is being performed towards this aim [109,148,149]. One major impediment to MCR-ALS analysis of LC-high resolution MS data is the immense size of the data. LC-DAD analysis collect UV-Vis spectra which typically contain 100-200 data points per spectrum corresponding to a wavelength range of approximately 190-600 nm sampled at 2-4 nm intervals. High resolution MS (HRMS) produces several orders of magnitude more data points. For example, a mass range of 100-1000 amu at a 0.001 amu precision, gives $9 \times 10^5$ data points per mass spectrum. In our lab, we implement a data reduction strategy that allows us to exclude many of these masses corresponding to background and noise [149]. By employing a binning strategy which initially reduces the precision to unit mass, MCR-
ALS can be performed to resolve masses corresponding to true chemical compounds and discard the remaining masses. Using this limited range of masses, a second binning process takes place, this time to 0.1 amu bins. MCR-ALS is performed on this data and once again any masses not corresponding to compounds are discarded. This process continues until the final precision level is reached as dictated by the instrument’s capabilities.

Automation should also be a key focus of curve resolution research moving forward. As with any technique, MCR-ALS can always be improved upon; however, if it is difficult to implement due to the need for experience and knowledge of computer programming, these techniques will never find widespread use. It is our belief that MCR-ALS is developed to a point that with automation of certain steps (component determination, application of constraints, etc) rather than optimization of the algorithm should be a major focus of research. Some work has been performed in other groups towards this goal, specifically on the determination of the number of components [150–152]; however, these methods can overestimate the number of components [97] and still may require several iterations of MCR-ALS with different numbers of components to find the optimal fit. While not completely automated, graphical user interfaces (GUI) have been developed for MCR-ALS analysis. One of the more popular MCR-ALS GUIs was developed by Jaumot, de Juan, and Tauler [118] which simplify the MCR-ALS process by eliminating the need for programming experience, but still requires a certain level of expertise and experience on the part of the user to determine the number of components and properly apply constraints. With further automation of these steps and subsequent commercialization of MCR-ALS packages, we believe MCR-ALS can become a standard tool in the analytical chemist’s arsenal.
List of References
<table>
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[144] R. Baleni, Z. Bekker, A. Walubo, J.B. Du Plessis, Co-administration of fresh grape fruit
juice (GFJ) and bergamottin prevented paracetamol induced hepatotoxicity after paracetamol overdose in rats, Toxicol. Reports. 2 (2015) 677–684. doi:10.1016/j.toxrep.2015.05.004.


Appendix A

The MCR-ALS analyses in this work was performed in MATLAB using personal code adapted from Robert Allen and Ernst Bezemer, previous graduate students in the Rutan group. This code is located in Virginia Commonwealth University’s network R drive (R:\CHEM\Rutan_lab\Dan\Dissertation\Cook_Toolbox) and is reproduced here.

als_DWCv5.m

This is the core function for applying MCR-ALS. It calls upon the function constrain_DWC.m which is located within the same directory. The inputs ig and options are obtained using the functions listed further in this Appendix.

function [r_opt,s_opt,IT,fit,conc_opt]=als_DWCv5(data,ig,iterations,options)
%Adapted from Robert Allen's als_Xv4
%data consists of 4 dimensions of data in the following order:
% 1st dim time, 2nd dim time, sample, spectra
%ig consists of spectra x components
%iterations is the maximum number of iterations
%nn is nonnegativity where the 1st row is concentrations and 2nd row is
% spectra
%u is unimodality in which the 1st row is 1st dim and 2nd row is 2nd dim
% u must also contain an extra column at the end specifying how to
% implement unimodality
%uTol is tolerance for unimodality
%ssel is chromatographic selectivity in the with the dimensions of:
% 1D x 2D x sample x component or 1D x sample x component
%ssel is spectral selectivity with the dimensions of:
% spectra x component

% Initialize data and options
if size(size(data),2)==3 & & size(data,2)>size(data,3)
    data=permute(data,[1 3 2]);
end

if size(size(data),2)==4 & & size(data,3)>size(data,4)
    data=permute(data,[1 2 4 3]);
end

nn=options.NN;
u=options.Uni;
uTol=options.Utol;
ssel=options.ssel;
csel=options.csel;
trilinearity=options.Trilinearity;
smoothness=options.Smoothness;

if exist('constrain_DWC','file') == 0
    error('Missing constrain_DWC file necessary to impose constraints')
end

if any(any(smoothness ~= 0)) && exist('whitsm','file') == 0
    error(['Missing whitsm function - needed for smoothing.'...
          'whitsm is part of Perfect Smoother'])
end

if numel(ssel)==1
    ssel=NaN(size(ig));
end

if numel(csel)==1
    csel=NaN(size(data));
end

if size(nn,2) == 1 %if only a single number, duplicate it for all components
    nn=ones(2,size(ig,2)).*nn;
end

if size(nn,1)==1 %If nn is only one row, duplicate it for second row
    nn(2,:)=nn(1,:);
end

if size(size(data),2)==3 % If the data is 1D data
    if size(u,2) == 1
        u=[ones(1,size(ig,2))).*u 1];
    end
end

if size(size(data),2)==4 % If the data is 2D data
    if size(u,2) == 1
        u=[ones(1,size(ig,2))).*u 1];
    end
    if size(u,1) == 1
        u(2,:)=u(1,:);
    end
end

%% Begin ALS
comp=size(ig,2);
S=ig;
md=3;
osq=inf;
bsq=inf;
dc=0;
ssqX=sum(sum(sum((data.^2))));

for n=1:size(size(data),2)
dimensions(1,n)=size(data,n);
end
reshapesize=prod(dimensions(1,1:size(size(data),2)-1));
refoldsize=dimensions(:,1:end-1);
if size(dimensions,2) ~= 3 && size(dimensions,2) ~=4
    error('dimensionality of data is incorrect')
end
datars=reshape(data,reshapesize,dimensions(1,end));

h= waitbar(0,sprintf('0 out of %d iterations',iterations));

for IT=1:iterations
    C=datars*pinv(S');
    C=reshape(C,[refoldsize,comp]);
    %Constrain concentration profiles
    for a=1:comp
        for b=1:dimensions(1,end-1) %samples
            %If this is LCxLC data, we need to constrain %dimensions differently
            if size(dimensions,2) == 4
                for c=1:dimensions(1,2)
                    C(:,c,b,a)=constrain_DWC(C(:,c,b,a),nn(1,a),u(1,a),... u(1,comp+1),uTol,csel(:,c,b,a),smoothness(1,a));
                end
            end
            % To constrain 1st dimension, uncomment the next 4 lines
            %for c=1:dimensions(1,1)
            %   C(c,:,b,a)=constrain_DWC(C(c,:,b,a),nn(1,a),u(2,a),... u(2,comp+1),uTol,csel(c,:,b,a),smoothness);
            %end
        end
        if any(trilinearity == 1)
            if size(dimensions,2) == 3
                [C,Ctrl,conc]=apply_trilinearity1D(C,trilinearity);
            else
                if IT == 1
                    warndlg(['Trilinearity constraint is not defined'... '4-way data. Press OK to continue without'... 'trilinearity','Warning:Trilinearity'])
                end
            end
        end
    end
    C=reshape(C,reshapesize,comp);
    S=(pinv(C)*datars)'; %Solve for spectral profiles
    %constrain spectral profiles
    for a=1:comp
        S(:,a)=constrain_DWC(S(:,a),nn(2,a),0,0,0,ones(size(ssel(:,a))),...
smoothness(2,a));
S(:,a)=S(:,a)/max(abs(S(:,a)));
S(:,a)=constrain_DWC(S(:,a),nn(2,a),0,0,0,ssel(:,a),...
      smoothness(2,a));

end

T=C*S';
res=datas-T;
ssq=sum(sum(sum(res.^2)));%
imp=(osq-ssq)/osq %Percent improvement
if IT==1
    imp=0;
    r_opt=C;
    s_opt=S;
    bsq=ssq;
end
if ssq<bsq
    dc=0;
    r_opt=C;
    s_opt=S;
    bsq=ssq;
end
if ssq<osq
    dc=0;
end
if ssq>osq
    dc=dc+1;
end
osq=ssq;
if dc>md  %If error increases 3 times, its divergent
    break
end
if abs(imp<1e-6)&imp>0
    break
end

waitbar(IT/iterations,h,sprintf(' %d of %d iterations',IT,iterations))
end

if exist('Ctril','var')  %If trilinearity was imposed
    r_opt=Ctril;
    conc_opt = conc;
end

r_opt=reshape(r_opt,[refoldsize,comp]);

fit=100*sqrt(bsq/ssqX);

waitbar(IT/iterations,h,sprintf('Completed!'))
pause(0.5)
close(h)

if size(ig,2)<5
    for n=1:size(ig,2)
function \([C\_out, C\_tril, conc]=apply\_trilinearity1D(C, components)\)

C\_out=C;
C\_tril=C;
tril\_comp = find(components == 1);
conc=NaN(size(C,2),size(C,3));

for k=tril\_comp %for each component
    [Cnorm,~,weights]=normalize(C(:,:,k),1);
    profile\_average=zeros(size(C,1),1);
    
    for j=1:size(C,2) %for each sample
        profile\_average = profile\_average + Cnorm(:,j)*weights(:,j);
    end
    profile\_average = profile\_average./sum(weights(:,j));

    C\_out(:,:,k)=profile\_average*weights;
    conc(:,k)=weights;
    C\_tril(:,:,k)=repmat(profile\_average,1,size(C,2));
end
end

**initialguess.m**

initialguess.m performs IOPA to extract a set number of spectra for use in als\_DWCv5.m. If the input variable ‘ncomponents’ is left blank a Scree plot is created and the user is prompted to choose the number of components. This function calls iopav2.m which is located in the same directory.
function [ig] = initialguess(data,ncomponents)
%Function will create initial guess for MCR-ALS using iopav2
%ncomponents is the number of components you wish to use, if left empty the
%program will perform SVD and allow you to choose based on Skree plot

if size(data,3)~=1
    data=reshape(permute(data,[1 3 2]),size(data,1)*size(data,3),... size(data,2));
end

%Create an initial guess using iopav2
if exist('ncomponents','var')==0
    [~,s,~]=svd(data,0);
    figure
    plot(log(diag(s)),'*')
    ncomponents=input('How many components are to be used?');
end
close
[ind_opt,max_det]=iopav2(data,ncomponents);
ig=data(ind_opt,:);
figure
plot(ig)
end

optionsALS.m

optionsALS.m creates the options structure for setting constraints in the als_DWCv5.m function. Inputting the number of components (NoC) prepopulates each field within the structure.

function [options] = optionsALS_edit(NoC)
%%
%The optionsALS function creates an options structure that is used for
% als_DWCv5

%INPUT:
% NoC is an optional input of the number of components to prepopulate
% some of the constraints with the correct number of elements

%nn is nonnegativity; it is input as a vector of 1 x number of components,
% with 1 setting component to non-negative and 0 leaving it unconstrained
% If one row is set, both concentration and spectra are constrained, else
% two rows can be defined, where 1st row= conc and 2nd row=spectra
%
%u is unimodality in which the 1st row is 1st dim and
% 2nd row is 2nd dim(if exists)
% u must also contain an extra column at the end specifying how to
% implement unimodality; 1 drops concentration to zero vertically, 2 cuts
% off horizontally, and 3 cuts intensity to half, which over the course
% of many iterations, sets second maximum to zero (3 is most common)
% uTol is tolerance for unimodality; 1 allow no increase after intial maximum, % 1.1 allows 10% increase, etc.
% csel is chromatographic selectivity in the with the dimensions of:
%   time x sample x component or 1D time x 2D time x sample x component
% ssel is spectral selectivity with the dimensions of:
%   spectra x component
% Trilinearity sets components to be trilinear. Row vector with sample
% number of elements as components. 1 sets trilinear, 0 leaves it
% unconstrained. Trilinearity is only defined at this point for 3-way
data
% Smoothness constrains components to be smooth, based on Eiler's perfect
% smoother. First row is concentration, second row is spectra

if ~exist('NoC','var')
    NoC=1;
end

field1='NN';
value1=zeros(1,NoC);

field2='Uni';
value2=zeros(1,NoC+1);
value2(:,end)=3;

field3='Utol';
value3=1;

field4='ssel';
value4=NaN;

field5='csel';
value5=NaN;

field6='Trilinearity';
value6=zeros(1,NoC);

field7='Smoothness';
value7=zeros(2,NoC);

options=struct(field1,value1,field2,value2,field3,value3,field4,value4,field5,value5,field6,value6,field7,value7);
end
Appendix B

The scripts here are the basic scripts used for the work in Chapter 4. These scripts and related functions are saved on Virginia Commonwealth University’s network R drive (R:\CHEM\Rutan_lab\Dan\Dissertation\Chapter 4).

MonteCarloALS.m

The script below was used for the design of experiments, simulation of data, and MCR-ALS resolution of the simulated data. In addition to the functions for MCR-ALS as listed in Appendix A, two additional functions were created and used in this script. They are reproduced in this Appendix as well.

```matlab
sfact=[12.54 6.27 0.627]';
position=[4.7 4.6 4.55]';
specind=[3 5 6]';

%create design of experiments
dFF=fullfact([size(sfact,1) size(position,1) size(specind,1)]);
clear DOJ
DOE(:,1)=sfact(dFF(:,1));
DOE(:,2)=position(dFF(:,2));
DOE(:,3)=specind(dFF(:,3));
%set the number of components to use for MCR-ALS
DOE(:,4)=3;
DOE(3:3:27,4)=5;
clearvars -except DOE
load(['C:\Users\CHEM_RUTANLAB\Documents\Dan\MatLab'...
   'MCR-ALSperformanceReview\18Jan16\Artificial_Spectra.mat'])
for k=1:50
    NoS=5; %number of samples in each dataset
    OptAll=struct;
    for i=1:size(DOE,1)
        [bkgd,time,wave]=genbkgd(NoS); %Generate random backgrounds
        OptAll(i).Opt=optSim(2); %Options for peak creation
        OptAll(i).Opt.Background=bkgd; % samples
        OptAll(i).Opt.Time=time;
        OptAll(i).Opt.Sigma(:)=0.05; % #peaks
        OptAll(i).Opt.Spectra=spectra(:,[DOE(i,3) 1]);
        OptAll(i).Opt.Position=[4.5 DOE(i,2)];
        OptAll(i).Opt.Intensity(1:NoS,1)=[28.2 21.9 25.1 23.5 26.6];
        OptAll(i).Opt.Intensity(1:NoS,2)=[0.1 0.25 0.5 0.75 1].*DOE(i,1);
    ```
end
clear bkgd

for i=1:size(DOE,1)
    %Simulate peaks
    [peaks,results(i).profiles,results(i).res]=peakSim(OptAll(i).Opt);
    results(i).spectra=OptAll(i).Opt.Spectra;
    results(i).intensity=OptAll(i).Opt.Intensity;
    %results(i).peaks=peaks;
    n=DOE(i,4);
    ig=initialguess(reshape(permute(peaks,[1 3 2]),size(peaks,1)*...  
                    size(peaks,3),size(peaks,2)),n);

    %number of components
    ALSopt=optionsALS(n);  %Options for MCR-ALS
    ALSopt.NN(1:2)=1;
    ALSopt.Uni(1:2)=1;
    ALSopt.ssel=NaN(n,113);
    ALSopt.ssel(40:end,1:2)=0;
    ALSopt.csel=NaN(size(peaks,1),size(peaks,3),n);
    ALSopt.csel([1:190 700:end],:,1:2)=0;
    ALSopt.Smoothness(1,3:end)=10e4;

    %Rearrange initial guess to match the order of the real spectra
    rearr=corrcoef(cat(2,ig,results(i).spectra)).^2;
    rearr=rearr(1:n,n+1:n+size(results(i).spectra,2));
    for j=1:size(rearr,2)
        [x,y]=find(rearr == max(max(rearr)));
        rearrx(y)=x;
        rearr(:,y)=0;rearr(x,:)=0;
    end

    %Rearrange ig to match spectra order
    x=1:size(ig,2);x(rearrx)=[];ig=ig(:,[rearrx x]);

    [results(i).r_opt,results(i).s_opt,~,results(i).fit]=...  
            als_DWCv5(peaks,ig,10000,ALSopt);

    %rearrange s_opt and r_opt to match real spectra order
    rearr=corrcoef(cat(2,results(i).s_opt,results(i).spectra)).^2;
    rearr=rearr(1:n,n+1:n+size(results(i).spectra,2));
    for j=1:size(rearr,2)
        [x,y]=find(rearr == max(max(rearr)));
        rearrx(y)=x;
        rearr(:,y)=0;rearr(x,:)=0;
    end
    x=1:size(results(i).s_opt,2);x(rearrx)=[];
    results(i).s_opt=results(i).s_opt(:,[rearrx x]);
    results(i).r_opt=results(i).r_opt(:,:,[rearrx x]);
    results(i).quan=squeeze(sum(results(i).r_opt));
end
eval(['results' num2str(k) ' =results;'])
clear results
end

**peakSim.m**

This function was created to simulate two chromatographic peaks on a real instrumental background. The input is an options structure defined using the optSim.m function reproduced below.

```matlab
function [peaks,profiles,res]=peakSim(options)
% INPUT
% options = an options structure defined by optSim program
%
%OUTPUT
%peaks = the output simulated chromatogram
%profiles = the pure peaks before background
%res = the resolution between the peaks

N=size(options.Spectra,2);
NoS=size(options.Intensity,1);

for n=1:N %Peaks
    temp=gausspeak(options.Time,options.Position(n),options.Sigma(n))*options.Spectra(:,n)';
    for k=1:NoS %Samples
        profiles(:,:,n,k)=temp.*options.Intensity(k,n);
    end
end
size(profiles)
peaks=squeeze(sum(profiles,3))+options.Background;

for j=1:N %peak 1
    for r=1:N %peak 2
        res(j,r)=2*abs(options.Position(j)-options.Position(r))/(4*options.Sigma(j)+4*options.Sigma(r));
    end
end
```

**optSim.m**

This function creates an options structure for use in peakSim.m.

```matlab
function [options]=optSim(N);

%INPUTS
% N = number of "compounds" to simulate

field1='Time';
value1=NaN;
```
field2 = 'Position';
value2 = NaN(1,N);

field3 = 'Sigma';
value3 = NaN(1,N);

field4 = 'Intensity';
value4 = NaN(1,N);

field5 = 'Spectra';
value5 = NaN;

field6 = 'Background';
value6 = NaN;

options = struct(field1, value1, field2, value2, field3, value3, field4, value4, field5, value5, field6, value6);
Vita

Daniel Wesley Cook was born on August 31, 1989, in Virginia Beach, Virginia and is an American citizen. He graduated from Salem High School, Virginia Beach, Virginia in 2011. He received his Bachelor of Science in Chemistry from Randolph-Macon College, Ashland, Virginia in 2011. After graduation he was employed as a stockroom assistant at Virginia Commonwealth University, Richmond, Virginia in the Department of Chemistry for one year. In the fall of 2012 he joined the chemistry PhD. program at Virginia Commonwealth University, where he joined the laboratory of Dr. Sarah Rutan in October of 2012. He has two first-author publications and has presented his research at three scientific meetings.

Publications


Presentations

*indicates poster presentation

