

Successful Flash Chromatography

A White Paper from Biotage



CONTENTS

3	Introduction
3	Normal Phase or Reversed phase?
3	Normal Phase
3	Normal Phase Solvents
3	Normal Phase Media
4	Reversed phase
4	Reversed Phase Media
4	Reversed Phase Solvents
5	Where to Begin – It's All About Chemistry
5	Solubility
5	So, to Summarize...
5	Compound Chemistry (Functionality)
5	Neutral, Acidic, Basic Compounds
5	Amines
6	Determining the Appropriate Stationary Phase
6	Dry loading
9	Method Development and Optimization
9	Normal Phase and Thin-layer Chromatography
9	Column Volume
9	Solvent Choices
10	Selectivity
10	Strength
10	Setting Purification Goals
11	Elution Modes
13	Organic Amines
13	Reversed Phase
13	Reversed Phase Column Conditioning
13	Method Development
14	Peptides
15	Scaling Up
16	Linear Velocity
18	Summary

Introduction

Flash chromatography is the preferred purification technique of organic, medicinal, natural product chemists, and more recently peptide chemists because it has the power to separate a broad variety of compounds more efficiently than other crude purification techniques such as crashing out of solution or liquid-liquid extraction. To deliver pure compounds, chemists can manipulate a large variety of variables to accomplish the desired level of purity. Here is a whitepaper which seeks to explain the factors that ensure successful purifications with flash column chromatography.

For organic and medicinal chemists, flash chromatography is simply part of synthesis workflow – *design* the synthesis, *perform* the synthesis, *purify* and *isolate* the desired product or intermediate. Generally, flash chromatography method development gets little thought and chemists rely on the “feel” gained through years of experience for how to purify their synthetic compound mixtures.

Chemists studying natural products face similar but different challenges. Instead of purifying and isolating milligrams to grams of a synthesized target compound, natural product chemists need to screen extracts for their specific needs. Flash chromatography helps by isolating groups of compounds to determine which have the desired activity. When a compound group shows a positive result, flash chromatography is used to separate the individual components for further testing.

Flash chromatography is commonplace in biochemistry research labs where it is used for peptide purification. Flash columns possess higher loading capacity than HPLC columns and purify peptides faster helping advance the development of novel peptide therapeutics.

What follows is information that can be put to use when purifications are more challenging.

Normal Phase or Reversed Phase?

The two dominant flash chromatography modes are **normal phase and reversed phase**. These modes differ in two ways – column media polarity and solvent polarity.

Normal Phase

Normal phase chromatography, sometimes known as straight phase chromatography, uses a polar column media such as silica with non-polar/moderately polar organic solvents. The separation mechanism is adsorption/desorption and compounds are retained based on their adsorptive affinity for the media and are desorbed with increasing solvent polarity. Hence, low polarity compounds elute earlier while those that are more polar are retained longer on the stationary phase.

Normal Phase Solvents

Typical normal phase solvent mixtures include hexane (or heptane) + ethyl acetate or dichloromethane + methanol. Other solvent mixtures are used when a different separation (aka alternate selectivity) is needed. Chromatography solvents are classified in various ways to assist with method development but the most frequently utilized are solvent strength and selectivity class, Table 1.

Table 1. Typical normal phase organic solvents and their relative elution strengths and selectivity classes.

Solvent	Polarity	Selectivity Class
Hexane	0.01	---
Heptane	0.01	---
Isooctane	0.01	---
Toluene	0.22	VII
Dichloromethane	0.32	V
Ether	0.40	I
Ethyl acetate	0.43	VIa
Acetone	0.50	VIa
Acetonitrile	0.51	VIb
THF	0.53	III
Isopropanol	0.60	II
Ethanol	0.68	II
Methanol	0.70	II

Normal Phase Media

While silica is the most commonly used stationary phase for normal phase flash chromatography others include:

- » Alumina
- » NH₂ – bonded silica
- » Diol – bonded silica
- » CN – bonded silica

Use of these other media is application specific and can often facilitate a separation or eliminate a mass-transfer issue when silica fails.

Table 2. Typical normal phase media, solvents, and applications.

Media	Solvents	Applications
Silica	Alkane + ethyl acetate or methylene chloride + methanol	Low to high polarity organic compounds
Alumina	Alkane + ethyl acetate	General organic purification, some organic amines
NH₂	Alkane + ethyl acetate	Organic bases (3°, 2°, N-heterocyclic)
CN	Alkane + ethyl acetate Alcohol + water	Polar organic molecules
Diol	Alkane + ethyl acetate Alcohol + water	Polar organic molecules

Reversed Phase

Reversed phase chromatography is essentially the opposite of normal phase as the name implies. In a reversed phase mode, the column media is non-polar (often silica bonded with an organic alkyl chain like C18) and the solvents are polar. The separation mechanism is solute partitioning between the media and the solvent (think liquid-liquid extraction). Compounds typically (but not always) elute in reversed (hence “reversed phase”) order compared to normal phase such that more polar compounds are retained less and elute ahead of less polar ones, Figure 1.

Reversed Phase Media

Reversed phase media include covalently bonded octadecyl silane (C18 or ODS), octyl silane (C8), butyl silane (C4) and other similar variations.

Reversed Phase Solvents

Solvent systems include water + methanol or water + acetonitrile. If sample components are ionizable or ionic, volatile pH modifiers (formic acid, acetic acid, TFA, ammonium hydroxide, etc.) and/or volatile buffers (ammonium formate, ammonium acetate, etc.) are beneficial and improve separation efficiency and peak shape.

Table 3. Typical reversed phase media, solvents, and applications.

Media	Solvents	Applications
C18 (ODS)	Water + methanol or acetonitrile. Add a volatile acid, base, or buffer if compounds are ionic or ionizable in solution	Low to high polarity organic compounds’ ionizable and ionic compounds; peptides
C8 (Octyl Silane)	Water + methanol or acetonitrile. Add a volatile acid, base, or buffer if compounds are ionic or ionizable in solution	Low to medium polarity organic compounds’ ionizable and ionic compounds; peptides
C4 (Butyl Silane)	Water + methanol or acetonitrile. Add a volatile acid, base, or buffer if compounds are ionic or ionizable in solution	Low polarity organic compounds’ ionizable and ionic compounds; peptides

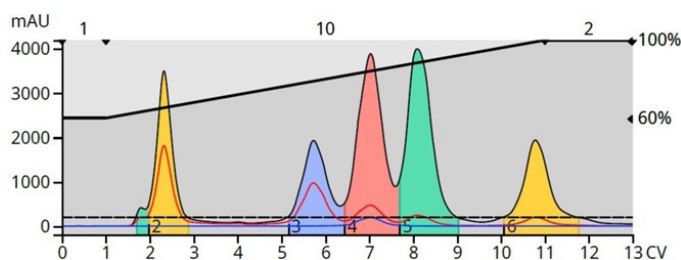
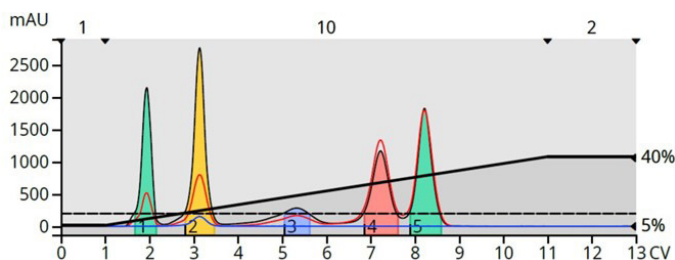


Figure 1. Comparison of normal phase (left) and reversed phase (right) chromatography using the same sample. Normal phase elution order: naphthalene, 1-nitronaphthalene, 3,5-dibenzoyloxyacetophenone, butyl paraben, methyl paraben. Reversed phase elution order: methyl paraben, butyl paraben, 1-nitronaphthalene, naphthalene, 3,5-dibenzoyloxy acetophenone.

Where to Begin – It's All About Chemistry

Getting the desired result with flash chromatography simply boils down to chemistry...

- » Sample solubility
 - » Which is the best solvent for the mixture being purified?
 - » Is the solvent used to dissolve the crude mixture compatible with the selection of elution solvents?
- » Targeted compound functional groups
 - » Is the target compound to be purified acidic, basic, or neutral?
 - » Could pH modifiers be needed to optimize mass-transfer kinetics during the separation?

Knowing these variables is a great start to a successful purification strategy. A great analogy to remember is...

Like Likes Like

Meaning that if your sample components are *polar*, they will be more attracted to a *polar* column media (i.e. silica). Likewise, if the compounds are non-polar they will be soluble in non-polar solvents and attracted to hydrophobic media (i.e. bonded silica).

Solubility

Reactions are often performed in an organic solvent but this solvent is not necessarily the best choice to load the crude reaction mixture onto a flash column. If a reaction is performed in a non-polar solvent such as dichloromethane, then normal phase purification using a silica column and a solvent system of an alkane (e.g. hexane) + ethyl acetate is likely the right choice because the reaction reagents have low to moderate polarity. One could argue the sample components in this example are non-polar, which is likely also true, and therefore should be purified using reversed phase. Absolutely right, reversed phase can be used but what about the sample's solubility? With reversed phase, compounds should be somewhat soluble in methanol or acetonitrile so they elute from the non-polar media. If not, reversed phase should not necessarily be ruled out, but compound solubility must be evaluated before proceeding.

So, to Summarize...

Mixtures containing non-polar, organic soluble target compounds are best purified with normal phase flash chromatography

Mixtures containing polar organic solvent and water-soluble target compounds are best purified with reversed phase flash chromatography.

Compound Chemistry (Functionality)

Neutral, Acidic, Basic Compounds

For neutral compounds, using solvents without the addition of modifiers is suitable, but what if your compounds have acidic or basic functionality? For separating target compounds that are acidic, adding a small amount of a volatile organic acid (formic, acetic) to the solvent system will help keep the acids protonated and minimize any secondary H-bonding interactions with the silica.

Amines

Organic amines, however, often require a competing amine in the solvent system. Ammonia (in methanol), ammonium hydroxide, and triethyl amine are among the most popular. Since silica is classified as a Brønsted acid, it attracts organic amines, sometimes to extremes causing peak broadening which can impede the ability to satisfactorily separate the mixture. The addition of an amine in the solvent helps to "neutralize" the silica surface, and will help not only with compound elution, but also with the compound's recovery or yield.

Organic soluble amines such as N-heterocycles, tertiary, and secondary amines can also be purified using [amine-functionalized silica](#). This type of modified silica eliminates or at least improves the solute-sorbent mass transfer kinetics and removes the need to add a competing amine.

Determining the Appropriate Stationary Phase

A comprehensive infographic to assist with determining the appropriate flash column stationary phase and loading technique is found in Figure 3.

To use the flow chart, begin with the upper section (green type), which focuses on solubility and sample chemistry. From this information, it is possible to decide whether normal phase or reversed phase chromatography is most appropriate for a given purification.

If it is determined that normal phase is most appropriate, then the column media should be selected based on the sample's chemistry. If the target molecule is basic (amine), an amine-bonded silica should be utilized (blue path), but if the molecule is neutral or acidic, then unbonded silica is the best choice (red path).

If the crude mixture containing the target compounds of interest is most soluble in a polar solvent, then reversed phase is an appropriate choice (grey path). Also, if the target compound is of high molecular weight (>1000 da), then using a wide pore reversed phase media (~ 300 Å) can work successfully (light grey).

Solvent polarity has a major impact on separation performance so using the infographic above is recommended. Additionally, the information in Table 4 is helpful to determine the best option for loading the crude mixture onto the column, knowing if a liquid load is practical or if dry loading should be used.

Dry Loading

Dry loading is a sample introduction technique with which a sorbent, Table 5, with the dissolved sample and then the solvent is evaporated. Dry loading has its advantages which include:

- » Improved chromatographic separation
- » Increased sample loading
- » Potential impurity removal

Table 4. Sample dissolution solvent and flash chromatography mode options. Solubility in some solvents may require dry loading as indicated.

Rank	Solvent Polarity	Normal Phase	Recommend Dry Loading	Reversed Phase	Recommend Dry Loading
1	Dimethyl sulfoxide (DMSO)			x	
2	N,N-Dimethylformamide (DMF)			x	
3	Water			x	
4	Methanol (MeOH)			x	
5	Acetonitrile (MeCN)			x	
6	Ethanol (EtOH)			x	
7	Propanol (PrOH)	x	x	x	x
8	Acetone	x	x	x	x
9	Tetrahydrofuran (THF)	x	x	x	x
10	Ethyl acetate (EtOAc)	x	x	x	x
11	Ether	x		x	x
12	Methyl chloride (DCM)	x	x	x	x
13	Hexane/Heptane/Alkane	x	x	x	x

If the crude mixture is soluble in a strong solvent (one that interferes with compound concentration at the head of the column), dry loading is required or a poor separation outcome (and low target compound purity) can result, Figure 2.

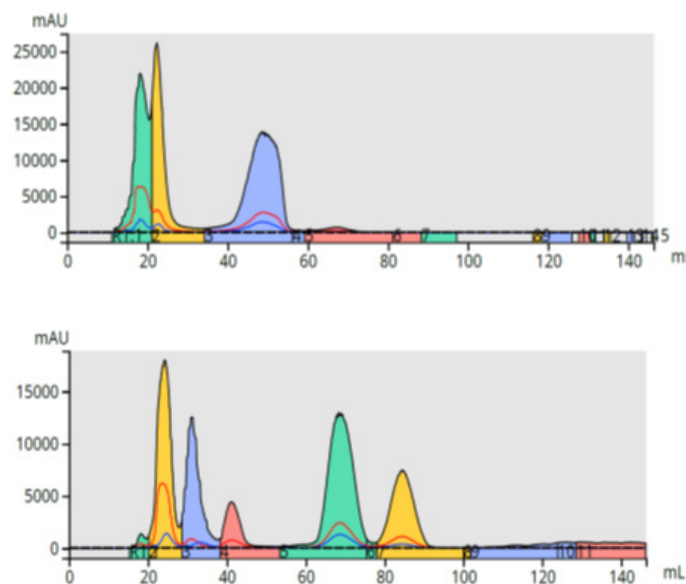


Figure 2. Purification of a reaction mixture using liquid loading (top) and dry loading (bottom). Dry loading removed polar solvent from the reaction mixture enabling more retention and an improved purification.

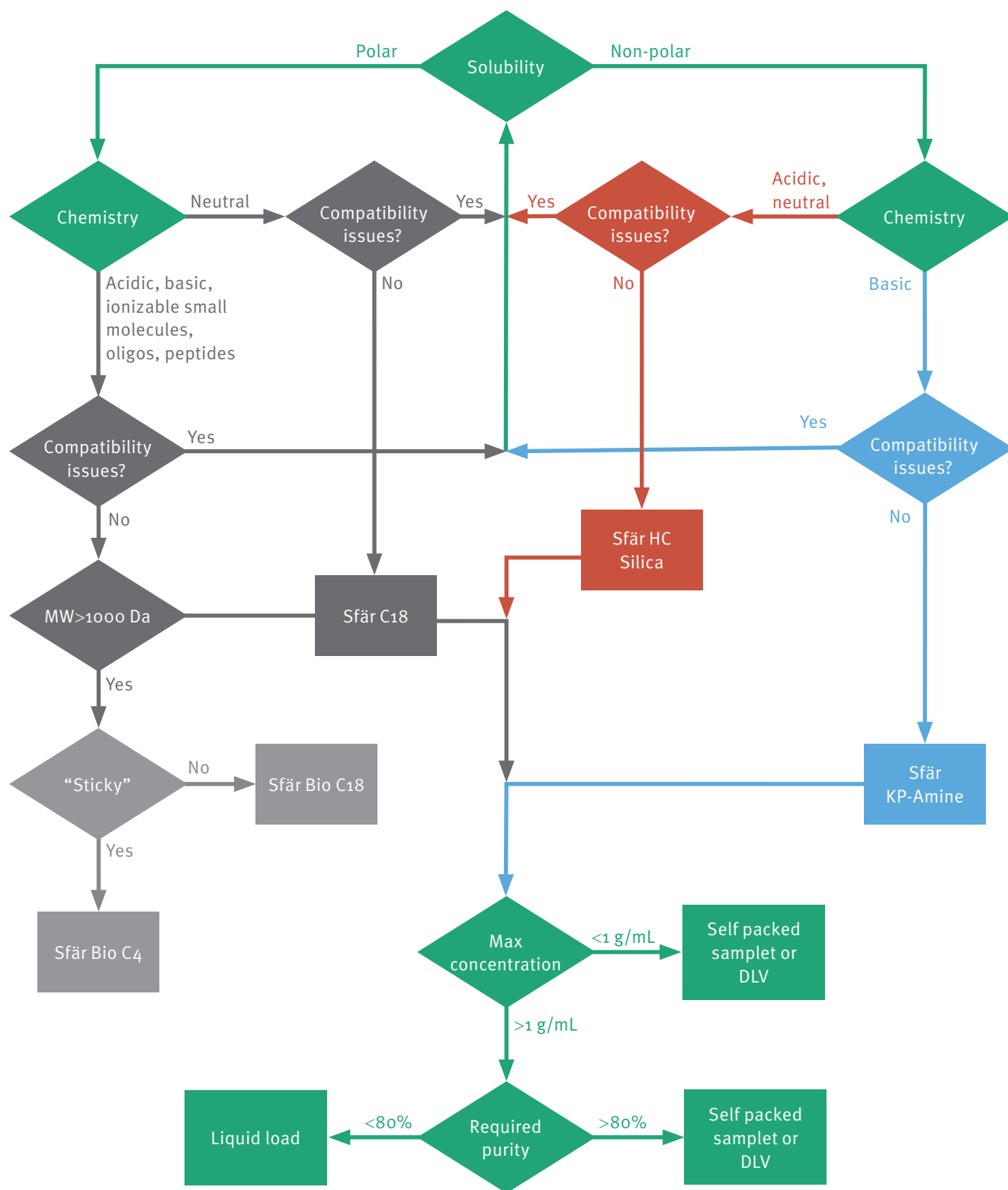


Figure 3. A stationary phase decision matrix to assist with choosing the correct flash column and loading technique.

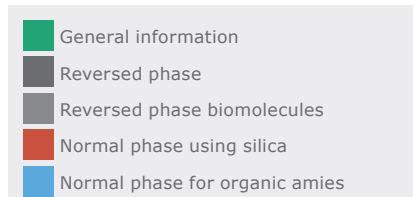


Table 5. Dry loading media options and their application areas.

Media	Mode	Application
Silica	Normal phase	General purpose, will bind highly polar compounds
Reversed Phase	Reversed phase, Normal phase	General purpose, will bind lipophilic compounds in reversed phase but will release all but the most hydrophobic compounds in normal phase
Diatomaceous Earth (Celite®)	Reversed phase, Normal phase	General purpose, inert, will release all loaded compounds
Alumina	Normal phase	General purpose, less non-specific binding than silica
Florisil®	Normal phase	General purpose, less non-specific binding than silica
Amine-bonded Silica	Reversed phase, Normal phase	Acts as an acid scavenger; good choice for normal phase organic amine sample loading
Ion Exchange	Reversed phase, Normal phase	Acts as ion scavenger (remove ionic/ionizable impurities) or as a catch & release media (concentrates ionic/ionizable component of interest and releases compound later)

Dry loading also has other uses including impurity removal or scavenging. Ion exchange media is one option for selectively attracting ionic/ionizable compounds (i.e. undesired reaction by-products) that co-elute with the target compound purification.

Dry load sorbent selection is important. Commonly used to remove unwanted particulates from crude mixtures, diatomaceous earth is a media that is chemically inert and allows for the entire sample to elute (think of this material as a sponge). Other media can be chemically interactive and may be useful in removing some impurities (a secondary benefit) or help increase target compound concentration.

Method Development and Optimization

Once a decision has been made as to which column media and purification mode (normal- or reversed phase) will work best for the flash separation, the next step is to determine the best separation conditions. The separation method conditions might include the solvent blend ratio and modifiers.

Normal Phase and Thin-layer Chromatography

Thin layer chromatography (TLC) is one of the better tools applied to developing purification methods. This technique applies to silica, alumina, and polar bonded silica (NH₂, diol, CN). With TLC, it is easy to evaluate various solvents in different ratios in parallel in order to determine which provides the best separation.

In normal phase chromatography, separations occur via an adsorption – desorption mechanism. Retention is based on compound polarity where polar compounds have greater affinity for the TLC sorbent than non-polar compounds. Compounds retain on the media until the elution solvent, otherwise known as the mobile phase, desorbs them. Once desorbed, the compounds do not resorb and will migrate unimpeded with the mobile phase.

With TLC, compound retention is determined by the ratio of the compound migration distance divided by the solvent migration difference and is known as R_f (retardation factor). R_f values range from 0 to 1 and correlate well to compound retention in flash chromatography.

$R_f = \text{Compound migration distance} / \text{Solvent migration difference}$

Column Volume

In flash chromatography, compound retention is measured in terms of **column volumes** (CV) rather than time as with gas chromatography (GC) or high performance liquid chromatography (HPLC). The reason for this is 2-fold...

- Using column volumes simplifies any scaling up of flash chromatography methods by eliminating the need to know the column size and flow rate.
- The number of column volumes required to elute a compound from a flash column is proportional to the reciprocal of the compound's R_f value when using the TLC solvent blend, Table 6.

$$CV = 1/R_f$$

Table 6. The CV/R_f relationship.

R _f	CV
0.1	10.00
0.2	5.00
0.3	3.33
0.4	2.50
0.5	2.00
0.6	1.67
0.7	1.43
0.8	1.25
0.9	1.11
1.0	1.00

The value of using CV for flash chromatography retention measurement is that it is independent of flow rate. Each column size has its own specific CV, which is the empty space within a packed column. Column packing materials are porous so a column volume is equal to the media's porosity plus the interstitial volume (volume between the media particles).

Solvent Choices

Solvents are classified in terms of **selectivity** and **strength**. Selectivity relates to a solvent's ability to separate a pair of compounds while strength deals with retention. Elution methods use a binary mixture of a weak solvent (low strength) and a strong solvent (higher strength than the weak solvent).

Typical flash chromatography solvents are hexane (weak) with ethyl acetate (strong) and dichloromethane (weak) with methanol (strong). To increase compound retention, the solvent blend can be weakened and to decrease retention, the solvent blend can be strengthened.

Table 7. Common flash chromatography solvents by selectivity group and strength.

Solvent	Group	Strength
Methanol	II	0.70
Ethanol	II	0.65
Propanol	II	0.60
Tetrahydrofuran	III	0.53
Acetone	VIa	0.50
Acetonitrile	VIb	0.50
Ethyl acetate	VIa	0.43
Ether	I	0.40
Dichloromethane	V	0.32
Toluene	VII	0.22
Heptane	---	0.01
Hexane	---	0.01

Selectivity and strength are independent variables. When developing methods it is best to evaluate solvents with different selectivity first followed by a strength adjustment. The selected solvents must be miscible, of course, in order to facilitate a separation.

Selectivity

Selectivity is measured in terms of ΔCV , which is the number of CV between two adjacent compounds on the TLC plate. The larger the ΔCV , the better the separation.

$$\Delta CV = CV_1 - CV_2$$

Strength

To maximize ΔCV , it is best to set TLC elution conditions where the target compound has an R_f value between 0.15 and 0.4. This is accomplished by adjusting the strong solvent/weak solvent ratio. R_f values decrease when the weak solvent percentage increases, Figure 4.

In this example, a strong solvent blend (left) separates the two compounds on the TLC plate but because the R_f values are high (low retention) there is little retention and minimal separation using flash chromatography.

Changing the TLC solvent mix so that it is weaker allows the two components to retain better on the plate, which decreased their respective R_f values and improved the flash chromatographic separation (middle and right).

Setting Purification Goals

With this background information, it is possible to begin to examine various solvent blends by TLC to see which will provide the best separation. However, it is up to the chemist to determine the criteria for a successful purification.

With any chromatographic separation, there are three competing variables – speed (flow rate), load (yield), resolution (purity), known as the Chromatographer's Triangle, Figure 5.

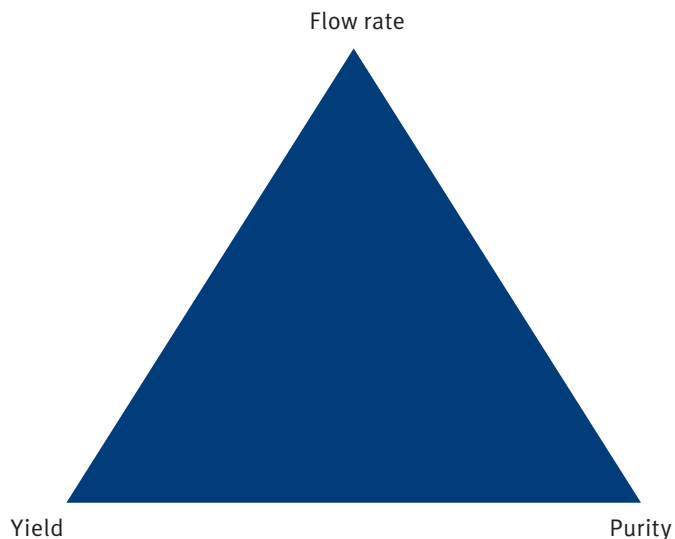


Figure 5. The Chromatographer's Triangle asks chemists to determine which of the three variables is most important; only two can be optimized.

Before starting on method development, one should know what outcome(s) the purification must achieve – highest purity, highest yield, fastest elution – or any combination of two of these. For synthetic chemists, optimizing purity and/or yield

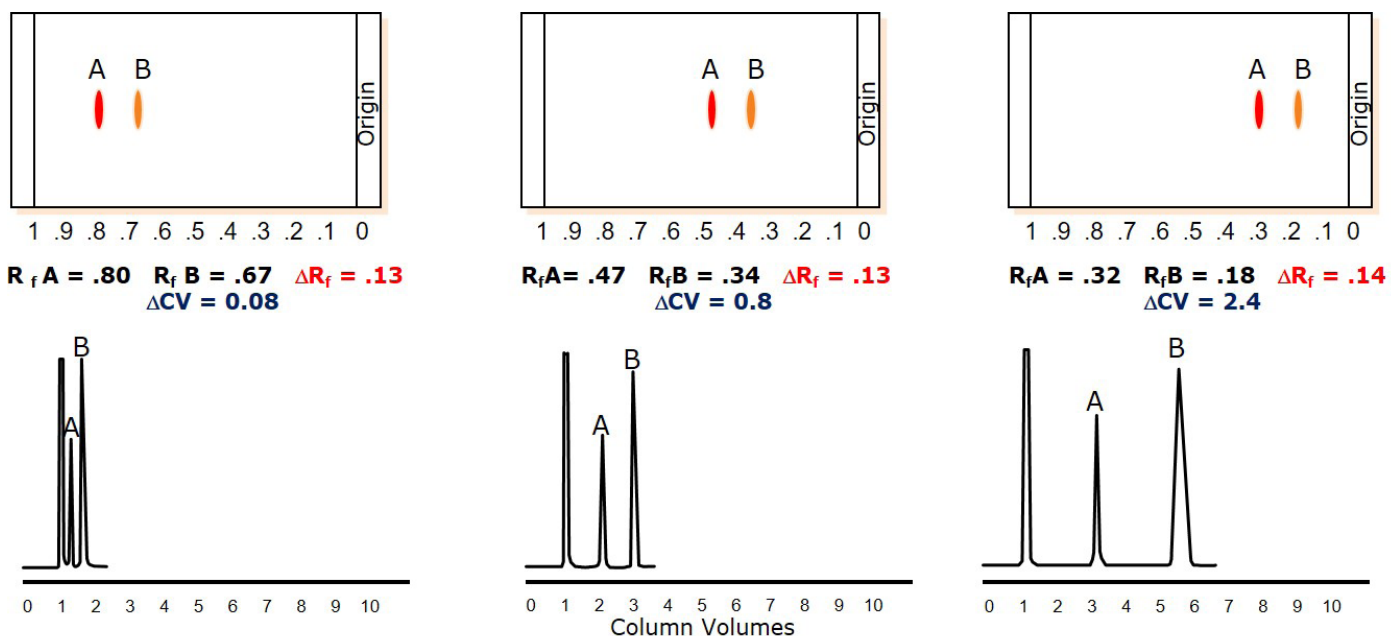


Figure 4. Solvent strength impact on separation performance. Left - strong solvent mixture elutes compounds with high R_f provides minimal separation ($\Delta CV = 0.08$), Middle - acceptable solvent strength (one compound within R_f range 0.15 - 0.4) provides a good separation ($\Delta CV = 0.8$), Right - optimal solvent conditions for maximum load ($\Delta CV = 2.4$).

is most important depending on whether the target compound to be purified is an intermediate (yield > purity > speed) or final product (purity > yield > speed).

Most normal phase methods use an alkane solvent (hexane, heptane, isooctane, or cyclohexane) and ethyl acetate. For more polar compounds, dichloromethane and methanol are most often used. Since these solvent mixtures are not always optimal, some experimentation with TLC and other solvents is recommended.

In this example, a 5-component mixture is separated by TLC with three different solvent selectivity classes – hexane/ethyl acetate (class VI), toluene (class VII), and dichloromethane (class V). The TLC solvents all separate the targeted compound (middle spot) but elute it at different R_f values and with different ΔCV from its nearest impurities. With this example, 100% toluene actually provides the best separation with a maximum ΔCV and an R_f within the optimal range of 0.15–0.40, Figure 6.

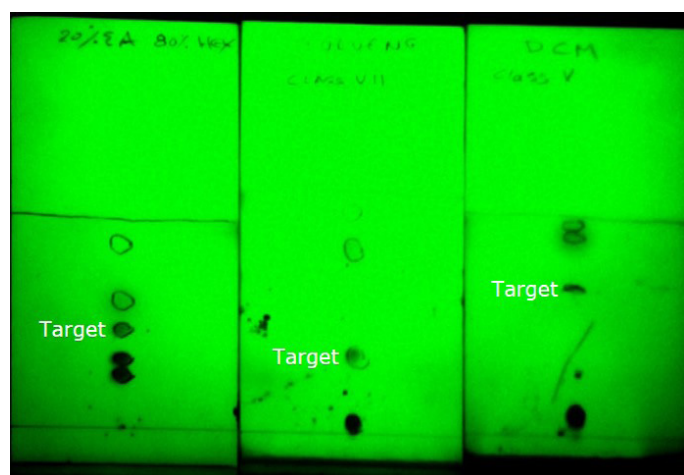


Figure 6. Solvent choice affects the separation selectivity of a 5-component sample mixture. Left - 20% EtOAc/hexane separates the target from its nearest impurities but with a higher R_f and lower ΔCV than what is ideal. Middle - 100% toluene fully isolates the target compound from the other four components with a suitable R_f value, Right - 100% DCM also provides a suitable separation but the target has a high R_f value.

Once a suitable solvent system has been found to separate the targeted compound from the other sample components, it is fine to move forward with using flash chromatography.

Elution Modes

Isocratic

As previously mentioned, TLC methods translate to flash chromatography well when the same solvents are used at the same ratio. This is an elution mode known as *Isocratic*. With isocratic elution, the mixture's components will elute in a predictable number of column volumes (refer to Table 6). For example, if a compound has an R_f of 0.2, it will require 5 CV to elute. Likewise, a compound with an R_f of 0.4 requires 2.5 CV to elute.

Isocratic methods are very good with simple purifications but low R_f (well retained) compounds elute in broad bands diluted in a lot of solvent, Figure 7. This can negatively affect detection, purity, and yield.

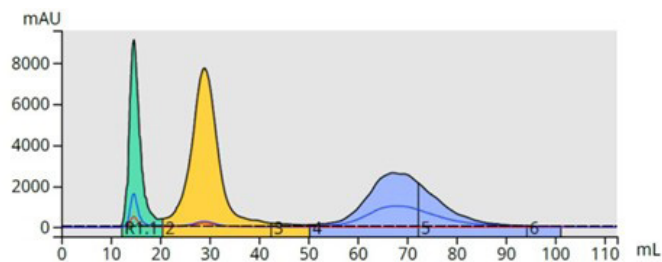


Figure 7. Isocratic elution method separated the three compounds but with major increases in peak volume which can impact sensitivity, purity, and yield.

Linear Gradient

Purification improves when the TLC method is converted to a *linear gradient*. With a linear gradient, the mass-transfer or adsorption-desorption kinetics are altered during the purification process. At the beginning of the gradient, a solvent ratio weaker than the TLC solvent blend is used but is changed, linearly, over time by increasing the strong solvent percentage. Early eluting (high R_f) compounds are retained more and separated better than with an isocratic elution and lower R_f compounds elute in higher concentration which enables detection and fractionation, Figure 8.

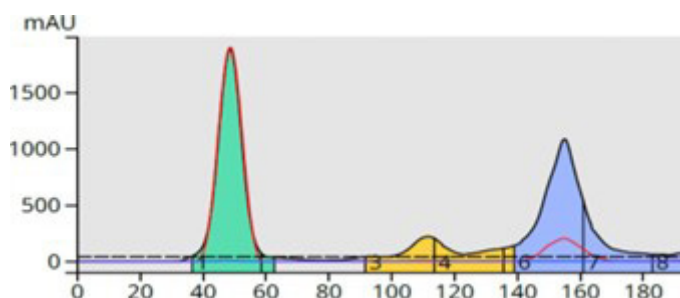


Figure 8. Linear gradient purification increased the separation between compounds 1 and 2 and eluted compound 3 in less volume thereby increasing its sensitivity and recovery. Also, an impurity has become visible between peaks 2 and 3 that was not easily seen with the isocratic run.

The gradient slope influences both the separation quality and purification time so this needs evaluation. However, a good, reliable algorithm to use when converting a TLC method to a linear gradient is...

$$\frac{1}{4} \text{ of the TLC strong solvent \% , hold 1 CV}$$

$$\frac{1}{4} \text{ TLC strong solvent to } 2x \text{ strong solvent \% over 10 CV}$$

$$2x \text{ TLC strong solvent \% , hold 2 CV}$$

Therefore, when we put the algorithm to use, a TLC method using 20% ethyl acetate (strong solvent) with 80% hexane converts to the following linear gradient method, Figure 9.

- » 5% ethyl acetate for 1 CV
- » 5% to 40% ethyl acetate over 10 CV
- » 40% ethyl acetate for 2 CV



Figure 9. TLC-based linear gradient. The TLC strong solvent % was 30 which converts to a linear gradient of 7-60% over 10 column volumes (CV).

Linear gradients using this algorithm to elute all compounds with an Rf of 0.1 and higher.

Step Gradient

There are times where a linear gradient fails to achieve the purification goals due to a less than ideal separation between the target compound and its nearest impurity, or a poor load capacity, or possibly to address a desire to less solvent and be a bit greener. In these cases, a [step gradient](#) may be the better option.

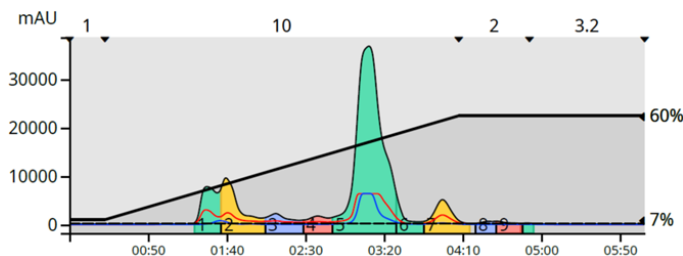


Figure 11. Using a step gradient provided a separation of the trailing shoulder from the product and reduced run time by over 30% as well.

Step gradients are comprised of a series of isocratic segments. Methods built with discrete steps can provide selective compound elution, increase sample load, and decrease solvent consumption. Figure 10 shows how a step gradient shortened the separation and reduced the volume between eluting peaks all while improving the overall separation compared to both the isocratic and linear gradient runs.

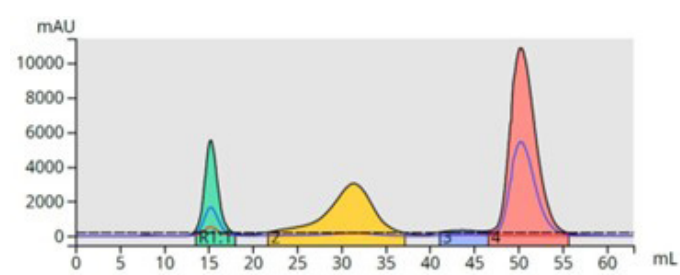


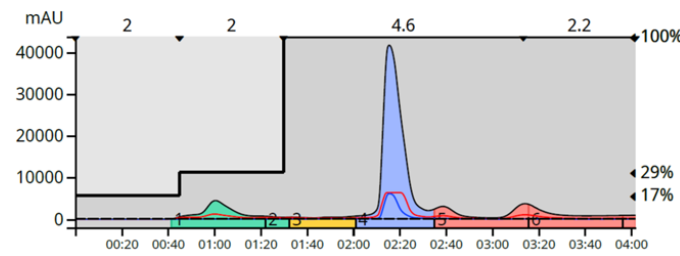
Figure 10. Step gradient elution optimized the separation of the three major compounds and the by-product and used less solvent and time than the other methods.

Step gradients can be challenging to develop, however. In the above example, after some trial and error, a gradient starting with 20% strong solvent (TLC was 30%) and increasing in 10% steps of two CV each was successful. Other separations will likely require different steps, both in height and length. So, often these gradients are most effective when a large amount (>10 grams) of material needs purification.

TLC can assist with step gradient building as well but this process does require a flash system suitably equipped with a “TLC to step gradient” function.

The separation, solvent consumption, and sample load all typically improve with a step gradient, Figure 11.

With a step gradient, purification strategy is optimized, and the trade-offs mentioned earlier with the Chromatographer’s Triangle, are minimized.



Organic Amines

Organic soluble amines pose a different challenge to chemists, especially if they have basicity. Lewis bases interact strongly with silica gel, which is a weak acid, making them difficult to elute and separate without introduction of a competing amine to the solvent system as a modifier. A competing amine helps to neutralize the acidic silica surface, which enables the basic amines to elute.

The typical solvent system used to purify these types of compounds on silica is a dichloromethane (DCM)/methanol (MeOH) gradient incorporating a small amount (on the order of 0.05 to 0.2%) competing amine (ammonium hydroxide, ammonia in MeOH, triethylamine, diethylamine, among others). Optimizing the separation conditions, however, still can be challenging as small changes in mobile phase polarity (MeOH or amine modifier) can often cause target amine compounds to elute very quickly without separating.

Other organic purification options include alumina and **amine-functionalized silica**. While alumina is a more amine-friendly media than silica, its large particle size (~150 μm) minimizes its usefulness. Amine-functionalized silica, on the other hand, is a silica bonded phase (~50 μm), and provides better separations.

The amine functionality takes the place of the competing amine in the mobile phase and simplifies the chromatography, Figure 12. Usual mobile phases include hexane/ethyl acetate or ethyl acetate/IPA for polar amines.

Reversed Phase

Prior to using a new reversed phase flash column for method development or purification, it needs conditioning. Why? Because reversed phase flash columns are dry and require pre-conditioning or wetting with solvents in order for them to function properly.

Reversed Phase Column Conditioning

The wetting process is simple and quick entailing only three steps.

1. Flush the column with 100% organic solvent (ACN or MeOH) for 3-5 column volumes (CV)
2. Flush with 50% organic solvent (ACN or MeOH)/50% aqueous for 3-5 column volumes (CV)

3. Flush with your initial gradient blend for 3-5 column volumes (CV)

Once a column is conditioned, this process does not need repeating.

Method Development

Reversed phase method development usually does not involve TLC for a variety of reasons including TLC media wettability limitations, sample loading limitations, and mass-transfer kinetics. It is best to develop reversed phase flash chromatography methods with an HPLC column (preferably packed with the same flash chromatography media as your flash column) or on a reversed phase flash column through trial and error. This process need not be long and tedious but does require an understanding of the potential ionizability your target compound and by-products may possess in solution.

Neutral compounds are separable using gradients of either water + methanol (MeOH) or water + acetonitrile (ACN). **Ionic and ionizable compounds**, however, require addition of either an organic buffer or pH modifier (acid or base) to the mobile phase to suppress ionization and improve the solutes' mass transfer kinetics, Table 8.

Table 8. List of solvent modifiers for ionic and ionizable compounds.

Additive	Concentration	Application
Formic acid	0.1%	Organic acids, ionizable hydroxyl groups, peptides
Acetic acid	0.1%	Organic acids, ionizable hydroxyl groups, peptides
Trifluoroacetic acid	0.1%	Organic acids, ionizable hydroxyl groups, peptides
Ammonium Hydroxide	0.1%	Organic amines, Lewis bases
Triethylamine	0.1%	Organic amines, Lewis bases
Ammonium formate	100 mM	Ionic compounds, zwitterions
Ammonium acetate	100 mM	Ionic compounds, zwitterions

With reversed phase chromatography, the separation mechanism is solute partitioning between the mobile phase and the non-polar stationary phase. Compounds retain on the stationary phase and elute based on their hydrophobicity. Ionic or ionizable compounds also follow this mechanism but their ionization level, Figure 13, influences retention and peak shape.

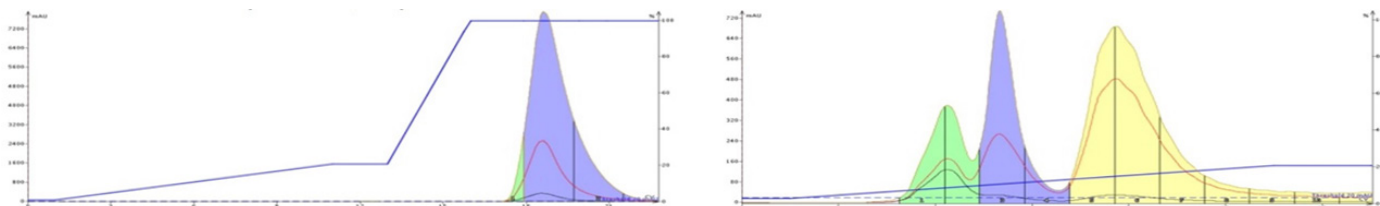


Figure 12. Purification of three tricyclic amines using silica with a DCM/MeOH gradient (left) and an amine bonded silica column (Biotage® Sfär KP-Amine) with a hexane/EtOAc gradient (right).

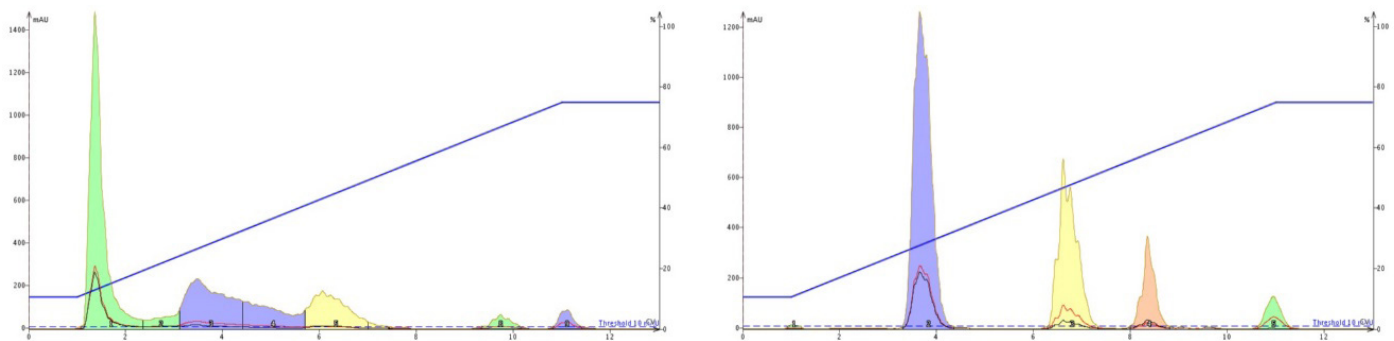


Figure 13. Reversed phase flash chromatography showing the impact of mobile phase buffering on a mix of ionic food colorings. Left – no added buffer. Right – 100 mM ammonium acetate, pH 7, added to mobile phase during the gradient. Compound elution order – yellow 5 (tartrazine), red 40 (allura red AC), blue 1 (brilliant blue FCF), red 3 (erythrosine).

For most sample mixtures, reversed phase compound elution order is exactly opposite that of normal phase (hence the term “reversed phase”). As mentioned earlier, with reversed phase, compounds elute based on the differences in hydrophobicity. The normal phase chromatographic mechanism is adsorption/desorption and not always based on polarity differences. This is an important distinction because if you need to convert a normal phase method to a reversed phase method (or vice-versa), the reversed phase elution order may not always be the exact reverse of the normal phase elution order, Figure 14.

Peptides

Peptide purification using reversed phase flash chromatography is gaining in popularity due to the improved loading capacity resulting from high performance silica columns, purification speed and simplicity, and cost compared to preparative HPLC. **High performance flash columns** designed specifically for peptides (20 µm particles with ~300Å pores) provide comparable performance to prep HPLC at a lower cost.

To ensure the peptides retain and separate, shallow gradients of water, ACN, and an **acid** (formic, acetic, trifluoroacetic (TFA)) are common. The added acid ensures the peptides are protonated, which helps with their retention and selectivity.

With most peptide syntheses, there are impurities, many of which are deletion sequences missing one or more amino acids compared to the synthesized peptide. Because the peptides are larger in size (>1000 MW) compared to other small molecules (up to 1000 MW), the deletion sequences often have similar hydrophobicity to the main peptide making purification challenging, Figure 15.

Incorporating the acidified, shallow gradients with high-performance flash columns with larger pore diameter (~300Å) does improve purification efficacy in terms of both peptide purity and yield.

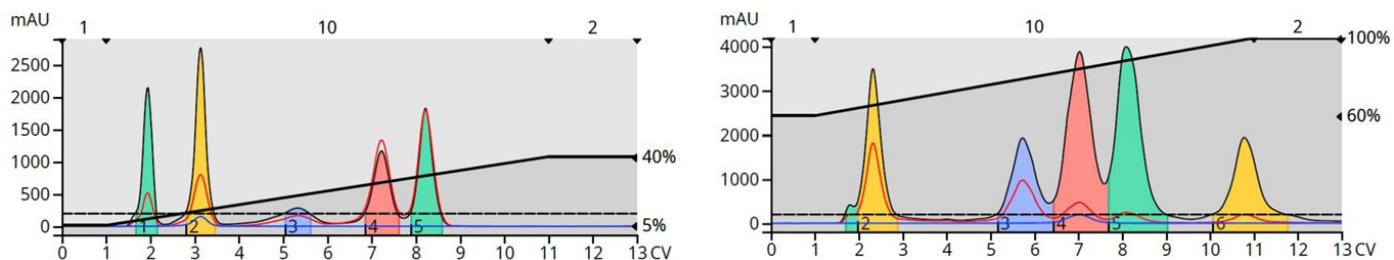


Figure 14. Example of non-predictable reversed phase elution order based on normal phase elution order. Left – normal phase separation (elution order naphthalene, 1-nitronaphthalene, 3, 5-dibenzoyloxyacetophenone, butyl paraben, methyl paraben). Right – reversed phase separation (elution order methyl paraben, butyl paraben, 1-nitronaphthalene, naphthalene, 3, 5-dibenzoyloxyacetophenone).

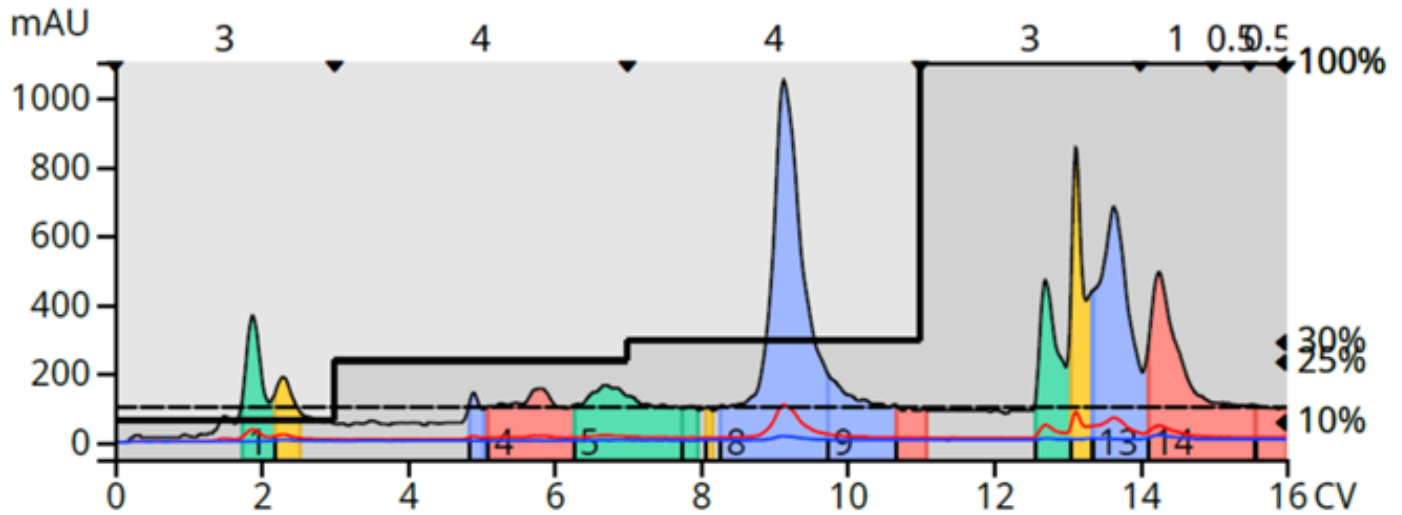


Figure 15. 10-mer peptide purification using a 10-gram Sfär Bio C18 flash column and an acetonitrile/water/formic acid step gradient. The peptide eluted at ~ 5:30.

Scaling Up

Scaling purification methods is not difficult as long as important parameters are maintained.

1. Sample concentration
2. Sample dissolution solvent
3. Sample introduction method (liquid/dry)
4. Column stationary phase/media
5. Solvent flow linear velocity

The variable to be controlled is flow rate. All of the other parameters listed are scalable based on column volume or column media mass ratios. These ratios or scale factors (SF) are easily determined.

$$SF = CV(Lg)/CV(Sm)$$

or

$$SF = (\text{Column Mass}(Lg))/(\text{Column Mass}(Sm))$$

Where Lg = the larger column and Sm = the smaller column.

As an example, a 25-gram column has 2.5 times the silica and volume that a 10-gram column has so the scale factor is 2.5.

$$SF = 25/10 = 2.5$$

For scale up this means that 2.5 times more sample mass can be purified using a 25-gram column compared to a 10-gram column.

Figure 16 shows a direct, linear scale up of a 7-component mixture using scaling factors. The 12-gram cartridge scale factor is 1, since that is the starting point or small column. The 60-gram column has a scale factor of 10 and the 400-gram column has a scale factor of 33.3. Table 9 provides scale factors from 5-gram to 40 kilogram columns.

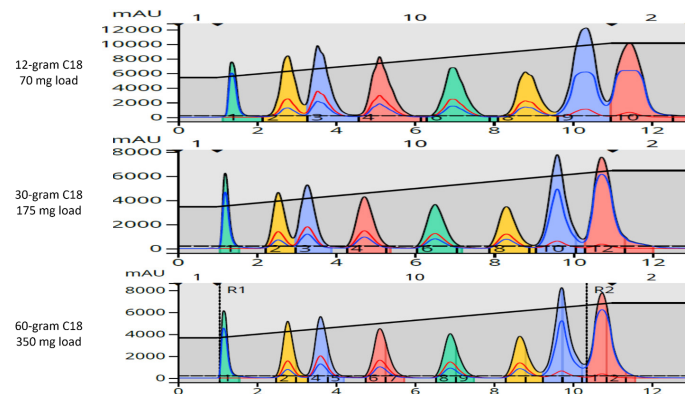


Figure 16. Direct, 10-fold purification scale-up. Top - 35 mg load on a 6-gram Sfär C18. Bottom - 350 mg load on a 60-gram Sfär C18.

Table 9. Purification scale factors from 5 g to 40 kg columns.

Column Size (g)	5	10	25	50	100	200	350	750	1500	2500	5000	20000	40000
5	1	2	5	10	20	40	70	150	300	500	1000	4000	8000
10		1	2.5	5	10	20	35	75	150	250	500	2000	4000
25			1	2	4	8	14	30	60	100	200	800	1600
50				1	2	4	7	15	30	50	100	400	800
100					1	2	3.5	7.5	15	25	50	200	400
200						1	1.75	3.75	7.5	12.5	25	100	200
350							1	2.1	4.3	7.1	14.3	57	114
750								1	2	3.3	6.6	27	54
1500									1	1.7	3.4	13.3	26
2500										1	2	8	16
5000											1	4	8
20000												1	2
40000													1

Linear Velocity

As previously mentioned, flow rates are not directly scalable so to maintain the same mass transfer kinetics, and therefore, separation performance, it is important to maintain solvent flow linear velocity. To determine the linear velocity, there are two calculations required.

The first equation determines the time required for a non-retained compound to elute and requires knowledge of the column's volume or CV. The CV values are often provided by column manufactures.

$$\text{Void time (min)} = (\text{Column volume (mL)}) / (\text{Flow rate (mL/min)})$$

The second calculation determines the linear velocity in cm/min and uses the column's packed bed depth in the equation. Most column manufacturers supply this information as well. Table 10 provides a listing of Biotage flash cartridges (silica), their column volumes, and packed bed depths.

$$\text{Linear velocity (cm/min)} = (\text{Bed depth (cm)}) / (\text{Void time (min)})$$

To make use of the linear velocity data, ensure the large column provides a similar linear velocity as used with the small column

Small column = Biotage® Sfär HC 10 gram with a CV of 17 mL and a bed depth of 5.5 cm. If the method ran at 20 mL/min, this column's void time was

$$\text{Void time} = (17 \text{ mL}) / (20 \text{ mL/min}) = 0.85 \text{ min}$$

and its linear velocity was

$$\text{Linear velocity} = (5.5 \text{ cm}) / (0.85 \text{ min}) = 6.47 \text{ cm/min}$$

If scaling up to a 100-gram column, calculate the flow rate, which provides the same linear velocity as the small column. Use the larger column's CV and bed depth as well as the smaller column's linear velocity.

Large column = Biotage® Sfär HC 100 gram with a CV of 180 mL and a bed depth of 15.7 cm, see Table 10.

To determine the flow rate to match linear velocity, multiply the larger column's CV by the ratio of the small column's linear velocity and the large column's bed depth, or

$$\text{Flow rate (Lg)} = \text{CV (Lg)} * ((\text{Linear velocity (Sm)}) / (\text{Bed depth (Lg)}))$$

For this example

$$\text{Flow rate (Lg)} = 180 * (6.47 / 15.7) = 74 \text{ mL/min}$$

Using these equations, determining linear velocity and required larger column flow rate are simplified and ensure matching chromatographic results.

Table 10. Biotage silica flash cartridge column volume and bed depth data by style and media.

Part Number	Column	Media	Particle Size (µm)	Media Quantity (g)	ID (cm)	Length (cm)	Column Volume (mL)
FSRD-0445-0005	Biotage® Sfär Silica D Duo 60 µm 5 g	Silica	60	5	1.64	4.42	9
FSRD-0445-0010	Biotage® Sfär Silica D Duo 60 µm 10 g	Silica	60	10	2.08	5.50	15
FSRD-0445-0025	Biotage® Sfär Silica D Duo 60 µm 25 g	Silica	60	25	2.91	7.40	42
FSRD-0445-0050	Biotage® Sfär Silica D Duo 60 µm 50 g	Silica	60	50	3.90	7.91	80
FSRD-0445-0100	Biotage® Sfär Silica D Duo 60 µm 100 g	Silica	60	100	3.90	15.75	150
FSRD-0445-0200	Biotage® Sfär Silica D Duo 60 µm 200 g	Silica	60	200	5.88	14.60	310
FSRD-0445-0350	Biotage® Sfär Silica D Duo 60 µm 350 g	Silica	60	350	5.88	24.60	530
FSRS-0445-0005	Biotage® Sfär Silica 60 µm 5 g	Silica	60	5	1.64	4.42	9
FSRS-0445-0010	Biotage® Sfär Silica 60 µm 10 g	Silica	60	10	2.08	5.50	15
FSRS-0445-0025	Biotage® Sfär Silica 60 µm 25 g	Silica	60	25	2.91	7.40	42
FSRS-0445-0050	Biotage® Sfär Silica 60 µm 50 g	Silica	60	50	3.90	7.91	80
FSRS-0445-0100	Biotage® Sfär Silica 60 µm 100 g	Silica	60	100	3.90	15.75	150
FSRS-0445-0200	Biotage® Sfär Silica 60 µm 200 g	Silica	60	200	5.88	14.60	310
FSRS-0445-0350	Biotage® Sfär Silica 60 µm 350 g	Silica	60	350	5.88	24.60	530
FSUD-0443-0005	Biotage® Sfär Silica HC D High Capacity Duo 20 µm 5 g	Silica	20	5	1.64	4.42	9
FSUD-0443-0010	Biotage® Sfär Silica HC D High Capacity Duo 20 µm 10 g	Silica	20	10	2.08	5.50	15
FSUD-0443-0025	Biotage® Sfär Silica HC D High Capacity Duo 20 µm 25 g	Silica	20	25	2.91	7.40	42
FSUD-0443-0050	Biotage® Sfär Silica HC D High Capacity Duo 20 µm 50 g	Silica	20	50	3.90	7.91	80
FSUD-0443-0100	Biotage® Sfär Silica HC D High Capacity Duo 20 µm 100 g	Silica	20	100	3.90	15.75	150
FSUD-0443-0200	Biotage® Sfär Silica HC D High Capacity Duo 20 µm 200 g	Silica	20	200	5.88	14.60	310
FSUD-0443-0350	Biotage® Sfär Silica HC D High Capacity Duo 20 µm 350 g	Silica	20	350	5.88	24.60	530
FSUS-0443-0005	Biotage® Sfär Silica HC High Capacity 20 µm 5 g	Silica	20	5	1.64	4.42	9
FSUS-0443-0010	Biotage® Sfär Silica HC High Capacity 20 µm 10 g	Silica	20	10	2.08	5.50	15
FSUS-0443-0025	Biotage® Sfär Silica HC High Capacity 20 µm 25 g	Silica	20	25	2.91	7.40	42
FSUS-0443-0050	Biotage® Sfär Silica HC High Capacity 20 µm 50 g	Silica	20	50	3.90	7.91	80
FSUS-0443-0100	Biotage® Sfär Silica HC High Capacity 20 µm 100 g	Silica	20	100	3.90	15.75	150
FSUS-0443-0200	Biotage® Sfär Silica HC High Capacity 20 µm 200 g	Silica	20	200	5.88	14.60	310
FSUS-0443-0350	Biotage® Sfär Silica HC High Capacity 20 µm 350 g	Silica	20	350	5.88	24.60	530

Part Number	Column	Media	Particle Size (µm)	Media Quantity (g)	ID (cm)	Length (cm)	Column Volume (mL)
FSAD-0909-0005	Biotage® Sfär KP-Amino D ↯ Duo 50 µm 5 g	Amine	50	5	1.64	4.3	9
FSAD-0909-0011	Biotage® Sfär KP-Amino D ↯ Duo 50 µm 11 g	Amine	50	11	2.00	5.6	15
FSAD-0909-0028	Biotage® Sfär KP-Amino D ↯ Duo 50 µm 28 g	Amine	50	28	3.02	7.5	33
FSAD-0909-0055	Biotage® Sfär KP-Amino D ↯ Duo 50 µm 55 g	Amine	50	55	3.94	7.8	66
FSAD-0909-0110	Biotage® Sfär KP-Amino D ↯ Duo 50 µm 110 g	Amine	50	110	3.94	15.8	132
FSAD-0909-0220	Biotage® Sfär KP-Amino D ↯ Duo 50 µm 220 g	Amine	50	220	5.92	14.5	264
FSAD-0909-0380	Biotage® Sfär KP-Amino D ↯ Duo 50 µm 380 g	Amine	50	380	5.92	25.0	510
FSUD-0401-0006	Biotage® Sfär C18 D ↯ Duo 100 Å 30 µm 6 g	C18	28	6	1.64	4.3	9
FSUD-0401-0012	Biotage® Sfär C18 D ↯ Duo 100 Å 30 µm 12 g	C18	28	12	2.00	5.6	17
FSUD-0401-0030	Biotage® Sfär C18 D ↯ Duo 100 Å 30 µm 30 g	C18	28	30	3.02	7.5	45
FSUD-0401-0060	Biotage® Sfär C18 D ↯ Duo 100 Å 30 µm 60 g	C18	28	60	3.94	7.8	85
FSUD-0401-0120	Biotage® Sfär C18 D ↯ Duo 100 Å 30 µm 120 g	C18	28	120	3.94	15.8	164
FSUD-0401-0240	Biotage® Sfär C18 D ↯ Duo 100 Å 30 µm 240 g	C18	28	240	5.92	14.5	328
FSUD-0401-0400	Biotage® Sfär C18 D ↯ Duo 100 Å 30 µm 400 g	C18	28	400	5.92	25.0	582

Summary

For success with flash chromatography ensure you have

- » Clear goals
- » Understand the sample's chemistry
 - » Solubility
 - » Functional groups
 - » Possible interactions with mobile phase solvents and stationary phase media
- » Spend a little time developing your method

For more information on flash chromatography, check out our blog at flash-purification.com

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