Introduction to DSP – Chromatography

Your Objectives:

At the end of the lesson, y			the process o	of Chromatogi	aphy fro	om a
One of the most importar	nt aspects of Biogen's	manufa	acturing prod	cesses is the i	recovery	and
	of a single target pro	otein mo	olecule. The	molecule typic	cally exis	ts in
solution and must be		from th	e other com	ponents in the	solution	١.
Chromatography provide	es a method fo	r sep	arating mo	olecules fro	n com	plex
	. By analyzing a	target i	molecule's	characteristics	, it car	ı be
differentiated from other i	molecules in a solutio	n. The t	target molec	ule can then l	be separ	ated
using the appropriate		proces	ses.			
The chromatography meth depend upon the chemic		of chroi]	olecule as we	ll as on	the
	of purity needed fo	or the	<u> </u>	necure as we	produc	
general, chromatography reversible. Chromatograph		-	• •	_		
distinguish proteins t	hat differ in			by	a s	ingle
	acid or a single atom	n, even.	Moreover, b	ecause the co	onditions	and
processes of chromatogra highly fragile components.	phy are gentle, it all	ows for	a reliable v	vay of separa	ting deli	cate
Chromatography has two		St	tationary &	mobile, each	moving	in a
definite direction, and are t	typically required for t	he regu	latory approv	val of protein	drugs.	

The simplest way to explain chromatography is by comparing it to a raging river:

A raging river can carry a lot of debris along with it. The speed at which floating debris is moved will depend upon (1) the types of floating debris (i.e., grains of sand are transported faster than pebbles), and (2) the nature of the riverbed (i.e., rough surfaces increase the friction of the floating debris, thus slowing or stunting the speed of removal) on the flow velocity.

In chromatography, various		= floating debris in the 'river') are
the "riverbed"). Due to the iphases, individual substances separated and distinguishable slightly larger 'pebbles' is intranalogy, after 100 metres all certain waiting period, all the strewn across a certain distanting river analogy is indeed if	nteractions* between the sare transported at differe from one another where oduced at a certain point in the sand will arrive first (symmetric) smaller pebbles follow and the sand will arrive first (symmetric).	t is the river') on a stationary phase (sample, stationary phase and mobile rent speeds and are, for that reason a mixture of 'sand', very small and in time into the 'river'; according to the pread over a few metres), and, after a d, much later, any larger particles, each initial understanding of the process of graphy is perhaps more evocative of a
digital process known as "stop	o-and-go traffic,") whereby	the sample
are either carried along with	in the mobile	(analogous to a 'ligh
raft' that is passively carried i	n a current) or adhering to	p the phase
(at 'zero speed'). The mole possibilities very rapidly and h		forth (vacillate) between these two
molecules suffer (through t	he chromatographic syste	ch as the delays that the various sample em) do not account for the friction ce is in the distribution of the differen
types of	(types A, B, C, etc.),	which correspond to differences in the
Chromatography allows for a	a conversion of such diffe	ules spend in the mobile phase) erences into 'speed differences', thus which, these minute (tiny) differences
could not be used, neither for		and cleaning processes nor for prope
/a	nalyses.	

^{* (}For interactions, see the "division under separation" principles.)

The Process

Four	nhases	of chi	romatogra	nhy are	annlied:
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1. Establishment of the flow of the mobile phase,		
Injection of the sample to be separated,		
3. Actual separation, and		
4. of components		
The flow of the mobile phase is achieved by any one of three means: by pressure (e.g. hydraulic pump, gas pressure), by capillary force, or then by applying an electrical		
The (i.e., introduction of the substance mixture into the		
chromatographic system) takes place either before the flow of the mobile phase is established		
(thin-layer chromatography) or while the mobile is already in flow.		
With a large number of samples, so-called autosamplers are used with automatable types of chromatography (together with their own data acquisition systems), which inject the samples fully automatically.		
The actual separation of the mixture follows on the separating		
section. Finally, (as with the separation or separate phase) chromatography would be inconceivable		
without (.:.making visible when a substance passes a certain section		
of the chromatography system or where a substance comes to a standstill after the process has		
ended). Different detection are applied for each type of		
chromatography, either by using physical properties (e.g., absorption of light, fluorescence,		
light scattering, and thermal) of the substances, or then by		
obtaining a signal through reactions by means of chemical		

reactions; for example, a	achieved in planar chromatography (e.g.,			
amino acids using ninhydrin), or	amino acids using ninhydrin), or then through reactions carried out before separation (pre-			
column derivatization) or other	erwise separation (post-column			
derivatization), in column chroma In the case of preparative chro collect the separated substance.	atography. matography, a fraction collector is additionally required to			
Due to design chromatographic p	purification processes are always batch processes. This means			
that only a certain	of substance can be applied (injected) and			
separated before proceeding with the next (equal) amount, making it particularly problematic when working up large amounts. Therefore, specific methods, all of which could not be otherwise be done by a simple column purification, have been developed so as to operate chromatography continuously:				
Continuous annular chromatogo simulated moving bed chromatogo	raphy (CAC), true moving bed chromatography (TMB) and graphy (SMB).			
CAC lends itself to the separationes. TMB is used as a cost-effective pr SMB is a theoretical concept	on of multi-component mixtures as well as of bi-component ocess.			
-	nethod was developed for the high-performance liquid nination of penicillin using fluorescence detection The			
resulting reaction	was injected directly onto a reversed-phase			
column and analysed by HPLC.				
Post-column derivatization,	also known as post-column reaction, renders			
certa	in compounds that are normally invisible. This trick is			
accomplished after the separation gives them an easily detectable p	on by performing a chemical reaction on the substances that hysical property.			

Aufgabe Lückentext:

Folgende Wörter bitte in den Lückentext einfüllen. Jedes Wort kommt einmal vor. Bitte Gross- und Kleinbuchstaben beachten.

amino, analysis, amount, after, chromatography, composition, coloration, conductivity, chemical, Detection, detection, degree, final, injection, isolated, molecules, molecules, mixture, makeup, phases, phase, purification, phase, solutions, stationary, steps, substances, separation, systems, substance, voltage, visible