Regeneration Things to consider + Sensor Regeneration condition overview

Selvakumar Dakshnamurthy Application Scientist





Regeneration

Regeneration is the process of restoring the sensor to its initial condition by removing the attached analyte without damaging the ligand on the sensor surface.





Regeneration of Current Biosensor Products

- This presentation will show:
 - Regeneration conditions for current biosensor products.
 - How to determine the right custom regeneration conditions.
 - How to use regeneration in the software.



Regeneration of Current Biosensor Products

- Biosensor regeneration is dependent on:
 - Capture chemistry and affinities
 - Assay requirements
- Regeneration of biosensors for quantitation applications must be more complete than for kinetics assays. This is because the quantitation results are more dependent on surface capacity of the sensor.
 - For example, a loss of 20% capacity over 10 regeneration cycles will not affect kinetic constants, but would affect precision of quantitation by 10-20%.



Protein A/G/L and FAB2G Biosensors



• Optimal Regeneration Buffer

ProA	10mM Glycine pH 1.0-1.5
ProG	10mM Glycine pH 1.7-2.0
ProL	10mM Glycine pH 1.5
FAB	10mM Glycine pH 1.7

- Be sure to use the pre-conditioning option for best precision
- 3 cycles of 5-10 seconds alternating with running buffer as a neutralizations step





Regeneration of ProA binding 500 ug/mL hlgG



Data shown by sensor. Each graph shows binding curves from 10 binding cycles (9 regenerations)



Regeneration efficiency of Protein A Biosensors

	Ave. calculated concentration of 8 channels (ug/mL)	%CV of 8 channels
REG0	485.0	5%
REG1	485.2	4%
REG2	496.1	5%
REG3	490.0	5%
REG4	490.5	5%
REG5	503.4	5%
REG6	505.1	5%
REG7	500.7	4%
REG8	501.4	4%
REG9	498.9	4%

CV over 9 regenerations

2%

Regeneration protocol:

3 Cycles of 10 mM Glycine pH 1 for 5 sec at 200 rpm, Sample Diluent for 5 sec at 200 rpm







- Regeneration not recommended due to:
 - High affinity of capture antibody. Requires harsh conditions that damage the surface.
 - Pfizer SSF is regenerating this sensor, but they are using it only for kinetic epitope binning assays since loss of surface capacity is evident.







- Optimal Regeneration Conditions:
 - 10mM Glycine pH = 1.7 (3 cycles of 5 seconds alternating with running buffer as a neutralizations step)
 - Pre-conditioning of the sensor surface is required for accurate kinetics.



Kinetics analysis using AHC Biosensors with regeneration



KD (M)	Kon (1/Ms)	Kdis (1/s)
3.66E-09	3.97E+0 5	1.45E-03
3.45E-09	4.06E+0 5	1.40E-03
3.75E-09	3.89E+0 5	1.46E-03
3.94E-09	3.84E+0 5	1.51E-03
3.99E-09	3.77E+0 5	1.50E-03



Note 10% loss after 3 cycles, yet kinetic constants stay the same



Residual ProA/CHO HCP kits

Bind analyte

Quantitate analyte (rProA)

2nd Ab (CHO HCP)

Residual Protein A/ CHO HCP



Residual ProA

- Regeneration not recommended due to:
- High affinity of capture antibody. ٠ Requires harsh conditions that damage the surface.
- Kit contains only enough reagents to ٠ support use of each biosensor once.

Regeneration not recommended due to:

Quantitate analyte (CHO HCP)

CHO HCP

- Kit contains only enough reagents to support use of each biosensor ones.
- DAB precipitates on sensor surface and is not regenerable





HIS1K and HIS2 (Anti-PentaHIS) Biosensors



- Regeneration possible if:
- Affinity for analyte is sufficiently low allowing full clearing of the surface without loss of surface capacity.
- Generally this is not possible for quantitation, however Centocor is regenerating for a kinetic characterization assay.



Ni-NTA Biosensor



- Optimal Regeneration Conditions:
- 10mM Glycine pH 1.7 3 cycles of 5-10 seconds alternating with running buffer as a neutralizations step
- After the final cycle, sensor must be recharged with 10mM NiCl2 in water
- DO NOT use preconditioning

CONFIDENTIAL

DO NOT regenerate for Q applications (surface is totally different after each recharge)



Anti-GST Biosensor



- Optimal Regeneration Buffer for kinetic applications
- 10mM Glycine pH 1.7-2.0
- DO NOT use pre-conditioning for best precision.
- 3 cycles of 5-10 seconds alternating with running buffer as a neutralization step.
- Note that there will be 2-3% surface capacity loss with each regen cycle (do not regenerate for quantitative applications).



APS Biosensor



- Optimal Regeneration Conditions:
- Are extremely variable and depend totally on what is attached.
- Generally detergent solutions work well with this biosensor (1% Triton X-100, Tween-20, SDS etc).
- If using a detergent it is very important to properly wash the biosensor clean before the next binding step.



AR2G Biosensor



- Optimal Regeneration Conditions:
- Are extremely variable and depend totally on what is attached.



Kinetics analysis using AR2G Biosensors with regeneration



Note 25% loss after 3 cycles, yet kinetic constants stay the same



7 CONFIDENTIAL

Chemical Stability of the AR2G Biosensor

Reagent	Maximum validated exposure time*
10 mM Acetate buffer pH 0.5, 1, 2, 3	15 minutes
10-100 mM Citrate buffer pH 2	15 minutes
КОН рН 9, 10, 11	15 minutes
50 mM NaOH	15 minutes
Phosphoric acid pH 2 + 5% Tween 20	15 minutes
SDS (0.5%, 0.1%, 0.05%)	5 minutes
5M NaCl	15 minutes
4M MgCl2	15 minutes
1mM HCl	15 minutes
Ethylene glycol (25%, 50%)	15 minutes
20 mM EDTA	5 minutes
Mixture of 0.46 M KSCN, 1.83M MgCl2, 0.92 M urea and 1.83M guanidin-HCl	5 minutes
CHAPS (1%, 0.3%, 0.05%)	Not recommended
Mixture of equal amount DMSO, formamide, ethanol, acetonitrile and 1-butanol	5 minutes

* Exposure times are for the biosensor itself. Stability of the protein studied will be protein dependent.



SSA (Super Streptavidin) biosensor



- Optimal Regeneration Conditions:
 - Neutral buffers only!
 - Small molecules of low affinity should dissociate fully in running buffer.
 - Chemical regeneration of this biosensor will damage the surface preventing proper use in small molecule analysis







- Optimal Regeneration Conditions:
- Are extremely variable and depend totally on what is attached.
- Note, if this biosensor is used in a multistep quantitation assay that utilizes precipitating substrate, it cannot be regenerated.





Chemical Stability of SA/SAX/SAX2 Biosensor

Reagent	Maximum validated exposure time*
HCl (pH 0.5, 1.0, 1.5)	15 minutes
NaOH (pH 10, 11)	15 minutes
NaOH (pH 12, 12.5, 13)	Not recommended
10 mM Glycine (pH 1, 2, 3)	15 minutes
NaCl (1, 2.5, 5 M)	15 minutes
MgCl2 (0.1, 0.5, 1 M)	15 minutes
Tween-20 (0.1%, 0.25%, 0.5%)	15 minutes
SDS (0.05%, 0.1%, 0.25%, 0.5%)	Not recommended
SDS (0.005%, 0.01%)	15 minutes
Phosphoric Acid (50, 100, 250, 500 mM)	15 minutes
EDTA (25, 50, 100 mM)	15 minutes
TritonX-100 (0.1%, 0.25%, 0.5%)	15 minutes

* Exposure times are for the biosensor itself. Stability of the protein studied will be protein dependent.



Regeneration of Current Biosensor Products

• Quick Overview:

A méile a dur	Intended			
hiosonsore	Application	Percentration	Pecommended buffer	Accay Paramotore
Diosensors ProA		Vec	10mM Glycine pH 1 0-1 5	Precord 3y 5-10s
ProG	0	Ves	10mM Glycine pH 1 7-2 0	Precond 3x 5-10s
Prol	0	Ves	10mM Glycine pH 1.5	Precond 3x 5-10s
		Voc	10mM Glycine pH 1.7	Precond 3x 5 10s
	Q/N	Ves for K	10mM Glycine pH 1.7	Precond 3x 5-10s
	ĸ	Vec for K	10mM Glycine pH 1.7	Precond 3x 5-10s
	0			Frecond, 5x 5-10s
			-	
	Q		-	
- :				
Biosensor	Intended	D		A D 4
KITS	Application	Regeneration	Recommended buffer	Assay Parameters
rProA	Q	No for Q	-	
сно нср	Q	No for Q	-	
Anti-TAG	Intended			
biosensors	Application	Regeneration	Recommended buffer	Assay Parameters
HIS1K	Q/K	Custom	Protein dependent	Protein dependent
HIS2	Q	Custom	Protein dependent	Protein dependent
NTA	O/K	Ves for K No for O		
	Gent	Testor R, No Tor Q	10mM Glycine pH 1.7	3x 5-10s
FLG	Q/K	Yes for K, No for Q	10mM Glycine pH 1.7 10mM Glycine pH 1.7	3x 5-10s Precond, 3x 5-10s
FLG GST	Q/K Q/K	Yes for K, No for Q Yes for K, No for Q	10mM Glycine pH 1.7 10mM Glycine pH 1.7 10mM Glycine pH 1.7	3x 5-10s Precond, 3x 5-10s 3x 5-10s
FLG GST	Q/K Q/K Q/K	Yes for K, No for Q Yes for K, No for Q	10mM Glycine pH 1.7 10mM Glycine pH 1.7 10mM Glycine pH 1.7	3x 5-10s Precond, 3x 5-10s 3x 5-10s
FLG GST Generic	Q/K Q/K Intended	Yes for K, No for Q Yes for K, No for Q	10mM Glycine pH 1.7 10mM Glycine pH 1.7 10mM Glycine pH 1.7	3x 5-10s Precond, 3x 5-10s 3x 5-10s
FLG GST Generic biosensors	Q/K Q/K Intended Application	Yes for K, No for Q Yes for K, No for Q Regeneration	10mM Glycine pH 1.7 10mM Glycine pH 1.7 10mM Glycine pH 1.7 Recommended buffer	3x 5-10s Precond, 3x 5-10s 3x 5-10s Assay Parameters
FLG GST Generic biosensors APS	Q/K Q/K Intended Application	Yes for K, No for Q Yes for K, No for Q Regeneration Custom	10mM Glycine pH 1.7 10mM Glycine pH 1.7 10mM Glycine pH 1.7 Recommended buffer Protein dependent	3x 5-10s Precond, 3x 5-10s 3x 5-10s Assay Parameters Protein dependent
FLG GST Generic biosensors APS AR2G	Q/K Q/K Intended Application K	Yes for K, No for Q Yes for K, No for Q Regeneration Custom	10mM Glycine pH 1.7 10mM Glycine pH 1.7 10mM Glycine pH 1.7 Recommended buffer Protein dependent Protein dependent	3x 5-10s Precond, 3x 5-10s 3x 5-10s Assay Parameters Protein dependent Protein dependent
FLG GST Generic biosensors APS AR2G SSA	Q/K Q/K Intended Application K K K	Yes for K, No for Q Yes for K, No for Q Yes for K, No for Q Regeneration Custom Custom Neutral pH only	10mM Glycine pH 1.7 10mM Glycine pH 1.7 10mM Glycine pH 1.7 Recommended buffer Protein dependent Protein dependent Protein dependent	3x 5-10s Precond, 3x 5-10s 3x 5-10s Assay Parameters Protein dependent Protein dependent Protein dependent
FLG GST Generic biosensors APS AR2G SSA SA	Q/K Q/K Intended Application K K K K	Yes for K, No for Q Yes for K, No for Q Yes for K, No for Q Regeneration Custom Custom Neutral pH only Custom	10mM Glycine pH 1.7 10mM Glycine pH 1.7 10mM Glycine pH 1.7 Recommended buffer Protein dependent Protein dependent Protein dependent Protein dependent	3x 5-10s Precond, 3x 5-10s 3x 5-10s Assay Parameters Protein dependent Protein dependent Protein dependent Protein dependent



Regeneration of Current Biosensor Products

- This presentation will show:
 - Regeneration conditions for current biosensor products.
 - How to determine the right custom regeneration conditions.
 - How to use regeneration in the software.



Rapid Regeneration Scouting Acquisition Template

Use of RegenerationConditionScouting Template embedded in the software:

Octet Dat	Acquisition 9.0.0.7						
<u>File</u> <u>V</u> iew	Experiment Instrument <u>W</u> indow	<u>H</u> elp					
🔌 🖄	New Experiment Wizard (Edit Assay Parameters	Ctrl+N					
	Edit Sensor Types						
	Set Plate Temperature						
	Templates	•	Epitope Binning	- + j			
	Skip Step		Kinetics	•	Biomolecule kinetics - AHC biosensor	+	
	Stop		Quantitation	•	Biomolecule kinetics - AMC biosensor	- F	
				1	Biomolecule kinetics - AR biosensor	_ → _	
					Biomolecule kinetics - SA biosensor	+	Kinetic Characterization_8CH_96W.fmf
					Small Molecule and Fragment Kinetics - SSA biosensor	•	RegenerationConditionScouting_8CH_96W.fmf

- Use 1 Ligand loading concentration and 1 high Analyte concentration and 8 different Regeneration Buffers.
- Copy Regeneration Buffer names in Sensor Info column on Sensor Assignment tab (3):

Well	Sensor Type	Lot Number	Information
A1	SA (Streptavidin) 👻		5 M NaCl
B1	SA (Streptavidin)		0.01% SDS
C1	SA (Streptavidin)		NaOH, pH 10
D1	SA (Streptavidin)		NaOH, pH 11
E1	SA (Streptavidin)		HCl, pH 0.5
F1	SA (Streptavidin)		10mM Glycine, pH 1
G1	SA (Streptavidin)		10mM Glycine, pH 2
H1	SA (Streptavidin)		500mM phosphoric acid





 Analyze data both with and without step Align Y axis to baseline:

Data Correction		
1 - Align Y Axis		
Shift all data in trace	by value as selected below:	
Align Data to:	Average of Baseline Step	~
Start:	115.01 🔹 End:	120.01



÷.

4-

3-

2

Jde

• Set Color by Cycle: Stacked





• Group data by Sensor Info:

View					Y Axis Scaling	Report Poir	nts		
Stack	ked	Gro	up	Always	Auto Scale	Time (sec):	100	Add to Table	Sav
Indiv	vidual	Optio	ns Refresh	Refresh	Full Scale	Use 20	Point Average	Remove All	Loa
			-			Fit	tina View		
0.8-				🚽 Group View O	ptions				x
0.7-				Group Graph	s Bv:	Graph	Size in Pixels		
0.6					, Info		# Graphs/Row:	4 🌩	
0.5-		1		Sensor	IIIO	• • AL	to Size: Width =	1.5 x Height	
= 0.4	1					- - - - - - - - - - - - - -	ed Size: Width:	300	
	Ale and a second					-	Height	200	
0.3-	184						noight.	200 🔻	
0.2-				Lege	nd by:	Graph	Options		
0.1-		6		Colo	r -		Title		
0						7	Individual Co	olumn Names	
-	1 1	1				-	Legend by	Column Name	
Ó			50	Data		XA	vie: 🔽 Labele	🔽 Titles	
				All Date	ata	YA	vie: 🔽 Labele		
1				🔘 Inclu	ded Traces Only			i nues	
0				Selection	ted Traces Only		Grid Lines	Step Dividers	
-1-	1 1	3	50	Data Optio	ns /as%.ofRmax IV Sl	how Curve Fits	Display Tr	aces in Table Color	
U			50	Additional (Granhe				
Include	Index	Color	Sensor Location	Resid	uals 📃 Stea	dy-State	🕅 X-Y	Iso-Affinity	
	0		A4						
~	1		A4		OF	(Cancel		
	2		A4	and the second	11/7.			200.000	





 When using Align Y axis, data shows overlay of association curves of different cycles and right regeneration buffer can be chosen.



 When not using Align Y axis, data gives information if regeneration was too harsh for the ligand (starting point the same, association curves decrease in nm-shift) or if analyte did not come of sensor (starting point increases in nmshift, association curves reach similar level in nm-shift).



Regeneration of Current Biosensor Products

- This presentation will show:
 - Regeneration conditions for current biosensor products.
 - How to determine the right custom regeneration conditions.
 - How to use regeneration in the software.



Regeneration in Quantification mode

- Regeneration possible in both
 - Basic Quantitation with Regeneration
 - Advanced Quantitation





Regeneration in Quantification mode

- Standard Assay parameters for regeneration are:
 - 5 secs Regeneration at 200/1000 rpm
 - 5 secs Neutralization at 200/1000 rpm
 - 3 cycles of above steps
- Differentiation can be made between amount of cycles for preconditioning and between assays.

- Regeneration		
Integeneration	Time (s):	Shake speed (rpm):
Regeneration:	5 🚔	1000 🚔
Neutralization:	5 🚔	1000 🚖
		Regeneration cycles:
Between assa	ay steps:	3 🚔
Pre-condition	3 🚔	
Post-condition	3	



Regeneration in Kinetic mode

• Add Regeneration and Neutralization column to your plate





Regeneration in Kinetic mode

CONFIDENTIAL

 Add Regeneration step to your assay step list (change parameters using Regeneration Params button)

-	Step	Data List			1.000	" ¹⁰ (0), 0110110 0p00	A in (ipin)	
44 49 11	A	dd C	Сору	Remove	Regeneration Param	ns Threshold	l Params	
		Name	Time	Shake speed	Туре	Threshold		
		Baseline	60	1000	🛌 Baseline			
<i>1</i>	→	Regeneration	30	1000	🔹 Regeneration			
		2						
								×
				F	Regeneration Paramet	ers		_
					Step Name:	Regeneration		
						True (c)	Chalus are and (see).	
					Receperation:	Time (s)	snake speed (rpm):	
		au No Cam	nla C	ton Nama	Negeneration.	3	1000	
	455	dy NU. Jali	ipie 5		Neutralization:	5	1000 🚔	
bles	Ľ	I AII	n	egeneration +	Regeneration cycles:	3		
					Total step time:	30 s	ОК	Cancel

- Double-click on Regeneration column to have Regeneration step added to the assay
- Note that this Regeneration Step defined by the software will put the nmshift after last Neutralization step automatically at 0 nm.



Thank You

