



# Post translational modification of proteins in the context of immune recognition

F. Hudecz<sup>1,2</sup>

<sup>1</sup>Research Group of Peptide Chemistry, Hungarian Academy of Sciences,  
Eötvös Loránd University (ELTE),

<sup>2</sup>Department of Organic Chemistry, Institute of Chemistry, ELTE, Budapest, Hungary

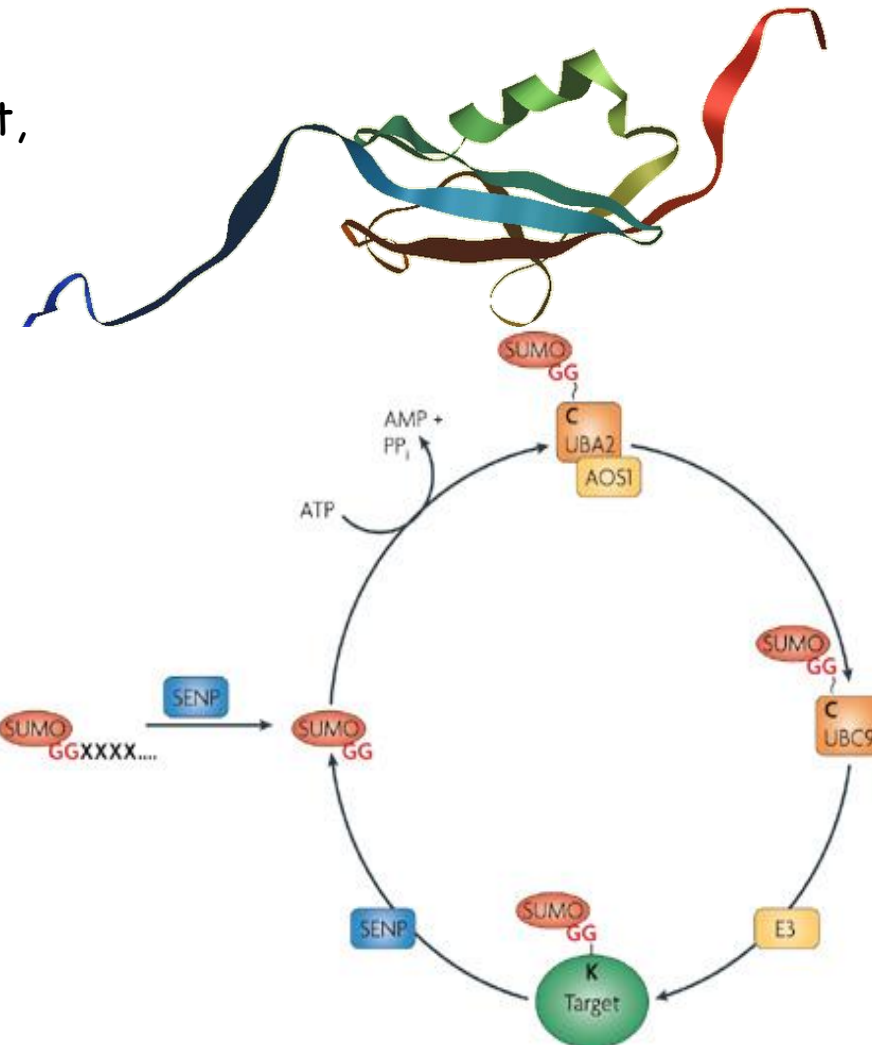
# In vivo post-translational modifications - diversity

alkyl	Cleavage of peptide bond
acyl (O-,N-,S-)	N-terminals Met or fMet signalpeptide precursor activation (proinsulin → insulin)
N-terminal, Lys, Ser, Thr	
amide	Disulphid bond formation
C-terminal	
phosphoric acid ester (Ser, Thr, Tyr)	Isomerisation (Pro, Asp)
sulphonic acid ester	
glycosylation	Coupling of nucleotide (e.g. flavine)
O- in Golgi (Ser, Thr)	
N- in RER (Asn)	
nitrosation	
desamidation	
decarboxylation	
Arg desamination, citrullination (Arg → citrullin)	Coupling of protein/peptide : sumoylation (SUMO protein) ubiquitination (ubiquitin) neddylation (Nedd)
hydroxylation (Pro, Lys)	
oxidation	
gamma-carboxylation (e.g. Glu )	
beta-elimination (e.g. Thr → alkene)	

# SUMOylation (Small Ubiquitin-like Modifier)

Involved in nuclear-cytosolic transport,  
transcriptional regulation,  
apoptosis  
protein stability,  
**but, not in degradation**

- SUMO proteins: 100 aa., 12 kDa, 4 isoforms
- Post-translational modification
- Activation: - cleavage of 4 residues at the C-terminal
- Attachment to target protein by using three enzymes .



R. Geiss-Friedlander & F. Melchior  
Nature Rev. Mol. Cell Biol. 8, 947-956 (2007)

# Ubiquitination

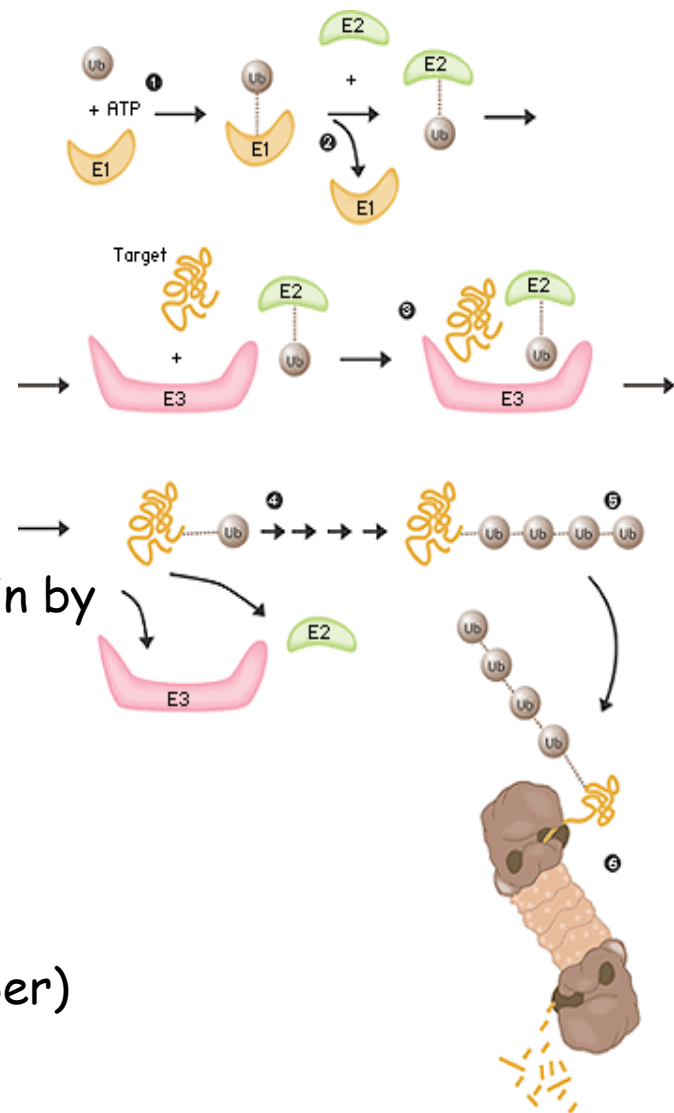


KUNGL.  
VETENSKAPS AKADEMIEN  
THE ROYAL SWEDISH ACADEMY OF SCIENCES



Nobel Prize in Chemistry, 6 October 2004  
A. Ciechanover, A. Hershko, I. Rose

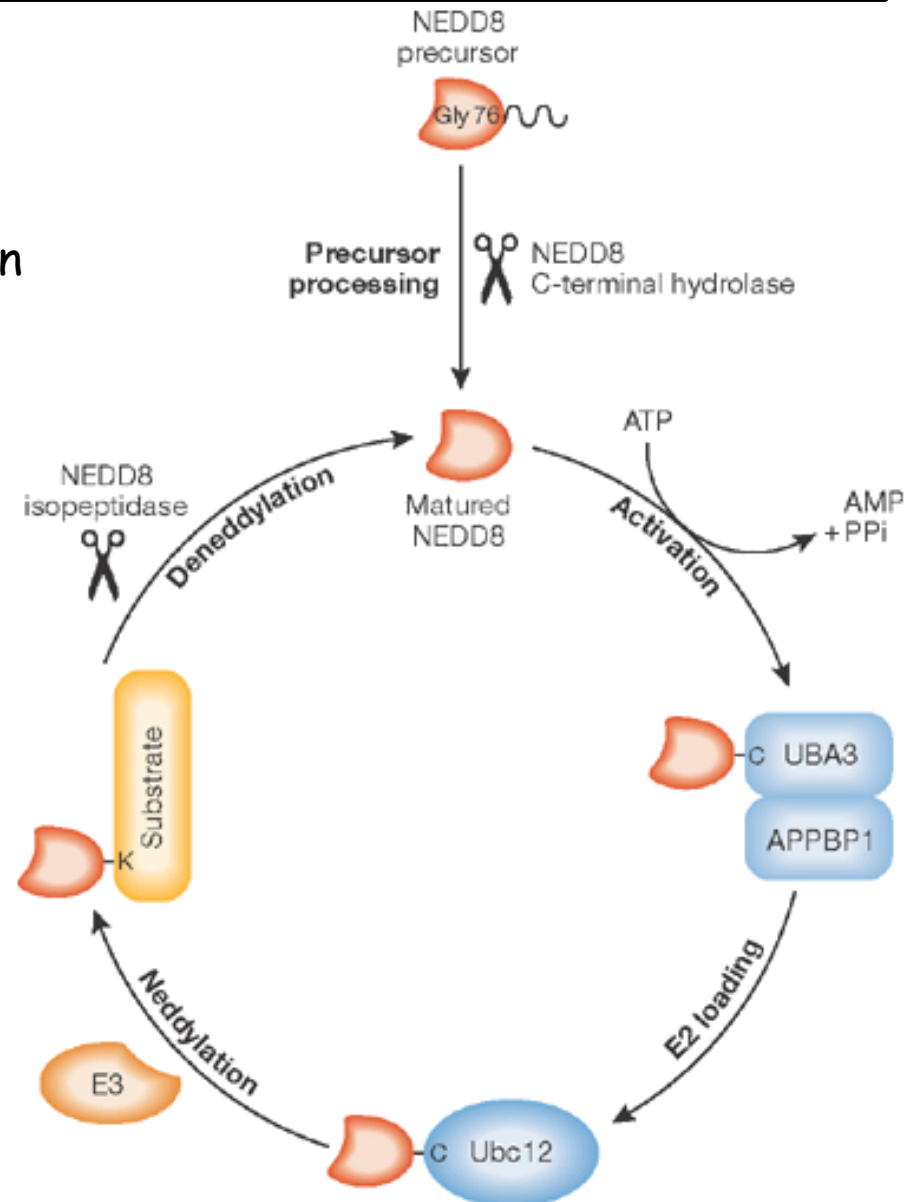
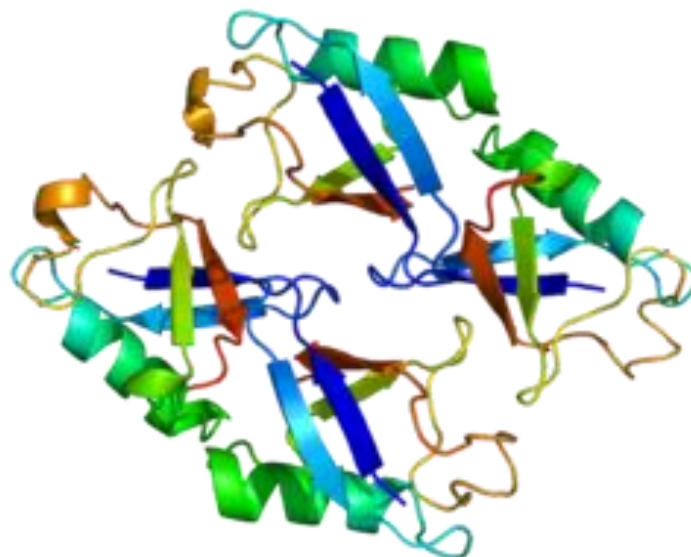
- Ubiquitin: protein (76 aa., 8.5 kDa)
- Almost all tissues of eukaryotic organisms
- „Label“ the degradation of the attached protein by transporting to proteasome.
- Isopeptide linkage (4)
- Enzymes involved:
  - E1 (ub activation)
  - E2 (ub conjugation to  $\epsilon$ -amino group (Lys), thiol (Cys) by thioester, and to OH (Thr/Ser) by ester formation
  - E3 ( ub ligation)



# NEDDylation

(Neural-precursor-cell-expressed developmentally down-regulated)

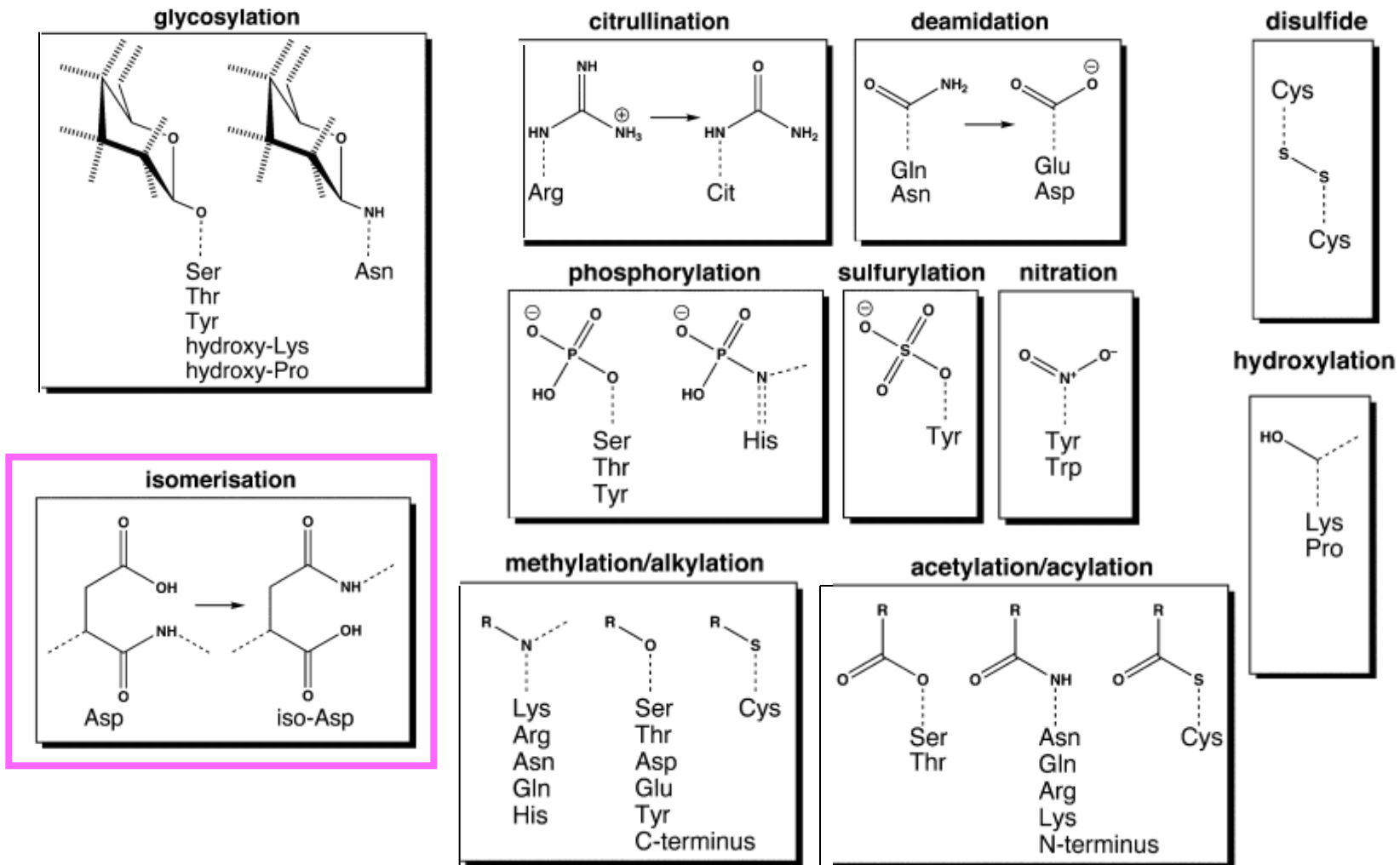
- Function:  
activation/regulation of ubiquitin



# Post-translational modification: Immune recognition related diseases

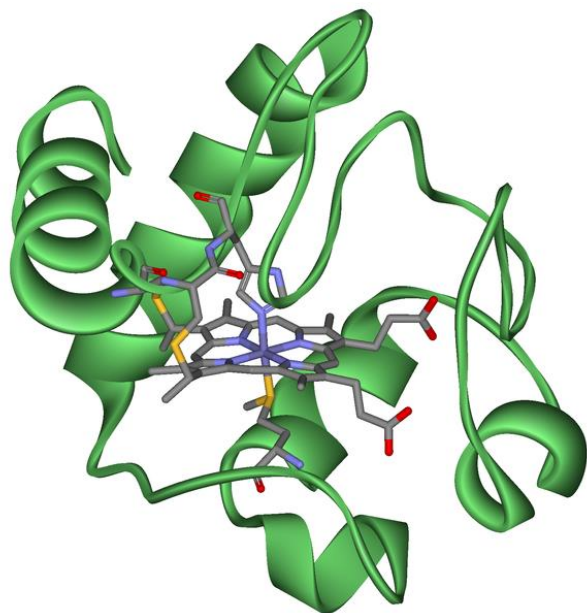
Modification	Autoantigen	Disease
Acetylation	Myelin basic protein	Multiple sclerosis
Citrullination	Collagen type II Myelin basic protein	RA Multiple sclerosis
Deamidation	Insulin	Type I diabetes
Glycosylation	Insulin proceptor Collagen type II Thyrotropin receptor Myelin oligodendrocyte glycoprotein Mucin glycoprotein (MUC2)	Diabetes RA Graves disease MS Colon carcinoma
Isoaspartylation	snRNP	Systemic lupus erythematosus
Lipoylation	PDC-E2	Primary biliary cirrhosis
Phosphorylation	Myelin basic protein	Multiple sclerosis
Methylation	Sm, D1,D3	Systemic lupus erythematosus
Transglutamination	Histone H2	Systemic lupus erythematosus
Tyrosine nitration	Mitochondrial proteins	Experimental autoimmune uveitis

# Post-translational modification: influence on immune recognition



# The effect of post-translational modification on T-cell immune recognition: Isomerisation of Asp to $\beta$ -Asp

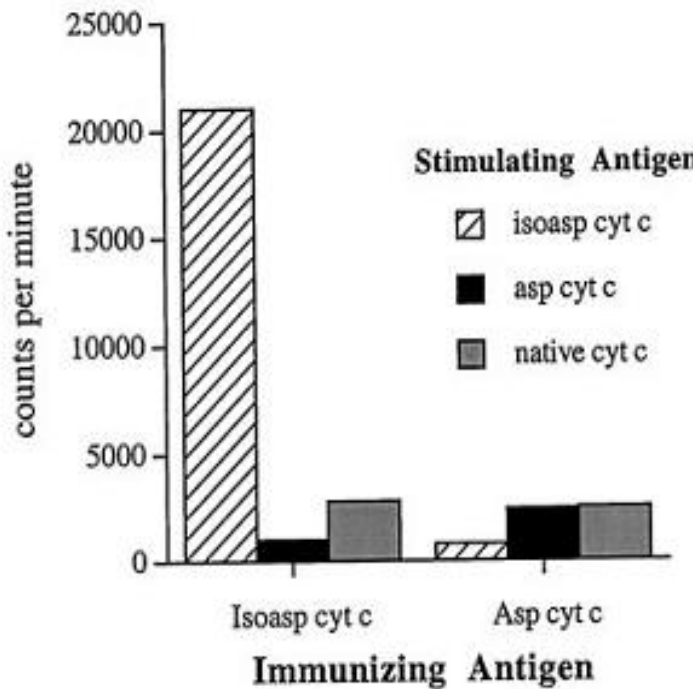
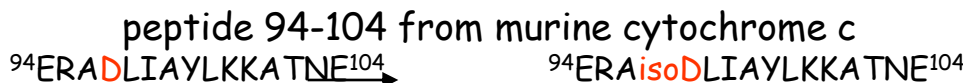
## T cell response



structure of horse heart cytochrome c (PDB:1HRC)

### Assay:

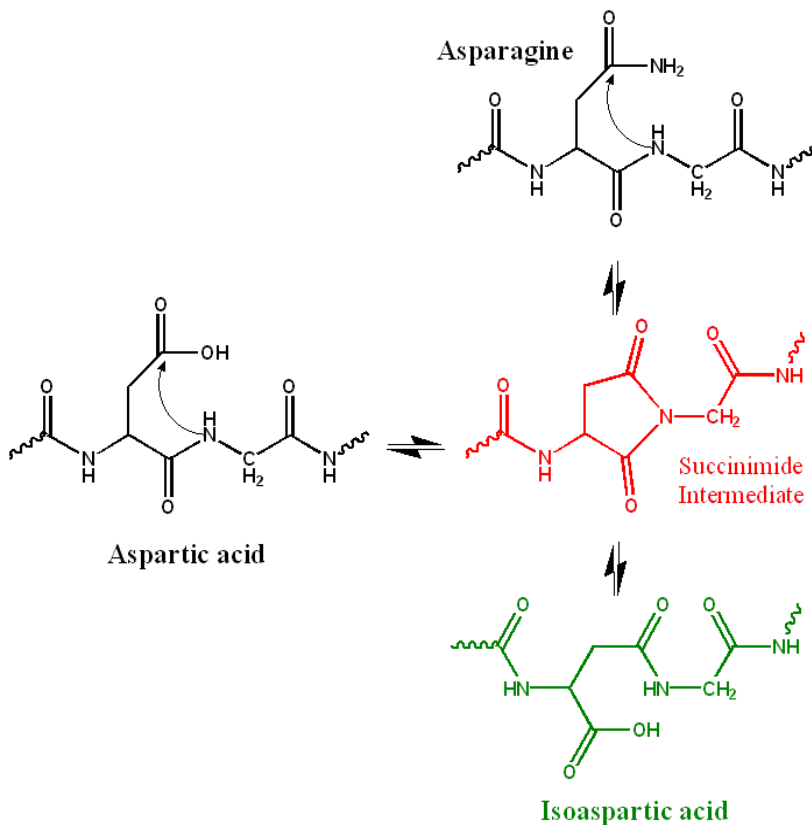
- B10.A mice immunized with 100  $\mu$ g peptide 94-104 with CFA
- after 10 days cell suspension from lymph nodes
- antigen stimulation with peptide a, b or full protein
- [ $^3$ H]thymidine incorporation assay



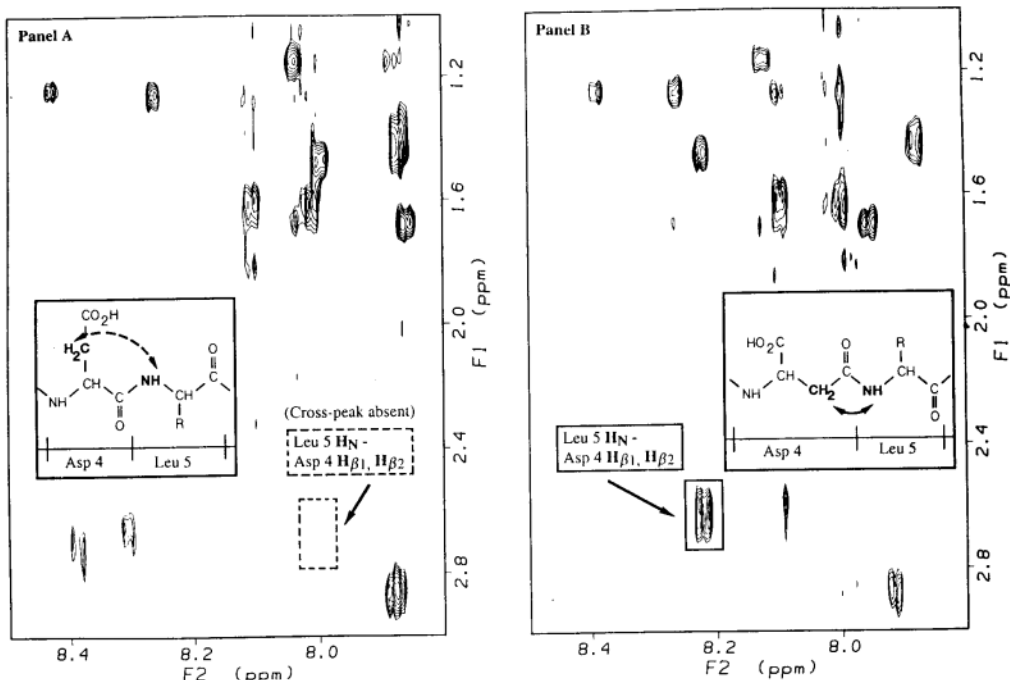
T cells respond to immunization with isoAsp self-peptide and fail to respond Asp self-peptide

# The effect of post-translational modification on T-cell immune recognition: Isomerisation of Asp to $\beta$ -Asp

peptide 94-104 from murine cytochrome c



NMR analysis  
of immunogenic/non-immunogenic self-peptide



$^{94}\text{ERA}\textcolor{red}{D}\text{LIAYLKKATNE}^{104}$

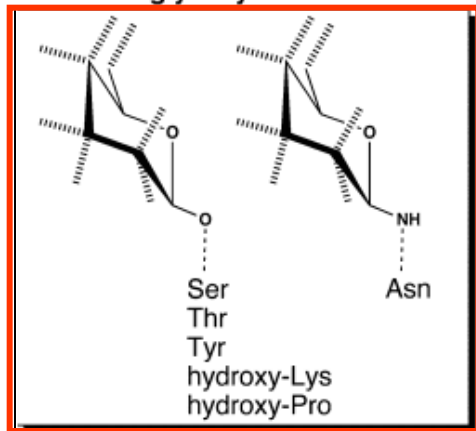
$^{94}\text{ERA}\textcolor{red}{isoD}\text{LIAYLKKATNE}^{104}$

Mamula, M.J. et al. J. Biol. Chem. 274: 22321-22327 (1999)

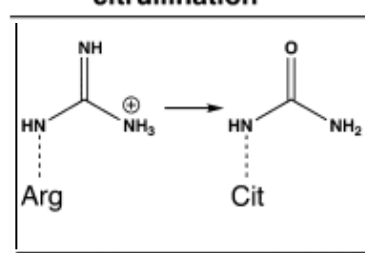
The effect of post-translational  
modification on antibody recognition:  
glycosylation

# Glycosylation

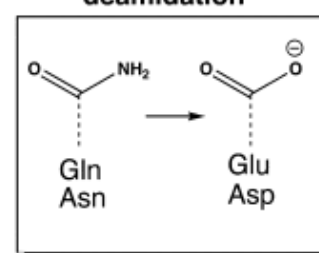
**glycosylation**



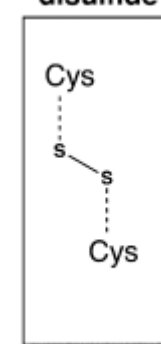
**citrullination**



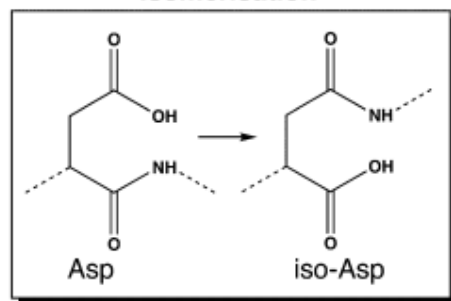
**deamidation**



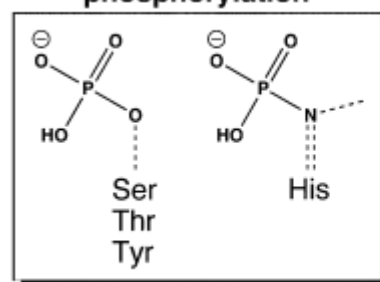
**disulfide**



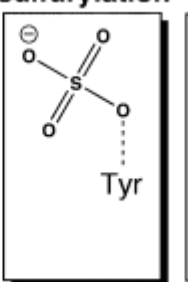
**isomerisation**



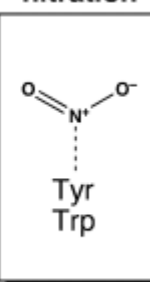
**phosphorylation**



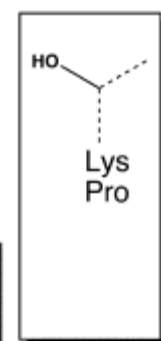
**sulfurylation**



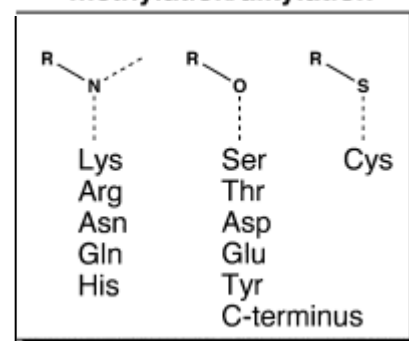
**nitration**



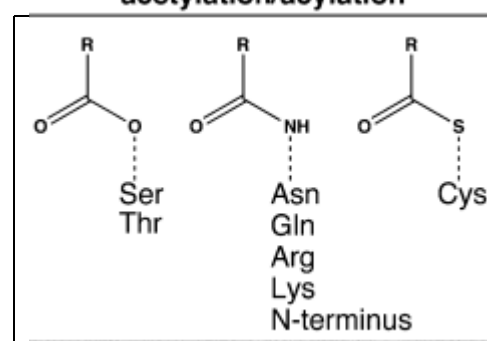
**hydroxylation**



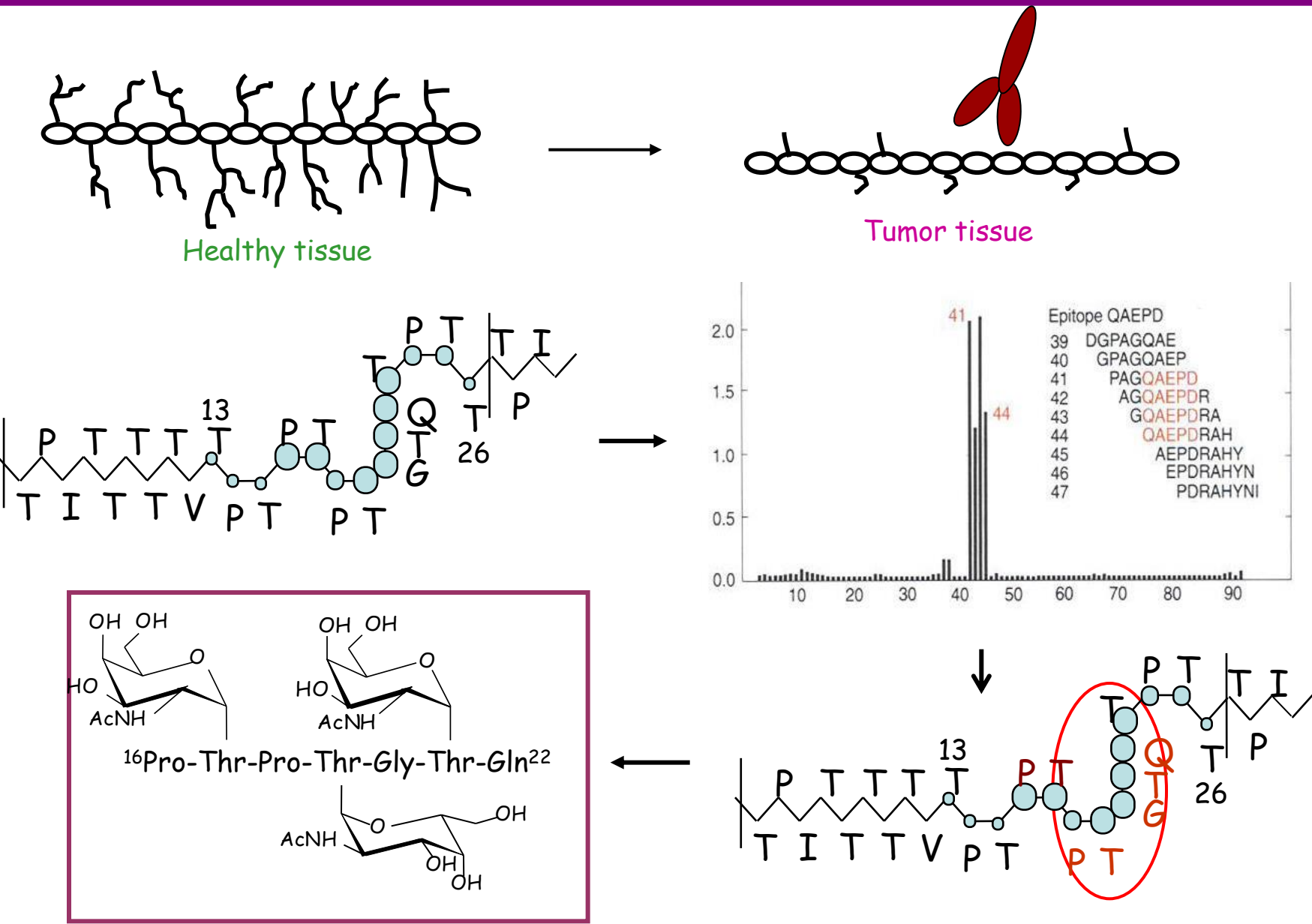
**methylation/alkylation**



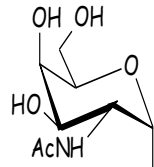
**acetylation/acylation**



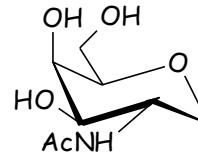
# Identification of antibody epitope of mucin-2



# The effect of carbohydrate moiety on MoAb binding

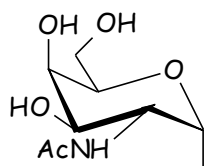
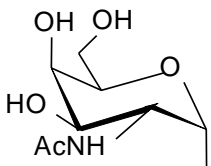


$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$

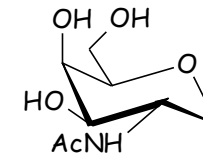
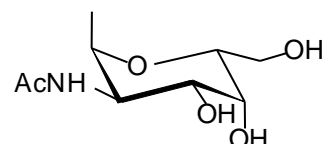


$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$

$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$

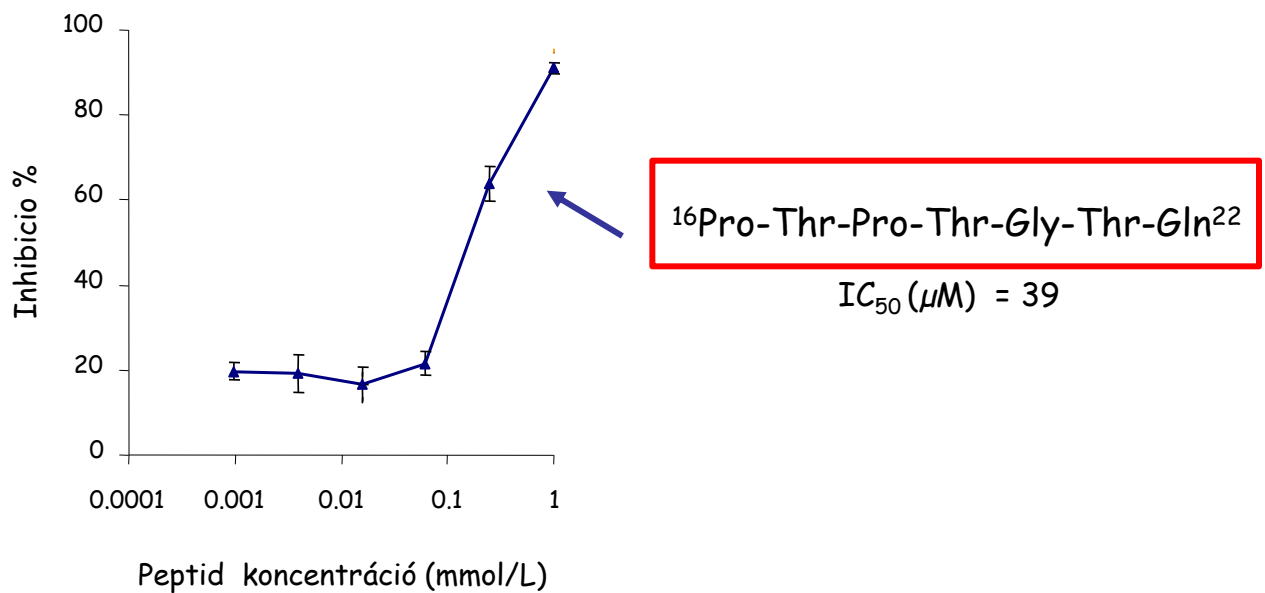


$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$



$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$

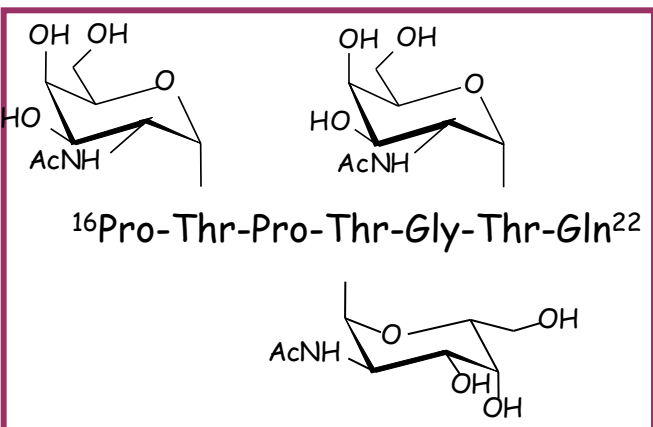
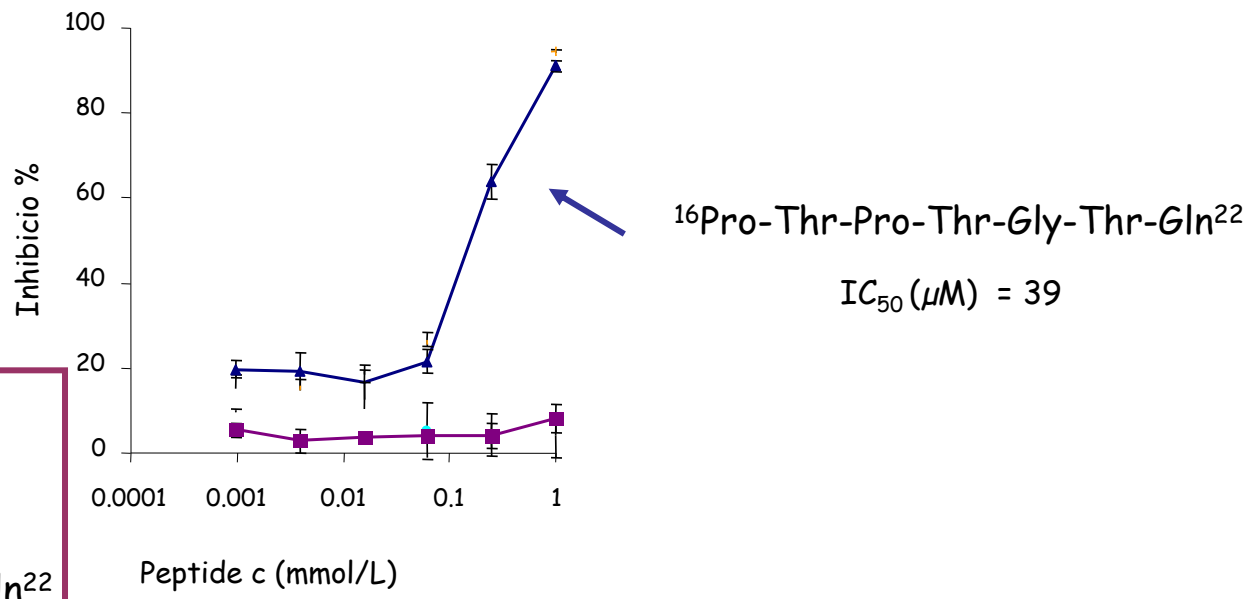
# The effect of carbohydrate moiety on MoAb binding



MAb 996:

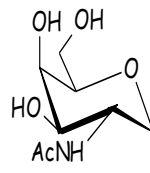
Anti-BSA-[K<sup>12</sup>VTPTPTPTGTQTP<sup>25</sup>]

# The effect of carbohydrate moiety on MoAb binding



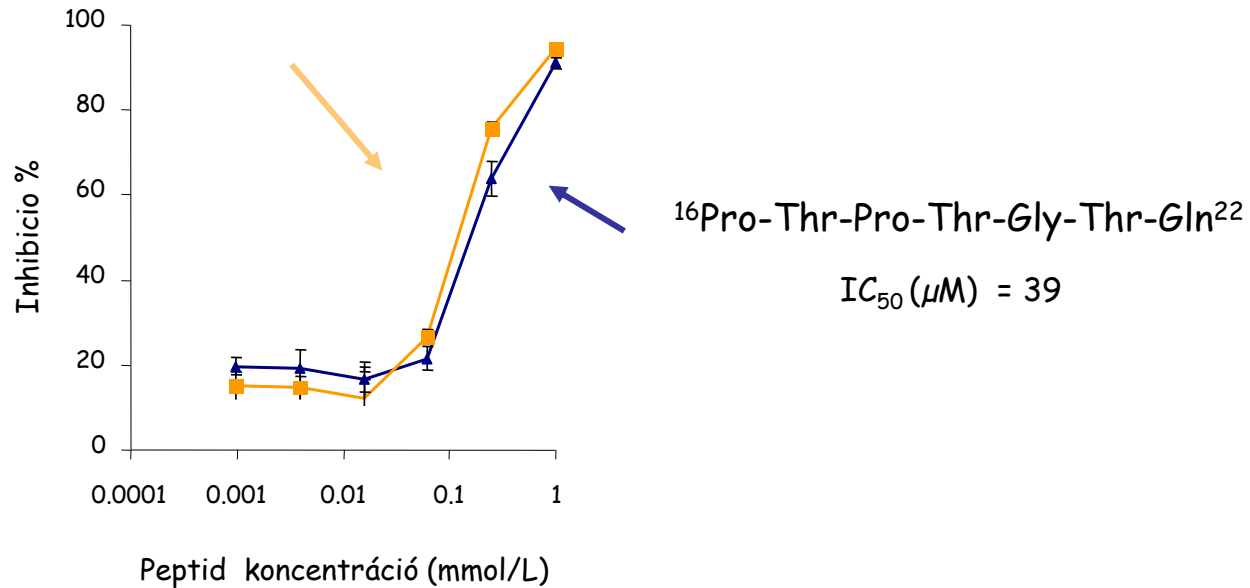
$\text{IC}_{50} (\mu\text{M}) = > 1000$

# The effect of carbohydrate moiety on MoAb binding



$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$

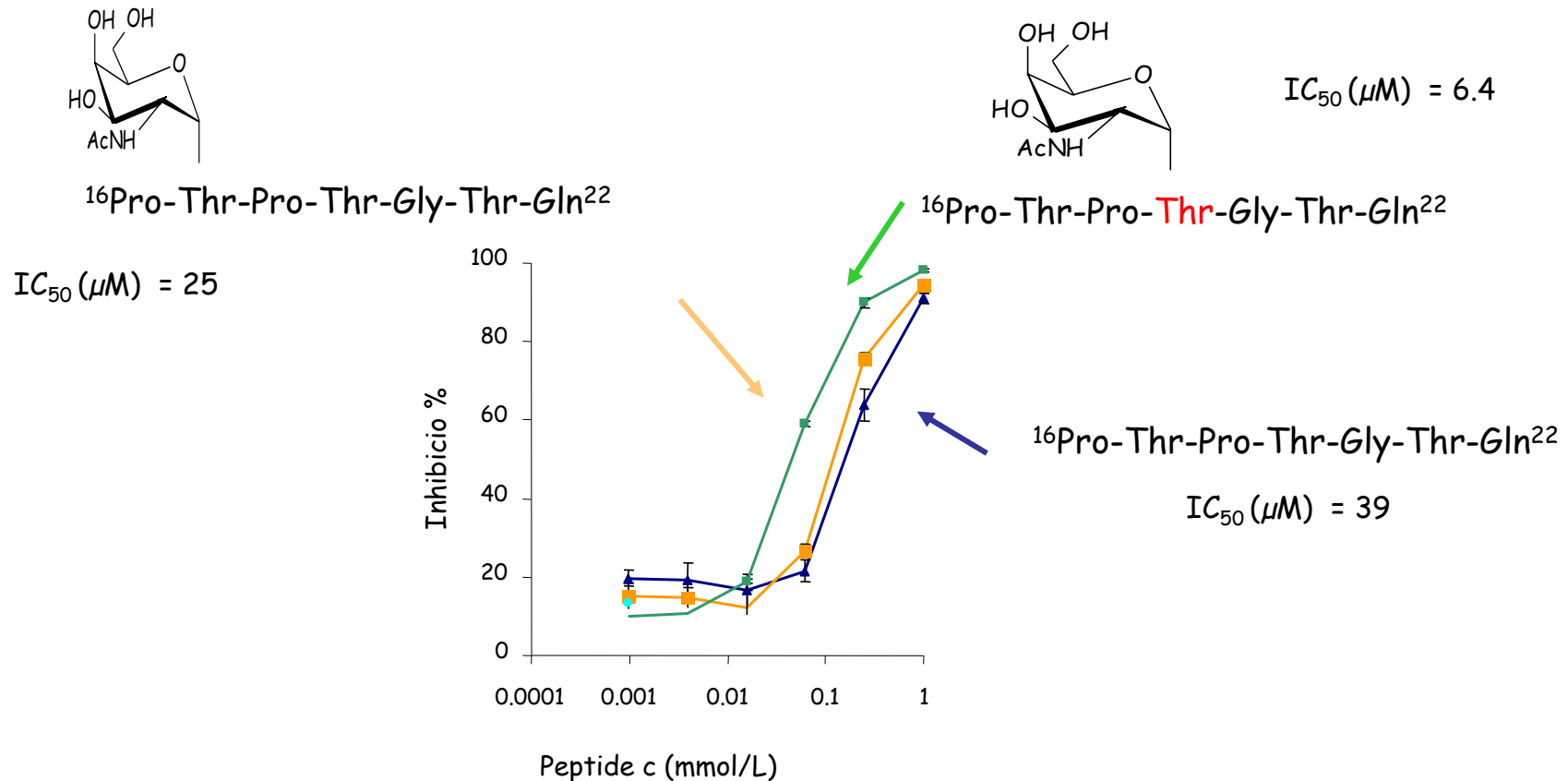
$IC_{50} (\mu\text{M}) = 25$



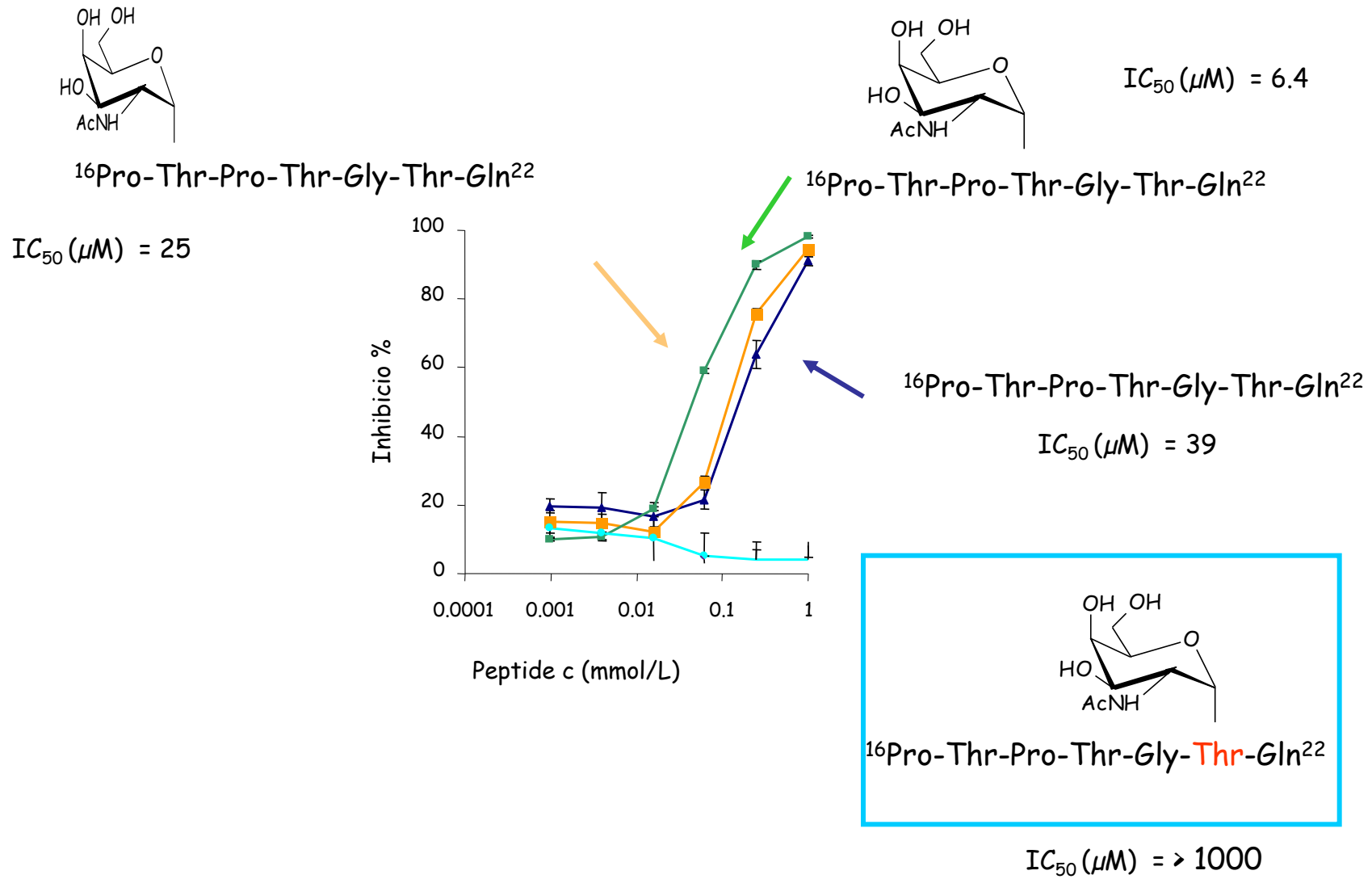
$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$

$IC_{50} (\mu\text{M}) = 39$

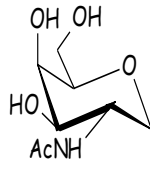
# The effect of carbohydrate moiety on MoAb binding



# The effect of carbohydrate moiety on MoAb binding

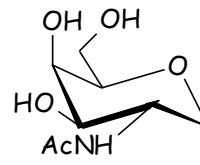


# The effect of carbohydrate moiety on MoAb binding

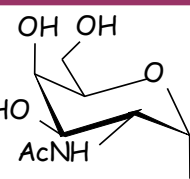
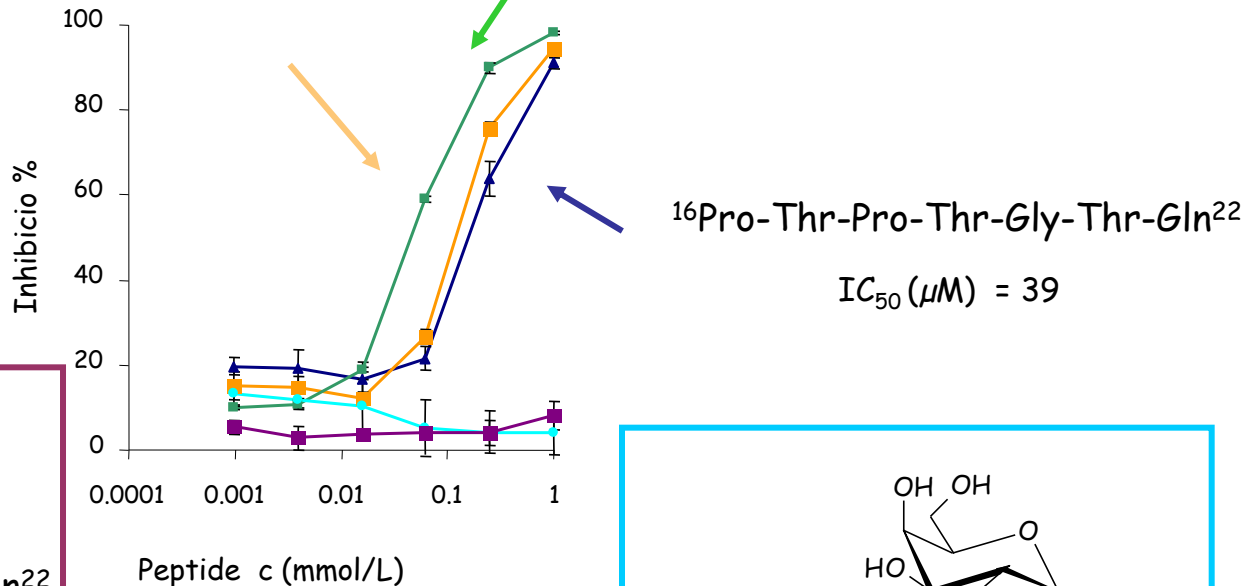


$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$

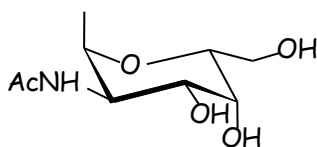
$IC_{50} (\mu\text{M}) = 25$



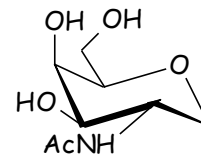
$IC_{50} (\mu\text{M}) = 6.4$



$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$



$IC_{50} (\mu\text{M}) = > 1000$



$^{16}\text{Pro-Thr-Pro-Thr-Gly-Thr-Gln}^{22}$

$IC_{50} (\mu\text{M}) = > 1000$

# ECD (Electronic Circular Dichroism)

Instrument: Jasco-810

Solvent:

- water
- TFE

Concentration: 0,5 mg/ml

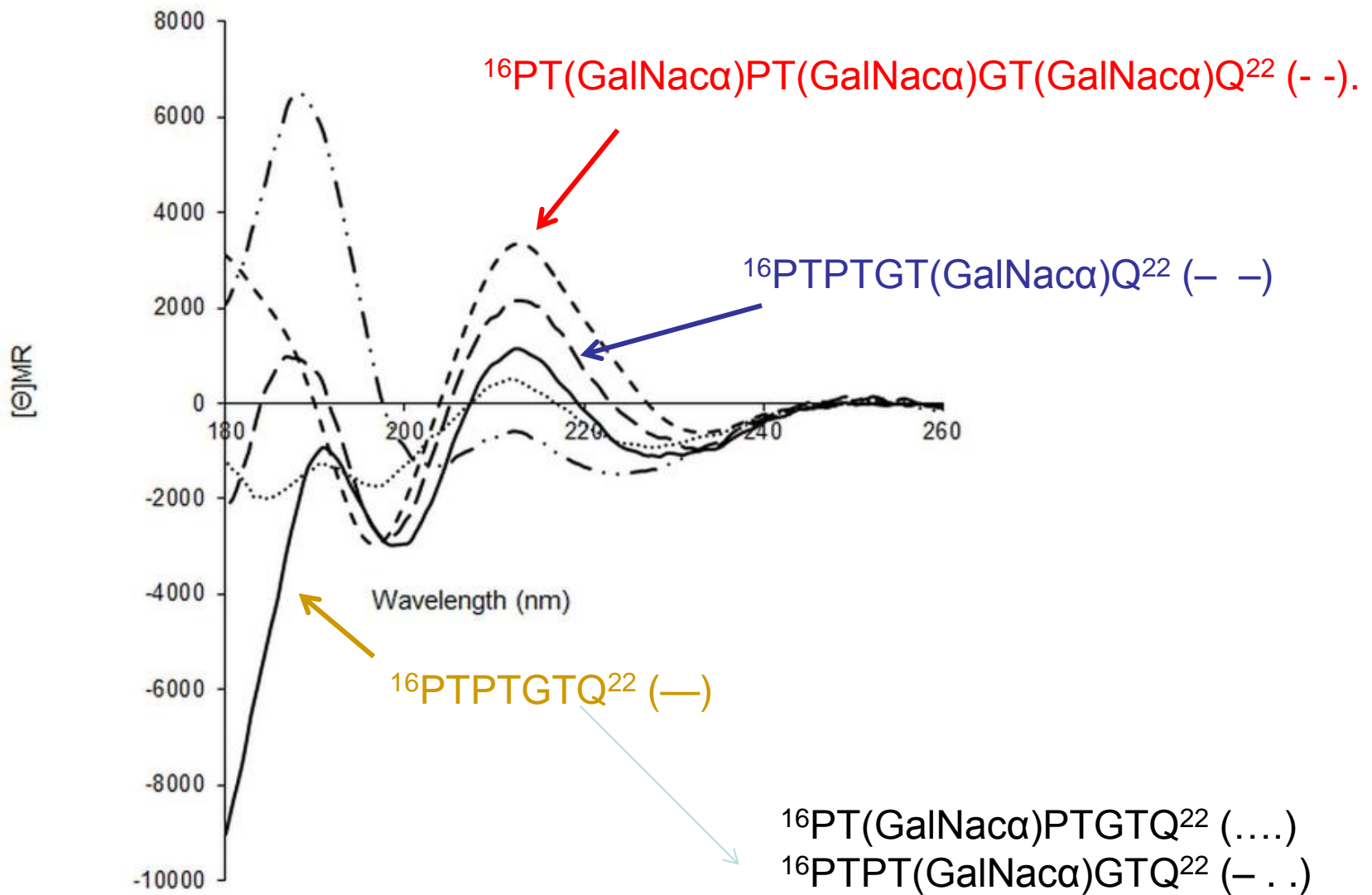
Wavelength:  $\lambda=180-300$  nm

0,02 cm quartz cuvette

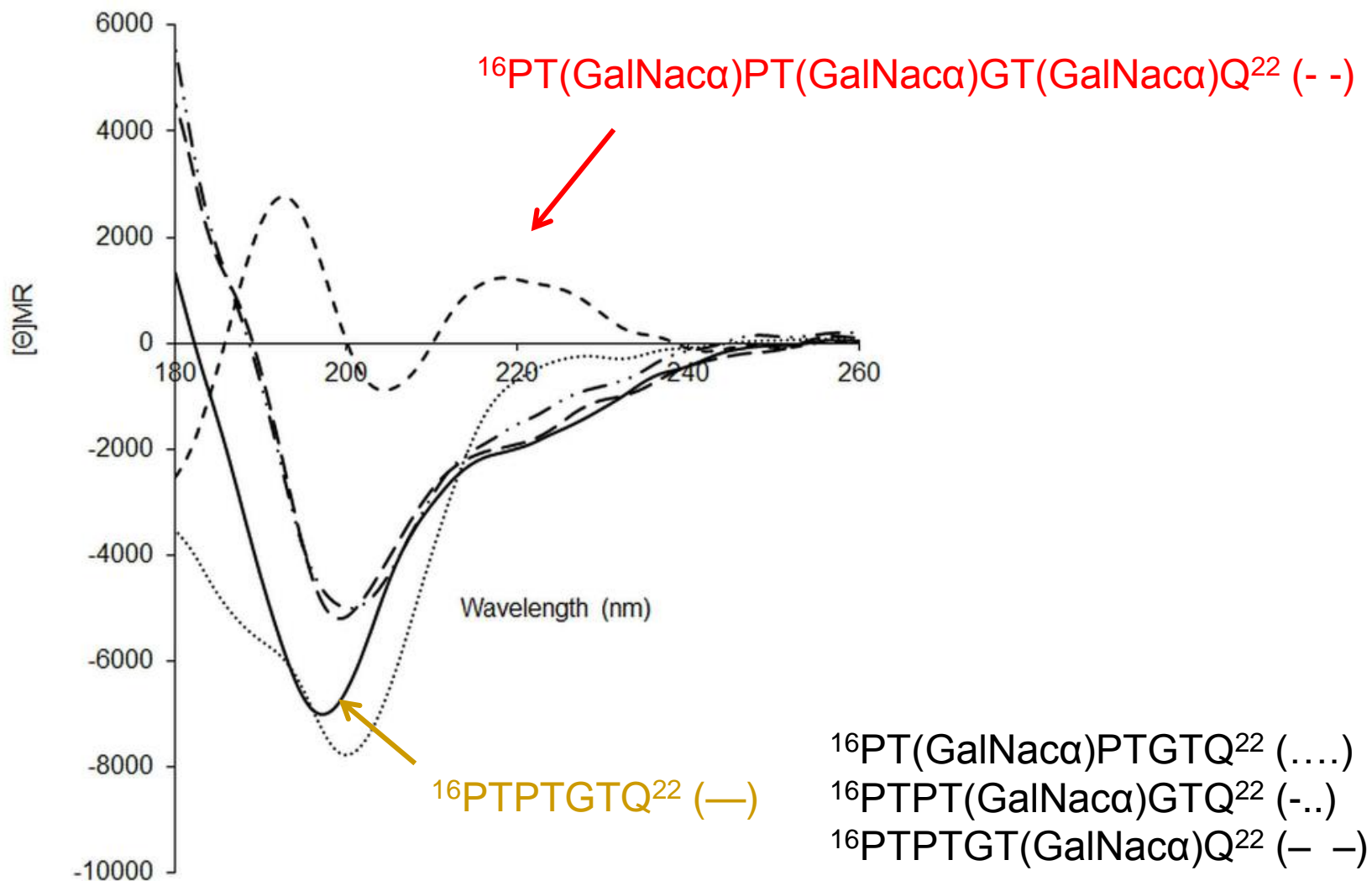


[http://www.andrew.cmu.edu/user/jamess3/  
JWSfac.html](http://www.andrew.cmu.edu/user/jamess3/JWSfac.html)

# The circular dichroism spectra of peptides in TFE

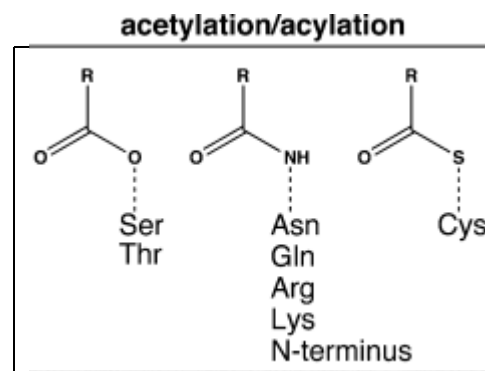
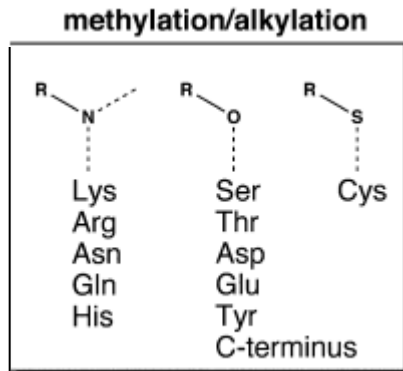
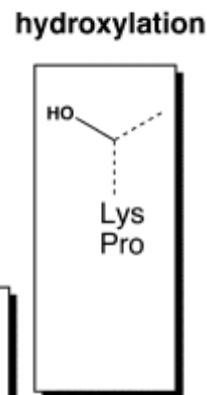
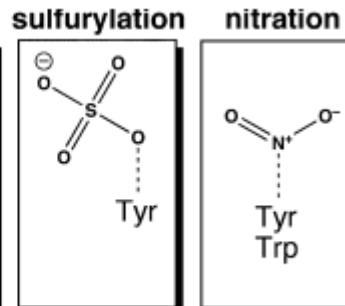
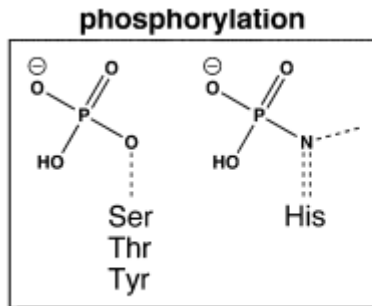
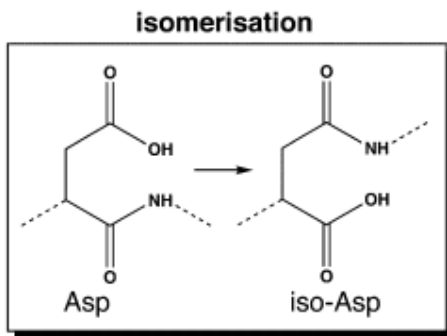
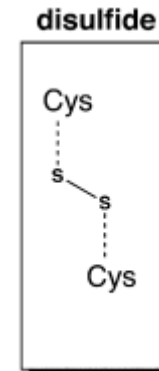
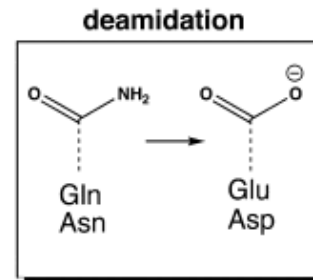
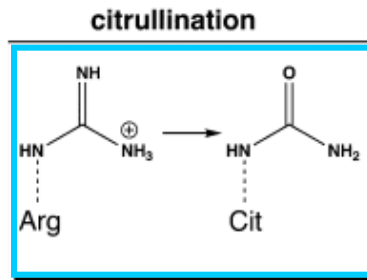
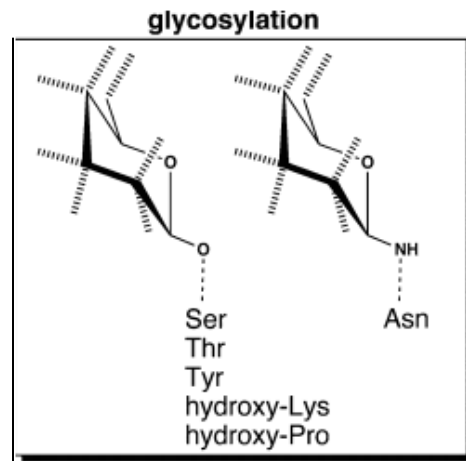


# The circular dichroism spectra of peptides in water

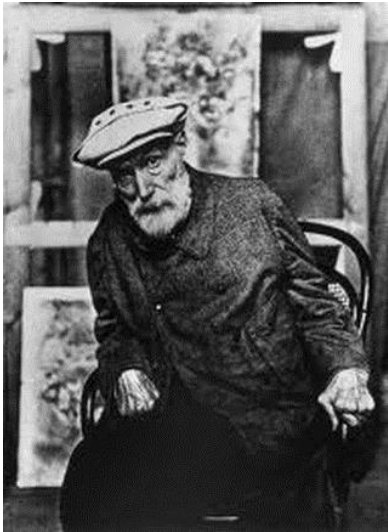


# The effect of post-translational modification on antibody recognition: citrullination

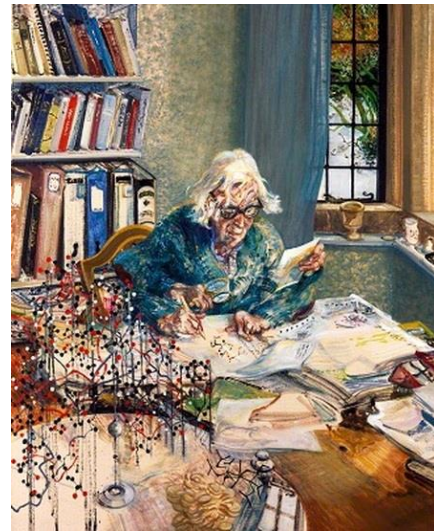
# Citrullination



# Rheumatoid arthritis



Pierre Auguste Renoir (1841 - 1919)



Dorothy C. Hodgkin (1910-1994)  
Nobel dij (1964)



Raoul Dufy (1877-1953)

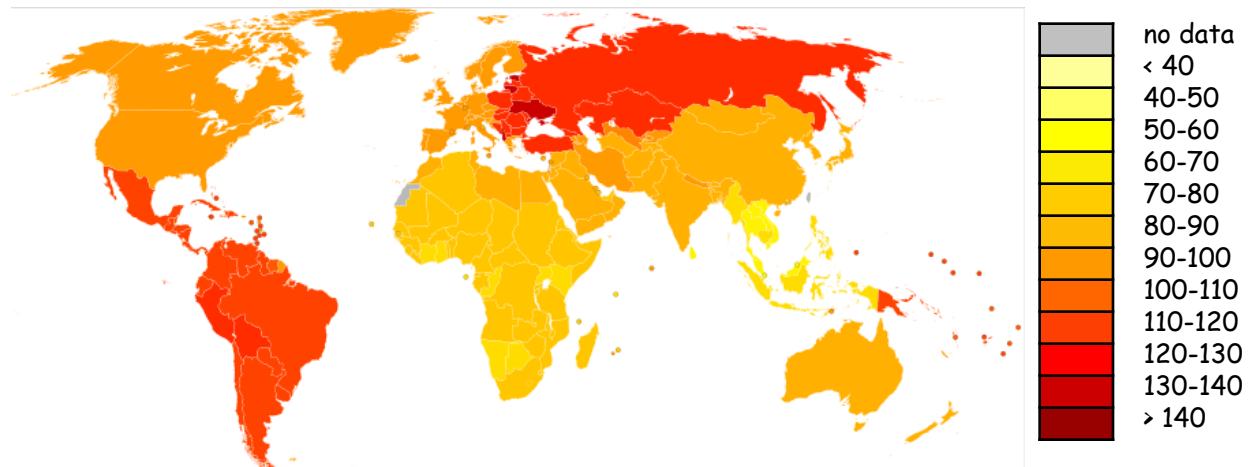
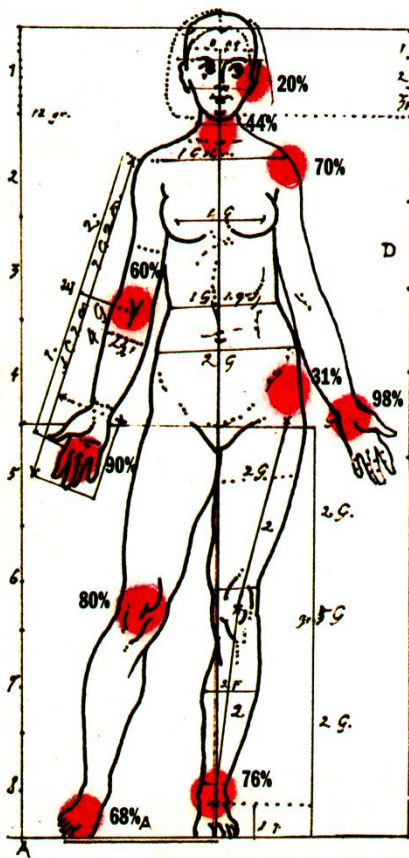


# Post-translational modification: Immune recognition related diseases

Modification	Autoantigen	Disease
Acetylation	Myelin basic protein	Multiple sclerosis
Citrullination	Collagen type II Myelin basic protein	RA Multiple sclerosis
Deamidation	Insulin	Type I diabetes
Glycosylation	Insulin proceptor Collagen type II Thyrotropin receptor Myelin oligodendrocyte glycoprotein	Diabetes RA Graves disease MS
Isoaspartylation	snRNP	Systemic lupus erythematosus
Lipoylation	PDC-E2	Primary biliary cirrhosis
Phosphorylation	Myelin basic protein	Multiple sclerosis
Methylation	Sm, D1,D3	Systemic lupus erythematosus
Transglutamination	Histone H2	Systemic lupus erythematosus
Tyrosine nitration	Mitochondrial proteins	Experimental autoimmune uveitis

# Rheumatoid Arthritis

- chronic, systemic inflammatory disorder
- systemic autoimmune disease
- attacks synovial joints
- hyperplasia of synovial cells,
- excess synovial fluid,



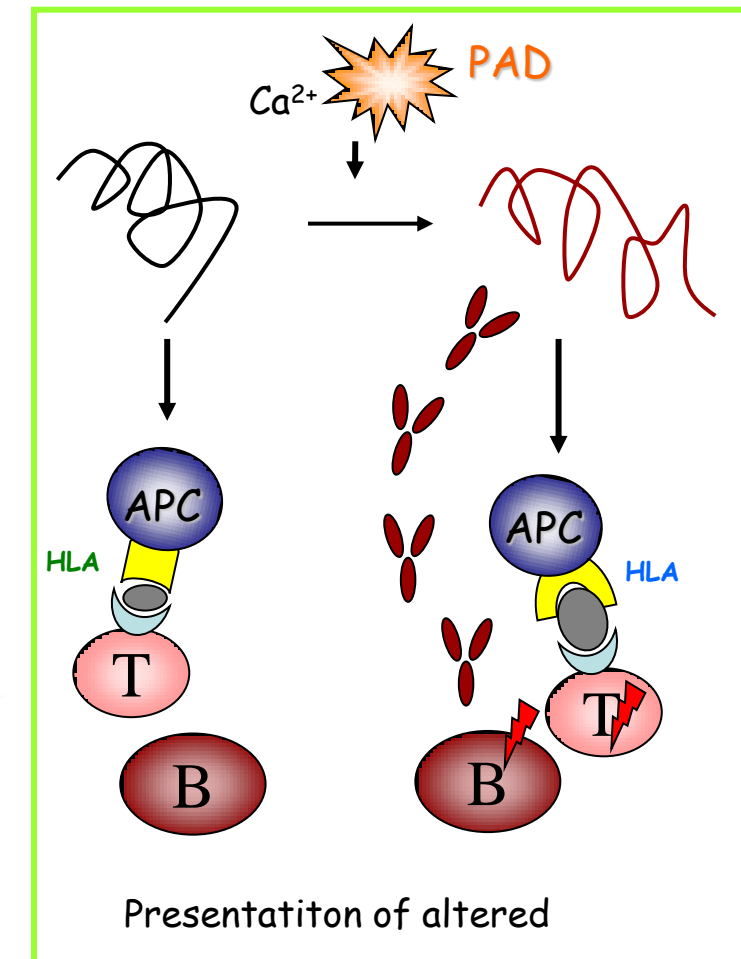
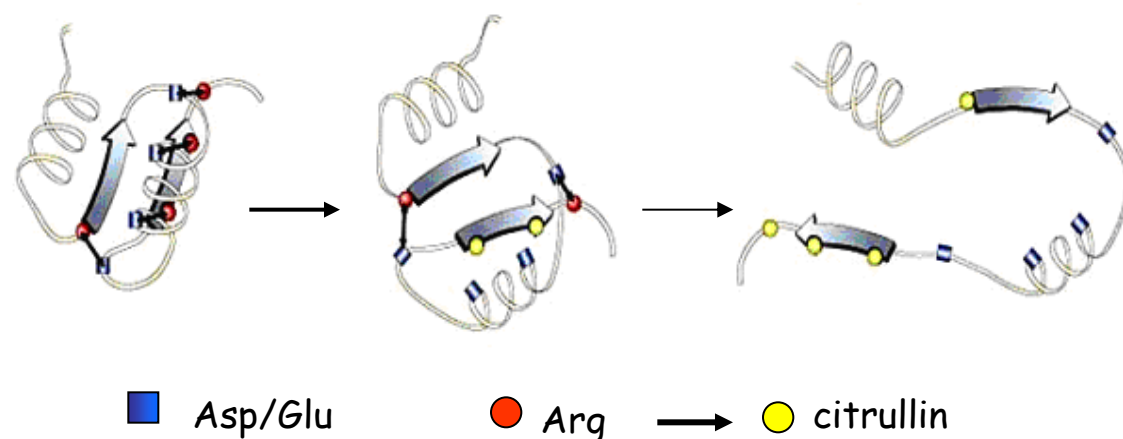
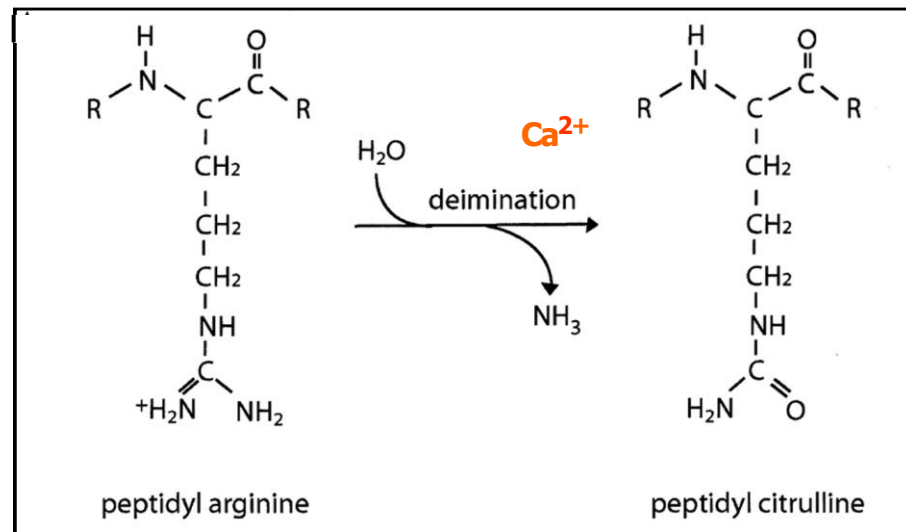
[http://en.wikipedia.org/wiki/File:Rheumatoid\\_arthritis\\_world\\_map\\_-\\_DALY\\_-\\_WHO2004.svg](http://en.wikipedia.org/wiki/File:Rheumatoid_arthritis_world_map_-_DALY_-_WHO2004.svg)



## Epidemiology

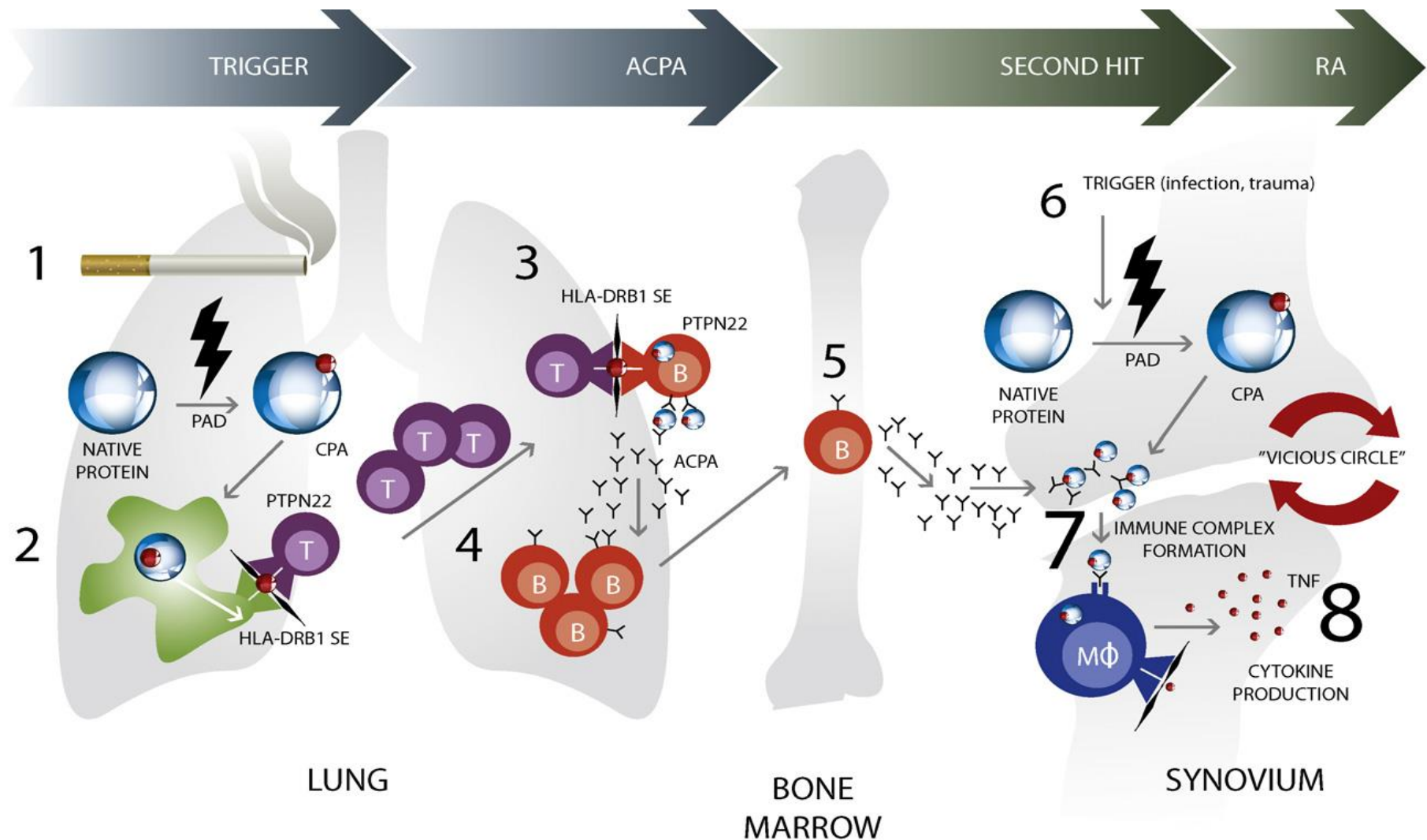
- 1% of the world's population
- women : man = 3:1
- most frequent ages 40 - 50

# The effect of post-translational modification on immune recognition: change in 3D structure of proteins



Yamada, R. et al. *Bioscience* 10: 54-64 (2005)  
 Yamada, R. *Autoimmunity Reviews* 4: 201-206 (2005)

# Immunity to citrullinated proteins in rheumatoid arthritis



# The effect of post-translational modification on immune recognition: proteins involved

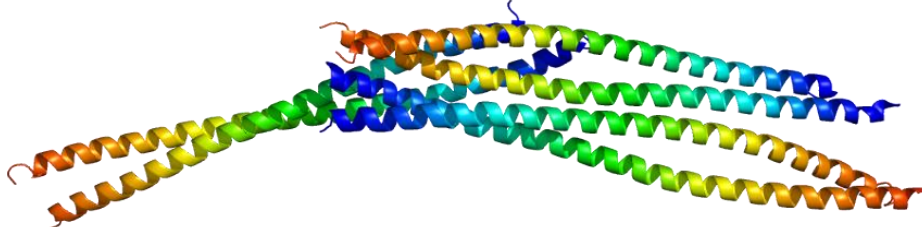
Disease	Modification	Antigen modified
RA	Hydroxylation	Type II collagen
	Glycosylation Oxidation Citrullination	Filagrin Fibrin Vimentin IgG
SLE	Glycosylation  Phosphorylation Deamidation Mannose modification Methylation Oxidation	Multiple snRNP D, H2B Multiple SM D1, D3 Cardiolipin, ox LDL, C1q, calreticulin

Eggleton, P. et al. *Rheumatology* 47: 567-571 (2008)



filaggrin<sup>1</sup>

vimentin<sup>2</sup>

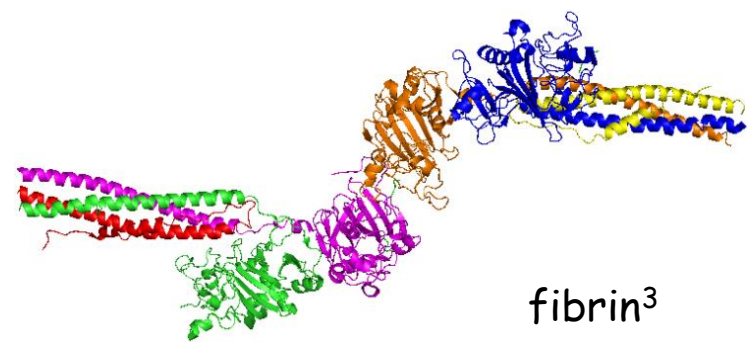


Based on [PyMOL](#) rendering of PDB [1gk4](#)

<sup>1</sup> Sebbag, M. et al. *Clin. Invest.* 95: 2672-2679 (1995)

<sup>2</sup> Vossenaar, E.R. et al. *Arthritis Res. Ther.* 6(2): 86-89 (2004)

<sup>3</sup> Masson-Bessiere, C. et.al. *J. Immun.* 166: 4177-4184 (2001)

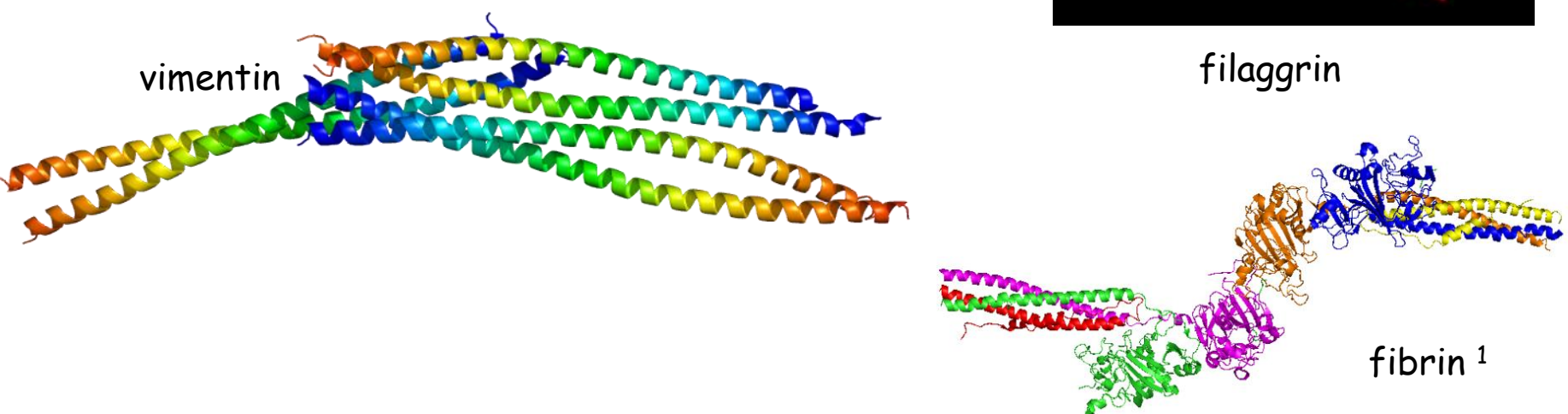
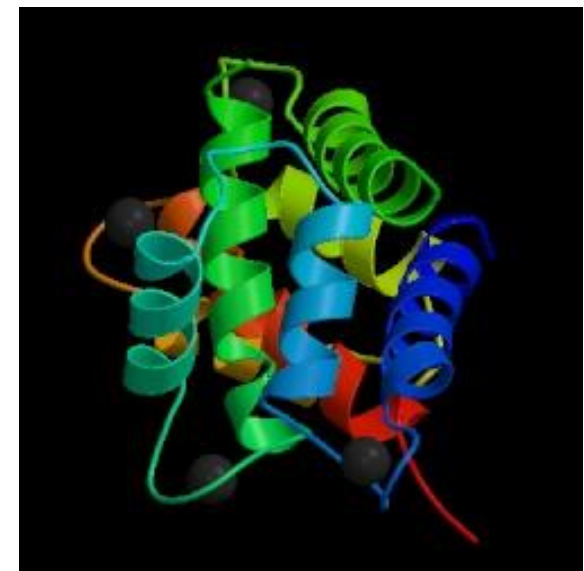


fibrin<sup>3</sup>

Crystal Structure of Fibrin from *Petromyzon marinus*, [1n73](#)  
<http://www.proteopedia.com/wiki/index.php/Image:1n73b.png>

# The effect of post-translational modification on immune recognition: epitope peptide

Vimentin	$^{65}\text{SAVRA}\textcolor{red}{R}\text{SSVPGV}\textcolor{red}{RK}^{77}$
Fibrin $\alpha$	$^{34}\text{GPRVV}\textcolor{red}{R}\text{HQSAKDS}^{48}$
Fibrin $\beta$	$^{60}\text{RPAPPPISSGGY}\textcolor{red}{RAR}^{74}$
Filaggrin (5-mer)	$^{311}\text{T}\textcolor{red}{RGRS}^{315}$
Filaggrin (19-mer)	$^{311}\text{SHQEST}\textcolor{red}{RGRSRGRSGRSGS}^{326}$



<sup>1</sup> Iobagiu C., Magyar, A. et al. *J. Autoimmunity* 37: 263-272 (2011)

## Aims

---

1. Identification of minimal and optimal antibody epitope of partially deimidated filaggrin by synthetic peptides based on 306-324 sequence using multi-pin approach and serum samples from diseased individuals.
2. Introduction of biotin label for soluble epitope peptide
3. Analyze
  - the effect epitope size and orientation on antibody recognition,
  - the effect the presence and position of biotin on solution conformation,
  - RA specificity in serum samples as compared with that of SLE and healthy individuals using the optimized peptide epitope by direct ELISA.

## Aims

---

1. Identification of minimal and optimal antibody epitope of partially deimidated filaggrin by synthetic peptides based on 306-324 sequence using multi-pin approach and serum samples from diseased individuals.
2. Introduction of biotin label for soluble epitope peptide
3. Analyze
  - the effect epitope size and orientation on antibody recognition,
  - the effect the presence and position of biotin on solution conformation,
  - RA specificity in serum samples as compared with that of SLE and healthy individuals using the optimized peptide epitope by direct ELISA.

# Filaggrin (filament aggregating protein)

(FILA\_HUMAN), <http://swissmodel.expasy.org/>

**profilaggrin: 4061 AA, 435170 Da**

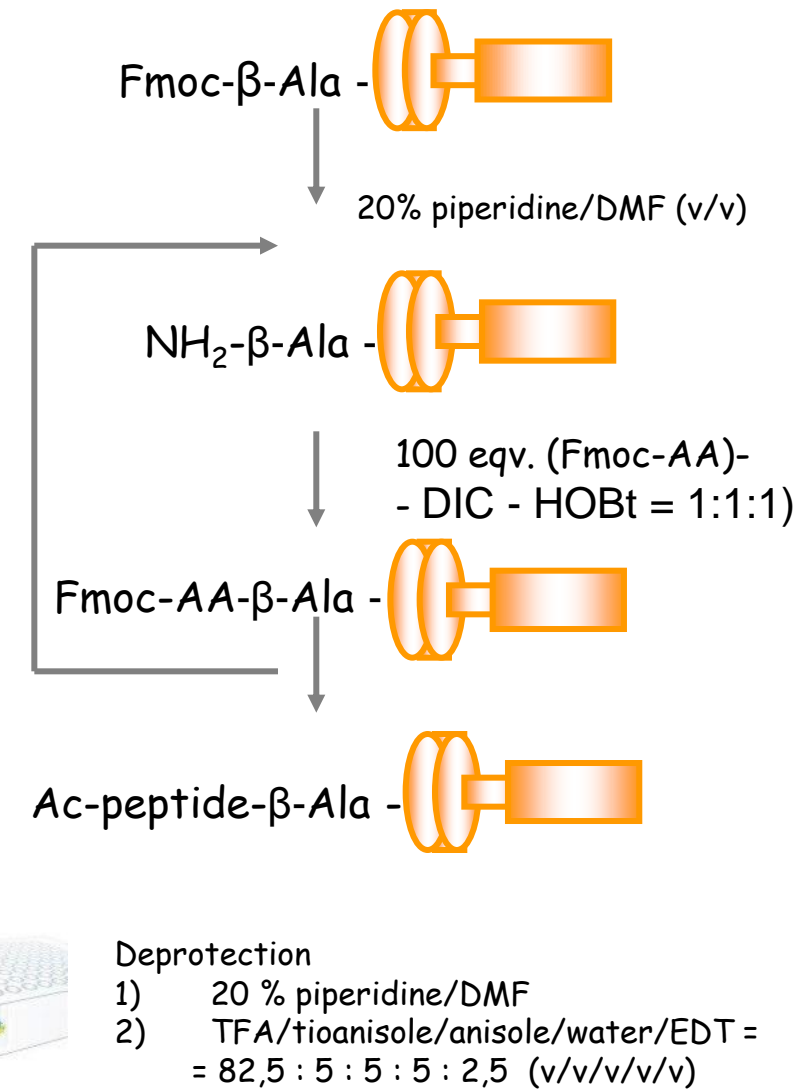
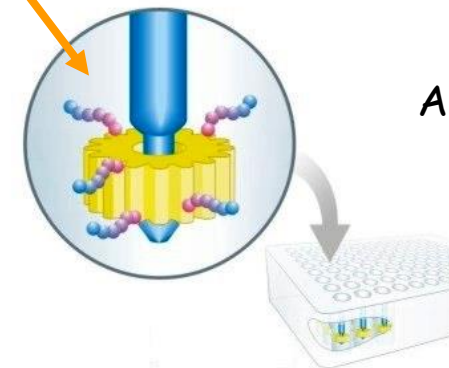
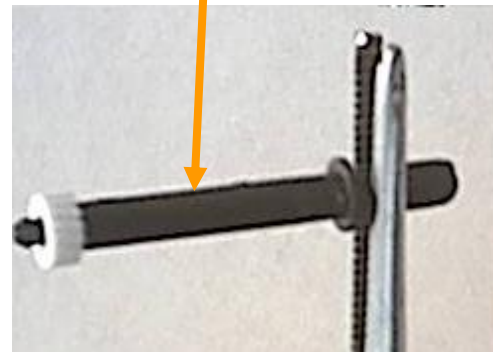
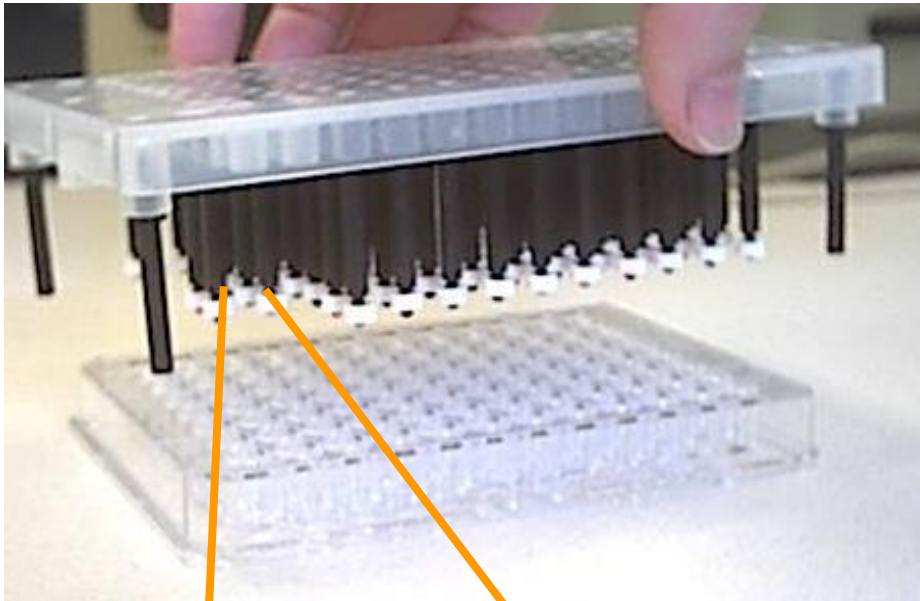
**10-12 filaggrin unit, 324 AA**

→ SHQESTRGRSRGRSGRSGS \*

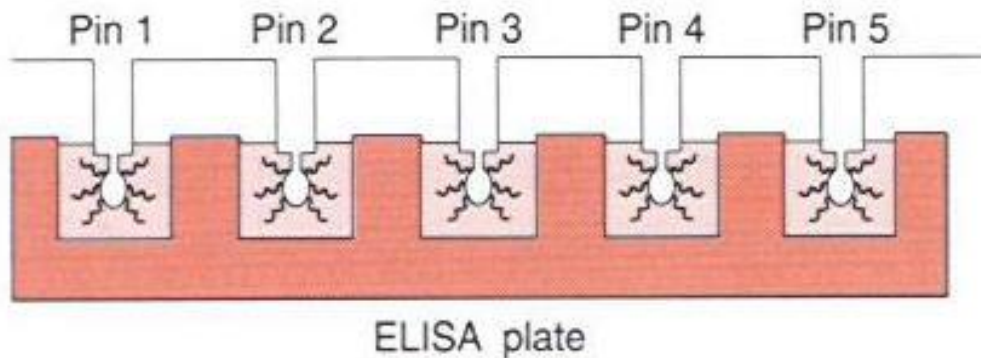
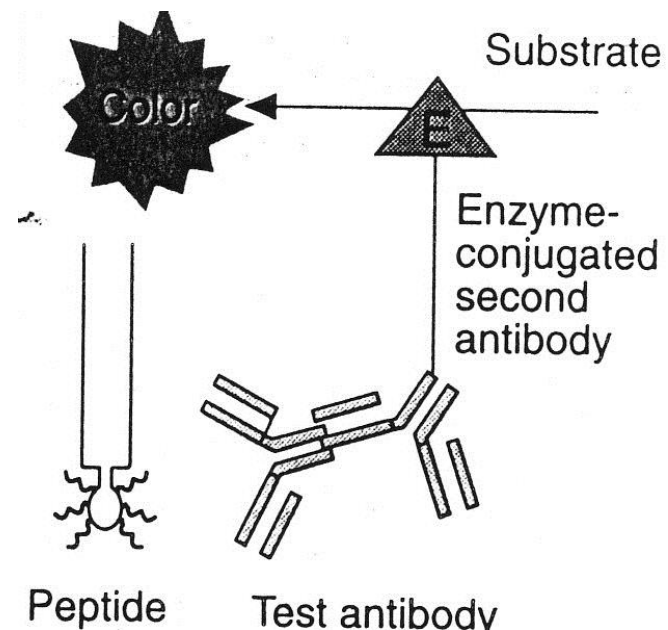
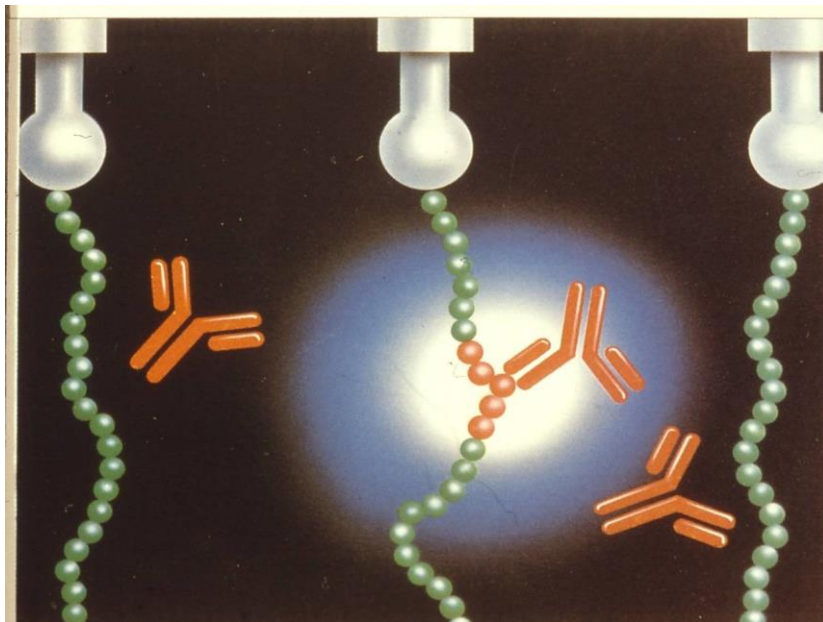


MSTLLENIFA IINLFKQYSK KDKNTDTLSK KELKELLEKE FRQILKNPDD PDMVDFMDH LDIDHNKKID FTEFLLMVFK LAQAYYESTR KENLPISGHK HRKHSHHDKH EDNKQEENKE NRKRPPSSLER RNNRKGNKGR SKSPRETGGK RHESSEKKE RKGYSPTHRE EEGYKHNHNS SKKEKNKTEN TRLGDNRKRL SERLEEKEDN EEGVYDYENT GRMTQKWIQS GHIATYYTIQ DEAYTTDSL LEENKIYERS RSSDGKSSSQ VNRSRHENTS QVPLQESRTR KRRGSRVSQD RDSEGHSEDS ERHSGSASRN HHGSAWEQSR DGSRHRPRSH EDRASTGHSA DSSRQSGTRH AETSSRGQTA SSHEQARSSP GERHGSHHQ SADSSRHSAT GRGQASSAVS DRGHGRGSSG QASDSEGHSE NSDTQSVSGH GKAGLRQSH QESTRGRSGE RSGRSGSSLV QVSTHEQPDS AHGRTGTSTG GRQGSHHEQA RDSSRHSASQ EGQDTIRGHP GSSRGGRQGS HHEQSVNRSG HSGSHHSHTT SQGRSDASHG QSGRSRSARQ TRNEEQSGDG TRHSGSRHHE ASSQADSSRH SQVGQGQSSG PRTSRNQGSS VSQDSDSQGH SEDSERWSGS ASRNHHGSAQ EQS RDGSRHP RSHHEDRAGH GHSADSSRK GTRHTQNSS GQAASSHEQA RSSAGERHGS RHQLQSDSS RHSGTGHGQA SSAVRDSGHR GSSGSQATDS EGHSEDSDTQ SVSGHGQHAG HQQSHQESAR DRSGERSRRS GSFLYQVSTH KOSESSHGW GPTSTGRQGS HHEQARDNSR HSASQDGQDT IRGHPGSSRR GRQGSHHEQS VDRSGHSGSH HSHTTSQGRS DASRGQSGSR SASRTTNEE QSRDGRSHSG SRHHEAASSA DISRHSQAGQ GQSEGSRSTS RQGSSVSQDS DSEGHSEDSE RWSGSASRNH RGSAQEFSQR GSRHPRSHHE DRAGHGSAD SSRQSGTPHA ETSSGGQAA SHEQARSSP ERHGSRHQHS ADSSRHSQIP RRQASSAVRD SGHWGSSGSQ ASDSEGHSE S DTQSVSGH QDGPHQHQSHQ ESARDWSSGR SGRSGSFYQ VSTHEQSEA HGRTRTSTGR RQGSHHEQAR DSSRHSASQE GQDTIRAHG RQGRRGQSH HEQSVDRSGH SGSHHSHTT QGRSDASHHQ SGSRSAQRT RDKQSGDGS RHSGSRHHEA ASWADSSRHS QVGQEQQSSG RTSRHQGSSV SQDSDSERHS DDSERLSGA SRNHGSSRE QSRDGRSHPG FHQEDRASHG HSADSSRQSG THHTESSSIHG QAVSSHEQAR SSPGERHGS HQQSDSSRH SGIGHRQASS AVRDSGHRGS SGSQVTNSEG HSEDSDTQSV SAHGOAGPHQ QSHKESARGQ SGESSGRSRS FLYQVSSHEQ SESTHGQTAP STGGRQGSRH EQARNSSRHS ASQDGQDTIR GHPGSSRGG QGSYHEQSV RSGHSGYHHS HTTPQGRSDA SHGQSGPRA SRQTRNEEQS GDGRSHGSR HEPSTRAGS SRHSQVGQGE SAGSKTSRRQ GSSVSQDRDS EGHSEDSEERR SESASRNHYG SAREQSRHGS RNPRSHQEDR ASHGHSAEQ RQSGTRHAET SSGGQAASSQ EQARSPGHER HGRHQQSAD SSTDSTGRR QDSSVVGDS NRGSSGSQAS DSEGHSEED TQSVAHGQA GPHQSHQES TRQSGSERG RSGSFLYQVS THEQSESAHG RTGPSTGGRQ RSRHEQARDS SRHSASQEGQ DTIRGHPGSS RGGRQGSHYE QSVDSGHSG SHHSHTTSQE RSDVSRQSG SRSVSRQTRN EKQSGDGSRH SGSRHHEASS RADSSRHSQV GQGQSSGPR SRNQGSSVSQ DSDSQGHSED SERWSGSASR NHLGSAWEQS RDGSRHPGSH HEDRAGHGH ADSSRQSGTR HTESSSRQQA ASSHEQARSS AGERHGSHHQ LQSAQDSRHS GIGHQASSA VRDSGHRYGS GSQASDSEGH SEDSDTQSVS AQGKAGPHQ SHKESARQGS GESSGRSGSF LYQVSTHEQ STHGQSQS TGGRQGSHY QAQDSSRHS ASEQGQDTIR HPGPSRGGQ GSHQEVSQDR SGHSGSHHS TTSQGRSDAS RGQSGRSRS RKTYDKEQSG DGSRHGS SHHEASSWDSS RHLVGQGQS SGRTSRPRG SSVSQDSDE GHSEDSEERR GSASRNHHGS ASEQRSRDGS HPRSHHEDRA GHGHSAEASSR QSGTHAENS SGGQASSHEA QARSSAGERH GSHHQSQADS SRHSGIGHHQ ASSAVRDGSH RGSSQSQASD SEGHSEDSDT QSVSAHGQAG PHQSHQEST RGRSAGRSGR SGFLYQVST HEQSESAHGR TGTTGGRQG SHHKQARDSS RHTSQEGQD TIHGPSSS GGRQGSHYEQ LVDRSGHSGS HHSHTSQGR SDASHGHSGS RSASRQTRND EQSGDGRHS GSRHHEASSR ADSSGHSQVG QQQSEGPRTS RNWGSSFSQD SDSQGHSEDS ERWGSASRN HHGSAQEQLR DGSRHPRSQH EDRASTGHSA DSSRQSGTRH TQTSSGGQAA SSHEQARSSA GERHGSHHQ SADSSRHSQI GHGQASSAVR DSGHRYGSGS QASDNEGHSE DSDTQSVSAH GQAGSHHQSH QESARGRSGE TSGHSGSFLY QVSTHEQSES SHGWTGPSTR GRQGSRHEQA QDSSRHSASQ DGQDTIRGHP GSSRGGRQGY HHEHSDSS HGSQHHSHTT SQGRSDASRG QSGRSRSASRT TRNEEQSGDG SRHSGSRHHE ASTHADISR SQAVQGQSEG SRRSRRQGSS VSQDSDSEGH SEDSERWSGS ASRNHHGSAQ EQLRDSRHP RSHQEDRAGH GHSADSSRQS GTRHTQTSSG GQAASSHEQA RSSAGERHGS HHQSQSDSS HSGIGHQAS SAVRDSGHRG YSGSQASDNE GHSEDSDTQS VSAHGQAGSH QSQHQSARG RSGETSGHSG SFLYQVSTHE QSESSHGWGWT PSTRGRQGSR HEQAQDSSRH SASQYQGDTI RGHPGSSRGG RQGYHHEHSV DSSGHGSHH SHTTSQGRSD ASRGQSGRS ASRTRNEEQ SGDSSRHSVS RHHEASTHAD ISRHSQAVQG QSEGSRRSRR QGSSVSQSD SEGHSEDSEER WSGSASRNHR GSVQEQRHG SRHPRSHHED RAGHGHSAD SRQSGTRHAE TSSGGQASSHE QOARSSPGE RHGSRHQQA DSSRHSQIP GQASSAVRDS RHGSSGSGQA SDSEGHSEES DTQSVSGHQQ AGPHQSQHQE SARDRSGGRS GRSGFLYQV STHEQSESAH GRTRTSTGR QGSHHEQARD SSRHSASQEG QDTIRGHPG SRRGRQGSHY EQSVDRSGHS GSHHSHTSQ GRSDASRQGS GRSASRQTR NDEQSGDGR HSWSHHEAS TQADSSRHSQ SGQGQSGAP TSRNQGSSV QDSDSQGH DSERWGSAS RNHGRQSAEQ SRDGRSHPTS HHEDRAGHGH SAESSRQGST HHAENSSGGQ AASSHEQARS SAGERHGSII QSQADSSRHS GIGHGQASSA VRDGSRHPG GSQASDSEGH SEDSDTQSVS AHQAGPHQ SHQESTRGRS RGRSGRSGSF LYQVSTHEQS ESAHGRAGPS TGGRQGSRHE QARDSSRHSQ SQEGQDTIRG HPGSRRGGQ GSYHEQSVDR SGHGSHHH TTSGRQSDAS HGQSGRSRSAS RETRNEEQSG DGSRHSGRH HEASTQADSS RHSQSGQGES AGSRRSRRQG SSVSQDSDE AYPEDSEERRS ESASRNHHGS SREQSRDGS RHPGSSHRTA SHVQSPVQS DSSTAKEHGH FSSLSQDSAY HSGIQSRGSP HSSSSYHYQS EGTERQKGQS HGSVS

# Search for minimal/optimal epitope: Multi-pin approach



# Identification of linear antibody epitopes



# *In vitro* analysis of antibody recognition

---

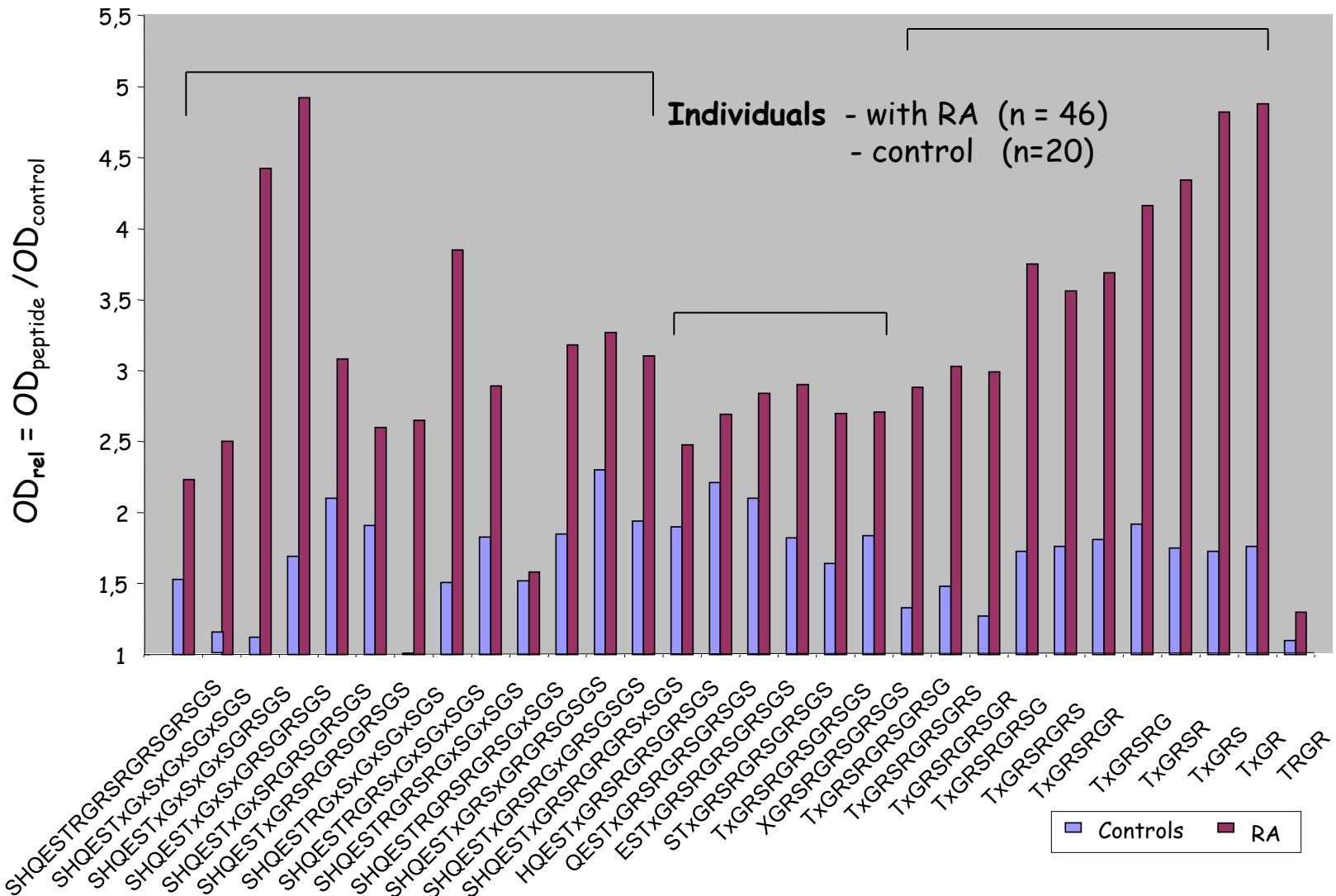
- Peptides
  - C-terminally pin-bound oligopeptides
  - free peptides labelled with biotin at the N-terminal
  - free peptides labelled with biotin at the C-terminal
- Serum samples:
  - from healthy CCP positive individuals,
  - from diseased CCP positive RA individuals,
  - from healthy CCP negative individuals,
- Synovial fluid samples:
- direct ELISA

# Search for minimal/optimal epitope

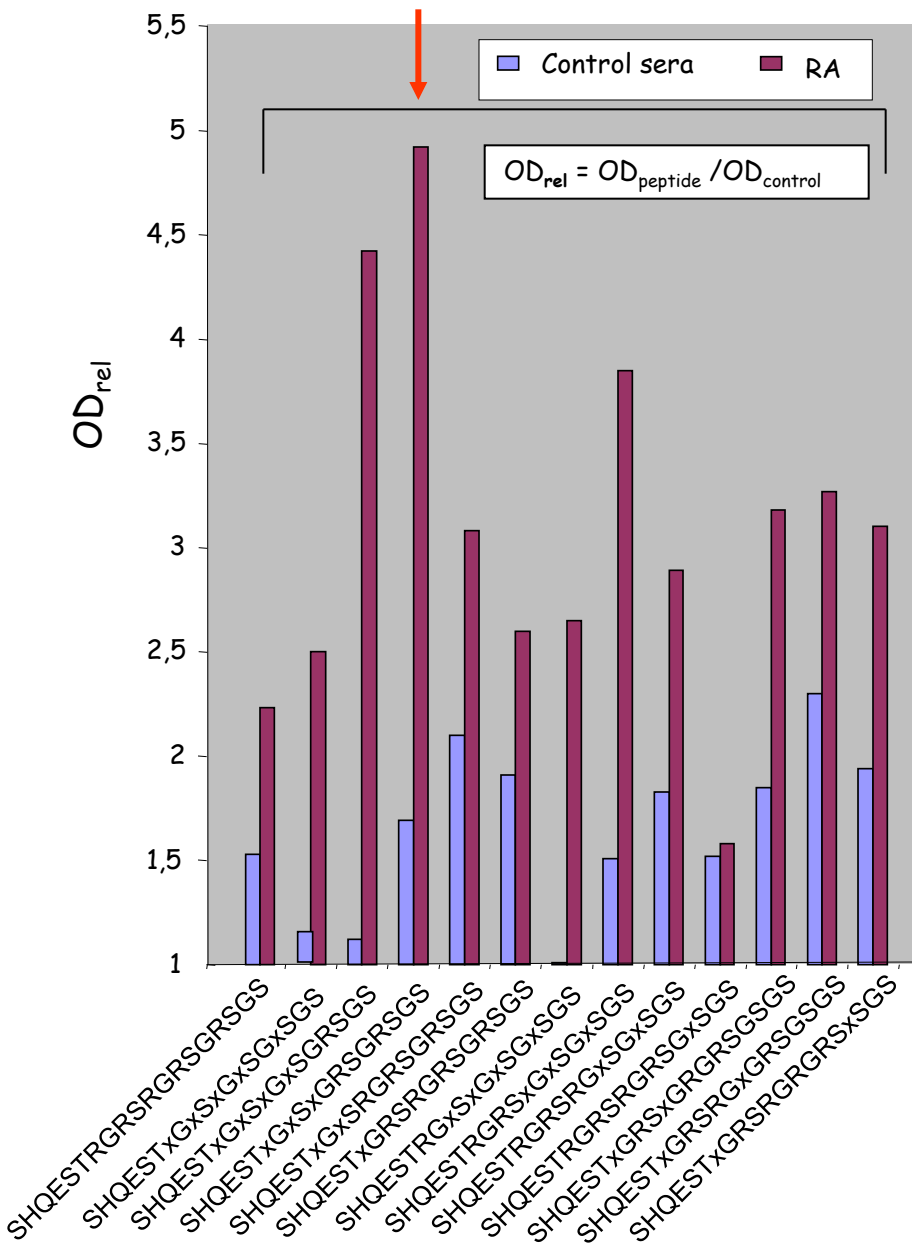
19-mer analogues	N-terminal truncation of peptide <i>Cit</i> <sup>312</sup> (306-324)	C-terminal truncation of peptide 311-324
306 SHQESTRGRSRGRSGRSGS <sup>324</sup>	306 SHQESTXGRSRGRSGRSGS <sup>324</sup>	311 TXGRSRGRSGRSGS <sup>324</sup>
306 SHQESTXGXSXGXSGXSGS <sup>324</sup>	307 HQESTXGRSRGRSGRSGS <sup>324</sup>	311 TXGRSRGRSGRSG <sup>323</sup>
306 SHQESTXGXSXGXSGRSGS <sup>324</sup>	308 QESTXGRSRGRSGRSGS <sup>324</sup>	311 TXGRSRGRSGRS <sup>322</sup>
306 SHQESTXGXSXGRSGRSGS <sup>324</sup>	309 ESTXGRSRGRSGRSGS <sup>324</sup>	311 TXGRSRGRSGR <sup>321</sup>
306 SHQESTXGXRGRSGRSGS <sup>324</sup>	310 STXGRSRGRSGRSGS <sup>324</sup>	311 TXGRSRGRSG <sup>320</sup>
306 SHQESTXGRSRGRSGRSGS <sup>324</sup>	311 TXGRSRGRSGRSGS <sup>324</sup>	311 TXGRSRGRS <sup>319</sup>
306 SHQESTRGXSXGXSGXSGS <sup>324</sup>	312 XGRSRGRSGRSGS <sup>324</sup>	311 TXGRSRGR <sup>318</sup>
306 SHQESTRGRSXGXSGXSGS <sup>324</sup>		311 TXGRSRG <sup>317</sup>
306 SHQESTRGRSRGXSGXSGS <sup>324</sup>		311 TXGRSR <sup>316</sup>
306 SHQESTRGRSRGRSGXSGS <sup>324</sup>	Control peptides	311 TXGRS <sup>315</sup>
306 SHQESTXGXRGRSGRSGS <sup>324</sup>	PLAQGGGGGG	311 TXGR <sup>314</sup>
306 SHQESTXGRSXGRSGRSGS <sup>324</sup>	GLAQGGGGGG	311 TRGR <sup>314</sup>
306 SHQESTXGRSRGRSGXSGS <sup>324</sup>		

(X=citrullin)

# Search for minimal/optimal epitope



# Analogues 19-mer peptides: Critical Cit residue(s)

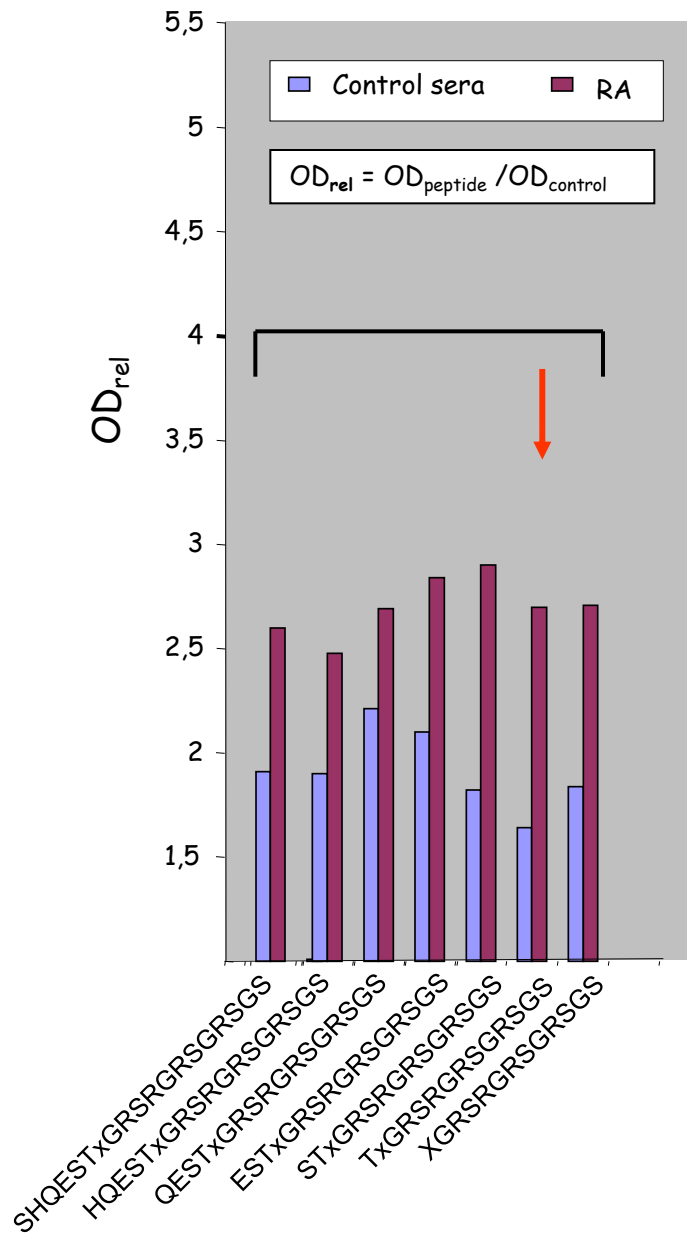


19-mer analogues
306 SHQESTRGRSXRGSRSGS <sup>324</sup> *
306 SHQESTXGSXGSXGS <sup>324</sup>
306 SHQESTXGSXGSXGRSGS <sup>324</sup>
306 SHQESTXGSXGSXGRSGS <sup>324</sup>
306 SHQESTXGSXSRGRSGS <sup>324</sup>
306 SHQESTXGRSRGRSGS <sup>324</sup>
306 SHQESTRGXGSXGSXSGS <sup>324</sup>
306 SHQESTRGRSXRGSXSGS <sup>324</sup>
306 SHQESTRGRSRGXSGXSGS <sup>324</sup>
306 SHQESTRGRSRGXSXSGS <sup>324</sup>
306 SHQESTRGRSRGSGXSGS <sup>324</sup>
306 SHQESTXGSXSRGRSGS <sup>324</sup>
306 SHQESTXGRSXGRSGRSGS <sup>324</sup>
306 SHQESTXGRSRGRSGXSGS <sup>324</sup>

(X = citrullin)

\*based on Hu-profilaggrin cDNA aa 306-324

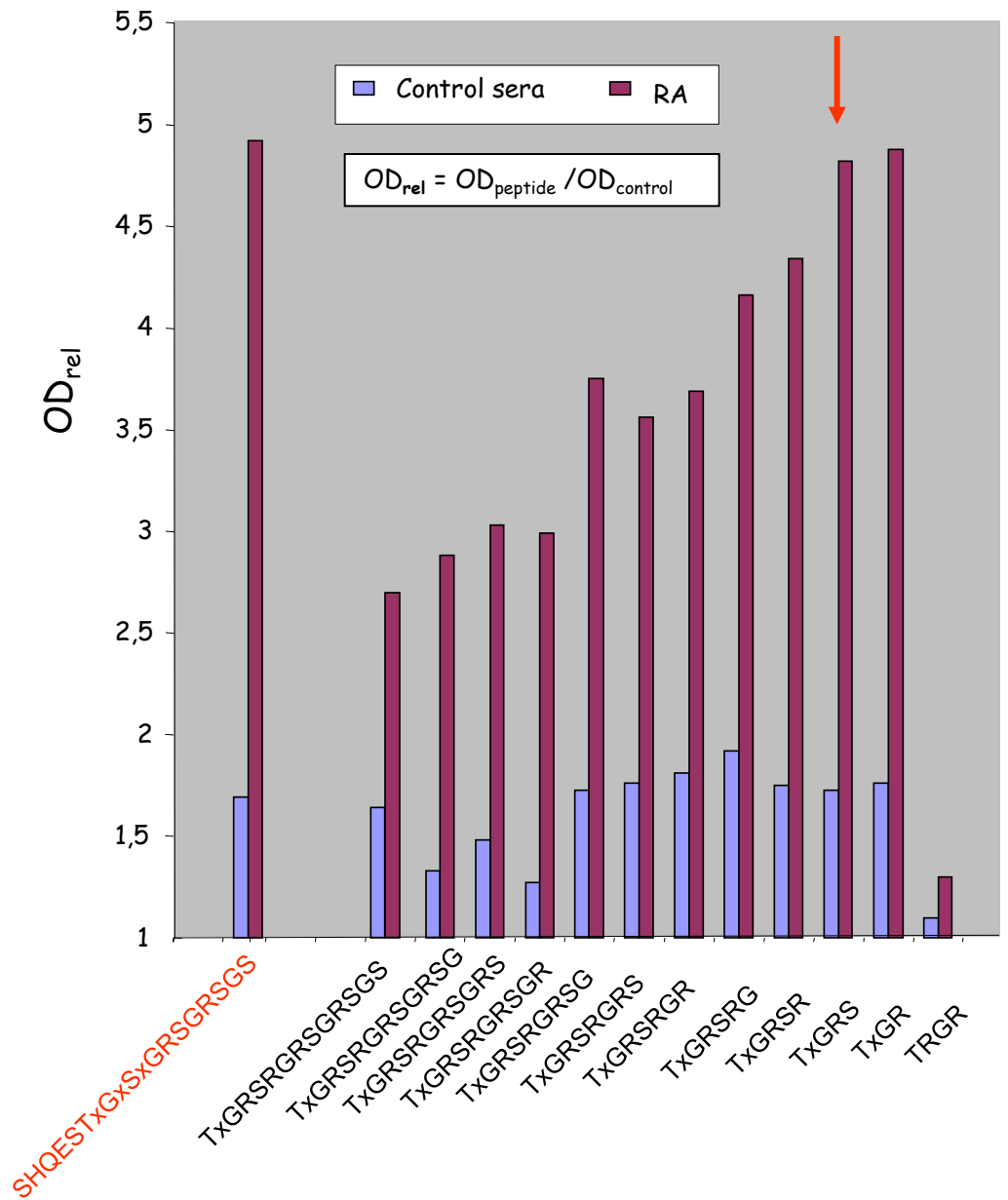
# N-terminal truncation of peptide Cit<sup>312</sup> (306-324)



N-terminal truncation of peptide Cit <sup>312</sup> (306-324)	
306	SHQESTXGRSRGRSGS <sup>324</sup>
307	HQESTXGRSRGRSGS <sup>324</sup>
308	QESTXGRSRGRSGS <sup>324</sup>
309	ESTXGRSRGRSGS <sup>324</sup>
310	STXGRSRGRSGS <sup>324</sup>
311	TXGRSRGRSGS <sup>324</sup>
312	XGRSRGRSGS <sup>324</sup>

(X = citrullin)

# C-terminal truncation of peptide Cit<sup>312</sup> (311-324)



C-terminal truncation of peptide 311-324
311 TXGRSRGRSGRS <sup>324</sup>
311 TXGRSRGRSGRS <sup>323</sup>
311 TXGRSRGRSGRS <sup>322</sup>
311 TXGRSRGRSGR <sup>321</sup>
311 TXGRSRGRSG <sup>320</sup>
311 TXGRSRGRS <sup>319</sup>
311 TXGRSRGR <sup>318</sup>
311 TXGRSRG <sup>317</sup>
311 TXGRSR <sup>316</sup>
311 TXGRS <sup>315</sup>
311 TXGR <sup>314</sup>
311 TRGR <sup>314</sup>

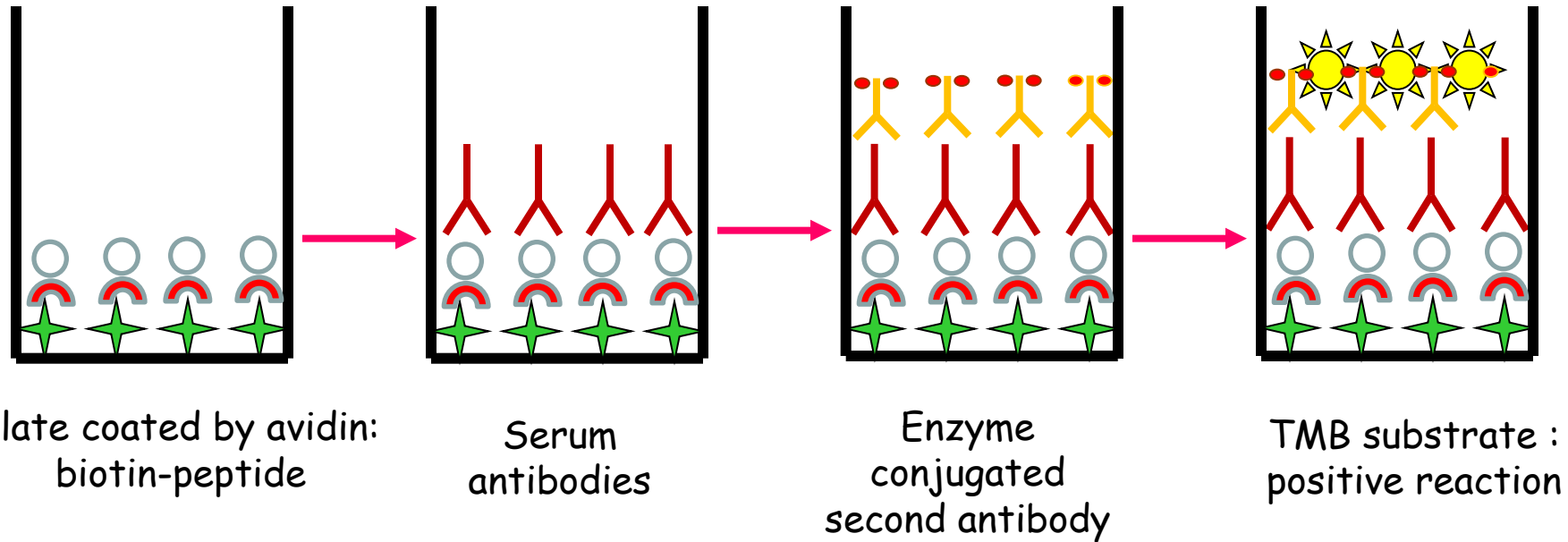
(X = citrullin)

# Aims

---

1. Identification of minimal and optimal antibody epitope of partially deimidated filaggrin by synthetic peptides based on 306-324 sequence using multi-pin approach and serum samples from diseased individuals.
2. Introduction of biotin label for soluble epitope peptide
3. Analyze
  - the effect epitope size and orientation on antibody recognition,
  - the effect the presence and position of biotin on solution conformation,
  - RA specificity in serum samples as compared with that of SLE and healthy individuals using the optimized peptide epitope by direct ELISA.

# Analysis of antibody binding to biot- $\beta$ 60-74Cit/Arg peptide by direct ELISA



Avidin



Serum antibodies  
(healthy/RA)



Tetramethylene  
benzidine (TMB)

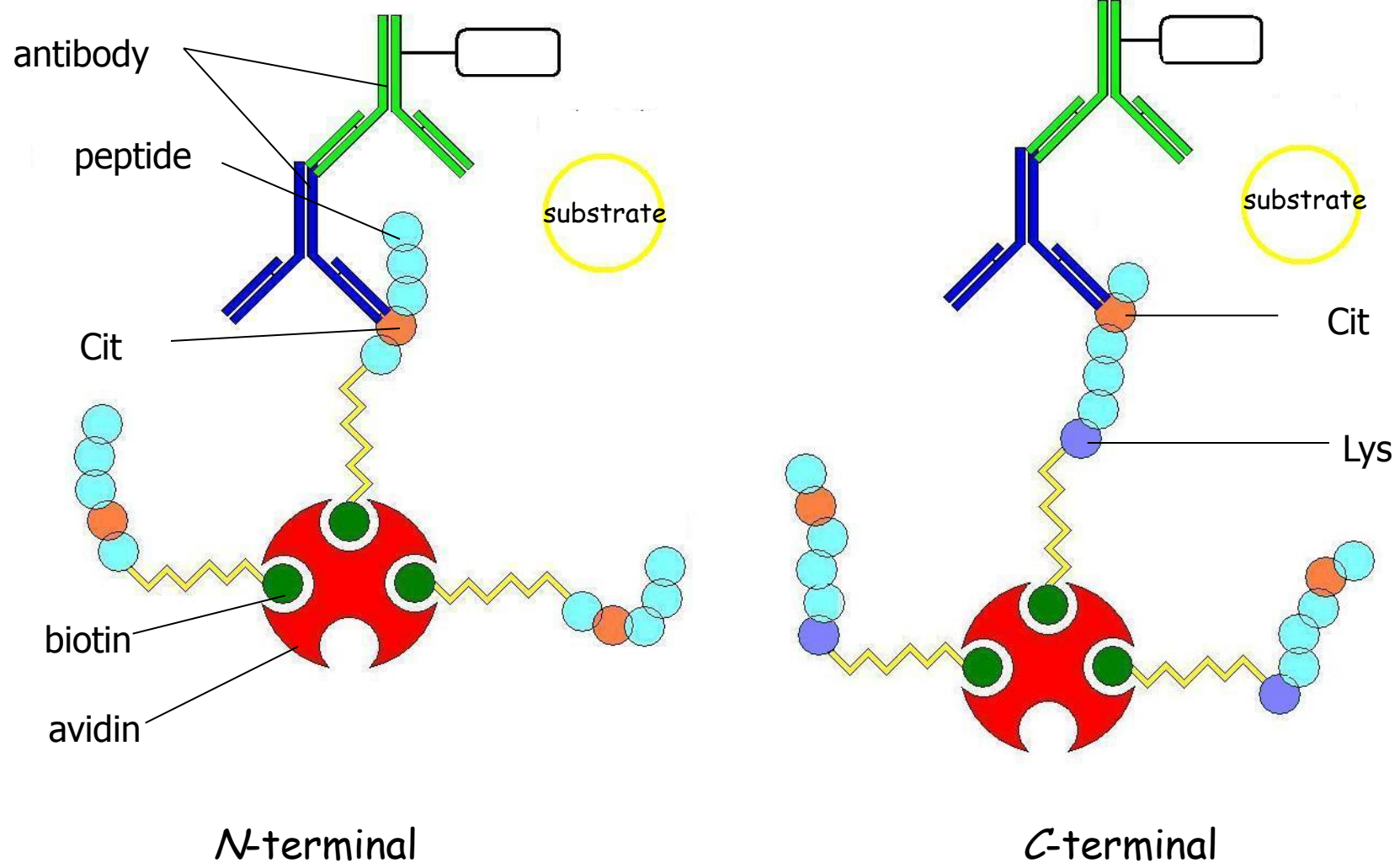


Biot- $\beta$ 60-74Cit/Arg peptide

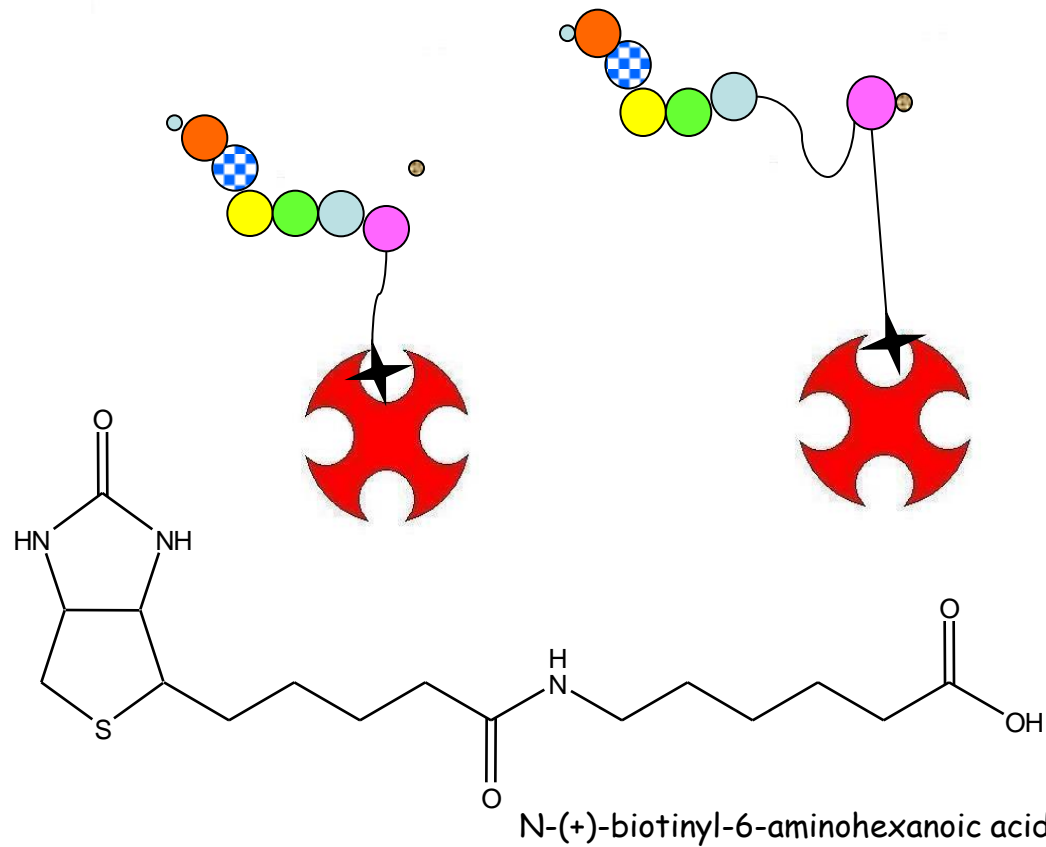
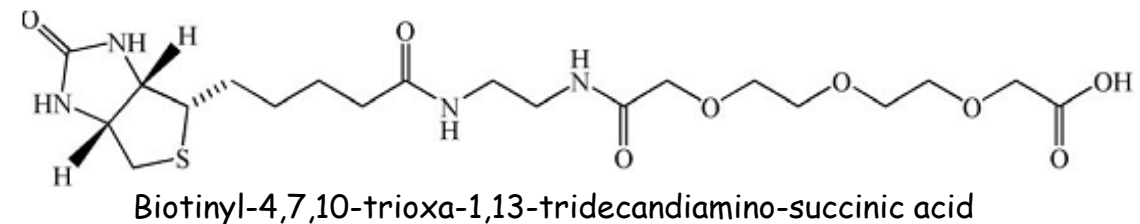
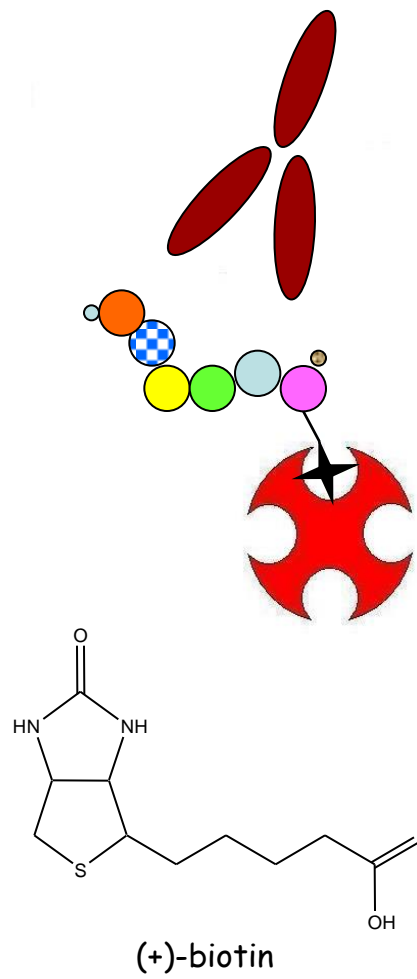


HRP enzyme conjugated  
anti human IgG antibodies

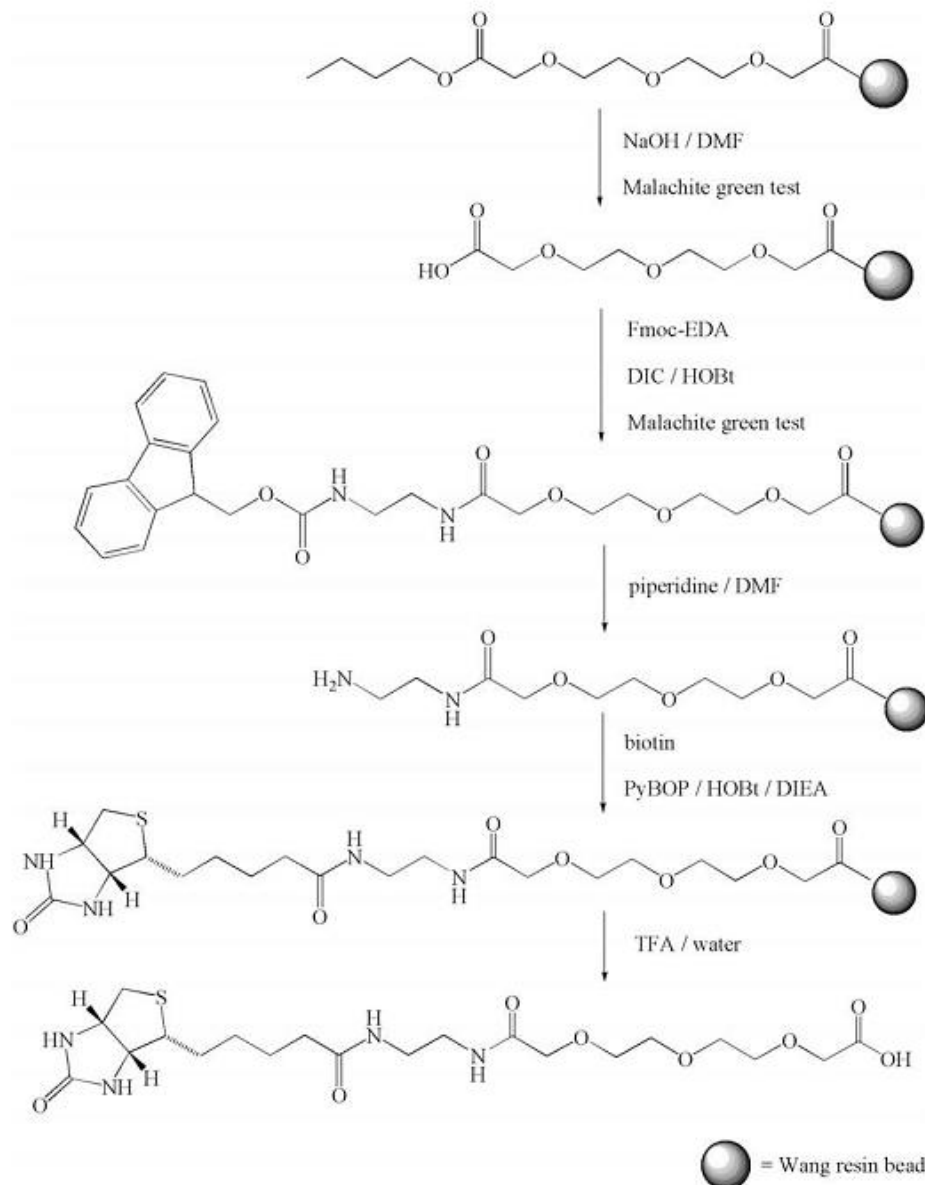
# The effect of epitope orientation: the position of biotin



# The effect of epitope accessibility: linkers



# Synthesis of biotinyl-4,7,10-trioxa-1,13-tridecandiamino-succinic acid



Bartos, Á. et al.  
*Biopolymers* 92: 110-115 (2009)  
Bartos, Á. et al.  
*Tetrahedron Letters* 50: 2661-2663 (2009)

## Aims

---

1. Identification of minimal and optimal antibody epitope of partially deimidated filaggrin by synthetic peptides based on 306-324 sequence using multi-pin approach and serum samples from diseased individuals.
2. Introduction of biotin label for soluble epitope peptide
3. Analyze
  - the effect epitope size and orientation on antibody recognition,
  - the effect the presence and position of biotin on solution conformation,
  - RA specificity in serum samples as compared with that of SLE and healthy individuals using the optimized peptide epitope by direct ELISA.

# Analysis of serum samples

## Samples:

- 263 RA patients with established disease,
- 46 CCP negative, non-RA patients with other autoimmune diseases
- 18 patients with systemic lupus erythematosus
- 152 age-matched healthy controls

The diagnosis of the disease was established on the basis of the revised ACR/EULAR classification criteria.<sup>1</sup>

## The baseline data of RA patients:

32 men/176 women; age: 58,4 +/- 14,3 years;  
rheumatoid factor (RF) +/-: 127/30; CCP2 +/-: 157/27; MCV +/-: 164/25;  
disease duration: 9,8 +/- 9,4 years.

## Statistical analysis:

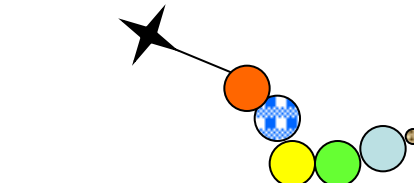
ANOVA, compared with Pearson's correlation analysis

1. Aletaha, D., Neogi ,T., Silman, A.J. et al. 2010 rheumatoid arthritis classification criteria: an American College of Rheumatology/European League Against Rheumatism collaborative initiative. *Ann Rheum Dis* 69:1580-8 (2010).

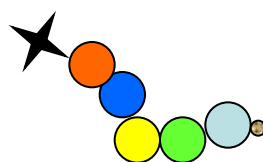
# Synthesis of 5-mer epitope peptide with *N*- terminal biotin



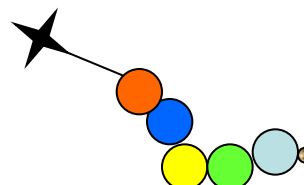
biotinyl-TXGRS-NH<sub>2</sub>



biotinyl-6-aminohexanoyl-TXGRS-NH<sub>2</sub>



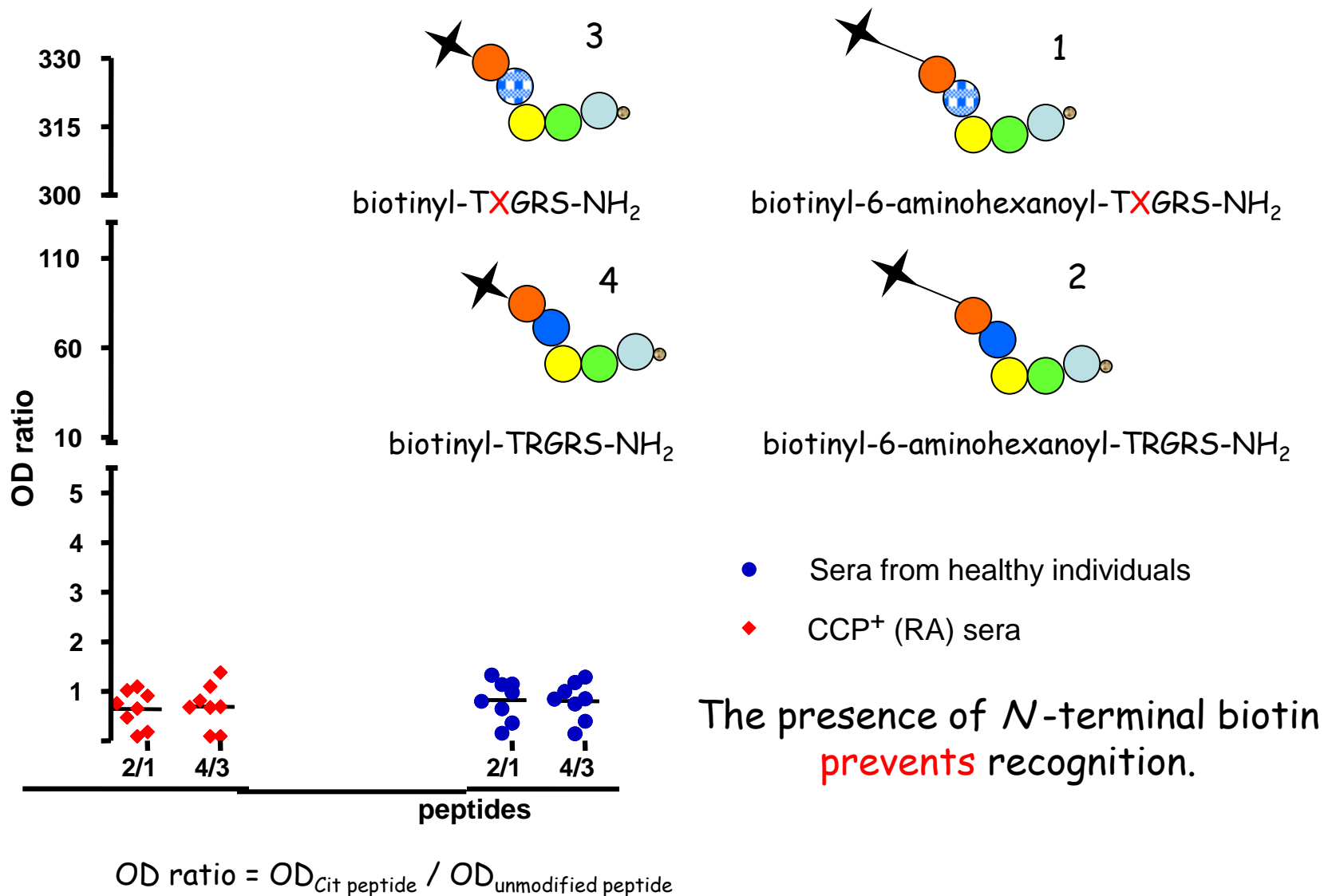
biotinyl-TRGRS-NH<sub>2</sub>



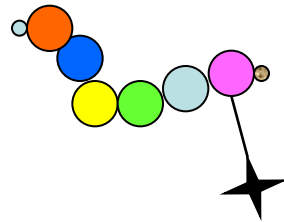
biotinyl-6-aminohexanoyl-TRGRS-NH<sub>2</sub>

Compound	M <sub>av</sub> (calc)	M <sub>av</sub> (meas)	R <sub>t</sub> (min)
biotinyl-TRGRS-NH <sub>2</sub>	802,9	802,8	14,23
biotinyl-TXGRS-NH <sub>2</sub>	803,9	803,7	14,12
biotinyl-6-aminohexanoyl-TRGRS-NH <sub>2</sub>	914,1	913,9	17,07
biotinyl-6-aminohexanoyl-TXGRS-NH <sub>2</sub>	915,1	914,9	16,50

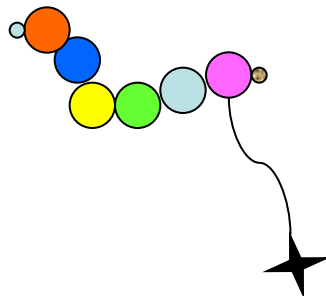
# Antibody recognition of 5-mer epitope peptides with *N*-terminal biotin



# Synthesis of 5-mer epitope peptides with *C*-terminal biotin

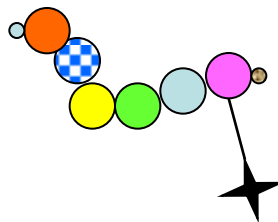


Ac-TRGRS**K**(biotinyl-hexanoyl)-NH<sub>2</sub>

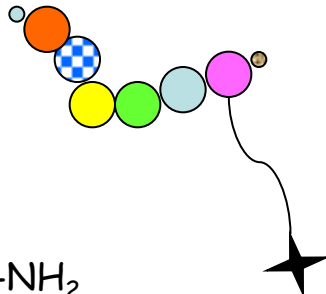


Ac-TRGRS-**Ttds**-**K**(biotinyl-hexanoyl)-NH<sub>2</sub>

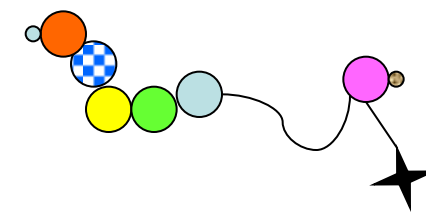
Ac-TRGRS**K**(biotinyl-Ttds)-NH<sub>2</sub>



Ac-TXGRS**K**(biotinyl-hexanoyl)-NH<sub>2</sub>



Ac-TXGRS**K**(biotinyl-Ttds)-NH<sub>2</sub>



Ac-TXGRS-**Ttds**-**K**(biotinyl-hexanoyl)-NH<sub>2</sub>

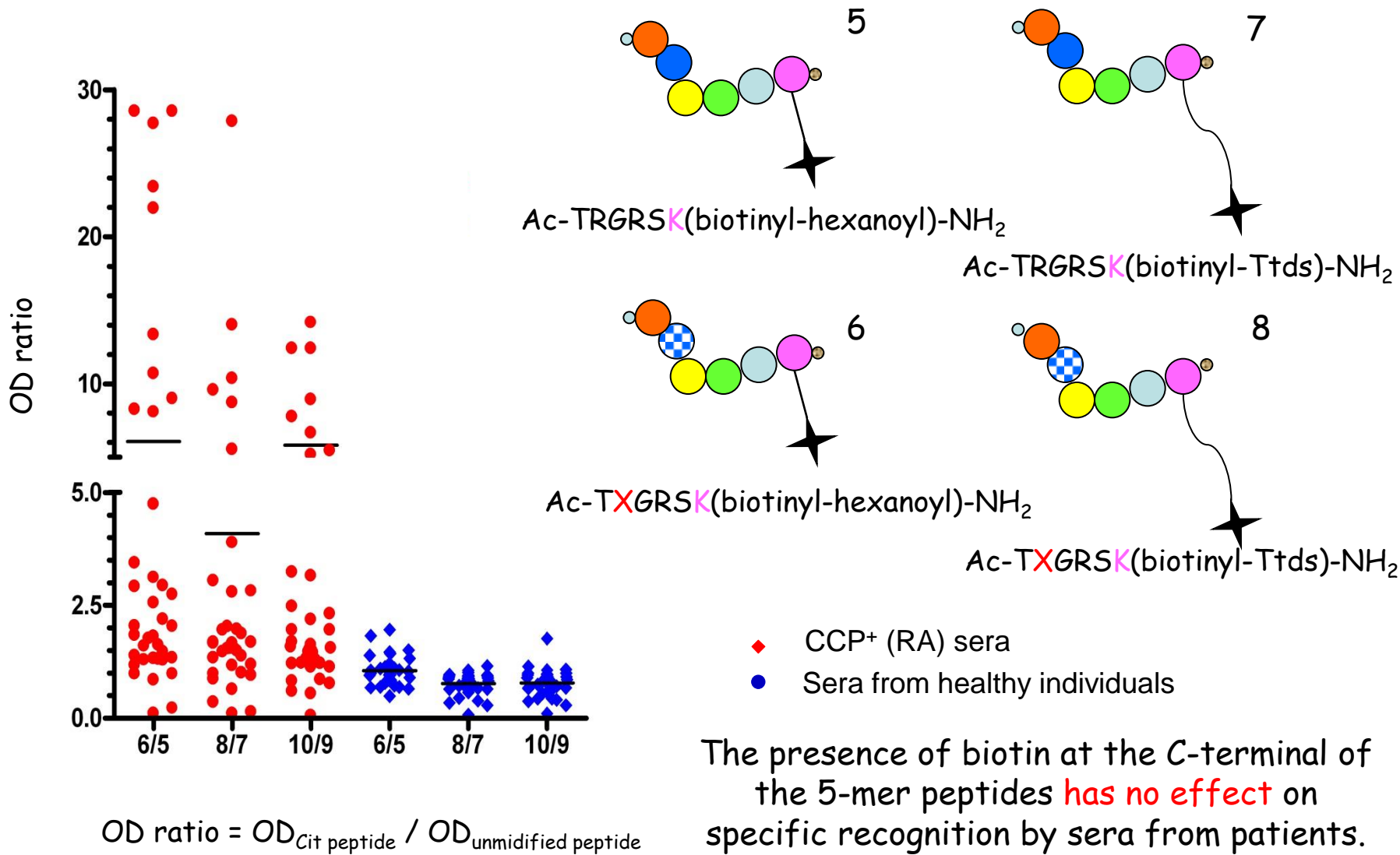
# Characteristics of 5-mer epitope peptide with C-terminal biotin

Compound	$M_{av}$ (calc)	$M_{av}$ (meas)	$R_t$ (min)
Ac-TRGRSK(biotinyl-hexanoyl)-NH <sub>2</sub>	1084,3	1084,6	17,08
Ac-TXGRSK(biotinyl-hexanoyl)-NH <sub>2</sub>	1085,3	1084,6	17,05
Ac-TRGRSK(biotinyl-Ttds)-NH <sub>2</sub>	1273,1	1273,2	17,97
Ac-TXGRSK(biotinyl-Ttds)-NH <sub>2</sub>	1274,1	1273,9	17,62
Ac-TRGRS-Ttds-K(biotinyl-hexanoyl)-NH <sub>2</sub>	1386,3	1386,4	18,16
Ac-TXGRS-Ttds-K(biotinyl-hexanoyl)-NH <sub>2</sub>	1387,3	1387,4	18,42

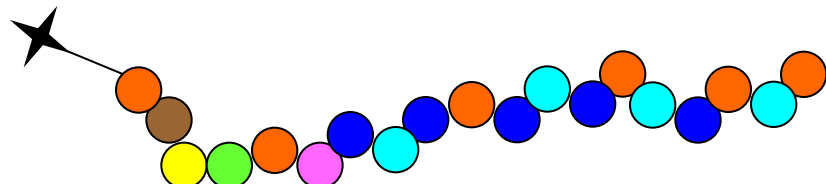
HPLC: KNAUER, Synergi MAX-RP, C12, 250 x 4mm, 5µm silica, 100 Å column, 5% B - 95 % B, 50 min,  
eluent A: 0,1% TFA/water (V/V); eluent B: 0,1% TFA/acetonitrile-water (80:20 V/V)

MS: Esquire 3000+

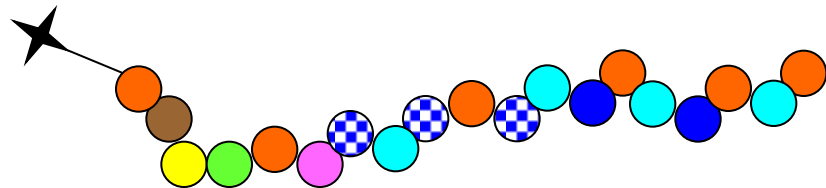
# Antibody recognition of 5-mer epitope peptides with C-terminal biotin



# Synthesis of 19-mer epitope peptide $(^{306}\text{SHQESTRGRSRGRSGRSGS}^{324})$ with N-terminal biotin



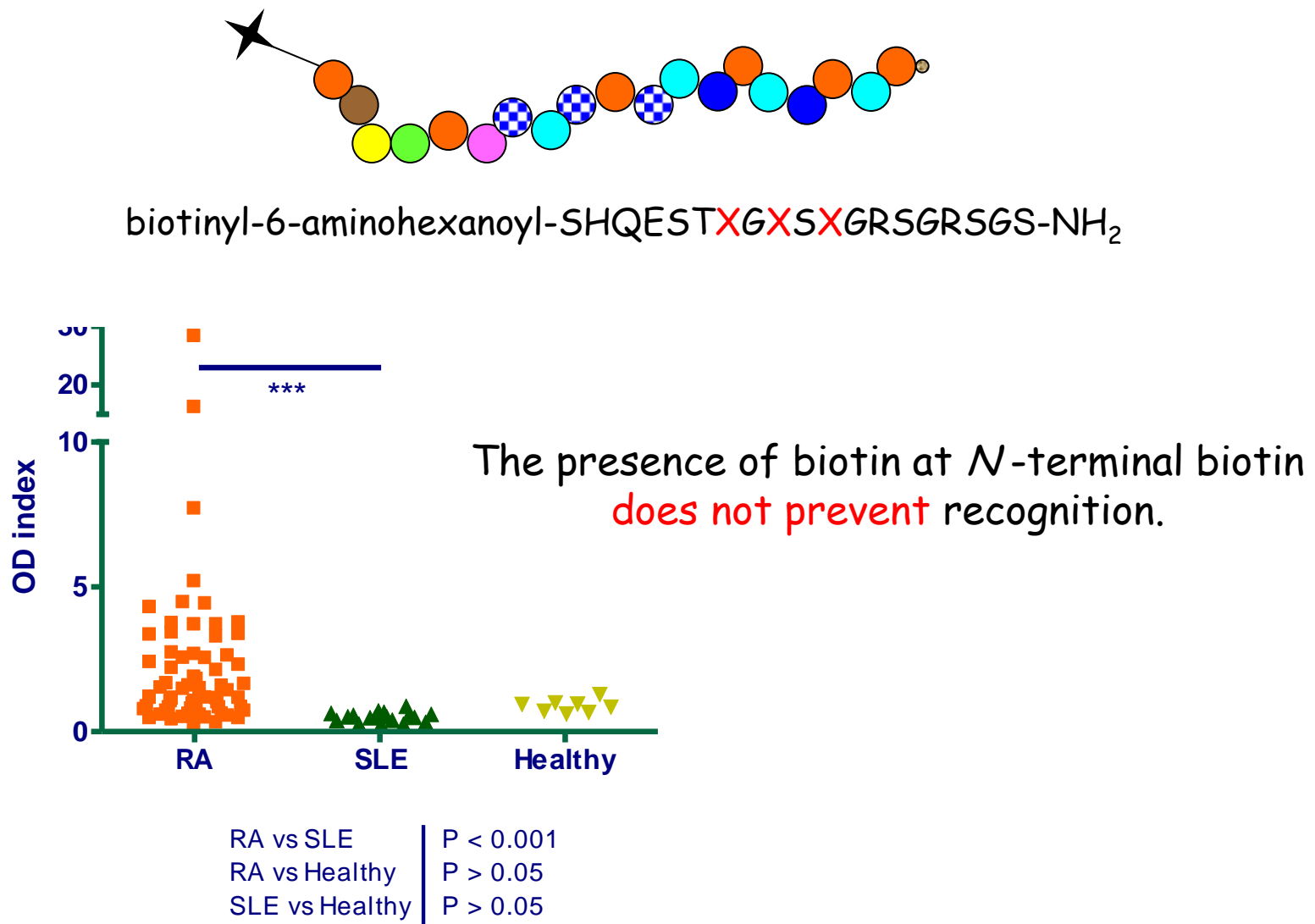
biotinyl-6-aminohexanoyl-SHQESTRGRSRGRSGRSGS-NH<sub>2</sub>



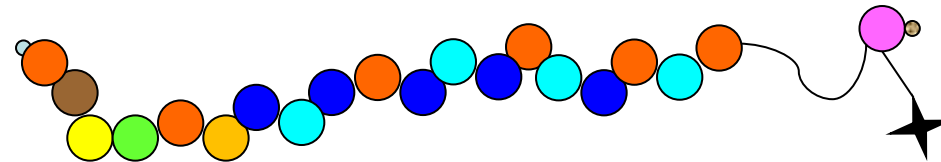
biotinyl-6-aminohexanoyl-SHQESTXGXSXGRSGRSGS-NH<sub>2</sub>

Compounds	M <sub>av</sub> (calc)	M <sub>av</sub> (meas)	R <sub>t</sub> (min)
biotinyl-6-aminohexanoyl-SHQESTRGRSRGRSGRSGS-NH <sub>2</sub>	2383,6	2383,8	13,27
biotinyl-6-aminohexanoyl-SHQESTXGXSXGRSGRSGS-NH <sub>2</sub>	2386,6	2386,7	12,95

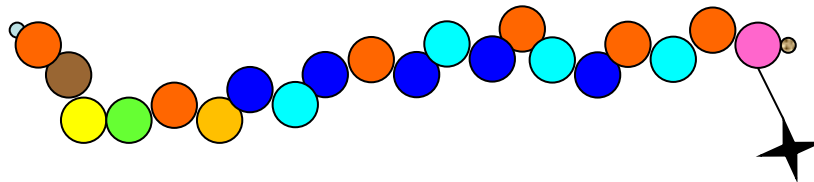
# Antibody recognition of 19-mer epitope peptide with *N*- terminal biotin by RA, SLE and healthy samples



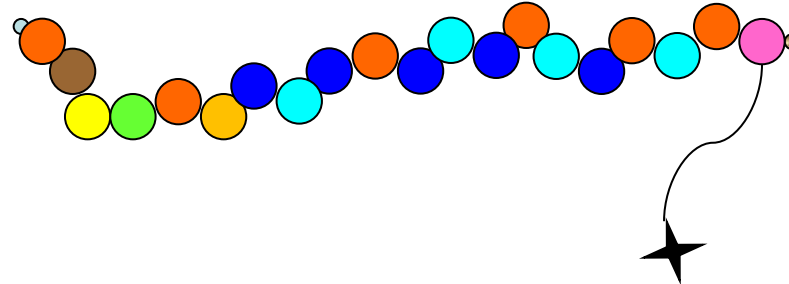
# Synthesis of 19-mer epitope peptide $(^{306}\text{SHQESTRGRSRGRSGRSGS}^{324})$ with C-terminal biotin



Ac-SHQESTRGRSRGRSGRSGS-Ttds-K(biotinyl-6-aminohexanoyl)-NH<sub>2</sub>



Ac-SHQESTRGRSRGRSGRSGSK(biotinyl-6-aminohexanoyl)-NH<sub>2</sub>



Ac-SHQESTRGRSRGRSGRSGSK(biotinyl-Ttds)-NH<sub>2</sub>

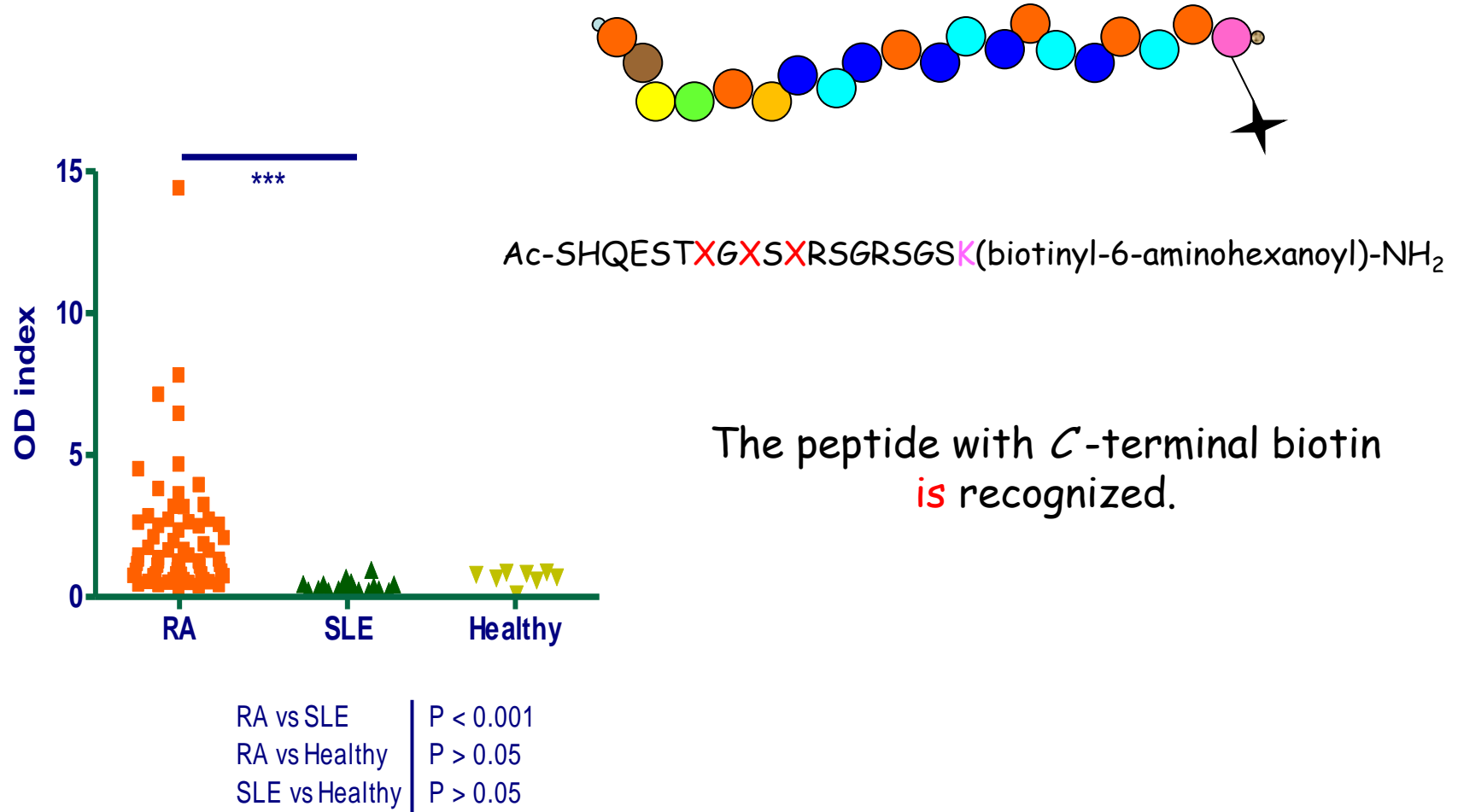
# Characteristics of 19-mer epitope peptide ( $^{306}$ SHQESTRGRSRGRSGS $^{324}$ ) with C-terminal biotin

Compound	$M_{av}$ (calc)	$M_{av}$ (meas)	$R_t$ (min)
Ac-SHQEST <b>RGRSRGRSGS</b> K(biotinyl-aminohexanoyl)-NH <sub>2</sub>	2553,8	2553,8	14,23
Ac-SHQEST <b>XGXSXGRSGRSGSK</b> (biotinyl-aminohexanoyl)-NH <sub>2</sub>	2556,8	2556,9	13,73
Ac-SHQEST <b>RGRSRGRSGSGSK</b> (biotinyl-Ttds)-NH <sub>2</sub>	2743,3	2743,4	15,25
Ac-SHQEST <b>XGXSXGRSGRSGSK</b> (biotinyl-Ttds)-NH <sub>2</sub>	2746,3	2746,5	14,90
Ac-SHQEST <b>RGRSRGRSGS-Ttds-K</b> (biotinyl-aminohexanoyl)-NH <sub>2</sub>	2856,4	2856,5	17,65
Ac-SHQEST <b>XGXSXGRSGRSGS-Ttds-K</b> (biotinyl-aminohexanoyl)-NH <sub>2</sub>	2859,4	2859,5	17,35

HPLC: KNAUER, Synergi MAX-RP, C12, 250 x 4mm, 5µm silica, 100 Å column, 5% B - 95 % B, 50 min,  
eluent A: 0,1% TFA/water (V/V); eluent B: 0,1% TFA/acetonitrile-water (80:20 V/V)

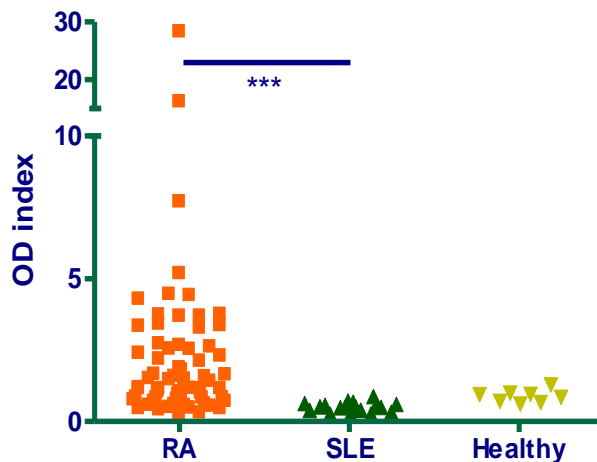
MS: Esquire 3000+

# Antibody recognition of 19-mer epitope peptide with *C*- terminal biotin by RA, SLE and healthy samples

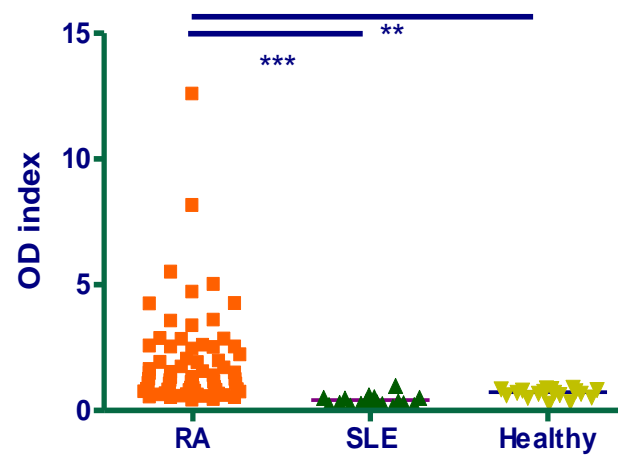


# Antibody recognition of 19-mer epitope peptide with *N*- or *C*- terminal biotin

biotinyl-6-aminohexanoyl-  
-SHQEST**XGXSXGRSGRSGS-NH<sub>2</sub>**

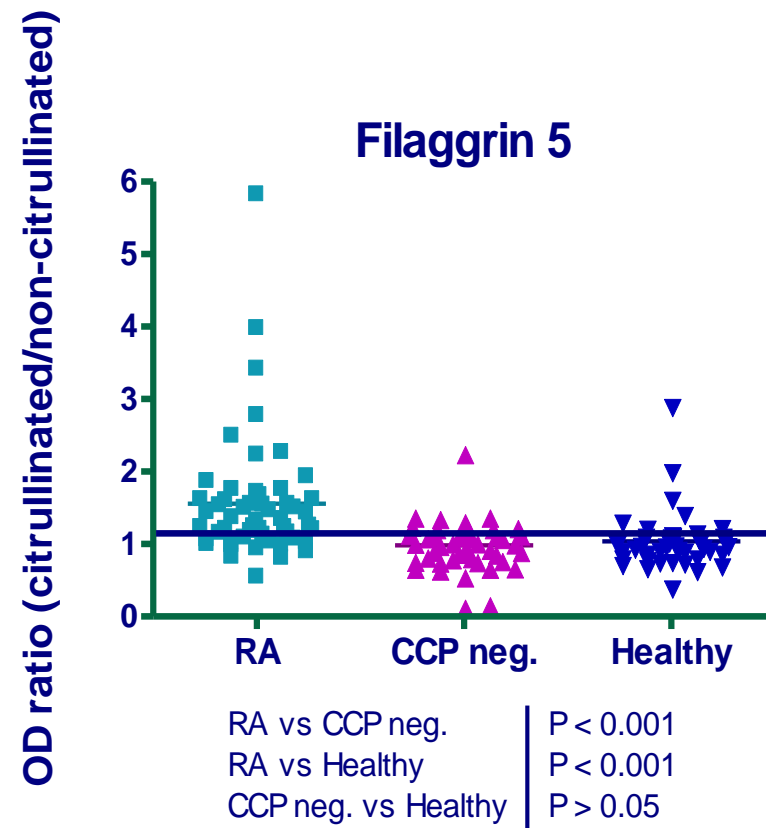
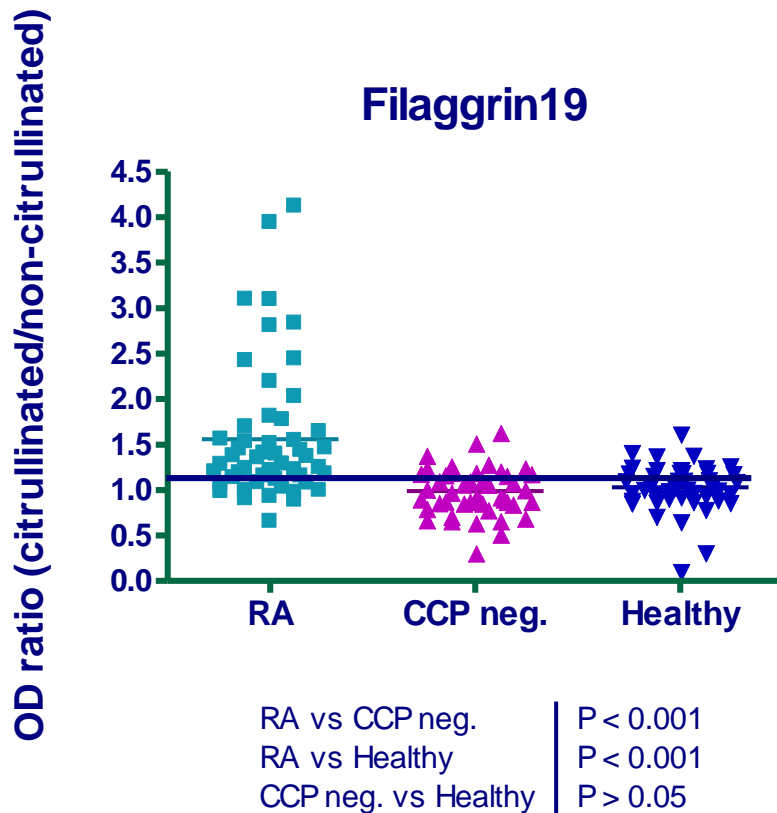


Ac-SHQEST**XGXSXGRSGRSGSK**  
(biotinyl-aminohexanoyl)-NH<sub>2</sub>



Both *C*- and *N*-terminal biotinylated 19-mer epitope peptides are recognized by RA sera samples

# Comparison of antibody recognition of the 5-mer and the 19-mer epitope peptide

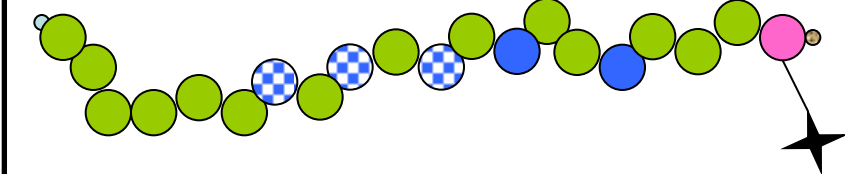
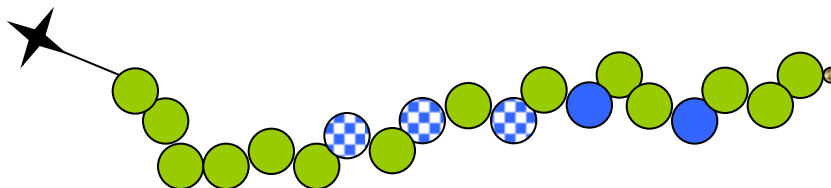
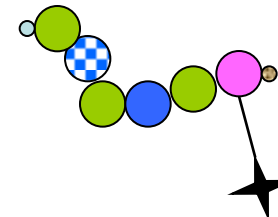


# Short summary

N-terminal biotinylation



C-terminal biotinylation



# ECD (Electronic Circular Dichroism)

Instrument: Jasco-810

Solvent:

- water
- TFE

Concentration: 0,5 mg/ml

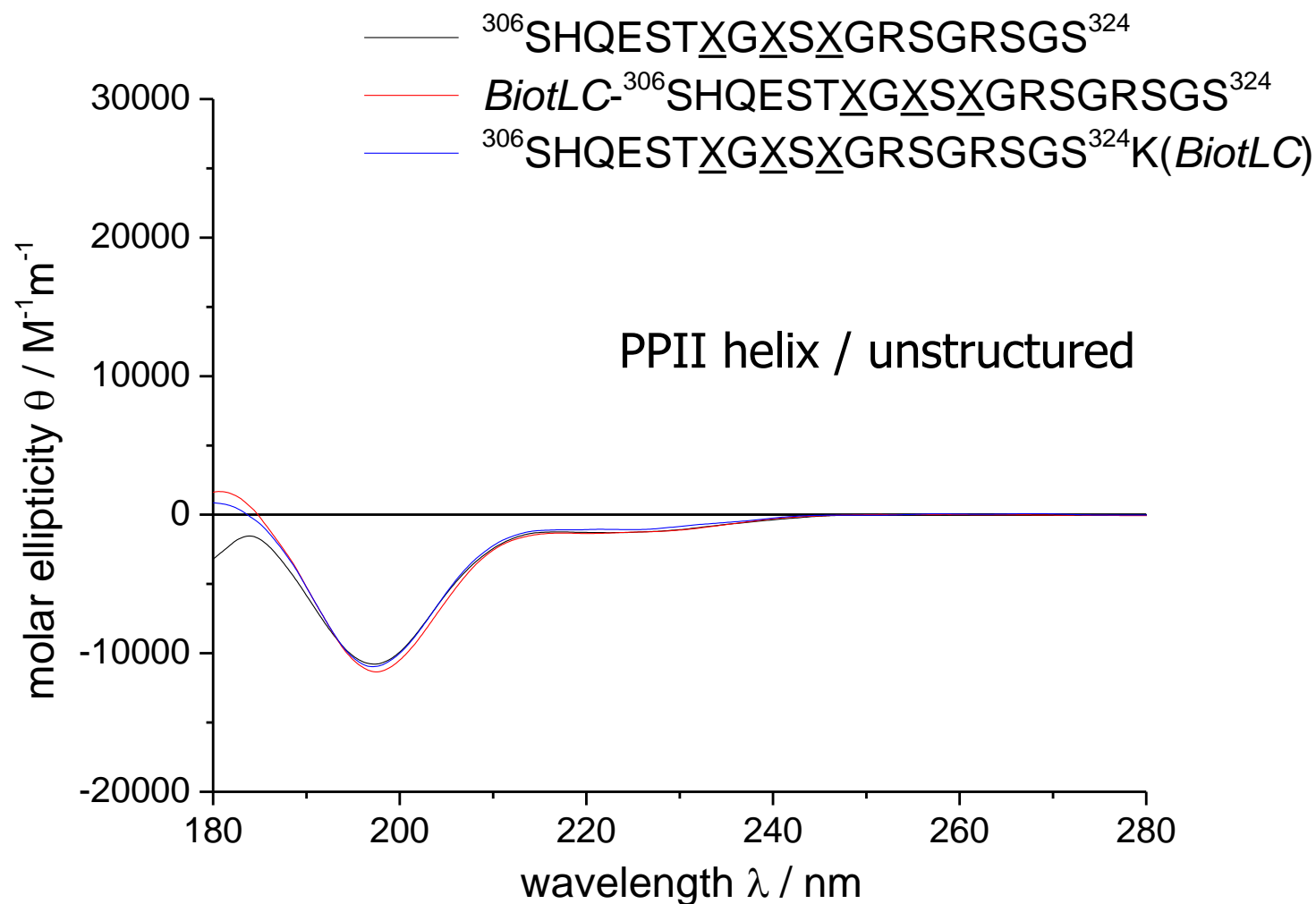
Wavelength:  $\lambda=180-300$  nm

0,02 cm quartz cuvette

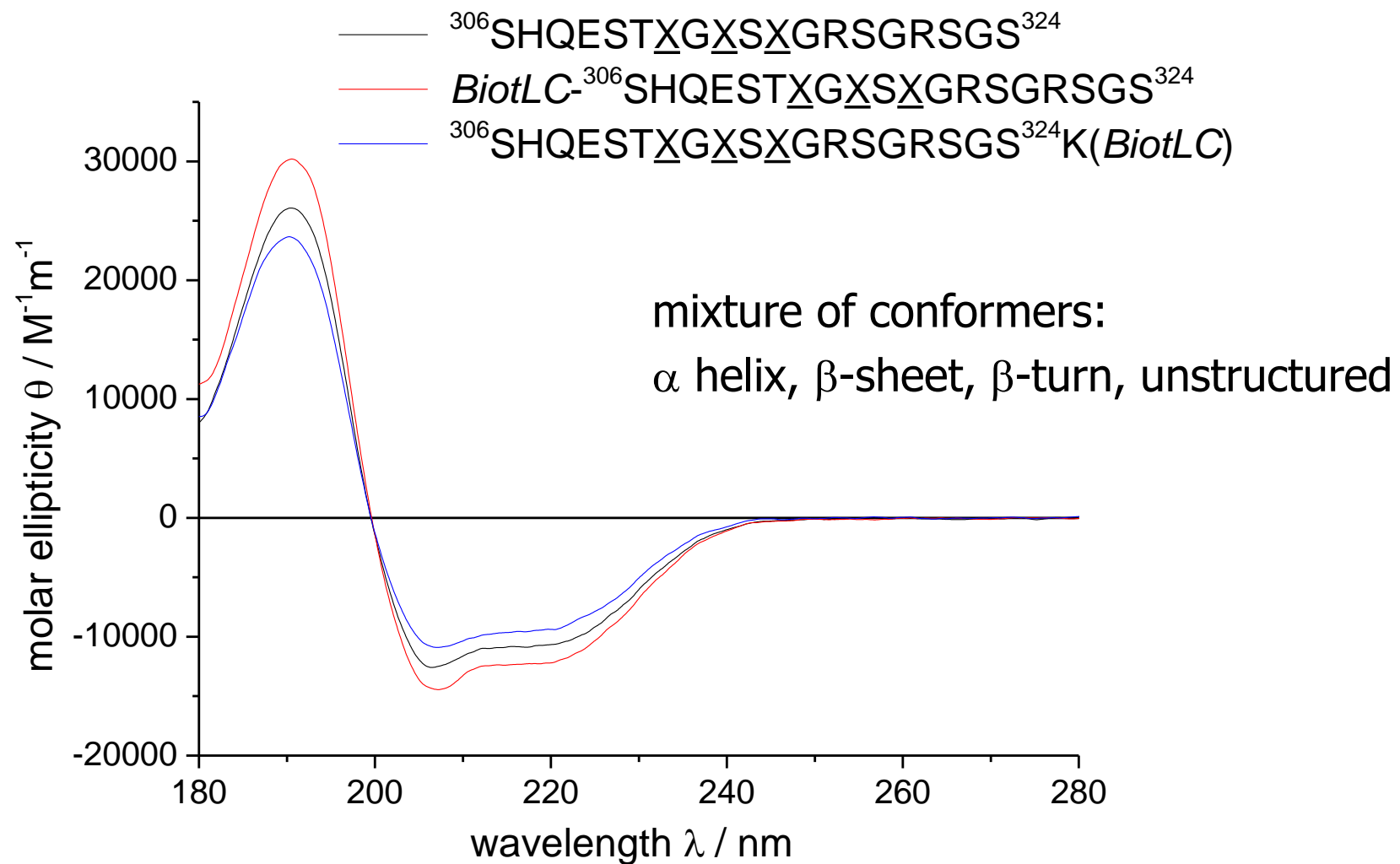


[http://www.andrew.cmu.edu/user/jamess3/  
JWSfac.html](http://www.andrew.cmu.edu/user/jamess3/JWSfac.html)

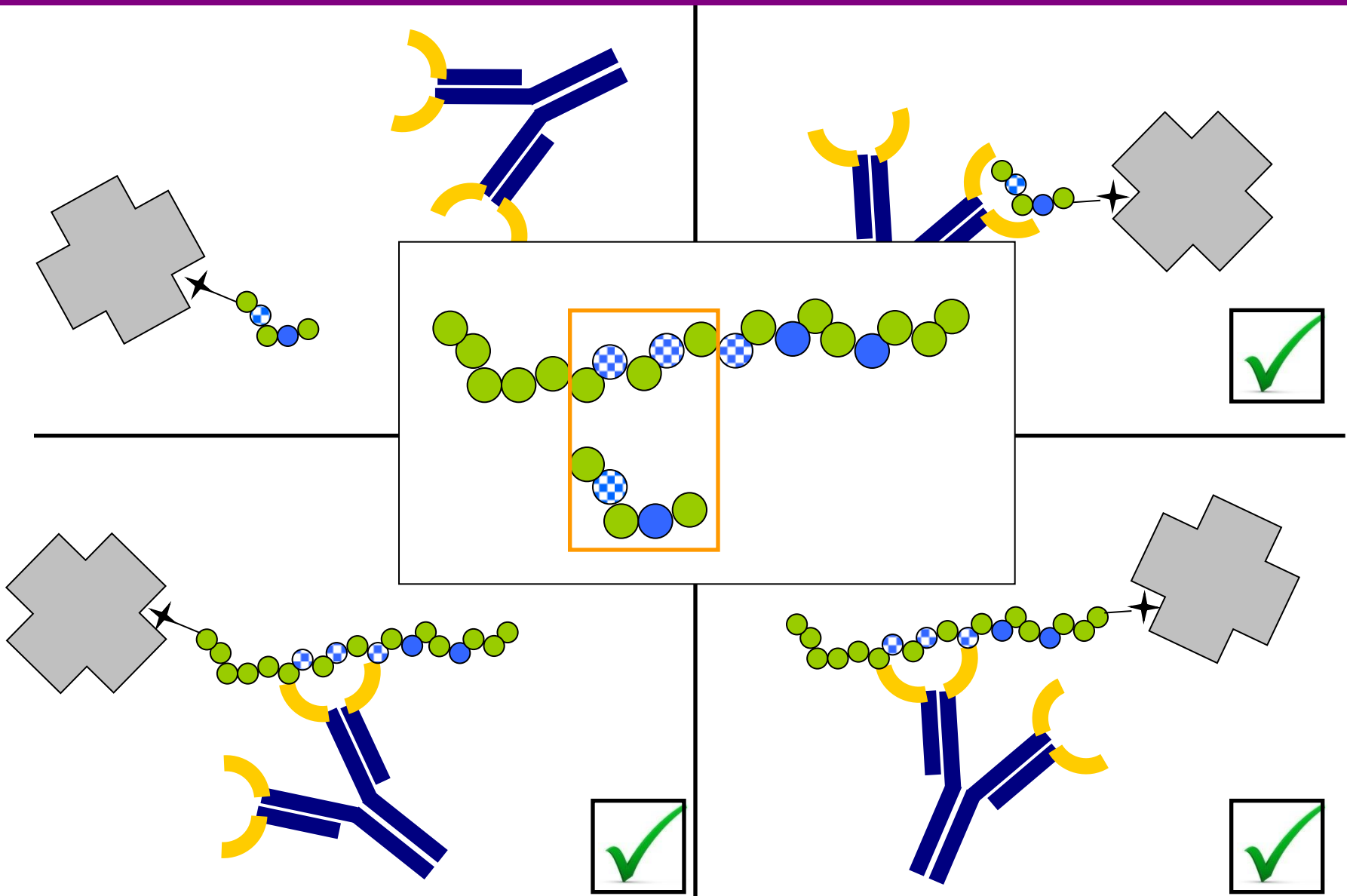
# The effect of biotin position on ECD spectra of 19-mer in water



# The effect of biotin position on ECD spectra of 19-mer in TFE



The position of the epitope core within the epitope region influence the antibody recognition



## Conclusions

---

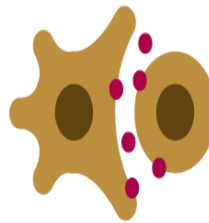
1. An epitope region (19-mer) and an epitope core (5-mer) were identified.
2. Introduction of biotin to the *N*-terminal of the 5-mer resulted in **no binding**. The presence of biotin at the *C*-terminal of the 5-mer had no effect on binding.
3. However, the **presence of biotin at *N*- or *C*-terminal of the 19-mer has no effect on epitope recognition** by serum antibodies.
4. The position of biotin markedly influences the solution conformation of the **epitope peptide (5-mer)**. In contrast, no influence of the biotin position could be detected in case of 19-mer epitope region peptide.
5. The 5-mer as well as the 19-mer citrullinated peptides have shown a significantly higher reactivity with  $CCP^+$  RA sera as compared to healthy controls,  $CCP^-$  serum samples.

# Acknowledgements

F. Babos<sup>1</sup>, E. Szarka<sup>2</sup>, A. Magyar<sup>1</sup>, G. Sármay<sup>2</sup>

<sup>1</sup> Research Group of Peptide Chem., Hungarian Academy of Sciences, Eötvös L. University,

<sup>2</sup> Department of Immunology, Eötvös L. University, Budapest



**CellKom**

Regionális Egyetemi Tudásközpont



Pázmány

[O] Pázmány Péter program

A projekt a Nemzeti Kutatási és Technológiai Hivatal támogatásával valósult meg.



## Support

Hungarian-French Intergovernmental Program (F-9/2010)

Hungarian Academy of Sciences

Hungarian National Research Fund (OTKA T045634)

Ministry of Health (ETT)