



# Analytical Applications of Aptamers

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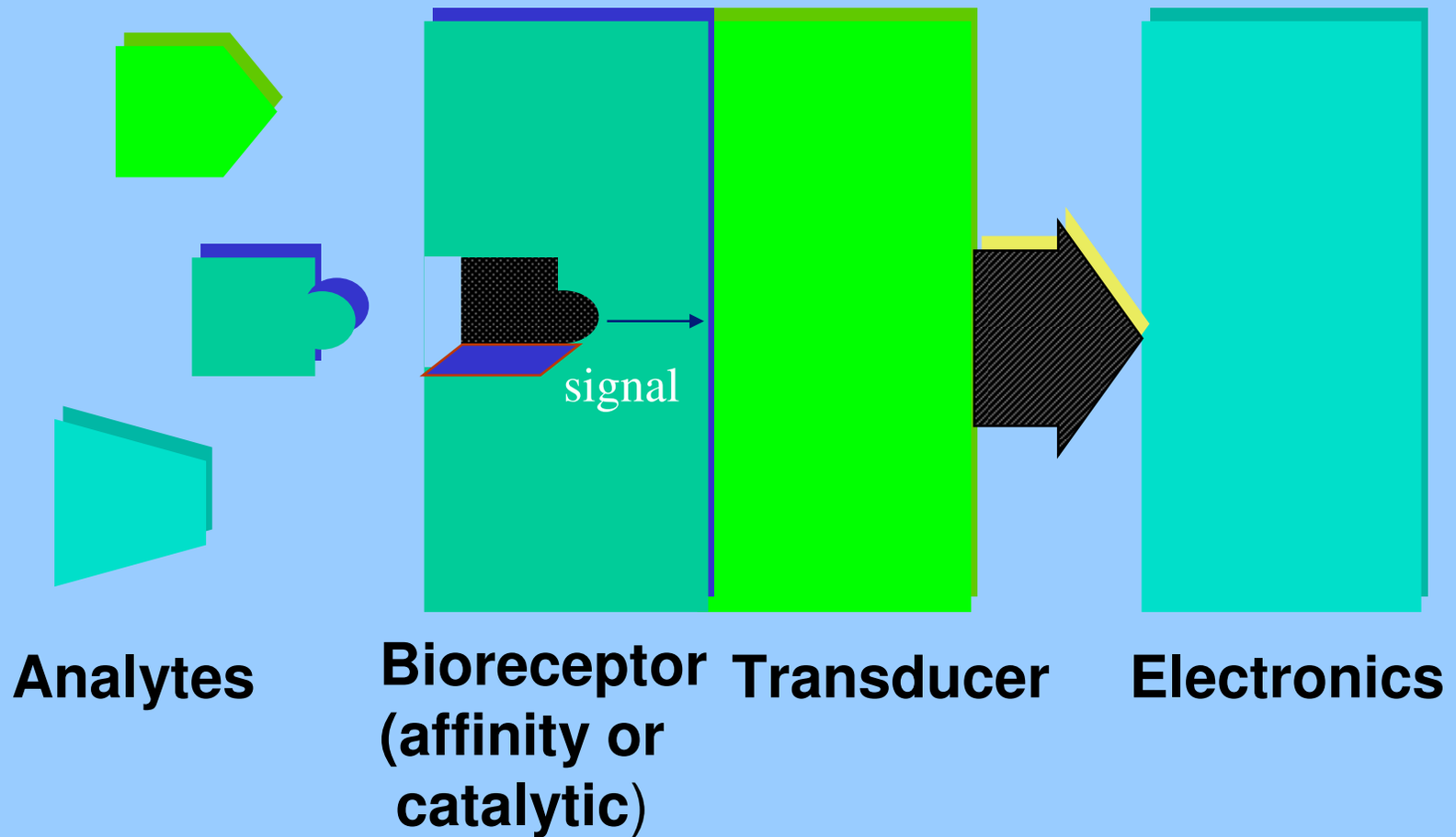
**[www.unifi.it/dclabi](http://www.unifi.it/dclabi)**

**“*Biosensors* are analytical devices incorporating a biological material (e.g. tissue, microorganisms, organelles, cell receptors, enzymes, antibodies, nucleic acids etc.), a biologically derived material **or biomimic** intimately associated with or integrated within a physicochemical transducer **or transducing microsystem**, which may be optical, electrochemical, thermometric, piezoelectric, or **magnetic.**”**

***Biosensors* usually yield a digital electronic signal which is proportional to the concentration of a specific analyte or group of analytes. While the signal may in principle be continuous, **devices can be configured to yield single measurements to meet specific market requirements.**”**  
**(One-shot biosensors)**

***Biosensors & Bioelectronics (2000)***

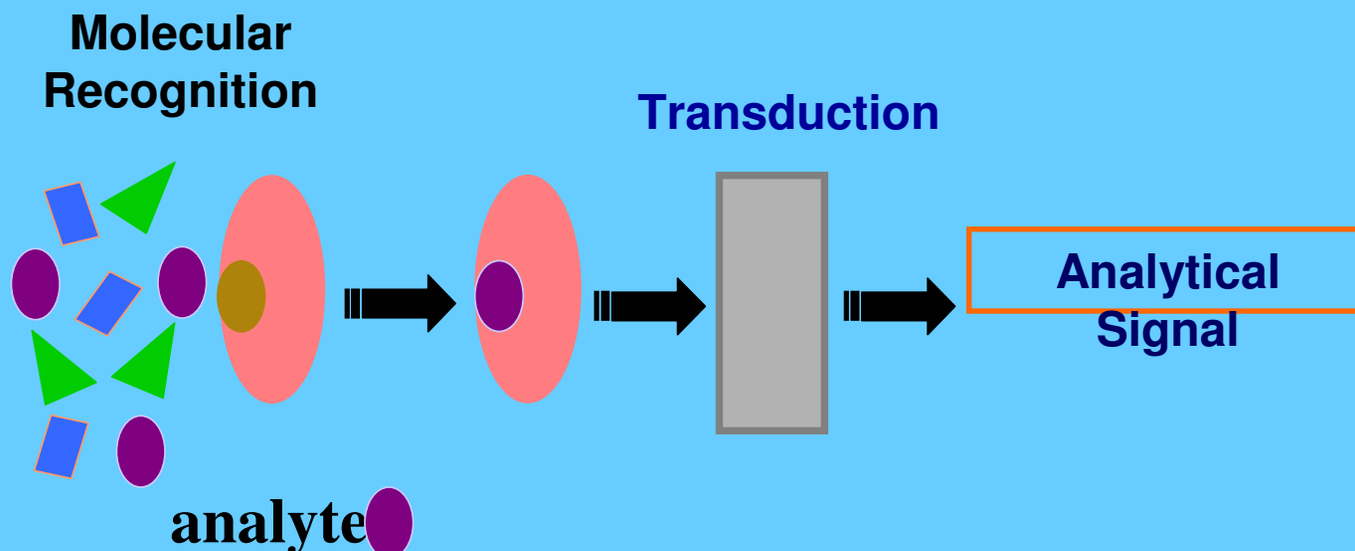
# *The Biosensor*



# DNA BIOSENSORS

- **Trace measurements of pollutants  
(intercalators, binders of DNA)**
- **Hybridization indicator  
(bacteria , virus , genetic inherited diseases)**
- **Biosensing of drugs**

# AFFINITY BIOSENSOR



## MOLECULAR RECOGNITION ELEMENTS

### BIOLOGICAL

- Antibodies
- Receptors
- Nucleic Acids

### BIOMIMETIC

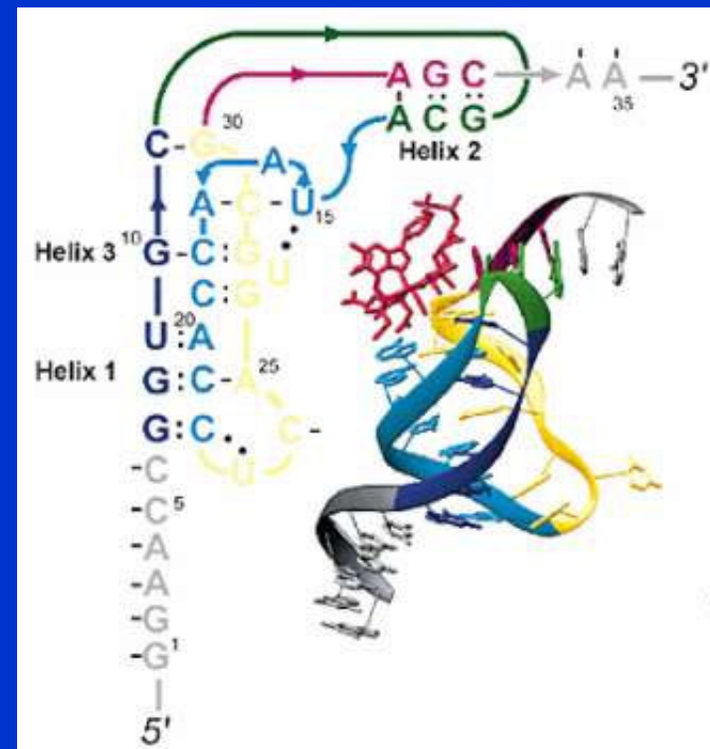
- Molecular Imprinted Polymers (MIP)
- Oligonucleotides
- Oligopeptides
- **Aptamers**

# Aptamers

**Aptamers** are oligonucleotides (DNA or RNA molecules) that can bind with high affinity and specificity to a wide range of target molecules (proteins, peptides, drugs, vitamins and other organic or inorganic compounds).

They were “discovered” in 1990 by the development of an *in vitro* selection and amplification technique, known as **SELEX** (Systematic Evolution of Ligands by Exponential enrichment).

Their name is derived from the Latin word “aptus” which means “to fit” and the Greek word “meros” part or region !



# APTAMERS

Aptamers are oligonucleotides that are identified through a combinatorial selection process for high affinity binding to target molecules. In the selection process, a combinatorial library of oligonucleotides is passed through a column containing the immobilized target.

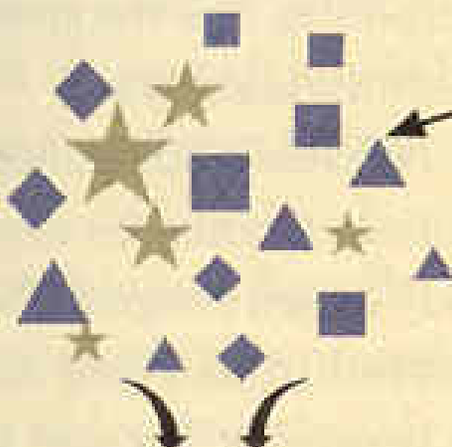
Those oligonucleotides that do not bind are discarded, while those that bind are collected and amplified.

This cycle is repeated several times until a small number of affinity binders, or aptamers, have been isolated from the combinatorial pool.

# APTAMERS

- Synthetic sequence (30-40 mer) of nucleic acids, single strand DNA or RNA, obtained by an *in vitro* selection (SELEX)
- Molecular recognition highly selective for the structures selected
- Possibility to obtain aptamers for a wide kind of structures
- Thermal stability and lifetime higher in comparison with protein receptors (antibodies)
- Any animal involved ; the procedure is suitable also for small molecule or any toxicants
- Time for obtaining : 2-3 months (v. monoclonal antibodies)

Oligonucleotide library



Amplification

Bound target



Elute retained oligomers

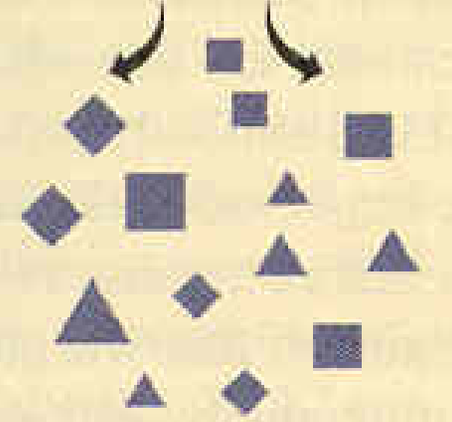


Analysis of binding sequences

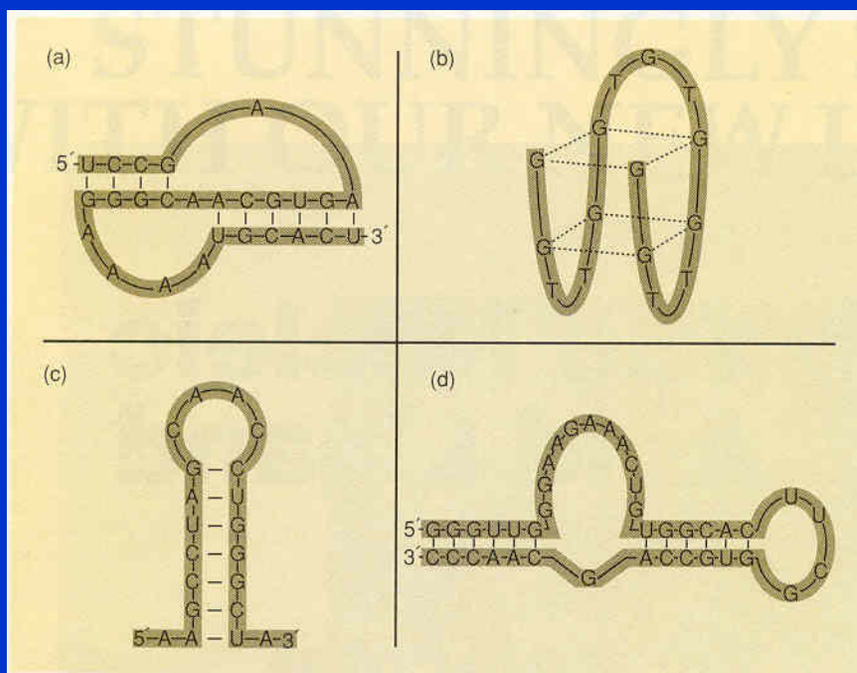
Nucleic acid ligand



Discard unretained oligomers



The majority of aptamer structures result from intramolecular base pairing to produce loops or bulges, forming structures such as the hairpin, the pseudo knot, and the stem-loop/bulge. A different type of structural motif is the G-quartet, also known as “quadruplex”, “tetraplex” or “G4” DNA.



A) Pseudoknot , b) G-Quartet, c) hairpin, d) stem-loop/bulge

From Anal. Chem. 1995, 664A

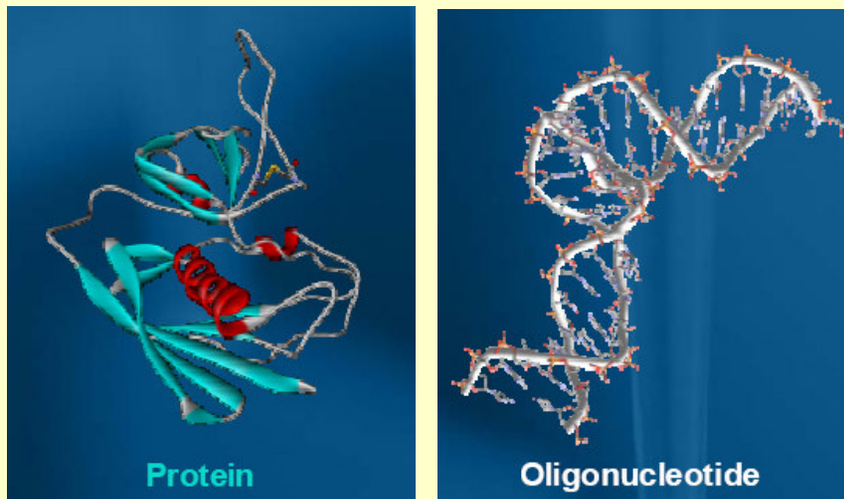
# Aptamers

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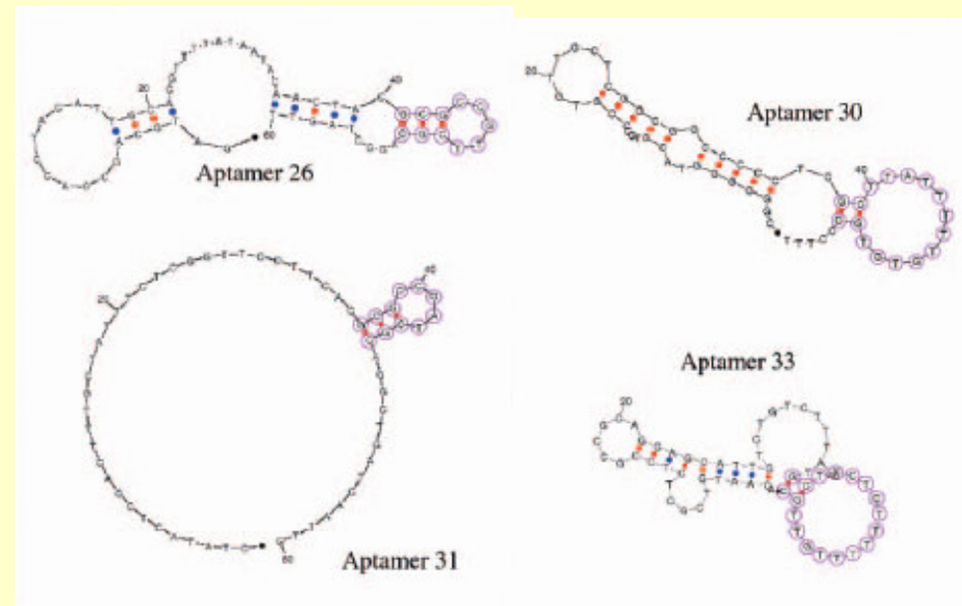
They were “discovered” in 1990 by the development of an in vitro selection and amplification technique, known as SELEX (Systematic Evolution of Ligands by Exponential enrichment).

(Ellington et al., **Nature** 346, 818; Tuerk and Gold, **Science** 249, 505)

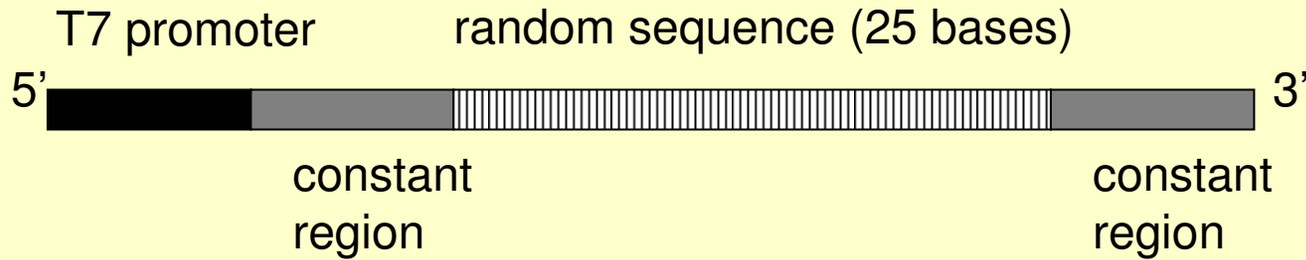
Their name is derived from the Latin word “**aptus**” which means “to fit”.



Similar to proteins short oligonucleotides can adopt complex three-dimensional structures



# Combinatorial oligonucleotide library



**A, G, C, U(T)**

$$4^1 = 4$$

$$4^2 = 16$$

$$4^3 = 64$$

$$4^4 = 256$$

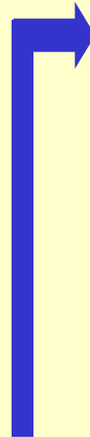
$$4^5 = 1024$$

.....

.....

.....

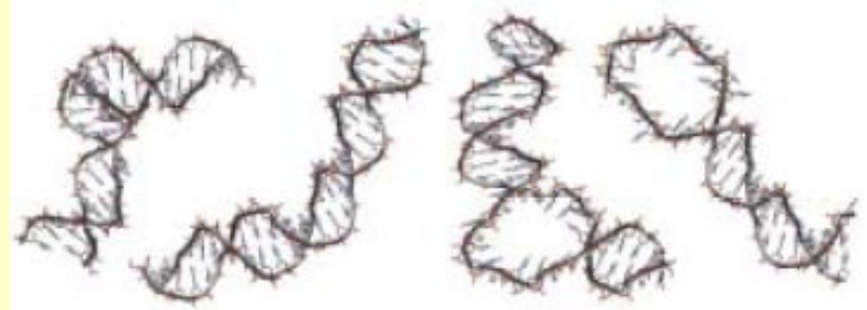
$$4^{25} = 1125899906842624$$

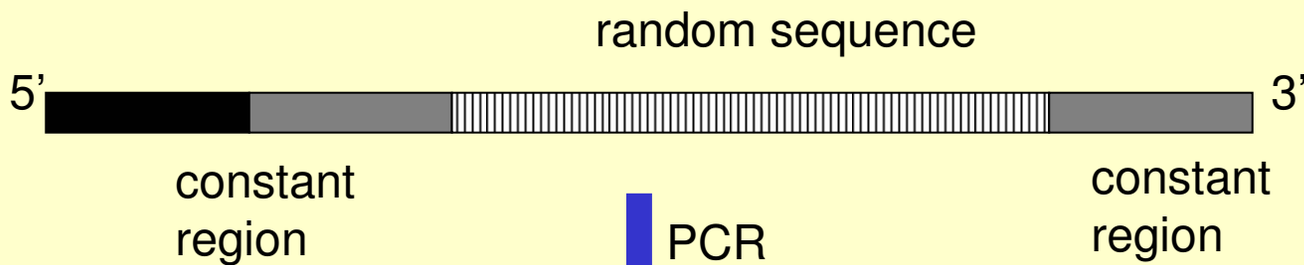


Pool of randomized DNA or RNA

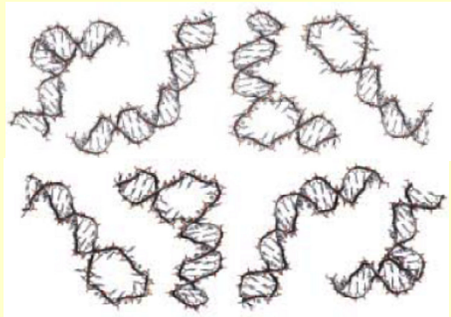
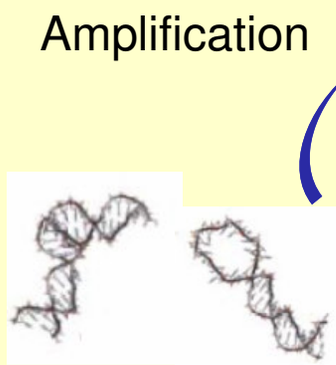


**10<sup>15</sup> different sequences!!!!**





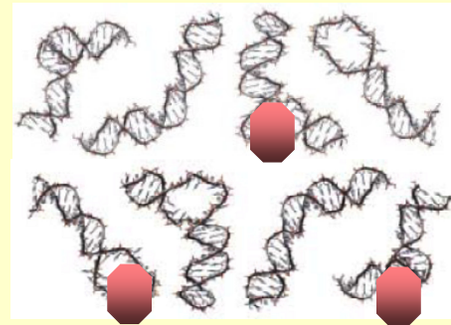
↓ PCR



DNA or RNA pool

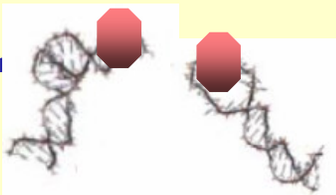


Target

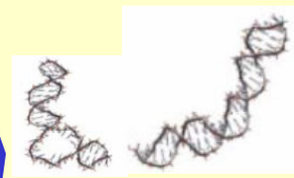


Incubation of target and RNA or DNA

Elution of RNA or DNA



Separation of binding from non-binding species



Patent Number	Year	Description
US5580737	1996	Counter SELEX
WO9833941	1998	Flow cell Selex
WO0056930	2000	Truncation SELEX
US5683867	1997	Blended Selex
US6387620	2002	Transcription Free SELEX
US5567588	1996	Solution Selex
WO9604403	1996	Chimeric Selex
US6376474	2002	Tissue SELEX
US6001577	1999	Photo SELEX
US7312325	2001	Toggle Selex
US5763595	1998	Covalent SELEX/ Chemi SELEX
US6261774	2001	Genomic SELEX
WO0224954	2002	SELEX without purified protein
WO03102212	2003	CE-SELEX
EP1386972	2007	Mirror-image SELEX- Spiegelmers
WO2007035532	2007	Laser SELEX, DeSELEX

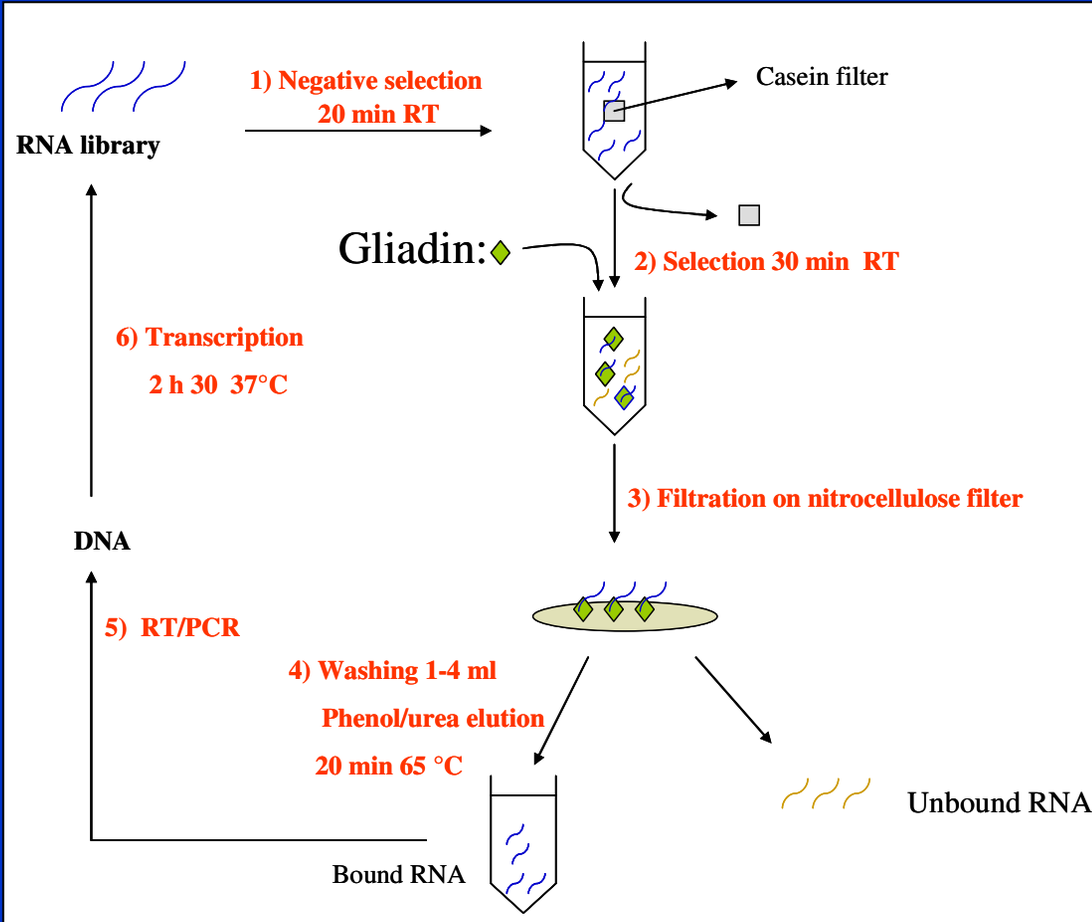
Tetranucleotides =  $4 \exp 4 = 264$

Tetrapeptides =  $20 \exp 4 = 160.000$

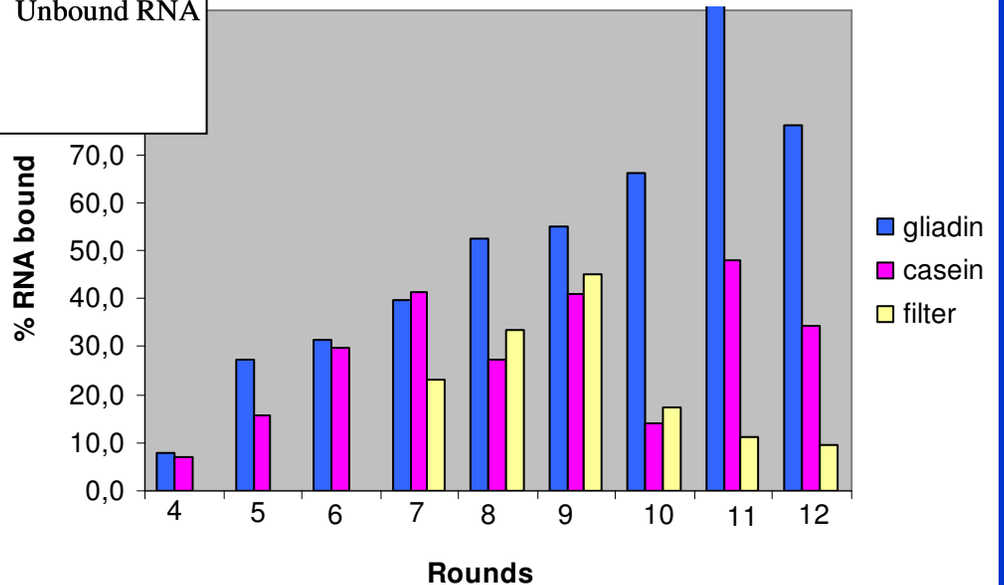
Tetrasaccharides = few millions of structures

# SELEX: an example

Target:  
 $\alpha$ -Gliadin (IRMM-480 from european wheat)



## SELEX evolution after 12 rounds



Sonia Centi

Institut Européen de Chimie et  
 Biologie, Bordeaux, France

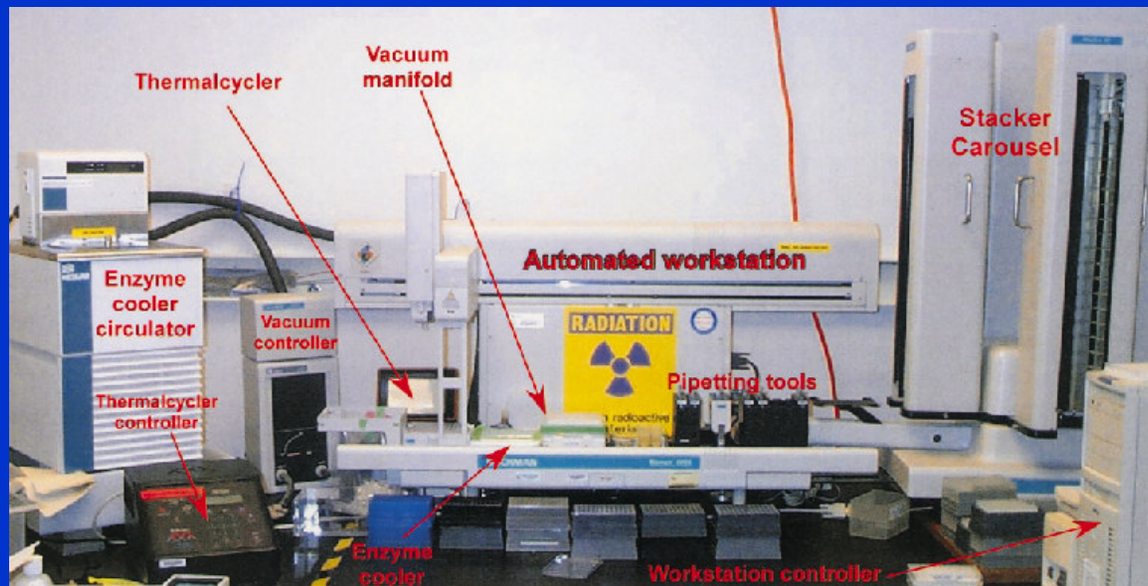
Prof. J.J. Toulmè

June-October 2004

# Automation and modification of the SELEX process

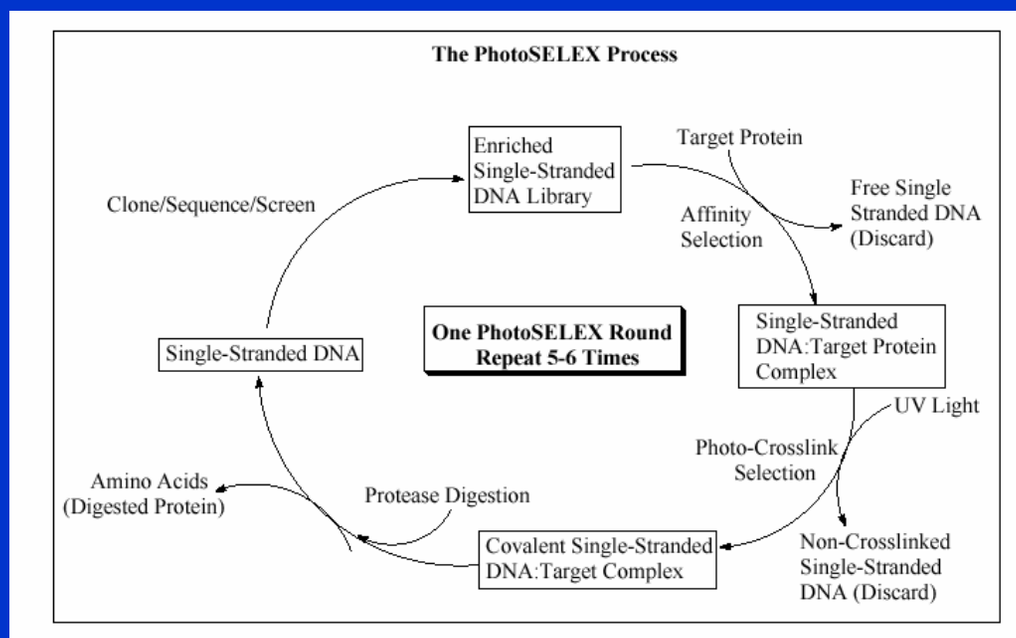
- Automated selection of aptamers

J.C. Cox, A.D. Ellington, *Bioorg. Med. Chem.* 9, 2525-2531, (2001)



- PhotoSELEX:** modified ssDNA aptamers capable of photocross-linking the target molecule.

M.C., Golden, B.D. Collins, M.C. Willis, T.H. Koch, *J. Biotechnol.* 81, 167-178, (2000)  
C. Bock et al., *Proteomics*, 4, 609-618, 2004



## Target molecules

### PROTEINS

Syrian golden hamster prion

Escherichia coli SelB

L-selectin

Tyrosine phosphatase

Ff gene 5

Thrombin

HIV-1 Tat

HIV-1 Rev

Vascular endothelial growth factor

Prostate specific antigen

Human IgE

Taq DNA polymerase

Iron regulatory protein

Human oncostatin M

Human neutrophil elastase

Human CD4 antigen

Lysozyme

C-reactive protein

Tumor necrosis factor  $\alpha$

NF- $\kappa$ B

Acetylcholine receptor

Thyroid transcription factor

### INORGANIC COMPOUNDS

Malachite green

Mg<sup>2+</sup>

### ORGANIC COMPOUNDS

ATP

FMN

Theophylline

Organic dyes

Cocaine

### VITAMINS

Cyanocobalamin

Biotin

### DRUGS

Neomycin B

Streptomycin

Tobramycin

Tetracyclin

Kanamycin A

Dopamine

### TOXINS

Cholera toxin

Staphylococcal enterotoxin B

### POLLUTANTS AND

### CARCINOGENIC COMPOUNDS

4-chloroaniline

2,4,6-trichloroaniline

Pentachlorophenol

Methylenedianiline

### OTHERS

*Bacillus anthracis* spores

# Applications

Applications based on molecular recognition:

Therapeutics: aptamers have been selected to disrupt the function of their targets and to inhibit or modify the metabolism associated with that target

Diagnostics: the impressive discrimination between two molecules of very similar structure has suggested that aptamers can be potential diagnostic reagents

Analytical tools:

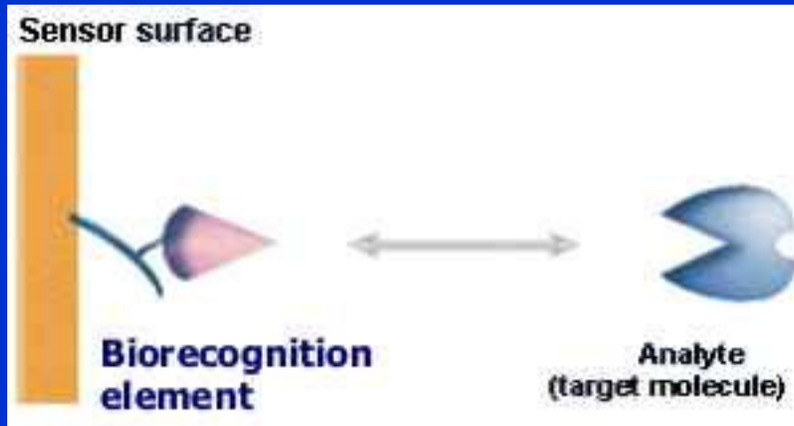
flow cytometry

capillary electrophoresis and electrochromatography

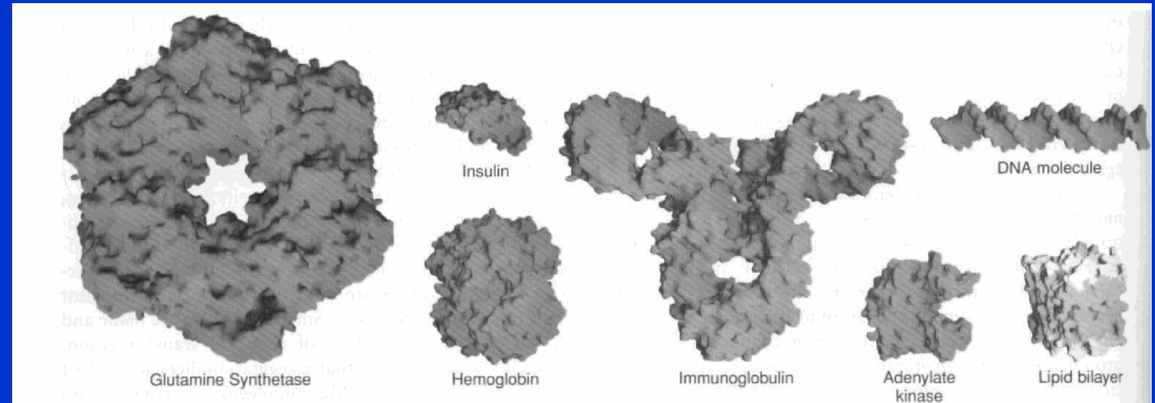
affinity chromatography

biocomponents in biosensors

# Affinity Biosensors



## Biomolecules (natural)

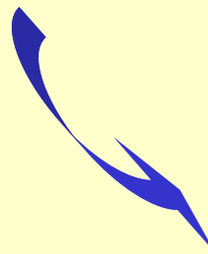


## Synthetic receptors:

- Peptide Nucleic Acids (PNAs)
- Molecularly Imprinted Polymers (MIPs)
- Oligopeptides
- Aptamers

## *Why aptamers can rival antibodies and other synthetic receptors?*

- Overcoming of the use of **animals** for their production
- After selection, aptamers are **produced by chemical synthesis** and purified to a very high degree by eliminating the batch-to-batch variation found when using antibodies
- By chemical synthesis, **modifications** in the aptamer can be introduced enhancing the stability, affinity and specificity of the molecules
- Higher **temperature stability**
- Because of their **small size**, denser receptor layers can be generated
  
- **Amplification** by PCR during their selection

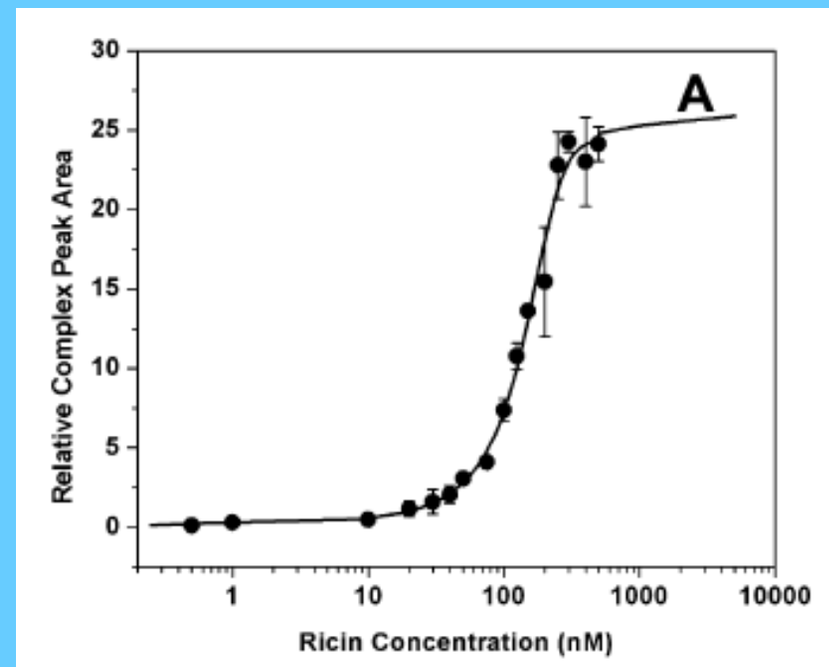
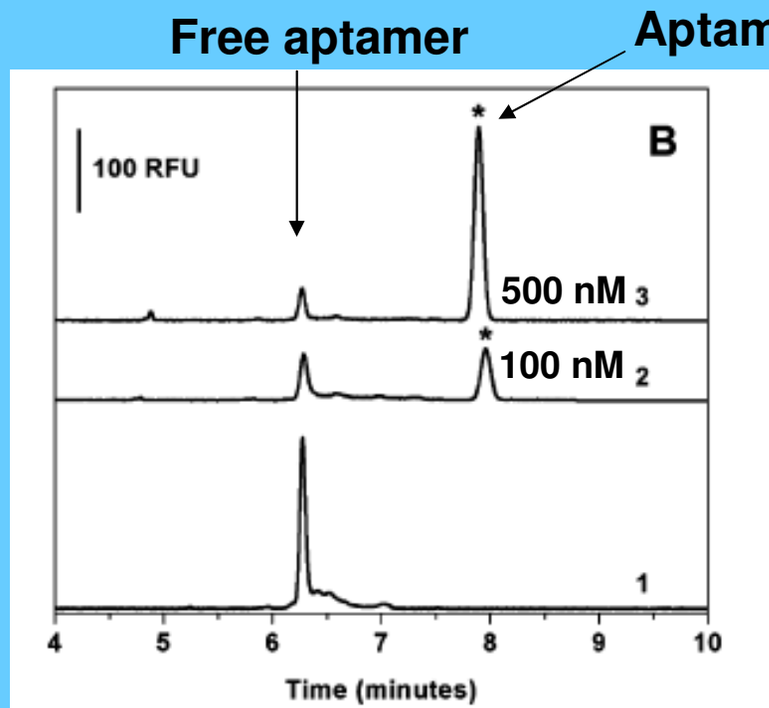


Advantage respect to other synthetic receptors such as **oligopeptides** which have a higher number of possible structures due to the higher number of “building blocks” (21 aminoacids), but they can not be amplified during the “production”

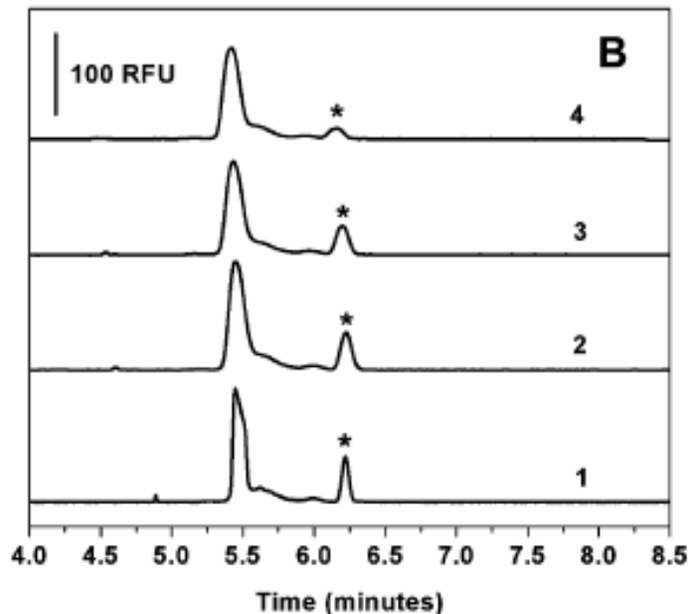
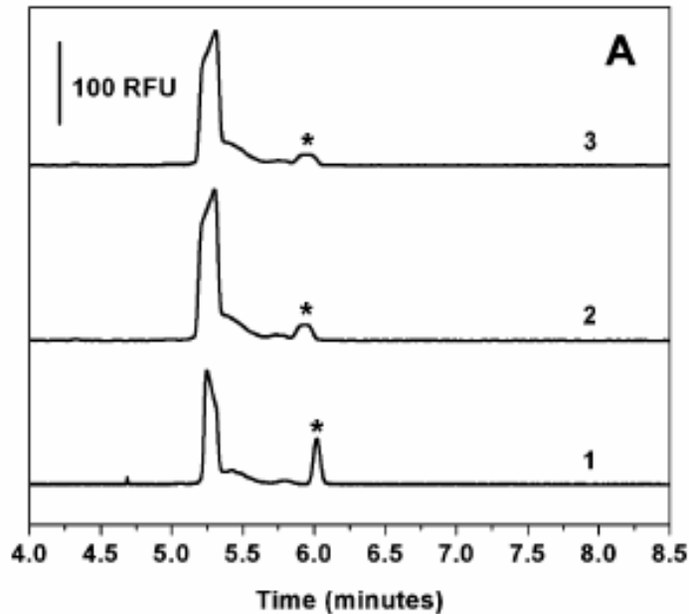
## Capillary electrophoresis

The integration of laser-induced fluorescence-based capillary electrophoresis with fluorescently labelled aptamers provides a novel approach for the detection of ricin. This free solution assay offers an alternative technique for protein detection in comparison to standard immunoassay and ELISA methods.

- Target molecule: Ricin
- RNA aptamer
- Transducer: affinity probe-based capillary electrophoresis



# Capillary electrophoresis



- Detection limit 500 pM
- Dynamic range low nM- low  $\mu$ M
- $K_d$   $K_d=134$  nM

## Detection of ricin in protein mixtures:

- A) (1) 50 nM ricin  
(2) 50 nM ricin and 50  $\mu$ g/mL BSA  
(3) 50 nM ricin and 100  $\mu$ g/mL BSA

- B) (1) 50 nM ricin  
(2) 50 nM ricin and 50  $\mu$ g/mL casein  
(3) 50 nM ricin and 100  $\mu$ g/mL casein  
(4) 50 nM ricin and 150  $\mu$ g/mL casein

# *Transducers*

- Acoustic sensors
- Cantilever-based biosensors
- Optical sensors
- Electrochemical sensors

# *Critical aspects to be considered when developing an aptamer-based biosensor*

## APTAMER IMMOBILIZATION

The procedure to fix the aptamer to the biosensor/bioanalytical device surface is of paramount importance to obtain an ordered layer able to exploit as much as possible the flexibility of the bioreceptor without altering its structure and its affinity for the target molecule.

The immobilization of the aptamer on a solid support must avoid any steric hindrance or constraint which could prevent the folding of the aptamer in the correct conformation

## BINDING PROTOCOL

From the examination of the different protocols employed in aptamer-based assays, one important point must be emphasized and that is the nature, conformation and sequence of each aptamer should be carefully considered and also stress that optimal working conditions can remarkably vary from one aptamer to another

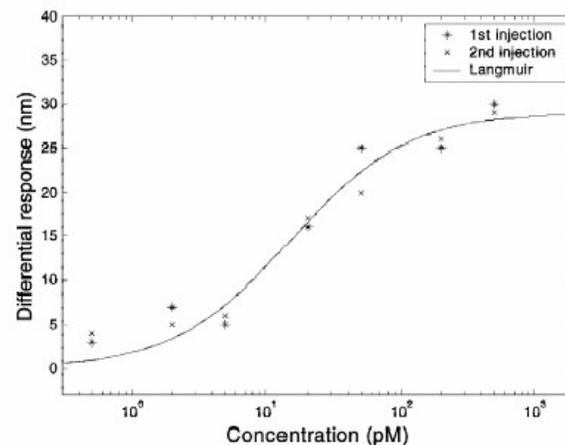
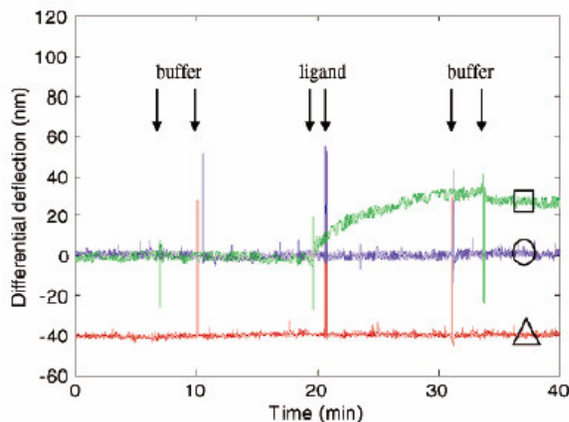
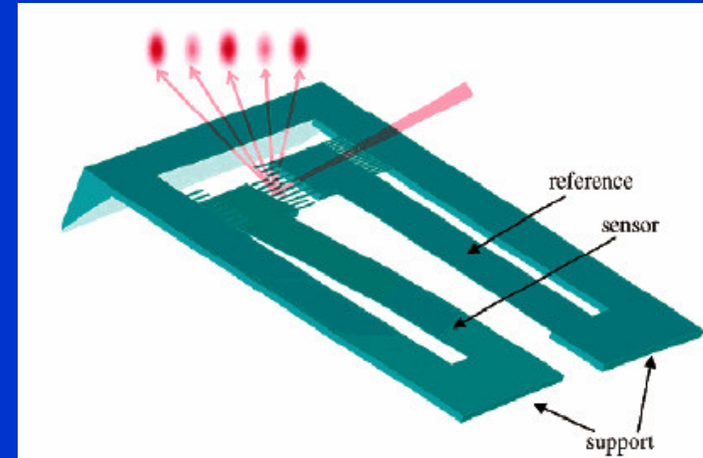
# Cantilever-based biosensor

Cantilever-based biosensing:

Label-free detection  
Batch-fabricated  
Small scale

Arrays can be used in parallel to detect various proteins simultaneously

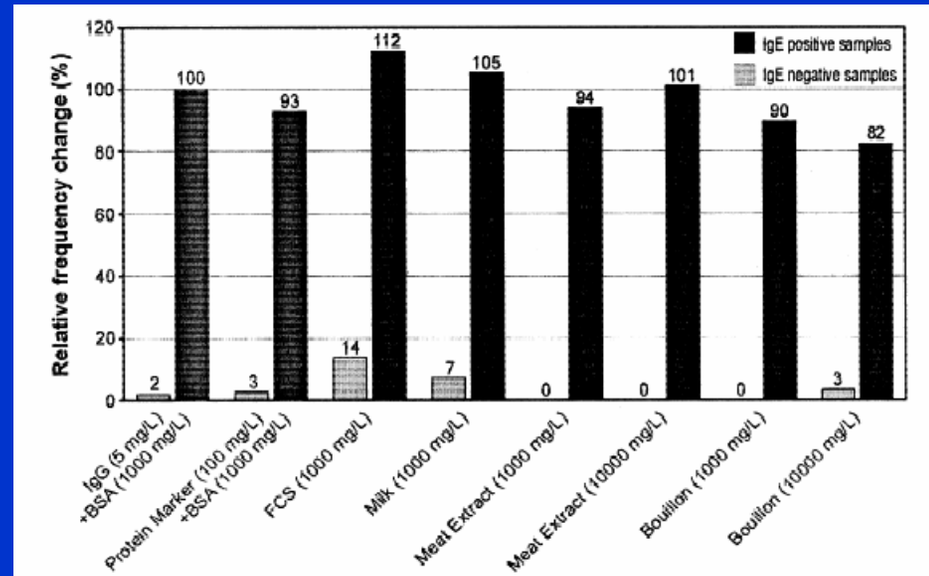
- Target molecule: Taq DNA polymerase
- DNA aptamer
- Transducer: cantilever
- Immobilisation of the aptamer on the sensor: 5' thiolated aptamer immobilised on gold



- Affinity:  $K_d = 15 \text{ pM}$

# Quartz crystal biosensor 1

- Target molecule: human IgE
- DNA aptamer compared with anti-IgE antibody
- Transducer: quartz crystal microbalance
- Immobilisation of the aptamer on the sensor surface: 5' biotinylated aptamer immobilised on streptavidin fixed on the gold surface with DSP.



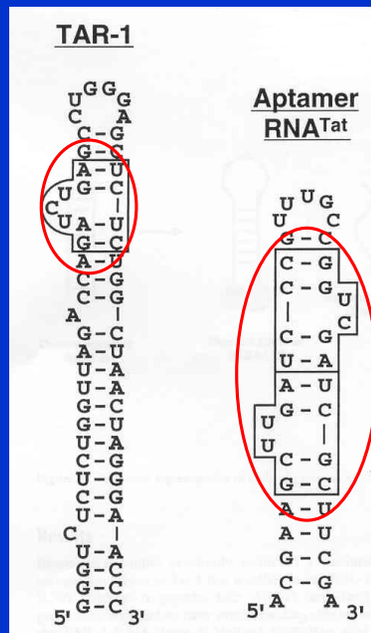
- **Detection limit** 100  $\mu\text{g/L}$  (Ab and aptamer)
- **Linear range** 0.1-1 mg/L (Ab)  
0.1-10 mg/L (aptamer)
- **Affinity**  $K_d = 1.9$  nM (Ab)  
 $K_d = 3.6$  nM (aptamer)
- **Stability** crystals modified with aptamers could be stored for several weeks

# Quartz crystal biosensor and Surface Plasmon Resonance biosensor

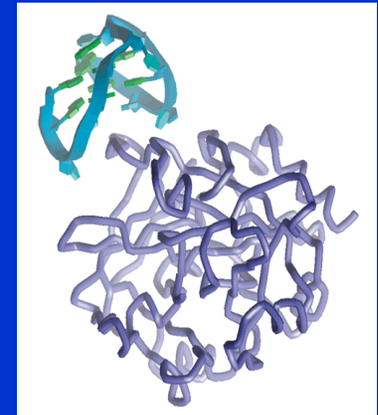
- Target molecule: HIV-1 tat protein and thrombin
- RNA aptamer (tat) and DNA aptamer (thrombin)
- Transducer: quartz crystal microbalance and SPR



Tat is a small polypeptide of 86-102 amino acids comprising a few functional regions, controlling the HIV-1 replication cycle. The arginine-rich region (49-57) of Tat is involved in binding the RNA trans-activation response element (TAR).



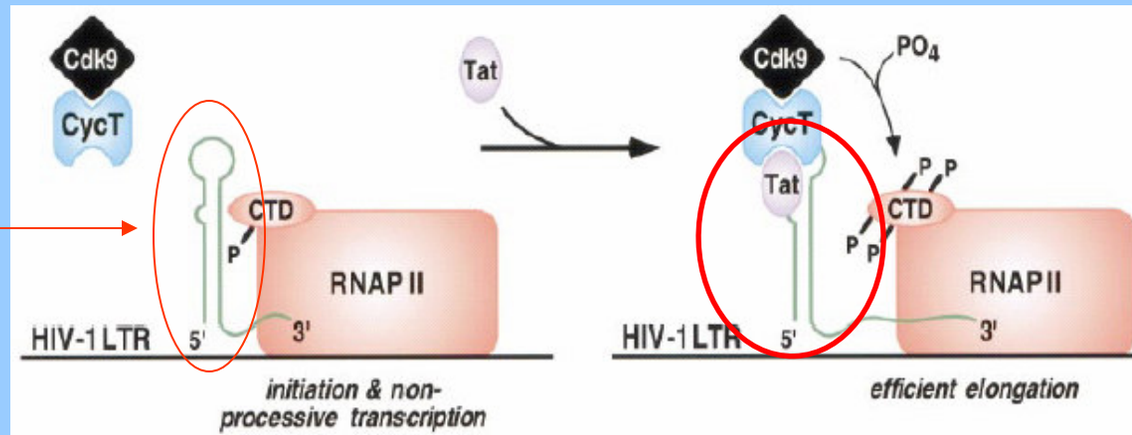
Thrombin is a serine protease and its function is to cut specifically the large protein fibrinogen into fibrin monomers. The conversion of the plasma precursor prothrombin (factor II) to  $\alpha$ -thrombin is one of the final steps in the blood cascade.



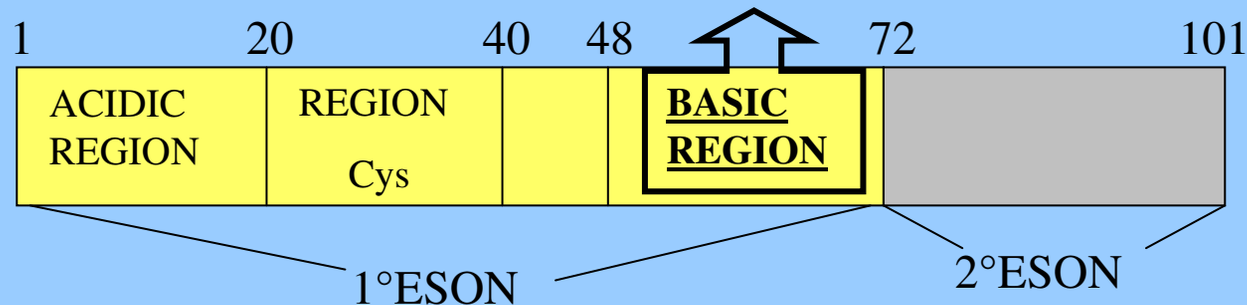
# Tat Protein (trans-activating protein)

- Protein containing 86-101 aa
- Replication HIV virus and HIV pathologies related
- Vaccin Experimental work
- Diagnostic interest

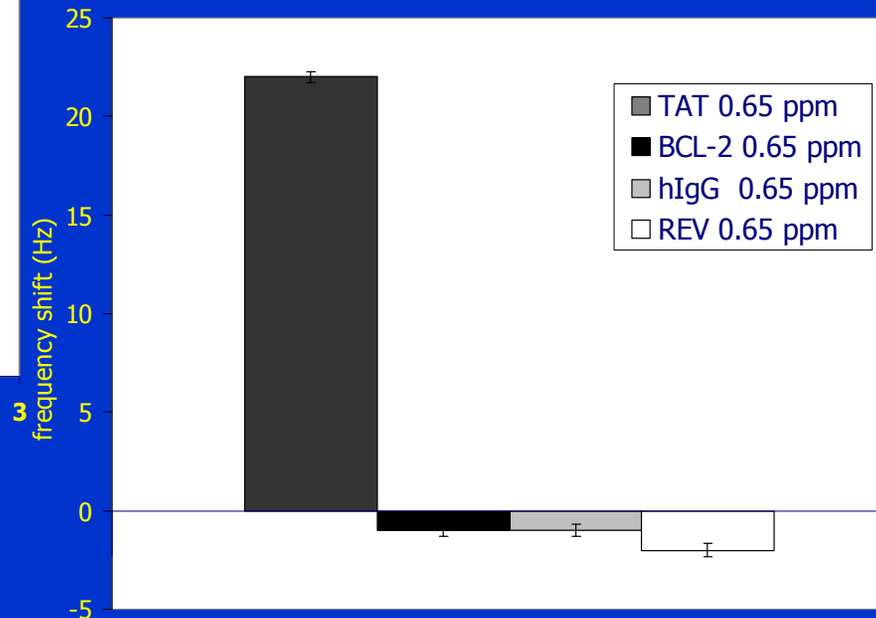
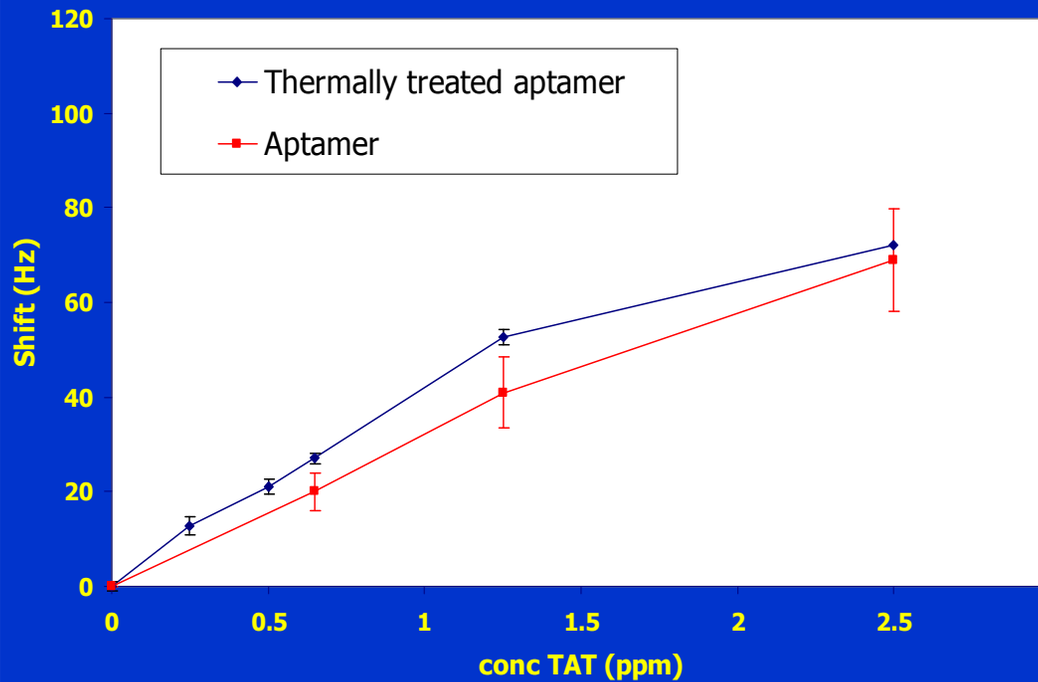
**TAR**  
(trans-activating region)



In the region 49-57 the domain for binding TAR is present



## Piezoelectric biosensor results



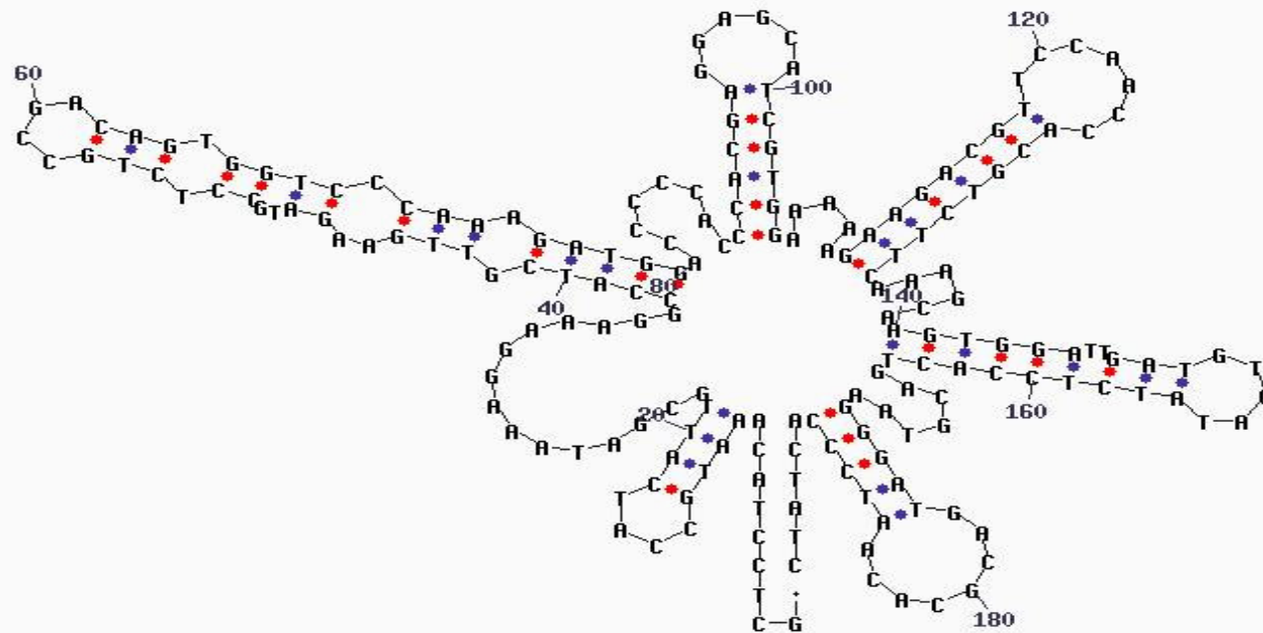
Improvements in reproducibility

Non-treated aptamer: CV%=16% (n=3 for each concentration) (1 crystal);  
CV%=21% (8 crystals)

Thermally treated aptamer: CV%=6% (n=3 for each concentration) (1 crystal);  
CV%=8% (8 crystals)

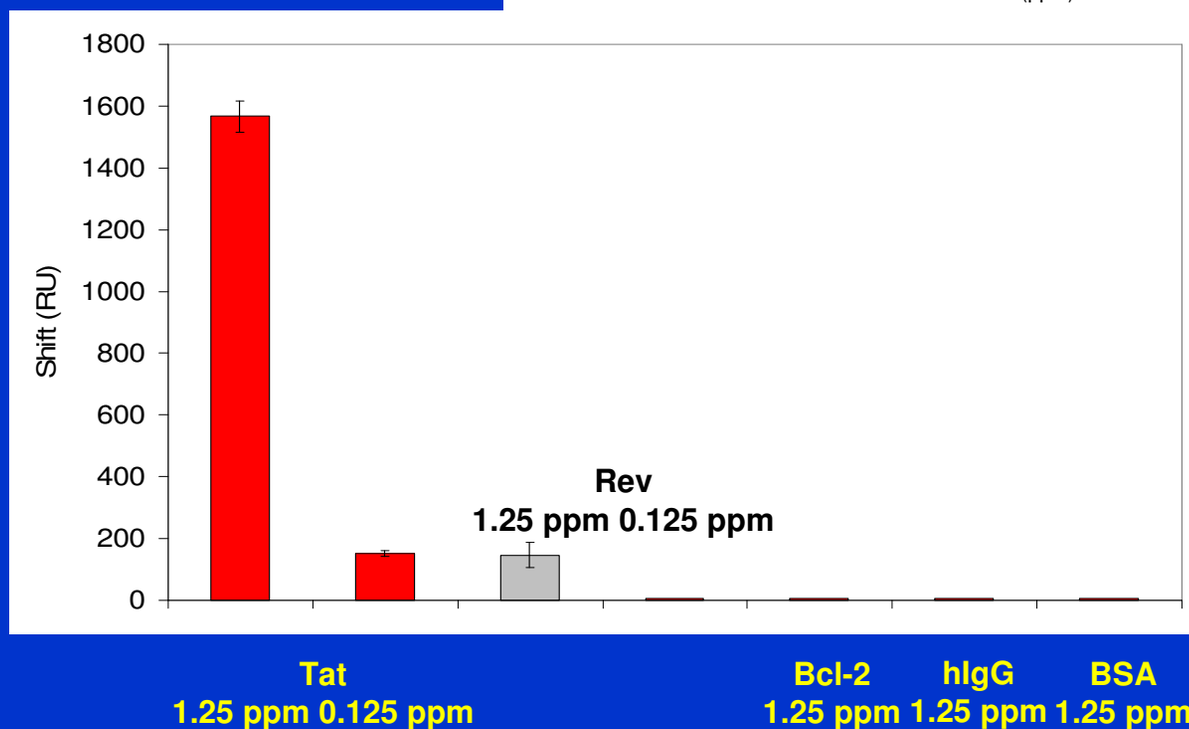
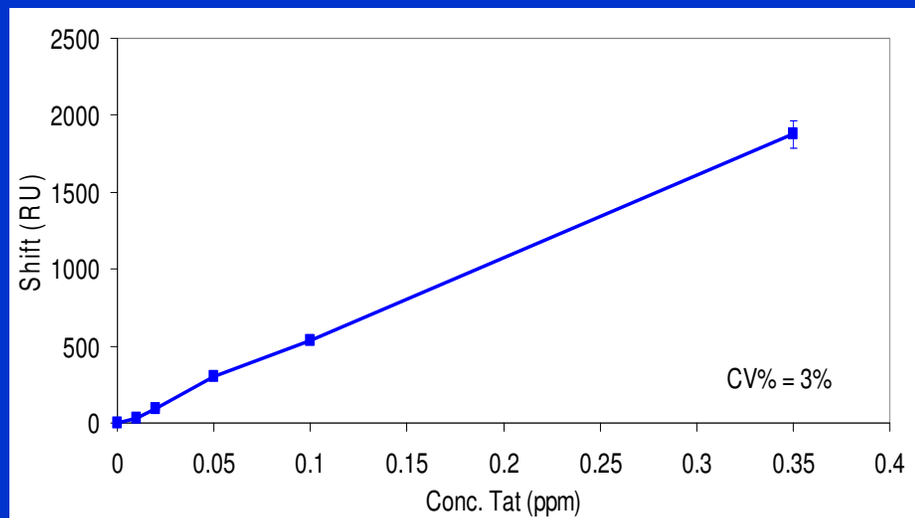
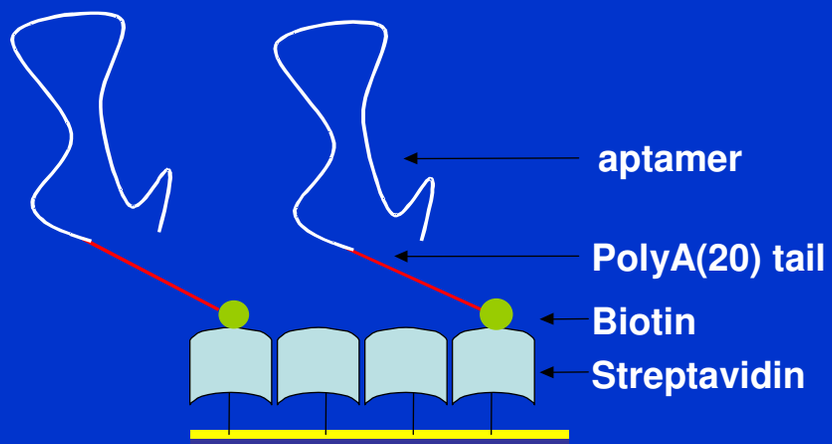
# Secondary structures of ssDNA of P35S

plt22.jpg by D. Stewart and M. Zuker  
© 2002 Washington University



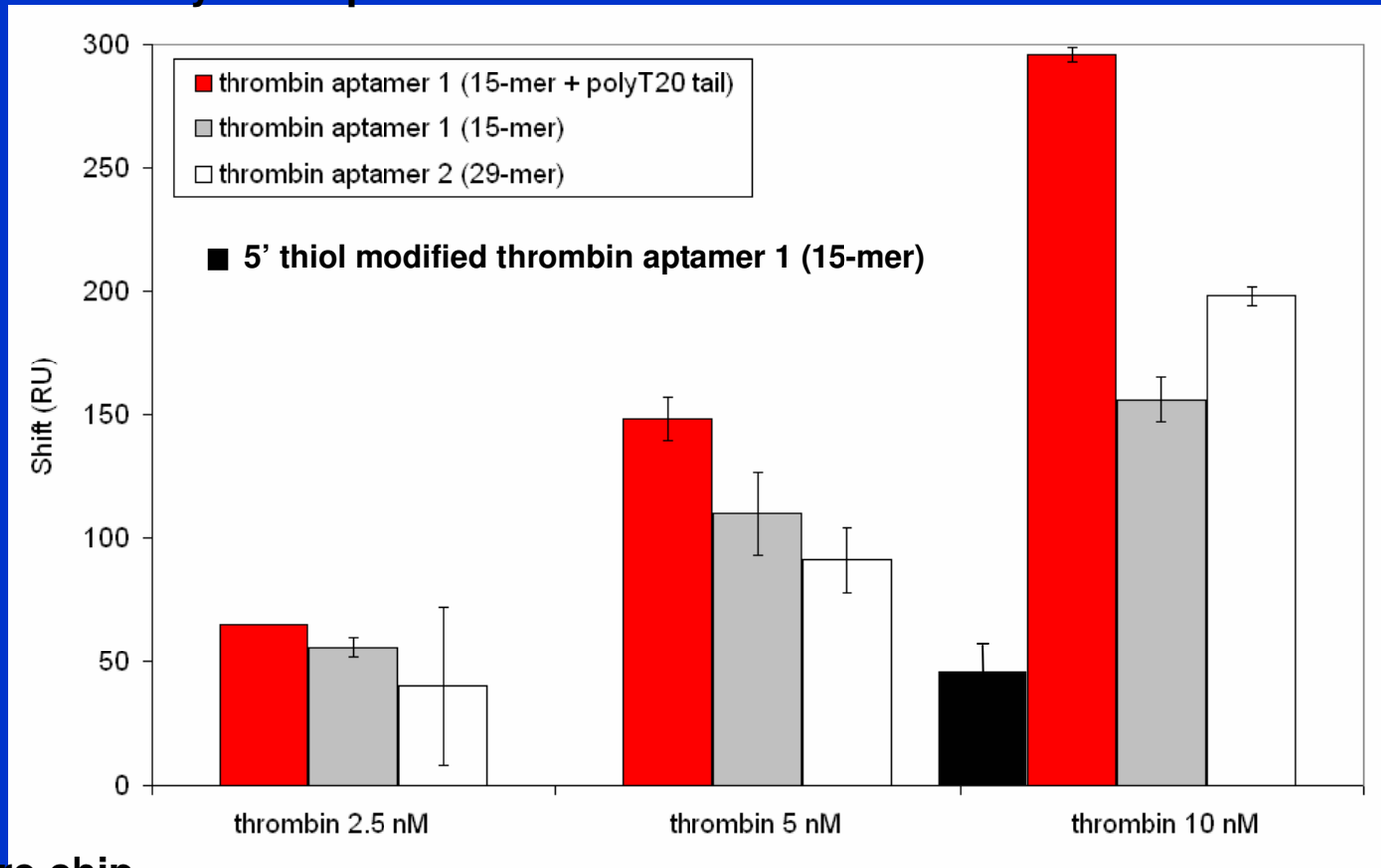
dG = -31.1 243bp

# SPR biosensor results (aptamer with polyA tail)

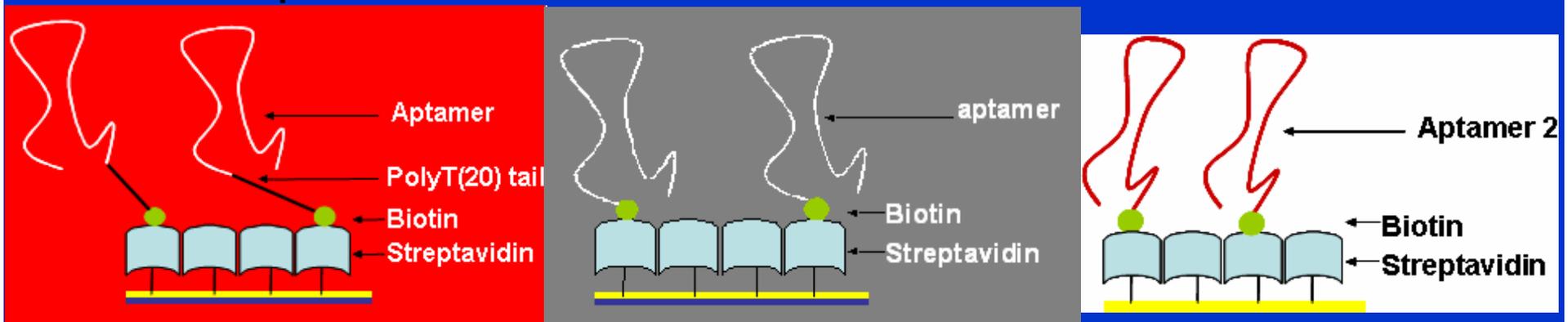




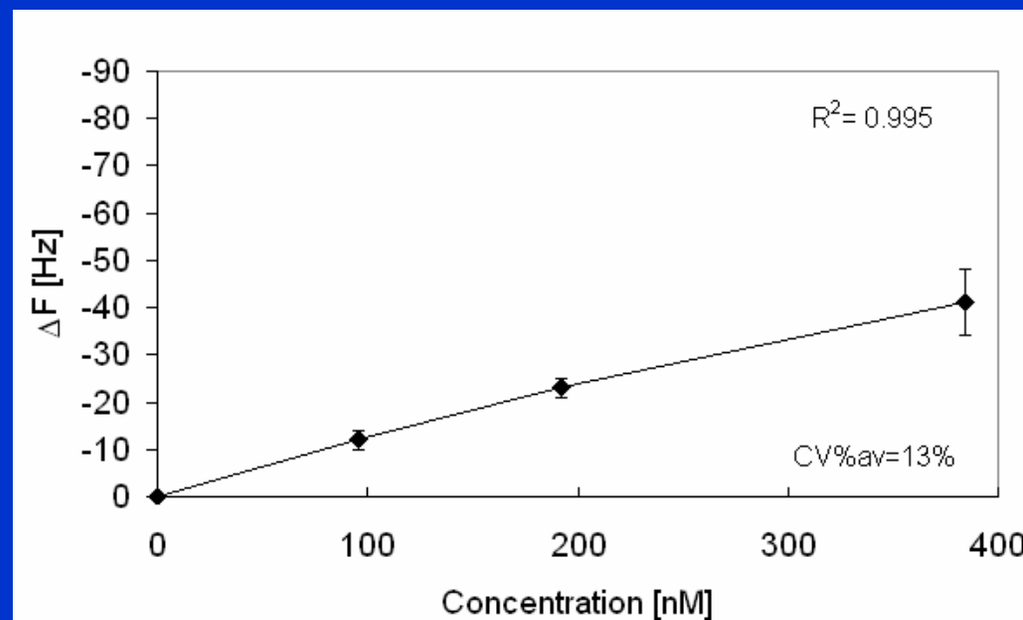
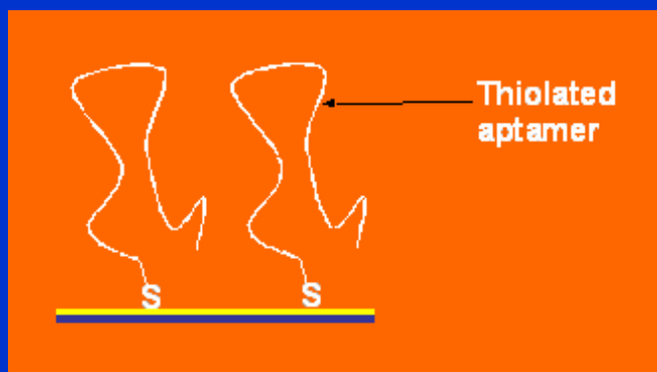
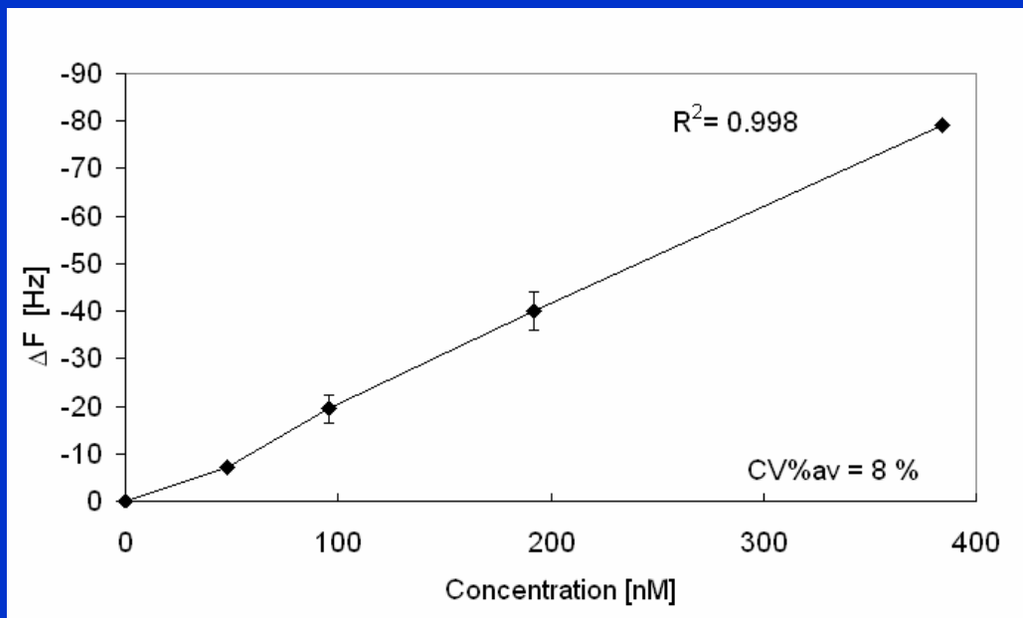
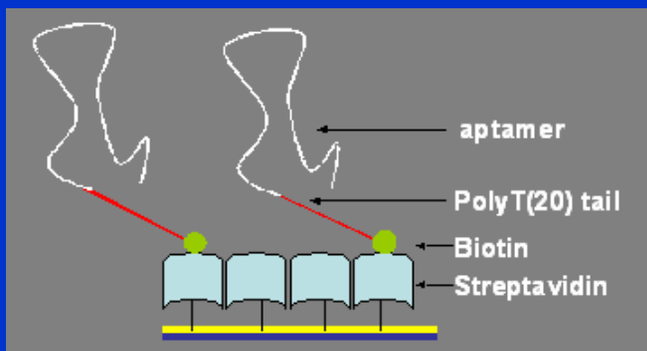
5' biotinylated aptamers



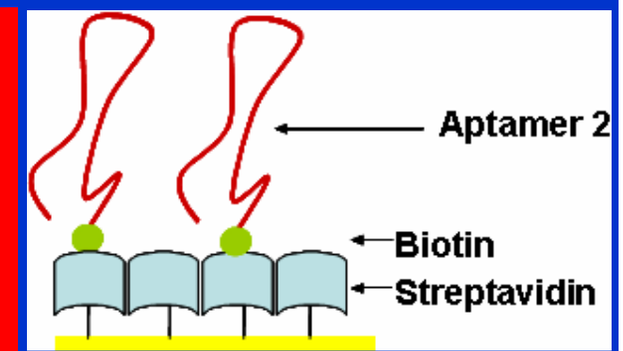
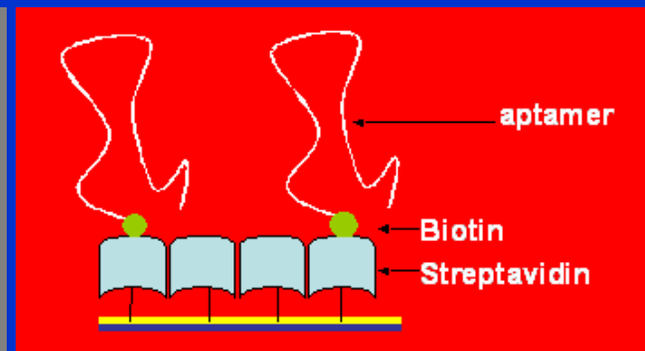
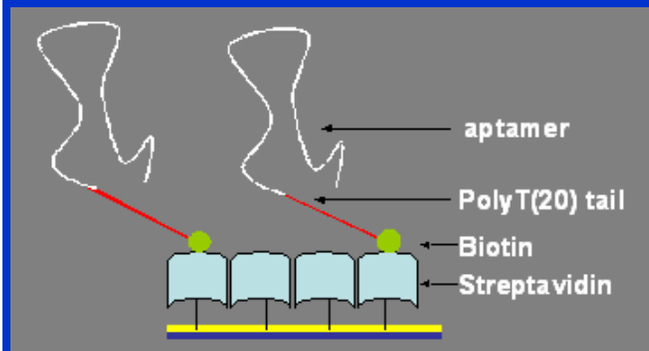
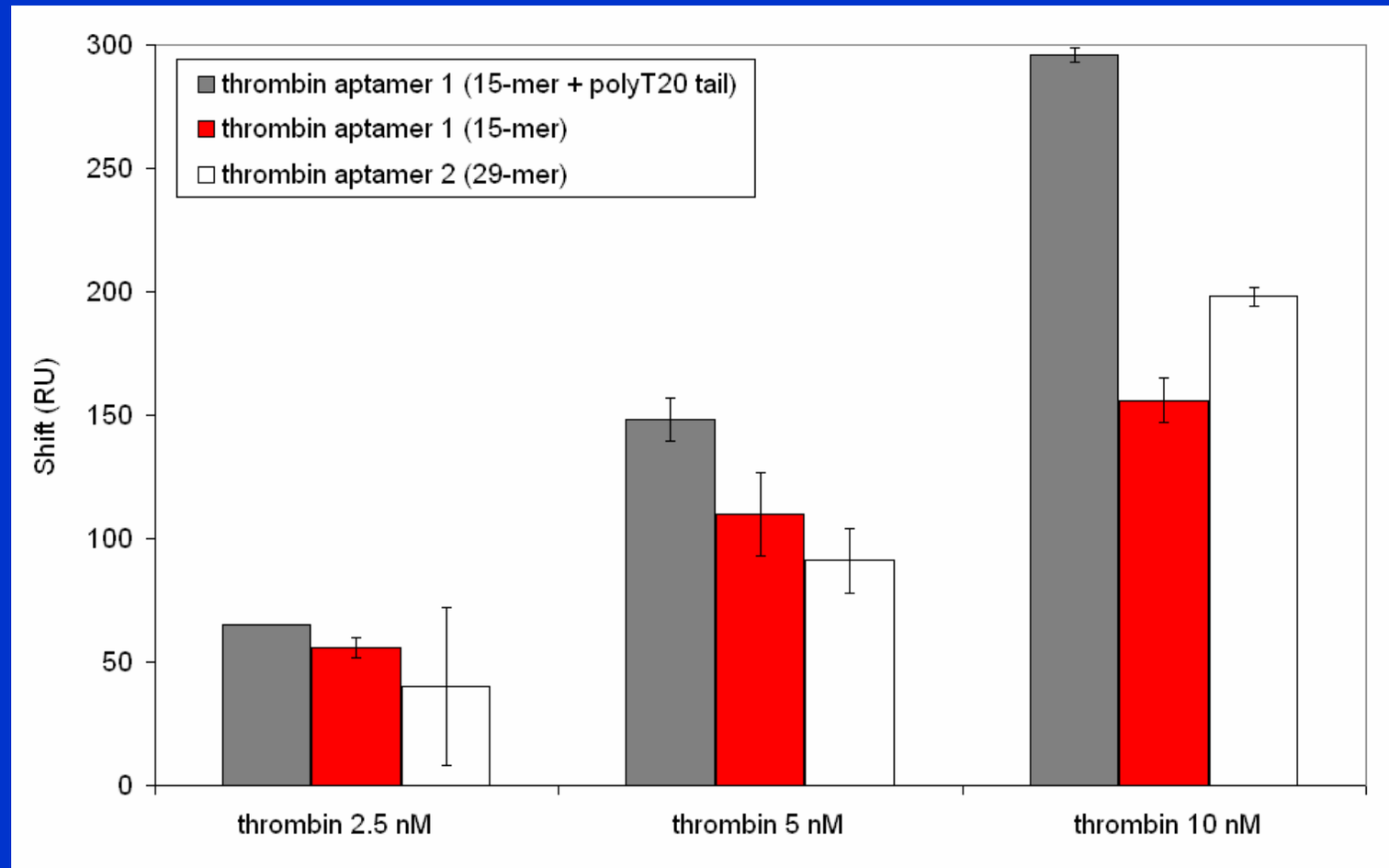
CM5 Biacore chip



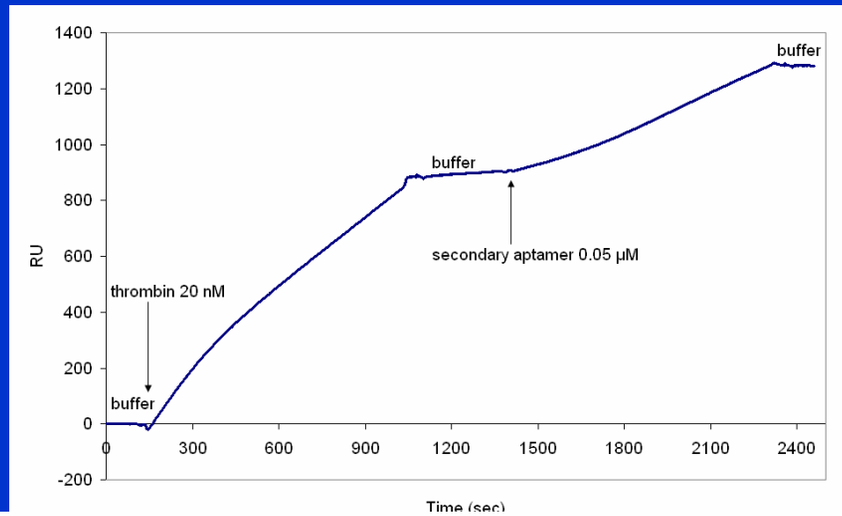
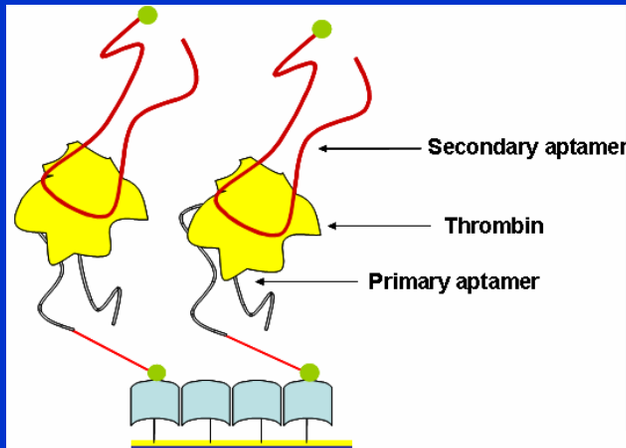
# Immobilisation optimisation QCM



# Immobilisation optimisation SPR



# Sandwich assay



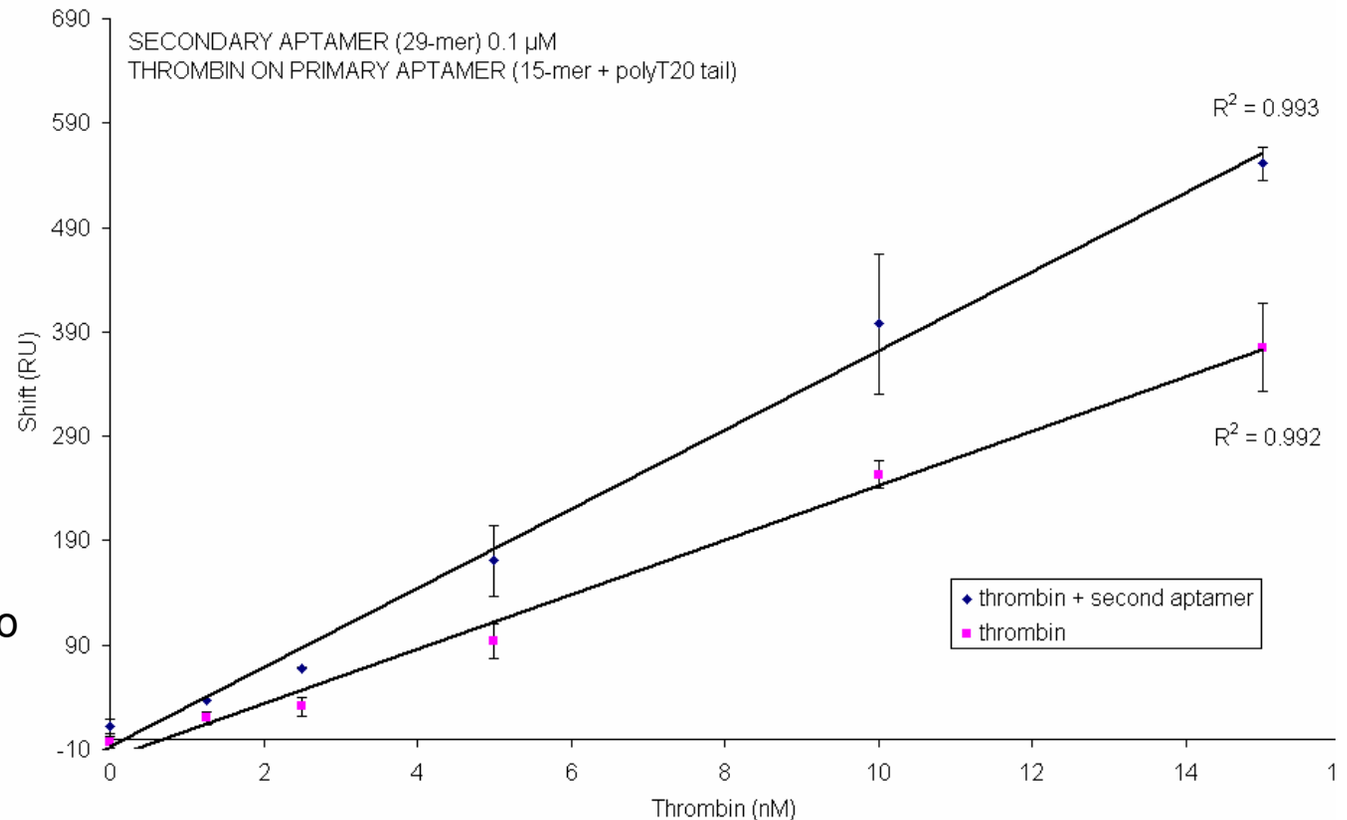
## Detection limit

1.4 nM

0.7 nM (sandwich)

(thrombin physiological conc.  
range: low nM – low μM)

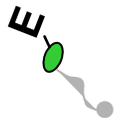
Protocol to be transferred onto  
an electrochemical biosensor  
coupled with magnetic beads



# Electrochemical sandwich assay

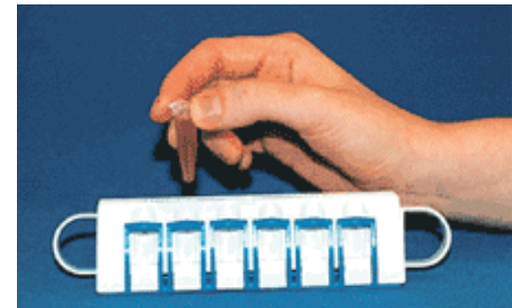
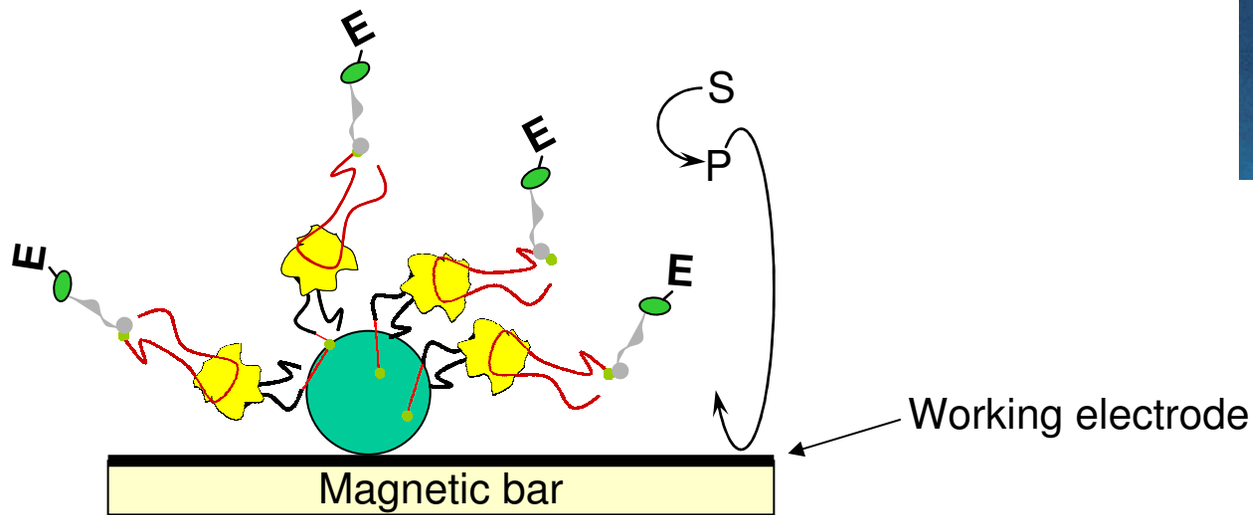
 Streptavidin-coated magnetic bead

 Thrombin

 Streptavidin-alkaline phosphatase conjugate

 5' biotinylated aptamer + polyT tail (20-mer)

 5' biotinylated secondary aptamer



Magnetic separator

# Immunochemical reactions and magnetic separation

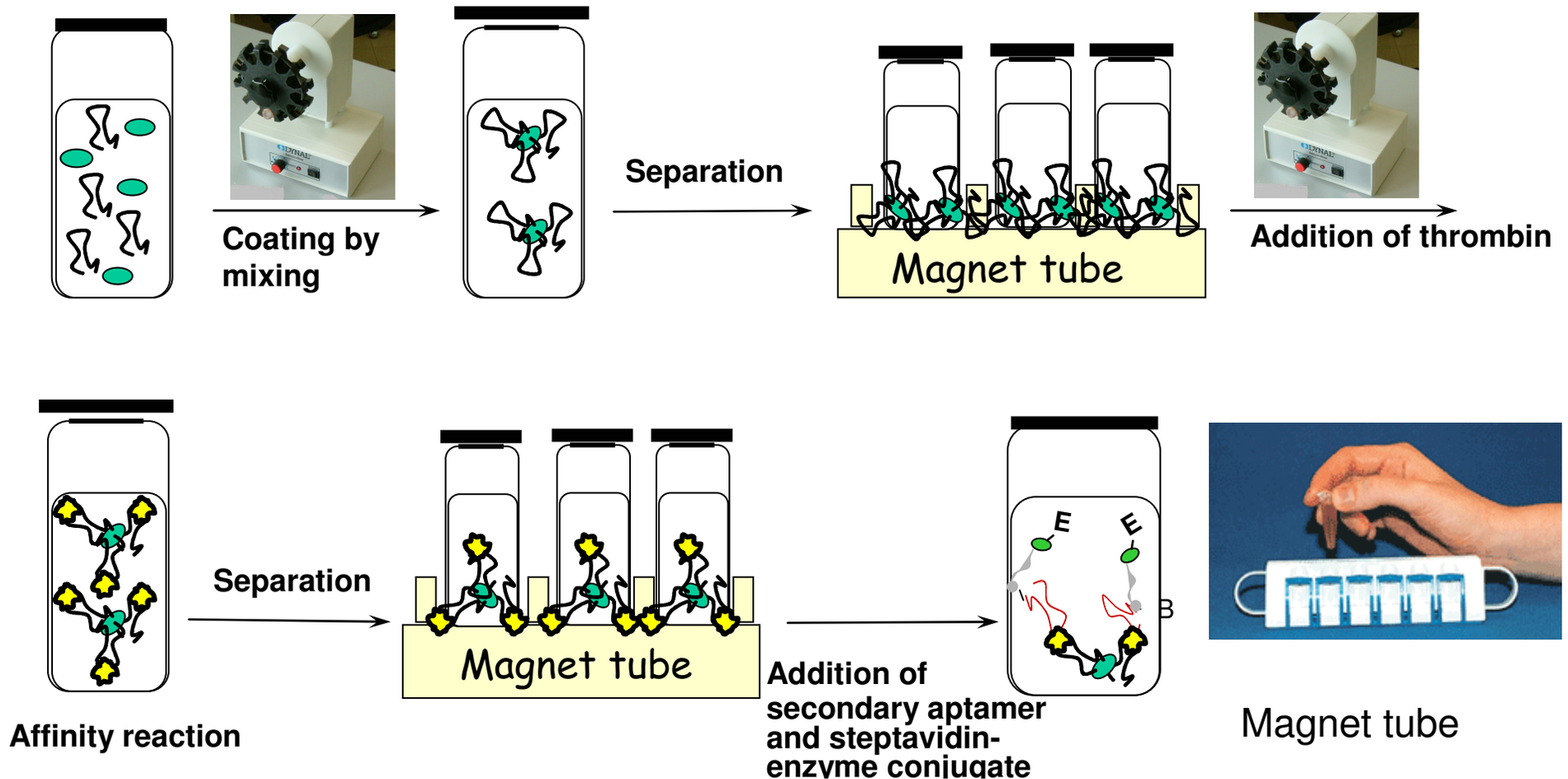
 Streptavidin-coated magnetic bead

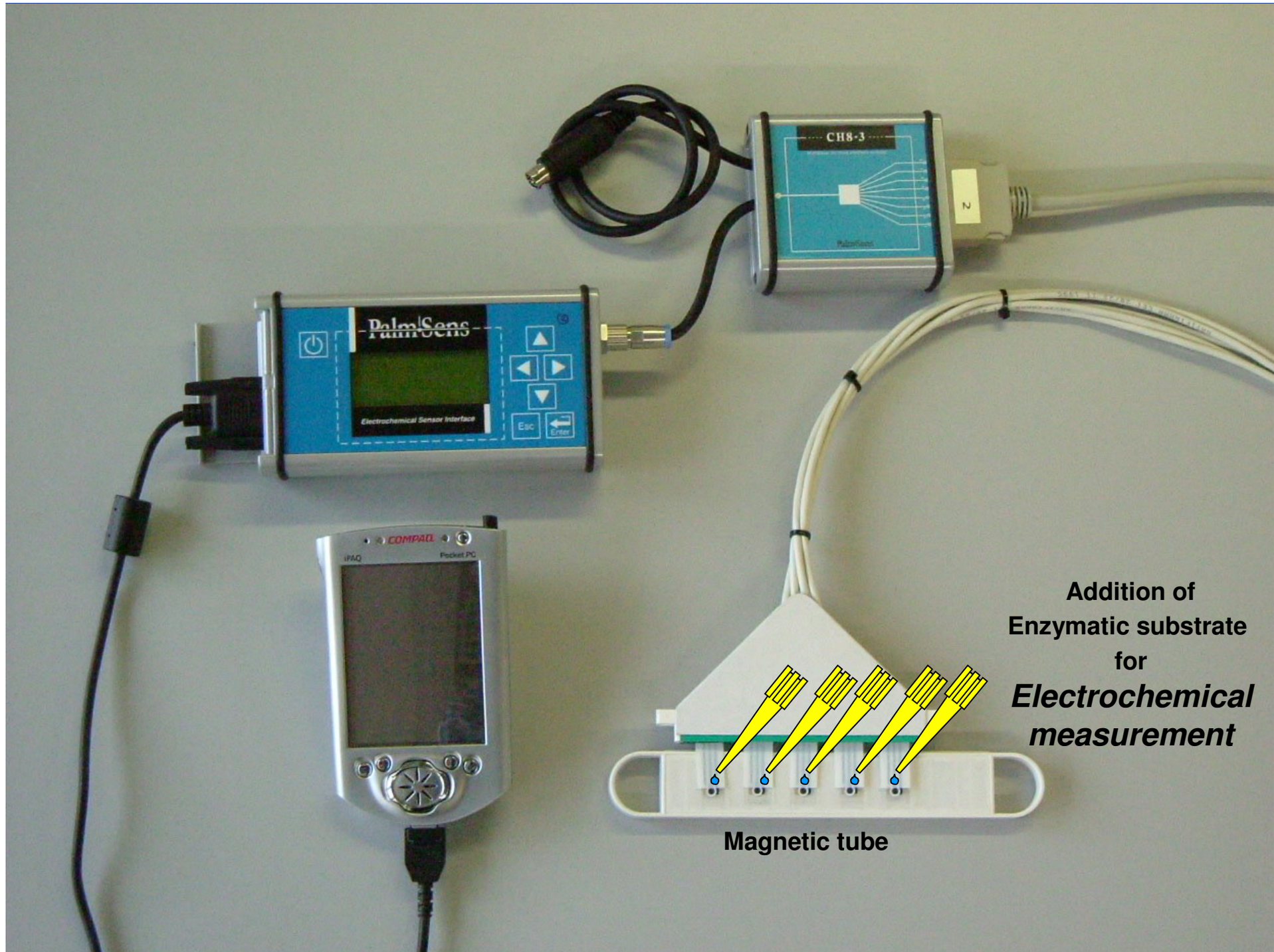
 Thrombin

 Streptavidin-alkaline phosphatase conjugate

 5' biotinylated aptamer + polyT tail (20-mer)

 5' biotinylated secondary aptamer

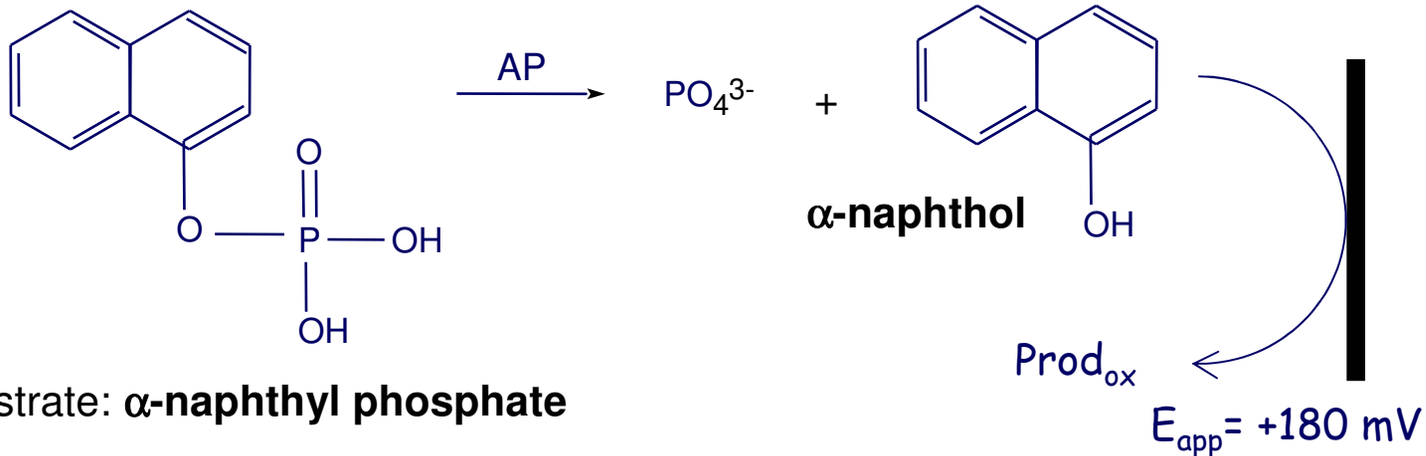




Addition of  
Enzymatic substrate  
for  
*Electrochemical  
measurement*

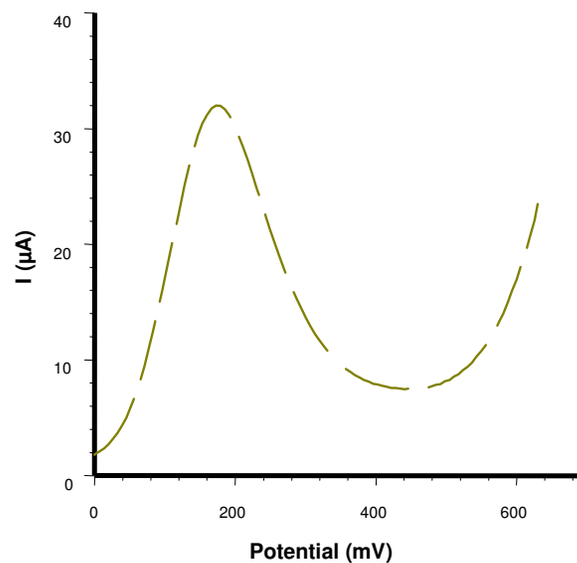
Magnetic tube

## Alkaline Phosphatase (AP)

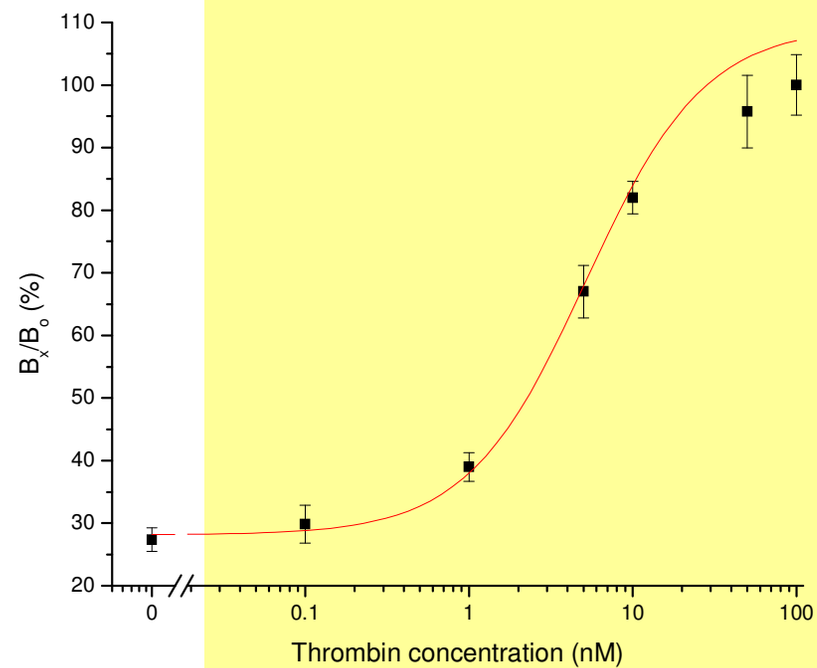
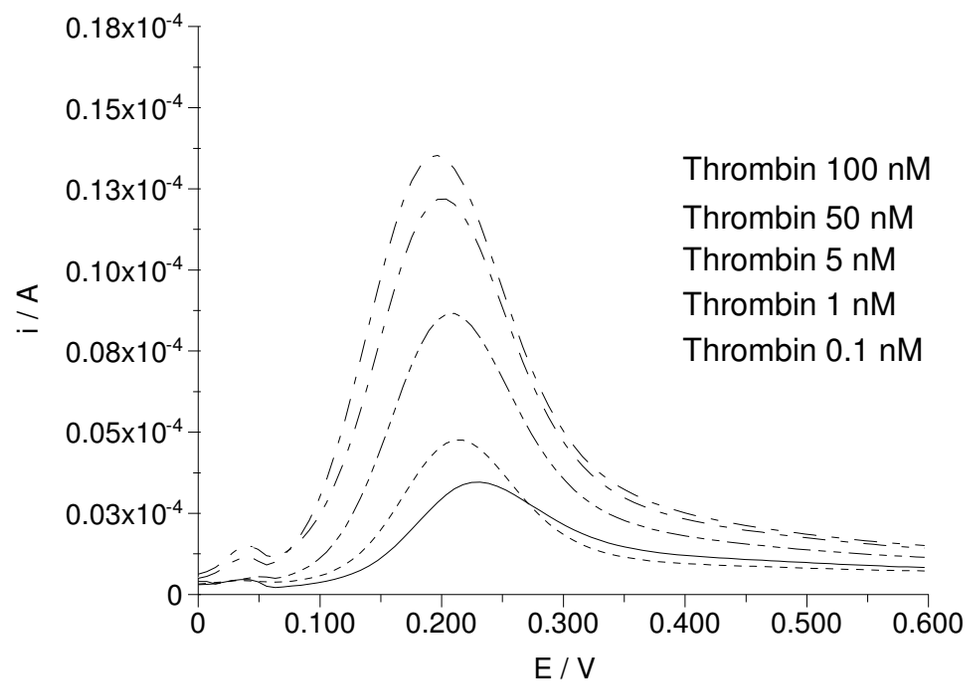


## Differential pulse voltammetry (DPV) measurements:

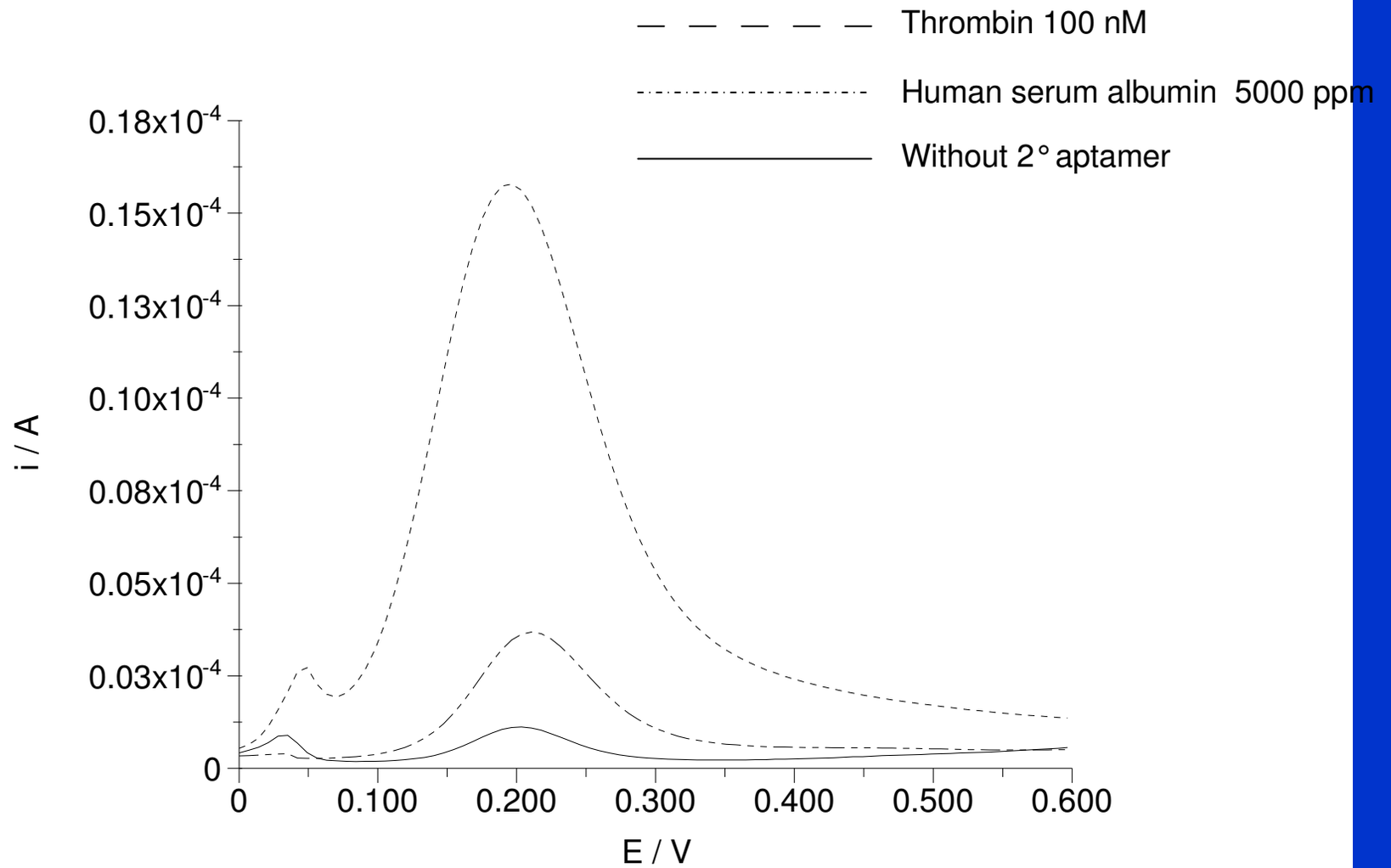
- Range potential: 0/+900 mV
- Scan rate: 70 mV/s
- Pulse amplitude: 70 mV
- Substrate: 1 mg/mL in DEA buffer



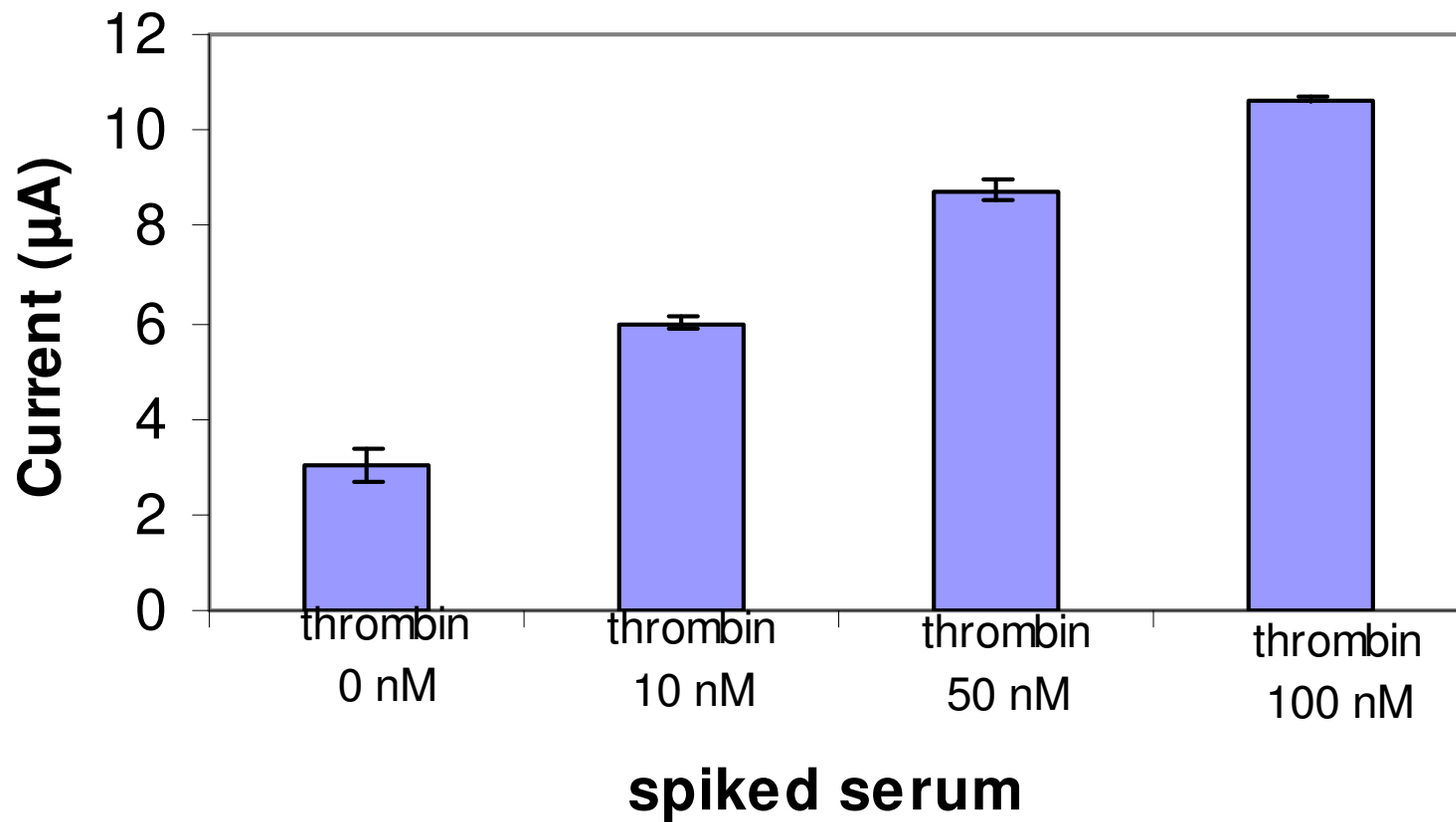
# *Dose-response curve*



# Specificity of the assay

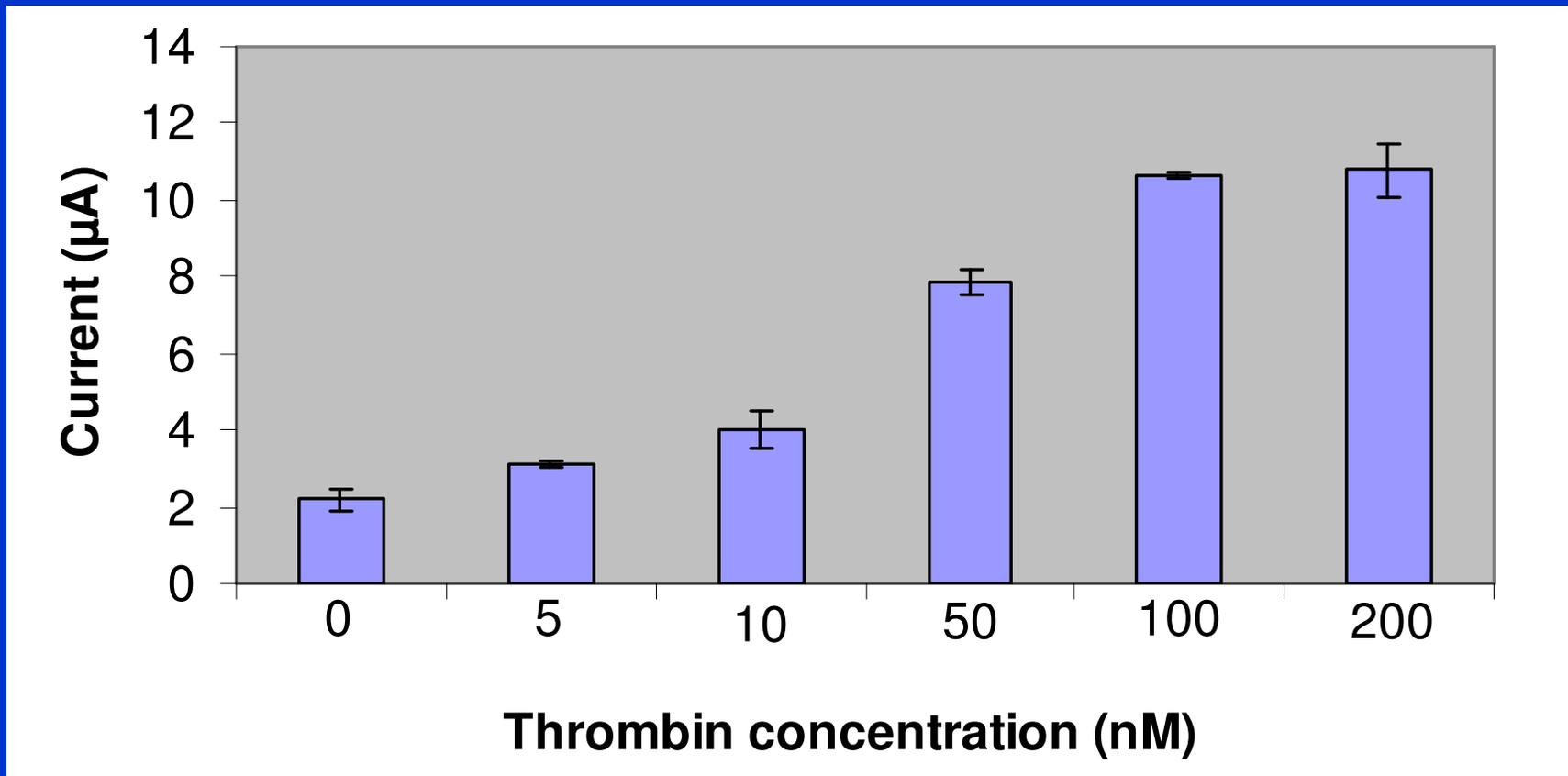


## *Measurements in serum*



## Measurements in plasma

- Precipitation of fibrinogen by  $(\text{NH}_4)_2\text{SO}_4$
- Addition of thrombin standard solutions to plasma



## *Conclusions*

Several applications based on aptamers have been reported, focusing on the parameters that need to be optimized when developing such assays (i.e. immobilization protocols, etc.). Different bioanalytical methods based on aptamers have been considered both for the detection of proteins or small molecules.

From the examination of the different protocols employed in such assays, one important point must be emphasized and that is the nature, conformation and sequence of each aptamer should be carefully considered and also stress that optimal working conditions can remarkably vary from one aptamer to another.

These important characteristics, together with the shortening of the time required for the selection process, demonstrate that aptamers can actually represent the alternative for the development of bioanalytical methods with the possibility of producing new multi-analyte aptamer-based arrays.

## Explore technologies and methods using aptamers in different applications

This book details bioanalytical technologies and methods that have been developed using aptamers in analytical, medical, environmental, and food science applications. Because aptamers have some advantages over their rival antibodies, this is a rapidly emerging field with vast potential in varied disciplines. After an introduction to aptamers, aptamer targets, and their general uses, *Aptamers in Bioanalysis*:

- Discusses different applications, with particular attention to the comparison between aptamer-based biosensors and methods versus the corresponding immunosensors
- Includes examples of aptamer-based diagnostic techniques such as whole-cell protein profiling (proteomics) and medical diagnostics for the distinction of diseased versus healthy states
- Includes chapters written by leading experts in their fields

With an overview of proven bioanalytical technologies and methods and numerous references for further study, this is a core reference for analytical chemists, electrochemists, pharmaceutical/medicinal chemists, and biotechnologists.

**MARCO MASCINI, PhD**, is a Full Professor of Analytical Chemistry in the Department of Chemistry at the University of Firenze, Italy. He has been one of the pioneers of biosensor technology. His research interests are related to electrochemical, piezoelectric, and optical biosensors. He pioneered several practical applications of these devices in industrial prototypes for use in bioanalysis, medicine, the environment, and food quality and safety. Some of these devices are commercialized today. Dr. Mascini is the author of more than 300 publications and has taken part in several European Commission projects and steering committees.

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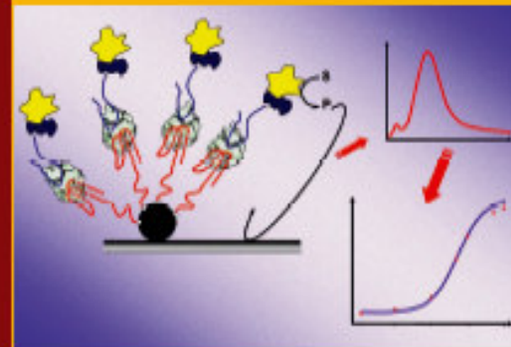
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MASCINI

Aptamers in Bioanalysis

# Aptamers in Bioanalysis



Edited by

**MARCO MASCINI**

*Thank.....*



*.....you!*