

Citation for published version: Jolly, P, Formisano, N & Estrela, P 2015, 'DNA Aptamer-based detection of prostate cancer', Chemical Papers, vol. 69, no. 1, pp. 77-89. https://doi.org/10.1515/chempap-2015-0025

DOI: 10.1515/chempap-2015-0025

Publication date: 2015

Document Version Early version, also known as pre-print

Link to publication

University of Bath

General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

1	DNA aptamer-based detection of prostate cancer
2	
3	Pawan Jolly, Nello Formisano, Pedro Estrela
4	
5	Department of Electronic & Electrical Engineering, University of Bath, Bath BA2 7AY,
6	United Kingdom
7	
8	
9	Corresponding author: Dr Pedro Estrela; P.Estrela@bath.ac.uk; Department of Electronic and
10	Electrical Engineering, University of Bath, Claverton Down, Bath, BA2 7AY,
11	United Kingdom
12	
13	Received [Dates will be filled in by the Editorial office]
14	
15	
16	The use of aptamers in biosensing have gained considerable attention as an attractive
17	alternative to antibodies because of their unique properties such as long term stability, cost
18	effectiveness and tunability to various applications. Among various cancers, early diagnosis
19	of prostate cancer (PCa) is one of the biggest concerns for ageing men worldwide. One of the
20	most commonly used biomarker for PCa is prostate specific antigen (PSA), which can be
21	found in elevated levels in patients with cancer. In this review, a presentation on the gradual
22	transition of research from antibody-based to aptamer-based biosensors is presented
23	specifically for PSA. A brief description on aptamer-based biosensing for other PCa
24	biomarkers is also presented. Special attention is given to electrochemical methods as
25	analytical techniques for development of simple, sensitive and cost effective biosensors. The
26	review also focuses on different surface chemistries exploited for fabrication and their
27	application with clinical samples. Utilization of aptamers provides a promising tool for
28	development of point-of-care biosensors for early detection of prostate cancer. In the view of
29	the unmatched upper hand of aptamers, future perspectives are also discussed, not only in the
30	point of care format but also in other novel applications.
31	

32 Keywords: DNA aptamer, biosensor, electrochemical detection, prostate specific antigen,
 33 prostate cancer, surface chemistry

35

36 37

Introduction

Prostate cancer (PCa) is a type of cancer that develops in the prostate gland, which is a part of a male reproductive system. PCa is the most commonly diagnosed cancer amongst men in Europe and the United States and is the second worldwide leading cause of morbidity. It has been reported that PCa is predominant in older men above the age of 50 (Kirk, 1997; Hoffman, 2011) and among black men (Stanford et al., 1999; Greenlee et al., 2000). It has been also projected that PCa will be the most common cancer by 2030 in the UK (Greenlee et al., 2000; Jeong et al., 2010).

Most of the PCa generate in the epithelium cells (Bostwick, 1989). As androgens regulate cell division of the gland epithelium (Ross et al., 1998), these hormones are believed to be the main cause of PCa. However, a study demonstrating a consistent correlation between androgens and prostatic carcinogenesis has not yet been reported to date and the precise causes that lead to PCa are still not well understood (Kufe et al., 2003).

PCa often develops very slowly and the lack of symptoms during the early stages of 50 the disease leads to a late diagnosis of the tumour. Moreover, if diagnosed at a late stage, no 51 effective treatments are currently available for its cure. In many cases PCa does not show any 52 clinical manifestation during the lifetime of a patient, who might die for non-related PCa 53 causes. However, for those patients that develop a more aggressive cancer form, PCa cells can 54 break away from a prostate tumour and metastasise. Since the prostate is well connected to 55 numerous lymph nodes, the spread is easy and some of the most common sites of PCa 56 metastatic process are bones (Chou & Simons, 1997). 57

- 58
- 59

60

- 00
- 61

There is no solitary test for the diagnosis of PCa. Moreover, all the tests which are used to diagnose have pros and cons which are usually discussed by the doctors with their patients. The most commonly used methods for PCa detection are: digital rectal examination (DRE), transrectal ultrasound (TRUS), biopsy and PSA blood test.

Current detection methods

In DRE, a doctor inserts a gloved finger into the rectum and examines for bumps or swelling of the prostate gland. It is an inexpensive method and can also detect PCa

irrespective of changes in the level of prostate specific antigen (PSA) in blood. Accuracy of 68 diagnosis can be increased when DRE is combined with PSA tests and biopsy results (Uzzo et 69 al., 1995; Basler & Thompson, 1998; Jeong et al., 2010). In comparison to DRE, in the TRUS 70 method an ultrasound probe is inserted into the rectum, emitting energy sound waves to image 71 the prostate gland. It is a very useful tool to understand pathology of tumours and in guiding 72 needle biopsies for sampling of tissue (Aus et al., 1996; Irani et al., 1997). For a biopsy, a 73 small section of the tissue is removed through the rectum using a needle and is 74 microscopically examined by pathologists. It requires a high number of samples from the 75 prostate making it a painful protocol. Not only the results from biopsies are controversial, 76 there is also a high risk of severe infections with subsequent biopsies (Jeong et al., 2010; Loeb 77 et al., 2013). 78

The most frequently used test for PCa screening is the quantification of levels of PSA 79 in blood. If PSA levels are higher than the cut off levels of 4 ng/ml, biopsy procedures are 80 considered (Catalona et al., 1991, Jeong et al., 2010; Savory et al., 2010). However, the levels 81 of PSA in blood in ageing men can also be raised due to other factors like benign prostatic 82 hyperplasia (BPH) and prostatis, which could lead to an over-diagnosis in men (Carter et al., 83 1992). Consequently, due to faulty diagnosis, patients undergo biopsy surgery making PSA 84 testing a controversial diagnostic tool. Due to these controversies with PSA testing, in May 85 2012 the US Preventative Services Task Force recommended against PSA screening in all 86 men. This emphasized the need for more reliable biomarkers for diagnosis of the disease 87 (Moyer, 2012). 88

89

90

91

Prostate specific antigen (PSA): a PCa biomarker

92

PSA belongs to the family of kallikrein proteins which are defined as serine proteases. There are about 15 kallikrein family members that have been identified in humans. PSA is the only kallikrein specific to prostate (hK3). Pancreatic renal kallikrein (hK1) and human glandular kallikrein (hK2), which are androgen regulated, are also expressed in the prostate (Balk et al., 2003).

PSA is synthesised in its inactive form: a 244 amino acid long protein called pro-PSA.
Pro-PSA is cleaved from the N terminus in the prostate by the hK2 enzyme leading to active
PSA which is a 237 amino acid long protein (Takayama et al., 1997). The active PSA is a
30 kDa protein which can be found in both serum and semen of men. PSA is present in semen

in the range of 0.5 - 2 mg/ml and its physiological role is to de-coagulate semen by breaking 102 down the proteins semenogelin I and II (Lilja et al., 1987; Lövgren et al., 1999). In prostate 103 cancer there is release of both active PSA and pro-PSA due to rupture of the basal membrane. 104 Moreover, internally cleaved forms of PSA (with no enzymatic activity) also enter the blood 105 stream but remain un-complexed and are taken into the free PSA (fPSA) count. However, 106 when active PSA enters the blood stream it becomes immediately complexed with protein 107 inhibitors. Most of the assays employing antibodies measure the total amount of PSA (tPSA) 108 (Takayama et al., 1997). 109

Many studies reported that PSA levels are directly proportional to the stage of the 110 cancer and to the volume of the tumour (Stamey et al., 1987; Grossklaus et al., 2002; Pinsky 111 et al., 2007; Lilja et al., 2008). PSA detection results are nowadays highly sensitive (Madu & 112 Lu, 2010) and reasonably inexpensive. Moreover PSA testing is a more accepted procedure 113 by patients compared to DRE and this has augmented the early detection of PCa (Balducci et 114 al., 1997). However, even though PSA testing induced a decrease in PCa mortality of 20% its 115 screening led to over-diagnosis and over-treatment (Andriole et al., 2009) of patients that 116 would have not been clinically affected by the tumour during their lifetime. Over diagnosis 117 can, in fact, lead to unnecessary treatments and increase the state of anxiety in patients. 118 Conversely, clinicians are not able nowadays to discriminate between a harmless or lethal 119 form of prostate cancer and so to decide whether the patient needs a treatment. Once a 120 prostate cancer has been definitively treated, PSA screening is the most reliable and fast 121 means that enable to detect a contingent recurrence of the tumour (Lilja et al., 2008). 122

With the shortcomings of the current tests for PCa, including PSA testing, there is a concerted effort to look for alternatives. However, it would be a challenge to replace PSA entirely due to its minimally invasive nature and low cost. Instead, there is a pressing need to look for other biomarkers to complement PSA that can increase the specificity and sensitivity of PCa screening and inform prognosis and treatment courses.

One path currently being looked at when a high level of PSA is detected in patients 128 with cancer, is to differentiate PSA into different forms namely free PSA (fPSA) and total 129 PSA (tPSA) and quantify them independently. One of the approaches is to measure the ratio 130 of free PSA to total PSA in the blood. It has been proven, in fact, that the levels of fPSA are 131 lower in patients with PCa than in patients with BPH (Christensson et al., 1993), which can 132 thus be an indication of the aggressiveness of the cancer. However, the method can cause 133 false negative results as the amount of fPSA can be higher in patients with larger prostate 134 volume (Stephan et al., 1997; Catalona et al., 1998). Nevertheless, the ratio of free to total 135

PSA when combined with the total PSA levels increases the confidence of the diagnosis(Velonas et al., 2013).

138

139 Pro-PSA

Several studies are also focused on the detection of a distinct form of free PSA, called 140 proenzyme PSA (pro-PSA). Pro-PSA is an enzymatically inactive precursor of PSA obtained 141 by co-translational removal of an amino-terminal leader. The N-terminal of pro-PSA can be 142 cleaved at various positions resulting in different forms of pro-PSA. Pro-PSA truncated 143 between the third and second amino acid is called [-2]pro-PSA and is believed to provide a 144 better discrimination between cancerous and benign form of prostate disorders (Mikolajczyk 145 et al., 2001; Mikolajczyk et al., 2004). Increased values of other forms of pro-PSA ([-5] and 146 [-7]) have also been associated to PCa. A truncated precursor form of prostate-specific 147 antigen is therefore a more specific serum marker of prostate cancer. 148

149

150 PSA density

A better discrimination of BPH from PCa might be achieved by measuring the ratio of PSA to prostate volume. However, this parameter called PSA density showed contradictory evidence on the tumour aggressiveness and malignity (Stamey et al., 1987; Ohori et al., 1995). Furthermore, in order to obtain prostate volume values, TRUS is required in addition to the standard PSA test with a consequent discomfort for patients as well as an increase in the cost and time required to perform the test. For these reasons PSA density has not been rest extensively employed as a routine test for PCa.

158

159 PSA velocity and PSA doubling time

PSA velocity refers to the rate of serum PSA increase over time while PSA double time refers to the time required for a given PSA level to be doubled. As the previous PSA derivatives, also PSA velocity can be used to distinguish a prostate cancer from a BPH (Carter et al., 1992). Both PSA velocity and PSA double time are used to monitor the recurrence of the tumour after treatment (D'Amico et al., 2004; D'Amico et al., 2005). Again, some studies compared the responses from PSA velocity and PSA double time with biopsy results demonstrating how these two PSA derivatives can fail the diagnosis (Melichar, 2012)

168 Age-specific PSA reference ranges

Since the level of PSA increases with the age of men, scientists studied this correlation in order to obtain a median value of PSA for given ranges of age. By comparing the PSA level with the median PSA for that patient's age (age-specific PSA) a better choice might been taken before ordering biopsies (Loeb & Catalona, 2007).

- 173
- 174
- 175
- 176

Oligonucleotide Aptamers

In recent years, a range of assays for PSA detection such as electrochemical assays 177 (Okuno et al., 2007; Panini et al., 2008), enzyme linked immunosorbent assays (Acevedo et 178 al., 2002), cantilever assays (Wee et al., 2005), and chemiluminescent immunoassays 179 (Albrecht et al., 1994; Seto et al., 2001) have been developed. These assays are mostly based 180 on antibodies as recognition elements. One of the alternatives to antibodies is aptamers which 181 can offer several advantages with respect to the former. However, an enormous research is 182 being carried out to prove if antibodies can be replaced by aptamers to develop a real 183 biosensor for clinical applications. The scope of this review is to highlight the major 184 developments on PSA aptasensors and their potential to be used with real clinical blood 185 samples. 186

Oligonucleotide aptamers are single stranded DNA or RNA sequences that can bind to 187 a target molecule with high specificity and affinity. Aptamers had already been widely used in 188 drug delivery applications and are now being extensively studied as new emerging 189 bioreceptors for biosensors (termed aptasensors) (Hianik & Wang, 2009; Iliuk et al., 2011). 190 Aptamers have shown comparable or even stronger binding than antibodies towards a broad 191 range of targets (e.g. proteins, peptides, amino acids, drugs, whole cells, etc.), especially with 192 the development of novel selection technologies (Xiao et al., 2005). The high affinity of the 193 aptamers towards the target molecule is defined by their capability of undergoing 194 conformation changes upon the binding event (Hermann & Patel, 2000; Song et al., 2008; 195 Hianik & Wang, 2009). Although using aptamers have many added advantages over 196 197 antibodies, they still need careful consideration while fabricating a biosensor. For instance, binding of an aptamer to protein might be affected by changing buffer conditions. Also, as 198 aptamers are oligonucleotide sequences, special care is needed as they are sensitive to DNase 199

200 and RNase activity. Furthermore, the k_d value of aptamers is often not as good as that for 201 antibodies.

Aptamers are developed using an in vitro selection process based on Systematic 202 Evolution of Ligands by EXponential enrichment (SELEX) (see fig. 1). Briefly, it consists of 203 three steps that are repeated systematically in order to identify the oligonucleotide sequence 204 that binds better to the target. The first step is called library generation, where a library 205 consisting of random DNA or RNA sequences (usually 30-40 base-pairs long) flanked by the 206 primer binding site are used. The library is then incubated with the target molecule. 207 Thereafter, the target bound library is separated from unbound library. Finally, the target-208 bound library is amplified using polymerase chain reaction (PCR) to create a new library to be 209 used in the next round. Aptamers binding and conformation characteristics are identified 210 using various biological assays (Syed & Pervaiz, 2010; Liu et al., 2012). 211

212

213



Fig. 1. The general SELEX protocol. Starting with a random library followed by incubation
with the target. Later the bound sequences are separated and further amplified for the
next round of selection. Adapted from Song et al. (2008).

- 219

There has been an intense interest in understanding the in-depth of ligand-binding and 221 conformational properties of aptamers. Aptamers have many advantages over antibodies, 222 making them very important molecular tools for both diagnostics and therapeutics. For 223 instance, selection of aptamers is an *in vitro* process and they can be raised to a wide variety 224 of targets ranging from small molecules and toxins to large proteins and even whole cells. 225 Secondly, aptamers, once selected, can be synthesised with high purity and reproducibility. 226 Also, as compared to antibodies, aptamers are usually highly chemically stable. Furthermore, 227 they can undergo significant conformational changes in their structure upon binding with the 228 target – a feature which can be exploited for biosensing applications. This offers great 229 flexibility to design novel biosensors (Clark & Remcho, 2002; Tombelli et al., 2005; Willner 230 & Zayats, 2007; Mairal et al., 2008; Song et al., 2008; Liu et al., 2012). 231

- 232
- 233
- 234
- 235

PSA detection

PSA is currently detected in dedicated laboratory settings using automated analysers 236 running antibody-based assays which are generally expensive and time consuming (Lin & Ju, 237 2005; Healy et al., 2007). Cost effective, easy to use and possibly portable devices are 238 required in order to allow more powerful tools for early detection of prostate cancer. To date, 239 researchers have exploited several techniques for PSA detection such as optical (Besselink et 240 al., 2004; Huang et al., 2005; Cao & Sim, 2007), piezoelectric (Weeks et al., 2003; Wee et al., 241 2005) and electrochemical (Sarkar et al., 2002; Fernández-Sánchez et al., 2004; Liu et al., 242 2013). 243

Although label-free-based biosensors can provide many advantages, label-based 244 approaches are still intensively studied and can offer interesting features such as low limit of 245 detection due to amplification strategies. An interesting magnetic bead-based detection system 246 for PSA detection was developed by Zani et al. (2009): paramagnetic microparticles were 247 adsorbed on an array of screen-printed electrodes and PSA was sandwiched in between two 248 antibodies on the beads; the alkaline-phosphatase-labelled secondary antibody could be 249 detected with differential pulse voltammetry (DPV) to achieve a detection limit of 1.4 ng/ml. 250 A limit of detection as low as 0.5 pg/ml in undiluted serum samples was obtained by Mani et 251 al. (2010) by combining a multienzyme-labelled immunoassay with gold nanoparticles 252 sensing surface: in this case the secondary antibody was bound to micromagnetic HRP-253 labelled beads, which massively amplified the current signals for a very low PSA detection 254

255 limit. A similar detection technique was improved and integrated in a microfluidic system by 256 Chikkaveeraiaha et al. (2011) reaching an even lower detection limit. A fascinating 257 electrochemiluminescence-based immunoassay was developed by Sardesai et al. (2011) for 258 both PSA and interlukin 6 (IL-6) by using single-wall carbon nanotubes (SWCNT) fabricated 259 on microwells and a sandwich assay where the secondary PSA antibody was functionalized 260 with RuBYP-Silica particles: the detection limit achieved was of 1 pg/ml for PSA.

261

262 Label-free electrochemical sensors for PSA detection

Electrochemical techniques are widely employed in biosensing devices as they can be 263 highly sensitive, simple to perform and cost effective. An electrochemical biosensor involves 264 an electrode surface that is functionalised with a molecular recognition element for sensing 265 biomolecules. Binding of an analyte to this element results in an electrical change in current 266 transfer (amperometric), voltage (potentiometric and field effect transistors), impedance 267 (impedimetric), conductivity (conductometric) or ion charge across the electrode, which can 268 be quantified and correlated to the amount of analyte captured. As mentioned in the previous 269 sections, most biosensors for PSA detection currently available are antibody-based. Amongst 270 the antibody-based electrochemical sensors, particularly important results are the ones using 271 label-free systems. Arya & Bhansali (2012) developed a gold biosensor modified with a 272 cysteamine self-assembled monolayer (SAM) for PSA detection. Li et al. (2005), on the hand, 273 employed In_2O_3 nanowires and carbon nanotubes. Electrochemical impedance spectroscopy 274 EIS) based sensors have been reported by Chiriacò et al. (2013) and Chornokur et al. (2011). 275 The former exploits a combined use of two different antibodies for both free and total PSA, 276 while the latter reported on a miniaturized sensor obtained with photolithographic techniques 277 using a single monoclonal antibody. Another label-free antibody-based sensor which uses a 278 polycrystalline silicon field-effect transistor was reported by Huang et al. (2013). 279

- 280
- 281
- 282
- 283

Aptasensor for PSA detection

An aptasensor biosensor comprises an aptamer as a biorecognition element (Lim et al., 285 2009). Aptasensors can be integrated with different sensing techniques such as 286 electrochemical, optical, and mass sensitive. Among these varied techniques, electrochemical 287 aptasensors have been fabricated using several detection techniques, namely EIS, 288 potentiometry and differential pulse voltammetry (DPV) (Cho et al., 2009; Clark & Remcho, 289 2002; Feng et al., 2008; Ikebukuro et al., 2005; Liu et al., 2012; Numnuam et al., 2008; Wang
290 et al., 2007; Xu et al., 2005). For detection of PSA, both RNA and DNA aptamers have been
291 developed, although there are only a handful of reports on PSA biosensors using aptamers. A
292 summary of aptamer-based biosensors for PCa detection is presented in table 1.

- 293
- 294

295 Table 1. Performance comparison of different aptasensors for PCa detection

296

Method	Material	Biomarker	Detection limit	Reference
QCM-D/EIS	Gold	PSA	-	Formisano et al., 2014
EIS	Gold	PSA	1 ng/ml	Jolly et al., 2014
Optical	AuNPs	PSA	32 pg/ml	Chen et al., 2012
DPV/CV	AuNPs@GMCs	PSA	0.25 ng/ml	Liu et al., 2012
EIS	Gold	PSMA cells	-	Min et al., 2010

297

298

The first aptamer developed was a RNA aptamer (Jeong et al., 2010) that has been used to demonstrate the recognition of active PSA. Following that, a DNA aptamer was developed using a genetic algorithm with post-SELEX screening against PSA (Savory et al., 2010). To date, there is no reported literature on the application of RNA aptamers for PSA biosensing, which could be due to the long length of the sequence making it difficult to synthesise commercially.

DNA based PSA aptamer has been combined with different sensing techniques with 305 sensitivities ranging from pg/ml to ng/ml. Chen et al. (2012) were the first to report the use of 306 PSA aptamer to develop an optical based aptasensor. The conjugation of gold nanoparticles 307 (AuNPs) with DNA aptamers were used to develop an aptasensor based on resonance light 308 scattering (RLS) spectral assay. The novel technique relied on changes in resonance light 309 scattering on binding of PSA to the aptamer, with a detection limit of 32 pg/ml. Thiolated 310 DNA aptamers were immobilized on AuNPs and then a blocking step with BSA was 311 performed prior the use of the complex AuNPs-aptamers with PSA samples. In this 312 configuration, the gold surface of the nanoparticles was covered by the flexible aptamer 313 structure and as a result no aggregation of particles occurred in absence of PSA. In the 314 presence of PSA, aptamer-PSA complexes were formed and the aptamers undergo a 315 316 conformational change in their structure from flexible to rigid. The changes in aptamer

conformation exposed some parts of the AuNPs that were thus available to form AuNPs 317 aggregates upon addition of potassium chloride. This resulted in an increase in the RLS 318 signal. The assay exhibited good sensitivity and selectivity towards PSA and tests made on 319 human blood samples showed results comparable to those obtained with ELISA (relative 320 deviation < 7%). 321



323





Fig. 2. Schematic illustration of fabrication process of the aptasensor based on gold 326 nanoparticles encapsulated by graphitized mesoporous carbon (a); PSA detection (b). 327 Adapted from Liu et al. (2012). 328

329 330

With regards to electrochemical aptasensor, modification of the electrode surface is 331 one of the biggest fields of investigation. Research is typically focused on finding the most 332 suitable recognition platform to give a stable organization to the sensor interface leading to 333 optimized binding efficiency and signal outcome (Lee et al., 2005; Putzbach & Ronkainen, 334 335 2013). Liu et al. (2012) applied aptasensors based on amplification via AuNPs and

graphitized mesoporous carbon (GMCs) combined with streptavidin-biotin system for 336 electrochemical detection of PSA (see fig. 2). GMCs encapsulated AuNPs formed the first 337 layer on cleaned pyrolytic graphite electrode followed by coating with streptavidin. All the 338 non-specific sites were blocked with bovine serum albumin (BSA). Finally, biotinylated DNA 339 aptamers were allowed to react with streptavidin immobilized on electrode surface. The 340 fabricated aptasensor was then used to capture PSA which was measured via differential pulse 341 voltammetry (DPV). The limit of detection of the aptasensor was 0.25 ng/ml with high 342 specificity to PSA. In spite of high sensitivity and specificity, the fabrication procedure which 343 is a layer-by-layer development of sensor surface is quite complex, which may be a drawback 344 in fabricating a cost effective sensor. The group also used Electrochemical Impedance 345 Spectroscopy (EIS) to characterize the layer-by-layer fabrication of the aptasensor. 346

Electrochemical Impedance Spectroscopy is one of the most promising 347 electrochemical techniques for DNA-based approaches but requires a careful design in order 348 to optimize its signal. Particularly important for EIS biosensors is the formation of a well-349 organized self-assembled monolayer (SAM) which allows an optimal charge transfer to occur. 350 For successful EIS measurements, it is necessary to have a good and reliable SAM layer on 351 the gold electrode surface. One of the most accepted approaches to achieve this goal is by 352 alkanethiol chemistry. Alkanethiols can be easily adsorbed and form SAMs (Love et al., 353 2005) on a clean gold surface through thiol bonds (see fig. 3). It has been reported that longer 354 alkane chains give a more compact structure with minimal defects (Campuzano et al., 2006). 355 Among different configuration of SAM, a mixed SAM of 11-Mercaptoundecanoid acid 356 (MUA), HS(CH₂)₁₀COOH, and 6-Mercapto-1-hexanol (MCH), HS(CH₂)₆OH, exhibited 357 reasonable starting impedance values and improved reliability (Herne & Tarlov, 1997). In 358 order to gather or to enhance the extent of a measureable signal of the recognition event 359 occurring on the working electrode, marker molecules such as redox couples, are exploited. 360 The recognition events that happen on the SAM not only modify the charge transfer processes 361 between redox couples present in the measurement solution and the sensor surface but also 362 affect the double layer at the sensor interface. Both these events cause a change in the system 363 charge transfer resistance (R_{ct}) which can then be measured by using an appropriate 364 equivalent circuit. 365

366



Fig. 3. Schematic illustration of fabrication process of the aptasensor with 6-mercaptohexanoland thiolated DNA aptamer.

372

373

In EIS measurements using PSA aptamers, Jolly et al. (2014) and Formisano et al. 374 (2014) reported a reduction in charge transfer resistance (R_{ct}) upon binding of PSA to the 375 immobilised DNA aptamers. This decrease is contradictory to what has been reported in the 376 literature for PSA where an increase of R_{ct} has been observed (Liu et al., 2012), even though 377 these studies used EIS mainly to characterize the bio-recognition layer and not for dose 378 response determination. A reduction of R_{ct} upon aptamer-analyte interaction has also been 379 reported for a different aptasensor using a lysozyme aptamer, where the reduction in charge 380 transfer resistance upon binding of lysozyme to its specific DNA aptamer was attributed 381 mainly due to screening of charges on DNA (Rodriguez et al., 2005). The reduction of R_{ct} 382 could arise from two reasons: firstly, upon binding, PSA might screen the charges of the DNA 383 aptamer; secondly, as PSA is also a charged protein, it could be that more positive charges are 384 exposed because of the protein architecture itself. Consequently, as there is screening of 385 charges of DNA, there is a reduction on electrostatic barrier to the ferro/ferricyanide anions 386 towards the electrode surface, leading to lowering of the $R_{\rm ct}$ value of the system. 387

Earlier reports on DNA detection using DNA (Keighley et al., 2008a) and PNA probes (Keighley et al., 2008b) have demonstrated the importance of optimization of the oligonucleotide probe surface coverage in order to have efficient binding. On the same grounds, Formisano et al. (2014) investigated for the first time the importance of optimization of surface coverage by DNA aptamer for efficient binding using Quartz Crystal Microbalance

with Dissipation mode (QCM-D). The aim of this study was to optimize the conditions of an 393 EIS aptamer-based sensor for PSA detection. In fact, EIS optimisation for DNA aptamers is 394 somewhat complex due to the different characteristics that induce a signal change: namely 395 DNA density, change in charge density close to the electrode upon DNA conformational 396 changes, size and charge of the analyte, screening of DNA charges upon analyte binding. The 397 use of QCM-D provided valuable information about conditions for maximum analyte binding 398 as well as the hydration, folding and behaviour of the aptamer distribution on the electrode. 399 The system comprised a gold surface functionalized with a mixed SAM made of DNA 400 aptamer and MCH which was used as spacer molecule. The best conditions in terms of buffer 401 solution and aptamer mole fraction (concentration of aptamers/total thiols) for the binding of 402 PSA to the aptamers were obtained by comparing the data from two techniques under similar 403 conditions. With regards to the buffer conditions, the study demonstrated how the DNA 404 aptamers' behaviour exhibits a strong dependence on the environment where it interacts with 405 PSA. 406

In order to investigate an optimum surface chemistry that not only has a good 407 antifouling effect but is also simple and cost effective, a new molecule has been investigated 408 by Jolly et al. (2014) as a spacer molecule replacing MCH: a thiol terminated sulfo-betaine 409 (fig 4). It was the first report on thiol terminated sulfo-betaine application for aptamer-based 410 sensor. Thiol terminated sulfo-betaine, which has a molecular mass of 398.6 g/mole, is a 411 zwitter ion due to presence of both positive and negative charges with a flexible chain that 412 makes it a good antifouling molecule (see Fig 5). It been reported that sulfo-betaine not only 413 reduces non-specific binding but also increases the sensitivity of the sensor (Bertok et al., 414 2013). 415

416

417



Fig. 4. Structure of thiol terminated sulfo-betaine. Image adapted from Bertok et al. (2013).
421
422



47.4

Fig. 5. Schematic of fabrication of thiol terminated sulfo-betaine based PSA aptasensor. (a)
First SAM layer by co-immobilizing 11-mercatoundecanoic acid with thiol terminated
sulfo-betaine. Image adapted from Jolly et al. (2014).

428 429

A comparison study between MCH and thiol terminated sulfo-betaine thiol chemistry 430 was carried out by monitoring non-specific binding using human serum albumin (HSA) as a 431 control protein. A schematic of the fabrication protocol for surface chemistry with thiol 432 terminated sulfo-betaine is presented in fig 5. Co-immobilization of 11-mercaptoundecanoic 433 acid (MUA) and thiol terminated sulfo-betaine formed the first SAM layer on clean gold 434 electrodes. The carboxyl group of MUA was then activated with conventional EDC/NHS 435 coupling reaction. The activated carboxyl groups were then used to immobilize amine 436 terminated DNA aptamers for PSA and finally the electrodes were treated with ethanolamine 437 to deactivate all the unreacted groups. The fabricated aptasensor with thiol terminated sulfo-438 betaine surface chemistry can discriminate PSA levels down to 1 ng/ml, which falls in the 439

lower clinical cut-off range of PSA in blood. The fabricated aptasensor with thiol terminated 440 sulfo-betaine also showed a significant reduction of the non-specific binding with HSA as 441 compared to the sensor where MCH was used instead as a spacer molecule. However, it has 442 also been reported the obstacles on the optimization of the amount of DNA aptamers 443 immobilized on the surface via EDC/NHS coupling. It was assumed that the charged thiol 444 terminated sulfo-betaine has an influence on the attachment of DNA aptamer to activated 445 MUA via EDC/NHS coupling leading to difference in amounts of DNA aptamers in different 446 electrodes fabricated under similar conditions. 447

- 448
- 449
- 450
- 451

Aptasensors for other PCa biomarkers

Besides PSA, other biomarkers for PCa are currently studied and can potentially be 452 used for DNA/RNA-based detection systems. One is the prostate-specific membrane antigen 453 (PSMA), which is a type II integral membrane glycoprotein found in human serum. It is 454 overexpressed on prostate tumour cells and may play an important role in the progression of 455 PCa. It can also differentiate between BPH and PCa (Feneley et al., 2000; Ghosh and Heston, 456 2004; Madu and Lu, 2010; Pircher et al., 2011). Furthermore, by analysing the expression of 457 PSMA, two cell lines can be distinguished among PCa cells: PSMA (-) and PSMA (+) cells 458 (Ghosh and Heston, 2004). Min et al. (2010) reported on an RNA/peptide dual-aptamer-based 459 biosensor able to detect both PSMA (-) and PSMA (+) cells by using EIS. The biosensor 460 comprises of an anti-PSMA RNA aptamer (Lupold et al., 2002) which can target PSMA (+) 461 cells and a DUP-1 peptide aptamer (Zitzmann et al., 2005) specific for PSMA (-) cells. 462

Another emerging biomarker is Alpha-methylacyl-CoA Racemase (AMACR), which is a racemace type of protein found in urine and blood. Its function is to metabolize fatty acids in the human body. It is also overexpressed in PCa and can be detected with a high sensitivity and specificity with a cut off value of 10.6 ng/ml. It also has the potential to differentiate between BPH and PCa. Currently AMACR aptamers have been independently developed by Base Pair Biotechnologies, Inc. (aptamer AM310_2) and by Yang et al. (2013). However, no reports on their application to biosensing have been published so far.

- 470
- 471
- 472

Future perspectives and conclusions

Recent work on the development of PSA aptasensors has enabled the transition from 474 using antibody to aptamers as a recognition layer. Surface modification plays an important 475 role in the development of promising biosensors which would be aided with the ongoing 476 revolution in fabrication techniques. Easier fabrication would enable these biosensors to be 477 mass produced and commercially viable. The inclination towards the development of 478 aptasensors for PSA still needs further investigation for its use as an alternative to antibodies. 479 Also, the sensitivity of an aptasensor is most likely to be influenced not only by the surface 480 chemistry but also by the analytical method used for the detection of the target molecule, and 481 so far no aptasensors have yet been used in complex samples such as blood. Overall, the 482 development of aptamer based biosensor will see increasing reported literature because of its 483 ease of synthesis and the possibilities of multiple modifications; it will always be a fresh field 484 for more scope of adaptation of methodologies that will finally drive their solicitations with 485 real blood samples. For early diagnosis of PCa, detection of different biomarkers would be 486 preferred; consequently, more work is expected on development of aptamers for different 487 isoforms of PSA and other biomarkers of PCa. An ideal biosensor for PCa detection would be 488 based on a parallel sensing of different biomarkers using an array of sensors for more accurate 489 diagnosis. In addition to the need for a simple surface chemistry, the scope of biosensor in 490 future point-of-care devices will majorly depend on the integration of the format into a device 491 that will enable easy and simple sample handling and an efficient read out system with rapid 492 and accurate sample analysis of minimal blood sample volumes. 493

494 495

496 Acknowledgement(s). This work was funded by the European Commission FP7 497 Programme through the Marie Curie Initial Training Network PROSENSE (grant no. 498 317420, 2012-2016).

- 499
- 500
- 501
- 502

References

Acevedo, B., Perera, Y., Ruiz, M., Rojas, G., Benítez, J., Ayala, M., & Gavilondo, J. (2002).
Development and validation of a quantitative ELISA for the measurement of PSA
concentration. *Clinica Chimica Acta*, *317*(1), 55-63. DOI: 10.1016/S0009-8981(01)00749-5.

- Albrecht, S., Brandl, H., Steinke, M., & Freidt, T. (1994). Chemiluminescent enzyme
 immunoassay of prostate-specific antigen based on indoxyl phosphate substrate. *Clinical Chemistry*, 40(10), 1970-1971.
- 509 Andriole, G. L., Crawford, E. D., Grubb III, R. L., Buys, S. S., Chia, D., Church, T. R.,
- 510 Fouad, M. N., Gelmann, E. P., Kvale, P. A., Reding, D. J., Weissfeld, J. L., Yokochi, L. A.,
- 511 O'Brien, B., Clapp, J. D., Rathmell, J.M., Riley, T. L., Hayes, R. B., Kramer, B. S.,
- Izmirlian, G., Miller, A. B., Pinsky, P. F., Prorok, P. C., Gohagan, J. K., & Berg, C. D.
- 513 (2009). Mortality results from a randomized prostate-cancer screening trial. New England
- 514 Journal of Medicine, 360(13), 1310-1319. DOI: 10.1056/NEJMoa0810696.
- 515 Arya, S. K., & Bhansali, S. (2012). Anti-prostate specific antigen (anti-PSA) modified
- interdigitated microelectrode-based impedimetric biosensor for PSA detection. Biosensors
- 517 Journal 1, H110601. DOI: 10.4303/BJ/H110601.
- 518 Aus, G., Ahlgren, G., Bergdahl, S., & Hugosson, J. (1996). Infection after transrectal core
- biopsies of the prostate. *British Journal of Urology*, 77(6), 851-855. DOI: 10.1046/j.1464410X.1996.01014.x.
- 521 Balducci, L., Pow-Sang, J., Friedland, J., & Diaz, J. I. (1997). Prostate cancer. *Clinics in* 522 *Geriatric Medicine*, *13*(2), 283-306.
- ⁵²³ Balk, S. P., Ko, Y. J., & Bubley, G. J. (2003). Biology of prostate-specific antigen. *Journal of* ⁵²⁴ *Clinical Oncology*, *21*(2), 383-391. DOI:10.1200/JCO.2003.02.083.
- 525 Basler, J. W., & Thompson, I. M. (1998). Lest we abandon digital rectal examination as a
- screening test for prostate cancer. Journal of the National Cancer Institute, 90(23), 1761-
- 527 1763. DOI: 10.1093/jnci/90.23.1761.
- 528 Bertok, T., Klukova, L., Sediva, A., Kasák, P., Semak, V., Micusik, M., Omastova, M.,
- 529 Chovanová, L., Vlček, M., Imrich, R., Vikartovska, A., & Tkac, J. (2013). Ultrasensitive
- impedimetric lectin biosensors with efficient antifouling properties applied in glycoprofiling
- 531 of human serum samples. *Analytical Chemistry*, 85(15), 7324-7332. DOI:
 532 10.1021/ac401281t.
- 533 Besselink, G. A. J., Kooyman, R. P. H., van Os, P. J., Engbers, G. H.,, & Schasfoort, R. B.
- 534 (2004). Signal amplification on planar and gel-type sensor surfaces in surface plasmon
- resonance-based detection of prostate-specific antigen. *Analytical Biochemistry*, 333(1), 165-
- 536 173. DOI: 10.1016/j.ab.2004.05.009.
- 537 Bostwick, D. G. (1989). The pathology of early prostate cancer. CA: a Cancer Journal for
- 538 Clinicians, 39(6), 376-393. DOI: 10.3322/canjclin.39.6.376.

- 539 Campuzano, S., Pedrero, M., Montemayor, C., Fatás, E., & Pingarrón, J. M. (2006).
 540 Characterization of alkanethiol-self-assembled monolayers-modified gold electrodes by
 541 electrochemical impedance spectroscopy. *Journal of Electroanalytical Chemistry*, 586(1),
- 542 112-121. DOI: 10.1016/j.jelechem.2005.09.007.
- Cao, C., & Sim, S. J. (2007). Double-enhancement strategy: a practical approach to a femtomolar level detection of prostate specific antigen-alpha1-antichymotrypsin (PSA/ACT
 complex) for SPR immunosensing. *Journal of Mcrobiology and Biotechnology*, *17*(6), 1031-
- 546 1035.
- 547 Carter, H. B., Pearson, J. D., Metter, E. J., Brant, L. J., Chan, D. W., Andres, R., Fozard J. L.,
- ⁵⁴⁸ & Walsh, P. C. (1992). Longitudinal evaluation of prostate-specific antigen levels in men
- ⁵⁴⁹ with and without prostate disease. Journal of the American Medical Association, 267(16),
- 550 2215.-2220. DOI: 10.1001/jama.267.16.2215.
- 551 Catalona, W. J., Smith, D. S., Ratliff, T. L., Dodds, K. M., Coplen, D. E., Yuan, J. J. J.,
- 552 Petros, J. A., Andriole, G. L. (1991). Measurement of prostate-specific antigen in serum as a
- screening test for prostate cancer. New England Journal of Medicine, 324(17), 1156-1161.
- 554 DOI: 10.1056/NEJM199104253241702.
- 555 Catalona, W. J., Partin, A. W., Slawin, K. M., Brawer, M. K., Flanigan, R. C., Patel, A.,
- 556 Richie, J. P., deKernion J. B., Walsh, P. C., Scardino, P. T., Lange, P. H., Subong, E. N.,
- 557 Parson, R. E., Gasior, G. H., Loveland, K. G., Southwick, P. C. (1998). Use of the
- ⁵⁵⁸ percentage of free prostate-specific antigen to enhance differentiation of prostate cancer
- 559 from benign prostatic disease: a prospective multicenter clinical trial. Journal of the
- *American Medical Association, 279*(19), 1542-1547. DOI: 10.1001/jama.279.19.1542.
- 561 Chen, Z., Lei, Y., Chen, X., Wang, Z., & Liu, J. (2012). An aptamer based resonance light
 562 scattering assay of prostate specific antigen. *Biosensors and Bioelectronics*, 36(1), 35-40.
 563 DOI: 10.1016/j.bios.2012.03.041.
- 564 Chikkaveeraiah, B. V.; Mani, V.; Patel, V.; Gutkind, J. S.; & Rusling, J. F. (2011).
 565 Microfluidic electrochemical immunoarray for ultrasensitive detection of two cancer
 566 biomarker proteins in serum. *Biosensors and Bioelectronics*, 26(11), 4477-4483. DOI:
- 567 10.1016/j.bios.2011.05.005.
- 568 Chiriacò, M. S., Primiceri, E., Montanaro, A., de Feo, F., Leone, L., Rinaldi, R., & Maruccio,
 569 G. (2013). On-chip screening for prostate cancer: an EIS microfluidic platform for
 570 contemporary detection of free and total PSA. *Analyst*, *138*(18), 5404-5410. DOI:
 571 10.1039/c3an00911d.

- 572 Cho, E. J., Lee, J. W., & Ellington, A. D. (2009). Applications of aptamers as sensors. *Annual* 573 *Review of Analytical Chemistry*, 2, 241-264. DOI:
- 574 10.1146/annurev.anchem.1.031207.112851.
- 575 Chornokur, G., Arya, S. K., Phelan, C., Tanner, R., & Bhansali, S. (2011). Impedance-based
- 576 miniaturized biosensor for ultrasensitive and fast prostate-specific antigen detection. Journal
- of Sensors, 2011, 983752. DOI: 10.1155/2011/983752.
- 578 Chou, E., & Simons, J. W. (1997). The molecular biology of prostate cancer morbidity and
- 579 mortality: accelerated death from ejaculate poisoning? Urologic Oncology: Seminars and
- 580 Original Investigations, 3(3), 79-84. DOI: 10.1016/S1078-1439(97)00041-0.
- 581 Christensson, A.s, Björk, T., Nilsson, O., Dahlén, U., Matikainen, M. T., Cockett, A. T.,
- Abrahamsson P. A., Lilja, H. (1993). Serum prostate specific antigen complexed to alpha 1antichymotrypsin as an indicator of prostate cancer. *Journal of Urology*, *150*(1), 100.
- 584 Clark, S. L., & Remcho, V. T. (2002). Aptamers as analytical reagents. *Electrophoresis*,
- 585 23(9), 1335-1340. DOI: 10.1002/1522-2683(200205)23:9<1335::AID-ELPS1335>3.0.CO;2586 E.
- ⁵⁸⁷ D'Amico, A. V., Chen, M. H., Roehl, K. A., & Catalona, W. J. (2004). Preoperative PSA
 ⁵⁸⁸ velocity and the risk of death from prostate cancer after radical prostatectomy. *New England*⁵⁸⁹ *Journal of Medicine*, *351*(2), 125-135. DOI: 10.1056/NEJMoa032975.
- 590 D'Amico, A. V., Renshaw, A. A., Sussman, B., & Chen, M. H. (2005). Pretreatment PSA
- velocity and risk of death from prostate cancer following external beam radiation therapy.
- 592 Journal of the American Medical Association, 294(4), 440-447. DOI: 593 10.1001/jama.294.4.440.
- 594 Feneley, M. R., Jan, H., Granowska, M., Mather, S. J., Ellison, D., Glass, J., Coptcoat, M.,
- 595 Kirby, R. S., Ogden, C., Oliver, R. T. D., Badenoch, D. F., Chinegwundoh, F. I., Nargund,
- 596 V. H., Paris, A. M. I., & Britton, K. E. (2000). Imaging with prostate-specific membrane
- ⁵⁹⁷ antigen (PSMA) in prostate cancer. Prostate Cancer and Prostatic Diseases, 3(1), 47-52.
- 598 DOI: 10.1038/sj.pcan.4500390.
- 599 Feng, K., Sun, C., Kang, Y., Chen, J., Jiang, J. H., Shen, G. L., & Yu, R. Q. (2008). Label-
- free electrochemical detection of nanomolar adenosine based on target-induced aptamer
- displacement. Electrochemistry Communications, 10(4), 531-535.
- 602 DOI:10.1016/j.elecom.2008.01.024.
- 603 Fernández-Sánchez, C., McNeil, C. J., Rawson, K., & Nilsson, O. (2004). Disposable
- ⁶⁰⁴ noncompetitive immunosensor for free and total prostate-specific antigen based on

- 605 capacitance measurement. *Analytical Chemistry*, *76*(19), 5649-5656. DOI: 10.1021/ac0494937.
- 607 Formisano, N., Jolly, P., Cromhout, M., Fogel, R., Limson, J. L., & Estrela, P. (2014).
- ⁶⁰⁸ Optimisation of an electrochemical impedance spectroscopy aptasensor by exploiting quartz
- 609 crystal microbalance with dissipation signals. *Submitted for publication*.
- 610 Ghosh, A., & Heston, W. D. W. (2004). Tumor target prostate specific membrane antigen
- (PSMA) and its regulation in prostate cancer. Journal of Cellular Biochemistry, 91(3), 528-
- 612 539. DOI: 10.1002/jcb.10661.
- G13 Greenlee, R. T., Murray, T., Bolden, S., & Wingo, P. A. (2000). Cancer statistics, 2000. CA: a
- 614 Cancer Journal for Clinicians, 50(1), 7-33. DOI: 10.3322/canjclin.50.1.7.
- 615 Grossklaus, D. J., Smith Jr, J. A., Shappell, S. B., Coffey, C. S., Chang, S. S., & Cookson, M.
- S. (2002). The free/total prostate-specific antigen ratio (% fPSA) is the best predictor of
- tumor involvement in the radical prostatectomy specimen among men with an elevated PSA.
- 618 Urologic Oncology: Seminars and Original Investigations, 7(5), 195-198. DOI:
- 619 10.1016/S1078-1439(02)00190-4.
- 620 Healy, D. A., Hayes, C. J., Leonard, P., McKenna, L., & O'Kennedy, R. (2007). Biosensor
- developments: application to prostate-specific antigen detection. *Trends in Biotechnology*,
 25(3), 125-131. DOI: 10.1016/j.tibtech.2007.01.004.
- Hermann, T., & Patel, D. J. (2000). Adaptive recognition by nucleic acid aptamers. *Science*, *287*(5454), 820-825. DOI: 10.1126/science.287.5454.
- Herne, T. M., & Tarlov, M. J. (1997). Characterization of DNA probes immobilized on gold
 surfaces. *Journal of the American Chemical Society*, *119*(38), 8916-8920. DOI:
 10.1021/ja9719586.
- Hianik, T., & Wang, J. (2009). Electrochemical aptasensors-recent achievements and
 perspectives. *Electroanalysis*, 21(11), 1223-1235. DOI: 10.1002/elan.200904566.
- 630 Hoffman, R. M. (2011). Screening for prostate cancer. New England Journal of Medicine,
- 631 *365*(21), 2013-2019. DOI: 10.1056/NEJMcp1103642.
- 632 Huang, L., Reekmans, G., Saerens, D., Friedt, J. M., Frederix, F., Francis, L., Muyldermans,
- 633 S., Campitelli, A., & Van Hoof, C. (2005). Prostate-specific antigen immunosensing based
- on mixed self-assembled monolayers, camel antibodies and colloidal gold enhanced
- 635 sandwich assays. Biosensors and Bioelectronics, 21(3), 483-490.
- 636 DOI:10.1016/j.bios.2004.11.016.
- 637 Huang, Y. W., Wu, C. S., Chuang, C. K., Pang, S. T., Pan, T. M., Yang, Y. S., & Ko, F. H.
- 638 (2013). Real-time and label-free detection of the prostate-specific antigen in human serum

- by a polycrystalline silicon nanowire field-effect transistor biosensor. *Analytical Chemistry*, *85*(16), 7912-7918. DOI: 10.1021/ac401610s.
- 641 Ikebukuro, K., Kiyohara, C., & Sode, K. (2005). Novel electrochemical sensor system for
- protein using the aptamers in sandwich manner. Biosensors and Bioelectronics, 20(10),
- 643 2168-2172. DOI: 10.1016/j.bios.2004.09.002.
- 644 Iliuk, A. B., Hu, L., & Tao, W. A. (2011). Aptamer in bioanalytical applications. *Analytical*645 *Chemistry*, 83(12), 4440-4452. DOI: 10.1021/ac201057w.
- 646 Irani, J., Fournier, F., Bon, D., Gremmo, E., Dore, B., & Aubert, J. (1997). Patient tolerance
- of transrectal ultrasound-guided biopsy of the prostate. British journal of urology, 79(4),
- 648 608-610. DOI: 10.1046/j.1464-410X.1997.00120.x.
- 649 Jeong, S., Han, S. R., Lee, Y. J., & Lee, S. W. (2010). Selection of RNA aptamers specific to
- active prostate-specific antigen. *Biotechnology letters*, 32(3), 379-385. DOI:
 10.1007/s10529-009-0168-1.
- 652 Jiang, Z., Fanger, G. R., Woda, B. A., Banner, B. F., Algate, P., Dresser, K., Xu, J., & Chu, P.
- 653 G. (2003). Expression of α -methylacyl-CoA racemase (p504s) in various malignant
- neoplasms and normal tissues: a study of 761 cases. Human Pathology, 34(8), 792-796.
- 655 DOI: 10.1016/S0046-8177(03)00268-5.
- Jolly, P., Formisano, N., Tkáč, J., Kasák, P., Frost, C. G., & Estrela, P. (2014). Label-free
 impedimetric prostate cancer aptasensor with antifouling surface chemistry. *Submitted for publication*.
- Keighley, S. D., Li, P., Estrela, P., & Migliorato, P. (2008a). Optimization of DNA
 immobilization on gold electrodes for label-free detection by electrochemical impedance
 spectroscopy. *Biosensors and Bioelectronics*, 23(8), 1291-1297. DOI:
 10.1016/j.bios.2007.11.012.
- 663 Keighley, S. D., Estrela, P., Li, P., & Migliorato, P. (2008b). Optimization of label-free DNA
- detection with electrochemical impedance spectroscopy using PNA probes. Biosensors and
- 665 Bioelectronics, 24(4), 906-911. DOI: 10.1016/j.bios.2008.07.041.
- 666 Kirk, D. (1997). MRC study: when to commence treatment in advanced prostate cancer.
- 667 Prostate Cancer & Prostatic Diseases, 1(1), 11-15.
- Kufe, D. W., Pollock, R. E., Weichselbaum, R. R., Bast, R. C., & Gansler, T. S. (2003). *Holland-Frei cancer medicine*: BC Decker Hamilton, ON.
- 670 Lee, J. W., Sim, S. J., Cho, S. M., & Lee, J. (2005). Characterization of a self-assembled
- monolayer of thiol on a gold surface and the fabrication of a biosensor chip based on surface

- plasmon resonance for detecting anti-GAD antibody. Biosensors and Bioelectronics, 20(7),
- 673 1422-1427. DOI: 10.1016/j.bios.2004.04.017.
- 674 Leman, E. S., & Getzenberg, R. H. (2009). Biomarkers for prostate cancer. *Journal of*675 *Cellular Biochemistry*, 108(1), 3-9. DOI: 10.1002/jcb.22227.
- 676 Li, C., Curreli, M., Lin, H., Lei, B., Ishikawa, F. N., Datar, R., Cote, R. J., Thompson, M. E.,
- & Zhou, C. (2005). Complementary detection of prostate-specific antigen using In₂O₃
- nanowires and carbon nanotubes. Journal of the American Chemical Society, 127(36),
- 679 12484-12485. DOI: 10.1021/ja053761g.
- 680 Lilja, H., Oldbring, J., Rannevik, G., & Laurell, C. B. (1987). Seminal vesicle-secreted
- 681 proteins and their reactions during gelation and liquefaction of human semen. Journal of
- 682 *Clinical Investigation*, 80(2), 281. DOI: 10.1172/JCI113070.
- Lilja, H., Ulmert, D., & Vickers, A. J. (2008). Prostate-specific antigen and prostate cancer:
 prediction, detection and monitoring. *Nature Reviews Cancer*, 8(4), 268-278. DOI:
 10.1038/nrc2351.
- 686 Lim, Y. C., Kouzani, A. Z., & Duan, W. (2009). Aptasensors design considerations
 687 Computational intelligence and intelligent systems (pp. 118-127): Springer. DOI:
- **688** 10.1007/978-3-642-04962-0_14.
- Lin, J., & Ju, H. (2005). Electrochemical and chemiluminescent immunosensors for tumor
 markers. *Biosensors and Bioelectronics*, 20(8), 1461-1470. DOI:
 10.1016/j.bios.2004.05.008.
- 692 Liu, B., Lu, L., Hua, E., Jiang, S., & Xie, G. (2012). Detection of the human prostate-specific
- antigen using an aptasensor with gold nanoparticles encapsulated by graphitized mesoporous
 carbon. *Microchimica Acta*, 178(1-2), 163-170. DOI: 10.1007/s00604-012-0822-5.
- 695 Liu, J., Lu, C. Y., Zhou, H., Xu, J. J., Wang, Z. H., & Chen, H. Y. (2013). A dual-functional 696 electrochemical biosensor for the detection of prostate specific antigen and telomerase
- activity. *Chemical Communications*, 49(59), 6602-6604. DOI: 10.1039/C3CC43532F.
- Loeb, S., & Catalona, W. J. (2007). Prostate-specific antigen in clinical practice. *Cancer Letters*, 249(1), 30-39. DOI: 10.1016/j.canlet.2006.12.022.
- 700 Loeb, S., Vellekoop, A., Ahmed, H. U., Catto, J., Emberton, M., Nam, R., Rosario, D. J., &
- ⁷⁰¹ Lotan, Y. (2013). Systematic review of complications of prostate biopsy. *European Urology*
- 702 64(6), 876-92. DOI: 10.1016/j.eururo.2013.05.049.
- 703 Love, J. C., Estroff, L. A., Kriebel, J. K., Nuzzo, R. G., & Whitesides, G. M. (2005). Self-
- assembled monolayers of thiolates on metals as a form of nanotechnology. Chemical
- 705 *Reviews*, 105(4), 1103-1170. DOI:10.1021/cr0300789.

- ⁷⁰⁶ Lövgren, J., Valtonen-André, C., Marsal, K., Liua, H., & Lundwall, Å. (1999). Measurement
 ⁷⁰⁷ of prostate-specific antigen and human glandular kallikrein 2 in different body fluids.
 ⁷⁰⁸ *Journal of Andrology*, 20(3), 348-355. DOI: 10.1002/j.1939-4640.1999.tb02528.x.
- 709 Lupold, S. E., Hicke, B. J., Lin, Y., & Coffey, D. S. (2002). Identification and characterization
- of nuclease-stabilized RNA molecules that bind human prostate cancer cells via the prostate-
- specific membrane antigen. *Cancer Research*, 62, 4029-4033.
- Madu, C. O., & Lu, Y. (2010). Novel diagnostic biomarkers for prostate cancer. *Journal of Cancer*, *1*, 150-177. DOI: 10.7150/jca.1.150.
- 714 Mairal, T., Özalp, V. C., Lozano Sánchez, P., Mir, M., Katakis, I., & O'Sullivan, C. K.
- (2008). Aptamers: molecular tools for analytical applications. *Analytical and Bioanalytical Chemistry*, *390*(4), 989-1007. DOI: 10.1007/s00216-007-1346-4.
- 717 Mani, V., Chikkaveeraiah, B. V., Patel, V., Gutkind, J. S., & Rusling, J. F. (2009).
- ⁷¹⁸ Ultrasensitive immunosensor for cancer biomarker proteins using gold nanoparticle film ⁷¹⁹ electrodes and multienzyme-particle amplification. *ACS Nano*, *3*(3), 585-594. DOI: ⁷²⁰ 10.1021/nn800863w.
- 721 Maraldo, D., Garcia, F. U., & Mutharasan, R. (2007). Method for quantification of a prostate
- cancer biomarker in urine without sample preparation. *Analytical Chemistry*, *79*(20), 76837690. DOI: 10.1021/ac070895z.
- Melichar, B. (2012). Tumor biomarkers: PSA and beyond. *Clinical Chemistry and Laboratory Medicine*, 50(11), 1865-1869. DOI: 10.1515/cclm-2012-0631.
- 726 Mikolajczyk, S. D., Marker, K. M., Millar, L. S., Kumar, A., Saedi, M. S., Payne, J. K.,
- Evans, C. L., Gasoir, C. L., Linton, H. J., Carpenter, P., & Rittenhouse, H. G. (2001). A
- truncated precursor form of prostate-specific antigen is a more specific serum marker of
- prostate cancer. Cancer Research, 61(18), 6958-6963.
- 730 Mikolajczyk, S. D., Catalona, W. J., Evans, C. L., Linton, H. J., Millar, L. S., Marker, K. M.,
- 731 Katir, D., Amirkhan, A., & Rittenhouse, H. G. (2004). Proenzyme forms of prostate-specific
- antigen in serum improve the detection of prostate cancer. Clinical Chemistry, 50(6), 1017-
- 733 1025. DOI: 10.1373/clinchem.2003.026823.
- 734 Min, K., Song, K. M., Cho, M., Chun, Y. S., Shim, Y. B., Ku, J. K., & Ban, C. (2010).
- 735 Simultaneous electrochemical detection of both PSMA (+) and PSMA (-) prostate cancer
- r36 cells using an RNA/peptide dual-aptamer probe. Chemical Communucations, 46, 5566-
- 737 5568. DOI: 10.1039/c002524k.

- Moyer, V. A. (2012). Screening for prostate cancer: US Preventive Services Task Force 738 recommendation statement. Annals of internal medicine, 157(2), 120-134. DOI: 739 10.7326/0003-4819-157-2-201207170-00459. 740
- Numnuam, A., Chumbimuni-Torres, K. Y., Xiang, Y., Bash, R., Thavarungkul, P., 741
- Kanatharana, P., Pretsch, E., Wang, J., & Bakker, E. (2008). Aptamer-based potentiometric
- measurements of proteins using ion-selective microelectrodes. Analytical Chemistry, 80(3), 743
- 707-712. DOI: 10.1021/ac701910r. 744

- Ohori, M., Dunn, J. K., & Scardino, P. T. (1995). Is prostate-specific antigen density more 745 useful than prostate-specific antigen levels in the diagnosis of prostate-cancer? Urology, 746
- 46(5), 666-671. DOI: 10.1016/s0090-4295(99)80298-2. 747
- Okuno, J., Maehashi, K., Kerman, K., Takamura, Y., Matsumoto, K., & Tamiya, E. (2007). 748
- Label-free immunosensor for prostate-specific antigen based on single-walled carbon 749
- nanotube array-modified microelectrodes. Biosensors and Bioelectronics, 22(9-10), 2377-750 2381. DOI: 10.1016/j.bios.2006.09.038. 751
- Panini, N. V., Messina, G. A., Salinas, E., Fernández, H., & Raba, J. (2008). Integrated 752
- microfluidic systems with an immunosensor modified with carbon nanotubes for detection 753
- of prostate specific antigen (PSA) in human serum samples. Biosensors and Bioelectronics, 754
- 23(7), 1145-1151. DOI: 10.1016/j.bios.2007.11.003. 755
- Pinsky, P. F., Andriole, G., Crawford, E. D., Chia, D., Kramer, B. S., Grubb, R., Greenlee, R., 756
- & Gohagan, J. K. (2007). Prostate-specific antigen velocity and prostate cancer gleason 757 grade and stage. Cancer, 109(8), 1689-1695. DOI: 10.1002/cncr.22558. 758
- Pircher, A., Hilbe, W., Heidegger, I., Drevs, J., Tichelli, A., & Medinger, M. (2011). 759
- Biomarkers in tumor angiogenesis and anti-angiogenic therapy. International Journal of 760 Molecular Sciences, 12(10), 7077-7099. DOI: 10.3390/ijms12107077. 761
- Putzbach, W., & Ronkainen, N. J. (2013). Immobilization techniques in the fabrication of 762 nanomaterial-based electrochemical biosensors: a review. Sensors, 13(4), 4811-4840. 763 DOI:10.3390/s130404811. 764
- Rodriguez, M. C., Kawde, A. N., & Wang, J. (2005). Aptamer biosensor for label-free 765 impedance spectroscopy detection of proteins based on recognition-induced switching of the 766
- surface charge. Chemical Communications, (34), 4267-4269. DOI: 10.1039/B506571B. 767
- Ross, R. K., Pike, M. C., Coetzee, G. A., Reichardt, J. K. V., Yu, M. C., Feigelson, H., 768
- Stanczyk, F. Z., Kolonel, L. N., & Henderson, B. E. (1998). Androgen metabolism and 769
- prostate cancer: establishing a model of genetic susceptibility. Cancer Research, 58(20), 770
- 4497-4504. 771

- 772 Rubin, M. A., Zhou, M., Dhanasekaran, S. M., Varambally, S., Barrette, T. R., Sanda, M. G.,
- Pienta, K. J., Ghosh, D., & Chinnaiyan, A. M. (2002). α-methylacyl coenzyme A racemase
- as a tissue biomarker for prostate cancer. Journal of the American Medical Association,
- 775 287(13), 1662-1670. DOI: 10.1001/jama.287.13.1662.
- 776 Sardesai, N. P., Barron, J. C., & Rusling, J. F. (2011). Analytical Chemistry, 83(17), 6698-
- 777 6703. DOI: 10.1021/ac201292q.
- 778 Sarkar, P., Pal, P. S., Ghosh, D., Setford, S. J., & Tothill, I. E. (2002). Amperometric
- biosensors for detection of the prostate cancer marker (PSA). *International Journal of Pharmaceutics*, 238(1), 1-9. DOI: 10.1016/S0378-5173(02)00015-7.
- 781 Savory, N., Abe, K., Sode, K., & Ikebukuro, K. (2010). Selection of DNA aptamer against
 782 prostate specific antigen using a genetic algorithm and application to sensing. *Biosensors*
- *and Bioelectronics*, 26(4), 1386-1391. DOI: 10.1016/j.bios.2010.07.057.
- 784 Seto, Y., Iba, T., & Abe, K. (2001). Development of ultra-high sensitivity bioluminescent
 785 enzyme immunoassay for prostate-specific antigen (PSA) using firefly luciferase.
 786 Luminescence, 16(4), 285-290. DOI: 10.1002/bio.654.
- ⁷⁸⁷ Song, S., Wang, L., Li, J., Fan, C., & Zhao, J. (2008). Aptamer-based biosensors. *Trends in* ⁷⁸⁸ Analytical Chemistry, 27(2), 108-117. DOI: 10.1016/j.trac.2007.12.004.
- 789 Stamey, T. A., Yang, N., Hay, A. R., McNeal, J. E., Freiha, F. S., & Redwine, E. (1987).
- Prostate-specific antigen as a serum marker for adenocarcinoma of the prostate. *New England Journal of Medicine*, *317*(15), 909-916. DOI: 10.1056/nejm198710083171501.
- 792 Stanford, J. L., Stephenson, R. A., Coyle, L. M., Cerhan, J., Correa, R., Eley, J. W., Gilliland,
- F., Hankey, B., Kolonel, L. N., Kosary, C., Ross, R., Severson, R., & West, D. (1999).
 Prostate cancer trends 1973-1995, SEER Program, National Cancer Institute. *NIH pub* 994543.
- 796 Stephan, C., Lein, M., Jung, K., Schnorr, D., & Loening, S. A. (1997). The influence of 797 prostate volume on the ratio of free to total prostate specific antigen in serum of patients 798 with prostate carcinoma and benign prostate hyperplasia. *Cancer*, *79*(1), 104-109. DOI:
- ⁷⁹⁹ 10.1002/(SICI)1097-0142(19970101)79:1<104::AID-CNCR15>3.0.CO;2-8.
- 800 Syed, M. A., & Pervaiz, S. (2010). Advances in aptamers. Oligonucleotides, 20(5), 215-224.
- 801 DOI: 10.1089/oli.2010.0234.
- 802 Takayama, T. K., Fujikawa, K., & Davie, E. W. (1997). Characterization of the precursor of
- prostate-specific antigen Activation by trypsin and by human glandular kallikrein. Journal of
- 804 Biological Chemistry, 272(34), 21582-21588. DOI: 10.1074/jbc.272.34.21582.

- Tombelli, S., Minunni, M., & Mascini, M. (2005). Analytical applications of aptamers. 805 Biosensors and Bioelectronics, 20(12), 2424-2434. DOI: 10.1016/j.bios.2004.11.006. 806
- Uzzo, R. G., Wei, J. T., Waldbaum, R. S., Perlmutter, A. P., Byrne, J. C., & Vaughan Jr., D. 807
- (1995). The influence of prostate size on cancer detection. Urology, 46(6), 831-836. DOI: 808
- 10.1016/S0090-4295(99)80353-7. 809
- Velonas, V. M., Woo, H. H., dos Remedios, C. G., & Assinder, S. J. (2013). Current status of 810
- biomarkers for prostate cancer. International Journal of Molecular Sciences, 14(6), 11034-811
- 11060. DOI: 10.3390/ijms140611034. 812

- Wang, X., Zhou, J., Yun, W., Xiao, S., Chang, Z., He, P., & Fang, Y. (2007). Detection of 813
- thrombin using electrogenerated chemiluminescence based on Ru (bpy)₃²⁺-doped silica
- nanoparticle aptasensor via target protein-induced strand displacement. Analytica Chimica 815
- Acta, 598(2), 242-248. DOI: 10.1016/j.aca.2007.07.050. 816
- Wee, K. W., Kang, G. Y., Park, J., Kang, J. Y., Yoon, D. S., Park, J. H., & Kim, T. S. (2005). 817
- Novel electrical detection of label-free disease marker proteins using piezoresistive self-818
- sensing micro-cantilevers. Biosensors and Bioelectronics, 20(10), 1932-1938. DOI: 819 10.1016/j.bios.2004.09.023. 820
- Weeks, B. L., Camarero, J., Noy, A., Miller, A. E., De Yoreo, J. J., & Stanker, L. (2003). A 821 microcantilever-based pathogen detector. Scanning, 25(6), 297-299. DOI: 822 10.1002/sca.4950250605. 823
- Wu, C. L., Yang, X. J., Tretiakova, M., Patton, K. T., Halpern, E. F., Woda, B. A., Young, R. 824
- H., & Jiang, Z. (2004). Analysis of α-methylacyl-CoA racemase (P504S) expression in high-825
- grade prostatic intraepithelial neoplasia. Human Pathology, 35(8), 1008-1013, 8. DOI: 826
- 10.1016/j.humpath.2004.03.019. 827
- Willner, I., & Zayats, M. (2007). Electronic aptamer-based sensors. Angewandte Chemie -828 International Edition, 46(34), 6408-6418. DOI: 10.1002/anie.200604524. 829
- Xiao, Y., Lubin, A. A., Heeger, A. J., & Plaxco, K. W. (2005). Label-free electronic detection 830 of thrombin in blood serum by using an aptamer-based sensor. Angewandte Chemie, 831
- 117(34), 5592-5595. DOI: 10.1002/ange.200500989. 832
- Xu, D., Xu, D., Yu, X., Liu, Z., He, W., & Ma, Z. (2005). Label-free electrochemical 833
- detection for aptamer-based array electrodes. Analytical Chemistry, 77(16), 5107-5113. DOI: 834
- 10.1021/ac050192m. 835
- Yang, D. K., Chen, C. S., & Chen, L C. (2013). Screening of functional aptamers against 836
- alpha-methylacyl-coA recemase. In Abstracts of the 13th American Institute of Chemical 837

- 838 Engineers Annual Meeting, November 3–8, 2013. Retrieved May 9, 2014, from
 839 https://aiche.confex.com/aiche/2013/webprogram/Paper332288.html.
- 840 Zani, A.; Laschi, S.; Mascini, M.; & Marrazza, G. (2011). A new electrochemical multiplexed
- 841 assay for PSA cancer marker detection. *Electroanalysis*, 23(1), 91-99. DOI:
- 842 10.1002/elan.201000486.
- 843 Zitzmann, S., Mier, W., Schad, A., Kinscherf, R., Askozylakis, V., Krämer, S., Altmann, A.,
- Eisenhut, M., & Haberkorn, U. (2005). A new prostate carcinoma binding peptide (DUP-1)
- for tumor imaging and therapy. *Clinical Cancer Research, 11*, 139-146.
- 846