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## Aptamer based tools for environmental and therapeutic monitoring: A review of developments, applications, future perspectives

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#### ABSTRACT

Nucleic acids in the form of aptamers play a growing and significant role in the targeted and rapid analysis of environmental sample composition and medical analyses. In this paper, the review of both aptamers synthesis methods as well as application of these short chain oligonucleotides (with critical comments on their strong and weak features) are given. The first ones include: systematic evolution of ligands by exponential enrichment (SELEX), high throughput aptamer identification screen (HAPIscreen), and a non-equilibrium capillary electrophoresis of equilibrium mixture (NECEEM). Afterwards, manuscript describes variety of sensors and biotests utilizing aptamers as active part of its action starting from electrochemical aptasensors, through optical to piezo-electric ones. Described biotests present basic developments in enzymelinked apta-sorbent assays (ELASA) that can be performed with different variations (enzyme-linked aptamer assay (ELAA), enzyme-linked oligonucleotide assay (ELONA) and aptamerlinked immobilized sorbent assay (ALISA)). Next, the review presents advantages and drawbacks of recent aptameric developments in versatile laboratory applications, namely medical ones, as well as analytical and bioassays. Utilitarian development of aptasensors and aptamers would strongly benefit from an assembly of interdisciplinary teams containing chemists, physicists, biologists, medical doctors, and material and electronic scientists, to determine the most effective application methodologies.

#### **KEYWORDS**

Aptamer; aptasensors; SELEX; HAPIscreen; NECEEM; ELONA; ELASA

### 1. Introduction – current challenges in analysis of environmental and biological samples

While over the past decades, significant advances have been made in the field of chemical and biological analysis, yet, low-cost, reliable, rapid, highly precise, sensitive, specific and targetoriented (to compounds bacteria,

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viruses, proteins, antibodies and others, at low concentration levels) analyses remain a challenge for researchers. Correct analysis usually begins with proper sampling and transportation, these stages are crucial and even the best-performed analysis will be of no value, if these steps are poorly carried out. Badly collected samples will not be representative and poorly preserved or examined after a significant period will not reflect the average composition of a given studied object (Csuros 2018).

Determination of different types of analytes (simple organic and inorganic chemicals as well as macromolecular structures such as proteins or even cells) requires subjecting the sample to additional preparation stages such as purification, extraction, isolation and enrichment. At each of these stages, contamination or alteration of analyte concentration may occur (e.g. by loss of sample volume, absorption of the analyts on the walls of laboratory vessels, aggregation of the determined proteins etc.), these phenomena can be a significant obstacle in obtaining the correct result in the trace analysis.

All these stages significantly extend time of the analysis, increase costs and consumption of reagents (often very toxic to the environment). Some samples require special treatment, their composition can be variable over time as in the case of body fluids such as blood, the longer the path of the samples from the point of collection to the laboratory, the greater is the risk of changes in their properties and therefore the risk of diagnostic error (Singh, Graber, & Hofer, 2016). To face this challenges there is a great dement for creating innovative analytical devices characterized by high precision, sensitivity, specificity, speed, and usability (e.g. portable devises) for monitoring a growing number of environmental pollutants as well as for medical applications (Long, Zhu, & Shi, 2013).

In this paper authors focused on a critical evaluation on how to the use aptamers as semi-universal tools for recognizing different types of analytes and their usefulness in defeating challenges related to sample analysis.

# 2. Aptamers – their properties, mode of actions and methods of production

Aptamers are short-chain oligonucleotides of RNA or singlestranted DNA (usually in the range of 40 to 100 nucleotides) and were simultaneously described for the first time by three groups of scientists: Tuerk and Gold (1990), Robertson and Joyce (1990), and Ellington and Szostak (1990). The uniqueness of aptamers lies in their high affinity, and is a result of their flexibility and the ability to fold upon binding with a target; hence their name, derived from the Latin *aptus* – fit, attached or joined and the Greek

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Table 1. Differences between antibodies and aptamers (Nezlin, 2014; Toh et al., 2015).

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Features	Aptamers	Antibodies
Molecule size	Minimal size is 10 kDa	Minimal size is ca. 180 kDa
Production/development	Via chemical synthesis and selection resulting in less batch variation an	Laboratory animals are used, what affects the batch to batch
	shorter time of production	of production
Immobilization	Through adsorption or chemical coupling	Immobilization by biotin/streptavidin
Production conditions	Necessity to use highly purified	Immunoantigens administered to
	immmunoantigen for selection	animals do not need to be purified
Costs	Less expensive than production of	Large costs result from the use of
	antibodies, however, it is still expensive	laboratory animals, the difference in batches and their storage conditions
Possibility of storage	More stable in higher temperatures and posse a longer shelf life compared to antibodies.	Can be irreversibly denatured at room temperature
Chemical modification of aptamer/antibody	Can be manipulated chemically and labeled without losing specificity	Labeling may reduce the affinity of antigens towards targets
Toxicity/immunotoxicity	There is no immunological response of the organisms to aptamers	Possibility of inducing an immune response in the patients
Activity in different environments	Active in different reaction environments	Active only in a physiological environment

*meros* – part. (Tuerk & Gold, 1990; Robertson & Joyce, 1990; Ellington & Szostak, 1990; Ng et al., 2006; Nezlin 2014).

Aptamers act as ligands, binding strongly with the target molecule as evidenced by the nano- or pico-molar dissociation constant  $\left(K_d = \frac{[free \ aptamer] \cdot [free \ target]}{[aptamer-target \ complex]}\right)$  of the resulting target-aptamer complex, which can exhibit a greater affinity for the target than the antibodies, that were so far considered highly selective. Unlike antibodies, aptamers are obtained during chemical synthesis (completely different than in the case of antibodies that are obtained using laboratory animals); in addition, they retain their properties both during storage and in different reactionary environments. Additional information on the differences between aptamers and antibodies are summarized in Table 1 (Nezlin 2014; Toh, Citartan, Gopinath, & Tang, 2015).

Molecular recognition of target molecules by aptamers may occur 119 between different secondary and tertiary structures. For example, small 120 molecule compounds, such as amino acids, combine with aptamers using 121 pseudoknots, internal loops, and one-sided bulges for interaction (Figure 1) 122 (Antunes, Jorge, Caffarena, & Passetti, 2018). In the case of large molecules, 123 different types of reactions and physical factors are involved in the forma-124 tion of aptamer-target complexes, including hydrogen bonds, polar groups, 125 shapes, and van der Waals forces as indicated by nuclear magnetic reson-126 ance studies (Figure 1) (Song, Wang, Li, Fan, & Zhao, 2008). All of these 127 interactions result in a permanent 'fitting' of the aptamer-ligand to a given 128 biomolecule, or vice versa, if the oligonucleotide is larger than its target 129

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**Figure 1.** Three-dimensional structure of aptamer with bonded analyte e.g. a) small molecule, b) large molecule and schematic secondary structures of aptamers: c) pseudoknot, d) G-quartet, e) hairpin, f) stem-loop/bugle, g) G-quadruplex, adapted from (Nezlin, 2014; McGown, Joseph, Pitner, Vonk, & Linn, 1995; Radom et al., 2013; Deng et al., 2014; Jo & Ban, 2016). G, guanine; C, cytosine; A, adenine; T, thymine; U, uracile.

Aptamers can bind to a wide range of enzymes, regulatory proteins, growth factors, mono- and polyclonal antibodies (Hayat & Marty, 2014), lectins as well as low molecular weight vitamins (Selvakumar & Thakur, 2012), antibiotics, organic pigments, amino acids, peptides, nucleotides (Tombelli, Minunni, & Mascini, 2005), tissues and organisms (Pereira et al., 2018), cells (Chen et al., 2016) and even some metal ions (Zhang, Liu, Guo, & Lin, 2018) (Figure 1; Tereshko, Skripkin, & Patel, 2003; Radom, Jurek, Mazurek, Otlewski, & Jeleń, 2013, Long, Long, White, & Sullenger, 2008).

Large quantities of aptamers can be obtained by biochemical synthesis most frequently by Systematic Evolution of Ligands by EXponential enrichment (SELEX). Additional methods include a high throughput aptamer identification screen (HAPIscreen), and a non-equilibrium capillary electrophoresis of equilibrium mixture (NECEEM) (Dausse et al., 2011; Toh et al., 2015; Berezovski et al., 2005). Aptamers, despite their advantages and versatility, also have disadvantages that may limit their usefulness; these limitations are enumerated in the Table 2.

It is possible to integrate aptamers with various platforms and transducers that will convert the signal from aptamers bounded with analytes to fluorescent, colorimetric, or electrochemical signals that are measurable and dependent upon the sample analyte concentration. Although most aptamer development is focused on medical applications (blood or human serum measurements), there are new achievements in environmental field. Commercially produced aptamers are available from multiple suppliers (Aptagen LLC, BasePair Biotechnologies Inc., AptaMatrix Inc.) which provide services for adjusting the sensitivity and structure of produced aptamers, but for now, there are no aptamer based commercial tests or biosensors dedicated to environmental pollution (Aptagen LLC, 2019; BasePair Biotechnologies Inc., 2019; AptaMatrix Inc., 2019). Table 2.Advantages and disadvantages of aptamers application (Keefe et al., 2010; Rimmele,2003;Cheng, Chen, Lennox, Behlke, & Davidson, 2013; Mascini, 2009; Crivianu-Gaita &Thompson, 2016;Nguyen et al., 2014, Palchetti & Mascini, 2008, Amiri et al., 2017).

Advantages	Disadvantages
<ul> <li>Small size: allows penetration of the tissue passing through barriers such as the blood-brain barrier.</li> <li>Nucleic acids prefer forming secondary strur resulting in significant number of receptor conformations.</li> <li>Specificity to the target molecule, which is at the antibody level.</li> <li>Possible <i>in vitro</i> tests as well as <i>in vivo</i>.</li> <li>Lack or low immune response and no toxic</li> <li>Lower cost of production compared to competing antibodies.</li> <li>The production proceeds <i>in vitro</i>, so there i possibility of contamination with biological factors that can induce immune response.</li> <li>Aptamers are chemical synthesis without co culture or animals, which eliminates ethical concerns.</li> <li>High molecular stability (DNA is more stabl proteins) – resistance to temperatures and damaging proteins and the ability to easily renaturation, after disturbing the spatial str</li> <li>A very large variety of target molecules rar from small inorganic ions to cells.</li> <li>Are able to renature and regain analyte bir ability after denaturation (possibility of multiple usages).</li> <li>Low detection limits (even at zmol L<sup>-1</sup> leve</li> </ul>	<ul> <li>Small size: aptamers can be easily rinsed out of the body, therefore, it is necessary to use modifications (or higher concentration) in order to prolong their presence in the human body.</li> <li>Sensitive to nuclease degradation, this forces the use of phosphate backbone modifications.</li> <li>Some targets due to the lack of functional groups or other properties will not bond with aptamers.</li> <li>Bonds with target molecules are usually weaker than antibodies.</li> <li>Aptamers can be easily digested by enzymes.</li> <li>Due to the fact that it is still constantly evolving branch of science costs can be large in comparison to conventional methods.</li> <li>Sequence determination via SELEX is still slow.</li> <li>Around one-third of the target surface may be lost or unavailable after immobilization of the target molecule.</li> </ul>

#### 2.1. Methods for generating aptamer sentences

#### 2.1.1. SELEX

SELEX *in vitro* selection is the most widely used method for obtaining RNA and single strand DNA (ssDNA) aptamers with the desired physical and chemical properties. In the first step, a very large library of oligonucleotides is synthesized, consisting of randomly generated sequences and shapes of constant length surrounded by 5' and 3' ends. During the synthesis of different regions, monomer concentrations (pyridine and purine bases A, T/U, G, and C) in the reaction mixture are equimolar, so the probability of occurrence for each oligonucleotide combination is the same. In the case of n-nucleotide fragments, this gives 4<sup>n</sup> different combinations; a pool of oligonucleotides that theoretically represent every possible nucleotide sequence (Groher & Suess, 2016).

The next step is incubation of the oligonucleotide library with the target molecule under specific conditions (pH, temperature and other) followed by elution. The target molecule is usually immobilized on a membrane. Sequences that specifically bind to the molecule are not removed by the following wash and are amplified by PCR (or by reverse transcription PCR in case of an RNA library) after elution from the target molecule. Obtaining



Figure 2. Schematic representation of basic SELEX (Klug & Famulok, 1994).

ssDNA after PCR amplification is the most critical step in the SELEX procedure. One method to prevent double strand DNA (dsDNA) formation is to use biotinylated reverse primers in the amplification step. Another method is asymmetrical PCR, which is performed with an excess amount of forward primer and very small amount of reverse primer. The selection process is repeated while gradually exacerbating the reaction conditions. During every amplification cycle and rinse, the participation of oligonucleotides that specifically recognize the target molecule increases (McKeague et al., 2015; Wei & Ling-Yun, 2009; Svobodová, Pinto, Nadal, & O' Sullivan, 2012). The last step of SELEX is duplication of the selected aptamers (see Figure 2 for details).

Then the oligonucleotides are tested for their binding strength ( $K_d$ ) of the resulting complex and oligonucleotide affinities are determined; those characterized by the best performance are introduced into plasmids, cloned in bacterial cells, and sequenced. After learning the unique nucleotide sequence, multiple copies of the given aptamer are made by chemical synthesis. SELEX stages are presented schematically in Figure 2 (Klug & Famulok, 1994).

There are many variants of the SELEX method, including genomic
SELEX to produce mirror aptamers (Spiegelmers), monoLEX, *in vivo*SELEX, photo-SELEX, whole cell-SELEX, GO-SELEX, and *in silico*-SELEX.

259 This process is constantly being improved in order to shorten the process 260duration and to obtain aptamers with a very high affinity for different tar-261 gets (Toh et al., 2015; Berezovski, Musheev, Drabovich, & Krylov, 2006; 262 Wondergem, Schiessel, & Tompitak, 2017; Nguyen, Kwon, Kim, & Gu, 263 2014; Sefah, Shangguan, Xiong, O'donoghue, & Tan, 2010; Nitsche et al., 264 2007). Additional information about SELEX types are presented in Table 3. 265 Unfortunatly, the SELEX approach is still relatively slow, can only be auto-266 mated to a small degree, and is strongly biased, since efficiently amplified 267 poor affinity binders may mask low copy/high affinity aptamers. The most 268time and labor-consuming step in developing aptamers is the affinity evalu-269 ation of the isolated oligonucleotide sequences prior to choosing the 270aptamer with the highest affinity. 271

## 2.1.2. Non-SELEX methods: HAPIscreen and NECEEM

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HAPIscreen method is a non-SELEX method, which combines the target molecule and the potential aptamer with donor and acceptor microbeads. The aptamer interacts with the target molecule while simultaneously releasing a fluorescent signal produced by the microbeads, which is detected using a fluorescence reader (Dausse et al., 2011).

A precursor to HAPIscreen is the AlphaScreen® method (refer to Figure 3). The biological interaction between molecules closes the beads and further initiates a cascade of chemical reactions to obtain a greatly enhanced signal. After laser excitation, photosensitizers in the donor bead transform the surrounding oxygen into a more excited singlet state. The oxygen molecules disperse and react with a chemiluminescencer in the acceptor bead, which additionally activates the fluorophores that start to emit light at 520–620 nm wavelength. This type of aptamer screening allows for the selection of high-affinity aptamers within two hours (Dausse et al., 2011).

Another non-SELEX method is NECEEM, which consists of two major steps: incubation of the randomized DNA/RNA library with the target molecule, and separation of the formed complexes from the unbound nucleic acid ligands without the need for amplification. This is usually repeated three times until a pool of nucleic acids containing high affinity binders is obtained. Method principles rely on the assumption that when using high voltage unbounded targets, oligonucleotides will migrate at a different velocity than the target-aptamer complex It is assumed that the target molecule migrates faster than the oligonucleotide, and that the complex of the two has an averaged velocity. After  $t_0$  time this initial mixture is divided into equivalent fractions (Figure 4; Krylov 2006; Kanoatov et al., 2015).

Table 3. Variatio	ns of SELEX processes used to produce aptamers.	
Name	Basic features	References
Whole cell-SELEX	In Whole cell-SELEX process ssDNA library is incubated with the target cells (i.e. cancer cell). Bounded sequences are recovered from the cells by heating the complexes up to 95°C, followed by centrifugation and amplified by PCR. The procedure, from design of oligonucleotides to enrichment of the selected pools. Takes around 3 months.	Sefah et al., 2010
TECS-SELEX Monol FX	TECS-SELEX utilizes of SELEX to select processions of the cell surface. TECS-SELEX utilizes cell-SELEX to select processions ectopically expressed on the cell surface. Method for the identification of a hinh-affinity DNA antamer. A sincle sten with use of affinity chromatronanhy followed by subsequent physical	Ohuchi, Ohtsu, & Nakamura, 2006 Nitsche et al 2007
	menod of the reconstruction of a right formuly over apparents of providence of point and the reconstruction of the afficiency providence of according to the reconstruction of t	Victor Locard, Discharz 0
	Anows setection of aptaineds without sequence of primers. After each new setection cycle, the inplaced with primers, reverse transcribed into complementary DNA (cDNA) and amplified by PCR. The reverse DNA strand is then cleaved by alkline cleavage at a special site.	Vater, Jarosch, Buchher, ه Klussmann, 2003
Conditional SELEX	Method enables the selection of aptamers that either can bind with the target only in the presence of a regulator molecule, or in contrary they can't bind to the target in the presence of regulator molecule.	Smith & Gold, 2004
MAKAS	Magnetic-Assisted kapid Apramer Selection. Largeted protein is attached to magnetic hanoparticle, incubated with oligonucleotide library and subjected to rotation motion produced by an external magnetic field. This enables selection of high-affinity aptamers that remain bound with the	Lai & Hong, 2014
GenomicSELEX	target despite magnetic starring. Method for studying the network of nucleic acid-protein interactions within organisms. GenomicSELEX starts from Ibraries derived from genomic	Zimmermann, Bilusic, Lorenz, &
Sol–gel SELEX	DNA. FIGH-BITHING DIRGING KINAS ARE ENTICIPED TOM THE INITIAL POOL TUTOUGN MUTCIPLE TOURDS OF DIRGING OF THE KINAS TO A GIVEN INGARD. The method uses a specially fabricated advanced device utilizing microfluidics for aptamer selection. For protein immobilization, there are sol-gel	schroeder, 2010 Bae et al., 2013
	arrays with droplets on the tops of individual microheaters of such microfluidics device used. After the target is immobilized, the solution containing oligonocceptie filtrary is injected into the device and inclusted with the target. Then each agarose droplet is individually heated and	
RAPID-SELEX	KINA aptamets are separately euted, reverse transcribed to CUVA and amplitied by P.C.K. Combination of conventional SELEX and non-SELEX approaches. Cycle performed by standard SELEX protocol is followed by next SELEX round where the amplification step is not included. This method reduces time of aptamer selection in comparison with conventional SELEX while retaining the	Szeto et al., 2013
MSD-SELEX-Particle	same attinity aptamer to the target molecule. Primer is immobilize on beads through NHS-ester. Beads are mixed with an oligonucleotide library in water-in-oil droplets and then incubated with a	Zhu, Song, et al., 2014
Display selection In vivo-SFLFX	target. Bound sequences are isolated, amplified followed by identification and analysis. In the method laboratory animals are used Bandom RNA library is injected to laboratory animals (in reference study mice). Then antamers are	Cheng et al. 2013
	recovered and purified with DNase and RNase, amplify and re-injected. After 12 rounds (study data) negative selection was conducted.	
IVIAI-SELEX	we not to generate aplamets that spectifically bind to dimerent subunits or a procein. In the first stage target molecule is immonitized on magnetic bead followed by magnetic separation of the beads (5 rounds). Next, the aptamer pool from the first stage is incubated with the target, in which	uawande et al, 2017
Tissue slide- based SELEX	only one or the protein suburints is presented. DNA library is incubated with slices of cancerous tissue. After incubation period, the complex of DNA with the cancer tissue is removed and the oligonucleotide sequences are used in PCR amplification. In the second round of selection, the PCR products are incubated with normal tissue	Zhang et al., 2015
SELEX-SAGE	slices as a counter selection step. The DNA library is incubated with proteins immobilized in 96-well plate, followed by washing, elution and PCR amplification of bound sequences. Monitor DNM one contractor with accounted into account or and active councers.	Djordjevic, 2007
GO-SELEX	Aniphined DAYs are sequenced using massively planets injuerinoecure sequencing. DNA library is pre-incubated with the target, then added to the graphene oxide (GO), which adsorb unbounded ssDNA while bound DNA remains in the drafting DNM is control from the transmission do not by prophere oxide (GO), which adsorb unbounded ssDNA while bound DNA remains in	Nguyen et al., 2014
In silico-SELEX	the solutionary is separated from the target and amplitied by refut averal system spread are subjected to selection, doming and sequencing. Computer modeling using special software, to predict the outcome of the experiment under given conditions.	Wondergem et al., 2017



**Figure 3.** Diagram of AlphaScreen® test, components A and B represent the target molecules and the oligonucleotide (AlphaScreen®, 2018).



Figure 4. Diagram of the process of non-equilibrium capillary electrophoresis of equilibrium mixture (NASEEM) for receiving aptamers (Toh et al., 2015).

#### 2.2. DNA and RNA libraries and their modifications

The vast amount and variety of nucleic acids make it possible to choose an aptamer for virtually any target. Calculations have shown that the optimal number of different sequences tested ranges from  $10^{13}$  to  $10^{15}$ . Both DNA and RNA can be used as aptamer bases; RNA enables intracellular expression, while DNA shows greater stability (Radom et al., 2013).

DNA and RNA libraries may contain sequences that surround the targeted aptamers to form a specific secondary structure, or include modified nucleotides that can greatly broaden the range of possible sequences. This may also affect aptamer properties, such as *in vivo* stability or resistance to RNAse and DNAse. Oligonucleotides can be easily modified by the substitution of one of the ribose hydroxyl groups with a methyl or an alkyl group 388 (Cummins et al., 1995). Alkylation of nucleic acids creates aptamers con-389 taining 2'-O-methylpyrimidine; such modified aptamers are more stable 390 and more resistant to the attacks of exonuclease enzymes; similar action 391 was observed with substitution of the ribose 2'-OH group with an amino 392 group or fluorine atoms, which increased resistance to degradation by 393 blood serum enzymes (Scaggiante et al., 2013; Nawrot & Sipa, 2006). 394 Dimerization of selected aptamers allows for a fold-increase in affinity of 395 the ligand to the molecule, and an increase in the durability of the complex 396 formed that enables an increased number of binding sites. Therefore, it is 397 possible to interact with more than one target molecule (Hasegawa, Taira, 398 Sode, & Ikebukuro, 2008). Modification of phosphate residues by replacing 399 them with sulfur atoms increases resistance to 3'-exonuclease digestion 400 (Kaur, Rob, Caton-Williams, & Huang, 2013). Oligonucleotides can also be 401 modified to show specificity to one of the enantiomers. For example, ribo-402 nucleases can recognize and hydrolyze only oligoribonucleotides based on a 403 series of L-riboses (so-called mirror aptamers) (Klussmann, Nolte, Bald, 404 Erdmann, & Fürste, 1996). Additionally, aptamers can be combined with 405 fluorescent dyes, the so-called 'bridging' molecules that enable the aptamer 406 to be labeled without losing its affinity, and preserving the fluorescent 407 properties of the dye. Labeling in the vicinity of the intermolecular recogni-408 tion region enables researchers to follow the conformational changes of the 409 aptamer; after binding to the target molecule, the aptamer itself acts as a 410 biosensor (Babendure, Adams, & Tsien, 2003). 411

# 3. Aptamer-based sensors and aptamer-based tests in environmental research and monitoring

#### 3.1. Aptamer-based sensors (aptasensors)

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Biosensor technology has developed quickly since the days of the first enzyme electrode for glucose detection described by Clark and Lyons in the 1960s; in particular, those containing aptamers suitable for binding very different targets, from micromolecules such as metals to macromolecules such as proteins, various test configurations have been designed and reported (Clark & Lyons, 1962). The combination of biological and electronic elements allows for quick, sensitive, and precise detection of not only small amounts of chemical compounds, but also dangerous microorganisms or toxins. The majority of these designs fall into either single-site or dual-site binding categories (Figure 5; Song et al., 2008).

Dual-site binding tests ('sandwich' tests) are one of the most commonly used formats, in which the analyte is surrounded by a pair of aptamers (Figure 5b). One is the capture probe, usually immobilized on the surface of solid supports (electrodes, glass chips, nanoparticles, or microparticles),



**Figure 5.** Single-site binding a) or dual-site binding b) aptamer-based types of test (Song et al., 2008).

and the other is a reporter probe, often conjugated to signal residues (fluorophores, enzymes, or nanoparticles) (Song et al., 2008). Measurement of biological phenomena occurring in the sensor can take place according to the transduction principle means of electrochemical (potentiometric, amperometric, and conductometric), optical (optical fiber and surface plasmon resonance biosensors, colorimetric, and fluorescent) and piezoelectric (mass sensitive including quartz crystal microbalance) detectors (Justino, Duarte, & Rocha-Santos, 2017).

#### 3.1.1. Electrochemical aptasensors

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Changes in the electrical field occur during redox reactions, and are measured by a transducer electrode. Electrochemical techniques are simple to use, fast, cost-effective, and are characterized by high selectivity and sensitivity. Most of them use label-free electrochemical impedance spectroscopy (EIS) or semiconductor field-effect sensors (FET), which show great promise, as they are not destructive. EIS is a frequency domain measurement made by applying a sinusoidal perturbation, often a voltage, to a system and in trasystors, in FET sensors, the interfacial potential between the sample and the gate is material depending on the target activity of the analyte measured (Kaisti, 2017; Muñoz, Montes, & Baeza, 2017). This is highly attractive for the detection of low-mass molecules and ions. In EIS, the change in electron transfer resistance is measured upon the aptamer binding to the target. The aptamer-target complex creates an isolating barrier on the electrode surface, and using FET sensors, the conductivity between two electrodes is gated by the change in surface potential that is induced by aptamer-target binding (Miranda-Castro, de-los-Santos-Alvarez, & Lobo-Castañón, 2017).

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474 Most aptasensor electrode surfaces are coated with microwires formed of platinum (PtNPs), silver (AgNPs) and gold (AuNPs) nanoparticles or dot-475 476 ted carbon nanoparticles (GO, Graphene Oxide), on which aptamers are 477 placed using various immobilization strategies and/or detection formats (Havat & Marty, 2014; Zhang, Liu, et al., 2018). Flexible aptamers, which 478 479 form three-dimensional structures upon binding with the target molecule, 480 are immobilized on the electrode surface through chemisorption of thiol 481 groups on Pt, Ag, Au, or C surfaces. This is followed by the creation of a 482 self-assembled monolayer by the attachment of an amine-terminated 483 aptamer to a thiol group, most often via the 3' end, as this configuration 484 seems to be more resistant to nuclease (Hayat & Marty, 2014; Yang, Yang, 485 Pang, Vara, & Xia, 2015; Arroyo-Currás et al., 2017; Liu, Luo, et al., 2017). 486 The aptamer may also be conjugated by covalent bonds to a chemically 487 modified surface, most often hydroxyl, amino, or carboxylic acids, and less 488 often with silicates or silicones (Zhu, Suter, White, & Fan, 2006; Actis 489 et al., 2011; Taghdisi, Danesh, Emrani, Ramezani, & Abnous, 2015). 490 Electroactive probes such as methylene blue (MB), ferrocene (Fc), 491  $Fe(CN)_6^{4/3}$  polymers containing ferrocene, and ruthenium complexes are 492 more often used for the transduction of biological information into electro-493 active signals (Song, Lee, & Ban, 2012, Arroyo-Currás et al., 2017). Most of 494 these methods have great potential however, there are, some limitations to 495 eletrochemine aptasensors that use nanomaterials that arise from ntrinsic 496 properties that may result in creating inefficient aptasensors (Jalalian, 497 Karimabadi, Ramezani, Abnous, & Taghdisi, 2018). 498

MB and Fc are used both in labeled and non-labeled aptasensors (Figure 6a and b). In labeled aptasensors, the electrochemical aptasensor target induction causes the aptamer to stiffen into its three-dimensional structure, which interrupts electron exchange between the MB and the electrode surface, resulting in a decreased signal proportional to the amount of analyte. In the label-free version, an aptamer structure contains many MB probes that allow the flow of electrons. Changes in aptamer conformation upon target binding result in the release of all MB probes affecting the signal (Cao et al., 2017; Tao, Zhong-Yuan, Lian-Zhe, & Guo-Bao, 2011). In 'sandwich' type aptasensors, the target first binds to the aptamer immobilized on the electrode surface, and then the second aptamer binds to the target. Therefore, this type of test is useful for detecting molecules with two or more binding sites, such as proteins (Walter et al., 2012; Figure 6d).

Electrochemical aptasensors are already used to detect impurities in environmental samples from low molecular weight heavy metals to endocrine disrupting chemicals (EDC), antibiotics, and pesticides. Methods for utilization are constantly improved, as rapid and targeted determination of small molecule compounds such as heavy metals in environmental samples



**Figure 6.** Labeled a) and label-free b) signal-off aptasensors.  $Fe(CN)_6^{4-/3-}$  probe c) is prevented from reaching the electron surface by stiffened structure of the aptamer bonded with the target molecule. 'Sandwich' aptasensor d) – two aptamer bonded with one target molecule with more than one active place (Hayat & Marty, 2014, Mishra et al., 2018).

are very important for public health reasons. Intensive research is ongoing to develop very sensitive and selective aptamer biosensors to detect heavy metals at very low concentration levels.

Electrochemical aptasensors are used for the detection of heavy metals such as  $Hg^{2+}$ ,  $Pb^{2+}$ ,  $Cd^{2+}$ , and  $As^{3+}$ . In one approach, an ultra-sensitive and highly selective electrochemical label-free aptasensor was proposed for the quantitative detection of  $Hg^{2+}$  in zeptomolar (zmol L<sup>-1</sup>) levels. In a customized sensor, Amiri, Navaee, Salimi, and Ahmadi (2017) substituted thiol groups in single-stranded DNA, which then self-assembled on a gold electrode surface through the S-Au interaction. Hybridization of ssDNA with complementary DNA (cDNA), and the consequences of dehybridization in the presence of  $Hg^{2+}$  were followed through differential pulse voltammetry responses using a redox probe ( $[Fe(CN)_6]^{3-/4-}$ ). An essential and desirable phenomenon for the detection of  $Hg^{2+}$  in environmental samples is the formation of a thymine– $Hg^{2+}$ –thymine complex (Amiri et al., 2017; Mishra, Sharma, & Mishra, 2018).

In another aptameric study, Pb<sup>2+</sup> ions were detected with sensors devel-oped on the base of a carboxylic acid group in functionalized multi-walled carbon nanotubes and directly electrodeposited gold nanoparticles (GNPs). The DNA capture probe was self-assembled onto the surface of the modi-fied electrode for hybridization with the guanine-rich aptamer probe to form a DNA double helix structure.  $Pb^{2+}$  ions added to the sample caused the DNA duplex to unwind, and formed a stabilized G-quadruplex due to the  $Pb^{2+}$ -induced G-rich DNA conformation. The aptasensor was able to detect lead ions at a 4.3 pmol  $L^{-1}$  detection limit (Zhu, Zeng, et al., 2014). 

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Furthermore, an ultrasensitive electrochemical aptasensor based on a glass carbon electrode modified with AuNPs, chitosan, and aptamers has been constructed to detect cadmium (II) ions. The presence of Cd<sup>2+</sup> triggers a change in aptamer conformation, causing the Cd/Cd-aptamer complex to adsorb more electrochemical signal indicator  $[Ru(NH_3)_6]^{3+}$  than poly-(diallyl dimethyl ammonium chloride) (PDDA), which is applied to neutralize the Cd-aptamer via electrostatic interaction. Using differential pulse voltammetry (DPV) detection, the peak current increases along with the increase in Cd<sup>2+</sup> concentration with a detection limit as low as  $0.05 \text{ pmol } \text{L}^{-1}$  (Liu, Lai, et al., 2017).

In another study, As<sup>3+</sup> ions were detected using new thermally stable polyarylidene (azomethine-ether)s and copolyarylidene (azomethine-ether)s (PAAP) hybrids based on diarylidenecycloalkanone synthesis using a solution polycondensation method. A thin layer of PAAP was coated onto a flat glassy carbon electrode (GCE) using a conducting nation (5%, CAS no. 66796-30-3) coating agent to fabricate a sensitive and selective  $As^{3+}$  ion aptamer with a short response time in a neutral buffer system. The detection limit was calculated at  $6.8 \pm 0.1$  nmol L<sup>-1</sup> (Rahman, Hussein, Aly, & Asiri, 2018). Tang, Wang, Yu, Zhang, and He (2018) presented an aptasensor containing a multi-channel one-time screen-operated carbon electrode (SPCE) using open circuit technology (OCP) for the simultaneous detection of three heavy metals,  $Hg^{2+}$ ,  $Cd^{2+}$  and  $As^{3+}$ , with detection limits of 2.0, 0.62, and 0.17 pmol  $L^{-1}$ , respectively. More detailed review on possibility of using aptamers and nanomaterials can be found in work of Farzin, Shamsipur, and Sheibani (2017).

585 In addition to heavy metal identification, aptasensors can be used to 586 identify pesticides, antibiotics, and even bisphenols. Fei et al. (2015) proposed a low-cost, selective, and sensitive electrochemical aptasensor using the direct impedimetric method to detect the pesticide acetamiprid in environmental samples. AuNPs with multiwalled carbon nanotube-reduced GO-nanoribbon composites were synthesized in a one-pot reaction (Au/ MWCNT-rGONR). This unique composite was used as a platform to immobilize aptamers. This aptasensor was successfully used to determine pesticides of interest even when the sample contained interfering substances with a detection limit of 0.17 fmol  $L^{-1}$  (Fei et al., 2015).

595 Another label-free electrochemical aptasensor was constructed by Zhan 596 et al. (2015) to detect tetracycline in dietary, environmental, and biological 597 samples in a cost-efficient, sensitive, and selective manner. A suitable 598 aptamer was immobilized on a composite consisting of reduced GO, mag-599 netite (Fe<sub>3</sub>O<sub>4</sub>), and sodium alginate on the surface of a screen-printed car-600 bon electrode (SPCE). Tetracycline could be quantified using DPV with a 601 detection limit of 0.6 nmol  $L^{-1}$  (Zhan et al., 2016). 602

Similarly Cui et al. (2015) presented a label-free, single-step method for the detection of bisphenol A (BPA) in aqueous samples. An aptamer based molecular probe was immobilized on a commercially available array of interdigitated aluminum microelectrodes. To quantify the amount of BPA in the sample, the interfacial capacitance rate change caused by the specific binding between the analyte and the immobilized aptamer was measured. The sensor reached a detection limit as low as 10 fmol  $L^{-1}$ , with a 20 s response time. This inexpensive, highly sensitive, and rapid method is promising technology for the on-site detection of BPA in food and water samples (Cui et al., 2015).

Electrochemical aptasensors have also been used in the detection of pathogens such as *Salmonella*. A biosensor based on a glassy carbon electrode modified with GO and AuNPs was combined with specific aptamers. In order to quantify the content of *Salmonella* sp., the electrochemical impedance spectrum was measured. The more *Salmonella* was added to the test solution, the more the current between the electrode and the electrolyte was reduced, which enabled a detection limit of 3 colony forming units (CFU) mL<sup>-1</sup>. This aptasensor is useful for detecting *Salmonella* species in water and food samples (Ma et al., 2014).

The use of eletrochermical aptasensors also can involve problems, unfortunately, these sensors are most often tetsted on spiked samples and there are few studies that would be carried out on real samples. The electrochemical methods themselves are sensitive to electromagnetic interference and there is a need for buffers to maintain the aptamer working environment (Zhang, Liu, et al., 2018).

## 3.1.2. Optical aptasensors (fluorescence and colorimetric sensors)

Fluorescence-based aptasensors are a type of optical sensor, which require a fluorophore (a fluorescent chemical compound) and a quencher (quenches fluorescence). These types of sensors arecurrently used with great success in indentification and quantification of environmental pollutants such as heavy metals ions (García-Gutiérrez, Huerta-Aguilar, Thangarasu, & Vázquez-Ramos, 2017).

In aptasensors the fluorophore part is attached to the other. In some cases, a surface made of a GO-sheet can act as quencher and adsorb the fluorophore-labeled aptamer in its unbound state. In other cases, the quencher is attached to a complementary DNA sequence (cDNA) which is unbound from the aptamer when the target molecule appears in the solution (Feng, Dai, & Wang, 2014). These types of aptasensors can be divided into signal-on and signal-off categories (Figure 7a-c), which increase or decrease the fluorescence signal upon binding with the target molecule. In signal-on sensors, the initial conformation of the aptamer maintains the



**Figure 7.** Schematic presentation of fluorescent aptasensor: signal-on presented in a), signal-off presented in b) and c) of fluorescent aptasensors. d) Binding targets by aptamers immobilized on AuNPs initiates their aggregation resulting in color change from red to blue (Song et al., 2012, Feng et al., 2014).

fluorophore and quencher very close to each other and results in a low fluorescence signal. Off and On fluorescent-based sensors had been for some time used with success in detection environmental pollution (García-Gutiérrez et al., 2017). Binding to the target molecule causes a change in the aptamer conformation and separates the fluorophore from the quencher, resulting in a higher fluorescence signal. In signal-off sensors, the coupling of fluorophore and quencher occurs when the target molecule appears in the solution. This decreases the fluorescence signal (correlated to the concentration of the target), which otherwise remains strong when the elements are separated (Akki & Werth, 2018).

Colorimetric tests are easy to perform, and their results can be measured and determined with uncomplicated equipment and in some cases, by the naked eye (Lu et al., 2013). Most colorimetric aptamer based tests are based on the principle that the visible color of AuNP suspensions change depending on the dispersion and aggregation in the solution. For example, if aptamers are immobilized on AuNPs stabilized in highly concentrated salts (e.g. NaCl), they are dispersed in the solution and prevent nanoparticle aggregation. This would turn the sample solution red. As soon as an aptamer recognized analyte appears in the solution, the aptamer is separated from the AuNPs, which then aggregate and cause the solution to turn blue (Figure 7d). Change and color intensity provide qualitative and 689quantitative information about the presence and concentration sample ana-690lyte, and the blue shift range is quantified as the ratio of absorbance from691 $620 \text{ nm to } 520 \text{ nm } (\sim A_{620}/A_{520} \text{ nm}) (Akki & Werth, 2018).$ 

692 The literature also describes colorimetric tests that change from blue to 693 red after the analyte is introduced. For example, aptamer thiol groups are 694 covalently bound to AuNPs and the secondary sequence of aptamer-AuNPs 695 cause the aggregation of these complexes and turn the solution blue. 696 Introduction of the target to the solution causes the molecules to disperse, 697 which manifests as a color change to red. In some cases, a reactive chromo-698 genic agent like tetramethylbenzidine (TMB) is used in conjunction with 699  $H_2O_2$ . TMB reacts on the surface of AuNPs, which serve as a catalyst to 700 promote peroxidase-like activity. When the target molecule binds with 701 aptamers, the AuNPs surface is released and can catalyze the oxidation of 702 colorless TMB to form TMB-diimine, which changes the color from red to 703 purplish-blue (Yuan, Zhao, Wang, & Quan, 2017; van den Kieboom et al., 704 2015). As with fluorescence-based aptasensors, colorimetric assays are suc-705 cessfully applied to detect analytes in tap and lake water, milk, soil, and 706 wastewater samples. The only potential analytical obstacle to using tests 707 containing AuNP particles as carriers is the sample matrix. This is why 708 environmental samples are usually diluted in special buffers, and only those 709 with a high salt content may be a problem (van den Kieboom et al., 2015). 710 Additionaly, despite continuous improvement, there are still many difficul-711 ties in the functioning of these aptasensors such as pH, ionic strength, and 712 temperature, viscosity and other external conditions. For some sensors 713 there is a need for using different types of buffers that can affect the quality 714 of the determination (Robati et al., 2016).

715 Small molecules such as heavy metal ions can be determined by optical 716 sensors; for example, Qian, Shan, Chai, Chen, and Feng (2015) developed a 717 fluorescent signal-on nanosensor for Pb<sup>2+</sup> based on biocompatible gra-718 phene QDs (quantum dots), GO and an aptamer in G-quadruplex forma-719 tion with a detection limit of  $0.6 \text{ nmol } \text{L}^{-1}$ . Another simple and reliable 720 detector for As<sup>3+</sup> in drinking water and blood serum was introduced by 721 Taghdisi, Danesh, Ramezani, Emrani, and Abnous (2018). The presented 722 sensor was the switch-on type. Upon introduction of As<sup>3+</sup> to the solution, 723 the label-free aptamer was immobilized on silica nanoparticles coated with 724 a streptavidin surface and formed a hairpin structure, which strengthened 725 the fluorescence signal with a very low detection limit of  $0.45 \text{ nmol } \text{L}^{-1}$ 726 (Taghdisi et al., 2018).Lin, Yu, Li, Cao, and Guo (2016) and Yang, Qian, 727 et al. (2015) presented two different fluorescent sensors to determine the 728 presence of acetamiprid (Lin et al., 2016; Yang, Qian, et al., 2015). Lin 729 et al. (2016) used an aptamer-modified ZnS:Mn probe constructed by com-730 bining ZnS:Mn QDs with an acetamiprid binding aptamer. Without the 731

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732 presence of acetamiprid, the signal is switched off by means of multi-walled 733 carbon nanotubes (MWCNTs) based on the fluorescence resonance energy transfer (FRET) between the ZnS:Mn-aptamer complex and the MWCNTs. 734 735 FRET is a mechanism describing energy transfer between two light-sensitive molecules called chromophores (or fluorophores). Donor chromo-736 737 phore, when in its electronic excited state, may transfer energy to an 738 acceptor chromophore through nonradiative dipole-dipole coupling in 739 short range < 10 nm (Zal & Gascoigne, 2004).

740 Introduction of acetamiprid to the sample switches on the fluorescence 741 signal by the specific combination of the aptamer and the target molecule with a detection limit of  $0.7 \text{ nmol L}^{-1}$ . This was satisfactorily used to detect 742 pesticides in environmental samples (Lin et al., 2016). Additionally, Yang, 743 744 Qian, et al. (2015) used hemin-functionalized reduced graphene oxide 745 (hemin-rGO) composites, which adsorb the aptamer and hemin that catalyze 3,3,5,5-tetramethylbenzidine (TMB) in the presence of H<sub>2</sub>O<sub>2</sub> to 746 747 identify acetamiprid. At a suitable salt concentration, the proposed 748 colorimetric aptasensor reached a detection limit of 40 nmol  $L^{-1}$  (Yang, 749 Qian, et al., 2015). 750

FRET is also used to determine the presence of  $17\beta$ -estradiol. An aptamer duplex consisting of a carboxyfluorescein (FAM)-labeled E2 and complementary DNA partially modified with Black Hole Quencher 1 (BHQ1) was used. After the introduction of  $17\beta$ -estradiol into the sample to compete for binding sites with cDNA, cDNA released from the aptamer/DNA duplexes FAM fluorescence was recovered, and the fluorescent signal was switched on. This sensor has been successfully applied to detect  $17\beta$ -estradiol in environmental, biological, and dietary samples with a 0.35 nmol  $L^{-1}$  detection limit (Zhang, Li, Zhang, & Chen, 2018). Samples tested with optical methods must maintain certain standards, they must be transparent and, if possible, colorless; additionally, the cost and durability of fluorescent markers must be taken into account.

## 3.1.3. Piezoelectric (mass-dependent) aptasensors

Piezoelectric aptasensors are less common than the previously mentioned types, however, they include sensors for detecting viruses, bacteria, and toxins. Mass-dependent sensors are label-free, have low noise and high sensitivity, and include acoustic wave-based sensors such as quartz crystal microbalance (QCM) and surface acoustic wave (SAW) sensors (Figure 8a), micromechanical cantilever-based sensors (Figure 8b), and surface-plasma resonance (SPR) sensors (Figure 8c) (van den Kieboom et al., 2015). The piezoelectric effect is based on reversible electro-mechanical interactions with substances that have crystalline materials and possesses an asymmetrical structure. The principle is that mechanical deformation causes voltage



Figure 8. Piezoelectric aptamer based-sensor with wave-detection (QCM and SAW) a) microcantilever sensor b) and with SPR detection c) (Song et al., 2008).

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tension on the material, and vice versa. The relationship between the two is always linear (Aksel & Jones, 2010; Song et al., 2008). Other piezoelectric aptasensors use SAWs. Such devices are constructed from piezoelectric materials on which electrodes made from inert metals or alloys are arranged in the shape of a comb. Applying voltage to the electrode generates a wave that propagates through the piezoelectric material, and this wave can be disrupted by an analyte bound with an aptamer (Mujahid & Dickert, 2017). The operational principle of cantilevered aptasensors is completely different from those described above. In cantilevered sensors, the aptamer is placed on the surface of a microcantilever, which deflects when the aptamer is weighted by the bound target molecule. Such deflection can be measured optically or electronically (Zhao, Jia, Wen, & Yu, 2017; Lim, Kouzani, & Duan, 2010). SPR devices are capable of registering mass changes by the associated change in the refractive index at the surface bound to the aptamer (Figure 8c). In this case, the sensor measures the resonance angle changes that occur on the molecular surface that are altered when the aptamer binds to the target molecule (Guo, 2012).

808 Wave based-aptasensors were used by Ozalp, Bayramoglu, Erdem, and 809 Arica (2015) to detect the presence of pathogenic Escherichia in food and 810 water samples at a 100 CFU mL<sup>-1</sup> detection limit. The sensor was based 811 on a QCM with an aptamer-based magnetic separation system for the rapid 812 enrichment of target pathogens for on-line monitoring (Ozalp et al., 2015). 813 More examples utilizing aptasensors for the detection of environmental 814 pollutants, as well as additional information are presented in Table 4. 815 Unfortunately, piezometric methods are characterized by low sensitivity 816 and high detection limits in relation to other methods described. 817

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Table 4. Lis	it of aptasenso	rs used fo	r different types of environmental pollution	ı analytical purp	ooses.		
Type	Analytes	Aptamer based on	Analytical method/transducer	LOD	Linear concentration range	Type of samples	References
Electrochemical : Polychlorinated	sensors PCB 77	DNA	DNA-aptamer-modified Au electrode modified with	0.01 µg L <sup>_1</sup>	0.2–200 µg L <sup>–1</sup>	Environmental samples	Wu et al., 2016
	C.4 <sup>2+</sup>	DNA	Generative TC RGO-AUNP-Ap/Au Electrode Annibe/CC and GCE	0.1 pg L <sup>-1</sup>	0.001–10,000 nmol L <sup>-1</sup>	Tap water	Wu, Lu, Fu, Wu, & Liu, 2017
Heavy metal	cu As <sup>3+</sup>	DNA	SAM on SPCE via Au-5 bond	0.15 nmol L <sup>-1</sup>	0.2-100 nmol L <sup>-1</sup>	uap water Water samples	ciu, cai, et al., 2017 Cui, Wu, & Ju, 2016
Heavy metal	Hg <sup>2+</sup>	DNA	DNA layer absorbent construct from nanocomposite of	0.035 nmol L <sup>-1</sup>	0.1–100 nmol L <sup>–1</sup>	Aquarius solutions	Yang, Kang, et al., 2015
		DNA	Thymine-Uniteritymine coordination chemistry and use of MDC for circual amolification chemistry and use	0.0036 nmol L <sup>-1</sup>	0.01–5000 nmol L <sup>-1</sup>	River and tap water, Isochate	Zeng et al., 2017
		DNA	Label-free based on the hybridization/dehybridization	0.6 zmol L <sup>-1</sup>	5–55,000 zmol L <sup>–1</sup>	Aqueous solutions	Amiri et al., 2017
EDCs	BPA	DNA	or gourde-stranged DNA on the Au electrode Nontarget-induced bridge assembly and aptamer extension reaction triggered by terminal	15 pmol L <sup>-1</sup>	0.08–15 nmol L <sup>-1</sup>	Tap water and food samples	Abnous, Danesh, Ramezani, Alibolandi, &
		DNA	deoxynucleotidyl transferase Photoelectrochemical AuNPs/ZnO	0.5 nmol L <sup>-1</sup>	1–1000 nmol L <sup>–1</sup>	Drinking water	Taghdisi, 2018 Qiao et al., 2016
		DNA	CNTs/G powders are significantly impeded by their poor 3D conductivity	0.3 nmol L <sup>-1</sup>	0.010–1000 μmol L <sup>-1</sup>	Lake water	Wang, Wang, et al., 2015
Antibiotic	Tetracyclines	DNA	Reduced GO, magnetite (Fe <sub>3</sub> O <sub>4</sub> ) and sodium alginate used to modify the surface of a screen-printed GCF	0.6 nmol L <sup>-1</sup>	0.001–5 µmol L <sup>-1</sup>	Water, food samples	Zhan et al., 2016
Antibiotic	Ofloxacin	DNA	AuNPs on the surface of GCE through sulfhydryl- terminated monolayer in which thiolated ssDNA morbes are immobilized	1 nmo L <sup>-1</sup>	50–20,000 nmol L <sup>–1</sup>	Tap and waste water	Pilehvar et al., 2017.
EDCs/hormone	Progesterone	DNA	SAM/Au/[Fe(CN) <sub>6</sub> ] <sup>3–/4–</sup> /EIS	900 ng L <sup>-1</sup>	10-60 µg L <sup>-1</sup>	Tap water, wastewater,	Contreras Jiménez et al., 2015
EDCs/hormone	17β-estradiol	DNA	Ni <sup>3+</sup> , Fe(CN) <sub>6</sub> l <sup>3-</sup> /AuNP CoS/AuNPs	0.8 pmol L <sup>-1</sup> 0.7 pmol L <sup>-1</sup>	1–60 pmol L <sup>–1</sup> 1–100 pmol L <sup>–1</sup>	yroundwater Wastewater samples Water samples	Fan, Zhao, Shi, & Liu, 2015 Huang, Liu, Zhang, Cao, & Liu, 2015
Pesticide	Carbendazim	DNA	SAM thiol-modified aptamer on AuNPs electrodes	8.2 ng L <sup>-1</sup>	0.01–10 µg L <sup>–1</sup>	Environmental and food samples	Eissa & Zourob, 2017
Pesticide	Acetamiprid	DNA	PtNPs were deposited in a bridge-like arrangement, in	1 pmol L <sup>-1</sup>	0.10-1 µmol L <sup>-1</sup>	Water samples	Madianos, Tsekenis, Skotadis,
Toxin	Auazine Cylindrosper- mopsin	DNA	TH-G nanocomposite through the cross-linker GA	юршы с 0.117 µg L <sup>-1</sup>	0.39–78 µg L <sup>–1</sup>	Lake water	Zhao, Chen, Ma, Liu, & Wang, 2015
	-	DNA	Label-free impedimetric aptasensor (fluorescence assay antamer on the Au curface)	100 pmol $L^{-1}$	0.1-80 nmol L <sup>-1</sup>	Fresh water	Elshafey, Siaj, & Zourob, 2014
Toxin	Anatoxin-a	DNA	SAM formed on a Au electrode using the disulfide	$0.5 \text{ nmol L}^{-1}$	1–100 nmol L <sup>–1.</sup>	Drinking water	Elshafey, Siaj, & Zourob, 2015
Toxin Plasticizer	Microcystin-LR Ethanolamine	DNA	GCE SDD-Co(II) AgNPs	0.37 nmol L <sup>-1</sup> 0.08 nmol L <sup>-1</sup>	7.5–500 μmol L <sup>-1</sup> 0.16–16 nmol L <sup>-1</sup>	Water samples Tap water	Li, Cheng, et al., 2016

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Table 4. C	ontinued.						
Type	Analytes	Aptamer based on	. Analytical method/transducer	LOD	Linear concentration range	Type of samples	References
		DNA	AuNPs and Rhodamine B as sensing probe, fluorescent guencher and fluorescent indicator	0.48 nmol L <sup>-1</sup>	0.48–200 nmol L <sup>–1</sup>	Water samples	Ni et al., 2017
EDCs/hormone	Progesterone	DNA	Fluorescence detection <sup>b</sup>	110 pg L <sup>_1</sup>	10–100 ng L <sup>-1</sup>	Tap water	Alhadrami, Chinnappan, Eissa, Rahamn, & Zourob, 2017
Pesticide	Malathion	DNA	Cationic peptide and unmodified AuNPs	1.94 pmol L <sup>-1</sup>	0.01-0.75 nmol L <sup>-1</sup>	Environmental samples	Bala et al., 2017
Pesticide	Acetamiprid and Atrazine	DNA	Sputtering and e-beam lithography techniques, Pt NPs deposited in a bridge-like arrangement, in between IDEs	1 pmol L <sup>-1</sup> for acetamiprid; 10 pmol L <sup>-1</sup> for atrazine	0.10–100 nmol for acetamiprid; 0.1–1000 nmol L <sup>–1</sup> for arrazine	Tap and mineral water	Madianos et al., 2018
Antibiotic	Oxytertra-cycline	DNA	GO hydrogel developed through a fast and facile delation. immersion and fluorescence determination	25 μg L <sup>-1</sup>	25-1000 μg L <sup>-1</sup>	Environmental samples	Tan, Zhao, Du, Gan, & Ouan. 2016
		DNA	Controllable peroxidase-like catalysis of a graphene/ gold nanoparticle colorimetric aptasensor	91 nmol L <sup>-1</sup>	0.17–0.50 µmol L <sup>-1</sup>	Water samples	Yuan, Zhao, et al., 2017
Antibiotic	Sulfadimetho-xine	DNA	Label-free tunable peroxidase-like activity of G/Ni-Pd hybrids – colorimetric aptasensor	0.7 µg L <sup>-1</sup>	1–500 μg L <sup>–1</sup>	Lake water	Wang et al, 2017
Antibiotic Piezoelectric ser	Ampicillin 1sors	DNA	AuNPs/MBs	70 ng L $^{-1}$	0.1–100 μg L <sup>–1</sup>	River water	Luo et al., 2017
Heavy metal	Pb <sup>2+</sup>	DNA	SAM of the AuNPs on the QCM	4 nmol L <sup>-1</sup>	5-200 nmol L <sup>-1</sup>	Water samples	Yuan, Song, et al., 2017
Toxin	Ochratotoxin A	DNA	EWA platform	3 nmol L <sup>-1</sup>	6–500 nmol L <sup>–1</sup>	Water and food samples	Wang, Xiang, Zhou, Liu, & Shi, 2015
Pathogen	Salmonella enterica	DNA	Aptamer-based QCM		100 CFU mL <sup>-1</sup>	Water and food samples	Ozalp et al., 2015
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phene; MIPMs, molecularly imprinted polymer microspheres; NPG, nanoporous gold; QCM, quartz crystal microbalance; PFP, poly(9,9-bis(6-N,N,n-trimethylammonium)hexyl)fluorine phenylene; PQC, piezoelectric quartz crystal; RGO, dotted reduction graphene oxide; RS, resonance scattering; SAM, self-assembled monolayer; SPCE, screen-printed carbon electrode; SERS, surface-enhanced Raman scattering; T, thymine; TH, thionine. wave all-fiber; FAM, 6-carboxyl fluorescein; FC, ferrocene; FRET, fluorescence resonance energy transfer; G, grapheme; GA, glutaraldehyde; GCE, glassy carbon electrode; GO, graphene oxide; IDEs, interdigitated electrodes; LFS, lateral flow strip; LOD, limit of detection; MBs, modified magnetic beads; MEF, metal-enhanced fluorescence; MGO, magnetic oxidation gra-3D, three dimension; AuNPs, gold nanoparticles; BHQ1, Black Hole Quencher 1; CD,  $\alpha$ -cyclodextrin; CFU, colony forming unit; CoS, cobalt sulfide nanosheet; CNTs, carbon nanotubes; CTAB, cetyltrimethylammonium bromide; EDCs, endocrine disrupting compounds; EIS, Aptamer-based label-free impedimetric biosensor for detection of progesterone; EWA, evanescent

Poince contained activities in the mean sourcempt, it is many intermed and  $L^{-1}$  (for two studied aptamers) "Dissociation constants Kd = 1.60 ± 0.16 µmol L<sup>-1</sup> and 0.76 ± 0.18 µmol L<sup>-1</sup> (for two studied aptamers)

#### 947 **3.2. Aptamer based tests**

948 Aptamer-based assays may one day compete with the currently popular 949 ELISAs (enzyme-linked immunosorbent assays) where antibodies are used 950 as the recognition element. ELISAs are very sensitive and may be selective 951 towards various analytes. However, antibodies are still required for ELISAs 952 and as previously mentioned, antibody procurement requires an animal 953 source. This factor may play a significant role in future preference for 954 aptamer-based assays vs. ELISAs. Enzyme-linked apta-sorbent assays 955 (ELASA) can be performed with different variations (enzyme-linked 956 aptamer assay (ELAA), enzyme-linked oligonucleotide assay (ELONA) and 957 aptamer-linked immobilized sorbent assay (ALISA)). Aptamers in these 958 tests may be labeled with fluorescein and replaced with a secondary anti-959 body, or they may directly bind the target molecule being attached to the 960 medium and act as a primary antibody. Comparisons of ELONAs with 961 ELISAs show that ELONAs are also a very sensitive and precise method 962 (Shi et al., 2014; Toh et al., 2015). Aptatests are performed mainly on 963 multi-welled test plates and the test endpoint is observed (by eye) or meas-964 ured (by special equipment) by a change in color or fluorescence signal. 965

Aptamers may also serve as recognition elements in lateral flow assays 966 (LFAs), which are a rapid format platform for the detection and quantifica-967 tion of analytes in complex mixtures. Currently, LFAs are usually used for 968 home-based or in situ diagnostics, such as popular pregnancy tests; these 969 tests are inexpensive, easy to use and utilize antibodies as a target recogni-970 tion element (Koczula & Gallotta, 2016). Despite the effectiveness of anti-971 body-based LFAs, alternative versions based on aptamers are sought after 972 (Chen & Yang, 2015). The LFA strip/dipstick consists of several overlap-973 ping membranes that are mechanically stabilized on a small backing card, 974 usually made from nitrocellulose. A few drops of sample are placed in the 975 designated area, which then migrate laterally to the membranes at the con-976 jugation pad by capillary force. There the fluid rehydrates a detection label 977 composed of a recognition element (antibody or aptamer) coupled to a 978 reporter molecule (AuNP), that form a complex with the target (Schüling, 979 Eilers, Scheper, & Walter, 2018). The target molecule connects to the cap-980 turing molecule (AuNPs), and as soon as the complex target-AuNPs crosses 981 the test line containing the aptamers or antibodies, the color of the strip 982 can be observed by the naked eye. To assure proper test operation (proper 983 fluid flow, activity of recognition, and capture molecules), there is also a 984 control strip placed on the test strip. This type of design is known as a 985 competitive design (Figure 9b) (Schüling et al., 2018). In a 'sandwich-LFA' 986 (Figure 9a), when an analyte has multiple binding sites, the detection labels 987 provided within the conjugation pad react with the target analyte to form a 988 complex. This complex is then captured in the test strip by an interaction 989



**Figure 9.** Sandwich LFA a) and competitive LFA with aptamer as recognition element (Bahadır & Sezgintürk, 2016; Jauset-Rubio, El-Shahawi, Bashammakh, Alyoubi, & O'Sullivan, 2017).

Table 5. Types of aptamer-based test and information about target molecules and lowest recorded concentration.

				The lowest concentration of	
Test	Aptamer	Target/		targets recorded in	
name	type	Analytes	Samples	the research	References
ELASA	DNA	SGIV virus	Biological material	As low as 125 nmol L <sup>-1</sup>	Li, Zhou, et al., 2016
	DNA	Basic fibroblast growth factor (bFGF)	Biological material	The minimum detection level of 7 pg mL <sup>-1</sup>	Golden, Collins, Willis, & Koch, 2000
ELONA	RNA	Cytokines	Biological material	The minimum detection level was 100 pg mL <sup>-1</sup>	Yan, Gao, Yao, & Zhang, 2004
	DNA	Zearalenone monoclonal antibody	Corn samples	_	Wang, Zou, et al., 2015
	DNA	Leishmania infantum H2A antigen	Biological material	_	Ramos et al., 2007
ALISA	DNA	Tulmeria antigen produced by Francisella tularensis	Bacterial cultures	Minimal concentration 100 ng mL <sup>-1</sup>	Vivekananda & Kiel, 2006
ELAA	DNA	Influenza A virus	Virus breeding	—	Shiratori et al., 2014
LFSA	DNA	Hg <sup>2+</sup>	Water samples	$LOD - 0.13 \text{ ng mL}^{-1}$	Wu, Shen, et al., 2017
	DNA	Chloramphenicol	Food samples	LOD – 0.0031 na mL <sup>-1</sup>	Yan et al., 2018

LOD, limit of detection; LOQ, limit of quantification; ELONA, enzyme-linked oligonucleotide assay; ELAA, enzyme-linked aptamer assay; LFA, Lateral Flow Assay; ALISA, aptamer-linked immobilized sorbent assay.

between the analyte and the corresponding immobilized aptamers. The fluid reaching the test zone contains labeled and unlabeled target molecules that compete to bind to the capture molecules immobilized in the test zone. Examples of this along with additional information are presented in Table 5.

The use of sensors (in which the recognition elements are aptamers) makes the number of purposes to be determined becomes virtually

1033unlimited. Aptamers are small, chemically stable and cheaper compared to1034other sensors, in particular those based on antibodies. On the one hand,1035the ability to generate an aptamer to determine different targets is benefi-1036cial, but many of process of generating such aptamers are quite problem-1037atic. Aptasensors, in relation to other sensors, are characterized with high1038selectivity and sensitivity and the aptamers themselves are very sta-1039ble molecules.

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## 4.1. Medical applications

4. Other aptamer applications

Aptamers are considered 'smart ligands', which are ligands specially selected for specific parameters that generate a known biochemical reaction (Drabovich, Berezovski, Musheev, & Krylov, 2009). The aptamers most commonly used in medical applications focus on RNA bases that are more structurally diverse than DNA based aptamers. Known groups of aptamers can be directed to specific factors (A30, A9, A10, TTA1, and MUC1) located on the cytoplasmic membranes of tumor cells. These factors are extremely overexpressed only on the surface of tumor cell membranes (Pfeiffer & Mayer, 2016; Ferreira, Cheung, Missailidis, Bisland, & Gariepy, 2009; Zhou et al., 2016; Marton, Reyes-Darias, Sánchez-Luque, Romero-López, & Berzal-Herranz, 2010; Hicke et al., 2006). Aptamers used in targeted anticancer therapy are most often carriers of polymeric nanoparticles that contain antimitotic drugs such as paclitaxel; these medicines inhibit cancerous cell division (Dhar, Gu, Langer, Farokhzad, & Lippard, 2008; Tong, Yala, Fan, & Cheng, 2010).

In 2004, the FDA (Food and Drug Administration) approved Macugen<sup>®</sup> (pegaptanib), an aptamer based drug used to treat adults who suffer from wet-form age-related macular degeneration. In this treatment, a modified pegylated oligonucleotide binds selectively and strongly to the extracellular form of endothelial growth factor (VEGF165) to inhibit its action. In January 2017, the FDA approved oligonucleotide therapies following assessment that aptamer-based drugs provided a clear benefit in rigorously controlled clinical trials (Stein & Castanotto, 2017).

Radiology methods are used to identify, locate, and assess the degree of disturbance in the structure and function of examined organs. These methods require the use of contrast-friendly agents with a suitable signal for detection and high tissue specificity. Such a role may be played by labeled aptamers capable of binding to proteins found on the surface of specific tissues or cells. Using established *in vitro* tumor cell lines, aptamers directed against tenascin C (a marker protein produced in large quantities by cancer cells) and the extracellular domain of the RTK receptor (a tyrosine kinase

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1076transmembrane protein involved in signal transmissions necessary for can-1077cer proliferation and growth) have been successfully isolated (Hicke1078et al., 2001).

In addition to cancer cell detection, aptamers can be used to detect viruses, bacteria, and even parasites. There are reports of aptamers specific to the hepatitis C virus, influenza virus, type 1 human immunodeficiency virus, *Staphylococcus aureus*, *Bacillus subtilis*, and parasites that cause malaria. These aptamers bind to proteins present on the target membrane, or to host antibodies due to the presence of a given bacterium, virus, or parasite (Misono & Kumar, 2005; Biroccio, Hamm, Incitti, De Francesco, & Tomei, 2002; Pavski & Le, 2001; Someya et al., 2012; Cao et al., 2009).

1087 Western blotting with aptamers instead of antibodies offers researchers 1088 multiple benefits including an increased number of fluorophores, which 1089 increase the signal-to-noise ratio, fewer nonspecific bands, direct conjuga-1090 tion to the reporter molecule, and lower costs. Aptameric Western blotting 1091 allows the selective staining of proteins transferred to a membrane and is 1092 more effective than antibodies. Although not common the using an RNA 1093 His-tag-specific aptamer (which was further conjugated to a QD) has been 1094 reported (Shin, Kim, Kang, Yang, & Hah, 2010; Radom et al., 2013). 1095 Aptamers are also currently used in imaging diagnostics as carriers of 1096 fluorescent dyes, such as fluorescein or rhodamine, which can be detected 1097 by spectroscopic techniques following activation (Xu, Yang, Ye, He, & 1098 Fang, 2006).

1099 One of the very promising options for using aptamers is associated with 1100 point-of-care-testing (POCT) and personalized medicine of individual 1101 patients (Vashist, Luppa, Yeo, Ozcan, & Luong, 2015, Dhiman, Kalra, 1102 Bansal, Bruno, & Sharma, 2017). Among the many possibilities colometric 1103 and fluorescence tests offer the possibility of a quick visual assessment, 1104 which facilitates diagnostics and does not require specialist personel or 1105 equipment. The use of highly selective target binding aptamers enables fast 1106 color development that may be assessed qualitatively and/or quantitatively 1107 on simple spectrometer devices (Dhiman et al., 2017).

1108 Three different generations of Point-Of-Care (POC) tests are especifically 1109 designed to work at different levels according to individual patient require-1110 ments. The first ones include maintenance-free rapid tests (e.g. LFAs), the 1111 second-generation cover cartridge-based tests (e.g. nucleic acidamplification 1112 tests), the third generation of aptamer-besed POC tests include personal 1113 hand-held equipments (Pai, Vadnais, Denkinger, Engel, & Pai, 2012, 1114 Dhiman et al., 2017). Aptamers are a promising tool in the POC test group, 1115 are easier to obtain than crosses, it is possible to introduce aptamers to the 1116 development of commercial technologies, some of the POC tests already 1117 described in the literature are summarized in Table 6. 1118

### 1119 4.2. Analytical applications

1120 Although aptamers are increasingly applied in medical science, the possibil-1121 ities for their use extend beyond the medical arena. Aptamers are used suc-1122 cessfully in forensic science for toxicological analysis, to target analytes, 1123 and for chemical and biological weapons (Yáñez-Sedeño, Agüí, Villalonga, 1124 & Pingarrón, 2014). Aptamers can reveal latent fingermarks (Wood, 1125 Maynard, Spindler, Lennard, & Roux, 2012) and a deoxyribonucleotide-1126 based aptamer is used to detect cocaine in body fluids (Stojanovic, De 1127 Prada, & Landry, 2001). 1128

In affinity chromatography used for protein purification, specific 1129 aptamers are immobilized on a chromatography support to create an affin-1130 ity column that separates the sample ingredients (Romig, Bell, & Drolet, 1131 1999). Aptamer based electrochemical sensors have great potential due to 1132 their ability to recognize a large number of targets simultaneously present 1133 in environmental samples (Wu, Liu, Zhan, Wang, & Zhou, 2012; Hoang, 1134 Oyama, Saito, Aono, & Nagao, 2013; Li, Dong, & Wang, 2009; Hayat & 1135 Marty, 2014; Lee et al., 2009). Compared to aptamers, antibodies used in 1136 affinity chromatography have to be thoroughly purified before sample sep-1137 aration, lest the impurities significantly impact target isolation and purifica-1138 1139 tion. Additionally, there significant costs associated with obtaining antibodies and quality differences in the received batches. 1140

## 4.3. Aptamer based tests

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Aptamer-based tests can be used to detect food contaminants, for quality control, and as a diagnostic tool for early and rapid detection of specific risk factors (Toh et al., 2015). ELONA and ELAA tests are being developed to rapidly identify various pathogens including influenza viruses H1N1, HPV and the *Mycobacterium tuberculosis* (protein MPT64) bacteria (Bai et al., 2018; Sypabekova et al., 2017).

Recently Mukherjee, Manonmani, and Bhatt (2018) introduced ultrasensitive ELASA to detect aflatoxin  $B_1$  at a detection limit of 10 pg mL<sup>-1</sup> in a buffer. The aptamer based-test was characterized by higher sensitivity and recovery compared to antibody-based ELISA. Wu et al. (2015) described an LFA for the rapid visual assessment of the presence of *Escherichia coli* O157:H7 where as low as 10 CFU mL<sup>-1</sup> could be detected in laboratory tests.

For the time being, biological tests with aptamers are not as developed as antibody-based analogs. However, as previously mentioned, aptamers are a good alternative due to their simple preparation methods and the wide environmental ranges in which aptamers are active. Figure 10 and Table 7

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**Figure 10.** Applications of aptamers (Keefe, Pai, & Ellington, 2010; Daniel, Mélaïne, Roupioz, Livache, & Buhot, 2013; Hicke et al., 2006. Wiedman et al., 2017; Pfeiffer & Mayer, 2016; Jin et al., 2016; Battig & Wang, 2014; Darmostuk, Rimpelova, Gbelcova, & Ruml, 2015; Cruz-Aguado & Penner, 2008; Joshi et al., 2009; Hayat & Marty, 2014; Bagalkot et al., 2007; Wood et al., 2012. Song, Jurng, Park, & Kim, 2016).

contain a summary of the different applications of aptamers in various scientific fields.

#### 5. Future perspectives

The emergence of artificial genetic polymers known as xeno nucleic acids (XNAs) offer several options for the creation and synthesis of polymers that function like DNA and RNA. However, XNAs are poorly recognized by organism nucleases, making them highly resistant to decay and resulting in longer half-lives *in vivo*. Modifications to the XNA backbone include sugar replacements, such as those in locked nucleic acids, arabinonucleic acids, cyclohexenyl nucleic acids, 1,5-anhydrohexitol nucleic acids, and 2'-fluoroarabinonucleic acids (Bruno, 2015; Østergaard et al., 2014; Istrate, Medvecky, & Leumann, 2015). XNAs can be used to develop ultra-stable aptamers, however, any such modification may increase the risk of an adverse patient reaction (Bruno, 2015).

Aptamers also demonstrate the potential for stem cell differentiation, possibility indicating the ability to replace expensive recombinant growth factors in regenerative medicine, or to discover unknown surface cell receptors (Bruno, 2017). Stem cell differentiation can be beneficial in targeted

248         249         250         251         252         253         254         255         256         257         258         259         260         261         262         263         264         265         266         267         268         269         270         271         272         273	on of aptamers in various fields of science.	of samples/active sites Comments Reference	an blood serum LOD – 100 pmol L <sup>-1</sup> Roushani & Shahdost-fard, 2015 igical material LOD – 0.9 µmol L <sup>-1</sup> Neves, Blaszykowski, Bokhari, & Thomson, 2015	samples 2.3 µg L <sup>-1</sup> Kim, Lee, Jin, Lin, Jeong, 2014 Learned Lorent and Landon 2014	Ly range to $DNA = 15.7 \text{ mg } L^{-1}$ , $RNA = 10.1 \text{ mg } L^{-1}$ Toh et al., 2015; Jeong & Rhee Paeng, 2012	l samples LOD – 60 mol L <sup>-1</sup> Liu, Ouyang, et al., 2018 acial material LOD – 0.05 ua L <sup>-1</sup> Farzin. Shamsipur. & Tabrizi, 2015	ojcal sample LOD 0.05 μg[ <sup>1</sup> , range 0.3–867.5 μg L <sup>-1</sup> Shamsipur, Farzin, Amouzadeh Tabrizi, & Sheibani 2017	m no data Park & Paeny 2011 m and huffer samples Datectable concentrations up to 5 mmol 1 <sup>-1</sup> Bacton Nouven & 6u 2017	substances/solutions — — Yang et al., 2017	ogical samples High sensitivity (250 ng L <sup>-1</sup> ), fast detection Su, Fong, Cheung, & Yang, 2017 (within 30 min) with samples (1 ul)	er tissue —Zhuang et al., 2016	an blood serum — Tombelli, Minunni, Luzi, & Mascini, 2005	igical fluids — — Chen, Zhou, Chen, & Chen, 2015	igical fluids LOD – 0.3 חס וגL <sup>-1</sup> Bai et al, 2018 LOD – 0.9 גס וגL <sup>-1</sup>	gical fluids – – 2017 Liu, Zhang, Liu, Sharma, & Ding, 2017	d samples Lowest detected concentration 0.31 ng mL <sup>-1</sup> Her, Jo, & Ban, 2017	samples — — — — — — — — — — — — — — — — — — —	samples LOD – 2 CFU mL <sup>-1</sup> Di, Du, Pan, & Wang, 2017		samples LOD – 6 ng mL <sup>-1</sup> Huang, Chen, et al., 2015	cal sputum samples —		L samples — — — JOD and the LOD are 0.82 and 2.5 md l <sup>= 1</sup> Costantini et al 2016	Limit of detection 1ng mL <sup>-1</sup> Barthelmebs, Jonca, Hayat,	Prieto-Simon, & Marty, 2011 water 2.4 ng mL <sup>-1</sup> Frohnmeyer, Frisch, Falke, Betzel, &
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cancer therapy (Shamaileh et al., 2016). Aptamers have also proved to be
promising biorecognition elements in the development of electrochemicalbased sensors. Aptameric identification capabilities with selective detection
capacities for electrochemical signal transduction enable an almost unlimited
number of biosensors. Unfortunately, the currently limited number of proven
aptamers hinders the progression of aptamer-based electrochemical sensor
development (Liu, Morris, Macazo, Schoukroun-Barnes, & White, 2014).

1298 Due to their small sizes and dimensions, aptamers may cross physio-1299 logical barriers, enabling the delivery of therapeutic and diagnostic agents 1300 to intracellular organelles. Targeted/tailored nano-delivery will allow cell-1301 based strategies for delivery of active substances to the mitochondria, 1302 nucleus, lysosomes, or endoplasmic reticulum. However, it is important to 1303 understand the basics of organelle-targeted drug delivery strategies, which 1304 are critical to the design and successful use of biomedical nanomaterials for 1305 therapeutic treatment (Ma, Gong, Zhong, Sun, & Liang, 2016). The com-1306 bination of liposomes and aptamers is characterized by high efficacy and 1307 selectivity in the treatment of many diseases, including many types of can-1308 cer. In addition, aptamer-liposome combinations can be used as highly 1309 selective diagnostic sensors (Ries & Vogel, 2016).

1310 It is very likely that aptamers will replace antibodies in certain scientific 1311 and medical areas in the future (Li, Ho, & Ding, 2015). Research on 1312 aptamers continues to increase, and the PubMed publication database con-1313 tains almost 8,000 articles on this subject (PubChem, 2018). In terms of 1314 environmental monitoring, aptamer potential is not yet fully utilized. 1315 Aptamer based biosensors or tests for the presence of environmental pollu-1316 tants are still primarily in the prototype phase of development and are usu-1317 ally tested on samples with simple matrices such as tap or ground water. 1318 However, current reports are very promising, especially regarding the abil-1319 ity of aptamers to recognize a wide range of analytes at very low levels, in 1320 addition to the possibility of regeneration and their considerable stability 1321 (Justino et al., 2017; Verna and Bhardwaj, 2015). 1322

Based on a review of the current data, one can conclude that the utilization of aptamers in the 21<sup>st</sup> century is limited only by their intrinsic properties and the imagination and skills of the researcher. Environmental protection and monitoring issues are of growing importance in developed, developing and underdeveloped countries, so intelligent solutions for rapid diagnostic measurements *in situ* are required. The ease of preparing arrays for quick-format biotests and detection systems makes aptameric testing a promising solution in the following nine areas:

- 1. Aptamers can be used for designing rapid-format tests and portable analyzers for environmental monitoring and pollutant screening. This
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includes mixtures and samples characterized by complex matrix effects (Hsieh, Dantzler, & Weigl, 2017) and the automated assessment of risk/ exposure to different organisms based on advanced big-data treatments growing more accessible to an increasing number of societies (Ali et al., 2016; Andreu-Perez, Poon, Merrifield, Wong, & Yang, 2015; Bellinger, Jabbar, Zaïane, & Osornio-Vargas, 2017; Lokers, Knapen, Janssen, van Randen, & Jansen, 2016; Romero, Hallett, & Jude, 2017).

- 1341 2. Aptamers can be used for performing quick and comprehensive certified 1342 patient diagnostics at the point-of-care (or even at an accident site) to 1343 facilitate the process of decision making and the undertaking of life-sav-1344 ing actions (Burbelo, Ramanathan, Klion, Iadarola, & Nutman, 2008; 1345 Hesterberg & Crosby, 1996). Certain decisions on e.g. patients TRIAGE 1346 (French: to segregate) in cases of poisoning with unknown substances 1347 (or their mixtures, e.g. boosters), infection with unknown rare bacteria/ 1348 viruses, detection of possible allergic reaction to some pharmaceuticals 1349 of first choice (e.g. anesthetics) or showing symptoms of unknown bio-1350 chemical disorders can be undertaken already at first glance with basic-1351 level medical personnel to facilitate efficiency of patients care and short-1352 ening, afterwards, recovery time. Rapid development in this area of sci-1353 ence justifies conclusion that it is worth to prepare separate scientific 1354 review paper on this topic of interest. 1355
  - 3. Aptamers can be used as a certified early-warning system in households or public institutions (e.g. schools, hospitals) to rapidly detect the malfunction of media delivering systems, or even accidents or attacks with unknown agents that pose a threat to exposed individuals (Collins & Kapucu, 2008; Macherera & Chimbari, 2016; Moffitt et al., 2011; Greenhalgh, Robert, Macfarlane, Bate, & Kyriakidou, 2004).
  - 4. Aptamers can be used to create tailored 3D printed closed-loop modules that combine aptamers with bioassays (and possibly instrumental detection systems) to conduct chronic exposure studies of single and combined pollutants in organisms from different trophic levels (Juraschek, Cerdas, Posselt, & Herrmann, 2017; Debski et al., 2018; Fang et al., 2017).
  - 5. Aptamers can be used for rapid screening for boosters or illegal products in patient or convict body fluids for first-aid units or police or border guards (Avtonomov & Kornienko, 2015; Bendahan, 2017).
  - 6. Anti-terror aptasensor arrays can be used for detecting attacks with nerve gases, explosives, and biological or chemical agents. These sensors could also identify the transport of these agents, which is currently a challenging task. This is especially true when dealing with air samples that need to be studied and treated, while simultaneously detecting active agents to undertake protective actions if required (Caygill, Davis, & Higson, 2012; Ma, Wang, & Wang, 2015; Wells & Bradley, 2012).

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 Aptamers can be used for food safety and packaging material stability research after elaborating universal and package oriented methodologies based on international standards (Szczepańska, Kudłak, & Namieśnik, 2018).

- 8. They can be used to screen for doping incidences in human or animals sports (Alvarez-Risco, Del-Aguila-Arcentales, Delgado-Zegarra, Yáñez, & Diaz-Risco, 2019; Ayotte, Miller, & Thevis, 2017; Thevis, Kuuranne, Geyer, & Schänzer, 2017). In case of consumers point of view, the application of aptamers may be helpful in authentication of food products by performing cheap rapid-format aptasensors-based quickly read-out by consumers themselves (or quality control personnel).
- 1387 Target oriented aptasensor arrays can be used for screening the restor-9. 1388 ation/remediation process efficiency (Bardos et al., 2015; Sevcu et al., 1389 2017). Automated read-out can be even performed by unskilled patients 1390 themselves or non-skilled personnel by taking photo (with smart- or 1391 wearable devices) of quick-assay result and transferring data to cloud; 1392 certainly, based on current developments in the field, results interpret-1393 ation can be easily done either automatically or by professionals for ini-1394 tial screening of any above mentioned applications or to observe 1395 progress of ongoing processes e.g. of medical treatment at non-hospital 1396 conditions. More and more tests can be analyzed using generally avail-1397 able devices such as a camera, scanner or smatrphone. Due to the possi-1398 bility of taking good quality photos and various image processing 1399 software and portability, one of the best smartphones is considered 1400 image acquisition tools. Literature provides examples of the use of 1401 aptamer-based tests that can be analyzed with the use of a smratphone 1402 e.g. in 2019 Umrao et al. propose a FRET aptasensor with dye pairs on 1403 DNA aptamers to rapid and sensitive kanamycin detection, which 1404 reached the limit of detections of 28 nmol L<sup>-1</sup> (Umrao, Anusha, Jain, 1405 Chakraborty, & Roy, 2019). 1406

Additionally, technological development will contribute to the rapid production of highly selective aptamers and a reduction of their cost, which is smaller than that of antibodies but still significant (Song, Zhang, Zhu, & Yang, 2015). The rapid development of artificial intelligence (AI), especially in terms of data interpretation and big data mining, as well as the potential application in designing/tailoring aptamers and correlating results with individual patients should also be considered.

- 6. Conclusions
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- i. Aptamers have unique intrinsic properties, manifesting in the specificity and selectivity of the interactions between aptamers and target

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molecules. These properties affect aptameric aptitude for therapeutic utilization, and environmental pollutant analyses.

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  ii. The widespread use of aptamers depends on conditions including the cost, development of specific aptamers, and incorporation with other available technologies. All of these tasks are challenging but achievable, assuming rational cooperation between scientists including biologists, chemists, biotechnologists, chemometricians, toxicologists, and doctors, as well as stakeholders, authorities (and their social representatives), and NGOs.
- 1429 iii. The review provides an overview of the current technologies and meth-1430 odologies for the creation and utilization of aptamers with a specific 1431 focus on environmental and medical fields. The development of a var-1432 iety of applications and methods of utilizing aptamers are thoroughly 1433 discussed and for these reasons would interest readers in the fields of 1434 biosensors, medicine, or environmental monitoring; it makes a signifi-1435 cant contribution to the literature by consolidating existing data on 1436 aptamers. This makes it possible to clearly see overlapping applications 1437 for this technology as well as the potential for cross-disciplinary 1438 collaboration. 1439

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## Disclosure statement

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