**Abstract**

**Introduction** Many bacteria are responsible for infections in humans and plants, being found in vegetables, water, and medical devices. Most bacterial detection methods are time-consuming and take days to give the result. Aptamers are a promising alternative for a quick and reliable measurement technique to detect bacteria present in food products. Selected aptamers are DNA or RNA oligonucleotides that can bind with bacteria or other molecules with affinity and specificity for the target cells by the SELEX or cell-SELEX technique. This method is based on some rounds to remove the non-ligand oligonucleotides, leaving the aptamers specific to bind to the selected bacteria. Compared with conventional methodologies, the detection approach using aptamers is a rapid, low-cost form of analysis.

**Objective** This review summarizes obtention methods and applications of aptamers in the food industry and biotechnology. Besides, different techniques with aptamers are presented, which enable more effective target detection.

**Conclusion** Applications of aptamers as biosensors, or the association of aptamers with nanomaterials, may be employed in analyses by colorimetric, fluorescence, or electrical devices. Additionally, more efficient ways of sample preparation are presented, which can support food safety to provide human health, with a low-cost method for contaminant detection.

**Key points**
- Aptamers are promising for detecting contaminants outbreaks.
- Studies are needed to identify aptamers for different targets.

**Keywords** Biosensor · DNA Aptamer · Cell-SELEX · Bacteria

**Introduction**

The growth of epidemiologic diseases has been increasing due to food production and consumption trends, such as production processes, globalization of consumer goods, and high demand for raw or undercooked foods (Cheung and Kam 2012). Gram-positive and gram-negative bacteria are responsible for infectious diseases even at low concentrations (Soundy and Day 2017). Many methods may be applied to identify these bacteria, such as conventional bacterial culture, immunofluorescence, and polymerase chain reaction (PCR). However, these techniques are often time-consuming, without quantitative response, and often they are high-cost analyses (Lavu et al. 2016). In the last decades, rapid and efficient methods of detection are being developed to decrease food contamination and improve human health (Amaya-González et al. 2013).

Alternative methods to the detection of microorganisms are the use of single strands of DNA or RNA, called aptamers, which are capable to bind to non-nucleic acid molecules. The technique known as systematic evolution of ligands by exponential enrichment (SELEX) combines different steps such as incubation of oligonucleotides library, separation, amplification by polymerase chain reaction (PCR), and purification (Liu and Zhang 2015). This approach has been used since the 1990s when Tuerk and Gold (1990) discovered an RNA aptamer capable to bind to the T4 DNA polymerase, and Ellington and Szostak (1990) studied the bind of RNA aptamer to organic dyes. Over the years many publications were published on the use of aptamers in different ways, eventually conjugated with metallic, oxide or polymeric nanoparticles, and carbon nanotubes (Liu and Zhang 2015).
This review aims to address the techniques for obtaining aptamers, and their use particularly for contaminants detection in food products. The structure and properties of aptamers, as well as nanomaterials commonly used to be conjugated with aptamers, are presented. Moreover, recent applications and market perspectives are also introduced.

**Aptamers structure and properties**

Aptamers are single-stranded DNA or RNA molecules (oligonucleotides) with 50 to 100 nucleotides bases, 1 to 2 nm in size, and 7.5 to 32 kDa, with high affinity and specificity to bind to a target molecule (Ellington and Szostak 1990; Stoltenburg et al. 2005). According to Radom et al. (2013), DNA aptamers are molecules more stable than RNA; however, no significant differences were found between their specificities and binding capacities.

Aptamers can assume stable three-dimensional configurations in the aqueous phase, such as lops, triplexes, pseudoknots, G-quadruplexes, and staples (Bing et al. 2017). Due to the ability to change their shape, they can bind to different targets such as amino acids, vitamins, nucleotides, proteins, pesticides, drugs, bacteria, and inorganic or organic compounds. Moreover, they can be used as enzyme inhibitors to recognize proteins and nucleic acids, as well as inhibitors of toxins and hormones, applied in the detection of molecules in complex mixtures, purification, and biosensors (Liu and Zhang 2015; Peterson et al. 2015).

The affinity with the target molecule is associated with the dissociation constant that ranges from picomoles per liter to nanomoles per liter, being calculated through thermodynamic stability (Takenaka et al. 2017). Besides, aptamers may be stored and transported at room temperature due to the stability in the environmental conditions (Missailidis and Perkins 2007). Also, modifications in the aptamers can make them more resistant, stable, and improve the targeting ability, as shown by Ni et al. (2017). According to the authors, aptamers for the therapeutic area are susceptible to nuclease degradation and can be excreted by renal filtration even before they bind to the target. Thus, to increase the resistance to nuclease degradation, aptamers can be modified in the 3’ end with inverted thymidine or conjugated with biotin or biotin-streptavidin. Another modification is at 5’ position, which consists of the addition of cholesterol to increase the aptamer stability in plasma (Ni et al. 2017).

In the same way, it is possible to combine aptamers with nanostructures, such as gold nanoparticles or carbon nanotubes (Radom et al. 2013). They can be labeled with compounds such as dye or biotin. This process is widely used as systematic evolution of ligands by exponential enrichment (SELEX) to control the binding properties (Missailidis and Perkins 2007; Radom et al. 2013).

**Systematic evolution of ligands by exponential enrichment**

SELEX is a technique used to obtain aptamers through the in vitro selection of oligonucleotides over many rounds (Radom et al. 2013). In the 1990s, the first reports using the SELEX technique were published with RNA aptamers bounded to T4 DNA polymerase with high affinity (Tuerc and Gold 1990). In addition, Ellington and Szostak (1990) studied in vitro selection of organic dyes-binding RNA, and Robertson and Joyce (1990) selected the tetrahymena ribozyme and through amplifications obtained an RNA sequence able to cleave a specific DNA sequence.

This technique allows the selection of specific oligonucleotides that can be chemically synthesized, without the need to be produced or selected in a living organism (Stoltenburg et al. 2007). As shown in Fig. 1, SELEX first step is to obtain a library of random oligonucleotides, which can be single strands of DNA or RNA, with $10^{14}$ to $10^{15}$ different sequences. This library, also called a bank, has a fixed region and a random region of 20 to 80 nucleotides, which allows the molecules to form different structures as already mentioned, ready to bind the desired molecules (Stoltenburg et al. 2005). The fixed region, which is the library extremities, allows the amplification (Stoltenburg et al. 2005).

The binding step occurs by the incubation of the bank of ligands (aptamer candidates) and the target molecule in a buffer solution. After this, the bounded oligonucleotides are separated from non-bounded by physical separation and PCR amplification is carried out on the recovered bounded oligonucleotides, thus the first SELEX cycle started (Radom et al. 2013). Different techniques can be used in the separation process, such as centrifugation (Soundy and Day 2017; Ramnal et al. 2018); capillary electrophoresis (Tang et al. 2006); ultrafiltration with nitrocellulose filter (Josh 2009); flow cytometry (Davis et al. 1996); and affinity chromatography using agarose, sepharose, magnetic beads and microwell plate, or sol-gel channels to immobilize the target (Mcqueag et al. 2009; Bae et al. 2013; Kim et al. 2014).

The separation of aptamer candidates from the target molecule can be carried out by heat treatment, competing for ligand elution, urea addition, EDTA, or SDS techniques (Weiss et al. 1997; Bianchini et al. 2001; Theis et al. 2004; Stoltenburg et al. 2005). According to Ellington and Szostak (1990), the affinity chromatography technique is used when it is desired to obtain aptamers for small targets.

For the second SELEX cycle, the amplified product (double-stranded DNA or RNA) should be converted by PCR to a single-stranded. According to Stoltenburg et al. (2005), if the library is DNA, the most used methods are biotin/streptavidin-added electrophoresis for strand distinction (desired and unwanted), size-difference primers where the unwanted strand is modified, modified primers at tip 3’ (addition
of a ribose), or hexamethylene glycol spacer primers. For the last two methods, electrophoresis is used to check the size difference and separate the unwanted tape. If the double strand is RNA, dsDNA is transcribed into RNA with T7 RNA polymerase. After the simple tapes are incubated again with the target, one more SELEX cycle occurs (Stoltenburg et al. 2005).

The cycles are performed until a binder with high affinity and specificity is obtained (Fig. 2), and in this case, 5 to 20 cycles can be performed (Tuerk and Gold 1990; Missailidis and Perkins 2007; Liu and Zhang 2015). This number depends on some factors such as the selected library, selection conditions of oligonucleotides, the concentration of incubated target, as well as the concentration of aptamer candidates (Radom et al. 2013). Once obtained, the high affinity and specificity aptamers are cloned into bacterial vectors, usually, Escherichia coli, sequenced and characterized (Tuerk and Gold 1990; Stoltenburg et al. 2005; Missailidis and Perkins 2007).

Several aptamers have already been selected for a range of targets as inorganic ions (Na⁺) (Zhou et al. 2016), dopamine, and other organic compounds such as amino acids like L-tryptophan (Idili et al. 2019), proteins (Deng et al. 2014), antibiotics (Smart et al. 2020), and microorganisms (bacteria, viruses, and fungi) (Soundy and Day 2017; Smart et al. 2020). When the aptamers are obtained by whole cells, the technique is called cell-SELEX (Radom et al. 2013).

Contrariwise to the SELEX approach, which uses purified targets, the cell-SELEX technique allows the use of whole cells where no knowledge of target conformation or protein purification is required, and whole cells remain in their natural state throughout the selection process. This technique is used because in some cases, when the target is purified, the native configuration can be lost, and the target is masked. So, the candidate aptamers may not bind since the natural structure of the targets is not recognized (Ye et al. 2012). The cell-SELEX cycle follows the same structure as the SELEX technique, but with the addition of negative selection. This approach uses different cells that are non-target for reducing the number of aptamers, which bind with non-specific cells, thus increasing aptamer specificity (Fig. 2). In the negative selection, Ye et al. (2012) described that non-binding aptamers are discarded and those targeting cell-binding are eluted and amplified by PCR. On the other hand, to the negative control, non-target cells are incubated with amplified library and non-binding cells are separated and amplified by PCR and so on until high affinity and specificity aptamers are obtained (Ye et al. 2012). In the cell-SELEX, the candidate aptamers can bind to the three-dimensional configuration of the target (Ye et al. 2012).

Cell-SELEX is reported, in the literature, to select aptamers, i.e., Salmonella enterica serovar Typhimurium (Duan et al. 2013; Lavu et al. 2016), Pseudomonas aeruginosa (Soundy and Day 2017), Neisseria meningitidis (Mirzakhani et al. 2018), Escherichia coli O157:H7 (Amraee et al. 2019).
et al. 2017), \textit{Streptococcus pyogenes} (Hamula et al. 2011), \textit{Staphylococcus aureus} (Ramlal et al. 2018), \textit{Haemophilus influenzae} (Bitaraf et al. 2016), \textit{Trypanosoma cruzi} (Ulrich et al. 2002), tumor liver cells (Mi et al. 2009), and mouse stem cells (Guo et al. 2007).

In this way, the chosen method depends on the target. Many works employed the cell-SELEX technique to obtain aptamers for bacteria detection, providing high affinity and selectivity. On the other hand, the SELEX technique can be used to identify bacteria and other compounds such as organic and inorganic molecules, viruses, and tumors.

### Aptamers conjugation

Aptamers can have diverse applications, from basic research in medicine, pharmaceuticals, diagnostics, therapy, and drug development to pathogen detection, which encompasses the medical field and the food industry (Tuerk and Gold 1990; Lavu et al. 2016). In therapy, aptamers act as inhibitors of targets, as nucleolin inhibition (Radom et al. 2013), while for food safety, aptamers are used to detect contaminants (Amaya-González et al. 2013).

To improve the application range, aptamers may be conjugated to nanostructures, which assist in the identification of the target compounds. Common conjugates for aptamers are metal or silica nanoparticles, hydrogels, and even carbon nanomaterials, due to their biocompatibility, controllable chemical and physical properties, and stability (Liu and Zhang 2015; Yang et al. 2015).

Among the conjugation applications, one can be the aptamer conjugation for colorimetric detection. This type of detection is the most attractive and widely used since the target is detected through visual observation with the aid of colored reagent without the use of analytical instruments as a spectrophotometer. For this kind of application, gold, magnetic, or cerium oxide nanoparticles, carbon nanotubes, graphene oxide, or even polymers may be conjugated to the aptamers (Sharma et al. 2015). These nanostructured supports have been commonly synthesized and applied (Almeida et al. 2017; Valério et al. 2017; Chiaradia et al. 2018; Hoelscher et al. 2018; Maass et al. 2019).

Gold nanoparticles are widely used because they decrease the distance between dispersed particles and increase the size after aggregation resulting in red to blue colors, as shown in Fig. 3 (Sharma et al. 2015). Some authors reported the use of silver ions to improve detection sensitivity. In this case, silver ions adhere to the surface of AuNPs reducing silver atoms by electrons released from the reducing agent around the gold nanoparticles. Thus, the nucleation reaction increases the gold nanoparticles size changing the color making possible a visible identification (Liu et al. 2014).

Fluorescence is the emission of light from an excited molecule, a dye or even a nanomaterial that then returns to its initial state (Sharma et al. 2015). Both colorimetric and fluorescence assays are widely used in aptamer studies, as they present high sensitivity, high efficiency, and easy operation. Those techniques require the use of fluorophore and chromophore dyes for measurable signal emission (Sharma et al. 2015).

According to Sharma et al. (2015), besides dyes, some nanomaterials provide fluorescence emission, which are economically viable but have a time consuming laborious process that can affect the selectivity of the aptamer to the target...
binding. Assays without fluorescent markers consist of the use of DNA intercalators, base site binding dyes, and metallic nanomaterials with the fluorescence emission.

Organic dyes such as FAM (fluorescein amidite) are commonly bounded to oligonucleotides but have some limitations such as broad emission range, low photostability, low absorption, and photodegradation (Li et al. 2008). To improve these limitations, inorganic fluorescents like quantum dots can be used, since their optical characteristics depend on the size (1 to 10 nm). They have a broad absorption spectrum, narrow emission spectrum, and long life fluorescence. However, their drawbacks are the high-cost synthesis and toxicity associated (Sharma et al. 2015). An example of quantum dots is cadmium selenide, which has high luminescence and good quantum yield (Xu et al. 2015) as already reported to detect the presence of E. coli O157: H7 by Xu et al. (2015).

Yang et al. (2011) reported the use of carbon nanoparticles for fluorescence testing, highlighting material advantages such as low cost, high quantum productivity, simple preparation, low toxicity, good biocompatibility, good aqueous solubility, and superior photoluminescence properties. In addition to fluorescent dyes, nucleic acid dyes may be used (SYBR green I, AccuBlue, and PicoGreen). The drawbacks of the nucleic acid dyes are the low fluorescence intensity and the requirement to interleaving with the aptamer DNA sequence (Duan et al. 2014a).

The chemiluminescence technique also uses optical detection and is mainly applied in food safety since it produces energy through chemical reaction without the need for excitation source as in the fluorescence technique (Sharma et al. 2015). Chemiluminescence signals can be increased using AuNPs as catalysts enhancing biocompatibility and stability (Yang et al. 2011). DNA aptamers that are rich in guanine (G) can react with 3,4,5-trimethoxyphenylglyoxal (TMPG), which forms an energy-rich compound that emits light or transfers energy to some aptamer-coupled dye, such as 6-FAM (6-carboxyfluorescein) with green light emission and can be employed in target detection studies (Kwon et al. 2015). Thus, aptamers can be conjugated to different nanoparticles to increase the selectivity to the target. Besides increasing the selectivity, these nanomaterials allow visual detection.

### Aptamers applications

Several aptamers have been developed for different applications, as reported by Chan et al. (2008) that employed PEG-conjugated aptamers RB006 against coagulation. Wu et al. (2008) developed a PO RO10–60 aptamer to stimulate the immune response against pathogens by delaying symptoms and allowing the use of antibiotics. There are also electrochemical sensors, which are aptamers that act in real-time detection of cocaine in fetal bovine serum (Swensen et al. 2009).

Aptamer-based nanostructures have also been widely used in medicine, biology, and nanoelectronics due to the high stability, as shown in Table 1. The AS1411 aptamer conjugated with gold nanoclusters was tested in mice cancer cells and demonstrated to be a good radiosensitizer (Ghahremani et al. 2018). Through cell-SELEX, the JHIT2 aptamer was selected and labeled with FAM and iodine-131 to detect human hepatoma cell line HepG2 by a fluorescent signal (Zhang et al. 2020). All these studies show that the aptamers can assist in the rapid detection of different targets, shortening the time to start treatments that are important for human health.

Lavu et al. (2016) studied the use of gold nanoparticles with aptamers for the detection of Salmonella enterica. The aptamer SAL 26 was conjugated with gold nanoparticles at room temperature and in the presence of NaCl keeping the solution red, and when in the presence of Salmonella enterica (10^2 to 10^6 CFU/mL), the solution turned blue after 30 min. According to the authors, the color change is associated with
### Table 1: Examples of aptamers application to different targets in different areas

| Target                              | Method     | Sample                                      | Reference                          |
|-------------------------------------|------------|---------------------------------------------|------------------------------------|
| Streptococcus pyogenes              | Cell-SELEX | Cooked chicken                              | (Huang et al. 2018)                |
| Salmonella typhimurium              | Cell-SELEX | Pasteurized milk                            | (Duan et al. 2014b)                |
| Salmonella typhimurium and Vibrio parahemolyticus | Cell-SELEX | Frozen shrimp, chicken breasts              | (Duan et al. 2014a)                |
| Salmonella                          | –          | Pork                                        | (Ma et al. 2014)                   |
| Escherichia coli                    | Cell-SELEX | Milk and tap water and pond                 | (Kim et al. 2013; Jin et al. 2017)  |
| Staphylococcus aureus               | –          | Fresh fish                                  | (Jia et al. 2014)                  |
| Staphylococcus aureus               | Cell-SELEX | Pork meat                                   | (Hao et al. 2017)                  |
| Staphylococcus aureus               | Cell-SELEX | Milk                                        | (Yuan et al. 2014)                 |
| Listeria monocytogenes              | SELEX      | Liced beef, chicken, turkey                 | (Ohk et al. 2010)                  |
| Campylobacter jejuni                | SELEX      | Live cell                                   | (Bruno et al. 2009)                |
| Lactobacillus acidophilus           | Cell-SELEX | Oxidized PSI Fabry-Pérot thin films         | (Urmann et al. 2016)               |
| Francisella tularensis             | SELEX      | *Bacterial antigen*                         | (Vivekananda and Kiel 2006)        |
| Mycobacterium tuberculosis         | SELEX      | Live cell                                   | (Chen et al. 2012)                 |
| Vibrio parahemolyticus             | Cell-SELEX | Live cell                                   | (Duan et al. 2012)                 |
| Shigella sonnei                     | Cell-SELEX | Live cell                                   | (Song et al. 2017)                 |
| C. jejuni                           | Cell-SELEX | Live cell                                   | (Dwivedi et al. 2010)              |
| Vaccinia virus                      | SELEX      | Vaccinia intacto                            | (Labib et al. 2012)                |
| herpes simplex virus                | SELEX      | Gd protein of HSV-1                         | (Gopinath et al. 2012)             |
| Hepatitis C and hepatitis B virus   | SELEX      | *Hepatitis C virus*                         | (Kumar et al. 1997)                |
| Human immunodeficiency virus        | In vitro selection | Human immunodeficiency virus type-1      | (Bozioua et al. 1999)             |
| Influenza virus                     | SELEX      | Hemagglutinin protein of human influenza virus B | (Gopinath et al. 2005) |
| Severe Acute Respiratory Syndrome (SARS) coronavirus | SELEX  | Live cell                                   | (Jang et al. 2008)                |
| Trypanosoma spp.                    | SELEX      | Plasma of *T. cruzi* infected mice          | (Nagarkatti et al. 2014)           |
| Leishmania spp.                     | SELEX      | Live cell                                   | (Guerra-Pérez et al. 2015)         |
| Plasmodium spp.                     | –          | *P. falciparum* para-sites                  | (Cheung et al. 2018)               |
| Cryptosporidium parvum              | SELEX      | Fresh fruits                                | (Iqbal et al. 2015)                |
| Entamoeba histolytica               | SELEX      | Live cell                                   | (Ospina-Villa et al. 2015)         |
| MCF-7 breast cancer cells           | –          | Target cancer cells                         | (Wang et al. 2015)                 |
| Leukemia CCRF-CEM cells             | –          | Human leukemia CCRF-CEM cells               | (Ye et al. 2015)                   |
| Metastatic tumor tissues            | Cell-SELEX | Colon cancer cell SW620                     | (Li et al. 2015)                   |
| Ochratoxin A (OTA)                  | SELEX      | Immobilized OTA                             | (Cruz-Aguado and Penner 2008)      |
| Bacterial endotoxins                | SELEX      | Lipopolysaccharide                          | (Kim et al. 2012)                  |
| Copper                              | –          | Lake samples                                | (Chen et al. 2011)                 |
| Arsenic                             | –          | Aqueous solution                            | (Orovval et al. 2017)              |
| Acetamiprid                         | –          | Wastewater and tomatoes                     | (Fan et al. 2013)                  |
| Herbicides                          | SELEX      | Atrazine                                    | (Sinha et al. 2010)                |
| Milk allergen                       | SELEX      | β-LG variants A and B                       | (Eissa and Zourob 2017)            |
| Bisphenol A                         | –          | Aqueous solution                            | (Chen et al. 2017)                 |
| Beta1-adrenoreceptor autoantibodies | –          | Serum of patients                           | (Wallukat et al. 2016)             |
| Lung cancer                         | SELEX      | Cells                                       | (Bates et al. 2009)                |
| Colorectal cancer                   | –          | Camptothecin loaded-pegylated dendrimer     | (Aliboland et al. 2017)            |
| Breast cancer                       | –          | Breast cancer tissues                       | (Wang et al. 2017)                 |
the formation of a tertiary structure with the target cell that has no affinity for gold nanoparticles, resulting in salt-induced aggregation.

De Girolamo et al. (2011) developed a DNA aptamer to detect OTA (Ochratoxin A) mycotoxin produced by Aspergillus ochraceus and Penicillium verrucosum, found in wheat. They showed a system able to detect OTA in a range from 0.4 to 500 ng. Chen et al. (2015) reported the direct detection of FB1 (fumonisin B1) in maize samples by using gold nanoparticles conjugated to modified aptamers (5′-SH-(CH₂)₆-AGCACGACAGGTTGATCGGATCTGGATATTATTTTGGATACCCCTTGTGGGAACATCC TATGCGTGCTACCCGTGA-3′). The authors reported an accrued detection after 40 min at room temperature for FB1 concentrations above 2 pM. Another mycotoxin that is toxic to humans is zearalenone (ZEN). It is found in cereal crops and produced by Fusarium graminearum (Luo et al. 2020). To detect ZEN, the mycotoxin was extracted of cereal crops and different solutions were prepared and analyzed by the aptamer conjugated with zinc oxide-nitrogen doped graphene quantum dots (ZnO-NGQDs), which was capable to detect 3.3 × 10⁻¹⁴ g.mL⁻¹ (Luo et al. 2020).

Heavy metals present in milk and dairy products, fish, eggs, oils, and seeds can be also detected by aptamers. Hazardous metals, such as arsenic and mercury, can affect human health by interfering with the central nervous system and endocrine system. Thereby, colorimetric detection by metallic nanoparticles, such as the use of biosensors. Fan et al. (2013) developed a DNA aptamer conjugated with gold nanoparticles to cadmium and reported high-affinity detection for an aqueous solution containing cadmium at lower concentration (4.6 nM).

The green malachite fungicide is widely used in aquaculture and can contaminate fish and their eggs, posing a risk to those who consume them (Stead et al. 2010). In 2010, it was reported the first malachite green (MG) detection by RNA aptamer (5′-GGAAUCCCGACUCGGCGAGGC CAGGUAACGAUGGAUCC-3′) in fish skin samples, and the developed approach was able to quickly confirm the contamination after 15–20 min at 2 μg.kg⁻¹ of salmon tissue (Stead et al. 2010).

Many types of pesticides are used to prevent contamination by bacteria, fungi, and viruses, and the detection methods should be efficient even at low pesticide concentrations. However, detection by liquid and gas chromatography are expensive and time-consuming (Fan et al. 2013). To reduce the costs and analysis time, new technologies are employed, such as the use of biosensors. Fan et al. (2013) developed a conjugated aptamer with a gold nanoparticle that generates a signal of impedance to identify acetamiprid with a detection limit of 1 nM in wastewater and tomatoes and the process takes up to 3 h.

For herbicides, widely used in the cultivation of corn and oilseed rape, with the consequent environment and human life issue, different DNA aptamers with affinity to atrazine were studied by Williams et al. (2014). The double-stranded DNA aptamer for fluorescence detection was drawn to detect fipronil insecticide in river water samples and showed high sensitivity (Hong et al. 2018).

Moreover, the C07 aptamer was developed to detect Sudan dye III in chili sauce. From the study, it was reported a fast and accurate binding to the target, and according to the authors, 100 nM of aptamer was enough to detect 400 ng of Sudan dye III (Wang et al. 2018). Besides, organic molecules, such as bisphenol A (BPA), present in some food products are harmful to the human endocrine system, and in 2010, the US and Canadian governments banned their use (Mckeague et al. 2009). In this sense, Lee et al. (2011) studied the detection of BPA by aptamers conjugated with carbon nanotube as a biosensor and showed a detection limit at low concentrations (10 fM).

In this context, Smart et al. (2020) reported several promising biosensors for agribusiness. In some cases, aptamers were conjugated to nanoparticles forming carbon electrodes for detection of pesticides, toxins, antibiotics, microorganisms, vitamins, fructose, and lactate. In addition, Yan et al. (2020) showed different photoelectrochemical and electrochemiluminescent apta-sensors capable to detect food contaminants and pollutants.

Conclusions and market perspectives

There are several studies in the literature related to the development of aptamers for different targets. They are widely studied in the therapeutic area, to identify cancer cells, bacterial contamination, and viruses. Moreover, aptamers are being developed for food safety since there is a high diversity of pathogens in food products from different origins (animal, vegetable, processed), as well as contamination by packaging or transportation.

Despite being basic, inexpensive and selective, most existing aptamers are still not currently used in the industries and agribusiness routine. Through the aptamer applications, a quick analysis system can be launched as a biosensor, bringing advantages to the market as visual detection, low-cost compared with conventional techniques, as well as delivering robustness and selectivity.

Chemiluminescence detection is one of the most studied technics since it does not require equipment for signal detection. However, for many targets, studies are needed to ensure that sensitivity and specificity are enhanced using
nanoparticles. Further investigation should be directed to sample preparation methods.

Many of the aptamers are not able to recognize samples in the raw phase and need to be prepared in aqueous solutions. To reduce the gap between lab-scale and industrial large-scale applications, advances in fast and efficient detection for food safety are increasing, but some aspects still need to be improved, such as sample preparation, concentration, and the presence of contaminants from raw materials.

Due to the difficulties related to food safety, the companies are looking for cheaper and faster alternatives. Whereas the world population is expected to reach 8.5 billion people by 2030 (United Nations 2019), the food industry faces problems with changes in food production and supply, increased imports, changes in the environment that lead to contamination, development of outbreaks, or pests on crops in different locations. Herewith, the market is turning to faster and more effective sensors to prevent detect contamination. In this context, aptamers come to the market as an alternative (Liu and Zhang 2015).

According to their advantages, aptamers can supply the agribusiness as well as food industrialization needs, being increasingly used worldwide to speed up the food safety analysis, to avoid products recall and unnecessary business expenses around the world (Amaya-González et al. 2013; Aptamer Group 2016).

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