CRITICAL REVIEW



Cite this: Analyst, 2022, 147, 1006

Paving the way towards continuous biosensing by implementing affinity-based nanoswitches on state-dependent readout platforms

Annelies Dillen 💿 and Jeroen Lammertyn 💿 *

Continuous biosensors provide real-time information about biochemical processes occurring in the environment of interest and are therefore highly desirable in research, diagnostics and industrial settings. Although remarkable progress has been made in the field of biosensing, most biosensors still rely on batch processes and, thus, are not suited to perform continuous measurements. Recently, however, it has been shown that by combining affinity-based nanoswitches with state-dependent readout platforms, the necessity for batch processes can be overcome and affinity-based continuous biosensing can be achieved. In this review, we first provide an overview of affinity-based continuous biosensing and discuss the required components to achieve this goal. More specifically, we summarize the strategies that have been applied to develop and tune both protein and nucleic acid-based switches, as well as readout strategies that can be applied in combination with the former. Afterwards, biosensors in which both elements were already integrated and hence enabled continuous measurements are reviewed. We also discuss the challenges and opportunities associated with each approach and therefore believe this review can help to encourage and guide future research towards continuous biosensing.

Received 22nd December 2021. Accepted 15th February 2022

DOI: 10.1039/d1an02308j

rsc.li/analyst

Introduction 1.

Real-time, continuous biosensing is gaining more and more attention in healthcare,^{1,2} biotechnology,^{3,4} and food and beverage processing,^{5,6} as it provides instant and continuous information about biochemical processes occurring in the

KU Leuven, Department of Biosystems - Biosensors Group, Willem de Croylaan 42, Box 2428, 3001 Leuven, Belgium. E-mail: jeroen.lammertyn@kuleuven.be



Annelies Dillen

Annelies Dillen obtained a Master in Bioscience engineering (KU Leuven) in 2018 for which she submitted a thesis on the development of digital assays for single exosome detection in cancer diagnostics. In the same year she started her PhD in the Biosensors group in the division Mechatronics, Biostatistics and of the Sensors (MeBioS) Biosystems department at KU Leuven, where she is currently working on fiber optic SPR

sensors combined with DNA nanotechnology, to develop a new concept for continuous biosensing.



Jeroen Lammertyn

Jeroen Lammertyn is full professor in bionanotechnology at the KU Leuven. He founded the Biosensors group in 2005. The group closely follows the emerging field of food and medical diagnostics and is active in bioassay development, optical biosensors, and microfluidics. He is (co-)author of >300 peer reviewed research papers, conference papers and book chapters. He acts as reviewer for many international peer reviewed jour-

nals, is co-inventor of 10 patent applications and founder of the spin-off company FOx Biosystems. He is also chairman of the DIATECH 2014 and 2020 International Conferences on Novel technologies for in vitro Diagnostics in Leuven.



View Article Online

Critical Review

environment of interest, thereby allowing direct and even feedback-controlled intervention.⁷ Furthermore, by continuously monitoring target molecules in their respective environment, the necessity for batch measurements can be overcome, which is often a tedious and time-consuming process, as samples frequently have to be withdrawn, processed, transported to the sensing setup, and analyzed before information about the target concentration can be obtained.⁷ Although remarkable progress has been made in the field of biosensing since the development of the first biosensor in 1956,⁸ most biosensors do not satisfy the requirements for continuous biosensing, being that they (i) must not require externally added reagents, (ii) cannot generate products, (iii) cannot rely on multistep processes such as washing, incubation or regeneration steps, (iv) must exhibit a dynamic response and (v) must be stable for a long time period in the environment of interest.^{7,9} Glucose and other enzymatic sensors are the best-known examples of biosensors that are capable of continuously monitoring specific target concentrations,¹⁰ but as their detection strategy relies on the reactivity of the target itself, it cannot be applied for non-reactive targets.^{7,9,11} Affinity-based biosensors on the other hand, represent a more general approach for biosensing. However, they generally rely on multistep processes including washing, incubation or regeneration steps to translate the binding of an analyte to the bioreceptors into a detectable output, which limits their use for continuous biosensing.^{7,9} Recently, it was shown that by combining affinity-based nanoswitches with a state-dependent readout platform, the aforementioned limitations could be overcome and affinity-based continuous biosensing could be achieved.¹¹⁻¹⁷ Therefore, in this review, we will discuss both crucial components separately and will explain how these can be applied to develop affinitybased continuous biosensors.

First, we explain the concept of affinity-based nanoswitches in terms of continuous biosensing and provide an overview of the approaches to transform affinity-binders into the former. Second, we review readout strategies that are suitable to be applied in combination with the affinity-based nanoswitches, to develop an integrated biosensor for continuous monitoring. Finally, we discuss the platforms that have already integrated both elements, thereby overcoming the abovementioned challenges and enabling continuous biosensing.

2. Elements to enable affinity-based continuous biosensing

2.1. Affinity-based nanoswitches

Affinity-based nanoswitches are entities, composed of affinitybinders, that undergo a reversible conformational change upon target binding. Due to the conformational change, target binding can directly be translated into a detectable output (by using a state-dependent readout platform as will be explained in the next section), without multistep processes or exogenous reagents. Furthermore, as this conformational change is reversible, the switch can react to changing target concentrations and a dynamic response can thus be achieved.^{7,9} In this manner, the abovementioned limitations can be overcome and therefore, affinity-based nanoswitches are key elements for continuous biosensors. The development and use of these switches is inspired by nature, where switching molecules sense and react to biological cues in a variety of ways. The amount of targets for which naturally occurring switches exist is however limited, and therefore, strategies have been described to develop synthetic switches from affinity binders, which we subdivide here in the following categories.¹⁸⁻²⁰

The first approach is the development of a clamp, as shown in Fig. 1A and 2A, for which two affinity-binders are tethered together, so the switch will shift from a loose to a more rigid state when both receptors are bound to the target. In order to develop a switch using this approach for multivalent targets, two copies of the same binder can be used, as they can bind to different subunits of the target molecule.²¹ For monovalent targets however, either two affinity binders that can bind to distinct epitopes of the target molecule, or one affinity binder recognizing the target molecule and a second one binding to the target-receptor-complex, are required.^{18,20,22} This strategy is highly modular and thus allows for straightforward design

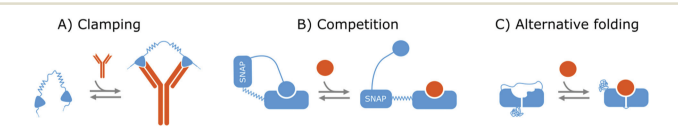


Fig. 1 Strategies to create protein switches. (A) Clamping: Two protein binders are tethered together *via* a flexible linker. When the target is present, both will bind the target and a rigid structure will be formed. (B) Competition: The protein binder is linked to an intramolecular ligand *via* a SNAP-tag and stiff linker. In this manner, the target will compete with the ligand, leading to its displacement and thus an extended conformation of the protein switch. (C) Alternative folding: Part of the protein binder is duplicated and linked to its opposite end so the protein can fold into two distinct conformations. Furthermore, a mutation is inserted into the duplicated part and as a result, the target can only bind one of both conformations. In this way, the protein binder can switch between its two conformations of which one is stabilized by target binding. A protein switch can also be developed *via* alternative folding by destabilizing the protein structure, so folding is only achieved in presence of the target molecule. For all strategies, the switches are depicted in blue while the target molecules are portrayed in orange.

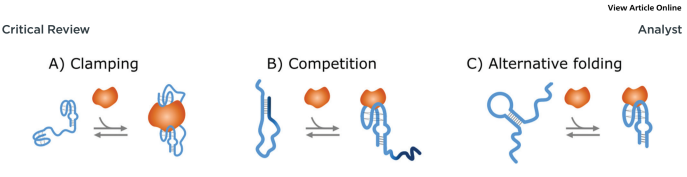


Fig. 2 Strategies to create NA switches. (A) Clamping: Two NA receptors are tethered together *via* a flexible linker. When the target is present, both will bind the target and a more rigid structure will be formed. (B) Competition: The NA receptor (light blue) is linked to a NA competitor (dark blue). In this way, the target will compete with the competitor, leading to its displacement and thus a conformational change of the NA switch. (C) Alternative folding: The secondary structure of the NA receptor to which the target binds is destabilized or an alternative folding of the receptor is introduced. In this manner, the NA receptor can switch between its two conformations of which one is stabilized by target binding. For all strategies, the switches are depicted in blue while the target molecules are portrayed in orange.

of affinity-based nanoswitches.²¹ However, as small ligands cannot easily be detected using clamping due to their limited recognition domains, affinity-based nanoswitches have also been engineered based on competition, which relies on the attachment of a competitor to the bioreceptor (Fig. 1B and 2B). The competitor will bind to the target recognition zone, so when the target is present, it will compete with and lead to the displacement of the competitor and thus conformational change of the switch. Next to the modularity and generalizability of this approach, the overall affinity of the switch can be rationally tuned without requiring changes to the native bioreceptor itself, by adjusting the binding strength of the competitor. However, by using a competitor, the overall affinity of the target towards the switch will be lower than its affinity for the native bioreceptor.^{18,20,23-26} Lastly, synthetic switches can be developed by introducing an alternative folding, as depicted in Fig. 1C and 2C. In this strategy, the affinity-binder needs to be adapted so it can fold into two well-defined structures, of which one is stabilized by target binding. As for this approach, the bioreceptor itself needs to be tailored to undergo a conformational change upon target binding, the design process is often challenging and requires empirical optimization, which is why this strategy is less generally applicable.^{18,20,24,25,27}

Using these strategies, both protein and nucleic acid (NA) switches have been developed. As both molecules are structurally and chemically different, the engineering of protein or NA switches requires a different expertise and is even conducted in different research fields (*i.e.* protein engineering and DNA nanotechnology, respectively). Therefore, the abovementioned strategies to create either protein or NA switches will be discussed in more detail separately. Afterwards, the development of NA-protein hybrid switches will be discussed shortly.

Importantly, for affinity-based switches suitable for continuous monitoring, all components of the switch need to be linked together, as no components can be added as external reagents. Therefore, we will only discuss the switches that fulfill this requirement.

2.1.1. Protein switches. Protein switches have become invaluable tools in both fundamental and applied research, as they provide a means to report or respond to desired biological cues in real-time. Therefore, in the field of protein engineer-

ing, much research is being performed to tailor-engineer protein switches with custom input and output functions.^{18,28} Although many protein switches have been described that are able to act as transducers or actuators,^{18,20} we will only discuss protein switches that undergo a conformational change as output since they fulfill the requirements for continuous biosensing.

To engineer such protein switches, the abovementioned strategies have been widely applied. For instance, proteinbased clamps have been constructed for antibody detection by the group of Maarten Merkx by connecting two copies of the respective antibody binding epitope, as presented in Fig. 1A. This construct was developed for antibodies against HIV (anti-HIV1-p17),²⁹⁻³² EGFR (cetuximab),^{30,33} hemagglutinin^{31,32} and dengue virus (type I),^{31,32} thereby demonstrating the suitability of this approach for antibody detection. Notably, the binding epitopes were connected by a glycine-serine linker, which are commonly applied flexible spacers, and are therefore desirable to develop a protein clamp that switches between a flexible and rigid state upon target binding.34 The abovementioned strategy was also applied by Adamson et al. by connecting two Affimer proteins³⁵ that bind to human C-reactive protein (a protein biomarker), Herceptin (a therapeutic antibody) or the cow pea mosaic virus (a plant virus) respectively, thereby further demonstrating the generalizability of this approach to develop protein switches for multivalent targets.²¹ Moreover, protein clamps have also been developed for various peptides, by sandwiching them between two distinct peptide binding interfaces that were selected through directed interface evolution.³⁶⁻³⁸ Additionally, a Zn(II)-binding clamp was developed by connecting two metal-binding domains (Atox1 and WD4).³⁹⁻⁴¹ Altogether, clamping proved to be a straightforward and general strategy to develop protein switches for multivalent targets, but remains less widespread for small and monovalent targets, due to the need for two distinct epitopes and affinity binders.

Therefore, protein switches have also been engineered based on competition as depicted in Fig. 1B. To achieve this, the group of Kai Johnsson covalently attached protein binders to their ligands *via* an intramolecular tether and SNAP-tag (a commercially available self-labeling protein tag), so when the

target molecule is present, the intramolecular ligand is displaced by the target and the protein switch adopted an extended conformation.⁴² Importantly, to ensure an extended conformation upon target binding, the linker between the binder and SNAP tag should not be too flexible. Therefore, although glycine/serine linkers proved useful for the development of clamps, polyproline linkers were applied in this format, as they are more rigid and thus better suited for the development of competitive protein switches. Furthermore, the intramolecular tethers should be long enough to allow binding of the intramolecular ligand and thus closing of the switch. Another important consideration is the affinity of the protein binder for its intramolecular ligand, as this should be high enough to ensure the closed state is predominant in the absence of target molecules, whereas it should be low enough to allow competition with the respective target, for the desired target concentration range. As a result, the overall affinity of the protein switch for its target depends on the affinity between the applied protein binder and the intramolecular ligand as well as the affinity between the protein binder and its target, of which the first can be tuned to adjust the concentration range to which the switch will respond. Using this strategy, the research group of Kai Johnsson was able to develop protein switches for the detection of small molecules such as glutamate,⁴³ acetylcholine,⁴⁴ sulfonamides,^{42,45} γ -aminobutyric acid⁴⁶ as well as for clinically relevant drugs including anti-epileptics (topiramate),⁴⁷ immunosuppressants (sirolimus, tacrolimus and cyclosporine A),⁴⁷ anti-arrhythmics (digoxin),⁴⁷ anticancer agents (methotrexate),^{47,48} bronchodilators (theophylline)⁴⁸ and drugs against malaria (quinine).⁴⁸

Furthermore, as many naturally occurring protein switches are based on alternative folding, this strategy has also been applied to develop their synthetic counterparts.¹⁸⁻²⁰ In particular, to engineer protein switches using alternative folding, either the structure of the protein can be destabilized, so the protein only folds upon target binding,⁴⁹ or the topology of the polypeptide chain of the respective protein binder itself can be altered, so the protein can fold into two distinct conformations, of which one is stabilized by target binding. For instance, Kevin Plaxco and coworkers inserted switching properties in the FynSH3 protein using the first alternative folding strategy. More specifically, they truncated the respective protein (four-residue carboxy-terminal truncation), which led to the destabilization of FynSH3 in absence of the target molecule. As a result, the population FynSH3 successfully shifted from an unfolded to a folded conformation upon binding of the VSL12 peptide target.^{49,50} To induce an alternative folding using the latter strategy, the research group of Stewart N. Loh duplicated part of the protein that recognizes the target molecule and attached it to the opposite side of the protein allowing its folding either into its native or alternative conformation. Furthermore, they inserted a mutation in the duplicated segment, so target molecules could only bind to the native state and as a result, an equilibrium was formed between the two conformational states of the protein binder, of which one (native) was stabilized by binding of the target

molecule, $^{28,51-54}$ as presented in Fig. 1C. Using this approach, protein binders were successfully converted into protein switches for the detection of Ca(II)^{51,53} and ribose.⁵⁴

Notably, when engineering protein switches, most often the reporter molecules and switching properties are engineered into the protein binder simultaneously. However, as the respective labels depend on the applied readout strategy, they will be discussed separately in section 2.2.

Although the broad chemical diversity of amino acids provides a wide functionality to proteins, which enabled the development of protein switches with various in- and outputs, due to this diversity, the behavior of proteins cannot be easily predicted. As a result, the development of protein switches can be challenging and commonly relies on extensive empirical optimization and is often limited to binders for which structural information is available.^{18,20}

2.1.2. Nucleic acid switches. Due to the emerging field of DNA nanotechnology in which NAs are used for engineering purposes rather than as a genetic carrier, NAs have extensively been used to develop affinity-based nanoswitches.⁵⁵ Although the chemistry and structure of nucleic and amino acids differs significantly, the same strategies to convert NA receptors into switching molecules can be applied.

In fact, NA switches have been engineered by tethering two aptamers or single-stranded NA (ssNA) probes together, via a flexible linker. This was achieved either through covalent attachment, using DNA,^{22,56-59} RNA,⁶⁰ PEG,^{59,61} phosphoramidite linkers⁶² or larger scale structures,¹¹ or through hybridization of ssNA linkers.⁶³ In this manner NA clamps were successfully formed that switched from a flexible to a more rigid state upon target binding,^{64–66} as shown in Fig. 2A. Using this clamping strategy, NA switches have been developed for protein targets including the hepatitis C virus non-structural protein 3,⁶⁰ vitronectin,⁶³ thrombin^{11,22,56–58,61,62} and vascular endothelial growth factor^{22,59} as well as DNA targets.¹¹ Although the design and development of NA clamps proved to be straightforward, their application remains limited and is mostly applied to thrombin, as this is the best-known example for which two well established aptamers targeting distinct epitopes are available.67

In contrast to clamping, competition is a widespread strategy for the development of NA switches from aptamers, as the predictability of Watson-Crick basepairing provides a straightforward approach to design and tune a competitor, consisting of ssNA that is partially complementary to the target binding region of the aptamer,^{23,26} as presented in Fig. 2B. Importantly, in contrast to many protein switches developed via competition, the competitors of these NA switches thus do not consist of the respective (modified) target itself. These switches were termed duplexed aptamers in a previous review paper by Munzar et al. (2019)²³ and have been developed by linking the competitor NA strands and aptamers again either covalently using ssNA spacers⁶⁸⁻⁹⁶ or PEG,⁹⁷⁻¹⁰⁰ or *via* hybridization to NA linkers^{101,102} or even DNA origami structures.¹⁰³ Using this strategy, NA switches have been developed for the detection of various targets including thrombin,68,71-74,102,104

protein tyrosine kinase-7,74-78,100 immunoglobin heavy mu chain,⁸⁹ nucleolin,^{86,103} theophylline,⁸⁸ platelet-derived growth factor (BB),^{84,85} lysozyme,^{81,82} IFN-Y,^{69,80,90} TNF- α ,⁶⁹ IgE¹⁰¹ ATP,^{70,87,91–99} AMP,^{79,81} ADP⁸³ and GTP.⁸³ This broad availability proves the generalizability of this approach for a wide range of targets. Note that these only represent the duplexed aptamers in which the competitor is linked to the aptamer itself, but that many more duplexed aptamers have been described in which the competitor is added as an external reagent which further demonstrates their widespread application.23,26,105 However, the latter do not fulfill the requirements for continuous biosensing and thus will not be discussed further. Importantly, as was also the case for protein switches developed by competition, the binding strength between the competitor and NA receptor needs to be carefully tuned. Therefore, in our previous work, we developed a model that predicts how the competitor affects the final switching properties of duplexed aptamers.²⁶ This model allows to rationally tune the performance of duplexed aptamers and thereby reduce the amount of empirical optimization required to develop new NA switches from NA bioreceptors based on competitive displacement. Furthermore, although most competitive NA switches were developed starting from a previously described aptamer, Nutiu et al. adapted the standard SELEX procedure (systemic evolution of ligands by exponentially enrichment), which is commonly used for the in vitro selection of aptamers for specific targets, to directly select NA switches based on competition from random sequences. Using this strategy, they successfully selected NA switches for the detection of ATP and GTP.¹⁰⁶ Altogether, the development of NA switches based on competitive displacement proved to be a straightforward and general strategy suitable for a variety of target molecules.

Finally, NA switches have been constructed based on the introduction of an alternative fold from both ssNA probes¹⁰⁷ and aptamers,^{24,27} commonly described as molecular beacons or structure-switching aptamers (SSAs) respectively, of which the latter is depicted in Fig. 2C. To develop SSAs, the group of Kevin Plaxco either destabilized the tertiary structure of the respective aptamer or introduced an alternative fold, similar as was described for protein switches.²⁷ Notably, to do this, the tertiary structure of the aptamer needs to be known. Using this strategy, they were able to develop NA switches for the detec-tion of proteins,^{108,109} drugs^{12,15–17,110–117} and other small molecules,¹¹⁸ including thrombin,¹⁰⁸ the TATA binding protein (transcription factor),¹⁰⁹ doxorubicin,^{12,17,112,113} kanamycin,^{12,17,114} tobramycin,^{16,17,115,117} gentamycin,¹⁷ irino-tecan,¹¹⁰ vancomycin,¹¹¹ cocaine^{15,112,114} and melamine.¹¹⁸ Although SSAs have successfully been developed using this strategy, the engineering process remains unpredictable and is mostly based on trial and error, thereby undermining the widespread adaptation of this type of NA switch.²³ Molecular beacons are commonly applied to detect NAs, due to their ease in design and development, as all interactions are based on hybridization. They were first described by Tyagi and Kramer in 1996¹¹⁹ and are composed of a ssNA probe equipped with hairpin formation sequences on each end. In this manner, a

hairpin loop is formed in absence of the NA target, which opens when the respective target sequence binds. This format has been widely applied throughout the years and proven its applicability for the detection of NA.¹²⁰⁻¹²⁸

Altogether, NAs are suitable components for the development of affinity-based nanoswitches due to their ease of synthesis and use, and their highly predictable thermodynamic behavior.^{23,129,130} Furthermore, due to the progress in aptamer selection processes (SELEX), the amount of NA receptors for various targets keeps increasing.^{131–133} Consequently, NA switches have already been widely applied for bioassay development^{55,134,135} and play an important role in the development of continuous biosensors.

2.1.3. Nucleic acid–protein hybrid switches. As explained in the previous sections, both protein and NA switches have been developed to respond to molecular cues in real-time. Importantly, as NA and proteins are inherently different, the opportunities and potential of both components for the development of affinity-based nanoswitches differs significantly. However, by merging NA and protein molecules, the advantages of both components can be combined and NA–protein hybrid switches can thus be engineered with a broad functionality and high predictability. For instance, the wide range of protein binders can be implemented for target recognition, whereas NA can be used to implement the conformational change after target binding.^{136,137}

In a similar format as the protein clamps developed by the group of Maarten Merkx (shown in Fig. 1A), the group of Francesco Ricci developed an NA-protein hybrid switch for the detection of the anti-HIV and anti-flag antibodies as presented in Fig. 3A. Therefore, they designed a DNA hairpin with ssDNA tails, that are hybridized with ssDNA equipped with the ligands of the respective antibodies. In this manner, when the target antibodies are present, they will bind to their respective ligands and thereby open the DNA hairpin.¹³⁸ With this format, they could achieve the same advantages as discussed in section 2.1.1. for the protein clamps developed by the group of Maarten Merkx. On top of that, their format allows (i) straightforward control over the conformational change and (ii) tuning of the switch thermodynamics, by altering the basepairing of the hairpin stem, which further demonstrates the potential of NA-protein hybrid switches.^{137,138} Furthermore, they showed that this format can be applied for the detection of monovalent targets (anti-Dig Fab fragment and DNAbinding transcription factor TBP), as well, on the condition that the simultaneous binding of two target molecules causes enough steric hindrance to open the DNA hairpin.138

Aside from the abovementioned NA–protein hybrid clamps, a competitive hybrid switch has also been developed by the research group of Menno Prins. Herein, DNA was applied both as a tether to link a microparticle to the sensor surface, as well as a linking group, to attach the competitor of the protein bioreceptors,^{14,139} as shown in Fig. 3B. This switch was developed for creatinine and will be elaborated further in section 3.2, where we will discuss biosensing strategies that applied affinity-based nanoswitches to enable continuous measurements.

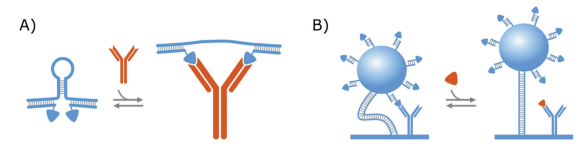


Fig. 3 Strategies to create NA-protein hybrid switches. (A) Clamping: Antibody ligands were attached *via* hybridization to a DNA hairpin. In this manner, when target antibodies were present, they would bind to their respective ligands and the DNA hairpin would be opened. (B) Competition: Switches are composed of microparticles, which were immobilized on the surface *via* dsDNA tethers. The microparticles were equipped with a (creatinine) competitor *via* hybridization, whereas the surface was equipped with protein bioreceptors (antibodies). In this manner, when the target molecule (creatinine) was present, they would bind to the protein bioreceptors and the switches became less restricted in their movement. For all strategies, the switches are depicted in blue while the target molecules are portrayed in orange.

The abovementioned examples clearly show how the advantages of proteins and NAs can be combined for the development of affinity-based nanoswitches. Notably, to do this, knowledge about both protein engineering and DNA nanotechnology is required, which might be a reason why the amount of NA-protein hybrid switches remains limited. In summary, although the potential of NA-protein hybrid switches is in our opinion currently still underexplored, we believe that this strategy can further help in broadening the library of available affinity-based nanoswitches and provides a highly versatile and modular technique for the development of affinity-based switches, suitable for continuous biosensing.

2.1.4. Dynamic response. Although many affinity-based nanoswitches have been developed and described in literature, most of them have only been studied in equilibrium conditions rather than in a continuous manner. In order to apply them for continuous biosensing applications however, it is important that the kinetic behavior of these switches is evaluated, as this will ultimately determine the dynamic response of the biosensor. Generally, there is a trade-off between the affinity for the target and the dynamic response of the biosensor. On one hand, a high affinity is considered advantageous as it generally provides a low detection limit.⁷ More specifically, the affinity between the switch and target will determine how many target molecules will bind to the switches and thus how many state-changes will occur for a certain target concentration. As for most readout platforms, the amount of state-changes is directly proportional to the obtained signal, a higher sensitivity and lower limit of detection (LOD) can usually be achieved for higher affinity capture elements. On the other hand, a high affinity generally provides a slower dynamic response. Indeed, as the dissociation constant, which is inversely correlated with the affinity, is defined as:140,141

$$K_{\rm D} = \frac{k_{\rm off}}{k_{\rm on}}$$

with $K_{\rm D}[{\rm M}]$ the dissociation constant and $k_{\rm on}\left[\frac{M}{s}\right]$ and $k_{\rm off}\left[\frac{1}{s}\right]$ the association and dissociation rate, respectively, a lower dis-

sociation constant is associated with a lower dissociation or higher association rate and thus less dissociation of the switch-target complex.¹⁴⁰ Consequently, for conditions in which the reaction rates are the limiting step of the (un)binding, the higher the affinity, the longer it will take for a new equilibrium to be formed and thus the slower the dynamic response. Notably, here, K_D represents the overall dissociation constant of the switch for its target, which depends on but is not the same as the dissociation constant of the applied bioreceptor for its target.

As both the sensitivity and dynamic response are important parameters for the performance of a continuous biosensor, attention should be paid to this trade-off during the design process of the affinity-based nanoswitches, for which the timeresolution and sensitivity that is required for the respective application should be taken into account. Notably, the choice of the strategy to achieve switching will affect the performance of the continuous biosensor and will even more determine how the abovementioned trade-off can be tuned. For example, when creating an affinity clamp, the overall binding strength between the switch and its target is generally increased compared to the binding strength of the stand-alone bioreceptors due to their increased avidity, which is aside from switching properties a main reason for the development of many clamps.^{22,64} Furthermore, for both protein and NA clamps, it was shown that by altering the length or stiffness of the linkers that attach both binders, the affinity of the switch can be tuned.^{34,39,142,143} More specifically, the length and stiffness of the linkers will affect both the magnitude of the conformational change as well as the local effective concentration of the respective binders, which both affect the final signal that is generated for a certain target concentration.^{34,39} This strategy was applied to enhance the affinity of protein and NA clamps for the detection of Ca(n),¹⁴⁴ K(n),¹⁴⁵ Zn(n)¹⁴² and immunoglo-bin heavy mu chain,¹⁴³ by screening different lengths and of glycine-serine compositions and **PEG-linkers** respectively.142-145 In this manner, the affinity could be enhanced from the pM to fM,¹⁴² low to mid nM,¹⁴⁴ low to mid mM and low to mid μM^{143} range for the Zn(II),¹⁴² Ca(II),¹⁴⁴

 $K(i)^{145}$ and immunoglobin heavy mu chain¹⁴³ binding switches respectively.

In contrast, by using a competitor, the overall affinity of the resulting competitive switches will be lower than the affinity of the applied native bioreceptor. However, this strategy allows rational and straightforward tuning of the final switch performance, as it was previously described how the final performance of competitive switches can be tuned by altering the binding strength between the bioreceptor and competitor. As a result, the overall affinity of competitive switches can be tuned by adapting the competitor's affinity for the bioreceptor.^{25,26} For example, Xue *et al.*⁴⁸ modified the structure of the intramolecular ligands in competitive protein switches for antibodies against methotrexate, theophylline and quinine, to lower their LOD. They did this by taking advantage of structural information available on the interaction between the antibodies and drugs, and in this manner could specifically remove or shift the position of the groups taking place in the binding.⁴⁸ As a result they were able to enhance the affinity from low to mid µM range for all three antibodies. Notably, the binding strength between the bioreceptor and competitor is not only dependent on the affinity between both, but also on their local effective concentration, meaning that the length or stiffness of the linker attaching both can also be changed to tune the overall affinity,²³ similar as for clamping. In fact, Wilson et al.¹⁴¹ successfully developed a range of competitive NA switches with distinct affinities for the detection of ATP (ranging from 10 µM to 40 mM) by screening both (i) different linker lengths between the aptamer and competitors (ranging from poly T(23) to poly T(43)) and (ii) different lengths of competing NA strands (ranging from 6 to 9 complementary nucleotides).141

Lastly, when an alternative fold is introduced to develop affinity-based nanoswitches, the overall affinity of the switch for the target can be tuned by adjusting the equilibrium between the native and the alternative conformational state of the switch, as both states have a distinct affinity for the target, by stabilizing or destabilizing either state (e.g. by inducing mutations in the oligo or peptide sequence).27,52 This was done by Porchetta et al.,¹⁴⁶ who destabilized the folded state of a SSA for cocaine, so the equilibrium would shift more to the unfolded state. Therefore, based on the secondary structure of the SSA, which is predicted to be a three-way junction, they designed three SSA variants in which either (i) mutations were inserted, (ii) the terminal stem was shortened or (iii) the sequence was circularly permutated. Notably, the target binding region was kept conserved throughout the variants to ensure target binding was not hindered. In this manner, they obtained three SSA variants with a distinct, lower affinity ($K_{\rm D}$ of 23, 82 and 1390 μ M) compared to the original SSA ($K_{\rm D}$ = 0.5 µM).146 Similarly, as the equilibrium of their FynSH3 protein switch shifted almost completely to the folded state at high ionic strength conditions (present in whole blood), Kurnik et al.49 destabilized the folded state of the respective switch by performing an I50L substitution in the hydrophobic core of the protein. In this manner, they successfully altered

the overall affinity and obtained target-responsiveness at high ionic strength conditions.⁴⁹ Importantly, by introducing an alternative fold, the overall affinity of switches developed through this strategy will be lower compared to the affinity of the applied bioreceptor,¹⁴⁷ similar as for competitive switches.

In summary, depending on the final application, particular care should be given to the trade-off between the sensitivity and dynamic response of the resulting biosensor, which will be partly determined by the applied strategy to transform affinity-binders into affinity-based nanoswitches. Importantly, although the affinity of the obtained switches is often characterized, this is not the case for the kinetics and dynamic response. Therefore, in order to apply these switches for continuous measurements, more research is required in the future to characterize and tune the kinetics as well.

Importantly, although we discussed the dynamic response and sensitivity of affinity-based switches, other performance parameters, such as the selectivity, are also important to take into account when developing a continuous biosensor. More specifically, as the selectivity of the affinity-based switches is mainly determined by the selectivity of the applied bioreceptors, care should be taken regarding this parameter when choosing which bioreceptor to apply.¹⁴⁸ Notably, as a thorough discussion about the selectivity of affinity-based bioreceptors is out of the scope of this review, it will not be discussed further.

2.1.5. Stability of affinity-based nanoswitches. Another important aspect of affinity-based nanoswitches for continuous biosensing is their stability. As the goal is to continuously measure target molecules directly in the environment of interest, it is needless to say that the sensor, and thus the affinitybased nanoswitches should be stable in that specific environment for the duration of the measurement.⁷ For most applications, this is composed of complex matrices such as body fluids,^{1,2,111} culture media^{3,4} or foodstuffs.^{5,6,118,149,150} Therefore, it is of paramount importance that the affinitybased nanoswitches are resistant to degradation or inactivation by the surrounding conditions or components present in the respective matrix, to avoid drift of the final output over time.¹⁴⁸ Importantly, various strategies have already been described to increase the stability of both protein and NA bioreceptors.^{151–156} More specifically, as proteins can generally fold in various intermediates, it needs to be ensured that the fold which renders the protein with the desired functionality is the most stable under the respective conditions. To achieve this, protein engineers both used rational and semi-rational approaches of which the first includes finding of the unstable regions, wherein structural features can be integrated to improve the stability of that certain region. Furthermore, sequence information can be used to design stable homologs of the protein of interest, or stable folds can be generated ab initio. In semi-rational approaches, protein stability can be improved by grafting from stable homologs or through directed evolution.156

Besides, to increase the stability of NA bioreceptors, both pre- and post-SELEX modification strategies have been

described. Notably, although an advantage of NAs compared to proteins is their high stability in various conditions, they are susceptible to nuclease degradation, and therefore, most of the reported modifications are meant to protect the NAs from the latter. More specifically, chemical modifications can be made post-SELEX to the sugar backbone, phosphate linkages or 3' and/or 5' ends, which all lead to an improved stability.^{155,157–162} Furthermore, SELEX can be performed starting from a modified NA library (pre-SELEX), such as xenobiotic nucleic acid (XNA), threose nucleic acid (TNA), locked nucleic acids (LNA), fluoroarabino nucleic acid (FANA) and hexitol nucleic acid (HNA) amongst others, which are more stable against nuclease degradation.¹⁶³⁻¹⁶⁶ Importantly, as NAs are chemically synthesized, they can be modified easily.¹⁵⁷ Notably, not only the inherent stability of NAs, but also the stability of aptamer folding is important, as their tertiary structure is responsible for target recognition. The tertiary structure of NAs is affected by the surrounding conditions such as pH, temperature or ionic strength and therefore, SELEX can be performed in those respective conditions to ensure target recognition in the environment of interest.^{131,160}

We will not provide a detailed description of the abovementioned strategies to improve both protein and NA stability here, as it is out of the scope of this review and already discussed elsewhere.¹⁵⁶⁻¹⁶¹

2.2. State-dependent readout platforms

In the previous section, affinity-based nanoswitches were discussed as an essential tool for continuous biosensing, as they respond to target binding by undergoing a conformational change.^{7,9} However, in order to translate this conformational change into a measurable output, a readout platform that is able to distinguish the different states of the switch (statedependent), is required. Furthermore, the switches need to be labeled with reporter groups, depending on the respective readout platform, as will be discussed further. In this manner, a distinct signal will be generated for the bound and unbound state, so the obtained output can be linked to the target concentration.⁹

2.2.1. State-dependent readout platforms combined with affinity-based nanoswitches. Fluorescence resonance energy transfer (FRET) is the most widely applied readout strategy in combination with affinity-based nanoswitches.55,167,168 This strategy is very well-suited to distinguish conformational changes, as the energy transfer between donor and acceptor fluorophores is highly dependent on the distance between both (effective at distances <10 nm).^{55,168,169} To utilize this approach, fluorescent reporters (donor and acceptor) have to be included in the protein or NA switch at locations known to participate in the conformational change, as represented in Fig. 4A. The total fluorescent signal is dependent on the amount of switches undergoing a conformational change and thus on the concentration of the target molecule.55,168 To perform a FRET readout, the biosensor should contain a light source to excite the respective donor fluorophore, a wavelength filter to isolate the photons emitted by the acceptor fluorophore and a detector that records the emitted photons and transduces it into a measurable output.¹⁷⁰ Consequently, the resulting biosensors rely on bulky, expensive optical equipment and therefore often are not suitable to be applied directly in the environment of interest (in situ). Furthermore, due to the optical readout, measurements cannot be performed in opaque samples (such as whole blood). As a result, although FRET proved to be a highly suitable 'state-dependent' readout strategy, it has mostly been applied in solution-based assays (NA switches)⁵⁵ or for intracellular imaging (protein switches),168 where the switches itself were applied as external reagents, and not for affinity-based continuous biosensing.

In contrast to FRET, electrochemical readout platforms have already been frequently applied in combination with

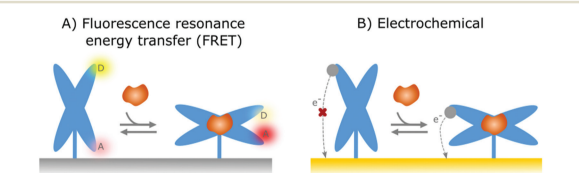


Fig. 4 Representation of the two most widely applied state-dependent readout strategies in combination with affinity-based nanoswitches. (A) Fluorescence resonance energy transfer (FRET): by labelling the switches with both a donor and acceptor fluorophore, a distinct fluorescent signal will be generated for the bound and unbound state of the switch, as the distance between both fluorophores gets altered due to the conformational change. Indeed, when the donor and acceptor fluorophores are in close proximity, energy will be transferred from the donor to the acceptor molecule, leading to a fluorescent signal originating from the acceptor molecule. As the energy transfer efficiency is highly dependent on the distance between both fluorophores, the energy transfer will decrease upon increasing distance between both and as a result, the fluorescent signal originating from the donor fluorophore will be released at higher distances. (B) Electrochemical: The affinity-based nanoswitches are labelled with a redox reporter. In this manner, when the switches undergo a conformational change after target binding, the redox reporter will be brought closer to the surface, thereby enhancing the electron transfer efficiency between the reporter and sensor surface and thus generating a detectable signal.

Critical Review

surface-immobilized, affinity-based nanoswitches (NA and switches).^{15–17,68,69,80,90,91,101,102,108–118,125,137,171,172} hybrid Using this platform, a state-dependent readout can be obtained by immobilizing the switches on the sensor surface and labelling them with a redox reporter, as depicted in Fig. 4B. Since the electron transfer efficiency between the reporter and sensor surface is highly dependent on the distance between both, a distinguishable signal can be obtained for the bound and unbound state of the respective switches, provided that the conformational change caused by target binding alters the distance between the reporter and sensor surface.²⁷ The respective signal can be obtained using various electrochemical interrogation methods, including square-wave voltammetry, 12,16,17,49,69,80,90,101,109-116,118,137,150,173,174 differenpulse voltammetry,^{68,125,171} tial alternating current voltammetry,^{15,102,108} impedance spectroscopy⁹¹ and chronoamperometry¹¹⁷ among others. Interestingly, as electrochemical sensors are easy and cheap to manufacture, can be miniaturized, and can easily be combined with further electronic readout and processing, they already have found widespread applications in the field of continuous biosensing.¹⁷⁵

In Table 1, the readout strategies that have currently been combined with affinity-based nanoswitches are summarized, together with the applied switching strategy, target molecule, obtained $K_{\rm D}$ and LOD values, as well as the matrices in which measurements were performed. Importantly, only readout strategies for which the switches were attached to the sensor surface are provided, as these are in accordance with the requirements for continuous biosensing applications. As can be seen in this table, this strategy proved to be suitable for the detection of a variety of target molecules including proteins, NA and small molecules in a wide range of complex matrices. Notably, until now, mostly NA switches have been applied, whereas the amount of protein switches that have been attached to a sensor surface remains limited. However, as protein switches already proved to be suitable for real-time intracellular imaging,¹⁶⁸ we believe that they are promising for surface-based continuous biosensing as well. Moreover, although many more state-dependent readout platforms have been combined with affinity-based nanoswitches, from this summary, it becomes clear that electrochemical sensors are most widespread in this field. Importantly, not all biosensors listed in Table 1 have been applied for continuous measurements. Indeed, although in theory they consist of the required components to develop a continuous biosensor, this often simply was not the final objective. In particular, reagentless biosensors, intended for batch measurements, are a major achievement in itself, and already entail significant improvements compared to conventional affinity-based assays.^{20,27} Furthermore, even when all components required for continuous biosensing are combined, important challenges still need to be tackled to turn a reagentless biosensor for batch measurements into a continuous biosensor, being the long-term stability of the sensor and its dynamic response, as discussed in the following and previous sections, respectively. The biosensors indicated in Table 1 that have been applied for continuous measurements will be discussed in more detail in section 3.

Notably, the readout strategies summarized in Table 1 are not the only possible choices when developing a continuous biosensor, but only represent an overview of what has been reported in literature. In fact, more already existing strategies or newly developed approaches could be applied for this purpose, on the condition that they provide a distinguishable signal when the switches are in their bound or unbound states respectively.

2.2.2. Stability of the state-dependent readout platforms. As explained before, stability is an important aspect for continuous biosensing. Not only does this relate to the affinityreagents, but also to the readout platforms, on which the affinity-based nanoswitches are immobilized. The main difficulty again arises from the application of the sensors in complex matrices, depending on the final application. For example, to perform continuous measurements for patient or bioreactor monitoring, the sensors should be functional in bodily fluids such as blood and urine, or culture media respectively. More specifically, significant fouling effects are often observed when exposing sensor surfaces to complex matrices, which cause drift in the output signal, thereby affecting the accuracy of the measurement.¹⁴⁸ Several lowfouling strategies have been proposed to block foulants from sticking to the sensor surface, such as polymers and copolymers, hydrogels, peptides, zwitterionic materials, polysaccharides, biomimetic membranes and even 3D NA structures, which proved to be successful when subjected to complex matrices.^{7,178-180} Next to low fouling-layers, the use of dialysis membranes also proved beneficial to increase sensor stability, by blocking large components present in complex matrices from the sensor surface.^{17,181,182} However, as this strategy is size-based, it cannot be used for detecting large target molecules such as large proteins or extracellular vesicles, since they cannot reach the surface.⁷ We will not discuss the strategies to reduce fouling further, as this is out of the scope of this review and as this is already discussed elsewhere^{7,178–180}

3. Innovative concepts for continuous biosensing

In the following section, we will discuss biosensors in which affinity-based nanoswitches have been successfully combined with a state-dependent readout platform to achieve continuous measurements of varying target concentrations.

3.1. Electrochemical aptamer-based sensors

The electrochemical aptamer-based sensor (EA-B), developed in the research group of Kevin Plaxco, was the first platform to be applied for affinity-based continuous biosensing. Herein, SSAs, labelled with a redox reporter, are applied on an electrochemical sensor (using square wave, alternating current or cyclic voltammetry). These probes undergo a reversible conformational change after target binding, thereby altering the distance between the redox reporter and the biosensor surface, as depicted in Fig. 5A, which provides a dynamic signal, dependent on the target concentration. Using this strategy, continu-

Table 1 Overview of the readout strategies applied in combination with affinity-based nanoswitches. The respective switching strategy and target molecules are provided as well, together with the obtained K₀, LOD and matrix in which measurements were performed. It is also indicated whether the respective biosensors were used for batch or continuous measurements

erg back ance pasch ance fhrombin (umani)Continuous back back back back back back $2.5 \mu M$ buffe back back back back backBuffer buf	tein Alternative fold Squarewave volammetry VSL12 Exch Continuous 56 ± 4 µM = 470 ± 60 µM = 40 ± 10 µM = 40 \pm 10 \muM = 40 \pm 10 \pm 10 \muM = 40 \pm 10 \pm 10 \pm 10 \muM = 40 \pm 10 \pm		Switch strategy	Readout	Target	Batch/ continuous	$K_{ m D}$	LOD	Matrix	Ref.
Camping Surface plasmon resonance Provide incluing Particle mobility sensing Provide incluing Tronnibility activity Provide incluing Particle mobility sensing Provide incluing Particle mobility sensing Provide incluing Particle mobility sensing Provide incluing Particle mobility sensing Provide incluing Provide incluing Provide incluing Rombinity sensing Thickness shear mode Thrombin (burnis) Continuous 32.4.10 0.4 nM Buffer Filtered blood plasma Competition Differential pulse Thrombin Batch - 11.1M Buffer Differential pulse Thrombin Batch - 11.1M Buffer Valuammetry Netmathery - - - Buffer Valuammetry Netmathery - - - - Buffer Valuammetry Netmathery - - - - - Buffer Valuammetry Netmathery - - - - - - - Netmathery Netmathery -	Clamping Surface plasmon resonance Pricombin (human) Continuous Continuous Continuous 256 cM	Protein	Alternative fold		VSL12	Continuous Botok	$76 \pm 4 \mu M$	2.5 µM	Buffer ^a , whole blood	49
Particle inobility sensing Thrombin (human) Continuous 170 ± 56 ml - Buffer SDNA Continuous 99 ± 10 pM - Buffer Buffer SDNA Continuous 99 ± 10 pM - Buffer SDNA Continuous 99 ± 10 pM - Buffer SDNA Continuous 99 ± 10 pM - Patter SDNA Thrombin Batch 2.3 ± 0.7 ml 0.4 ml Buffer Alternating current Thrombin Batch - 11 ml Buffer Alternating current Continuous 5.3 ± 0.7 ml 0.4 ml Buffer Alternating current Batch - 11 ml Buffer Alternating current Continuous 5.3 ± 0.7 ml 0.4 ml Buffer Alternating current Batch - 11 ml Buffer 0.6 ml Alternating current FN-Y Batch - - Buffer Alternating current FN-Y Batch - 0.3 ml Buffer Alternating current FN-Y Batch - - Buffer Alternating current FN-Y Batch - 0.3 ml Buffer Imped	Particle inobility sensing sibNAThrombin (bovine) sibNAContinuous 170 ± 56 nM-SandarSandaThrombin (hurnan)Continuous 230 ± 60 pM-SandarThrombin (hurnan)Continuous 230 ± 60 pM-SandarThrombin (hurnan)Continuous 230 ± 60 pM-SandarThrombinThrombinBatchSandarThrombinBatch-11 nMSandarThrombinBatchSandarThrombinBatchSandarThrombinBatchAlternating currentTNF-aBatchVoltammetryFRNBatch0.58 nMSandarTNF-aBatch0.58 nMSandarTNF-aBatch0.1 pmolSandarSandarSandarSandarSandarTNF-aBatchSandarSandarSandarSandarSandarSandarSandarSandarSandarSandarSandar <td>NA</td> <td>Clamping</td> <td>Surface plasmon resonance</td> <td>Pooa-2 Thrombin</td> <td>Batch</td> <td>$\frac{4}{100} \times 0.06 \text{ nM}$</td> <td> </td> <td>Buffer</td> <td>176</td>	NA	Clamping	Surface plasmon resonance	Pooa-2 Thrombin	Batch	$\frac{4}{100} \times 0.06 \text{ nM}$		Buffer	176
Incomput (uturn)Continuous 32.4 ± 0 pM	tion bifferential pulse thrombin (utunat) continuous $230 \pm 0.0 \text{ PM} = -2.30 \pm 0.0 \text{ PM} = -1.0 $		2	Particle mobility sensing	Thrombin (bovine)	Continuous	$170 \pm 56 \text{ nM}$		Buffer	11
Notice 100 ± 208 $-$ PSD outree 110 Itickness shear modeThrombinBatch -3 ± 0.7 M 0.4 MBuffer 23 ± 3.7 $-$ Tiltered blood plasma 23 ± 3.7 $ 11$ MBufferactive intertod $ 11$ MBuffer 110 Miterential pulseThrombinBatch $ 11$ M 110 Miterential pulseThrombinBatch $ 110$ Miterential pulseFNYBatch $ 110$ Miterential pulseFNYBatch $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ 110$ M $ -$ <td>tion Thickness shear mode Thrombin Batch 2.3 ± 0.7 mM 0.4 mM 0.1 mM</td> <td></td> <td></td> <td></td> <td>Infombin (numan) ssDNA</td> <td>Continuous</td> <td>$49 \pm 10 \text{ pM}$</td> <td> </td> <td>Buffer^a, filtered blood plasma</td> <td>11</td>	tion Thickness shear mode Thrombin Batch 2.3 ± 0.7 mM 0.4 mM 0.1 mM				Infombin (numan) ssDNA	Continuous	$49 \pm 10 \text{ pM}$		Buffer ^a , filtered blood plasma	11
534 ± 257 -Filtered blood plasmafitonDifferential pulseThrombinBatch 23 ± 0.7 nM 0.4 nMBuffera coustic methodThrombinBatch $-$ 11 nMBufferAternating currentNotammetryFN-YBatch $ -$ BuffervoltammetryFN-YBatch $ -$ BuffervoltammetryFN-YBatch $ -$ voltammetryFN-YBatch $ -$ voltammetryFN-YBatch $ -$ voltammetryFN-YBatch $ -$ voltammetryFN-YBatch $ -$ fieldFN-YBatch $ -$ fieldFN-YBatc	tion Differential pulse Thrombin Batch $23 \pm 0.7 \mathrm{nM}$ 0.4 nM acoustic method Thrombin Batch -1 11 nM 0.4 nM voltammetry IFN-Y Batch -1 11 nM -1 -1 -1 -1 -1 -1 -1 -1						680 ± 268 pM		PBS DUITET	139
Thickness shear modeThrombinBatch 2.3 ± 0.7 nM 0.4 nMBufferitionDifferential pulseThrombinBatch $-$ 11 nMBuffervoltammetryThrombinBatch $-$ 11 nMBuffervoltammetryFN-YBatch $-$ 11 nMBuffervoltammetryFN-YBatch $ -$ BuffervoltammetryFN-YBatch $ -$ BuffervoltammetryFN-YBatch $ -$ BuffervoltammetryFN-YBatch $ -$ BuffervoltammetryFN-YBatch $ 0.58$ nMBuffervoltammetryFR-YBatch $ 0.58$ nMBufferinpedance spectroscopyIgEBatch $ 0.58$ nMBufferinpedance spectroscopySTABatch $ 0.58$ nMBufferinpedance spectroscopyStAContinuous 150 ± 50 nM $ -$ inpedance spectroscopyStADiloid $ -$ inpedance spectroscopyStADiloid $ -$ inpedance spectroscopyStADiloid $ -$ intervalitiper site $ -$ inpedance spectroscopyStADiloid $ -$ inpedance spectroscopyStADiloid $ -$ inpedance spectr	Thickness shear modeThrombinBatch $\frac{2^{MA}}{3.3\pm0.7}$ nM 0.4 nMtionDifferential pulseThrombinBatch $ 11$ nMvoltammetryIFN-YBatch $ 11$ nMAlternating currentIFN-YBatch $ -$ voltammetryIFN-YBatch $ -$ Square wave voltammetryIFN-YBatch $ -$ NemetryIFN-YBatch $ -$ Square wave voltammetryIFN-YBatch $ -$ NemetryITNF- α Batch $ -$ NetITNF- α Batch $ -$ Impedance spectroscopyIPBatch $ -$ Impedance spectroscopySSDNAContinuous $ -$ Impedance spectroscopySSDNABatch $ -$ Impedance spectroscopySSDNABatch $ -$ Intromosing sensingSSDNASSDNA $ -$ Intromosing sensingSSDNASSDNA $ -$						524 ± 257		Filtered blood plasma	
tion Different Incode outsammetry Thrombin Batch - 11 nM Buffer Voltammetry IFNY Batch - 11 nM Buffer Voltammetry New voltammetry IFNY Batch - Buffer Voltammetry IFNY Batch - - Buffer Square wave voltammetry TNFa Batch - 0.58 nM Buffer ⁴ , ired blood cell-depleted whole Impedance spectroscopy ATP Batch - 0.58 nM Buffer ⁴ , ired blood, cell-depleted Impedance spectroscopy ATP Batch - 0.58 nM Buffer ⁴ , ired blood, cell-depleted Impedance spectroscopy ATP Batch - 0.58 nM Buffer ⁴ , ired blood, cell-depleted Impedance spectroscopy ATP - 0.58 nM Buffer ⁴ , ired blood, cell-depleted Buffer ⁴ , ired blood, cell-depleted Impedance spectroscopy ATP - 0.58 nM Buffer ⁴ , ired blood, cell-depleted	tion acoust fraction acoust of the sector of the sector fraction of the sector of the			Thickness shear mode	Thrombin	Batch	$2.3 \pm 0.7 \text{ nM}$	0.4 nM	Buffer	177
Vortammetry Notammetry Environ - Buffer Alteranding current voltammetry FN-Y Batch - - Buffer Square wave voltammetry FN-Y Batch - - Red blood cell depleted whole blood Square wave voltammetry FN-Y Batch - - Red blood cell depleted whole Impedance spectroscopy ATP Batch - 0.58 nM whole blood, cell suspension Impedance spectroscopy ATP Batch - 0.58 nM Buffer ⁴ , pol od, cell suspension Particle mobility sensing sSDNA Continuous 150 ± 50 nM - Buffer ⁴ , pol od, cell suspension Particle mobility sensing sSDNA Continuous 150 ± 50 nM - 0.1 pmol defleted Particle mobility sensing sSDNA Continuous 150 ± 50 nM - - Buffer ⁴ , pol od, cell suspension Particle mobility sensing sSDNA Continuous 150 ± 50 nM - - Buffer ⁴ , pol od, cell suspension Batch - - 0.0 pM Buffer ⁴ , pol od, cell suspension - - Buffer ⁴ , pol od, cell suspension Particle mobility sensing sSDNA SSD M - 0.1 pmol Buffer <	Alternating current voltammetryIFN-YBatch-Alternating current voltammetryIFN-YBatchSquare wave voltammetryIFN-YBatchSquare wave voltammetryTNF- α BatchAlternating currentTNF- α Batch-0.58 nMImpedance spectroscopyIgEBatch-0.58 nMParticle mobility sensings5DNAContinuous35 \pm 9 nM-iveFRETs5DNABatch-0.1 pmolbatchs5DNABatch-0.1 pmolvoltammetrys5DNABatch0.1 pmolvoltammetrycontinuous150 \pm 9 nM100 pMvoltammetrycoaineVoltammetrycoaineChrono-amperometryTobramycinContinuousChrono-amperometryCocaineChrono-amperometryTobramycinContinuousContinuous100 \pm 9 nMFRETsSDNAContinuous100 \pm ContinuousContinuous<		Competition	bifferential pulse	Thrombin	Batch	I	11 nM	Buffer	68
Square wave voltammetryFN-YBatchBed blood cell-depleted whole blood, cell suspensionSquare wave voltammetryFN-YBatchRed blood cell-depleted whole blood, cell suspensionTNF-aBatch-0.58 nMButfer's red blood cell-depleted whole blood-Impedance spectroscopyIfFe0.58 nMButfer's red blood cell-depleted whole blood, cell suspensionImpedance spectroscopyIfFe0.58 nMButfer's red blood cell-depleted whole blood, cell suspensionImpedance spectroscopyIfFe0.58 nMButfer's red blood, cell suspensionImpedance spectroscopyIfFe0.01 pm0Butfer's red blood, cell suspensionImpedance spectroscopyIfFe0.1 pm0Butfer's red blood, cell suspensionImpedance spectroscopyIfFe0.1 pm0Butfer's red blood, cell suspensionImpedance spectroscopyIfFe0.1 pm0ButferImpedance spectroscopyIfFe0.1 pm0ButferImpedance spectroscopyIfFe0.1 pm0ButferImpedance spectroscopyIfFe0.1 pm0ButferImpedance spectroscopyIfFe0.1 pm0ButferImpedance spectroscopyIfFeIfFeIfFeImpedance spectroscopyIfFeIfFeIfFeImpedance spectroscopyIfFeIfFeIfFeImpedance spectroscopyIfFeIfFeIfFeImpedance spectroscopyIfFeIfFeIfFeImpedance spectroscopyIfFe <td>Square wave voltammetryFN-YBatchSquare wave voltammetryFN-YBatch-$<60 \text{ pM}$Square wave voltammetryTNF-αBatch-$<60 \text{ pM}$TNF-αBatch-0.58 nM$=150 \text{ sm}$The dance spectroscopyIgEBatch-0.58 nMParticle mobility sensingS5DNAContinuous$150 \pm 50 \text{ nM}$-NameFRETssDNAContinuous$35 \pm 9 \text{ nM}$-NameFRETssDNABatch-0.1 pmolStringS5DNABatch-0.1 pmolNameFRETssDNABatch-0.1 pmolNameStringS5 ± 9 nM0.1 pmolNameFRETS5DNABatch-0.1 pmolNameString0.1 pmol-StringString0.1 pmol-NameString0.1 pmol-NameString0.1 pmolNameStringNameStringStringStringStringString<t< td=""><td></td><td></td><td>voltammetry Alternating current</td><td></td><td></td><td></td><td>I</td><td>Buffer</td><td>102</td></t<></td>	Square wave voltammetryFN-YBatchSquare wave voltammetryFN-YBatch- $<60 \text{ pM}$ Square wave voltammetryTNF- α Batch- $<60 \text{ pM}$ TNF- α Batch-0.58 nM $=150 \text{ sm}$ The dance spectroscopyIgEBatch- 0.58 nM Particle mobility sensingS5DNAContinuous $150 \pm 50 \text{ nM}$ -NameFRETssDNAContinuous $35 \pm 9 \text{ nM}$ -NameFRETssDNABatch- 0.1 pmol StringS5DNABatch- 0.1 pmol NameFRETssDNABatch- 0.1 pmol NameStringS5 ± 9 nM 0.1 pmol NameFRETS5DNABatch- 0.1 pmol NameString 0.1 pmol -StringString 0.1 pmol -NameString 0.1 pmol -NameString 0.1 pmol NameStringNameStringStringStringStringString <t< td=""><td></td><td></td><td>voltammetry Alternating current</td><td></td><td></td><td></td><td>I</td><td>Buffer</td><td>102</td></t<>			voltammetry Alternating current				I	Buffer	102
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	ive FRET TNF-a Batch - 60 pM ive FRET TNF-a Batch - 0.58 nM ive FRET Batch - 0.58 nM ive FRET SSDNA Continuous 35 ± 9 nM - ive FRET SSDNA Continuous 35 ± 9 nM - ive FRET SSDNA Continuous 35 ± 9 nM - ive FRET SSNA Continuous 35 ± 9 nM - ive FRET SSNA Batch - - 0.1 pmol ive FRET SSNA Batch - - - - ive FRE Batch - - - - - ive SSNA Batch - - - - -			sourcements Square wave voltammetry	IFN-Y	Batch	Ι		Red blood cell-depleted whole	69
$ \begin{array}{ccccccccccccc} & & & & & & & & & & & & &$	ive FRET TNF-a Batch - 0.58 nM - 0.18 nch -							<60 pM	blood, cell suspension Red blood cell-depleted whole blood	80
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	TNF- α TNF- α Batch-0.58 nMIgEIgEBatch-0.58 nMParticle mobility sensings5DNAContinuous150 \pm 50 nM-Particle mobility sensings5DNAContinuous35 \pm 9 nM-Particle mobility sensings5DNABatch-0.1 pmolParticle mobility sensings5DNAContinuous35 \pm 9 nM-Particle mobility sensings5DNABatch-0.1 pmolParticle mobility sensings5DNABatchDifferential pulses5DNABatch-0.1 pmolNoltammetrydsDNABatchVoltammetrycontinuous100 \pm 9 μ MChrono-amperometryThrombinBatchChrono-amperometryTobramycinContinuous100 \pm 9 μ M							5 ng ml^{-1}	Cell suspension	06
Impedance spectroscopyIgEBatch-60 pMBufferDunder appendationParticle mobility sensingssDNAContinuous 150 ± 50 nM-BufferBufferParticle mobility sensingssDNAContinuous 35 ± 9 nM-BufferBufferParticle mobility sensingssDNAContinuous 35 ± 9 nM-BufferParticle mobility sensingssDNAContinuous 35 ± 9 nM-BufferParticle mobility sensingssDNABatch-0.1 pmolBufferState-0.1 pmolBuffer-100 pMBufferStrNA100 pMBufferStrNA100 pMBufferDifferential pulsessDNA275 pMBufferVoltammetry275 pMBuffer-VoltammetryVoltammetryVoltammetryVoltammetryVoltammetryVoltammetryVoltammetryVoltammetryVoltammetryVoltammetry <t< td=""><td>Impedance spectroscopy IgE Batch - 60 pM Particle mobility sensing ssDNA Continuous 150 ± 50 nM - Particle mobility sensing ssDNA Continuous 35 ± 9 nM - ive FRET ssDNA Batch - - Continuous 35 ± 9 nM - - - Impedance spectroscopy ssDNA Batch - - Differential pulse ssRNA Batch - - 10 nM Notammetry dsDNA Batch - - - - Voltammetry Coraine Continuous 100 ± 9 µM - - - Chrono-amperometry Tobramycin Continuous 100 ± 9 µM - - -</td><td></td><td></td><td></td><td>TNF-α</td><td>Batch</td><td>I</td><td>0.58 nM</td><td>Buffer^a, red blood cell-depleted</td><td>69</td></t<>	Impedance spectroscopy IgE Batch - 60 pM Particle mobility sensing ssDNA Continuous 150 ± 50 nM - Particle mobility sensing ssDNA Continuous 35 ± 9 nM - ive FRET ssDNA Batch - - Continuous 35 ± 9 nM - - - Impedance spectroscopy ssDNA Batch - - Differential pulse ssRNA Batch - - 10 nM Notammetry dsDNA Batch - - - - Voltammetry Coraine Continuous 100 ± 9 µM - - - Chrono-amperometry Tobramycin Continuous 100 ± 9 µM - - -				TNF-α	Batch	I	0.58 nM	Buffer ^a , red blood cell-depleted	69
Impedance spectroscopy $\stackrel{\mathrm{JgE}}{\mathrm{ATP}}$ bacculor-00 pmbuttlet, too dunced setundParticle mobility sensingssDNABatch-00 pmbuttlet, too dunced setundParticle mobility sensingssDNAContinuous150 ± 50 nM-BufferBufferParticle mobility sensingssDNAContinuous $35 \pm 9 nM$ -BufferRETssDNABatch-0.1 pmolBufferSRNABatch-0.1 pmolBuffersrNAssRNA-0.1 pmolBuffersrNAssRNA-10 nMBuffersrNA20 pMBuffersrNA10 nMBuffersrNA100 pMbufferential pulsessDNAvoltammetry275 pMMitternating currentThrombinvoltammetryvoltammetryvoltammetryvoltammetryvoltammetryvoltammetryvoltammetryvoltammetryvoltammetryvoltammetryvoltammetryvoltammetryvoltammetry<	Impedance spectroscopy $\stackrel{\rm Agr}{ATP}$ Batch00 pmParticle mobility sensingssDNAContinuous150 ± 50 nMParticle mobility sensingssDNAContinuous35 ± 9 nMRETssDNABatch0.1 pmolRetrssNABatch10 nMSistab10 nMSistab10 nMSistab10 nMSistabSistabSistabSistabSistabSistabSistabSistabSistabSistabSistabSistabSistabSistabSistabSistabSistabSistab				T.e.T.	Dotoh		1 m 03	WIDIE DIOOU, CEII SUSPERISION	101
ive FRET solutions $150 \pm 50 \text{ nM}$ - butter Particle mobility sensing sSDNA Continuous $150 \pm 50 \text{ nM}$ - butter Particle mobility sensing sSDNA Continuous $35 \pm 9 \text{ nM}$ - Buffer Continuous $35 \pm 9 \text{ nM}$ - Buffer Continuous $35 \pm 9 \text{ nM}$ - Buffer Continuous $35 \pm 9 \text{ nM}$ - Buffer - $ -$	ive FRET ssDNA Continuous 150±50 nM Particle mobility sensing ssDNA Continuous 150±50 nM - Particle mobility sensing ssDNA Continuous 150±50 nM - Particle mobility sensing ssDNA Continuous 150±50 nM - Particle mobility sensing ssDNA Batch - 0.1 pmol Particle mobility sensing ssDNA Batch - 0.1 pmol Particle mobility sensing ssRNA Batch - 100 nM Particle mobility sensing ssDNA Batch - - Particle mobility sensing ssDNA Batch - - Particle mobility sensity Thrombin Batch - - Poltammetry Continuous 100±9 µM - - Chrono-amperometry Tobramycin Continuous - -			Immediance croatmoorn	IgE	Batch		60 pM	Buffer', 10× anuted Serum	101
reducte filter ssDNA Continuous LOUE 35 ± 9 nM - builter ive FRET ssDNA Batch - - Buffer Continuous 35 ± 9 nM - Buffer - Buffer SRNA Batch - 0.1 pmol Buffer ssRNA - 0.1 pmol Buffer bifferential pulse ssDNA Batch - - Differential pulse ssDNA Batch - - Voltammetry dsDNA - 275 pM Buffer Voltammetry Cocaine - - Buffer Voltammetry - - - -	ive FRET SSDNA Continuous $135 \pm 9 \text{ nM}$ – ive FRET SSDNA Batch – 0.1 pmol – 0.1 pmol – 0.1 pmol – 100 nM – $100 \text$			nupeuance spectroscopy		Continuent			Duiter	4 F
ive FRET SSDNA Batch - buttet	ive FRET sSDNA Batch			Particle modulity sensing	SSDINA	Continuous			Builter Diiffor	14 120
$ \begin{array}{cccccc} - & 0.1 \mbox{ pmol} & \mbox{ buffer} \\ \mbox{ ssRNA} & - & 0.1 \mbox{ pmol} & \mbox{ buffer} \\ \mbox{ ssRNA} & \mbox{ ssRNA} & - & 100 \mbox{ pM} & \mbox{ buffer} \\ & 100 \mbox{ pM} & \mbox{ buffer} \\ & - & \mbox{ buffer} \\ \mbox{ ssDNA} & \mbox{ ssDNA} \\ \mbox{ ssDNA} & \mbox{ buffer} \\ \mbox{ ssDNA} & \mbox{ buffer} \\ \mbox{ ssDNA} & \mbox{ ssDNA} \\ \mbox{ ssDNA} & \mbox{ serum} \\ \$	$ \begin{array}{ccccc} - & 0.1 \mbox{ pmol} \\ \mbox{ssRNA} & - & 0.1 \mbox{ pmol} \\ \mbox{ssRNA} & - & 0.1 \mbox{ pmol} \\ \mbox{ pmol} \\ \mbox{bifferential pulse} & \mbox{ssDNA} & \mbox{ Batch} & - & - \\ \mbox{ voltammetry} & \mbox{ dsDNA} & \mbox{ Batch} & - & - \\ \mbox{ voltammetry} & \mbox{ continuous} & 100 \pm 9 \mbox{ mod} \\ \mbox{ chrono-amperometry} & \mbox{ Tohramycin} & \mbox{ continuous} & 100 \pm 9 \mbox{ mod} \\ \mbox{ mod} \mbox{ mod} \\ \mbox{ mod} \\ \mbox{ mod} \mbox{ mod} \\ mo$		Alternative	FRET	SSDNA	Batch	TMTT 6 ∓ CC		Buffer	121
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$ \begin{array}{c cccc} & & & & & & & & & & & & & & & & & $		folding					0.1 pmol	Buffer	122
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$ \begin{array}{cccccc} \text{ssRNA} & - & 100 \ \text{pM} \\ - & - & 0 \ \text{ulse} & \text{ssDNA} & \text{Batch} & - & - \\ \text{asDNA} & \text{Batch} & - & - \\ \text{dsDNA} & - & 275 \ \text{pM} \\ - & - & 275 \ \text{pM} \\ - & - & - \\ \text{coraine} & \text{Continuous} & 100 \pm 9 \ \mu\text{M} & - \\ \text{erometry} & \text{Tobramycin} & \text{Continuous} & - & - \\ \end{array} $							10 nM	Buffer	124
oulsessDNABatch-Intracellular (HeLa cell)oulsessDNABatch-BufferdsDNA-275 pMBufferurrentThrombinBatch-DocaineContinuous $100 \pm 9 \mu M$ -Etail bovine serum	oulsessDNABatch $dsDNA$ $dstch$ -275 pM $drent$ Thrombin $Batch$ -275 pM $urrent$ Thrombin $Batch$ $cocaine$ Continuous $100 \pm 9 \mu M$ -erometryTobramycinContinuous-				SSRNA			100 pM	Buffer	123
bulse ssDNA Batch – – Buffer $-$ Buffer dsDNA $ -$	oulse ssDNA Batch $ -$ dsDNA $-$ 275 pM urrent Thrombin Batch $ -$ 275 pM - $ -$								Intracellular (HeLa cell)	120
display $ 2/5$ pmButterurrentThrombinBatch $ -$ Buffer, $2\times$ diluted fetal calf serumCocaineContinuous $100 \pm 9 \mu M$ $-$ Fetal bovine serum	dsDNAdsDNA $ 2/3$ pMurrentThrombinBatch $ -$ CocaineContinuous $100 \pm 9 \ \mu M$ $-$ erometryTobramycinContinuous $-$			Differential pulse	SSDNA	Batch			Buffer	125
Cocaine $100 \pm 9 \mu M$ – Fetal by the sector	$ \begin{array}{cccc} & & & & \\ & & & & \\ & & & & \\ & & & & $			voltammetry Alternating current	dsDNA Thromhin	Batch		Mq c/2	Buffer 2× diluted fetal calf serum	1/1
	Iobramycin Conunuous — —			voltammetry	Cocaine	Continuous	$100\pm9~\mu M$		Fetal bovine serum	12

Table 1 (Contd.)

Published on 15 2022. Downloaded on 2025/5/21 17:15:45.

	Switch strategy	Readout	Target	Batch/ continuous	K_{D}	LOD	Matrix	Ref.
		Square-wave voltammetry	Doxorubycin	Continuous	$824 \pm 18 \text{ nM}$	10 nM	Buffer ^a , human whole blood, rat	12, 17 and
					69.2 µM		Whole blood	112
				Batch	3.98 JM		Serum	
			Kanamycin	Continuous	920 µM		Buffer ^{<i>a</i>} , rat whole blood (<i>ex</i> and	12, 17, 112
							in vivo)	and 114
			Cocaine	Continuous	395 µM		Whole blood	112
				Batch	182 μM		Serum	
				Continuous			Whole blood	114
			Tobramycin	Continuous	250 µM		Rat whole blood (<i>ex</i> and <i>in vivo</i>)	16, 17, 115 and 116
			Irenotican	Continuous	$58 \pm 17 \ \mu M$		Buffer	110
					331 ± 57		Rat whole blood (<i>ex</i> and <i>in vivo</i>)	
			Gentamycin	Continuous			Rat whole blood (<i>in vivo</i>)	17
			Melamine	Batch	$490 \pm 10 \ \mu M$		Buffer	118
				Continuous	$463 \pm 10 \ \mu M$		Milk	
			Vancomycin	Continuous	$45.5 \pm$		Rat whole blood (ex and $in vivo$)	111
					2.2 μM			
			Transcription factor (TATA-binding protein)	Batch	$121 \pm 20 \text{ nM}$		Buffer ^{a} , HeLa nuclear extract ^{a}	109
			NGAL	Continuous		2/3.5 nM	Urine (artificial/human)	173
			Ochratoxin A	Batch	$6.5 \pm 0.6 \ \mu M$		Buffer	150
					$19 \pm 2 \mu M$		Unprocessed iced coffee	
			Phenylalanine	Continuous			Buffer, whole blood	174
		Surface plasmon	SSDNA	Batch		500 pM	Buffer	126
		Rudescence spectroscopy Surface-enhanced Raman	ssDNA	Batch	I	I	Buffer	127
NA-protein	Clamning	scauer mg Square wave voltammetry	Anti-DNP antibody	Batch	3.6 nM	0.6 nM	Buffer	137
hybdrid	Quitter		Anti-Dig antibody Anti-HIV antibody		8 nM 33 nM	2 nM 9 nM	Whole blood Whole blood	
	Competition	Particle mobility sensing	Creatinine	Continuous	$25 \pm 7 \mu M$	13.2 μM 	Buffer Buffer	14 130
					TATH $o \pm aa$		Dullet	LOL

 a The respective matrix in which the K_{D} and LOD values were obtained, in case measurements were performed in multiple matrices.

Analyst

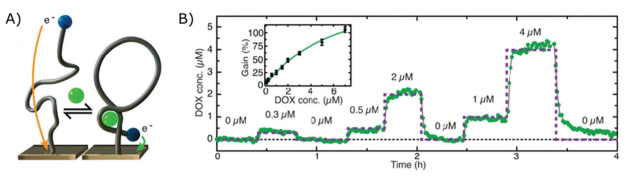


Fig. 5 (A) Working principle of the electrochemical aptamer-based sensor (EA-B): SSAs, equipped with a redox reporter, are immobilized on an electrochemical sensor. Upon target binding, the SSAs will undergo a conformational change, thereby bringing the redox reporter closer to the sensor surface. This conformational change enables electron transfer, which provides a detectable signal.¹² (B) Dynamic signal obtained by the EA-B platform for Doxorubicin detection. Adapted with permission from ref. 12. Copyright (2013) The American Association for the Advancement of Science.

ous monitoring of small molecules, including drugs such as Doxorubicin,^{12,17,112,113} Kanamycin,^{12,17} Cocaine,^{15,112,114} Tobramycin,^{16,17,115-117} Irenotican,¹¹⁰ Gentamycin,¹⁷ and Vancomycin,¹¹¹ could be achieved, of which an example can be seen in Fig. 5B.

In their first report on the use of this platform for continuous biosensing in 2009, cocaine concentrations were monitored in undiluted blood serum.¹⁵ Since then, much progress has been made to further enhance the performance and application of these sensors, such as the inclusion of a second reporter¹¹⁴ and the application of a biomimetic monolayer to increase the stability of the respective sensor.¹¹³ In this manner, they could successfully reduce drift due to fouling of the sensor surface when performing multi-hour measurements in whole blood.^{113,114} Furthermore, they tuned the affinity of SSAs as elaborated in section 2.1.4. and in this manner could tune, extend and narrow the dynamic range of the final EA-B sensor.146,183,184 Moreover, by using chronoamperometrical interrogation, they achieved measurements with a 300 ms time resolution.¹¹⁹ The group also increased the surface area of the working electrode by roughening the gold surface, which resulted in a higher signal to noise ratio.¹¹⁷ Later, the surface area was further increased by employing nanoporous gold, allowing miniaturization of the sensor.185 Due to these improvements, the EA-B sensor could be applied in situ (in vivo) for multi-hour measurements of therapeutic drug concentrations with (sub)second resolution.^{110,117} On top of that, to close the gap between drug administration and therapeutic drug monitoring, the EA-B sensor was successfully applied for feedback-controlled drug administration.¹¹¹ Furthermore, aside from the application of the EA-B sensor for therapeutic drug monitoring, the sensor was also applied to continuously monitor a protein biomarker for acute renal injury (NGAL) in urine¹⁷³ and toxic substances (melamine, ochratoxin A) for molecular quality control in food and beverage streams.^{118,150}

In 2020, the Plaxco group supplemented their arsenal of SSAs applied on the electrochemical platform for continuous biosensing with a protein switch as well, to further expand the range of targets to be detected. The protein switch was engineered through the inclusion of an alternative fold from the SH3 domain from human Fyn kinase (FynSH3), as discussed in section 2.1.1, so a binding-induced conformational change could be achieved.⁴⁹

Altogether, the EA-B sensor enables continuous biosensing directly in the environment of interest (*in situ*) with a fast time-resolution (subsecond). However, as mentioned before, a trade-off between dynamic response and detection limit can generally be expected. Indeed, although the dynamic response of the EA-B sensor is fast, the detection limit of the EA-B sensors is relatively high (μ M-mM). This is however not necessarily a limitation, as the detection limit suffices for reported applications of the EA-B sensor.

Currently, there are 2 companies working on the commercialization of the EA-B sensors, being Nanogenecs and Eccrine Systems, Inc.,¹⁸⁶ of which the latter is commercializing these sensors for continuous biosensing, and aims to 'individualize the prescription of medicine with a breakthrough platform for remote, non-invasive, and quantitative medication monitoring through sweat'.¹⁸⁶

3.2. Biomarker monitoring by particle mobility sensing

A second biosensor in which affinity-based nanoswitches have been combined with a state-dependent readout system to achieve continuous biosensing, was developed by the group of Menno Prins. The switches applied on their sensor are NAbased affinity clamps, composed of a microparticle tethered to a surface *via* dsDNA. Both the particle and surface are equipped with NA bioreceptors (ssDNA probes or aptamers), so when the target is present, a sandwich will be formed and the switch will undergo a conformational change, as presented in Fig. 6A. Digital state-change events of the particles are observed by tracking the mobility of each particle through microscopy. More specifically, when the switches are unbound, the particles will move according to Brownian motion and are only restricted in their movement by the dsDNA tether, with which they are linked to the surface. When

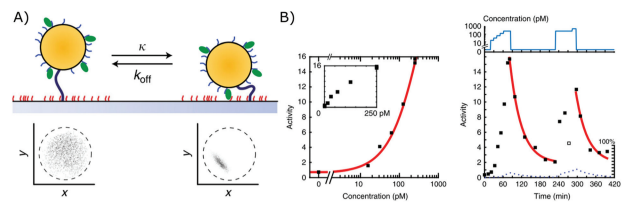


Fig. 6 (A) Affinity clamps composed of a microparticle, tethered to a sensor surface, both equipped with bioreceptors (top). The bound and unbound states of the switch can be discriminated through their mobility patterns (below), which are obtained by following the position of a single particle over time.¹¹ (B) The target concentration is linked to the activity of the sensor as shown in the left graph. Herein, the activity is determined as the average amount of state-changes occurring per switch, during a certain time-interval. By following the activity of the sensor over time, a dynamic signal, dependent on the target concentration, can be obtained (right graph). Adapted with permission from ref. 11 (CC BY 4.0).¹⁸⁷

they are however bound to the target, a sandwich is formed, and the particles are more restricted in their movement as they are linked to the surface both *via* the tether and the formed sandwich. In this manner, the mobility patterns, which are shown in Fig. 6A for both the bound and unbound state, can be clearly discriminated. Furthermore, by following the activity of the sensor (which is defined by the average amount of statechanges occurring per switch during a certain time-interval), the analyte concentration can be determined and continuously followed, as shown in Fig. 6B.¹¹

Using this approach, continuous monitoring of both ssDNA and protein (thrombin) targets could be achieved in filtered blood plasma, obtained by applying a porous membrane to the sensor setup (as discussed in section 2.2.2), of which the results can be seen in Fig. 6B for ssDNA detection.¹¹ Furthermore, by utilizing the dissociation rate of each single particle as a signature to identify a kinetic subpopulation, multiplexing of target molecules with a significantly different dissociation rate (ssDNA with different length and sequence) could be achieved.¹³

Although the proposed combination of the particle mobility sensing platform with affinity clamps proved to be suitable for continuous biosensing of both ssDNA and protein targets, it does not allow detection of small molecules, as the size of such analytes restricts the use of a sandwich conformation. Therefore, similar NA and NA-protein hybrid switches were developed via competition,¹⁴ for the detection of both creatinine and a short ssDNA oligo. The switches were again composed of a microparticle linked to the sensor surface. However, only one was equipped with a bioreceptor (antibody or ssDNA probes), whereas the other was linked to the competitor, being (i) a creatinine molecule or (ii) a DNA strand containing the same sequence as the short oligonucleotide target, for the detection of creatinine and DNA respectively. The resulting NA-protein hybrid switch for creatinine was depicted in Fig. 3B. Using these competitive switches, they were able to

successfully detect creatinine and DNA with μ M and nM detection limits respectively.¹⁴ Furthermore, corresponding to the importance of long-term stability of a continuous biosensor as discussed above, they improved the control and durability of their platform by functionalizing the respective switches *via* click-chemistry to a bottlebrush polymer (PLL-g-PEG).¹³⁹

Overall, by tracking the mobility of affinity-based switches, continuous biosensing of DNA, proteins and small molecules could be achieved with a low detection limit ($pM-\mu M$), due to the single molecule resolution of the platform.¹¹ Importantly, although it was stated before that a lower target affinity generally provides a lower sensitivity, this is not the case for this readout strategy. Indeed, low affinity capture elements can be applied to achieve a low detection limit, as the obtained signal is not determined by the average amount of bound switches for a certain target concentration, but by the amount of statechanges the switches will undergo during a certain time interval. Moreover, information about the binding kinetics could be obtained, which allowed multiplex detection.¹³ A limitation of the proposed strategy however arises from the fact that the readout strategy is based on microscopy. As a result, measurements cannot be performed in opaque samples (such as whole blood) or directly in the environment of interest (in situ). Rather part of the sample needs to be transported over the sensor and can potentially be transported back to the environment of interest (in-line). Furthermore, in order to determine the activity of the sensor (and thus the target concentration), the state-changes of the switches need to be followed over a certain timeframe and therefore, the time-resolution is limited by the duration this timeframe.¹¹

The biosensor for biomarker monitoring by particle mobility sensing is currently in the process of being commercialized by Helia Biomonitoring, a spin-off company of Eindhoven University of Technology, with a vision of 'real-time biomolecular sensing for closed-loop control in healthcare and industry'.¹⁸⁸

4. Conclusions and future perspectives

Continuous biosensors can provide real-time information about biochemical processes occurring in the environment of interest, which is highly desirable in both research, diagnostic, as well as industry settings. However, the amount of commercially available biosensors for this purpose remain limited and are all based on the enzymatic conversion of the target molecules. Affinity-based biosensors are not dependent on the reactivity of the target itself and can therefore be applied for a wider range of targets, such as DNA, proteins, small molecules $etc.^7$ Recently, it was shown that affinity-based continuous biosensors can be developed by combining affinity-based nanoswitches, with a state-dependent readout platform.^{11,15}

Three main strategies have been applied to develop affinitybased nanoswitches from affinity binders, being (i) the development of a clamp,^{18,22} (ii) linking the affinity binder to a competitor^{18,23-25} and (iii) the introduction of an alternative fold.^{18,24,25,27} Using these approaches, both DNA and protein switches have successfully been developed. Importantly, as both components are structurally and chemically different, they require a different expertise and provide distinct opportunities for switch development. Generally, due to its chemical simplicity, the thermodynamic behavior of NA is predictable, which eases the design and development of NA switches.55 On the other hand, as amino acids have a broader chemical diversity than nucleic acids, a broader functionality can be obtained using protein switches.¹⁸ Furthermore, by combining both components, their strengths can be combined, and NAprotein hybrid switches can thus be engineered with a broad functionality and high predictability.^{136,137} However, in our opinion, the potential of hybrid switches currently remains underexplored. Moreover, further improvements of current predictive software for both DNA and protein engineering are required to allow rational tuning of affinity-based nanoswitches and thereby ease their design and development of affinity-based nanoswitches for a specific application in the future.

To convert the conformational change of affinity-based nanoswitches into a detectable signal, they should be combined with a state-dependent readout strategy, which provides a distinct signal for both conformational states of the affinitybased nanoswitches. In this manner, the conformational changes of the switches can be linked to the target concentration.9 Furthermore, as the switches itself cannot be applied as external reagents, due to the requirements for continuous biosensing, all components should be attached to the sensor surface.7,9 Although many affinity-based nanoswitches have already been combined with state-dependent readout platforms, resulting in reagentless biosensors,²⁷ up to now, only 2 research groups have integrated both components for continuous biosensing.^{11,15} Indeed, although in theory it might seem easy to combine the required components to develop a continuous biosensor, in practice, it takes more than that. More

specifically, depending on the final application, the affinitybased nanoswitches must be carefully tuned to obtain the required sensitivity and dynamic response.^{7,9} Furthermore, the switches and sensor surface must be stable in the environment of interest for the duration of the measurement, so the provided signal does not suffer from drift over time.^{7,9,148} Although much progress has already been made to overcome each of the aforementioned difficulties, it remains challenging to integrate all solutions into one continuous biosensor. We believe however, that due to the ever-improving fields of protein engineering and DNA nanotechnology and by bringing together expertise from those and other fields, the field of continuous biosensing holds much more potential in the future.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

This work has received funding from the Research Foundation Flanders (FWO SB/1SC8519N) and European Union's Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie grant agreement no. 955623 (H2020-MSCA-CONSENSE).

References

- 1 M. C. Frost and M. E. Meyerhoff, *Annu. Rev. Anal. Chem.*, 2015, **8**, 171–192.
- 2 O. S. Alwan and K. P. Rao, *Healthc. Technol. Lett.*, 2017, 4, 142–144.
- 3 N. D. Lourenço, J. A. Lopes, C. F. Almeida, M. C. Sarraguça and H. M. Pinheiro, *Anal. Bioanal. Chem.*, 2012, 404, 1211–1237.
- 4 A. Guerra, M. Von Stosch and J. Glassey, *Crit. Rev. Biotechnol.*, 2019, **39**, 289–305.
- 5 R. Desmarais, D. Anderson, J. Downey, O. Wesstrom, M. Rice, A. Worley, S. Wise, F. Tappen, J. L. Cawley and N. Andersen, Inline monitoring aids in food safety and quality Available online: https://www.foodengineeringmag.com/articles/90659-inline-monitoring-aids-in-foodsafety-and-quality (accessed on May 27, 2020).
- 6 J. Meseguer and J. Quevedo, *Real-time monitoring and* control in water systems in: Real-time monitoring and operational control of drinking-water systems, Springer, 2017.
- 7 J. Qu, A. Dillen, W. Saeys, J. Lammertyn and D. Spasic, *Anal. Chim. Acta*, 2019, **1104**, 10–27.
- 8 R. Renneberg, D. Pfeiffer, F. Lisdat and G. Wilson, *Frieder Scheller and the Short History of Biosensors in: Biosensing for the 21st Century*, Springer, Berlin, 2008.
- 9 K. W. Plaxco and H. T. Soh, *Trends Biotechnol.*, 2011, **29**, 1–5.
- 10 K. Pandit, Indian J. Endocrinol. Metab., 2012, 16, 263–266.

- 11 E. W. A. Visser, J. Yan, L. J. Van Ijzendoorn and M. W. J. Prins, *Nat. Commun.*, 2018, 9, 1–10.
- 12 B. S. Ferguson, D. A. Hoggarth, D. Maliniak, K. Ploense, R. J. White, N. Woodward, K. Hsieh, A. J. Bonham, M. Eisenstein, T. E. Kippin, K. W. Plaxco and H. T. Soh, *Sci. Transl. Med.*, 2013, 5, 213–165.
- 13 R. M. Lubken, A. M. De Jong and M. W. J. Prins, Nano Lett., 2020, 20, 2296–2302.
- 14 J. Yan, L. Van Smeden, M. Merkx, P. Zijlstra and M. W. J. Prins, ACS Sens., 2020, 5, 1168–1176.
- 15 J. S. Swensen, Y. Xiao, B. S. Ferguson, A. A. Lubin, R. Y. Lai, A. J. Heeger, K. W. Plaxco and H. Tom Soh, *J. Am. Chem. Soc.*, 2009, **131**, 4262–4266.
- 16 N. Arroyo-Currás, G. Ortega, D. A. Copp, K. L. Ploense, Z. A. Plaxco, T. E. Kippin, P. Hespanha and K. W. Plaxco, *ACS Pharmacol. Transl. Sci.*, 2018, 1, 110–118.
- N. Arroyo-Currás, J. Somerson, P. A. Vieira, K. L. Ploense, T. E. Kippin and K. W. Plaxco, *Proc. Natl. Acad. Sci. U. S. A.*, 2017, **114**, 645–650.
- 18 V. Stein and K. Alexandrov, *Manifesting Synth. Biol. Synth.*, 2015, 33, 101–110.
- 19 A. Vallée-Bélisle and K. W. Plaxco, Curr. Opin. Struct. Biol., 2010, 20, 518–526.
- 20 H. Adamson and L. J. C. Jeuken, ACS Sens., 2020, 5, 3001– 3012.
- 21 H. Adamson, M. O. Ajayi, E. Campbell, E. Brachi, C. Tiede, A. A. Tang, T. L. Adams, R. Ford, A. Davidson, M. Johnson, M. J. Mcpherson, D. C. Tomlinson and L. J. C. Jeuken, *ACS Sens.*, 2019, 4, 3014–3022.
- 22 H. Hasegawa, K.-I. Taira, K. Sode and K. Ikebukuro, *Sensors*, 2008, **8**, 1090–1098.
- 23 J. D. Munzar, A. Ng and D. Juncker, *Chem. Soc. Rev.*, 2019, 48, 1390–1419.
- 24 T. A. Feagin, N. Maganzini and H. T. Soh, ACS Sens., 2018, 3, 1611–1615.
- 25 H. Farrants, J. Hiblot, R. Griss and K. Johnsson, *Rational Design and Applications of Semisynthetic Modular Biosensors: SNIFITs and LUCIDs in: Synthetic Protein Switches*, 2017.
- 26 A. Dillen, W. Vandezande, D. Daems and J. Lammertyn, *Anal. Bioanal. Chem.*, 2021, **413**, 4739–4750.
- 27 L. R. Schoukroun-Barnes, F. C. Macazo, B. Gutierrez, J. Lottermoser, J. Liu and R. J. White, *Annu. Rev. Anal. Chem.*, 2016, 9, 163–181.
- 28 M. M. Stratton and S. N. Loh, Protein Sci., 2011, 20, 19–29.
- 29 M. V. Golynskiy, W. F. Rurup and M. Merkx, *ChemBioChem*, 2010, **11**, 2264–2267.
- 30 Y. Ni, R. Arts and M. Merkx, ACS Sens., 2018, 4, 20–25.
- 31 R. Arts, I. Den Hartog, S. E. Zijlema, V. Thijssen,
 S. H. E. Van Der Beelen and M. Merkx, *Anal. Chem.*, 2016,
 88, 4525–4532.
- 32 K. Tomimuro, K. Tenda, Y. Ni, Y. Hiruta, M. Merkx and D. Citterio, *ACS Sens.*, 2020, **5**, 1786–1794.
- 33 M. Van Rosmalen, Y. Ni, D. F. M. Vervoort, R. Arts, S. K. J. Ludwig and M. Merkx, *Anal. Chem.*, 2018, 90, 3592–3599.

- 34 E. Ceballos-Alcantarilla and M. Merkx, *Understanding and applications of Ser/Gly linkers in protein engineering in: Linkers in Biomacromolecules*, Academic Press Inc., 2021, vol. 647.
- 35 C. Tiede, R. Bedford, S. J. Heseltine, G. Smith, I. Wijetunga, R. Ross, D. Alqallaf, A. P. E. Roberts, A. Balls, A. Curd, R. E. Hughes, H. Martin, S. R. Needham, L. C. Zanetti-Domingues, Y. Sadigh, T. P. Peacock, A. A. Tang, N. Gibson, H. Kyle, G. W. Platt, N. Ingram, T. Taylor, L. P. Coletta, I. Manfield, M. Knowles, S. Bell, F. Esteves, A. Maqbool, R. K. Prasad, M. Drinkhill, R. S. Bon, V. Patel, S. A. Goodchild, M. Martin-Fernandez, R. J. Owens, J. E. Nettleship, M. E. Webb, M. Harrison, J. D. Lippiat, S. Ponnambalam, M. Peckham, A. Smith, P. K. Ferrigno, M. Johnson, M. J. McPherson and D. C. Tomlinson, *eLife*, 2017, 6, 1–35.
- 36 J. Huang, K. Makabe, M. Biancalana, A. Koide and S. Koide, *J. Mol. Biol.*, 2009, **392**, 1221–1231.
- 37 J. Huang, A. Koide, K. Makabe and S. Koide, *Proc. Natl. Acad. Sci. U. S. A.*, 2008, **105**, 6578–6583.
- 38 J. Huang and S. Koide, ACS Chem. Biol., 2010, 5, 273-277.
- 39 S. J. Aper and M. Merkx, *ACS Synth. Biol.*, 2016, 5, 698–709.
- 40 E. M. W. M. Van Dongen, L. M. Dekkers, K. Spijker, E. W. Meijer, L. W. J. Klomp and M. Merkx, *J. Am. Chem. Soc.*, 2006, **128**, 10754–10762.
- 41 J. L. Vinkenborg, T. J. Nicolson, E. A. Bellomo, M. S. Koay, G. A. Rutter and M. Merkx, *Nat. Methods*, 2009, 6, 737.
- 42 M. A. Brun, K.-T. Tan, E. Nakata, M. J. Hinner and K. Johnsson, J. Am. Chem. Soc., 2009, 131, 5873–5884.
- 43 M. A. Brun, K.-T. Tan, R. Griss, A. Kielkowska, L. Reymond and K. Johnsson, *J. Am. Chem. Soc.*, 2012, **134**, 16.
- 44 A. Schena, K. Johnsson, A. Schena and K. Johnsson, Angew. Chem., Int. Ed., 2014, 53, 1302–1305.
- 45 M. A. Brun, R. Griss, L. Reymond, K.-T. Tan, J. Piguet, R. J. R. W. Peters, H. Vogel and K. Johnsson, *J. Am. Chem. Soc.*, 2011, **133**, 16235–16242.
- 46 A. Masharina, L. Reymond, D. Maurel, K. Umezawa and K. Johnsson, J. Am. Chem. Soc., 2012, 134, 56.
- 47 R. Griss, A. Schena, L. Reymond, L. Patiny, D. Werner, C. Tinberg, D. Baker and K. Johnsson, *Nat. Chem. Biol.*, 2014, 10, 598–604.
- 48 L. Xue, Q. Yu, R. Griss, A. Schena and K. Johnsson, Angew. Chem., Int. Ed., 2017, 56, 7112–7116.
- 49 M. Kurnik, E. Z. Pang and K. W. Plaxco, Angew. Chem., Int. Ed, 2020, 59, 1–5.
- 50 J. E. Kohn and K. W. Plaxco, Proc. Natl. Acad. Sci. U. S. A., 2005, 102, 10841–10845.
- 51 M. M. Stratton, D. M. Mitrea and S. N. Loh, *ACS Chem. Biol.*, 2008, **3**, 723–732.
- 52 J.-H. Ha and S. N. Loh, Construction of Allosteric Protein Switches by Alternate Frame Folding and Intermolecular Fragment Exchange in: Synthetic Protein Switches Methods and Protocols, Humana Press, 2017.
- 53 A. J. Degrave, J.-H. Ha, S. N. Loh and L. T. Chong, *Nat. Commun.*, 2018, **9**, 1–9.

- 54 J.-H. Ha, S. A. Shinsky and S. N. Loh, *Biochemistry*, 2013, 52, 600–612.
- 55 S. G. Harroun, C. Prévost-Tremblay, D. Lauzon, A. Desrosiers, X. Wang, L. Pedro and A. Vallée-Bélisle, *Nanoscale*, 2018, **10**, 4607–4641.
- 56 K. M. Ahmad, Y. Xiao and H. Tom Soh, *Nucleic Acids Res.*, 2012, **40**, 11777–11783.
- 57 J. Müller, D. Freitag, G. Mayer and B. Pötzsch, J. Thromb. Haemostasis, 2008, 6, 2105–2112.
- 58 J. Müller, B. Wulffen, B. Pötzsch and G. Mayer, *ChemBioChem*, 2007, **8**, 2223–2226.
- 59 C. Riccardi, D. Musumeci, C. Platella, R. Gaglione, A. Arciello and D. Montesarchio, *Int. J. Mol. Sci.*, 2020, **21**, 2–24.
- 60 T. Umehara, K. Fukuda, F. Nishikawa, M. Kohara, T. Hasegawa and S. Nishikawa, *J. Biochem.*, 2005, 137, 339–347.
- 61 Q. W. Hughes, B. T. Le, G. Gilmore, R. I. Baker and R. N. Veedu, *Molecules*, 2017, **22**, 1–8.
- 62 Y. Kim, Z. Cao and W. Tan, Proc. Natl. Acad. Sci. U. S. A., 2008, 105, 5664–5669.
- 63 C. H. Stuart, K. R. Riley, O. Boyacioglu, D. M. Herpai,
 W. Debinski, S. Qasem, F. C. Marini, C. L. Colyer and
 W. H. Gmeiner, *Mol. Ther. Nucleic Acids*, 2016, 5, 1–9.
- 64 C. Riccardi, E. Napolitano, D. Musumeci and D. Montesarchio, *Molecules*, 2020, **25**, 7–15.
- 65 D. Musumeci and D. Montesarchio, *Pharmacol. Ther.*, 2012, **136**, 202–215.
- 66 M. Vorobyeva, P. Vorobjev and A. Venyaminova, *Molecules*, 2016, 21, 14–16.
- 67 B. Deng, Y. Lin, C. Wang, F. Li, Z. Wang, H. Zhang, X.-F. Li and X. C. Le, *Anal. Chim. Acta*, 2014, 837, 1–15.
- 68 G. S. Bang, S. Cho and B.-G. Kim, *Biosens. Bioelectron.*, 2005, 21, 863–870.
- 69 Y. Liu, T. Kwa and A. Revzin, *Biomaterials*, 2012, 33, 1-18.
- 70 V. C. Özalp and T. Schäfer, *Chem. Eur. J.*, 2011, 17, 9893– 9896.
- 71 N. Hamaguchi, A. Ellington and M. Stanton, *Anal. Biochem.*, 2001, **294**, 126–131.
- 72 L. Xue, X. Zhou and D. Xing, Anal. Chem., 2012, 84, 3507– 3513.
- 73 X. Shi, J. Wen, Y. Li, Y. Zheng, J. Zhou, X. Li and H.-Z. Yu, *ACS Appl. Mater. Interfaces*, 2014, **6**, 21788–21797.
- 74 H. Shi, X. He, K. Wang, X. Wu, X. Ye, Q. Guo, W. Tan,
 Z. Qing, X. Yang and B. Zhou, *Proc. Natl. Acad. Sci. U. S. A.*, 2011, 108, 3900–3905.
- 75 Y.-M. Wang, Z. Wu, S.-J. Liu and X. Chu, *Anal. Chem.*, 2015, 87, 6470–6474.
- 76 S. Lin, W. Gao, Z. Tian, C. Yang, L. Lu, J.-L. Mergny, C.-H. Leung and D.-L. Ma, *Chem. Sci.*, 2015, 6, 4284–4290.
- 77 L. Li, Q. Wang, J. Feng, L. Tong and B. Tang, Anal. Chem., 2014, 86, 5101–5107.
- 78 X. Zhang, K. Xiao, L. Cheng, H. Chen, B. Liu, S. Zhang and J. Kong, *Anal. Chem.*, 2014, 86, 5567–5572.
- 79 G. Pelossof, R. Tel-Vered, J. Elbaz and I. Willner, *Anal. Chem.*, 2010, **82**, 4396–4402.

- 80 Y. Liu, J. Yan, M. C. Howland, T. Kwa and A. Revzin, *Anal. Chem.*, 2011, 83, 8286–8292.
- 81 C. Teller, S. Shimron and I. Willner, Anal. Chem., 2009, 81, 9114–9119.
- 82 C. Chen, J. Zhao, J. Jiang and R. Yu, *Talanta*, 2012, 101, 357–361.
- 83 C. Furutani, K. Shinomiya, Y. Aoyama, K. Yamada and S. Sando, *Mol. BioSyst.*, 2010, **6**, 1569–1571.
- 84 X. Tan, Y. Wang, B. A. Armitage and M. P. Bruchez, Anal. Chem., 2014, 86, 10864–10869.
- 85 L. Yang, C. W. Fung, E. J. Cho and A. D. Ellington, *Clin. Microbiol. Rev.*, 2006, **79**, 3329.
- 86 F. J. Hernandez, L. I. Hernandez, A. Pinto, T. Schäfer and V. C. Özalp, *Chem. Commun.*, 2013, **49**, 1285–1287.
- 87 R. M. Dirks and N. A. Pierce, *Proc. Natl. Acad. Sci. U. S. A.*, 2004, **101**, 15275–15278.
- 88 S. Xie and S. P. Walton, Anal. Chim. Acta, 2009, 638, 213– 219.
- 89 J. F. Georges, X. Liu, J. Eschbacher, J. Nichols, M. A. Mooney and A. Joy, *PLoS One*, 2015, **10**, 1–12.
- 90 Q. Zhou, T. Kwa, Y. Gao, Y. Liu, A. Rahimian and A. Revzin, *Lab Chip*, 2014, 14, 276.
- 91 T. Goda and Y. Miyahara, *Biosens. Bioelectron.*, 2011, 26, 3949–3952.
- 92 R. E. Armstrong and G. F. Strouse, *Bioconjugate Chem.*, 2014, 25, 1769–1776.
- 93 S. Liu, Y. Wang, C. Zhang, Y. Lin and F. Li, *Chem. Commun.*, 2013, 49, 2337.
- 94 J. Zhu, L. Zhang, Z. Zhou, S. Dong and E. Wang, *Chem. Commun.*, 2014, 50, 3323.
- 95 X. Cong and M. Nilsen-Hamilton, *Biochemistry*, 2005, 44, 7945–7954.
- 96 Y. Shen, W. Chiuman, J. D. Brennan and Y. Li, *ChemBioChem*, 2006, 7, 1343–1348.
- 97 Z. Tang, P. Mallikaratchy, R. Yang, Y. Kim, Z. Zhu, H. Wang and W. Tan, *J. Am. Chem. Soc.*, 2008, **130**, 11268– 11269.
- 98 C. Wu, T. Chen, D. Han, M. You, L. Peng, S. Cansiz, G. Zhu, C. Li, X. Xiong, E. Jimenez, C. James Yang and W. Tan, *ACS Nano*, 2013, 7, 5724–5731.
- 99 L. J. Nielsen, L. F. Olsen and V. C. Ozalp, ACS Nano, 2010, 4, 4361–4370.
- 100 H. Shi, D. Li, F. Xu, X. He, K. Wang, X. Ye, J. Tang and C. He, *Analyst*, 2014, **139**, 4181–4184.
- 101 B. Jiang, F. Li, C. Yang, J. Xie, Y. Xiang and R. Yuan, Anal. Chem., 2015, 87, 3094–3098.
- 102 Y. Xiao, B. D. Piorek, K. W. Plaxco and A. J. Heeger, J. Am. Chem. Soc., 2005, 127, 17990–17991.
- 103 S. Li, Q. Jiang, S. Liu, Y. Zhang, Y. Tian, C. Song, J. Wang, Y. Zou, G. J. Anderson, J.-Y. Han, Y. Chang, Y. Liu, C. Zhang, L. Chen, G. Zhou, G. Nie, H. Yan, B. Ding and Y. Zhao, *Nat. Biotechnol.*, 2018, **36**, 258–268.
- 104 J. Zhuang, Y. He, G. Chen and D. Tang, *Electrochem. Commun.*, 2014, 47, 25–28.
- 105 A. Dillen, A. Mohrbacher and J. Lammertyn, ACS Sens., 2021, 6, 3684.

- 106 R. Nutiu and Y. Li, *Angew. Chem., Int. Ed.*, 2005, 44, 1061–1065.
- 107 G. Goel, A. Kumar, A. K. Puniya, W. Chen and K. Singh, J. Appl. Microbiol., 2005, 99, 435–442.
- 108 Y. Xiao, A. A. Lubin, A. J. Heeger and K. W. Plaxco, Angew. Chem., Int. Ed., 2005, 44, 5456–5459.
- 109 A. J. Bonham, K. Hsieh, B. S. Ferguson, A. Valleé-Bélisle,
 F. Ricci, H. T. Soh and K. W. Plaxco, *J. Am. Chem. Soc.*,
 2012, 134, 3346–3348.
- 110 A. Idili, N. Arroyo-Currás, K. L. Ploense, A. T. Csordas, M. Kuwahara, T. E. Kippin and K. W. Plaxco, *Chem. Sci.*, 2019, **10**, 8164–8170.
- 111 P. Dauphin-Ducharme, K. Yang, N. Netzahualcóyotl, A.-C. Currás, K. L. Ploense, Y. Zhang, J. Gerson, M. Kurnik, T. E. Kippin, M. N. Stojanovic and K. W. Plaxco, ACS Sens., 2019, 4, 2832–2837.
- 112 H. Li, P. Dauphin-Ducharme, G. Ortega and K. W. Plaxco, J. Am. Chem. Soc., 2017, 139, 11207–11213.
- 113 H. Li, P. Dauphin-Ducharme, N. Arroyo-Currás, C. H. Tran, P. A. Vieira, S. Li, C. Shin, J. Somerson, T. E. Kippin and K. W. Plaxco, *Angew. Chem., Int. Ed.*, 2017, 56, 7492–7495.
- 114 H. Li, N. Arroyo-Currás, D. Kang, F. Ricci and K. W. Plaxco, J. Am. Chem. Soc., 2016, 138, 15809–15812.
- 115 N. Arroyo-Currás, K. Scida, K. L. Ploense, T. E. Kippin and K. W. Plaxco, *Anal. Chem.*, 2017, **89**, 12185–12191.
- 116 P. A. Vieira, C. B. Shin, N. Arroyo-Currás, G. Ortega, W. Li, A. A. Keller, K. W. Plaxco and T. E. Kippin, *Front. Mol. Biosci.*, 2019, 6, 1–10.
- 117 N. Arroyo-Currás, P. Dauphin-Ducharme, G. Ortega,K. L. Ploense, T. E. Kippin and K. W. Plaxco, ACS Sens., 2018, 3, 360–366.
- 118 H. Li, J. Somerson, F. Xia and K. W. Plaxco, *Anal. Chem.*, 2018, **90**, 10641–10645.
- 119 S. Tyagi and F. R. Kramer, *Nat. Biotechnol.*, 1996, **14**, 303–306.
- 120 T. Kihara, N. Yoshida, T. Kitagawa, C. Nakamura, N. Nakamura and J. Miyake, *Biosens. Bioelectron.*, 2010, 26, 1449–1454.
- 121 G. Yao and W. Tan, Anal. Biochem., 2004, 331, 216–223.
- 122 U. Uddayasankar and U. J. Krull, Anal. Chim. Acta, 2013, 803, 113–122.
- 123 X. Dai, W. Yang, E. Firlar, S. A. E. Marras and M. Libera, *Soft Matter*, 2012, **8**, 3036.
- 124 H. Du, M. D. Disney, B. L. Miller and T. D. Krauss, *J. Am. Chem. Soc.*, 2003, **125**, 4012–4013.
- 125 H. Nasef, V. Beni and C. K. O'sullivan, J. Electroanal. Chem., 2011, 662, 322–327.
- 126 Q. Su, D. Wesner, H. Schö and G. Nö, *Langmuir*, 2014, **30**, 14360–14367.
- 127 H. T. Ngo, H.-N. Wang, A. M. Fales and T. Vo-Dinh, Anal. Chem., 2013, 85, 6378–6383.
- 128 Molecular Beacons, ed. C. J. Yang and W. Tan, Springer, 2013.
- 129 N. C. Seeman and H. F. Sleiman, *Nat. Rev. Mater.*, 2017, 3, 1–23.

- 130 S. Tombelli, M. Minunni and M. Mascini, *Biosens. Bioelectron.*, 2005, 20, 2424–2434.
- 131 Z. Zhuo, Y. Yu, M. Wang, J. Li, Z. Zhang, J. Liu, X. Wu, A. Lu, G. Zhang and B. Zhang, *Int. J. Mol. Sci.*, 2017, 18, 2– 19.
- 132 J. Tan, M. Zhao, J. Wang, Z. Li, L. Liang, L. Zhang, Q. Yuan and W. Tan, Angew. Chem., Int. Ed., 2019, 58, 1621–1625.
- 133 D. Shangguan, Y. Li, Z. Tang, Z. C. Cao, H. W. Chen, P. Mallikaratchy, K. Sefah, J. Yang and W. Tan, *Proc. Natl. Acad. Sci. U. S. A.*, 2006, **103**, 11838–11843.
- 134 R. Nutiu and Y. Li, J. Am. Chem. Soc., 2003, 125, 4771-4778.
- 135 F. Wang, X. Liu and I. Willner, Angew. Chem., Int. Ed., 2015, 54, 1098-1129.
- 136 Y. Zhang, C. Ge, C. Zhu and K. Salaita, *Nat. Commun.*, 2014, 5, 1–10.
- 137 A. Valleé-Beíisle, F. Ricci, T. Uzawa, F. Xia and K. W. Plaxco, J. Am. Chem. Soc., 2012, 134, 15197–15200.
- 138 S. Ranallo, M. Rossetti, K. W. Plaxco, A. Vallée-Bélisle and F. Ricci, Angew. Chem., Int. Ed., 2015, 54, 13214–13218.
- 139 Y.-T. Lin, R. Vermaas, J. Yan, A. M. De Jong and M. W. J. Prins, ACS Sens., 2021, 6, 1980–1986.
- 140 J. Corzo, Biochem. Mol. Biol. Educ., 2006, 34, 413-416.
- 141 B. D. Wilson, A. A. Hariri, I. A. P. Thompson, M. Eisenstein and H. T. Soh, *Nat. Commun.*, 2019, **10**, 1–9.
- 142 E. M. W. M. Van Dongen, T. H. Evers, L. M. Dekkers, E. W. Meijer, L. W. J. Klomp and M. Merkx, *J. Am. Chem. Soc.*, 2007, **129**, 3494–3495.
- 143 P. R. Mallikaratchy, A. Ruggiero, J. R. Gardner, V. Kuryavyi, W. F. Maguire, M. L. Heaney, M. R. McDevitt, D. J. Patel and D. A. Scheinberg, *Nucleic Acids Res.*, 2011, 39, 2458–2469.
- 144 K. Horikawa, Y. Yamada, T. Matsuda, K. Kobayashi,
 M. Hashimoto, T. Matsu-ura, A. Miyawaki, T. Michikawa,
 K. Mikoshiba and T. Nagai, *Nat. Methods*, 2010, 7, 729–732.
- 145 H. Bischof, M. Rehberg, S. Stryeck, K. Artinger, E. Eroglu, M. Waldeck-Weiermair, B. Gottschalk, R. Rost, A. T. Deak, T. Niedrist, N. Vujic, H. Lindermuth, R. Prassl, B. Pelzmann, K. Groschner, D. Kratky, K. Eller, A. R. Rosenkranz, T. Madl, N. Plesnila, W. F. Graier and R. Malli, *Nat. Commun.*, 2017, 8, 1–12.
- 146 A. Porchetta, A. Vallée-Bélisle, K. W. Plaxco and F. Ricci, J. Am. Chem. Soc., 2012, 134, 20601–20604.
- 147 S. Bissonnette, E. Del Grosso, A. J. Simon, K. W. Plaxco, F. Ricci and A. Valleé-Bélisle, *ACS Sens.*, 2020, 5, 1937–1942.
- 148 N. Bhalla, P. Jolly, N. Formisano and P. Estrela, *Essays Biochem.*, 2016, **60**, 1–8.
- 149 H.-B. Huang, J.-M. Peng, L. Weng, C.-Y. Wang, W. Jiang and B. Du, *Ann. Intensive Care*, 2017, 7, 1–10.
- 150 J. Somerson and K. W. Plaxco, *Molecules*, 2018, 23, 1–7.
- 151 H. Ma, C. Ó'Fágáin and R. O'kennedy, *Biochimie*, 2020, 117, 213–225.
- 152 S. Warszawski, A. B. Katz, R. Lipsh, L. Khmelnitsky, G. Ben Nissanid, G. Javittid, O. Dym, T. Unger, O. Knop,

S. Albeck, R. Diskinid, D. Fass, M. Sharon and S. J. Fleishmanid, *PLOS Comput. Biol.*, 2019, **15**, 1–24.

- 153 S. Wang, M. Liu, D. Zeng, W. Qiu, P. Ma, Y. Yu, H. Chang and Z. Sun, *Proteins*, 2014, **82**, 2620–2630.
- 154 S. Ni, H. Yao, L. Wang, J. Lu, F. Jiang, A. Lu and G. Zhang, *Int. J. Mol. Sci.*, 2017, **18**, 1–21.
- 155 C. Kratschmer and M. Levy, *Nucleic Acid Ther.*, 2017, 27, 335–344.
- 156 P. G. Chandler, S. S. Broendum, B. T. Riley, M. A. Spence, C. J. Jackson and A. M. Buckle, *Strategies for Increasing Protein Stability in: Protein Nanotechnology*, Humana, New York, 2020, vol. 2073.
- 157 J. P. Elskens, J. M. Elskens and A. Madder, *Int. J. Mol. Sci.*, 2020, **21**, 1–31.
- 158 S. Ni, H. Yao, L. Wang, J. Lu, F. Jiang, A. Lu and G. Zhang, *Int. J. Mol. Sci.*, 2017, **18**, 1–21.
- 159 K. E. Maier and M. Levy, *Mol. Ther. Methods Clin. Dev.*, 2016, 3, 16014.
- 160 S. Y. Tan, C. Acquah, A. Sidhu, C. M. Ongkudon, L. S. Yon and M. K. Danquah, *Crit. Rev. Anal. Chem.*, 2016, 46, 521–537.
- 161 S. Gao, X. Zheng, B. Jiao and L. Wang, *Anal. Bioanal. Chem.*, 2016, **408**, 4567–4573.
- 162 S. Ni, Z. Zhuo, Y. Pan, Y. Yu, F. Li, J. Liu, L. Wang, X. Wu, D. Li, Y. Wan, L. Zhang, Z. Yang, B.-T. Zhang, A. Lu and G. Zhang, *Appl. Mater. Interfaces*, 2021, **13**, 9500–9519.
- 163 M. A. Vorobyeva, A. S. Davydova, P. E. Vorobjev, D. V. Pyshnyi and A. G. Venyaminova, *Int. J. Mol. Sci.*, 2018, **19**, 1–21.
- 164 E. Eremeeva, A. Fikatas, L. Margamuljana, M. Abramov,
 D. Schols, E. Groaz and P. Herdewijn, *Nucleic Acids Res.*, 2019, 47, 4927–4939.
- 165 M. R. Dunn, C. M. Mccloskey, P. Buckley, K. Rhea and J. C. Chaput, J. Am. Chem. Soc., 2020, 142, 7721–7724.
- 166 M. Varada, M. Aher, N. Erande, V. A. Kumar and M. Fernandes, *ACS Omega*, 2020, **5**, 498–506.
- 167 M. Merkx, M. V. Golynskiy, L. H. Lindenburg and J. L. Vinkenborg, *Biochem. Soc. Trans.*, 2013, 41, 1201–1205.
- 168 Fluorescent Protein-Based Biosensors Methods and Protocols, ed. J. Zhang, Q. Ni and R. H. Newman, Humana Press, 2014.
- 169 W. M. Keck, J. Cell Biol., 2003, 160, 629-633.

- 170 Principles of Fluorescence Spectroscopy, ed. J. R. Lakowicz, Springer, 3rd edn, 2006.
- 171 X. Miao, X. Guo, Z. Xiao and L. Ling, *Biosens. Bioelectron.*, 2014, **59**, 54–57.
- 172 B. S. Ferguson, D. A. Hoggarth, D. Maliniak, K. Ploense, R. J. White, N. Woodward, K. Hsieh, A. J. Bonham, M. Eisenstein, T. Kippin, K. W. Plaxco and H. T. Soh, *Sci. Transl. Med.*, 2013, 5, 213–165.
- 173 C. Parolo, A. Idili, G. Ortega, A. Csordas, A. Hsu, N. Netzahualcóyotl, A.-C. Currás, Q. Yang, B. S. Ferguson, J. Wang and K. W. Plaxco, ACS Sens., 2020, 5, 1877– 1881.
- 174 A. Idili, J. Gerson, T. Kippin and K. W. Plaxco, *Anal. Chem.*, 2021, **93**, 4023–4032.
- 175 D. Grieshaber, R. MacKenzie, J. Vörös and E. Reimhult, Sensors, 2008, 8, 1400-1458.
- 176 H. Hasegawa, K.-I. Taira, K. Sode and K. Ikebukuro, *Sensors*, 2008, **8**, 1090–1098.
- 177 T. Hianik, I. Grman and I. Karpisova, *Chem. Commun.*, 2009, 6303–6305.
- 178 S. Campuzano, M. Pedrero, P. Yáñez-Sedeño and J. M. Pingarrón, *Int. J. Mol. Sci.*, 2019, **20**, 1–19.
- 179 P.-H. Lin and B.-R. Li, Analyst, 2020, 145, 1110–1120.
- 180 J. Xu and H. Lee, Chemosensors, 2020, 8, 66.
- 181 J. Breault-Turcot and J.-F. Masson, *Chem. Sci.*, 2015, 6, 4247–4254.
- 182 G. Rong, S. R. Corrie and H. A. Clark, ACS Sens., 2018, 2, 327–338.
- 183 A. Idili, K. W. Plaxco, A. Vallée-Bélisle and F. Ricci, ACS Nano, 2013, 7, 10863–10869.
- 184 F. Ricci, A. Valleé-Beíisle, A. J. Simon, A. Porchetta, K. W. Plaxco, B. Science and E. Program, *Acc. Chem. Res.*, 2016, 49, 21.
- 185 A. M. Downs, J. Gerson, M. Nur Hossain, K. Ploense, M. Pham, H.-B. Kraatz, T. Kippin and K. W. Plaxco, ACS Sens., 2021, 6, 2299–2306.
- 186 Eccrine Systems, Inc., https://www.eccrinesystems.com/.
- 187 Creative Commons Attribution 4.0, http://creativecommons.org/licenses/by/4.0/.
- 188 Helia Biomonitoring, https://www.heliabiomonitoring.com/