

# Re-engineering Electrochemical Biosensors To Narrow or Extend Their Useful Dynamic Range\*\*

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The use of electrode-immobilized biomolecules, such as proteins and nucleic acids, is a common feature among many emerging biotechnological applications. For example, the specificity, affinity, and versatility of biomolecular recognition has been exploited for the development of a wide range of electrochemical biosensors that show promise for the detection of many clinically and industrially important analytes.<sup>[1,2]</sup> Such “bioelectronic interfaces” also form the basis of biofuel cells<sup>[3]</sup> and molecular logic gates,<sup>[4]</sup> applications that have attracted significant recent attention. Interest in the applications of surface-electrode-bound biomolecular systems is thus rapidly growing.

Despite their often impressive performances, technologies based on biomolecular recognition suffer from the inherent limitation of single-site binding represented by its fixed dose–response curve. That is, single-site binding is almost invariably characterized by a fixed, hyperbolic relationship between the target concentration and receptor binding (the Langmuir isotherm) for which the dynamic range (here defined as the range of target concentrations corresponding to receptor occupancies between 10% and 90%) spans an 81-fold range of target concentrations.<sup>[5]</sup> This fixed dynamic range reduces the utility of electrochemical biosensors in applications such as viral load monitoring, in which the concentration of the target molecule can vary over many orders of magnitude. It likewise limits the usefulness of biosensors in applications requiring high sensitivity (a steep relationship between target concentration and output signal), such as in the monitoring of drugs with narrow therapeutic windows. Thus, the possibility

to arbitrarily extend or narrow this fixed dynamic range would prove advantageous in several biosensing applications. For example, the ability to extend the dynamic range of biorecognition would likely improve the efficiency of biofuel cells,<sup>[3]</sup> and the ability to narrow the dynamic range would reduce noise in molecular logic gates.<sup>[4]</sup>

Recently we have shown that some of the mechanisms employed by nature to alter the otherwise fixed dynamic range of single-site binding can also be used to broaden and narrow the dose–response curves of solution-phase, optical biosensors.<sup>[6]</sup> For example, by combining biosensors of identical specificity but differing affinity we have expanded the useful 81-fold range of a molecular beacon, a model solution-phase optical biosensor, by more than 10 000-fold.<sup>[6]</sup> In parallel we have also adapted the sequestration mechanism, often employed by nature to generate “ultrasensitive” genetic networks, to narrow the dynamic range of the same biosensor down to 5-fold, thus greatly increasing the sensitivity of this category of biosensors.<sup>[6]</sup>

Following the previously mentioned work we demonstrate here the application of these approaches to modify the dynamic range of reagentless, electrochemical biosensors. Specifically, we have used these approaches to arbitrarily narrow and broaden the useful dynamic ranges of electrochemical “E-DNA” sensors,<sup>[7]</sup> a class of structure-switching DNA probes that enable the single-step detection of specific oligonucleotides directly in complex media such as blood serum and environmental samples (Figure 1).<sup>[7b,8]</sup>

E-DNA sensors comprise a redox-reporter-modified stem–loop DNA probe (receptor) attached to an interrogating electrode.<sup>[7b]</sup> In the absence of target, the formation of the stem holds the redox reporter in proximity to the electrode, supporting efficient electron transfer (ET). Upon hybridization with a complementary oligonucleotide target, the terminus of the probe is pushed away from the electrode, which, in turn, hinders the efficiency with which electrons are transferred to the electrode and reduces the observed Faradaic current (Figure 1A). The first strategy we have employed to narrow or extend the dynamic range of this sensor requires the availability of probes directed against the same target molecule but differing in affinity.<sup>[6]</sup> For E-DNA sensors this can be achieved by using a set of stem–loop probes that share a common recognition loop, and thus target the same DNA sequence, but differ in the stability of their double-stranded stems. In this way we can arbitrarily vary the target–probe dissociation constant ( $K_d$ )—here over three orders of magnitude—without affecting the target-recognizing loop, and thus without changing the probe’s sequence specificity (Figure 1B).<sup>[6,9]</sup>


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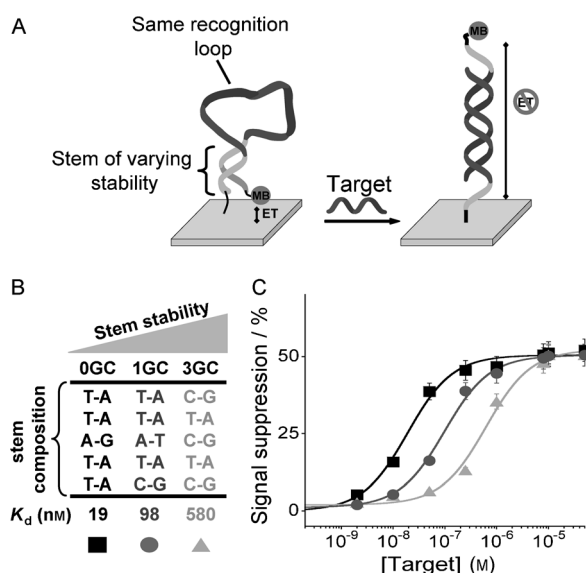
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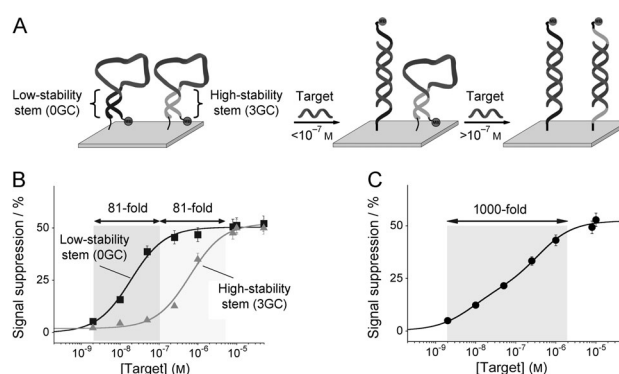


**Figure 1.** A) E-DNA sensors consist of a stem-loop DNA modified with a redox reporter (here methylene blue, MB) and attached to an interrogating gold electrode through an introduced thiol group.<sup>[7b]</sup> This probe undergoes a large-scale conformational switch upon hybridization with a DNA complementary to the loop, leading to large change in Faradaic current from the redox reporter. B) The affinity of such “switch-based” probes can be rationally tuned by many orders of magnitude, without affecting their specificity, by simply altering the stability of their nonbinding, nonsignaling state (e.g., by varying the stability of the E-DNA probe’s stem by changing the number of GC base pairs).<sup>[9]</sup> C) Here we have employed a set of three E-DNA probes sharing a common recognition element but with target affinities spanning almost three orders of magnitude. Error bars in this figure and in the following figures represent the average and standard deviations of measurements performed on at least three independently sensors.

While the affinity of E-DNA sensors is easily tuned by changes in their stem stability, reaching this objective can be more challenging for recognition elements displaying more complex structures. A number of strategies have, however, been reported for engineering (and tuning) similar switching mechanisms for aptamers, aptazymes, and even proteins.<sup>[10]</sup> Loh and co-workers, for example, have recently demonstrated a generic strategy to design novel protein-based switches, termed “alternate frame folding”, in which a portion of a protein’s sequence is duplicated in order to stabilize an alternative, nonbinding, circularly permuted conformation.<sup>[10d]</sup> Proteins and nucleic acids can also be engineered to undergo folding-induced conformational changes by the introduction of destabilizing mutations (typically remote from the target binding site so as to ensure that specificity is retained) that push the folding equilibrium toward the nonbinding, unfolded state. In this way, binding is coupled to a conformational change (folding) and simultaneously binding affinity is coupled to folding stability.<sup>[10]</sup>

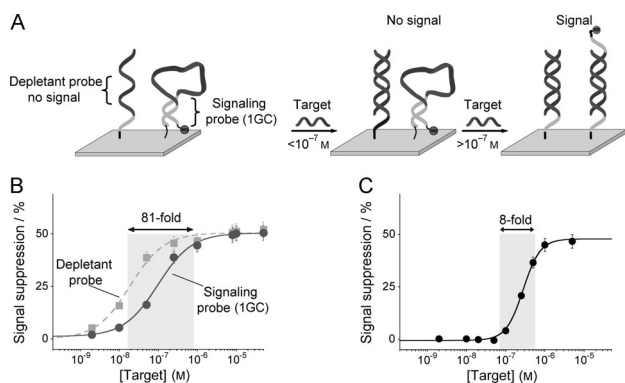
As noted above, traditional E-DNA sensors exhibit a useful dynamic range of 81-fold (Figure 1 C). We can extend this useful dynamic range by co-immobilizing two E-DNA probes differing in affinity for their (common) target

DNA onto a single electrode. (Of note, the E-DNA probes we have employed are both modified with the same methylene blue redox reporter and thus they both signal at the same redox potential and with the same relative signal change at saturating target concentrations; see Figure 1 A). To achieve optimal log-linear behavior in the modified sensor, the affinities of the two probes should differ by approximately 30-fold.<sup>[6]</sup> For example, by combining on the same electrode surface an equimolar concentration of the low-stability 0GC stem-loop probe (Figure 1 B), which exhibits a dissociation constant of 19 nm, with the more stable 3GC stem-loop probe, exhibiting a dissociation constant of 580 nm, we expand the normally 81-fold dynamic range to approximately 1000-fold (corresponding to a target concentration between 2 nM and 2000 nM) and achieve excellent linearity on a log[concentration] plot ( $R^2 = 0.978$ ; Figure 2 C).



**Figure 2.** By employing a pair of signaling probes differing in affinity we can broaden the dynamic range of E-DNA sensors. A) We co-immobilized a relatively low-affinity E-DNA probe (probe 3GC,  $K_d = 580$  nm) and a higher affinity E-DNA probe (probe 0GC,  $K_d = 19$  nm) in a 1:1 ratio on a single-electrode surface. B) The useful dynamic range of these individual probes spans an 81-fold range of target concentrations over two distinct concentration regimes. C) With this strategy the resulting dose–response curve is extended and spans a 1000-fold range of target concentrations.

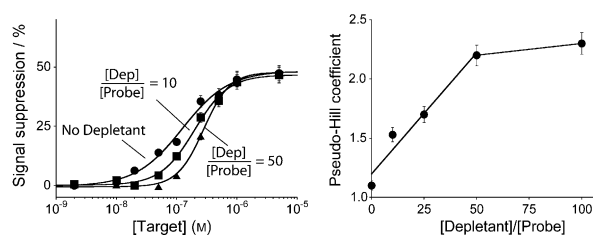
The availability of probes retaining a common specificity profile but differing in affinity also provides a means of narrowing the useful dynamic range of E-DNA sensors, thus enhancing their sensitivity (the steepness of the input/output curve) and improving their ability to measure small changes in concentration. Specifically, we adapted here the sequestration mechanism used by nature to improve the sensitivity of many regulatory cascades through the competition between a high-affinity, but not signaling, recognition element (the depletant) and a low-affinity signaling receptor.<sup>[11]</sup> To demonstrate this, we co-immobilized two E-DNA probes, the stem-loop sequence 1GC and an equivalent, fully linear probe lacking a complementary stem, both of which are complementary to the same 13-base target sequence (Figure 3 A). Because the linear probe does not undergo a binding-induced conformational change, its affinity for the DNA target is significantly greater than that of the stem-loop 1GC probe. In this application the higher-affinity linear probe lacks any redox reporter (methylene blue) and thus the hybrid-



**Figure 3.** The sequestration mechanism can be used to dramatically narrow the useful dynamic range of an E-DNA sensor, thus greatly improving its sensitivity. A) A low-affinity, signaling E-DNA probe is co-immobilized on a single-electrode surface with a higher affinity probe (depletant) which, lacking the redox reporter, does not signal upon binding its target. B) At low concentrations the target preferentially binds the depletant, which removes (sequesters) target from the sample without generating a signal. When the total amount of the target exceeds that of the depletant, a threshold response is achieved above which further addition of target dramatically raises the relative concentration of free target. This gives rise to a much steeper dose–response curve than would occur in the absence of a depletant. C) Using this approach we have narrowed the useful dynamic range of an unmodified E-DNA sensor (81-fold) to a mere 8-fold, thus increasing its sensitivity by an order of magnitude.

ization of the target to this probe does not produce any measurable signal change. This linear probe therefore acts as the depletant, “silently” sequestering the target until the threshold concentration is surpassed.<sup>[11c]</sup> The lower-affinity signaling probe (1GC) is only activated (and thus only signals the presence of the target) when the depletant is saturated and this threshold is surpassed (Figure 3B). Using this approach we convert the hyperbolic dose–response curve of a traditional E-DNA sensor into an ultrasensitive response with a dynamic range spanning only an 8-fold range of target concentrations, an order of magnitude narrower than the dynamic range of a traditional E-DNA sensor (Figure 3C).

The sensitivity achieved by means of the sequestration mechanism depends on the relative amounts of depletant and signaling probe,<sup>[11c]</sup> and thus on the relative density of the two on the sensor’s surface. To demonstrate this we have altered the ratio of probe and depletant on our sensors by altering the depletant/probe concentration ratio employed during sensor fabrication.<sup>[12]</sup> To compare sensors fabricated using differing depletant/probe ratios we fitted their input–output curves to the Hill equation, which, although physically meaningful only when used to describe the ultrasensitivity associated with allosteric cooperativity,<sup>[13]</sup> provides a convenient means of quantifying the steepness of a binding curve. As expected, we observe a “pseudo-Hill” coefficient near unity ( $1.1 \pm 0.1$ ) for sensors lacking the depletant. Upon the addition of the depletant probe, this coefficient increases monotonically with increasing depletant/probe ratios until it plateaus at 2.3 for ratios above 50 (Figure 4, right). The highest pseudo-Hill coefficient we have achieved compresses the 81-fold useful dynamic range of an unmodified E-DNA sensor to only 8-

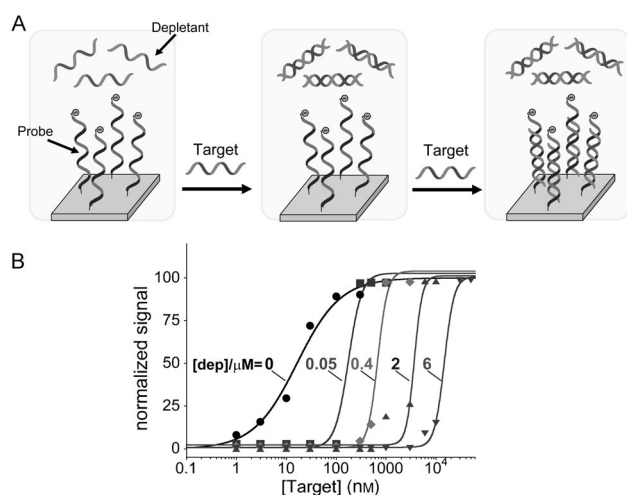


**Figure 4.** The sensitivity (i.e., steepness of the dose–response curve) achieved using the sequestration mechanism depends on the ratio of depletant to probe employed during sensor fabrication. To show this we have fitted our data to obtain pseudo-Hill coefficients, which, although our system is not classically cooperative, are analogous to the Hill coefficients commonly used to describe cooperative enzymatic systems.<sup>[13]</sup> We find that the pseudo-Hill coefficient increases monotonically with this ratio until plateauing at values above 50.

fold, significantly increasing the steepness of the dose–response curve of the sensor and, in turn, improving its ability to detect smaller relative changes in target concentration.

We must note that this strategy is not without limitations. Specifically, the sequestration approach only works for fixed, small sample volumes (here we have employed 3  $\mu\text{L}$  samples) in order to avoid the “premature” saturation of the fixed number of depletant molecules on the electrode surface. Moreover, as discussed before, the probe/depletant ratio on the electrode surface is a key factor which must be carefully controlled. We did so by assuming that the density ratios on the surface are linearly correlated with the concentration ratios deployed in solution during deposition. This (seemingly reasonable) assumption appears to be confirmed by the linear dependence of the absolute current signals (which are correlated to surface density<sup>[14]</sup>) versus [probe]/[depletant] ratio (Figure S11). However, we note that this correlation could be more complicated for less defined recognition elements which can induce a nonlinear immobilization of probe and depletant (Figure S12). Finally, the approach proposed is limited to [depletant]/[probe] ratios below 100:1; over this limit, the density of the probe, and therefore its signal, becomes so low that it is not possible to observe a measurable current (Figure S13). In order to circumvent these limitations, we also propose an alternative strategy where a fixed concentration of depletant is exogenously added to the mixture solution (Figure 5A) thus overcoming possible problems arising from uncontrolled density ratios. Moreover, because the depletant is now free to diffuse in solution, its affinity for the target is greatly increased over that of the surface-bound probe. This makes it possible to use the same recognition element for both the depletant and the signaling probe, thus making the approach also suitable for more complex biorecognition elements whose affinity cannot be easily tuned.

The unattached “non-signaling” depletant probe sequesters the target DNA until a threshold level is reached (fixed by the depletant concentration) above which further increase in target concentration results in a steep dose–response curve (Figure 5B). To improve the convenience of this approach, the specific amount of depletant was added by noncovalently



**Figure 5.** To overcome the limitations inherent to the surface-attached depletants (which are easily saturated), we also show that the depletant probe can be simply added in solution at a fixed concentration. A) Here we use an unlabeled nonsignaling probe (with the exact same sequence of the signaling redox-labeled probe) that sequesters the target DNA up to a threshold level (fixed by the depletant concentration in solution) over which further increase in target concentration results in a steep dose–response curve. Because the depletant is free in solution, it rapidly reacts with the target (and with higher affinity) before this later can diffuse to the electrode surface and “activate” the signaling probe. B) By using different concentrations of depletant in the reaction mix (0, 0.05, 0.4, 2, 6  $\mu\text{M}$ ) we achieve steeper transitions than those observed with the depletant co-immobilized with the probe and we can also easily tune the threshold level at which we observe the sharp digital-like response of the sensor.

absorbing it on the electrode surface. As soon as the sample is applied on the electrode surface, the depletant diffuses in solution, which maintains the single-step convenience of the reagentless sensor. With this strategy we have built an array of electrodes, each of which containing various concentration of depletant, and thus various target threshold with pseudo-Hill coefficient values between 3 and 4 and a dynamic range spanning only 2- to 3-fold of target concentration (Figure 5).

Here we have demonstrated convenient methods to extend and narrow the useful dynamic range of a model electrochemical DNA sensor. We did so by combining DNA probes of different target affinities but with similar specificity on the same electrode.<sup>[6]</sup> Employing a pair of signaling probes with dissociation constants differing by approximately an order of magnitude, we produced a pseudo-log linear response spanning three orders of magnitude in target concentration. And, by employing a pair of probes in which the higher-affinity probe is nonsignaling we have narrowed the useful dynamic range of an E-DNA sensor to only an 8-fold range of target concentrations, significantly improving its sensitivity. Moreover, because the relevant probes are all strongly chemisorbed onto their interrogating electrodes, the modified sensors remain reagentless, reusable, highly selective electrochemical devices readily amenable to lab-on-a-chip applications and point-of-care use.<sup>[7b]</sup> To overcome possible limitations in the application of the strategy employed to narrow the sensor’s dynamic range, we have

also demonstrated an alternative “sequestration” approach where the depletant is added in solution. A great advantage of this strategy is that it doesn’t require variants of the receptor with different affinities: the depletant displays a higher affinity than the probe itself since it is free to diffuse in solution.

Our work is not the first to rationally extend the useful dynamic range of an electrochemical biosensor. Our approach, however, appears to be easier to implement than other, previously reported approaches to this end. These include approaches based on the use of multiple sensors combined with a chemometric strategy<sup>[15]</sup> and on the use of diffusion barrier membranes.<sup>[16]</sup> In addition, our use of sets of recognition elements differing only in affinity, and not specificity, represents an advantage over other approaches, such as those utilizing combinations of enzymes differing in both affinity and specificity,<sup>[17]</sup> because we could achieve a fixed specificity profile across the sensor’s entire dynamic range.

In contrast to broadening the useful dynamic range of electrochemical biosensors, an established goal in the literature, we are not aware of any prior reports regarding the narrowing of their dynamic range. The steep dose–response curves we achieved open the door to a number of sensing applications requiring high sensitivity and a low signal-to-noise ratio at certain specific target concentrations. Of note, compared to a sensor that responds gradually to target inputs, an ultrasensitive electrochemical sensor would be far more useful to generate electrochemical logic gates, ideas that have attracted significant recent interest.<sup>[18]</sup>

The approaches demonstrated here are general, and can be applied to extend or narrow the dynamic range of other electrochemical biosensors provided that the affinities of the biomolecular recognition elements upon which they are based can be appropriately tuned. This is the case of, for example, structure-switching ribozymes and aptamers whose affinity have been rationally modulated through quantitative and predictive model to meet certain performance requirements.<sup>[19]</sup> Despite being a more challenging task, rational and semirational engineering strategies are also available to tune the affinity of proteins or more complex recognition elements.<sup>[10]</sup> Indeed, several examples have been reported, which suggest that our approach to affinity tuning may be broadly applicable.<sup>[10]</sup>

The ability to broaden or narrow the dynamic range of biomolecular recognition could also be of utility in biotechnologies beyond biosensing. The fixed dynamic range of single-site binding, for example, limits the utility of biomolecular recognition in biofuel cells, for which wider dynamic range equates with better power efficiencies.<sup>[3]</sup> It also limits the performance of bio-electronic “logic gates” used in biocomputing, as a steeper, nearly all-or-none “digital” response could significantly reduce the noise floor in such systems.<sup>[18a,4]</sup>

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- [1] a) J. Wang, *Chem. Rev.* **2008**, *108*, 814–825; b) J. Wang, *Biosens. Bioelectron.* **2006**, *21*, 1887–1892.
- [2] a) T. G. Drummond, M. G. Hill, J. K. Barton, *Nat. Biotechnol.* **2003**, *21*, 1192–1199; b) E. Palecek, M. Fojta, *Anal. Chem.* **2001**, *73*, 74a–83a.
- [3] R. A. Bullen, T. C. Arnot, J. B. Lakeman, F. C. Walsh, *Biosens. Bioelectron.* **2006**, *21*, 2015–2045.
- [4] a) V. Privman, V. Pedrosa, D. Melnikov, M. Pita, A. Simonian, E. Katz, *Biosens. Bioelectron.* **2009**, *25*, 695–701; b) J. Wang, E. Katz, *Isr. J. Chem.* **2011**, *51*, 141–150.
- [5] a) D. E. Koshland, *The molecular basis for enzyme regulation, Vol. 1*, Academic Press, New York, **1970**; b) A. Goldbeter, D. E. Koshland, *Proc. Natl. Acad. Sci. USA* **1981**, *78*, 6840–6844; c) D. E. Koshland, A. Goldbeter, J. B. Stock, *Science* **1982**, *217*, 220–225; d) J. E. Ferrell, Jr., *Trends Biochem. Sci.* **1996**, *21*, 460–466.
- [6] A. Vallee-Belisle, F. Ricci, K. W. Plaxco, *J. Am. Chem. Soc.* **2012**, *134*, 2876–2879.
- [7] a) C. H. Fan, K. W. Plaxco, A. J. Heeger, *Proc. Natl. Acad. Sci. USA* **2003**, *100*, 9134–9137; b) F. Ricci, K. W. Plaxco, *Microchim. Acta* **2008**, *163*, 149–155.
- [8] A. A. Lubin, K. W. Plaxco, *Acc. Chem. Res.* **2010**, *43*, 496–505.
- [9] A. Vallée-Bélisle, F. Ricci, K. W. Plaxco, *Proc. Natl. Acad. Sci. USA* **2009**, *106*, 13802–13807.
- [10] a) A. Vallee-Belisle, K. W. Plaxco, *Curr. Opin. Struct. Biol.* **2010**, *20*, 518–526; b) M. V. Golynskiy, M. S. Koay, J. L. Vinkenborg, M. Merckx, *ChemBioChem* **2011**, *12*, 353–361; c) G. Guntas, T. J. Mansell, J. R. Kim, M. Ostermeier, *Proc. Natl. Acad. Sci. USA* **2005**, *102*, 11224–11229; d) M. M. Stratton, S. N. Loh, *Protein Sci.* **2011**, *20*, 19–29; e) D. Strickland, X. L. Yao, G. Gawlak, M. K. Rosen, K. H. Gardner, T. R. Sosnick, *Nat. Methods* **2010**, *7*, 623–626; f) A. E. Palmer, M. Giacomello, T. Kortemme, S. A. Hires, V. Lev-Ram, D. Baker, R. Y. Tsien, *Chem. Biol.* **2006**, *13*, 521–530; g) J. E. Kohn, K. W. Plaxco, *Proc. Natl. Acad. Sci. USA* **2005**, *102*, 10841–10845; h) J. S. Marvin, H. W. Hellinga, *Nat. Struct. Biol.* **2001**, *8*, 795–798.
- [11] a) N. E. Buchler, M. Louis, *J. Mol. Biol.* **2008**, *384*, 1106–1119; b) N. E. Buchler, F. R. Cross, *Mol. Syst. Biol.* **2009**, *5*, 272; c) F. Ricci, A. Vallee-Belisle, K. W. Plaxco, *PLoS Comput. Biol.* **2011**, *7*, e1002171.
- [12] a) Y. Xiao, R. Y. Lai, K. W. Plaxco, *Nat. Protoc.* **2007**, *2*, 2875–2880; b) A. A. Rowe, R. J. White, A. J. Bonham, K. W. Plaxco, *J. Vis. Exp.* **2011**, e2922; c) D. Kang, X. L. Zuo, R. Q. Yang, F. Xia, K. W. Plaxco, R. J. White, *Anal. Chem.* **2009**, *81*, 9109–9113.
- [13] A. V. Hill, *J. Physiol.* **1910**, *40*, iv–vii.
- [14] F. Ricci, R. Y. Lai, A. J. Heeger, K. W. Plaxco, J. J. Sumner, *Langmuir* **2007**, *23*, 6827.
- [15] E. Chow, E. L. S. Wong, O. Pascoe, D. B. Hibbert, J. J. Gooding, *Anal. Bioanal. Chem.* **2007**, *387*, 1489–1498.
- [16] W. H. Mullen, F. H. Keedy, S. J. Churchouse, P. M. Vadgama, *Anal. Chim. Acta* **1986**, *183*, 59–66.
- [17] T. Yamazaki, K. Kojima, K. Sode, *Anal. Chem.* **2000**, *72*, 4689–4693.
- [18] a) M. N. Stojanovic, D. Stefanovic, *Nat. Biotechnol.* **2003**, *21*, 1069–1074; b) A. Saghatelian, N. H. Volcker, K. M. Guckian, V. S. Y. Lin, M. R. Ghadiri, *J. Am. Chem. Soc.* **2003**, *125*, 346–347; c) L. M. Adleman, *Science* **1994**, *266*, 1021–1024; d) F. Xia, X. L. Zuo, R. Q. Yang, R. J. White, Y. Xiao, D. Kang, X. O. Gong, A. A. Lubin, A. Vallee-Belisle, J. D. Yuen, B. Y. B. Hsu, K. W. Plaxco, *J. Am. Chem. Soc.* **2010**, *132*, 8557–8559.
- [19] a) N. Hamaguchi, A. Ellington, M. Stanton, *Anal. Biochem.* **2001**, *294*, 126–131; b) X. Chen, A. D. Ellington, *PLoS Comput. Biol.* **2009**, *5*, e1000620; c) C. L. Beisel, C. D. Smolke, *PLoS Comput. Biol.* **2009**, *5*, e1000363; d) P. S. Lau, B. K. Coombes, Y. F. Li, *Angew. Chem.* **2010**, *122*, 8110–8114; *Angew. Chem. Int. Ed.* **2010**, *49*, 7938–7942; e) K. Sefah, J. A. Phillips, X. L. Xiong, L. Meng, D. Van Simaey, H. Chen, J. Martin, W. H. Tan, *Analyst* **2009**, *134*, 1765–1775.