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Protected DNA strand displacement for enhanced single nucleotide discrimination in double-stranded DNA

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Single nucleotide polymorphisms (SNPs) are a prime source of genetic diversity. Discriminating between different SNPs provides an enormous leap towards the better understanding of the uniqueness of biological systems. Here we report on a new approach for SNP discrimination using toehold-mediated DNA strand displacement. The distinctiveness of the approach is based on the combination of both 3- and 4-way branch migration mechanisms, which allows for reliable discrimination of SNPs within double-stranded DNA generated from real-life human mitochondrial DNA samples. Aside from the potential diagnostic value, the current study represents an additional way to control the strand displacement reaction rate without altering other reaction parameters and provides new insights into the influence of single nucleotide substitutions on 3- and 4-way branch migration efficiency and kinetics.

S pecificity of nucleic acid hybridization underpins the precise transfer of genetic information in living organisms through the generations. At the same time, diversity and evolution of life is closely related to DNA sequence alterations. In medical diagnostics and human identification, precise nucleic acid hybridization and sequence discrimination between single nucleotides (or a single nucleotide polymorphism (SNP)) is of great importance and significance. Much effort has been made in order to increase the discrimination power of currently available nucleic acid tests (for review see Ref. 1). Among these attempts are the developments of molecular beacons²⁻⁸, conformationally constrained binary probes^{9–11}, probes modified with artificial nucleotides such as peptide nucleic acids (PNAs)^{8,12-14} and locked nucleic acids (LNAs)¹⁴⁻¹⁷ etc. Thermodynamic approaches such as an elevated hybridization temperature (close to the duplex melting point)^{18,19} or high concentrations of chaotropic agents in hybridization buffers^{19–21} are often used to weaken non-complementary strand interactions in order to increase specificity.

Modern achievements in dynamic DNA nanotechnology have also significantly contributed to the development in DNA genotyping approaches. These advances are predominantly based on toehold-mediated strand displacement reactions²²⁻²⁴ enabling unidirectional DNA re-hybridization between a double stranded DNA duplex (dsDNA) and a displacing single stranded oligonucleotide (displacing sequence (DS)) via a 3-way branch migration mechanism (3-WBM). The vast majority of genotyping research published in recent years has described the application of toehold-mediated strand exchange for SNP discrimination^{25–38}. The publications in this field showed successful discrimination of a single nucleotide substitution where the nucleotide in question was located within a toehold sequence ("external" SNP), thereby preventing docking of the DS to the overhanging part of the dsDNA²⁵⁻³¹. Zhang et al. have investigated a large variety of 'internal' SNPs, including both deletions and insertions within short DNA sequences, and showed that during displacement the thermodynamic gain of many correctly paired bases can often override the thermodynamic penalty of a few mismatches; thus leading to complete "false-positive" displacement^{26,32}. In order to minimize non-specific displacement, Zhang et al. proposed truncating the displacing sequence by several nucleotides from the opposite end of the toehold termini³², which allowed for full thermodynamic control of the displacement outcome. A similar approach with doublestranded toehold exchange and transition through a Holliday junction structure (4-way branch migration, 4-WBM) was shown to be successful in the detection of point mutations in the E. Coli rpoB gene³³.

SNP discrimination using a 3-WBM mechanism can be significantly improved by introducing steric hindrance during the displacement process. For example, reliable discrimination of 'internal' SNPs through a 3-WBM strand displacement has been achieved using a unique DNA origami scaffold by Subramanian *et al.*³⁴ They

observed non-specific displacement (up to 40%) for the interactions resulting in a dT-dG mismatched base pair. Other reactions leading to the formation of a mismatched nucleotide pair in the displaced product showed non-specific displacement at significantly lower level (<20%). The likely reason for this lack of displacement specificity in these experiments is the anchoring of the dsDNA duplexes to a semi-rigid DNA origami scaffold, which restricted flexible rotation (i.e., provided conformationally constrained rotation) of the interacting DNA molecules. Another example of conformationally constrained toehold-mediated SNP sensing was demonstrated by Xiao *et al.*³⁵ The authors used a combination of molecular beacons (MBs) with binary probes, which in turn resulted in the creation of a single sterically hindered triple-stem DNA structure capable of SNP specific discrimination.

Previous work by our group has shown a simple method for producing toehold-PCR products (TPPs) amplified from real-life DNA samples³⁶. The dsPCR products have an overhanging toehold sequence and therefore are perfectly suitable for 3-WBM toehold-mediated strand displacement reactions. Successful discrimination between two closely related 106 and 112 bp long toehold-PCR products, differing by a 6-nt insertion, as well as 7 single nucleotide substitutions, was shown. However, discrimination between 80 bp long samples with only one nucleotide difference was unpredictable and kinetically compromised³⁶.

Here, we present an approach for the reliable discrimination of SNPs using real-time PCR instrumentation within 80 bp real-life dsDNA sequences, in this case TPPs, using a combination of both 3-WBM and 4-WBM mechanisms. This approach (3/4/3-WBM) uses the 4-WBM stage as a topological filter for enhanced mismatch discrimination. The present research also reveals new insights into how SNPs influence 3- and 4-way branch migration efficiency and kinetics. Moreover, performing melting curve analysis of the displacement reaction products allows for visual monitoring of the foregoing displacement.

Results

Figure 1 shows the idealized molecular mechanism of the proposed method. The first two stages, namely the docking of the DS to a TPP followed by 3-WBM exchange, are the standard steps for toehold-mediated strand displacement. Then the mechanism enters its 4-WBM stage. This is simply achieved by initial hybridization of the DS with an auxiliary short oligonucleotide ('protector'). The pro-

tector sequence is fully complementary to the region of the DS carrying an internal discriminatory nucleotide (Fig. 1 •), i.e., a nucleotide responsible for SNP discrimination (which in turn is complementary to the nucleotide in question (Fig. 1 \blacksquare) in the TPP). Finally, the displacement finishes with another 3-WBM step, resulting in the formation of two double-stranded displacement products (the main displacement product representing a fully complementary DNA duplex and the displacement 'side' product).

The choice of protecting sequence length was determined by two main constraints: 1) the minimal length of the sequences was restricted by the integrity/stability of the double-stranded structure formed between the displacing sequence and the protector at 30° C (reaction temperature); and 2) the longer the protector sequence the slower the reaction rate of the 4-WBM step since the elementary step for 4-WBM process is at least one order of magnitude slower than that for the 3-WBM process^{41,42}. The protecting sequences of 19, 15, and 11 nucleotide length were therefore evaluated. In all cases the protecting sequences were designed in such a way that the discriminatory nucleotide (Fig. 1 •) in each DS was localized in the middle of the protector sequence.

Kinetic discrimination. Figure 2 shows the results of the toehold-PCR products (TPP(C), TPP(T), TPP(A) and TPP(G)) discrimination using four displacing sequences (DS(A), DS(T), DS(G) and DS(C)), where the letter in brackets stands for the nucleotide in question for TPPs and the discriminatory nucleotide for DSs, respectively) with and without protectors.

Among the four TPPs, the easiest to discriminate using nonprotected $DS(A)_{np}$, $DS(T)_{np}$, $DS(G)_{np}$ and $DS(C)_{np}$ (subscripted " $_{np}$ " stands for no protector) was TPP(C) (Fig. 2A(I–IV) blue traces). The complementary reaction of TTP(C) with $DS(C)_{np}$ reached the 70% plateau of displacement in ≈ 25 min (Fig. 2AIV, blue trace), having a rate constant of $1.47 \times 10^3 M^{-1} s^{-1}$ (Fig. 3II, Table S2). In all cases for the non-complementary reactions of TPP(C) with $DS(A)_{np}$, $DS(T)_{np}$ and $DS(G)_{np}$ (Fig. 2A(I–III), blue traces) there was a similar degree of conversion of 25–40% over a reaction time of 1 h. These three non-complementary processes showed an average constant rate of $1.62 \times 10^2 M^{-1} s^{-1}$ (varying in a range from 7.88×10^1 to $2.23 \times 10^2 M^{-1} s^{-1}$, Fig. 3II, Table S2). Thus, R_k s of 6.6, 7.9 and 18.6 were calculated for the reactions of the TPP(C) with nonprotected $DS(A)_{np}$, $DS(T)_{np}$ and $DS(G)_{np}$, respectively (Fig. 2A(I–III) blue traces, Fig. 3II, Table S3). It is worth noting that the mismatch



Figure 1 | Simplified representation of the approach consisting of 3-WBM and 4-WBM mechanisms for SNP discrimination. F and Q are fluorophore and fluorescent quencher, respectively, used for real-time strand displacement kinetic monitoring.



Figure 2 | Toehold-mediated strand displacement kinetic curves showing the influence of the protecting sequence length on (A) TPP(C), (B) TPP(T), (C) TPP(A) and (D) TTP(G) discrimination with the displacing sequences DS(A), DS(T) DS(G), and DS(C).

pairs being made during these non-complementary reactions are dC-dT, dC-dA and dC-dC, which in turn are the three weakest mismatches according to SantaLucia *et al.*⁴³ Among these values the largest R_k of 18.6 (TPP(C)-DS(G)_{np}, Fig 3II, Table S3) is related to the formation of the weakest (dC-dC) mismatch, supporting the finding that the dC nucleotide is the most discriminative⁴³.

As can be seen, the discrimination of TPP(T), TPP(A) and TPP(G) using non-protected DSs (Figs. 2B(I-IV), 2C(I-IV) and 2D(I-IV) blue traces) can barely be performed. The reactions of TPP(T), TPP(A) and TPP(G) with their complementary non-protected DSs reached the 80-90% level of conversion in just 25-30 minutes, with rate constants of 2.8 \times 10 3 M^{-1} s $^{-1}$, 2.13 \times 10 3 M^{-1} s $^{-1}$ and 2.67 \times 10³ M⁻¹ s⁻¹, respectively (Fig. 3I, 3III, 3IV, Table S2). On the other hand, for all cases of non-complementary reactions a high degree of conversion up to 80% was also achieved over 1 h of reaction. Among the three non-complementary processes the reaction of TPP(T) with DS(A)_{np} (Fig. 2BI, blue trace) was the fastest, showing a rate constant of 1.14×10^3 M⁻¹ s⁻¹ (Fig. 3I, Table S2). R_k for TPP(T)-DS(A)_{np} was determined to be 2.5 (Fig. 3I, Table S3). The rate constant for the reaction of the TPP(A) with the DS(C)_{np} (Fig. 2CIV blue trace) was fitted to be 6.74×10^2 M⁻¹ s⁻¹, which was only 3.2 (R_k) times slower than the TPP(A)-DS(A)_{np} complementary reaction (Fig. 2CI blue trace, Fig. 3III). An Rk of 2.4 (Fig. 3IV) was calculated for the fastest

non-complementary reaction of TPP(G) and DS(A)_{np} (1.12 \times 10³ M⁻¹ s⁻¹, Fig. 2DI blue trace, Table S2).

In order to increase the kinetic discrimination ratios for non-complementary reactions, short auxiliary oligonucleotide 'protector' sequences complementary to the displacing sequence were employed. This modification introduces an additional internal slow 4-WBM step. Given that both the toehold length and sequence are identical through all the strand displacements carried out, it was believed that the introduction of the protector sequence would significantly reduce the overall rate of the strand displacement. Moreover, such a modification of the traditional 3-WBM mechanism would decrease the rate of the non-complementary reaction much more than that of the complementary. This would be achieved through the formation of two mismatched base pairs in the main and 'side' displacement products, whereupon there would be an increase in the energetic barrier to overcome during the 4-WBM step.

As expected, a monotonic decrease of the rate constant of the reaction between TPPs and complementary 'protected' displacing sequences was observed along with an increase in the protector length (Fig. 3). The decrease of the non-complementary reaction rate constants was even more significant. This therefore allowed for better kinetic discrimination between the complementary and non-complementary reactions. For example, TPP(C) was discriminated at least twice as well as better



Figure 3 | Influence of protector length on rate constants of toehold-mediated strand displacement reactions. Brackets on the plots show the discrimination ratios (R_k) between a complementary reaction rate constant (k_{comp}) and a non-complementary reaction constant ($k_{non-comp}$).

using DS(A)₁₅ (where the subscripted number shows the length of the protector sequence) than with non-protected DS(A)_{np}, with R_k s of 12.8 vs 6.6, respectively (Fig. 2AI red and blue traces, respectively, Fig. 3II, Table S3). Better performance in kinetic discrimination was shown for the TPP(A), where the R_k of 3.2 for the reaction with DS(C)_{np} was improved up to 14.5, 33.0 and 53.9 using DS(C)₁₁, DS(C)₁₅, and DS(C)₁₉, respectively (Fig. 2CIV, green, red and black traces, Fig. 3III, Table S3).

The introduction of the protector sequences greatly affected the rate constant ratios for the reaction of TPP(T) with either DS(G) or DS(C) (Fig. 2BIII and Fig. 2BIV) and increased R_k from 6.8 (DS(G)_{np}) and 5.0 (DS(C)_{np}) up to 23.1 (DS(G)₁₅) and 22.1 (DS(C)₁₁) (Fig. 3I, Table S3). However, the discrimination ratio for the reaction between TPP(T) with the protected displacing sequences $DS(A)_{11}$, $DS(A)_{15}$ and DS(A)19 (Fig. 2BI) was not significantly improved compared with the non-protected $DS(A)_{np}$ (Fig. 2BI blue trace). Thus, the largest R_k of 5.0 was achieved for $DS(A)_{19}$, which was twice as high as for DS(A)_{np} (Fig. 2BI black and blue traces, respectively, Fig. 3I, Table S3). A similar situation was observed for the discrimination of TPP(G) with the protected displacing sequences $DS(A)_{11}$, $DS(A)_{15}$ and DS(A)19 (Fig. 2DI). Two protected displacing sequences were shown to be able to efficiently discriminate TTP(G) with a R_k of at least 26.6 and 32.7 for DS(T)₁₉ and DS(C)₁₁ (Fig. 2DII black trace and Fig. 2DIV red trace, respectively, Table S3). However, discrimination of TPP(G) using $DS(A)_{19}$ (Fig. 2DI black trace) gave the largest R_k of only 3.6 (Fig. 3IV, Table S3), which was 1.5 times higher than that for the non-protected DS(A) sequence (Fig. 2DI, blue trace). It is notable that in both cases (TPP(T) and TPP(G)) the lowest discrimination ratios were observed for the reactions with DS(A) regardless of protector length (Fig. 2DI, 2DI, 3I, 3IV), which form relatively strong dT-dT and dG-dT mismatches⁴³ in the respective main displacement products.

Thermodynamics of DNA base pairing and kinetic discrimination. The correlation between the level of kinetic discrimination and the thermodynamics of DNA base pairing is illustrated in Fig. 4, in which the logarithm of the kinetic discrimination ratio for all of the reactions is plotted as a function of the standard free energy change for the strand displacement process relative to that of the complementary reaction at 30°C, ΔG°_{30} , calculated from the Santa-Lucia nearest-neighbor model of DNA hybridization thermodynamics⁴³. The figure shows that, in general, a larger thermodynamic penalty for base-pairing mismatches in the displacement process leads to a greater discrimination ratio between complementary and noncomplementary reactions. For this reason, the reactions using protector sequences and 4-WBM are generally more discriminatory, as the thermodynamic penalty for forming two base-pair mismatches is typically greater than that for forming a single mismatch; this is as discussed above and as shown in the figure. The correlation between the kinetic discrimination ratio and DNA hybridization thermodynamics is not perfect, as a one-to-one correspondence between the energetic barriers for the 3- and 4-WBM steps and the free energy differences between the stable states that are connected by these branch migration steps does not necessarily exist. Also the stability of the nucleotide mismatch within 4-WBM migration point does likely depend on the structure of the mismatch and, even more crucially, on its interaction with flanking nucleotides⁴⁴. In addition, although the discrimination ratios in figure 4 for the 19-nt protector



Figure 4 | Correlation between the level of kinetic discrimination and the thermodynamics of DNA base pairing.

show a generally increasing trend with ΔG°_{30} , the slope of the trend is significantly greater than that for the other protector lengths. The changes in the 'dynamic' interactions between the four DNA strands during the 4-way branch migration steps are not well understood and complex, involving the concerted interaction of four different strands. It is therefore difficult to attribute the changes in the discrimination ratios to specific interactions between the bases in the four strands. To capture the variation in the discrimination ratio with protector length would require a more sophisticated kinetic model that accounts for branch migration steps other than those involving the mismatch.

Discrimination using melting analysis. In addition to real-time monitoring of the kinetics of the displacement reaction, a second approach based on fluorescent DNA melting analysis of the strand displacement products was used. We have previously reported that if any displacement reaction between the displacing sequence and the TPP occurs, a 'positive' melting peak (pointing up) appears when the change of the fluorescence signal intensity is plotted as a positive derivative with respect to temperature (dF/dT) against the temperature $(T)^{45}$. If displacement does not occur the melting peak points down ('negative' melting peak or trough). We therefore used this approach to assess the reaction products obtained in the current work. Figure 5 shows the results of the melting analysis of the displacement products. Melting analysis of the TPPs and any nonprotected DSs, including non-complementary sequences, resulted in only 'positive' melting peaks (Fig. 5, blue traces). Thus, melting curve-based discrimination using non-protected DSs is unfeasible. Interestingly, only 'positive' melting peaks were observed for the melting analysis of the displacement products obtained in the reactions between TPPs and their complementary protected DSs regardless of the protector length (Figs. 5AIV, 5BII, 5CI, 5DIII black, red and green traces). In contrast, the melting curves associated with noncomplementary protected DSs irrespective of protector length, had a sine-like curve shape, with a peak and a trough clearly visible (e.g., Figs. 5AI, 5BIII, 5CIII, 5DIV black, red and green traces) except for the reaction of TPP(G) with DS(A)₁₁ and DS(A)₁₅ (Fig. 5DI green and red traces), which showed only "positive" melting peaks.

The trough is likely explained by completion of the displacement reaction at the elevated temperatures during the melting analysis. The use of a protector sequence slows down the whole strand displacement process. This is particularly the case for the reaction of a TPP with non-complementary 'protected' DSs, in which two mismatched nucleotide pairs form (one in the main displacement product and the other in the 'side' displacement product). It has been shown previously that a temperature increase facilitates the strand displacement reaction³⁶. Thus we assume that the increase in the temperature during the melting analysis makes the slow "protected" displacement occur regardless of the mismatches formed. This may be due to dehybridization of the protector sequence, which leads to completion of the displacement reaction. Subsequent melting of the formed displacement product results in the positive peak on the melting analysis plot. The reaction between a TPP with its complementary "protected" DS always gives a single positive peak due to the displacement reaching completion long before melting starts. This data is in full accordance with the kinetic data obtained for the reactions, again proving that formation of relatively strong mismatched nucleotide pairs for TPP(G) with $DS(A)_{11}$, $DS(A)_{15}$ and $DS(A)_{19}$ allows nonspecific displacement to occur. In the case of the 19 nt protector sequence (Fig. 5DI black trace) the preliminary displacement reaction was significantly slowed down, which in turn allowed the displacement to occur only at elevated temperature.

Calculation of the normalized area under the curve (AUC) allows easy numerical assessment of the shape of the melting curve. Thus, the AUC for strictly positive melting curves always tends to give a positive value approaching 1 (for normalized data). On the other hand, the AUC for melting curves with a peak and a trough approaches zero. Figure 6 shows an example of the discrimination of the TPPs using the melting curve shape analysis approach. As stated previously the discrimination of TPP(T), TPP(A) and TPP(G) with non-protected displacing sequences was not possible since all the areas under the melting curves nearly equal 1 (Figs. 6I, 6III and 6IV, respectively). Introducing protector sequences allowed for a significant decrease of the AUC value with increasing protector length, down to 0.1 for the reaction of TPP(T) with DS(A) (Fig. 6I, black bars) and even slightly negative values for non-complementary displacement reactions with DS(C) and DS(G) (Fig. 6I, blue and green bars, respectively). The normalized AUC values for the complementary reactions of TPP(T) with $DS(T)_{np}$, $DS(T)_{11}$, $DS(T)_{15}$, and DS(T)₁₉ varied from 1 to 0.78 (Fig. 6A, red bars). Similar trends were observed for the rest of the toehold-PCR products (Fig. 6II, 6III and 6IV), so setting up a cut off value of 0.3 allows for reliable discrimination using all displacing sequences with 19 nt long protectors. Non-parametric Spearman correlation analysis showed a statistically significant relatedness between the reaction rate constants and the normalized AUC ($\rho = 0.71, p < 1 \times 10^{-3}$).

Discussion

We have demonstrated a new method for SNP identification and discrimination based on a toehold- mediated strand displacement reaction proceeding through both 3- and 4-way branch migration mechanisms. Additional short oligonucleotide sequences of various lengths were hybridized to single-stranded displacing sequences, which introduced a 4-way branch migration step into the overall reaction mechanism.

Our results suggest for the first time that, for most base pairs, there is a reasonable correlation between sequence-dependent 3- and 4-WBM migration barriers and DNA hybridization free energies. If the rate-determining step for the displacement process is branch migration through the base mismatch (as values of the kinetic discrimination ratio generally a factor of 5 or more would suggest) and if there is much greater variation between the rates of complementary versus non-complementary reactions than between complementary reactions for a given protector length (as the experimental data indicates), the discrimination ratio would be expected to scale approximately with $\exp(\Delta G^*/k_BT)$, where ΔG^* is the activation free energy difference between the non-complementary and complementary reactions for the SNP branch migration step. A linear correlation between the activation free energy difference, ΔG^* , and the difference between the free energy change of strand displacement, ΔG°_{30} , would give a



Figure 5 | Melting temperature analysis of toehold-mediated strand displacement products produced by the reaction of (A) TPP(T), (B) TPP(C), (C) TPP (A) and (D) TPP(G) with both non-protected (blue traces) and protected (black, red and green traces) DS(A), DS(T), DS(G), and DS(C).

linear relationship between the logarithm of the kinetic discrimination ratio and ΔG°_{30} , as Fig. 4 clearly shows (except for a few outlying points). The slope of the linear fit to the data in Fig. 4 is 0.25, indicating that the activation free energy differences due to base mismatches for 3- and 4-WBM are approximately 1/4 the thermodynamic free energy differences. This result is expected, as interactions between bases are partially formed or broken in the transition state for the branch migration step and therefore likely to be less sensitive to base mismatches than the free energies of the stable states. The outliers below the linear regression line in Fig. 4 correspond to the TPP(T) and TPP(G) reactions with protected DS(A), while the outliers above the line correspond to TPP(T) and TPP(C) reactions with protected DS(C). The lack of correlation suggests that simple thermodynamic considerations are not completely sufficient for describing the energetics of these processes and that further investigation will be necessary to elucidate the mechanisms of 3- and 4-WBM transition through sequence heterology.

In summary, the method presented here allows for reliable kinetic SNP discrimination. Among the four toehold-PCR products of 80 bp, which carried four different SNPs at the same position, the easiest to discriminate was the one with the SNP dC, which forms the weakest and the most destabilizing 4-WBM junction structure with mismatched pairs^{43,44}. Conversely, the poorest discriminative displacing sequence was the one carrying a dT discriminatory nucleotide. This is likely due to the smaller size of the dT nucleotide, which results in less steric hindrance during the branch migration transition through the SNP site and moderate stability of the newly formed mismatches. Also, the use of the protector sequence represents an additional way to controllably decrease the strand displacement rate without altering other reaction parameters such as the strength of toehold binding^{26,46} or the reaction temperature^{36,42,46}. It is worth to note that the control of the displacement rate with no changing the reaction conditions was also previously shown using the remote toehold approach of Genot *et al.*³⁹

In addition, performing melting analysis of the displacement products allows simple visual discrimination of all tested SNPs based only on the assessment of the shape and area of the obtained melting curves.

Methods

Nucleic acid isolation and PCR amplification. Human genomic and mitochondrial DNA were co-extracted from the authors' own blood samples using a QIAamp DNA blood mini kit (Qiagen, Germany), according to the recommendations of the manufacturer. PCR primers, including dU-modified and fluorescently labeled primers (FAM) (see, Table s1), were purchased from IDT DNA Technology, USA and used at a concentration of $0.2 \ \mu$ M.

In order to evaluate the proposed method for SNP discrimination, in addition to the SNPs found in the authors' own DNA samples (C16223 (SNP-C) and T16223 (SNP-T) within the Hyper Variable Region 1 (HVR-1) of the mtDNA), two recom-



Figure 6 | Normalized area under the curve for the curves obtained during the melting temperature analysis of the toehold-mediated strand displacement reaction of (I) TPP(A), (II) TPP(T), (III) TPP(G), (IV) TTP(C) with both protected and non-protected DSs.

binant plasmid DNA carrying SNP-A and SNP-G were engineered (for more details see SI).

PCRs were performed using a HotStar Taq DNA polymerase (Qiagen, Germany) within 1× HotStar Taq PCR buffer with a final MgCl₂ concentration of 2.5 mM. The concentration of each of the standard deoxynucleotide triphosphates (dATP, dTTP, dGTP, dCTP) was 0.2 mM. A PCR amplification regime of 95°C for 15 min, 30 cycles of 94°C for 15 s, 61°C for 15 s, 72°C for 25 s, and a final elongation of 72°C for 1 min was used. A template of 5 ng of the extracted human genomic DNA or 1 ng of the plasmid DNA carrying PCR product with either SNP-A or SNP-G was used. For the detailed protocol of the plasmid DNA preparation see SI.

Preparation of the toehold-PCR products (TPPs). Directly after the PCR, Uracil-DNA glycosylase (NEB, USA) (2.5 U) was added to the PCR solution and the entire mixture was then gently mixed via pipetting. After incubation at room temperature for 5 min the mixture was heated to 95°C for 5 min and then cooled back to room temperature; this resulted in the formation of a 9 nucleotide long toehold sequence. The PCR-UDG mixtures were purified using an Amicon Ultra-0.5 30K (Millipore, USA) according to the recommendations of the manufacturer. The nucleic acids were then eluted with a displacement 1× TEM buffer (pH 8), consisting of Tris•HCl (10 mM), EDTA (1 mM) and MgCl₂ (12.5 mM). Quantification of the eluted PCR products was carried out using a NanoDrop 1000 spectrophotometer (Thermo Scientific, USA).

Strand displacement reaction. Displacing sequences DS(A), DS(T), DS(C) and DS(G) (Table S1), 80 nt in length and labelled at the 3' termini with the fluorescent quencher carboxytetramethylrhodamine (TAMRA), were purchased from Eurogentec, Belgium.

Displacing sequences were mixed with corresponding protector sequences in stoichiometric ratios and then annealed from 95 to 20° C at a constant rate of 0.5° C/min.

All displacement reactions were carried out with 5 times excess of the DS over the TPP at 30°C in a reaction volume (15 μ L) using a RotorGene Q 6-plex real-time PCR thermocycler (Qiagen, Germany). Prior to the displacement reaction the toehold-PCR product (2 pmol, dissolved in 1 \times TEM buffer) was placed in a thin-wall PCR tube (0.2 mL) and briefly centrifuged. Then, the displacing sequence (10 pMole, dissolved in 2 μ L of 1 \times TEM buffer) was carefully applied onto the lid of the same PCR tube. The tube lid was then carefully closed and the tube placed into the thermocycler avoiding mixing of the TPP and the DS before the run protocol started.

Acquisition of fluorescent signals was performed within the SYBR Green/FAM channel at a gain of 7 within a time interval of 20 sec between fluorescent reads. Normalization of the raw fluorescence was made by dividing the signal values by the initial signal value, as described previously³⁹.

Melting curve analysis was performed according to the RotorGene Q 6-plex builtin protocol, which included a temperature ramp from 30 to 95° C with an increase of 1° C each step of 5 s waiting time.

Curve-fitting of the kinetic data was carried out using the non-linear curve fitting function in OriginPro 9 software (Origin Corporation, USA). Assuming that an excess of the displacing sequence shifts the displacement kinetics towards the first order behavior the obtained data were fit to the pseudo-first-order kinetic equation as described by Baker *et al.*⁴⁰,

$$F = F_1 + (F_0 - F_1)e^{-kt[DS]_o}, \qquad (1)$$

where *F* is the fluorescence intensity at time t, F_0 is the initial fluorescence intensity, F1 is the intensity after reaction completion, *k* is the rate constant, $[DS]_o$ is the initial concentration of the displacing sequence, and the *t* is the time in seconds.

The ratio between a complementary reaction rate constant ($k_{\rm comp}$) and a noncomplementary reaction constant ($k_{\rm non-comp}$) with no protector sequence, or with a protector of the same length as that of the complementary reaction, was used as a kinetic discrimination ratio,

$$R_{\rm k} = \frac{k_{\rm comp}}{k_{\rm non-comp}}.$$
 (2)

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Author contributions

D.A.K. and A.S.K. conceived, designed and performed the experiments; D.A.K. and D.M.H. analyzed the data; D.A.K., A.S.K. and D.M.H. wrote the paper draft; A.L. and A.V.E. edited the paper draft; A.V.E. guided the research.

Additional information

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