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Non-Equilibrium Hybridization Enables Discrimination of a Point Mutation within 5-40°C

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Supporting Information Placeholder

ABSTRACT: Detection of point mutations and single nucleotide polymorphisms in DNA and RNA has a growing importance in biology, biotechnology, and medicine. For the application at hand, hybridization assays are often used. Traditionally they differentiate point mutations only at elevated temperatures (>40°C) and in narrow intervals (∆T = 1-10°C). The current study demonstrates that a specially designed multi-stranded DNA probe can differentiate point mutations in the range of 5-40°C. This unprecedentedly broad ambient-temperature range is enabled by a controlled combination of (i) non-equilibrium hybridization conditions and (ii), a mismatch-induced increase of equilibration time in respect to that of a fully matched complex which we dub ‘kinetic inversion’.

Analysis of single nucleotide substitutions (SNS) has a growing importance in the diagnostics of genetic and infectious diseases, genome-wide association studies, forensics, and other applications. Hybridization probes have been extensively used in SNS analysis. The probes consist of nucleic acid oligomers of 15-25 nucleotides (or longer) that hybridize to DNA/RNA analytes containing an SNS site of interest (Fig. 1A). The duplex is subsequently destabilized by heat to differentiate fully matched hybrids from the mismatched ones. However, these probes enable SNS differentiation only within a relatively narrow temperature range (∆T), which is above the ambient range. Therefore, expensive instrumentation, such as e.g. qPCR thermocyclers with high-resolution melting capabilities, is required for heating and temperature control. Other approaches for SNS differentiation at ambient temperatures employ DNA-binding proteins, taking advantage of the differences in the 3D recognition of matched or mismatched base pairs. While these techniques are well-recognized and extensively used, they require protein production and storage as well as specific assay conditions to maintain protein activity, and thus potentially more resource intensive than hybridization-based assays. On the other hand, the development of hybridization probes with SNS selectivity at a broad ∆T downshifted to ambient temperatures would eliminate the need for specialized equipment and aid RNA analysis in living cells and molecular diagnostics in instrument-free settings. An active search for such hybridization probes is ongoing in the field.

Figure 1. Three types of hybridization probes: designs and fluorescent melting profiles in the presence of the matched (solid line) or mismatched (dotted line) analytes. A) Linear probe: an unfolded DNA probe hybridizes to a nucleic acid analyte. In the study, the analytes were labelled with a quencher dye (Q), whereas L1 probe was labelled with a fluorophore (F). B) Molecular beacon (MB) probe. The dash-dotted line (right panel) indicates the melting profile of the MB1 alone. C) X probe: strands m and f bind both the analyte and the universal MB (UMB) probe to form a four-stranded fluorescent complex (AS). Right panel: The melting profiles for the X1_m7 (non-equilibrium conditions at low temperatures, see main text). The dash-dotted line: melting of UMB1 alone; dashed line: melting of the X1_m7, no analyte. For experimental details see Fig. S1.
One approach uses ‘molecular beacon’ (MB) probe, a fluorophore- and a quencher-conjugated DNA stem-loop structure (Fig. 1B). When bound to a complementary DNA, the MB probe switches from the folded conformation with quenched fluorescence to extended, highly fluorescent conformation. Compared to linear probes, MB probes exhibit a broader ΔT and lower melting temperatures (Tm) of the MB-analyte complexes. The changes in Tm and ΔT result from the equilibrium shift towards the dissociated state (DS), stabilized by the base pairing of the stem portion of the unbound MB probe. We further advanced the idea of broadening ΔT by conformational constraints through using a multicomponent X probe (Fig. 1c). In this study, we found that the X probe differentiates between the matched and mismatched analytes in the range of 5-40°C, contrary to the predictions of equilibrium thermodynamics. This unprecedentedly broad differentiation range of the X probes results from the non-equilibrium operation mode and the ‘kinetic inversion’ effect observed in this study for the 1° time.

The X probe consists of a universal MB (UMB) probe and two adapter DNA strands m and f. The three strands form a tetrameric complex with the analyte (Fig. 1C). The sequences of the matched (T_G) and mismatched (T_A) analytes, as well as of the linear (L1) and the MB (MB1) probes were identical to those reported by Tsourkas et al. Our goal was to directly compare the performance of the X probe with that of the linear and MB probes studied earlier. A series of four X probes was designed to be fully complementary to T_G. All the X probes used the same universal MB probe (UMB1) and strand f (X1-f in Table S1), but differed in the length of the analyte-binding arm of strand m, which varied from 6 to 9 nucleotides (Table S1).

For L1, the difference in the probe-analyte Tm (ΔTm), the commonly used characteristics of SNS differentiation efficiency, was 9.6°C, which was identical to the value reported by Tsourkas et al. (cf. Table S2 and Fig. 1A). As expected, the ΔTm for the MB1 was broader (11.2°C) and shifted by 2-3°C towards lower values (Table S2), in agreement with the previous data. For the X probes, however, the downshift of Tm by 30°C and the broadening of the ΔTm by 5-7°C (in comparison with L1 and MB1 probes) were observed (Table S2). This performance exceeds that of other conformationally constrained probes. We explained this effect by the greater stabilization of the DS for the X probe as compared to the L1 or the MB1 (Table S3, Fig. S3, S4). The DS of the X probe is more stable than that of the MB probe due to (i) the residual hybrid between strand f and the analyte; and (ii) higher entropy of the DS resulting from the complex dissociating into three rather than two fragments, as would be the case for the MB probe.

To better visualize the SNS differentiation of the three probes, we plotted the Fm/Fm,n ratio as a function of temperature, where Fm and Fm,n are the fluorescence intensities of the probes in the presence of matched and mismatched analytes, respectively (Fig. 2A). The Fm/Fm,n for the X probe was greater than 1 down to 5°C (Fig. 2A). Similar results were observed for X probes with different lengths of m strands (cf. X1_m6, 8, 9 in Fig. S2) as well as at lower analyte concentrations (Fig. S5). We describe this practically important property of the X probe by a new parameter, ΔT1.5, the temperature interval in which the fully matched analyte produces the signal 1.5 times greater than the mismatched analyte (Table S2). The ΔT1.5 differentiation intervals were 14.8°C and 17.1°C for the L1 and the MB1 probes, respectively, and ca. 35°C for the X1_m7 probe (Fig. 2A and Table S2). The discrimination curve for the X1_m7 probe is asymmetric, with a SNS differentiation even at the temperatures below 15°C (Fig. 2A), which contradicts the theoretical curve predicted under the assumption of thermodynamic equilibrium (Fig. 2B). These observation strongly suggest that the X-probe operated under non-equilibrium conditions at low temperatures.

![Figure 2](image2.png)  Figure 2. Discrimination profiles for the three hybridization probes. A) The ratio of fluorescence produced by each probe in the presence of fully matched analyte (Fm) to that of mismatched analyte (Fm,n) plotted against temperature for L1 (dotted line), MB1 (dashed line) and X1_m7 (solid line). The threshold of Fm/Fm,n ~ 1.5 is indicated by the red dotted line. B) The nearest-neighbour (NN) model prediction of the profiles in the assumption of thermodynamic equilibrium (see SI for details).

![Figure 3](image3.png)  Figure 3. Equilibration of the probe-analyte complexes at different rates of the cooling-heating cycle. A) Temperature dependence of the fluorescent signal for the MB1 probe at the equilibration times of 19 s or 600 s allowed per each 1°C heating/cooling step. The solutions were first heated to 95°C, then cooled to 5°C, and then heated to 95°C again. B) Same as panel A for the X1_m7 probe. C) The fluorescent signal for the equilibration time of 19, 60, 600 or 1800 s/1°C observed for the MB1 probe-analyte complex at 10°C during the cooling cycle. D) Same as panel C for the X1_m7 probe. The orange line corresponds to X1_m7, no analytes. Solid and dashed lines are used for the complexes of the probes with T_G (matched) or T_A (mismatched) analytes, respectively.

To prove the latter point, we varied the heating and cooling rates in the hybridization experiments (Fig. 3). The MB1 probe reached equilibrium relatively fast, which is evident from the overlap of the fluorescent melting profiles obtained for the different equilibration times allowed for each 1°C step of the cooling/heating cycle (Fig. 3A). The fluorescent signal profiles are nearly symmetric, reflecting that both cooling and heating bring the hybridization ensemble to the same equilibrium state. Furthermore, the equilibrium was reached at the cooling rate of 60 °C/s of the matched and 19 °C/s for the mismatched analyte (Fig. 3C). Such hybridization kinetics is in agreement with the prior observations that mismatched duplexes equilibrate faster than the matched ones. In contrast, the X1_m7 probe demonstrated a strong hysteresis between the cooling and heating profiles (Fig.
3B), indicating the absence of equilibrium. At 10°C, the fluorescence of the fully matched X1_m7-analyte complex was somewhat stabilized at the cooling rate of 60 s°C (Fig. 3D). For the mismatched analyte, however, the fluorescent signal kept monotonously increasing (Fig. 3D) even when the equilibration time reached 16 hrs (Fig. S6) indicating that the probe-analyte hybridization remained incomplete. We therefore concluded that the complex of the X1_m7 probe with the matched analyte equilibrated faster than with the mismatched analyte. To the best of our knowledge such a phenomenon, which we dub here kinetic inversion, has not been previously reported. Kinetic inversion explains the broadening of the ΔT for the X probe towards lower temperatures: the lower the temperature, the more time is required for the equilibrium, which effects the mismatched complex greater than the matched one. To prove that the observed effect is not unique for a particular analyte sequence, we designed an X2 probe for the analysis of SNS in an arbitrary chosen DNA sequence of human RASSF1A gene (T2_C and T2_T in Table S1). The obtained melting profiles and ΔT differentiation range were similar to those observed for the X1 probe (Fig. S7).

Figure 4. Hypothetical reaction diagrams for hybridization of A) the MB probe and B) the X probe at low temperatures. MB# and UMB# are the high energy random coil conformations with destabilized base-pairing of the stem.

We explain the origin of the kinetic inversion phenomenon using the hybridization diagrams shown in Fig. 4. For the MB probe-analyte complex, the AS formation can proceed via the transition state MB# wherein both the MB and the analyte strands presumably adopt random coil conformations. Thus, the equilibration rate of AS ≈ DS is limited by the unfolding of the MB strand and the latter is faster for the mismatched complex due to the lower activation energy. Thus, the MB probe equilibrates faster with mismatched than with matched analytes in agreement with others and our own (Fig. 3) observations. In contrast, the hybridization for the X probe proceeds presumably via one or several intermediate states. In Fig. 4B, one possible intermediate (Ix) is shown as a complex of the m and f strands with the analyte. We assume that the equilibration rate for both matched and mismatched hybridization is limited by the rate of equilibration of AS ≈ Ix. The rate of AS → Ix should be the same for both matched and mismatched complex, since it depends on the stability of UMB stem loop, which is the same in both cases. On the other hand, the Ix → AS rate depends also on the steady-state concentration of the Ix intermediate: Rate_{IX → AS} = k_{IX → AS} × [Ix] × [UMB]. The [Ix] should be greater for the matched than for the mismatched probe-analyte complex considering that the latter is affected by the mismatch penalty. Thus, the rate of Ix → AS production, and the overall equilibration rate should be greater for the matched than for the mismatched complex. For other conceivable intermediate states, the argument remains similar. This explanation corresponds to the ‘kinetic proofreading’ model suggested by Hopfield to explain the high accuracy of strongly but non-specifically driven biochemical reactions (e.g. replication, protein synthesis). The model builds on intermediate states that are populated by increased fractions of the matching molecule with respect to equilibrium.

In conclusion, we report a novel phenomenon: the complex of the multicomponent X probe with mismatched analytes equilibrates slower than with matched analytes. To the best of our knowledge, this phenomenon was not observed for other hybridization probes. The kinetic inversion enables differentiation of a point mutation in DNA in an unprecedentedly broad temperature range of 5-40°C. This phenomenon can be exploited for the design of hybridization probes with high mismatch selectivity at low temperatures, such as the X probe studied here. Such designs open the possibility of detecting point mutations in RNA in living cells and developing room-temperature diagnostic assays that operate without precise temperature control.

Supporting Information
Details of experimental procedures, DNA sequences, computer modeling and probe design, fluorescent response of different X probes and phase diagrams fitting the thermodynamic parameter are included in Supporting Information.

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Notes
The authors declare no competing financial interests.

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