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Sensitive, Simultaneous Quantitation of Two Unlabeled DNA Targets Using a Magnetic Nanoparticle-Enzyme Sandwich Assay

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Abstract

We report herein the development of a simple, sensitive colorimetric magnetic nanoparticle (MNP)-
enzyme based DNA sandwich assay that is suitable for simultaneous label-free quantitation of two DNA
targets down to 50 fM level. It can also effectively discriminate single-nucleotide polymorphisms (SNPs)
in genes associated with human cancers (KRAS codon 12/13 SNPs). This assay uses a pair of specific
DNA probes, one being covalently conjugated to a MNP for target capture while the other being linked
to an enzyme for signal amplification, to sandwich a DNA target, allowing for convenient magnetic
separation and subsequently efficient enzymatic signal amplification for high sensitivity. Careful
optimization of the MNP surfaces and assay conditions greatly reduced the background, allowing for
sensitive, specific detection of as little as 5 attomole (50 fM in 100 μL) of target-DNA. Moreover, this
sensor is robust, it can effectively discriminate cancer specific SNPs against the wild-type non-cancer
target, and works efficiently in 10% human serum. Furthermore, this sensor can simultaneously
quantitate two different DNA targets by using two pairs of unique capture- and signal- DNA probes
specific for each target. This general, simple and sensitive DNA sensor appears to be well-suited for a
wide range of genetics based biosensing and diagnostic applications.

Keywords: biosensing, magnetic nanoparticle, enzymatic assay, simultaneous DNA quantitation,
single-nucleotide polymorphism
**Introduction**

The development of biosensors capable of rapid, sensitive detection of specific genetic biomarkers is critical to healthcare, allowing diagnosis of diseases, prediction of patients’ responses to treatment and risk of relapse of disease.\(^1\)\(^2\) The polymerase chain reaction (PCR) is the most widely used technique for DNA detection due to its great, exponential amplification capability.\(^3\) However, as a paradox of its great amplification power, even a tiny amount of contaminant can result in non-negligible false positive which can affect diagnostic accuracy. Moreover, PCR requires a relatively clean lab environment and long assay period, making it less well-suited for rapid, on-site diagnosis. Therefore, alternative, PCR-free based DNA sensing approaches have been actively exploited over the past decade. Among which, several gold nanoparticle (GNP) based methods, e.g. silver-amplified scannometric assay\(^4\), silver-amplified electric detection\(^5\), silver-amplified Raman finger printing\(^6\), and magnetic microparticle-assisted DNA nano-barcode assay\(^7\), have exhibited exceptionally promises. They displayed excellent sensitivities and specificities for DNA detection, down to the fM to aM level, making them potentially suitable for direct target detection without the need of PCR pre-amplification. Despite these, most assays use surface-immobilized DNA probes for soluble target capture, a heterogeneous process often suffer from slow binding kinetics and low capture efficiency. As a result, a relatively long period for target capture and GNP sandwich binding (ca. 6-8 h) is required,\(^4\)\(^-\)\(^6\) making them less well-suited for rapid detection.

Meanwhile, functional nanoparticles, such as quantum dots, metal nanoparticles and magnetic nanoparticles (MNPs), have unique, size-dependent optical and electrical properties that are well-suited for biosensing. In this regard, MNPs are extremely well-suited because of several attractive properties, e.g. ease of synthesis, versatile surface modification strategies, good stability, low toxicity, and moreover
their super-paramagnetic properties. As a result, they form stable, uniform dispersions in the media for homogenous, rapid and efficient target capture without an external magnetic field; but are readily collected and separated from the media upon applying an external magnetic field. Moreover, a large excess of MNP-capture probes can be used to push the equilibrium of the target-probe binding toward the captured state, allowing efficient target detection at concentrations far below the equilibrium dissociation constant ($K_d$). These make them extremely well-suited for biosensing, bioseparation and biocatalysis. Meanwhile, enzymes are extremely versatile, efficient biocatalysts with great substrate turnover (signal amplification) power, making them attractive for ultrasensitive biosensing. In fact, some of most widely used commercial diagnostic assays, e.g. the enzyme linked immunosorbent assay (ELISA), are based on enzymes such as horseradish peroxidase (HRP) and alkaline phosphatase (ALP). Enzymes have been combined with electrochemical, electrochemiluminescence, colorimetric, and fluorimetric readout strategies in sensing. Among which, electrochemical readout is widely used in point-of-care (PoC) diagnostics (e.g. the famous personal glucose meter). Despite high convenience, such PoC diagnostics are mostly suitable for high abundant targets due to limited sensitivity (e.g. $\mu$M-mM). For early diagnosis, a much higher sensitivity is needed. Furthermore, most DNA sensors have been demonstrated with a single target, which can limit diagnostic accuracy because “no tumor marker identified to date is sufficiently sensitive or specific to be used on its own to screen for cancer”. Therefore, the ability of detecting multiple analytes simultaneously is important to high diagnostic accuracy. In this regard, several multiplexed biodetection methods have been reported, including the giant magnetoresistive (GMR) sensor, multi-color GNP SERS finger-printing, GNP multicolor nanobeacons; multi-color molecular beacons (MBs) and graphene quenched multi-color sensors. Nevertheless, the sensitivity
of most approaches (e.g. high pM-low nM for the later three) needs to be improved considerably to make them competitive against existing clinical diagnostic assays. By combining the advantageous properties of both MNP and enzymes, here we have developed a simple, sensitive MNP-enzyme sandwich assay suitable for label-free quantitation of two DNA targets. We show this sensor is sensitive (50 fM); robust (works in 10% human serum), specific (can efficiently discriminate cancer specific single-base mutants from the wild-type non-cancer target).

Experimental Section

Materials: HPLC purified DNA probes and target strands were purchased commercially from the IBA GmbH (Germany). Their sequences are given in Table 1. HRP-neutravidin (HRP-NAV) and ALP-neutravidin (ALP-NAV) conjugates were purchased from Thermo Scientific (UK). Amplex red and fluorescein diphosphate (FDP) were purchased from Invitrogen Life Technologies (UK). The hetero-functional cross linker SM(PEG)$_{12}$ was purchased from Fisher Scientific Ltd (UK). All other chemicals and reagents were purchased from Sigma-Aldrich (UK). PBS (137 mM NaCl, 10 mM Na$_2$HPO$_4$, 2.7 mM KCl and 1.8 mM KH$_2$PO$_4$, pH 7.4) and Tris buffer (100 mM Tris.HCl, 100 mM NaCl, pH 8.5) were made with ultra-pure MilliQ water (resistance > 18 MΩ.cm$^{-1}$). The MNPs were in house synthesized and modified (see Supporting Information, SI, for details).

Instruments: UV-vis spectra were recorded on a Cary 50 Bio UV-Vis spectrophotometer with 1.0 cm optical path length.$^{15}$ The spectra were corrected by their respective buffer background. The HRP based DNA detection limit assay was monitored by fluorescence time-trace on an Envision plate reader using BODIPY TMR FP 531 as excitation filter and Cy3 595 as emission filter.$^{18}$ Powder X-ray
diffractogram was performed on PANalytical's X'Pert PRO Materials Research Diffractometers (MRD) using a scan range (2θ) of 10-90 ° at increments of 0.0332 °. The magnetisation measurement was carried out on a MagLab vibration sample magnetometer (Oxford Instruments) at 55 Hz with amplitude of 1.5 mm. The samples were measured at room temperature (RT) at a scan rate of 9.17 mT.s⁻¹.¹⁵a

**Table 1.** The DNA sequences and their abbreviations. T-DNA4 and T-DNA5 are two cancer-specific single-base mutants of the KRAS codon 12/13, while T-DNA3 is the wild-type target.

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence (5'→3')</th>
</tr>
</thead>
<tbody>
<tr>
<td>cDNA1</td>
<td>HS-TTT TTT TCC GAC CTG GGG</td>
</tr>
<tr>
<td>sDNA1</td>
<td>GAG TAT TGC GGA GGA-TTT TTT-Biotin</td>
</tr>
<tr>
<td>T-DNA1</td>
<td>TCC TCC GCA ATA CTC CCC CAG GTC GGA</td>
</tr>
<tr>
<td>cDNA2</td>
<td>HS-TTT TTT GGC AGT CCG TGG TAG</td>
</tr>
<tr>
<td>sDNA2</td>
<td>GGC AGG TTG GGG TGA TTT TTT-Biotin</td>
</tr>
<tr>
<td>T-DNA2</td>
<td>TCA CCC CAA CCT GCC CTA CCA CGG ACT GCC</td>
</tr>
<tr>
<td>cDNA3</td>
<td>TGG CGT AGG CAA GAG TTT TTT TT-HS</td>
</tr>
<tr>
<td>sDNA3</td>
<td>Biotin-TTT TTT GTG GTA GTT GGA GCT GG</td>
</tr>
<tr>
<td>T-DNA3</td>
<td>ACT CTT GCC TAC GCC ACC AGC TCC AAC TAC CAC</td>
</tr>
<tr>
<td>T-DNA4</td>
<td>ACT CTT GCC TAC GCC ATC AGC TCC AAC TAC CAC</td>
</tr>
<tr>
<td>T-DNA5</td>
<td>ACT CTT GCC TAC GCC AAC AGC TCC AAC TAC CAC</td>
</tr>
</tbody>
</table>

**General assay procedure:** The capture-DNAs (cDNAs) were covalently conjugated to amine-modified MNPs via a hetero-functional cross-linker SM(PEG)₁₂ (SI). The PEGylated cross-linker is used because it can effectively resist non-specific adsorption, greatly reducing background. The cDNA loading on the MNP was estimated by our previous method.¹⁸ The signal-DNA (sDNA) was linked to neutravidin- (NAV-) HRP or ALP conjugate via the strong biotin-NAV interaction at 1:1 molar ratio.¹⁵ For a typical UV-vis assay: MNP-cDNA (20 µg), HRP-sDNA (5 pmol) and various amounts of T-DNA were mixed to make a series of samples (final volume: 500 µL in PBS containing 1 mg/mL BSA). After incubation for 1 h at RT, the MNP-dsDNA-En sandwiches were separated magnetically and the clear supernatants were discarded. The MNPs were washed with PBS once, PBS with 0.1% Tween-20 twice,
and finally PBS once again to remove any unbound species. The MNP sandwiches were then dispersed in PBS (380 µL), and enzymatic amplification was initiated by adding Amplex red/H$_2$O$_2$ (50 µL each, both 0.2 mM). After a certain period, 20 µL N$_3$Na (1 M) was added to stop further amplification and the UV-vis spectra of the supernatants were recorded.

For ALP-based assay, the procedures were the same as above, except where sDNA2-ALP and FDP were used. All assays and washing steps were performed in Tris buffer. The termination of enzyme amplification was achieved by adding 20 µL PBS to the assay solution. For simultaneous quantitation of two DNA targets, the assay was carried out in Tris buffer using a mixture of MNP-cDNA1 and MNP-cDNA2 (15 µg each). After sandwich binding and washing steps, FDP and Amplex red were added simultaneously to initiate enzymatic amplifications. All other assay details are given in the SI.

**Results and Discussion**

The principle of the MNP-enzyme sandwich assay for simultaneous detection of two T-DNAs is shown schematically in Scheme 1. Two sets of unique MNP-capture and En-signal probes are used to target two specific T-DNAs (e.g. MNP-cDNA1/HRP-sDNA1 for T-DNA1; MNP-cDNA2/ALP-sDNA2 for T-DNA2). In the presence of T-DNA1, HRP-sDNA1 will bind to MNP-cDNA1 via specific sandwich hybridization, while T-DNA2 will link ALP-sDNA2 to MNP-cDNA2. All these are performed in homogenous solutions, allowing for efficient, rapid T-DNA capture and conversion of each captured T-DNA1 or T-DNA2 into a HRP or ALP for signal amplification. A large excess of MNP-cDNA and sDNA-En probes (10-200,000 fold molar equivalent of T-DNA, depending on T-DNA concentration) are used to push the equilibrium of the MNP-cDNA/T-DNA/sDNA-En
sandwich hybridization toward the hybridized state. A high cDNA loading on the MNP (each conjugated to several hundred copies of cDNAs, only one is shown here for simplicity) is used to enhance T-DNA capture efficiency and binding-affinity.\textsuperscript{14} Moreover, we have previously found that MNP-immobilized enzymes retained much higher activities over those on flat surfaces (~5-fold) from improved substrate accessibility.\textsuperscript{15} All these lead to significantly improved sensitivity. After magnetic separation, followed by washing steps, Amplex red and fluorescein diphosphate (FDP), which can be efficiently turned over by HRP and ALP into resorufin ($\lambda_{\text{max}} = 571$ nm) and fluorescein ($\lambda_{\text{max}} = 485$ nm), respectively, are added simultaneously. This allows the HRP/ALP catalyzed reactions to be used for T-DNA1/T-DNA2 detection, respectively.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{scheme.png}
\caption{Scheme 1. Principle of the MNP-enzyme sandwich assay for simultaneous detection of two DNA targets using two sets of unique MNP-cDNA and sDNA-En probes. The cDNA and sDNA probes are complementary to each half of their DNA targets for specific sandwich hybridization, linking HRP (with T-DNA1) or ALP (with T-DNA2) to the MNP. Therefore HRP/ALP amplified enzyme products: resorufin ($\lambda_{\text{max}} = 571$ nm) and fluorescein ($\lambda_{\text{max}} = 485$ nm) are used to quantitate T-DNA1/T-DNA2, respectively.}
\end{figure}

**Enzyme activity and termination**

This assay uses specific colored products generated by HRP/ALP catalyzed reactions for T-DNA1/
T-DNA2 detection. It is useful to be able to terminate the enzymatic reaction after a certain amplifying period before measurement. Here, NaN₃ and PBS are found to effectively stop the HRP and ALP catalyzed reactions, respectively (SI, Fig. S1). Moreover, the substrate turnover rate (TR) for HRP-NAV and ALP-NAV can be estimated from the assay slopes and ε of resorufin (54,000 M⁻¹.cm⁻¹) and fluorescein (50,400 M⁻¹.cm⁻¹) via, TR = slope/([enzyme] * ε). This yields 65 s⁻¹ in PBS and 56 s⁻¹ in Tris buffer for HRP-NAV; and 65 s⁻¹ for ALP-NAV in Tris, consistent with the literature.¹⁵

**Characterization of MNP and MNP-NH₂.**

The MNP is found to be mainly made of magnetite from the X-ray diffraction (Fe₃O₄, SI, Fig. S2A). A strong IR absorption at ~569 cm⁻¹, corresponding to the νFe-O of the magnetite core,¹⁶ is found in both the MNP/MNP-NH₂ (SI, Fig. S2B). A new band at ~1085 cm⁻¹ in the MNP-NH₂ may be assigned to the νSi-O of the silica shell. In addition, a broad band at ~ 3440 cm⁻¹ in the MNP-NH₂ may be assigned to νN-H or νO-H. These results are consistent with the MNP being successfully coated with an aminenated silica shell. Hydrodynamic diameters (HDs) of the MNP and MNP-NH₂ were measured as 28.9 ± 5.3 and 178±26 nm, respectively, by dynamic light scattering¹⁷ (SI, Fig. S2C). The MNP-NH₂ shows no apparent hysteresis with a saturated magnetization of ~40 emu/g in its magnetization curve (SI, Fig. S2D), lower than the MNP core (ca. ~90 emu/g),¹⁵b but is consistent with the coating of a non-magnetic shell. This confirms the MNP-NH₂ is super-paramagnetic and well-suited for biosensing: they form a stable, uniform dispersion in water but can be rapidly retrieved (1 min) with a magnet (SI, Fig. S3).

**Assay optimization**

A series of experiments are performed to optimise this assay.
First, MNP surface passivation: the MNP-NH$_2$ was first reacted with excess SM(PEG)$_{12}$ to introduce maleimides for cDNA conjugation, where any unreacted maleimides on the MNP could lead to non-specific enzyme adsorption and increasing background. A treatment with 2-mercaptoethanol (to cap unreacted maleimides) followed by bovine serum albumin blocking (1 mg/mL) was found highly effective, leading to a greatly improved signal to background (S/B) ratio (from ~4.4 to ~62, Fig. S4A).

Second, amount of MNP-cDNA: Despite of surface passivation, this cannot eliminate non-specific adsorption of enzymes on the MNPs completely. Fig. S4B showed assay signal increased with the increasing amount of MNP-cDNA1, indicating more efficient T-DNA capture, however, the background also increased. As a result, the net signal arising from T-DNA1 actually decreased when $> 20 \, \mu$g of MNP was used. Therefore, 20 $\mu$g MNP-cDNA was used for subsequent colorimetric sensing. Despite giving a weaker net T-DNA1 signal, 10 $\mu$g MNP-cDNA actually yielded the highest S/B ratio, so 10 $\mu$g MNP was used to assess the detection limit via fluorescence (both enzymatic products are fluorescent).

Third, temperature: Fig. S4C revealed that assay sensitivity was strongly temperature dependent: a 140% increase of sensitivity was observed as temperature was increased from 4 to 24 °C (~ RT), but showed no further increase as temperature was increased to 37 °C. Hence all assays were carried out conveniently at RT.$^{19}$

Fourth, cDNA loading: Increasing the cDNA loading on the MNP can increase the affinity and amount of cDNAs available for T-DNA hybridization, and hence may increase capture efficiency. Fig. S4D (SI) shows that the sensitivity increases first with the increasing cDNA loading, reaching the maximum value at ~0.5 nmol/mg (MNP). This loading was used for all subsequent assays.
DNA1 quantitation using HRP based signal amplification

The optimized conditions were used to quantitate T-DNA1 using MNP-cDNA1/HRP-sDNA1. Fig. 1 shows the absorption spectra of assay samples and the resulting calibration curve (1 h amplification). An extremely low background (sample without T-DNA1, $A_{571} = 0.0045\pm0.0003$) is observed. The signal is increased greatly with the T-DNA1 concentration, confirming the success of our assay strategy. In fact, the signal at 1 nM T-DNA1 (0.801±0.043) is ~177 times that of the blank, demonstrating an excellent S/B ratio and further confirming the success of our assay optimizations. A high S/B ratio is very beneficial for biosensing, allowing for direct, accurate target quantitation without the need of background correction.

![Absorption spectra and calibration curve](image)

**Fig. 1.** (A) Absorption spectra of the assay samples with different amounts of T-DNA1 using 1 h amplification, inset: a photograph of the corresponding samples; (B) the resulting calibration curve for T-DNA1 quantification, inset: amplified region over 0-200 pM T-DNA1 range fitted to a linear function.

Like typical enzymatic reactions, the whole calibration curve initially displays a rapid linear increase with T-DNA1 concentration up to 200 pM ($R^2 = 0.998$, Fig. 1B inset). As T-DNA1 concentration is increased further, the calibration curve deviates from linear and gradually becomes saturated, due to depleted Amplex red at higher target DNA (hence HRP) concentrations. This overall calibration curve could be fitted well ($R^2 = 0.991$) by the Hill equation, yielding an apparent k of 394 nM ($K_M$ minic of
enzyme activity) and n of 1.06, suggesting no strong target-binding cooperation.

**Fig. 2.** (A) Calibration curves for T-DNA1 quantitation using different enzymatic amplification times (0 – 300 mins, the amplification times are indicated on each curve). (B) The amplified region over 0 – 100 pM T-DNA1 range, and data were fitted to linear relationships. (C) The relationship between the corresponding slopes of calibration curves in (B) and the enzymatic amplification times.

We have evaluated the effect of amplification time on assay sensitivity. As shown in Fig. 2, over the whole 0 – 1 nM range, the calibration curves showed good linearity up to 120 mins of amplification time. As amplification time is extended further to >180 mins, the calibration curves deviate from linearity, due to depletion of the substrates under such conditions (10 μM Amplex red/H2O2). A careful examination of the calibration curves over 0-100 pM range (Fig. 2B) reveals all curves are linear. Besides, the corresponding slopes (representing sensitivity) are found to increase linearly with the amplification time (R² = 0.996, Fig. 2C), suggesting the sensitivity can be increased by extending the amplification time.

A large and systematically tunable dynamic range is important for biosensing. Here, we show that the amplification time can be used to tune the dynamic range and sensitivity: the dynamic range can be enlarged by reducing amplification time, while increasing the amplification time can be used to improve assay sensitivity. For example, extending the amplification time from 1 h to overnight (~15 h), this assay can directly detect 100 fM T-DNA unambiguously by UV-vis spectra with a good linearity over 0 -10
pM range ($R^2 = 0.990$, Fig. 3).

**Fig. 3.** Absorption spectra of samples containing different amounts of T-DNA3 detected by MNP-cDNA3 and HRP-sDNA3 with overnight amplification, inset: calibration curve with linear fit: $y = 0.0417 + 0.0314x$, $R^2 = 0.990$.

Moreover, by adapting a more sensitive readout using a conventional fluorescence plate reader,$^{18}$ this assay can detect 50 fM T-DNA3 directly without target pre-amplification over 2 h (SI, Fig. S5). Such a sensitivity is higher than many other sensitive, direct DNA sensing techniques, e.g. GNP amplified surface plasmon resonance (~10 pM)$^{20}$, microcantilever based nanomechanical sensing (~10 pM)$^{21}$, electrochemical DNA sensing (10-50 pM)$^{22}$, enzyme amplified electrochemical DNA sensing with a DNA tetrahedron (1 pM)$^{23}$, and silver nanoparticle amplified surface enhanced Raman scattering (1.1-33 pM)$^{24}$ It is also comparable to some ultrasensitive DNA assays, e.g. silver amplified GNP-DNA based scannometric assay/electric detection (~50 fM)$^{4,5}$, electrochemical DNA sensing via enzymatic amplification (~10 fM)$^{22a}$ or via a DNA supersandwich assembly (~100 fM)$^{22c}$ Another advantage here is that the MNP-target-En sandwiches can be easily retrieved magnetically, allowing MNP bound species to be concentrated or diluted where required, which can be further combined with the tuning of
amplification time to achieve the most desirable dynamic range and sensitivity for each specific assay needs.

The ability of performing assays in complex, clinical media is important for real world applications. To demonstrate this potential, T-DNA1 detection was carried out in 10% human serum (a clinical media, 1 h amplification). Fig. S6 (SI) reveals the T-DNA1 detection is unaffected in 10% human serum, yielding a good linear calibration curve ($R^2 = 0.999$). This demonstrates the MNP-enzyme assay is highly robust and works efficiently in complex media. We attribute this to the careful management of the MNP surfaces and assay conditions, leading to greatly reduced background and excellent assay robustness.

Stability of MNP-cDNA probe is another important parameter. Here we have performed T-DNA detection using MNP-cDNA probes after being stored (in pure water) for up to 5 months at 4 °C and compared the result with the freshly prepared probes. Results show that not only have the 5-month old MNP-cDNA probes remained magnetically responsive, but also given similar sensitivities (calibration slopes) to the fresh MNP-cDNA (SI, Fig S7), confirming the MNP-cDNA probe has good long-term stability.

**T-DNA2 detection using ALP based signal amplification**

The ALP catalyzed turnover of FDP into fluorescein ($\lambda_{\text{max}} = 485$ nm) was used to detect T-DNA2 using MNP-cDNA2 and ALP-s-DNA2 probes. The assay was done in Tris buffer because PBS inhibits the ALP activity (Fig. S1)$^{26}$. The resulting absorption spectra of assay samples (1 h amplification) and the corresponding calibration curve are shown in Fig. 4. A good positive linear calibration curve ($R^2 = 0.994$) over 0 - 200 pM range is obtained, suggesting that ALP is just as powerful as HRP in signal amplification, allowing direct quantitation of low pM T-DNA2 with 1 h amplification.
**Fig. 4.** (A) UV-vis absorption spectra of assay samples with different concentrations of T-DNA2 using ALP based signal amplification. (B) The corresponding $A_{485}$ versus concentration relationship of the assays and corresponding linear fit (red line).

### 3.7 Simultaneous quantification of two T-DNAs

The ability of simultaneously detect multiple DNA targets is important for high clinical diagnostic accuracy. To demonstrate this potential, two sets of MNP-cDNA and En-sDNA probes are used to detect two specific T-DNAs via HRP and ALP catalyzed reactions, respectively (Scheme 1).

**Fig. 5.** Simultaneous quantification of two DNA targets using the MNP-enzyme sandwich assay. (A) Absorption spectra of the assay samples with different amount of T-DNAs, T-DNA1:T-DNA2 molar ratio = 1:1. (B) The corresponding calibration curves for both DNA targets: data within the 0-500 pM range were both fitted to linear functions.
Fig. 5A shows the absorbance of both resorufin (A\textsubscript{571}) and fluorescien (A\textsubscript{485}) increased significantly with the increasing T-DNA1/T-DNA2 concentration (1 h amplification). Moreover, the correlations between the absorbance and T-DNA concentrations are both linear ($R^2 = 0.975$ for T-DNA1 and 0.994 for T-DNA2) over 0 - 500 pM range (Fig. 5B). Interestingly, both plots yield very similar slopes (sensitivities), in agreement with their similar substrate turnover rates.

These results thus established a simple, general versatile, doublet DNA assay that can be applied to target any DNA of interest by designing a pair of unique MNP-cDNA and En-sDNA probes for each specific T-DNA. Additional advantages here include, 1) the use of MNP based homogeneous, rapid target capture; 2) convenient magnetic separation for low assay background; 3) the use of large excesses of MNP-cDNA/En-sDNA probes to push T-DNA hybridization equilibrium towards the captured state, allowing efficient capture and detection of T-DNA at concentrations well below the $K_d$. As a result, this simple DNA assay can offer superior sensitivity and specificity over many other DNA sensing methods (Table S1) and appears well-suited for broad DNA based biosensing and diagnostics.

**Discrimination of cancer-specific SNP**

Genetic single-nucleotide polymorphism (SNP) is associated with numerous human diseases, e.g. cancer, diabetes, vascular disease, and some mental illness. To demonstrate the potential of our assay in SNP discrimination, the KRAS mutations (codon 12/13) associated with several human cancers, e.g. colorectal\textsuperscript{30}, pancreas\textsuperscript{31} and lung\textsuperscript{32}, are chosen as the DNA targets. This involves three T-DNAs (Table 1), the wild-type (T-DNA3) and two cancer-specific SNPs, T-DNA4 (17C$\rightarrow$T) and T-DNA5 (17C$\rightarrow$A). Here MNP-cDNA3 and HRP-sDNA3, fully complementary to T-DNA3, are used ($C_{T-DNA} = 100$ pM, 1 h amplification). Fig. 6 shows that the signal of the full-match T-DNA3 is considerably higher than the two
cancer-specific SNPs, confirming this assay can distinguish the normal gene from cancer-specific SNP mutants. The discrimination ratios (DRs) obtained here are 2.1 and 2.5 for T-DNA3/T-DNA4 and T-DNA3/T-DNA5 pairs, respectively. Reducing the assay buffer NaCl content from 150 to 100 mM improves the above SNP DRs to 3.3 and 4.2, respectively (SI, Fig. S8), comparable to many reported DNA SNP assays. Presumably because the formed MNP-dsDNA sandwiches containing a single mismatch (with cancer-specific SNPs) are affected more severely than the full-match wide-type target, in agreement with the literature. Further reducing the NaCl content to 50 mM produced significantly reduced signals for all T-DNAs (data not shown), due to greatly reduced stability of the dsDNA sandwich structure.

![UV-vis spectra](image)

**Fig. 6.** UV-vis spectra of samples showing the discrimination of cancer-specific SNPs (T-DNA4/TDNA5) against the wild-type non-cancer gene, inset: $A_{571}$ values of the DNA targets.

**Conclusion**

In summary, we have successfully developed a simple, sensitive and general MNP-enzyme sandwich assay that can be used for simultaneous quantitation of two specific DNA targets in clinically relevant media (e.g. 10% human serum). Combining the MNP-based rapid, efficient target capture and
convenient magnetic separation, and enzyme-based great signal amplification together with the careful optimization of the MNP surface and assay conditions, this assay can achieve direct, label-free quantitation of T-DNA down to 50 fM level within a total assay time of < 3 h (via fluorescence readout). Such a sensing performance places it among the very best of PCR- and label-free DNA assays in terms of both sensitivity and specificity (Table S1). Moreover, this assay is highly specific: it can effectively discriminate between a normal KRAS gene from its two cancer-specific SNPs. Therefore, we believe this robust, general, sensitive DNA sensing technology should have great potential in a wide range of DNA based biosensing and diagnostic applications. Currently, we are focused on further improving the assay sensitivity and extending its application to clinical samples.

Acknowledgement

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Supporting Information

Experimental details on MNP synthesis, surface modification, cDNA conjugation, and surface passivation as well as detailed assay procedures and supporting figures. This material is available free of charge via the Internet at http://pubs.acs.org
References and Notes


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