

Cis-regulatory hairpin-shaped mRNA encoding a reporter protein: catalytic sensing of nucleic acid sequence at single nucleotide resolution

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DNA sensing at a single nucleotide resolution is achieved using a hairpin-shaped, unmodified (unlabeled) RNA probe or the precursor double-stranded DNA (dsDNA) in a prokaryotic cell-free translation medium. The molecular-beacon-like probe consists of a loop region that is complementary to the target sequence and a stem composed of a ribosome-binding site (RBS) and its docking domain; the RBS is followed by the gene for a reporter protein such as luciferase or β -galactosidase. Target binding at the loop opens the hairpin to make RBS accessible by the ribosome to start translation of the reporter protein. This sensing system is signal amplifying by virtue of catalytic DNA-to-RNA transcription when using a dsDNA probe, catalytic RNA-to-protein translation, catalytic signal transduction by the enzymatic reaction of the translated reporter protein and, in the presence of RNase H, catalytic or even irreversible translation-activation of the target-probe heteroduplex. Preparation of a probe takes 1–3 d and gene sensing using the probe takes 1–3 h.

INTRODUCTION

Coupling of molecular beacon strategy to *cis*-acting RNA-controlled protein translation

As the number of identified genes continues to grow, the need increases for rapid and simple gene sensing with high sequence selectivity^{1,2}. Gene sensing of any type is based on selective target-probe hybridization and its output, which should be easily monitored. Molecular beacons (MBs) with a fluorescence resonance energy transfer (FRET) pair (Fig. 1a)³ utilize hybridization-induced conformational change of the probe from the light-off hairpin structure to a light-on open form. Much recent attention has also been paid to catalytic gene sensing^{4–13} through coupling to a signal-amplifying enzymatic^{4–8}, chemical^{9,10}, electrochemical¹¹ or magnetochemical¹² process. On the other hand, we reported a new strategy with MB-mRNA systems for selective genotyping¹³. This is based on naturally occurring¹⁴ or engineered¹⁵ hairpin-shaped or MB-like RNAs capable of riboregulator-dependent conformational change to control the translation activity, so that the sensing of the target oligodeoxynucleotide (ODN) as a regulator can be performed using a genetically encodable unmodified RNA as a probe in a typical prokaryotic translation system.

Design of MB-mRNA probes

The artificial translation regulation system (MB-mRNA; Fig. 1b) is composed of a *cis*-acting MB-like RNA structure, wherein the loop region (blue) is complementary to the target (green) and the stem consists of sequences for a ribosome-binding site (RBS; red) and the anti-RBS or RBS-docking domain complementary thereto (pink)¹³. The RBS is followed by a reporter gene sequence starting with an AUG start codon. The anti-RBS domain is preceded by another hairpin structure¹⁶. This is to endow an anti-RNase stability. Binding of the target (green) at the loop is expected to result in the opening of the MB structure, making the RBS domain accessible by the ribosome and hence initiating the translation of the reporter gene, such as luciferase.

The balance of the loop and the stem in terms of base-pair (bp) length is critical for sensitivity and selectivity. If the stem is long, the hairpin structure will be more stabilized and will be opened only by a longer target capable of more extensive target-probe hybridization. This will give a high sensitivity—that is, high target-on/target-off signal ratio—but low sequence selectivity. If, on the other hand,

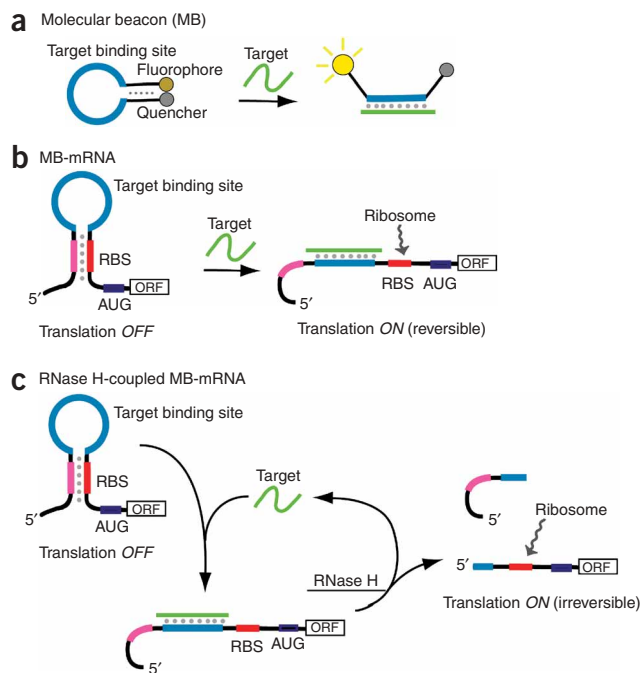


Figure 1 | Illustration of gene sensing systems. (a) MB, (b) MB-mRNA and (c) RNase H-coupled MB-mRNA sensing systems. The ribosome-binding site (RBS), anti-RBS or RBS-docking site, start codon, target-binding domain and target are shown in red, pink, purple, blue and green, respectively.



PROTOCOL



Figure 2 | Oligodeoxynucleotide targets. An 18-nt or 16-nt loop binding sequence is shown in green.

the stem is short, the hairpin structure will be less stabilized and easily opened even by a shorter target. This will give high sequence selectivity but a low sensitivity. Actually, we chose the commonly encountered 6-base AGGAGA sequence as RBS and set, after trial and error, an 8-bp stem and 18- or 16-bp target-probe hybridization at the loop to optimize sensitivity and selectivity.

Advantage of the method

The construction of MB-mRNA probes is apparently complex. The actual sensing takes time because it includes many steps of enzymatic conversion. The sensitivity or detection limit, 9 fmol (3.6 nM) as shown below, is not very high compared with those achieved recently by other methods¹⁷. Nevertheless, the present system has a number of unique aspects. (i) The unmodified (unlabeled) probe can be provided not only as RNA (MB-mRNA) but also in the form of much more stable and directly PCR-amplifiable double-stranded DNA (dsDNA), as the former is transcribed *in situ* from the latter in the cell-free translation system. (ii) The essential transcription and translation are natural processes of the cells. (iii) The sensing is multiply catalytic or signal amplifying, as all of the respective steps—that is, transcription of DNA to RNA, translation of RNA to protein and transduction of signal (chemiluminescence, for example) from the enzymatic reaction of the translated reporter protein—are catalytic. These aspects suggest that the present system is potentially applicable to in-cell gene sensing, as there are established protocols for the cellular uptake of dsDNA and expression therein of the encoded gene into protein. PCR-based methods, on the other hand, cannot be used in cells. The applicability of artificial signal-amplifying systems in cells is highly questionable. The usefulness of labeled probes in cells also remains to be elucidated as regards to the efficacy of cellular uptake, extent of toxicity and compatibility of the sensing reaction, such as FRET, with cellular environments.

Another merit of the present system is that the reporter protein can, in principle, be of any type. This is particularly important in terms of single nucleotide polymorphism (SNP) detection, since we can easily construct a set of allele-sensitive probes using a class of otherwise closely related reporter proteins that can be distinguished from each other.

Outline of protocol

Although our ultimate goal is to establish *in vivo* gene sensing using the present system, the scope of this protocol is to describe its *in vitro* performance at a quantitative level, thus providing a basis for its in-cell application on one hand and, on the other hand, a guideline for the *in vitro* use of RNA systems for selection/sensing of translation-controlling small molecules (riboswitches)¹⁸. The targets are ODNs (Fig. 2) copying the 18-nt (620–637) or 16-nt (620–635) sequence (green) of the HIV 1-related human CC chemokine receptor 5 (CCR5) with a mutation at position 627 (ref. 19). The actual targets with variation at the corresponding

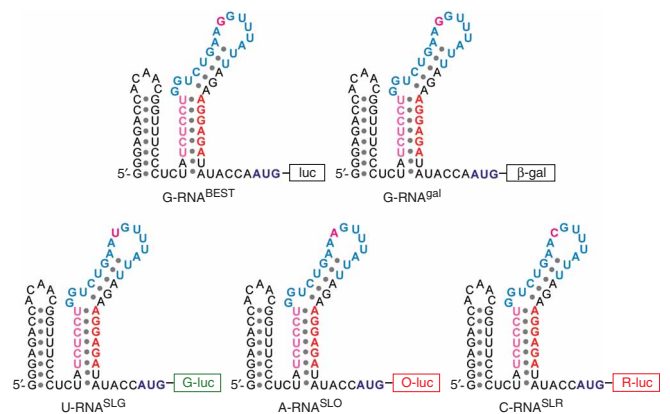


Figure 3 | Predicted secondary structures of the MB-mRNA probes. These contain the ORF domain for the reporter protein (either luciferase or β -galactosidase), RBS (red), target-binding site (blue) with 1-nt mutation shown in red and anti-RBS sequence (pink).

position shown in red are designated 18X and 16X (X = A, T, C or G) (Fig. 2). We also used a doubly mutated target 18TA.

Luciferase was our first choice as sensing output because of its high sensitivity, the good linearity of the related chemiluminescence assay and the ready color tuning (see below). Although this system requires luciferin as an external assay reagent, the latter is readily taken up into the cell and can thus be used for *in vivo* sensing²⁰. A prototypical 18C-targeting MB-mRNA probe, designated G-RNA^{BEST} (Fig. 3), was obtained by sequential PCRs on the luciferase gene (*luc*) encoded in plasmid vector pBESTluc and subsequent transcription (Fig. 4). The first PCR added the RBS (red), an 18-nt target-binding site (blue) with a target-complementary G base, and an intervening spacer. The second PCR added an anti-RBS domain (pink) and an additional hairpin structure preceded by the T7 promoter region. The primer sequences used are summarized in Figure 5. For sequence-selective production of multicolor reporter proteins, we took advantage

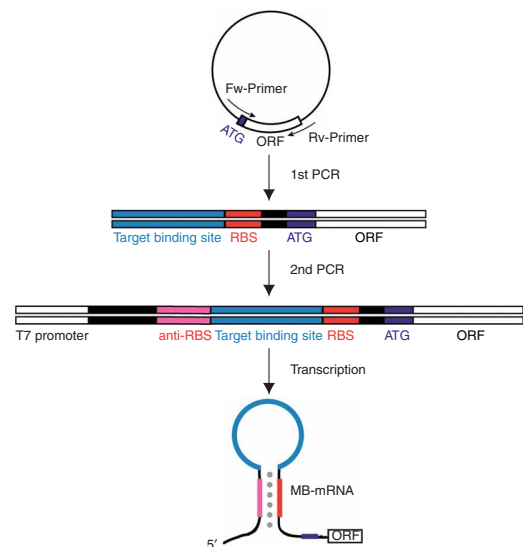


Figure 4 | Illustration of the preparation of MB-mRNA probes. This starts from a plasmid vector encoding a reporter protein.

of the MultiReporter Assay System. Three plasmid vectors, pSLG-test, pSLO-test and pSLR-test, encode green-luminescing wide-type *Rhagophthalmus ohbai* firefly luciferase (G-luc; emission maximum, 550 nm), orange-luminescing mutated (T226N) *R. ohbai* firefly luciferase (O-luc; 580 nm) and red-luminescing wild-type *Phrixothrix hirtus* firefly luciferase (R-luc; 630 nm), respectively^{21–23}. They were PCR-amplified as above to incorporate an A-sensitive, T-sensitive or G-sensitive target-binding site, respectively, to give U-RNA^{SLG}, A-RNA^{SLO} and C-RNA^{SLR} (Fig. 3). We also prepared an 18C-targeting probe, G-DNA^{gal} (Fig. 3), encoding β-galactosidase (β-gal). The latter cleaves *o*-nitrophenyl β-galactoside as a substrate to give colored *o*-nitrophenol for colorimetric or visual assay.

To further enhance catalytic performance, we also use RNase H. The translation-activated mRNA probe reversibly formed from stoichiometric binding of the target to the probe (structure on the right-hand side of Fig. 1b) is, at best, equimolar to the target. This would lead to low signal intensity, especially when the target is present in a tiny amount. RNase H is an endonuclease that specifically cleaves RNA in the DNA/RNA heteroduplex²⁴. It thus irreversibly digests the target-bound loop region of the probe to release the anti-RBS domain from the mRNA body, thus allowing

1st-fw-G-BEST	5'-d (GGTCTGAAGTTTATTAGAAGGAGATATACCAATGGAAAGCCAAAAACATA) -3'
1st-rv-G-BEST, 2nd-rv-G-BEST	5'-d (TATTCATTACAATTTGGACTTTCGCC) -3'
2nd-fw-G-BEST	5'-d (GAAATTAATACGACTCCTACTATAGGGAGACCACAACCGTTTCCCTCTATCTCCTGGTCTGAAGTTTA) -3'
1st-fw-U-SLG	5'-d (GGTCTGAAGTTTATTAGAAGGAGATATACCAATGGCTAACGAGATCATCC) -3'
1st-fw-A-SLO	5'-d (GGTCTGAAGTTTATTAGAAGGAGATATACCAATGGCTAACGAGATCATCC) -3'
1st-fw-C-SLR	5'-d (GGTCTGAAGTTTATTAGAAGGAGATATACCAATGGAAAGAGACATCGTAATGG) -3'
1st-rv-U-SLG, 1st-rv-A-SLO, 2nd-rv-U-SLG	5'-d (TATTCATTACAGCTTGGACTTCTCTG) -3'
2nd-rv-A-SLO, 3rd-rv-U-SLG, 3rd-rv-A-SLO	5'-d (TATTCATTACAGCTTGGACTTCTCTG) -3'
1st-rv-C-SLR, 2nd-rv-C-SLR, 3rd-rv-C-SLR	5'-d (TATTCATTACAGCTTGGACTTGGCC) -3'
2nd-fw-U-SLG	5'-d (GAAATTAATACGACTCCTACTATAGGGAGACCACAACCGTTTCCCTCTATCTCCTGGTCTGAAGTTTA) -3'
2nd-fw-A-SLO	5'-d (GAAATTAATACGACTCCTACTATAGGGAGACCACAACCGTTTCCCTCTATCTCCTGGTCTGAAGTTTA) -3'
2nd-fw-C-SLR	5'-d (GAAATTAATACGACTCCTACTATAGGGAGACCACAACCGTTTCCCTCTATCTCCTGGTCTGAAGTTTA) -3'
3rd-fw-U-SLG, 3rd-fw-A-SLO, 3rd-fw-C-SLR	5'-d (GAAATTAATACGACTCCTACTATAGG) -3'
1st-fw-G-gal	5'-d (GGTCTGAAGTTTATTAGAAGGAGATATACCAATGACCATTGACGGATTAC) -3'
1st-rv-G-gal, 2nd-rv-G-gal	5'-d (GGATTAGTTTATTCATTTTGTGACACCAGACCAAC) -3'
2nd-fw-G-gal	5'-d (GAAATTAATACGACTCCTACTATAGGGAGACCACAACCGTTTCCCTCTATCTCCTGGTCTGAAGTTTATTAG) -3'

Figure 5 | Sequences of primers. The T7 promoter domains are underlined.

catalytic use of target DNA (RNase H-coupled MB-mRNA; Fig. 1c) with improved selectivity and enhanced sensitivity²⁵, although, given the nature of RNase H, it can be used only *in vitro* for DNA targets and not for in-cell gene sensing, where the targets should be mRNAs.

The translation- or transcription/translation-coupled sensing of ODN targets can be carried out in a normal *Escherichia coli* S30 extract system such as RTS HY100 (Roche), but we use a reconstituted prokaryotic cell-free transcription/translation system (Pure System; Post Genome Institute)²⁶ throughout this work. Colorimetric assay of β-galactosidase cannot be conducted in the *E. coli* extract, which itself contains the enzyme. The effect of RNase H can also be most clearly evaluated in the reconstituted medium, which is otherwise free from RNase H.

MATERIALS REAGENTS

- Agarose for 50–800-bp fragments (Nacalai Tesque, cat. no. 01147-96) for gel electrophoresis (see REAGENT SETUP)
- β-Galactosidase enzyme assay system (Promega, cat. no. E2000) accompanied by the reporter lysis buffer
- DNA oligomers as primers in PCR (made to order by Gene Design Inc.)
- DNA stable marker 1-kbp ladder (Sigma Genosys, cat. no. MBMA1KBP-S)
- DNeasy tissue kit (Qiagen, cat. no. 69504) for purification of total DNAs
- dNTPs (accompanied by respective polymerase samples)
- *E. coli* K-12 MG1655 strain (ATCC, for example)
- Ethidium bromide solution (Nippon Gene, cat. no. 315-90051) for gel electrophoresis **! CAUTION** Toxic.
- Ethanol (Wako Pure Chemical Industries, cat. no. 053-06531) **! CAUTION** Highly flammable.
- GFX PCR DNA and gel band purification kit (GE Healthcare, cat. no. 27-9602-01) for purification of PCR products
- Luciferase assay kit (Promega, cat. no. E1501)
- Luria broth base powder (Invitrogen, cat. no. 12795-027)
- MEGAShortScript T7 (Ambion, cat. no. 1354) or MEGAScript T7 Kit (Ambion, cat. no. 1333) for transcription
- pBESTluc vector (Promega)
- pSLG-test vector (Toyobo, cat. no. MRV-101)
- pSLO-test vector (Toyobo, cat. no. MRV-102)
- pSLR-test vector (Toyobo, cat. no. MRV-103)
- *PfuUltra* high-fidelity DNA polymerase (Stratagene, cat. no. 600380) for PCR
- Pure system standard classic II (Post Genome Institute, cat. no. PURE2030C) or pure system classic I (Post Genome Institute) as a reconstituted prokaryotic transcription/translation medium
- *Pyrobest* DNA polymerase (Takara, cat. no. R005A) for PCR
- RNase-free water (molecular biology reagent) (Sigma, cat. no. W4502)
- **▲ CRITICAL** RNase-free water should be used throughout the procedure.
- RNeasy MinElute Cleanup kit (Qiagen, cat. no. 74204) for purification of RNA

- TaKaRa Ex *Taq* hot start version (TaKaRa, cat. no. RR006A) for PCR
- Tris-Acetate-EDTA (TAE) buffer (50×) (Nacalai Tesque, cat. no. 32666-81)
- Tth RNase H (Toyobo, cat. no. RNH-201)
- Buffer RPE (see REAGENT SETUP)
- LB medium (see REAGENT SETUP)
- Buffer AW1 (see REAGENT SETUP)
- Buffer AW2 (see REAGENT SETUP)

EQUIPMENT

- 96-well assay plate (Costar)
- Bio-shaker (Taitec, cat. no. BR-300LF)
- Centrifuge (Waken, model 2320)
- Densitograph (ATTO, cat. no. AE-6920V-FX)
- GeneQuant *pro* (GE Healthcare)
- iCycler thermalcycler dual block (Bio-Rad) for PCR
- Lumat LB9507 luminometer (Berthold Detection System)
- *i*-Mupid (COSMO BIO) for electrophoresis
- Multilabel counter (Wallac, cat. no. 1420) for luciferase/galactosidase assay
- 25-ml reagent reservoir (VistaLab) for multichannel pipetting
- Surgical stainless steel blades and handles (Feather Safety Razor) for gel cutting

REAGENT SETUP

1% Agarose gel Weigh 0.5 g agarose, and dissolve in 50 ml MilliQ-grade deionized hot water. When the solution has cooled to 50 °C, add 1 ml 50× TAE buffer and 5 μl 10 μg μl⁻¹ ethidium bromide solution. Mix thoroughly, and pour into gel tray.

Buffer RPE Combine 11 ml of the supplied concentrated buffer RPE (RNeasy MinElute Cleanup kit) and 44 ml ethanol.

Buffer AW1 Combine 19 ml of the supplied concentrated buffer AW1 (DNeasy Tissue kit) and 25 ml ethanol.

Buffer AW2 Combine 13 ml of the supplied concentrated buffer AW2 (DNeasy Tissue kit) and 30 ml ethanol.

LB medium Combine 7.5 g Luria broth base powder (Invitrogen) and 300 ml MilliQ-grade deionized water, and heat the mixture in an autoclave at 121 °C for 20 min.



PROTOCOL

PROCEDURE

Preparation of template dsDNA (G-DNA^{BEST}) ● TIMING 5 h

1| Prepare a first PCR mix containing the following:

Reagents	Volume (μl)
10× <i>PfuUltra</i> reaction buffer	2.5
dNTPs (2 mM each)	2.5
Forward primer (1st-fw-G-BEST; Fig. 5) (2 μM)	2.5
Reverse primer (1st-rv-G-BEST; Fig. 5) (2 μM)	2.5
<i>PfuUltra</i> DNA polymerase (2.5 U μl ⁻¹)	0.5
H ₂ O	13.5
Total volume	24

▲ **CRITICAL STEP** To prevent contamination, use nuclease-free filter tips.

2| Add 1 μl pBESTluc solution (5 ng μl⁻¹) or 1 μl water (for negative control) to each mix in a PCR tube.

3| Run the following PCR cycles:

Cycle number	Denaturation	Annealing	Polymerization	Final
3 min at 95 °C				
2–31	45 s at 95 °C	45 s at 54 °C	1 min at 72 °C	
32			10 min at 72 °C	
Hold				4 °C

4| Check the PCR by running 5 μl of each reaction mixture on a 1% agarose gel with a 1-kbp ladder as a marker.

■ **PAUSE POINT** The PCR product can be stored at –80 °C for 3 months.

5| Dilute the PCR product with water to give a 1:99 dilution.

6| Prepare a second PCR mix containing the following:

Reagents	Volume (μl)
10× <i>PfuUltra</i> reaction buffer	2.5
dNTPs (2 mM each)	2.5
Forward primer (2nd-fw-G-BEST; Fig. 5) (2 μM)	2.5
Reverse primer (2nd-rv-G-BEST; Fig. 5) (2 μM)	2.5
<i>PfuUltra</i> DNA polymerase (2.5 U μl ⁻¹)	0.5
H ₂ O	13.5
Total volume	24

▲ **CRITICAL STEP** To prevent contamination, use nuclease-free filter tips.

7| Add 1 μl of the diluted first PCR solution or 1 μl water (for negative control) to each mix in a PCR tube.

8| Perform the PCR in the same manner as in Step 3, and check as in Step 4 to ensure the formation of the desired, allele C-targeting dsDNA template (G-DNA^{BEST}).

▲ **CRITICAL STEP** If the purity (as judged by gel electrophoresis) is not satisfactory (more than one spot), purify the PCR product with a GFX PCR DNA and gel band purification kit for subsequent transcription to give a single spot.

■ **PAUSE POINT** The PCR product can be stored at –80 °C for 3 months.

Preparation of RNA probe G-RNA^{BEST} ● TIMING 15 h

9| Prepare the following MEGAshortscript T7 transcription mix in a PCR tube (MEGAscript T7 can be used in place of MEGAshortscript):

Reagents	Volume (μl)
T7 10× reaction buffer	2.0
T7 NTP solution (75 mM)	2.0 each
T7 enzyme mix	2.0
G-DNA ^{BEST} (2nd PCR solution; Step 8)	2.0
H ₂ O	6.0
Total volume	20

- 10| Mix briefly, spin down and tap the sample.
 ■ **PAUSE POINT** Incubate the mixture overnight at 37 °C.
 11| To decompose the DNA template, add 1 µl TURBO DNase solution, mix well and incubate at 37 °C for 15 min.
 12| Purify the transcribed RNA probe (G-RNA^{BEST}) with an RNeasy MinElute Cleanup Kit according to the manufacturer's manual.
 13| Determine the concentration of the purified probe by the absorbance at 260 nm, and dilute to give a stock solution of G-RNA^{BEST} (1 µg µl⁻¹).
 ▲ **CRITICAL STEP** Avoid repeated freezing and thawing of the sample, because this results in decomposition of the RNA. For repeated use, divide the stock solution into 10-µl aliquots and freeze separately.
 ■ **PAUSE POINT** The transcription product (G-RNA^{BEST}) can be stored at -80 °C for 2 weeks.

RNase H–uncoupled sensing of 18-nt targets with probe G-RNA^{BEST} ● TIMING 1.5 h

- 14| Prepare a reconstituted prokaryotic translation medium (solution AB) by mixing 35 µl of solution A and 5 µl of solution B of pure system classic I.
 ▲ **CRITICAL STEP** A normal *E. coli* S30 extract system (RTS HY100, Roche, cat no. 3 186 148) can be used with similar results in place of the reconstituted translation system.
 15| Mix probe G-RNA^{BEST} and 18-nt full-match target (18C) in a PCR tube, as follows:

Reagents	Volume (µl)
Solution AB	8.0
Probe G-RNA ^{BEST} (Step 13) (1 µg µl ⁻¹ or 1.8 pmol µl ⁻¹)	1.0
Target 18C (1.8 pmol µl ⁻¹)	1.0
Total volume	10

For negative control, add 1 µl water in place of target 18C.

▲ **CRITICAL STEP** Negative control without target is critical, as the target-on chemiluminescence intensity should always be evaluated in reference to that in the absence of the target.

- 16| Spin down and tap the mixture gently, and incubate for 1 h at 37 °C.
 17| Add 10 µl water into each sample in the tube.
 18| Take three 5-µl aliquots from each sample, and put them separately in a 96-well assay plate set in a multilabel counter (Wallac 1420).
 19| Add 100 µl luciferase assay reagent to each well, and read the chemiluminescence intensity (*I*). Confirm that the presence of the probe gives rise to approximately threefold higher intensity than is seen in its absence ($I_{on}/I_{off} \cong 3$).
 ▲ **CRITICAL STEP** To standardize conditions for all samples, it is critical to use a multichannel pipette and reagent reservoir and to read the chemiluminescence intensity immediately after mixing the sample with the reagent by pipetting. Chemiluminescence intensity gradually decreases in a time-dependent manner, giving approximately 50% loss of intensity in 10 min.

? TROUBLESHOOTING

RNase H–coupled sensing of 18-nt and 16-nt targets with probe G-RNA^{BEST} ● TIMING 2 h for each set of targets

- 20| For RNase H-coupled sensing, mix probe G-RNA^{BEST}, 18-nt target (fully matched 18C, singly A-mismatched 18A, singly T-mismatched 18T or doubly TA-mismatched 18TA) and RNase H in a PCR tube, as follows:

Reagents	Volume (µl)
Solution AB	8.0
Probe G-RNA ^{BEST} (Step 13) (1 µg µl ⁻¹ or 1.8 pmol µl ⁻¹)	1.0
Target 18C, 18A or 18TA (1.8 pmol µl ⁻¹)	1.0
Tth RNase H (0.1 U µl ⁻¹)	1.0
Total volume	11

For negative control, add 1 µl water in place of target ODN.

▲ **CRITICAL STEP** Negative control without target is essential, as the target-on chemiluminescence intensity should always be evaluated in reference to that in the absence of the target.

- 21| Follow the translation (Step 16), workup (Steps 17 and 18) and assay (Step 19) procedures as above with 9 µl water in place of 10 µl in Step 17. Confirm that the I_{on}/I_{off} ratio (target-on to target-off signal ratio) for the full-match target 18C is enhanced from 3 (in the absence of RNase H) to 6 (in its presence) with more (for 18A and 18TA) or less (for 18T) efficient base discrimination.



PROTOCOL

▲ CRITICAL STEP To standardize conditions for all samples, it is critical to use a multichannel pipette and reagent reservoir and to read the chemiluminescence intensity immediately after mixing the sample with the reagent by pipetting. Chemiluminescence intensity gradually decreases in a time-dependent manner, giving approximately 50% loss of intensity in 10 min.

▲ CRITICAL STEP To see the clear effect of RNase H, it is critical to use the reconstituted translation system, which is free from RNase H.

? TROUBLESHOOTING

22| To check the effect of base-pair length in target-probe hybridization, use 16-nt targets (fully matched 16C and singly T-mismatched 16T) in place of the 18-nt counterparts under RNase H-uncoupled (Steps 15–19) and RNase H-coupled (Steps 20 and 21) conditions. Confirm that a significant on/off ratio ($I_{\text{on}}/I_{\text{off}} \cong 6$) is achieved only for the full-match target 16C, with a satisfactory T-allele discrimination capacity, under the RNase H-coupled conditions.

▲ CRITICAL STEP To standardize conditions for all samples, it is critical to use a multichannel pipette and reagent reservoir and to read the chemiluminescence intensity immediately after mixing the sample with the reagent by pipetting. Chemiluminescence intensity gradually decreases in a time-dependent manner, giving approximately 50% loss of intensity in 10 min.

? TROUBLESHOOTING

RNase H-coupled sensing with G-DNA^{BEST} as a probe ● TIMING 1.5 h

23| Mix dsDNA template G-DNA^{BEST} (Step 8), 18-nt full-match target 18C and RNase H in a PCR tube, as follows:

Reagents	Volume (μl)
Solution AB	8.0
Probe G-DNA ^{BEST} (Step 8) (0.2 pmol μl^{-1})	1.0
Target 18C (1.8 pmol μl^{-1})	1.0
Tth RNase H (0.1 U μl^{-1})	1.0
Total volume	11

For negative control, add 1 μl water in place of target 18C.

▲ CRITICAL STEP Negative control without target is essential, as the target-on chemiluminescence intensity should always be evaluated in reference to that in the absence of the target.

24| Follow the translation (Step 16), workup (Steps 17 and 18) and assay (Step 19) procedures as above with 9 μl water in place of 10 μl in Step 17. Confirm that the on/off ratio is much enhanced, to approximately 13.

▲ CRITICAL STEP To standardize conditions for all samples, it is critical to use a multichannel pipette and reagent reservoir and to read the chemiluminescence intensity immediately after mixing the sample with the reagent by pipetting. Chemiluminescence intensity gradually decreases in a time-dependent manner, giving approximately 50% loss of intensity in 10 min.

? TROUBLESHOOTING

Preparation of template dsDNAs of multicolor reporter proteins (T-DNA^{SLG}, A-DNA^{SLO} and C-DNA^{SLR}) ● TIMING 6 h for each template

25| For the preparation of T-DNA^{SLG}, which is transcribable into A-targeting U-RNA^{SLG}, prepare a first PCR mix containing the following:

Reagents	Volume (μl)
10 \times <i>Pyrobest</i> buffer II	2.0
dNTPs (2.5 mM each)	2.0
Forward primer (1st-fw-U-SLG; Fig. 5) (2 μM)	2.0
Reverse primer (1st-rv-U-SLG; Fig. 5) (2 μM)	2.0
<i>Pyrobest</i> DNA polymerase (1.0 U μl^{-1})	1.25
H ₂ O	8.75
Total volume	18

▲ CRITICAL STEP To prevent contamination, use nuclease-free filter tips.

26| Add 2 μl solution of green luciferase-encoding vector pSLG-test (5 ng μl^{-1}) or 2 μl water (for negative control) to each mix in a PCR tube.

27| Run the following PCR cycles:

Cycle number	Denaturation	Annealing	Polymerization	Final
1–20	10 s at 98 °C	30 s at 56 °C	1 min at 72 °C	
Hold				4 °C

28| Check the PCR by running 5 μ l of reaction mixture on a 1% agarose gel with a 1-kbp ladder as a marker. If the purity (as judged by gel electrophoresis) is not satisfactory (more than one spot), purify the PCR product with a GFX PCR DNA and gel band purification kit according to the manufacturer's manual.

■ **PAUSE POINT** The PCR product can be stored at -80°C for 3 months.

29| Dilute the purified PCR product with water to give a 1:199 dilution.

30| Prepare a second PCR mix containing the following:

Reagents	Volume (μ l)
10 \times <i>Pyrobest</i> buffer II	2.0
dNTPs (2.5 mM each)	2.0
Forward primer (2nd-fw-U-SLG; Fig. 5) (2 μ M)	2.0
Reverse primer (2nd-rv-U-SLG; Fig. 5) (2 μ M)	2.0
<i>Pyrobest</i> DNA polymerase (1.0 U μ l $^{-1}$)	1.25
H ₂ O	9.75
Total volume	19

▲ **CRITICAL STEP** To prevent contamination, use nuclease-free filter tips.

31| Add 1 μ l of the diluted purified first PCR solution or 1 μ l water (for negative control) to each mix in a PCR tube.

32| Perform the PCR in exactly the same manner as in Step 27.

33| Check the purity and, when it is not satisfactory, purify the product as in Step 28.

■ **PAUSE POINT** The PCR product can be stored at -80°C for 3 months.

34| Dilute the purified second PCR product as in Step 29.

35| Prepare a third PCR mix containing the following:

Reagents	Volume (μ l)
10 \times <i>Pyrobest</i> buffer II	2.0
dNTPs (2.5 mM each)	2.0
Forward primer (3rd-fw-U-SLG; Fig. 5) (2 μ M)	2.0
Reverse primer (3rd-rv-U-SLG; Fig. 5) (2 μ M)	2.0
<i>Pyrobest</i> DNA polymerase (1.0 U μ l $^{-1}$)	1.25
H ₂ O	8.75
Total volume	18

36| Add 2 μ l of the diluted purified second PCR solution or 2 μ l water (for negative control) to each mix in a PCR tube.

37| Perform the PCR in exactly the same manner as in Step 27.

38| Check the purity, and purify the product dsDNA, that is, T-DNA^{SLG}, as in Step 28.

■ **PAUSE POINT** The PCR product can be stored at -80°C for 3 months.

39| For the preparation of A-DNA^{SLO} transcribable into T-targeting A-RNA^{SLO}, follow Steps 25–38 using orange luciferase-encoding vector pSLO-test and A-SLO relevant primers (1st-fw-A-SLO, 1st-rv-A-SLO, 2nd-fw-A-SLO, 2nd-rv-A-SLO, 3rd-fw-A-SLO and 3rd-rv-A-SLO) in place of pSLG-test (Step 26) and T-SLG relevant ones (1st-fw-U-SLG, 1st-rv-U-SLG, 2nd-fw-U-SLG and 2nd-rv-U-SLG, 3rd-fw-U-SLG and 3rd-rv-U-SLG) (Steps 25, 30 and 35).

40| For the preparation of C-DNA^{SLR} transcribable into G-targeting C-RNA^{SLR}, follow Steps 25–38 using red luciferase-encoding vector pSLR-test and C-SLR relevant primers (1st-fw-C-SLR, 1st-rv-C-SLR, 2nd-fw-C-SLR, 2nd-rv-C-SLR, 3rd-fw-C-SLR and 3rd-rv-C-SLR) in place of pSLG-test (Step 26) and U-SLG relevant ones (1st-fw-U-SLG, 1st-rv-U-SLG, 2nd-fw-U-SLG, 2nd-rv-U-SLG, 3rd-fw-U-SLG and 3rd-rv-U-SLG) (Steps 25, 30 and 35).

Preparation of RNA probes, U-RNA^{SLG}, A-RNA^{SLO} and C-RNA^{SLR} ● TIMING 15 h for each probe

41| For U-RNA^{SLG}, prepare the following MEGAscript T7 transcription mix in a PCR tube (MEGAscript T7 can be used in place of MEGAscript):



PROTOCOL

Reagents	Volume (μl)
T7 10 \times reaction buffer	1.0
T7 NTP solution (75 mM)	1.0 each
T7 enzyme mix	1.0
T-DNA ^{SLG} (third PCR solution) (Step 38)	3.0
H ₂ O	1.0
Total volume	10

42| Carry out transcription as in Steps 10–12. Determine the concentration of purified product by the absorbance at 260 nm, and dilute to give a stock solution of probe U-RNA^{SLG} (1 $\mu\text{g } \mu\text{l}^{-1}$).

▲ CRITICAL STEP Avoid repeated freezing and thawing of the sample, because this results in decomposition of the RNA. For repeated use, divide the stock solution into 10- μl aliquots and freeze.

■ PAUSE POINT The transcription product (U-RNA^{SLG}) can be stored at -80°C for 2 weeks.

43| For the preparation of A-RNA^{SL0}, follow Steps 41 and 42 using A-DNA^{SL0} in place of T-DNA^{SLG}.

44| For the preparation of C-RNA^{SLR}, follow Steps 41 and 42 using C-DNA^{SLR} in place of T-DNA^{SLG}.

Sensing of 16-nt targets with allele-selective RNA probes ● TIMING 2 h for each set of targets

45| Prepare a reconstituted prokaryotic translation medium (solution AB) by mixing 25 μl solution A and 10 μl solution B of pure system classic II.

46| Mix probe U-RNA^{SLG}, target (16A, 16T, 16C or 16G) and RNase H in a PCR tube, as follows:

Reagents	Volume (μl)
Solution AB	8.0
Probe U-RNA ^{SLG} (Step 42) (1 $\mu\text{g } \mu\text{l}^{-1}$ or 1.8 pmol μl^{-1})	1.0
Target 16A, 16T, 16C or 16G (1.8 pmol μl^{-1})	1.0
Tth RNase H (1.0 U μl^{-1})	1.0
Total volume	11

For negative control, add 1 μl water in place of target ODN.

▲ CRITICAL STEP Negative control without target is essential, as the target-on chemiluminescence intensity should always be evaluated in reference to that in the absence of the target.

47| Carry out target-dependent translation of green luciferase as in Step 16. Follow the workup and assay procedures as in Steps 17–19, but by adding 11 μl water in place of 10 μl in Step 17, taking 5.5- μl aliquots in place of 5- μl aliquots in Step 18 and using a luminometer in place of a multilabel counter. Confirm that only A-allele target 16A gives rise to a significant increase in the chemiluminescence intensity, that is, on/off ratio.

▲ CRITICAL STEP To standardize conditions for all samples, it is critical to read the chemiluminescence intensity immediately after mixing the sample with the reagent by pipetting. Chemiluminescence intensity gradually decreases in a time-dependent manner, giving approximately 50% loss of intensity in 10 min.

? TROUBLESHOOTING

48| Mix probe A-RNA^{SL0}, target (16A, 16T, 16C or 16G) and RNase H in a PCR tube as in Step 46. Follow the translation and assay procedures in Step 47 for orange luciferase. Confirm that only T-allele target 16T gives rise to a significant increase in the chemiluminescence intensity, that is, on/off ratio.

49| Mix probe C-RNA^{SLR}, target (16A, 16T, 16C or 16G) and RNase H in a PCR tube as in Step 46. Follow the translation and assay procedures in Step 47 for red luciferase. Confirm that only G-allele target 16G gives rise to a significant increase in the chemiluminescence intensity, that is, on/off ratio.

▲ CRITICAL STEP For multicolor SNP detection in a single tube, it is critical to optimize the quantities of the three probes or use $I_{\text{on}}/I_{\text{off}}$ criteria for evaluation, as the translation efficiencies and emission coefficients for the three types (green, orange and red) of luminescence are different.

Preparation of dsDNA template G-DNA^{gal} for a colorimetric assay based on a β -galactosidase reporter protein ● TIMING 25 h

50| Inoculate *E. coli* K-12 MG1655 strain in 5 ml LB medium in a plastic tube.

51| Incubate the mixture for 16 h at 37 $^\circ\text{C}$ with vigorous shaking.

52| Purify total DNAs in the strain with a DNeasy tissue kit according to the manufacturer's manual.

53| Determine the concentration of the purified total DNAs by the absorbance at 260 nm.

■ **PAUSE POINT** The DNAs can be stored at $-80\text{ }^{\circ}\text{C}$ for 3 months.

54| Prepare the first PCR mix containing the following:

Reagents	Volume (μl)
10 \times <i>Ex Taq</i> buffer	2.5
dNTPs (2.5 mM each)	2.0
Forward primer (1st-fw-G-gal; Fig. 5) (2 μM)	2.5
Reverse primer (1st-rv-G-gal; Fig. 5) (2 μM)	2.5
<i>Ex Taq</i> DNA polymerase (0.5 U μl^{-1})	1.25
H ₂ O	13.25
Total volume	24

▲ **CRITICAL STEP** To prevent contamination, use nuclease-free filter tips.

55| Add 1 μl of the total DNA solution obtained above (5 ng μl^{-1}) or 1 μl water (for negative control) to each mix in a PCR tube.

56| Run the following PCR cycles:

Cycle number	Denaturation	Annealing	Polymerization	Final
1	2 min at 94 $^{\circ}\text{C}$			
2–24	15 s at 94 $^{\circ}\text{C}$	15 s at 54 $^{\circ}\text{C}$	2 min at 72 $^{\circ}\text{C}$	
25			5 min at 72 $^{\circ}\text{C}$	
Hold				4 $^{\circ}\text{C}$

57| Check the PCR by running 5 μl of each reaction on a 1% agarose gel with a 1-kbp ladder as a marker.

■ **PAUSE POINT** The PCR product can be stored at $-80\text{ }^{\circ}\text{C}$ for 3 months.

58| Dilute the PCR product with water to give a 1:99 dilution.

59| Prepare the second PCR mix containing the following:

Reagents	Volume (μl)
10 \times <i>Ex Taq</i> buffer	2.5
dNTPs (2.5 mM each)	2.0
Forward primer (2nd-fw-G-gal; Fig. 5) (2 μM)	2.5
Reverse primer (2nd-rv-G-gal; Fig. 5) (2 μM)	2.5
<i>Ex Taq</i> DNA polymerase (0.5 U μl^{-1})	1.25
H ₂ O	13.25
Total volume	24

▲ **CRITICAL STEP** To prevent contamination, use nuclease-free filter tips.

60| Add 1 μl of the diluted first PCR solution or 1 μl water (for negative control) to each mix in a PCR tube.

61| Run the following PCR cycles:

Cycle number	Denaturation	Annealing	Polymerization	Final
1	2 min at 94 $^{\circ}\text{C}$			
2–20	15 s at 94 $^{\circ}\text{C}$	15 s at 55 $^{\circ}\text{C}$	2 min at 72 $^{\circ}\text{C}$	
21			5 min at 72 $^{\circ}\text{C}$	
Hold				4 $^{\circ}\text{C}$

62| Check as in Step 57 to make sure of the formation of the desired dsDNA template, that is, G-DNA^{gal}.

■ **PAUSE POINT** The PCR product can be stored at $-80\text{ }^{\circ}\text{C}$ for 3 months.

RNase H–coupled sensing with G-DNA^{gal} as a probe ● **TIMING 2.5 h**

63| Mix full-match 18-nt target 18C, dsDNA probe G-DNA^{gal} and RNase H in a PCR tube, as follows:

PROTOCOL

Reagents	Volume (μl)
Solution AB	8.0
Probe G-DNA ^{gal} (Step 62) (0.2 pmol μl^{-1})	1.0
Target 18C (1.8 pmol μl^{-1})	1.0
Tth RNase H (1 U μl^{-1})	1.0
Total volume	11

For negative control, add 1 μl water in place of target 18C.

▲ CRITICAL STEP Negative control without target is essential, as the target-on chemiluminescence intensity should always be evaluated in reference to that in the absence of the target.

▲ CRITICAL STEP It is critical to use the reconstituted translation system. A normal *E. coli* extract cannot be used, because it contains β -galactosidase.

64| Spin down and tap the mixture gently, and incubate for 1 h at 37 °C.

65| Mix the transcription/translation solution with 39 μl 1 \times reporter lysis buffer and 50 μl assay 2 \times buffer containing *o*-nitrophenyl β -galactoside of the β -galactosidase assay system.

66| Incubate the resulting solution in a 96-well plate for 1 h at 37 °C.

67| Read the absorbance at 405 nm with a multilabel counter (Wallac 1420), and take a photographic image of each sample.

? TROUBLESHOOTING

Troubleshooting advice can be found in **Table 1**.

TABLE 1 | Troubleshooting table.

Step	Problem	Possible reason	Solution
19, 21, 22, 24, 47, 67	Low signal intensity	Mis-handling of the translation medium	Handle the translation solution on ice and avoid vortexing. When mixing the solution, use gentle taps or pipetting
		Wrong translation medium possibly owing to damage during storage	Try a standard protein translation in the medium. If this does not work, use a new one
		Wrong MB-mRNA probe owing to mis-transcription or damage during storage	Check the purity by denaturing agarose gel electrophoresis (formaldehyde, MOPS). When the purity is not good enough, prepare a new MB-mRNA probe by using a freshly prepared and isolated template double-stranded DNA (dsDNA) for transcription
19, 21, 22, 24, 47, 67	Low on/off ratio ($I_{\text{on}}/I_{\text{off}}$)	Probe in high excess of the target. As the probe retains a target-independent translation activity, a large excess amount of the probe lowers the sensitivity ($I_{\text{on}}/I_{\text{off}}$)	Check the concentrations of target and probe carefully
22, 47	Low sequence selectivity	Poor RNase H activity	Check the RNase H activity. If it does not work, use a new one
		Contamination by MB-mRNA probes targeting different alleles	To prevent contamination of inadequate templates, use disposable filter tips for all PCRs and transcription reactions. Negative control without template is indispensable to check the contamination during the PCR step

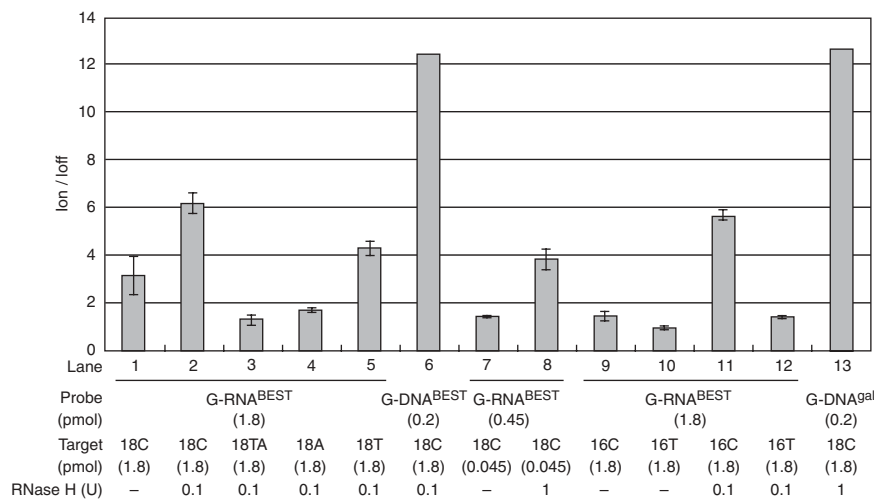


Figure 6 | Sensing of various targets with G-RNA^{BEST}, G-DNA^{BEST} or G-DNA^{gal} as a probe in the absence or presence of RNase H in 10–11 μl (lanes 1–6 and 9–13) or 2.5 μl (lanes 7 and 8) of a reconstituted translation medium after treatment with a luciferase or β -galactosidase assay solution. Data are averages of three or two independent experiments and error bars for triplicate runs represent s.d.

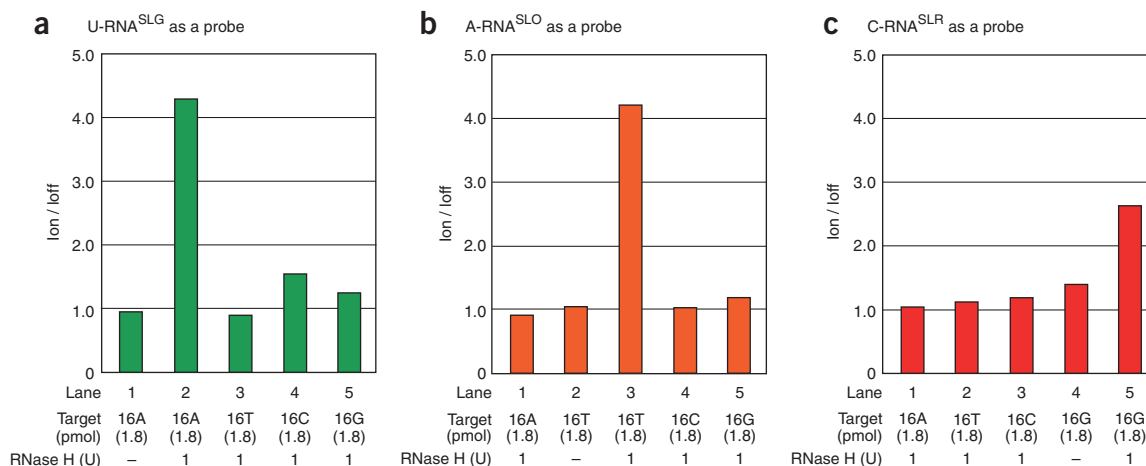


Figure 7 | RNase H-coupled sensing of targets 16A, 16T, 16C and 16G with (a) U-RNA^{SLG}, (b) A-RNA^{SLO} or (c) C-RNA^{SLR} as a probe in 11 μ l of a reconstituted translation medium after treatment with a luciferase assay solution.

ANTICIPATED RESULTS

Steps 19, 21, 22 and 24

As translation of reporter protein is not intended to be completely suppressed in the absence of target (see INTRODUCTION), it is most appropriate to refer to the on/off ratio of the chemiluminescence intensity (I_{on}) in the presence of a particular target to that (I_{off}) in its absence. Anticipated I_{on}/I_{off} ratios are summarized in **Figure 6**. In the absence of RNase H, target 18C can be sensed by the full-match probe G-RNA^{BEST} with $I_{on}/I_{off} = 3.1$ (lane 1). In the presence of RNase H, the sensitivity becomes higher ($I_{on}/I_{off} = 6.2$; lane 2) with good discrimination of doubly mutated (18TA; $I_{on}/I_{off} = 1.3$; lane 3) and singly A-mutated (18A; $I_{on}/I_{off} = 1.7$; lane 4) targets. This is, however, not the case for singly T-mutated target (18T; $I_{on}/I_{off} = 4.3$; lane 5), which would form a relatively stable GT mismatch in the target-probe heteroduplex. The on/off ratio for target 18C becomes even more pronounced when using G-DNA^{BEST} ($I_{on}/I_{off} = 12.5$; lane 6) in place of G-RNA^{BEST} ($I_{on}/I_{off} = 6.2$; lane 2). The advantage of the RNase H-coupled system can also be clearly demonstrated when using a smaller amount of target. Target 18C in 45 fmol in 2.5 μ l of the medium (18 nM) can scarcely be detected ($I_{on}/I_{off} = 1.4$; lane 7) in the absence of RNase H, but in its presence it is readily sensed, with $I_{on}/I_{off} = 3.8$ (lane 8), when a sensitive luminometer (Lumat LB 9507) is used. The detection limit with respect to full-match target 18C seems to lie in the range of 9 fmol ($I_{on}/I_{off} = 1.7$).

In the case of short 16-nt targets, even 16C ($I_{on}/I_{off} = 1.4$; lane 9), in addition to 16T ($I_{on}/I_{off} = 0.94$; lane 10), cannot be sensed effectively in the absence of RNase H. However, in the presence of the latter, the full-match C-allele target gives rise to a significant signal enhancement ($I_{on}/I_{off} = 5.7$; lane 11), whereas the T-mismatch remains inactive ($I_{on}/I_{off} = 1.4$; lane 12).

Steps 47–49

The chemiluminescence intensity (I_{on}) for any mismatch target-probe combination—that is, 16A/A-RNA^{SLO}, 16A/C-RNA^{SLR}, 16T/U-RNA^{SLG}, 16T/C-RNA^{SLR}, 16C/A-RNA^{SLO}, 16C/U-RNA^{SLG}, 16C/C-RNA^{SLR}, 16G/U-RNA^{SLG} and 16G/A-RNA^{SLO}—is hardly distinguishable from that (I_{off}) in the absence of target ($I_{on}/I_{off} \cong 1$ in most cases). On the other hand, matched target/probe combinations, that is, 16A/U-RNA^{SLG} and 16T/A-RNA^{SLO} ($I_{on}/I_{off} \cong 4$) and 16G/C-RNA^{SLR} ($I_{on}/I_{off} \cong 3$), give rise to higher signal intensities, as shown in **Figure 7**. Thus, any probe sensitively responds to its partner target on a complementarity basis.

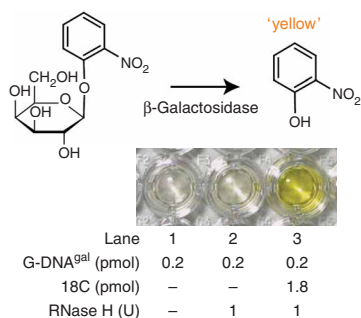


Figure 8 | RNase H-coupled sensing of target 18C with G-DNA^{gal} as a probe in 11 μ l of a reconstituted translation medium after treatment with a β -galactosidase assay solution. Reaction scheme and photographic images of the assay solutions.

Step 67

The anticipated target-on/target-off ratio of the absorbance at 405 nm with RNase H is $I_{on}/I_{off} = 12.6$ (**Fig. 6**, lane 13). o-nitrophenol formed can be easily detected even visually (**Fig. 8**).



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