

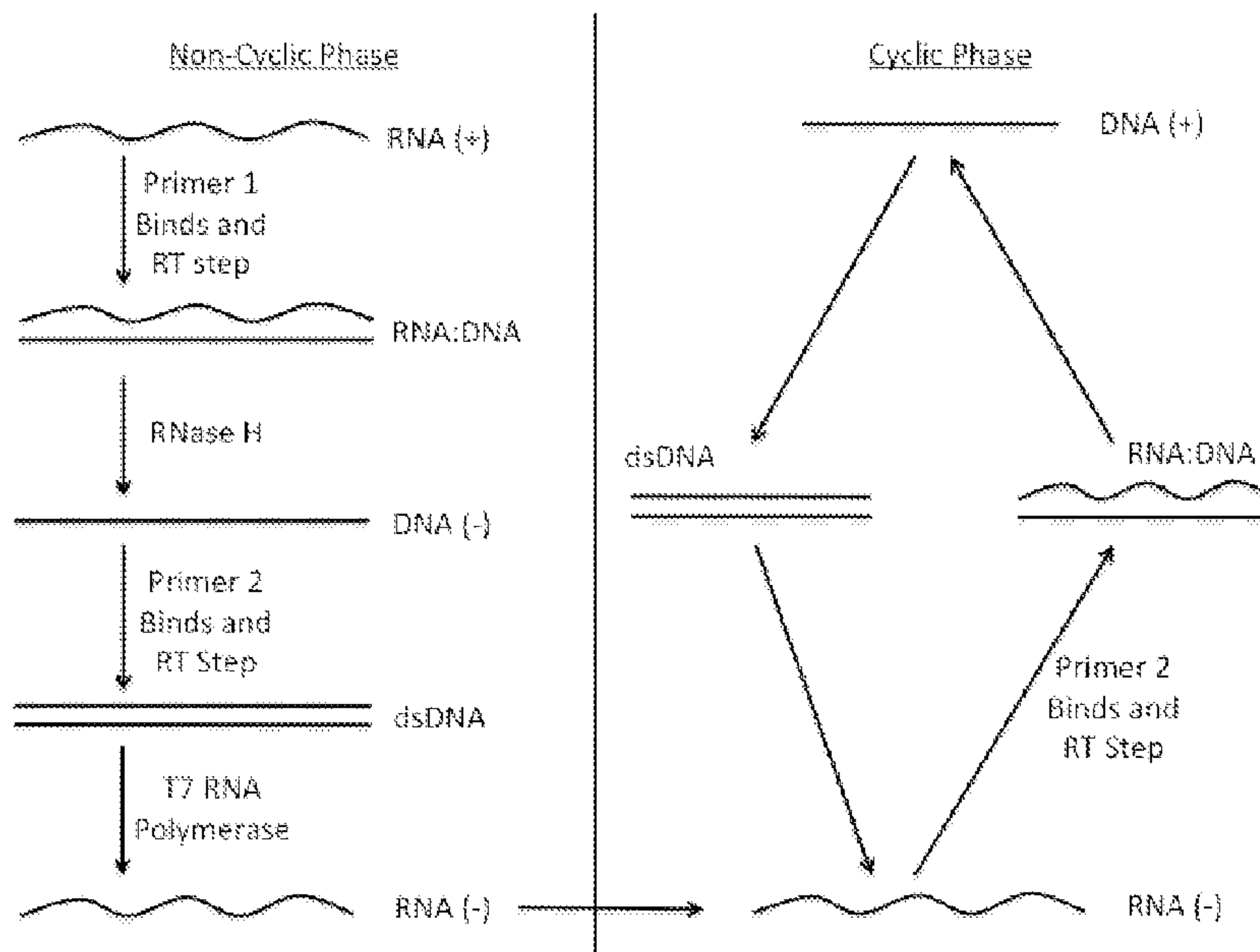


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Fig. 1



(57) Abrégé/Abstract:

An assay for detection of short sequences of RNA in a synthetic or clinically isolated sample is presented herein. Particular reference is made to detecting RNA based pathogens, such as H5 influenza.

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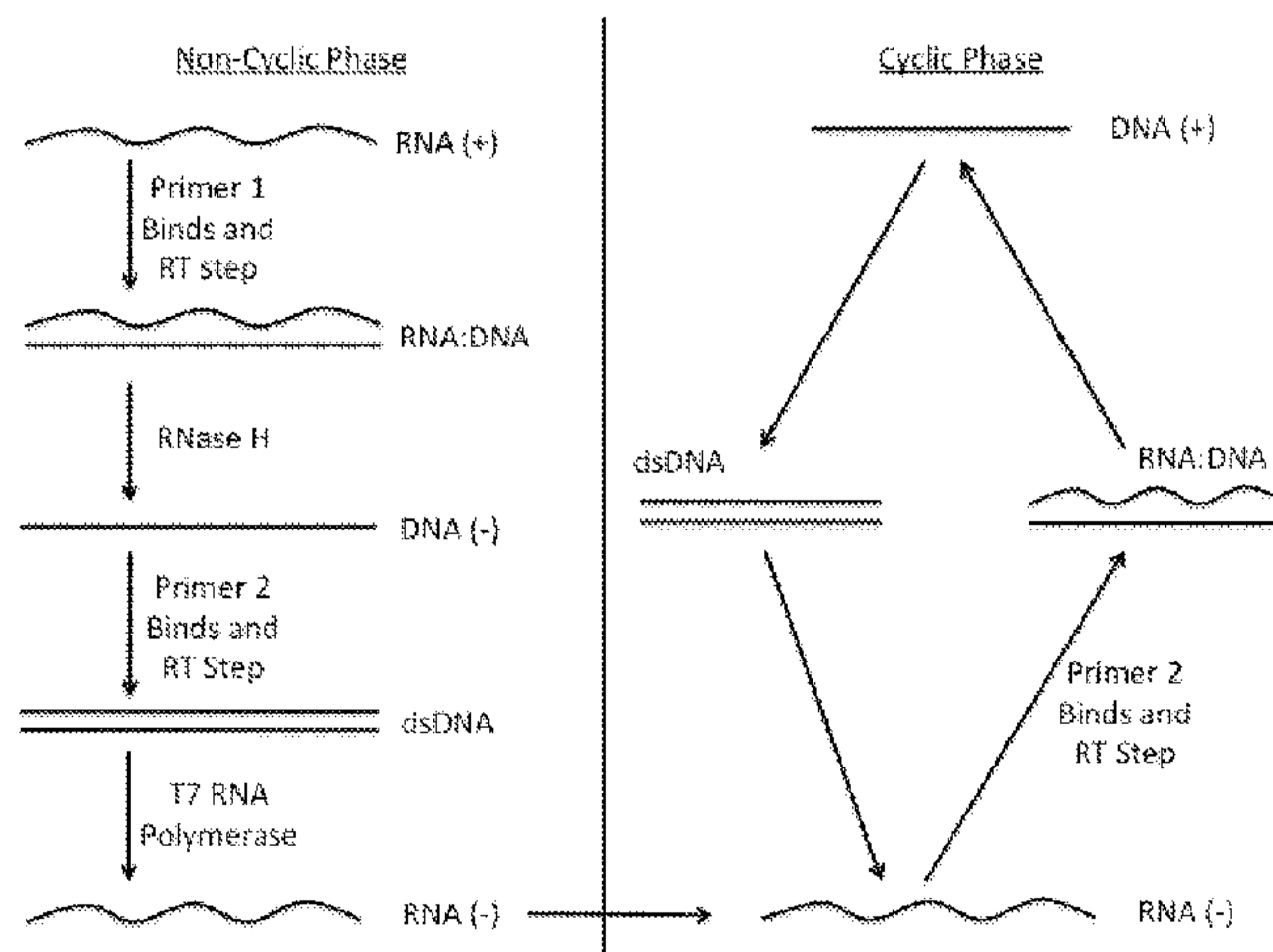
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(54) Title: DETECTION OF SHORT RNA SEQUENCES

Fig. 1



(57) Abstract: An assay for detection of short sequences of RNA in a synthetic or clinically isolated sample is presented herein. Particular reference is made to detecting RNA based pathogens, such as H5 influenza.

Detection of short RNA sequences

Sequence Listing

The instant application contains a Sequence Listing which has been submitted via EFS-
5 Web and is hereby incorporated by reference in its entirety. Said ASCII copy, created on August 2,
2010, is named B0197022.txt, and is 2,634 bytes in size.

Field of the Invention

Disclosed is an assay for detection of short sequences of RNA in a synthetic or clinically
10 isolated sample. Particular reference is made to detecting RNA based pathogens, focusing on H5
influenza.

Background of the Invention

A current assay used widely is Nucleic Acid Sequence-Based Amplification (NASBA).
15 A drawback to the NASBA technique is the secondary structure of target molecules. These can
either hinder or completely stop an amplification reaction.

A major problem in medicine is creating a sensitive, point of care device for RNA
pathogens. Faster and more efficient diagnosis would lead to more efficient treatment of
patients. Polymerase chain reaction (PCR) assays require precise equipment that is not
20 conducive to a point of care solution. Nucleic Acid Sequence-Based Amplification (NASBA)
may currently be the best choice for a point of care device as it involves using an isothermal
amplification step lasting only 90 minutes. NASBA, however, has previously not worked as
expected in our laboratory, and it may be due to secondary structure in the RNA that prevents
efficient primer binding and enzyme progression.

25 *Diagnosis of Influenza A H5N1*

Many techniques have been utilized in order to diagnose influenza. Historically, there
are two ways to confirm the presence of the virus: detection of a person's immune response to
the virus or detection of the virus itself. Four techniques that cover antibody detection, or the
immune response, are virus neutralization test (NT), hemagglutination inhibition (HI), enzyme
30 immunoassay (EIA), and complement fixation. These tests check for influenza antibodies in an
individual. These antibodies have their peak levels occur between four to seven weeks after

weeks before results are reached. Thus, there would be many problems to overcome if one were to try to utilize viral antibody detection [33].

Detection of the virus could be considered a more important approach for clinically relevant applications. There are three general approaches in this branch of detection:

5 immunospecific assays for viral antigen detection, viral isolation, and nucleic acid testing. The first group, immunospecific assays, encompasses two categories: rapid antigen tests and immunofluorescence microscopy [33].

Rapid antigen tests provide a very quick result and currently serve as one option for point of care detection. Commercial kits, such as Directigen Flu A and QuickVue influenza test, are
10 already on the market and ready to use [33]. Some kits are reported to detect subtype H5N1 virus [34]. These tests use specimens such as nasopharyngeal aspirates, nasopharyngeal swabs, and throat swabs. Many factors have an impact upon the sensitivity of the test. Reports suggest that sensitivity is useful at about two days after symptoms appear when viral shedding is maximal [33]. These tests, however, can have a wide range of specificity and sensitivity. The
15 range of reported sensitivity is between 39% and 100%, and the range for specificity is between 51% and 100%, varying with the kit as well as from where the sample was obtained. This varied range of sensitivity coupled with the lack of ability to subtype the different HA groups are the main drawbacks of this technique [35].

Immunofluorescence microscopy, which includes direct fluorescent antibody tests (DFA)
20 and immunofluorescent antibody tests (IFA), works by placing respiratory epithelial cells onto a slide and adding a series of specific antibodies. The slide is then viewed via fluorescence microscopy. These tests can give a result within about four hours. There are, however, some drawbacks in comparison to rapid antigen tests as a fluorescence microscope is needed, and a trained technician must carry out the test in order to perform the experiment as well as interpret
25 the results. Despite the drawbacks, its higher sensitivity than rapid antigen testing and ability to subtype make this technique a valuable asset to influenza diagnosis [33].

Viral isolation is used as it has a very high sensitivity level, down to about 10 pfu/mL (plaque forming units). Thus, the sensitivity for this assay is greater than the rapid antigen tests. Furthermore, it allows for laboratories to increase their stocks of virus for further studies [37]. In
30 addition, viral culture continues to be an important method in providing critical information about circulating strains and subtypes of influenza. In conventional test culture, a patient's

sample material is added to a cell culture. Then, the culture is monitored for signs of cytopathic effect. This alone, however, does not confirm that the culture is infected by influenza as the effect can be due to a number of viruses. Confirmation is typically performed via antibody staining and analyzing the culture with a fluorescence microscope. Disadvantages of this
5 strategy include the length of time to receive a result, which can take up to 14 days, the need for a specialized technician, the requirement of live, viable virus, and the need for highly certified laboratories (BSL-3) in cases where one is dealing with highly pathogenic strains of influenza [33].

It has been reported that low-speed centrifugation increases the viral infectivity of cells.
10 It is thought that this step disrupts cells and allows foreign viruses to enter more efficiently. In turn, this enhances the sensitivity of the culture, decreasing the time required for a diagnosis to between 18 and 48 hours. This technique may cause the virus to become nonviable. Thus, passaging cells becomes an issue, especially when a lab is utilizing cell culture to increase viral RNA for nucleic acid techniques.

15 A test directly detecting viruses is nucleic acid testing (NAT). In general, NAT works by specifically amplifying DNA or RNA or both in the presence of a specific sequence of nucleic acid. For influenza, typically amplification occurs in the presence of viral influenza RNA. NAT is considered more sensitive and specific than virus isolation, and in some cases it has replaced viral isolation as a reference standard. In addition, because nucleic acid is targeted and
20 amplified, both viable and nonviable virus can be used in an assay. These techniques also give investigators information about not only the subtype but also allows for sequence analysis that can be done after amplification. Results can be obtained within four to six hours. Two methods within this category that are used and researched widely are reverse transcription polymerase chain reaction (RT-PCR) and nucleic acid sequence based amplification (NASBA) [33].

25 PCR is a cyclic process consisting of three steps: denaturing, annealing, and extending. Denaturing the DNA involves separating the two strands. This usually involves heating the sample to 95°C. Heat denaturation of nucleic acids is reversible, unlike other methods such as chemical denaturation. The denaturation step allows the two primers to anneal to the DNA. Primers are short, single-stranded sequences of DNA that are reverse complementary to the DNA
30 strands to be amplified. This step occurs at a lower temperature, which varies depending on the DNA and primers, in order to allow for annealing to occur [39]. The last step, extension of

primers, utilizes the thermostable DNA polymerase *Thermus aquaticus*, also known as *Taq* polymerase. *Taq* polymerase extends the primers in a 5' to 3' direction. This step occurs at about 72°C, which is a higher temperature than the annealing step but a lower temperature than the denaturing step. The thermostable property of *Taq* is extremely important as it allows for the cyclic nature of the reaction to take place without needing to add reagents [40]. A typical cycle length is between three and five minutes, with a total of 20 to 40 cycles, thus the total length for PCR is usually a little over three hours [39].

A noted variation of PCR utilized for detecting RNA sequences is RT-PCR. Reverse transcription is an extra step needed before the PCR cycle because the starting material is viral RNA rather than DNA. Generally, an ssDNA primer hybridizes to a specific section of RNA and then is extended by an enzyme, such as avian myeloblastosis virus reverse transcriptase (AMV-RT) or Moloney murine leukemia virus reverse transcriptase (MMLV-RT). Another variation to RT-PCR that has been developed is multiplex PCR. In these assays, multiple primer sets are used either for detecting multiple genes of a single pathogen or subtype or for detecting multiple subtypes of influenza at the same time [41]. The former has been shown for H5N1 in a multiplex RT-PCR that, in a single tube, can detect the genes coding for M, H5, and N1. This could become useful in surveillance as mutations could be noted in a current strain [42]. The latter type of multiplex RT-PCR has been shown to be able to differentiate between H1, H3, and H5; N1 and N2; and virus types A and B [43].

Another method for amplification is NASBA. It is reported that NASBA will exponentially amplify targets without temperature cycling, between 37°C and 41°C. Thus, less sophisticated equipment is necessary [44, 45]. NASBA also generally requires fewer cycles than PCR to achieve the same amplification, which reduces the length of the reaction to between 1.5 and 2 hours [46]. Finally, NASBA has been utilized in detection of human papilloma virus (HPV) [47], HIV-1 [48], and influenza A virus of all HA subtypes [49].

Typically, NASBA requires three enzymes (a reverse transcriptase, an RNA polymerase that catalyzes the formation of RNA in the 5'→3' direction, and a non-specific endonuclease and catalyzes the cleavage of RNA. Examples of each are AMV-RT (also e.g., Moloney murine leukemia virus (MMLV-RT) and HIV-RT), T7 RNA polymerase (also, e.g., T3, and SP6 polymerases), and RNase H. NASBA also requires nucleoside triphosphates (both dNTPs and rNTPs), two DNA primers, and the correct buffer conditions. A diagram of the reaction is

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5 polymerases), and RNase H. NASBA also requires nucleoside triphosphates (both dNTPs and rNTPs), two DNA primers, and the correct buffer conditions. A diagram of the reaction is in Fig. 1. The first part runs non-cyclically. The target RNA, RNA (+), binds to primer 1, and AMV-RT extends the primer. It is important to note that primer 1 contains a promoter region for T7 RNA polymerase. RNase H, which selectively degrades RNA in RNA-DNA hybrids,
10 degrades the RNA, leaving the extended DNA, DNA (-). Primer 2 binds to the DNA and AMV-RT extends this primer, creating a double stranded DNA (dsDNA). T7 RNA polymerase then creates many copies of the negative RNA strand, RNA (-), from the dsDNA template. This triggers the cyclic phase. Primer 2 then binds RNA (-) and is extended by AMV-RT. RNase H degrades the RNA in the RNA-DNA hybrid, leaving DNA (+). Primer 1 then binds to DNA (+)
15 and AMV-RT extends the primer, creating dsDNA. T7 RNA polymerase completes the cycle by transcribing more RNA (-). Thus, the negative strand, RNA (-), is exponentially amplified. [46].
Noted also is locked nucleic acid (as well as other analogues such as TNA, GNA, and PNA). Locked nucleic acid (LNA), often referred to as inaccessible RNA, is a modified RNA
nucleotide. The ribose moiety of an LNA nucleotide is modified with an extra bridge connecting
20 the 2' oxygen and 4' carbon. The bridge "locks" the ribose in the 3'-endo (North) conformation, which is often found in the A-form of DNA or RNA. These nucleotides LNA, TNA, GNA, and PNA are collectively termed "Other Nucleotides."

Like PCR, there have been variations on NASBA developed to make this procedure more flexible. There have been some successful attempts at multiplex NASBA, where multiple
25 NASBA primers are used in the same reaction. One utilizes the method to amplify enteric viruses [50] while another has been used to detect hepatitis A virus and rotavirus simultaneously [51]. A protocol has also been reported that amplifies DNA using NASBA. Because the strands of DNA must be separated first, an initial denaturing step at 95°C is needed.

Fluorescence in detection is utilized with molecular beacons [54]. Molecular beacons are
30 single stranded DNA designed to be reverse complementary to itself on its two ends. This creates a hairpin shape with a stem region and a loop region. The ends are modified such that

Summary of the Invention

The method described is an extension of the Nucleic Acid Sequence Based Amplification (NASBA) protocol. The method disclosed is useful to exponentially amplify strands of RNA, complementary to at least a portion of a Target nucleotide sequence (*i.e.*, RNA, DNA, and Other Nucleotides) if the Target nucleotide sequence is present in a sample. "Sample" is used in reference to an aliquot, suspension, or fraction that contains the nucleotide (*e.g.*, RNA or DNA) under investigation.

The modified protocol in one embodiment employing ligation of contiguous probe fragments relies on four short DNA oligonucleotide sequences. The four sequences are:

ProbeLeft "5' - TargetRCLeft - HybridSeqRC -3'",

ProbeRight "5' - Primer2-TargetRCRight - 3'",

Primer1 "5' - T7 - HybridSeq - 3'", and

Primer2.

Fig. 2A shows a diagrammatic view of TargetRCRight and TargetRCLeft as hybridized to a sample RNA. Left and Right as used herein with reference to target sequences shall be understood to mean Left as to a nucleotide or nucleotide sequence positioned closer to the 5' end, while Right is used to indicate a nucleotide or nucleotide sequence positioned closer to the 3' end.

TargetRCLeft and TargetRCRight are short sequences that are reverse complementary (RC) to adjacent sequences in a sample being assayed. HybridSeq and Primer2 are designed to minimize the secondary structure of the RNA produced in this reaction, while maintaining a strong binding energy to their reverse complements.

The step of washing away Probe can be avoided with an additional reaction. ProbeLeft and ProbeRight are single-stranded DNA sequences designed such that when the 5'-end of ProbeRight is ligated to the 3'-end of ProbeL, the completed ligated sequence forms a sequence identical to the Probe sequence that would be used in the reaction. ProbeLeft and ProbeRight can be placed in a reaction mix containing a DNA ligase that ligates the fragments if and only if they are adjacent to each other while hybridized to a complementary DNA or RNA sequence. (Fig. 2A) To facilitate this juxtaposition, both ProbeLeft and ProbeRight usefully contain sequences that are reverse complements of the sequence being detected. Examples of ligases

include *E. coli* DNA ligase, *Taq* DNA ligase, and T4 DNA ligase. For clarity, hybridizing conditions are conditions that permit reannealing of nucleotides with their complementary bases.

ProbeLeft and ProbeRight are usefully added to a reaction mix containing the reagents described herein along with a ligase, as the ligation step may be carried out in the solution conditions described above and optimally at about the functional temperature for the ligase. A reaction time of about 10-90 minutes at such temperature (e.g., about 16-65°C, depending on the ligase). As a caution, it is noted that various enzymes are heat inactivated (e.g., T7 polymerase at about 41°C.) It will be understood if the reaction mix is heated above the inactivation point these heat labile enzymes should be added after the ligation step. Alternatively, Sample and ProbeLeft and ProbeRight can be reacted under the optimal reaction conditions for the ligase in a separate tube or container, and some of this reaction mix can be placed in place of the Sample in the previously described reaction conditions.

In one embodiment, the oligonucleotides are added to a Tris-buffered pH 8.3 solution containing MgCl₂, dithiothreitol, nucleoside triphosphates, deoxynucleoside triphosphates, DMSO, and isolated RNA. The solution is heated to 65°C for 5 minutes to disrupt the RNA secondary structure, and then cooled to 16°C. Once the solution has cooled, an enzyme mix containing AMV reverse transcriptase (AMV-RT), T7 RNA polymerase (T7Pol), RNaseH, and T4 DNA Ligase is added to the solution, and the solution is held at 16°C for 10 minutes. T4 DNA ligase forms a phosphodiester bond between adjacent DNA fragments that are hybridized to DNA or RNA strands. This ligates ProbeLeft and ProbeRight if, and only if, the Target sequence is present to make one combined oligonucleotide ("5' - Primer2-TargetRC-HybridSeqRC - 3'", "Probe"). The sequence is then exponentially amplified at 41°C in the following cycle of reactions. RNaseH selectively degrades RNA hybridized to DNA, so the Probe sequence is separated from the original RNA, which is destroyed. Primer 1 hybridizes with Probe (HybridSeq +HybridSeqRC). AMV-RT reads primed single stranded RNA and DNA sequences to synthesize DNA in the 5'→3' direction, and creates a double stranded DNA with one strand with the sequence 5,-T7-HybridSeq-TargetRC-Primer2RC-3'. The T7 region of the sequence is the highly conserved T7 polymerase promoter sequence. This acts as a recognition site for the T7 polymerase to begin in vitro transcription of an RNA sequence. Transcription is believed to occur by reading the DNA strand that the T7 promoter is hybridized to and transcribing the RNA sequence complementary to the DNA strand. In the case of double-

stranded DNA templates, this means the RNA strand has the same sequence as the original top strand of DNA. The polymerase does not copy its own promoter sequence, so the polymerase will make the following RNA strand "5'- HybridSeq-TargetRC-Primer2RC-3'". Since the transcription is not primer-dependent, the polymerase makes multiple copies of the RNA for each template that is made. The RNA sequences are templates for reverse transcription by AMV-RT after priming by Primer2, which makes a DNA strand with the sequence 5' - Primer2- Probe-HybridSeqRC-3', which is the Probe sequence. This exponentially amplifies RNA as a growing number of dsDNA templates are created. The progression of this reaction is monitored with molecular beacons, which are short DNA sequences that form a hairpin structure. A fluorophore and a fluorescence quencher are on the ends of the beacon sequence. As a result, fluorescence is not observable when the hairpin is in its closed state, but is observed when the sequence is open. The loop of the sequence is complementary to a target RNA or DNA sequence, while the 5'-end of the sequence is complementary to the 3'-end. The sequence is designed so that the binding energy of the loop to its complement is slightly greater than the binding energy of the stem to itself. This creates a very specific and sensitive reporter for the complement to the loop sequence. In this reaction, the beacon is designed to detect the presence of Primer2RC. Primer2RC is produced when the reaction is successful. A non-sequence specific method of assaying RNA synthesis (like RiboGreen) may also be used to evaluate the progression of the reaction.

This protocol detects multiple target sequences (such as different genotypes associated with the same clinically relevant phenotype) with the same molecular beacon by varying the Target sequence while keeping the Primer2 and HybridSeq sequences unchanged. This is believed to provide more favorable amplification thermodynamics than typical multiplex NASBA reactions, which are less efficient due to the number of primers present. In addition, the use of a single beacon lowers costs and simplifies detection.

This invention comprises A method of amplify Target RNA comprising the steps of introducing at least one Target RNA (*e.g.*, H5 influenza) to a sample containing probe nucleotide under hybridizing conditions;

wherein said Target RNA comprises three regions,

said first region being a Hybrid Seq RC region; and,

a second region, being a Target RC regions contiguous with said first region; and

a third region being a Primer 2 region contiguous with said second region; and, selectively amplifying the RNA of said Target RNA, and optionally detecting and/or quantifying the amplified RNA. In some embodiments detection and/or quantification is by gel electrophoresis or by fluorescence. Particularly noted is detection and/or quantification using molecular beacons. Multiple molecular beacons are useful to detect and/or quantify multiple Target RNAs.

In one embodiment of the method the amplifying of step (ii) comprises transcribing hybridized Target RNA into double-stranded DNA. The instant method also usefully employs a primer containing a T7 promoter sequence which binds to the probe RNA. The instant method further employs transcribing it by means of a reverse transcriptase such as AMV-RT. AMV-RT is useful to transcribe a reverse complementary DNA sequence which creates a DNA-RNA hybrid. Noted is an embodiment including transcribing said resulting double stranded DNA into RNA. One noted method of transcribing is by means of an RNA polymerase that catalyzes the formation of RNA in the 5'→3' direction. An aspect of the method further includes only the RNA strand of a DNA-RNA hybrid being degraded, optionally by RNase H. The resulting DNA strand is then primed with a primer at the 3' end and the transcription process is repeated with AMV-RT. Transcription produces a double stranded DNA (a template) from which T7 RNA polymerase transcribes an RNA.

In the claimed method the Hybrid Seq RC region, Target RC region and said Primer 2 region each comprise from about 8 to about 35 bases which number as to each region may be the same or different. In addition and optionally within the method a molecular beacon may be hybridized to the amplified RNA.

The invention further includes a method of detecting target nucleotide comprising

- (i) exposing ProbeLeft nucleotides and ProbeRight nucleotides under hybridizing conditions to a Probe-specific ligase;
- (ii) permitting ligating of said ProbeLeft and ProbeRight nucleotides if and only if they are adjacent to each other while said ProbeLeft and ProbeRight nucleotides are hybridized to a complementary nucleotide sequence; and,
- (iii) detecting and/or quantifying the presence or absence of said ligated ProbeLeft with ProbeRight nucleotide.

Brief Description of the Drawings

Fig. 1. Fig. 1 is a schematic of both the non-cyclic and cyclic phases of the NASBA reaction. Wavy lines represent RNA and straight lines represent DNA.

Fig. 2. Fig. 2 is a schematic of the capture step to the SMART assay.

5 Fig. 2A shows a diagrammatic view of TargetRCRight and Target RCLeft as hybridized to a sample RNA.

Fig. 3. Fig. 3. is a schematic of the SMART Probe with the variable arms on either side of the center portion.

Fig. 4. Fig. 4 is a schematic of the amplification step to the SMART assay.

10 Fig. 5. Fig 5 is a diagram showing where the molecular beacon binds in the SMART assay for real-time detection.

Fig. 6. Fig. 6 is a picture of a gel from Gel electrophoresis of the initial testing of the amplification step.

15 Fig. 7. Fig. 7 is a gel from an experiment varying DMSO concentration. Four conditions of different concentrations of probe, 100 nM, 10 nM, 1 nM, and 0 nM, were run for each DMSO concentration, 5% and 15%. The concentrations of probe go from left to right with each DMSO condition group.

20 Fig. 8. is a gel plot of an experiment varying Tris-HCl pH. Four conditions of different concentrations of probe, 100 nM, 10 nM, 1 nM, and 0 nM, were run for each pH condition, 8.0 and 8.3. The concentrations of probe go from left to right with each pH group.

Fig. 9 is a gel plot of electrophoresis of various nucleic acids in the SMART Assay using the Small RNA Assay.

Fig. 10. Fig. 10 is an electropherogram plot of a mix of Primers and Probe, cDNA, and cDNA with T7 RNA polymerase.

25 Fig. 11 is an electropherogram plot of three reactions. The first reaction (light gray) shows a reaction with AMV-RT only. The second line (dark gray) shows the reaction with AMV-RT and T7. The final line (black) represents the reaction involving all three enzymes.

Fig. 12. is a plot of relative fluorescence versus time for many dilutions of the Probe from Set 1.

30 Fig 13 is a design of a chip for SMART assay utilizing electrophoresis detection. Red areas denote a heating implement for a constant temperature.

Detailed Description of the Invention

The assay presented here, coined a Simple Method to Amplify RNA Targets (SMART), involves amplifying a probe engineered for improved binding to primers as well as minimizing the secondary structure of nucleic acids to be amplified. In some embodiments, this technique utilizes isothermal, cyclic amplification. These parameters are useful in point of care settings as such conditions minimize the equipment needs. In particular embodiments, the disclosed assay is employed in a microfluidic chip platform.

10 The work presented here particularly notes H5 influenza as a target. In addition and without limitation, the method is useful with any RNA based pathogen.

In particular embodiments, the SMART assay binds engineered ssDNA probes to an RNA target. Then the engineered probes are selectively amplified rather than the target RNA itself being amplified as is typical in the NASBA protocol. Selectively amplifying shall be understood to mean that at least about 80% or more and preferably 90% or more (by base count) of the RNA synthesized in this cycle of reactions is Target RNA and contiguous regions (*i.e.*, first region and third region) or fragments thereof.

One advantage of these probes is that they are short and can be engineered to have minimal secondary structure. This is distinct from the sample nucleotide. Detection of the amplified nucleic acid is usefully performed by any method but note is made of two standard methods: gel electrophoresis or fluorescence via molecular beacons.

This invention will be better understood with resort to the following definitions:

A. "Short" as to RNA sequences, shall mean from about 8 to about 35 bases (also termed nucleotides or nt). with particular reference to about 10 or about 18 to 25 nucleotides. This term is particularly directed to the TargetRC portion of the SMART probe.

B. SMART probe is an oligonucleotide as shown diagrammatically in Fig. 3. having variable arms on either side of the center portion ("Target RC"). The variable arms vary from about 5, or about 10 or about 18 to 25 or 35 nucleotides each. One said variable arm is termed Hybrid Seq. RC and one is termed Primer 2. As to such variable arms, it is understood and contemplated that the sequences used for hybridizing Primer 1 or Primer 2 to the Probe sequence may include part of the sequence being tested for in the Probe sequence.

C. Hybrid Seq. RC shall mean a sequence on the 3' end of the Probe sequence that is sufficiently complementary to the 3' end of Primer 1 to hybridize under reaction conditions.

D. Primer 1 shall mean a primer comprising a 5' promoter region for an RNA polymerase that catalyzes the formation of RNA in the 5'→3' direction (*e.g.*, T7 RNA polymerase) and a HybridSeq on the 3' end that hybridizes to the 3' end of the Probe sequence to enable the creation of a double-stranded DNA sequence by a DNA-dependent DNA polymerase (*e.g.*, AMV RT) to be used as template for a DNA-dependent RNA polymerase.

E. Primer 2 shall mean an oligonucleotide sequence complementary to the 3' end of RNA transcribed from the DNA template synthesized in "D", which will hybridize to this sequence to permit the transcription of a DNA:RNA hybrid catalyzed by an RNA-dependent DNA polymerase (such as AMV-RT).

F. Target RC shall mean a short oligonucleotide sequence that is reverse complementary (RC) to a sequence on a Target RNA sequence.

G. ProbeLeft and ProbeRight shall mean single-stranded DNA sequences designed such that when the 5'-end of ProbeRight is ligated to the 3'-end of ProbeLeft, the completed ligated sequence forms a sequence identical to the Probe sequence that would be used in the reaction. The point of ligation shall be within the region described as TargetRC. It is noted that in some embodiments of the instant invention the step of using magnetic-streptavidin coated beads bound to biotinylated capture probes a useful first step where some method of hybridizing the probes to the target RNA and washing away the unhybridized probes is required. Beads are one such method. Other useful methods include capturing the target RNA on another surface, like a microfluidic channel.

Two SMART assay sets for H5 influenza were tested. Set 1 was more efficient than Set 2. Set 1 had more favorable Primer 1 dimer formation (-8.5 kcal/mol for Set 1 vs. -4.2 kcal/mol for Set 2) while Set 2 had more favorable RNA self-binding (-4.2 kcal/mol for Set 1 vs. -5.2 kcal/mol for Set 2). Without being bound by any particular theory, it is believed that this indicates that secondary structure of binding sites has a greater effect on reaction efficiency than primer-primer interactions. In addition, it appeared that the optimal Tris-HCl buffer pH was about 8.0. This is lower than many literature sources for NASBA. Similarly, it appeared that optimal DMSO concentration was about 15% v/v.

Using gel electrophoresis, it was confirmed that the expected products were produced. Molecular beacons were shown to be a viable option for detection at concentrations in the femtomolar range.

5 The SMART assay is an improvement over nucleic acid methods for detection and permits a point of care device. NASBA is isothermal and typically runs at a relatively low temperature (41°C) and reportedly down to about 37°C. Also noted are available heat-stable equivalents for all the enzymes in the reaction which function up to about 70°C. Indeed, in some
10 embodiments, exact temperatures are not necessary to NASBA as compared to PCR [44, 45]. In addition, the amplification step for NASBA lasts for 90 minutes. Experiments have shown that it is difficult to find NASBA primers that yield a good result. Without being bound by any particular theory it is believed that the complex conformation that an RNA strand can take, especially in a sequence as long as influenza which is approximately 1700 bases, makes it difficult for primers to consistently bind at some places.

15 The characteristics of a short amplification time and an isothermal reaction are useful for a point of care device. To overcome the problem of NASBA assays that are ineffective, the instant method amplifies an engineered probe rather than the RNA target itself. This probe binds to the RNA targets, if present, and will be substantially washed away otherwise.

20 In a specific embodiment, magnetic, streptavidin-coated beads are bound to biotinylated capture probes via the streptavidin-biotin bond. The capture probe is reverse complementary to the RNA target in one region. The engineered NASBA probe is reverse complementary to another region, which is selected to be a more favorable binding site. With a series of washes, the unbound probe is substantially washed away and thus will have limited amplification. The separation step is presented in Fig. 2.

25 In Fig. 2, RNA is shown by a wavy line, and DNA is shown by a straight line. Streptavidin-coated magnetic beads are added to a solution containing biotinylated capture oligonucleotides, or capture probes. These capture probes are reverse complementary to the target RNA strand.

30 In a specific embodiment of the invention, the engineered NASBA type probe has two, 25 nucleotide long arms on each end with a middle section that is the reverse complementary

part to the RNA of interest. The middle section, which can vary from about 5, or about 10 or about 18 to 25 or 35 nucleotides due to the specificity and binding efficiency desired, is small in comparison to the two, variable arms. The arms can also vary about 5, or about 10 or about 18 to 25 or 35 nucleotides each. Engineering the length takes into account three concepts. If the segment is overlong it may fold onto itself thus slowing down reactions. Its design also reflects a length to have binding energy sufficient to stay bound through various washing steps, but not so long so as to unduly degrade binding specificity.

Thus, arm length is selected to improve/permit sufficient amplification speed. They are optimized for efficient binding in order to push the reaction forward. Furthermore, they are optimized to reduce its own secondary structure, which addresses one the concerns about traditional NASBA methods. A schematic is shown as Fig. 3.

In Fig. 3, one variable end has the same sequence as Primer 2. The other variable end is called "Hybrid Seq RC" as it is reverse complementary to a segment of Primer 1. Useful aspects of these regions are further delineated in the amplification steps.

The design of the amplification step is similar to the NASBA reaction. However, in distinction, the starting target is an engineered probe strand rather than the positive strand RNA. In some instances, the same enzymes as NASBA are used. AMV-RT extends DNA primers that are hybridized to either DNA or RNA segments. T7 RNA polymerase transcribes double stranded DNA downstream from a specific promoter site. RNase H selectively degrades the RNA in an RNA-DNA hybrid [46]. A design of the SMART version of the reaction is shown in Fig. 4.

As seen in Fig. 4, during the first step of the reaction, Primer 1 binds to the Probe, and AMV-RT turns this hybrid into double stranded DNA. In this example, Primer 1 contains the T7 promoter sequence. The double stranded DNA serves as the template for T7 RNA polymerase, and the enzyme transcribes multiple strands of RNA. The RNA created has a different sequence than the original RNA in that region except for the segment in the center. Primer 2 then binds to the RNA strand and is extended by another AMV-RT step, creating a DNA-RNA hybrid. The RNA strand of this hybrid is then degraded by RNase H, yielding another Probe DNA strand and creating a cyclic exponential reaction.

Real-time detection is achieved with molecular beacons or another sequence-specific fluorescent probe. As an example of a molecular beacon, one is created to bind at the Primer 2

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and correspond to the ends of the probe, the primers, and the stem of the beacon. Lowercase letters are dependent on the target sequence of the probe,

Table 1: Self-binding thermodynamics for single stranded nucleic acids in the SMART assay.

Identification	Sequence (5' to 3')	Length	dG (kcal/mol)	T _m (°C)
<i>Set 1</i>				
Probe	TCAAGAGTAGACACAGGATCAGCATaggcaatagatggagtcacGTAATCAGATCAGAGCAATAGGTCA	69	-0.6	49.2
Primer 1	TAATACGACTCACTATAGGTGACCTATTGCTCTGATCTGATTAC	44	-1.4	56.4
RNA made	GGUGACCUAUUGCUCUGAUCUGAUUACgugacuccaucuauugccuAUGCUGAUCCUGUGUCUACUCUUGA	71	-4.2	50.6
Beacon 1	/56-FAM/CGCGtcaagagtagacacaggatcCGCG/3IABlk_FQ/	28	-1.4	54.5
<i>Set 2</i>				
Probe	TCAAGAGTAGACACAGGATCAGCATaggcaatagatggagtcacAGGCATATAGAGAGTCAGACAGGAG	69	-0.6	49.2
Primer 1	TAATACGACTCACTATAGGCTCCTGTCTGACTCTCTATATGCCT	44	-1.0	50.5
RNA made	GGCUCCUGUCUGACUCUCUAUAUGCCUgugacuccaucuauugccuAUGCUGAUCCUGUGUCUACUCUUGA	71	-5.2	50.3
Beacon 2	/56-FAM/CGTCGtcaagagtagacacaggatcaCGACG/3IABlk_FQ/	31	-2.7	60.1
<i>Both Sets</i>				
Primer 2	TCAAGAGTAGACACAGGATCAGCAT	25	0.7	30.3

- 5 Table 2 characterizes the binding energies between the beacons and other single strands to ensure that the beacon would substantially bind to the RNA target. The data for both sets are shown with the far left column as the strand binding to the beacon. Note that the last line is the beacon binding to another strand of beacon and not itself, which is shown in Table 1.

10 Table 2: Binding energies between the molecular beacon and other single strand nucleic acids in the assay. The data for Set 1 are shown with Beacon 1, and the data for Set 2 are shown for Beacon 2.

	Set 1		Set 2	
Identification	dG (kcal/mol)	T _m (°C)	dG (kcal/mol)	T _m (°C)

and correspond to the ends of the probe, the primers, and the stem of the beacon. Lowercase letters are dependent on the target sequence of the probe,

Table 1: Self-binding thermodynamics for single stranded nucleic acids in the SMART assay.

Identification	Sequence (5' to 3')	Length	dG (kcal/mol)	T _m (°C)
<i>Set 1</i>				
Probe	TCAAGAGTAGACACAGGATCAGCATaggcaatagatggagt cacGTAATCAGATCAGAGCAATAGGTCA (SEQ ID NO: 1)	69	-0.6	49.2
Primer 1	TAATACGACTCACTATAGGTGACCTATTGCTCTGAT CTGATTAC (SEQ ID NO: 2)	44	-1.4	56.4
RNA made	GGUGACCUAUUGCUCUGAUCUGAUUACgugacuccauc uauugccuAUGCUGAUCCUGUGUCUACUCUUGA (SEQ ID NO: 3)	71	-4.2	50.6
Beacon 1	/56-FAM/CGCGtcaagagtagacacaggatcCGCG/3IABlk_FQ/ (SEQ ID NO: 4)	28	-1.4	54.5
<i>Set 2</i>				
Probe	TCAAGAGTAGACACAGGATCAGCATaggcaatagatggagt cacAGGCATATAGAGAGTCAGACAGGAG (SEQ ID NO: 5)	69	-0.6	49.2
Primer 1	TAATACGACTCACTATAGGCTCCTGTCTGACTCTCT ATATGCCT (SEQ ID NO: 6)	44	-1.0	50.5
RNA made	GGCUCCUGUCUGACUCUCUAUAUGCCUgugacuccauc uauugccuAUGCUGAUCCUGUGUCUACUCUUGA (SEQ ID NO: 7)	71	-5.2	50.3
Beacon 2	/56-FAM/CGTCGtcaagagtagacacaggatcaCGACG /3IABlk_FQ/ (SEQ ID NO: 8)	31	-2.7	60.1
<i>Both Sets</i>				
Primer 2	TCAAGAGTAGACACAGGATCAGCAT (SEQ ID NO: 9)	25	0.7	30.3

- 5 Table 2 characterizes the binding energies between the beacons and other single strands to ensure that the beacon would substantially bind to the RNA target. The data for both sets are shown with the far left column as the strand binding to the beacon. Note that the last line is the beacon binding to another strand of beacon and not itself, which is shown in Table 1.

thermal cycler (MyCycler, Bio-Rad). The reaction mix and nucleic acid mix are added together and heated to 65°C for 5 min, and then the temperature is lowered to 41°C for 5 minutes to allow the temperature to reach the target temperature. 5 µL of enzyme mix is prepared with 1.3 U/µL AMV-RT, 0.5 U/µL RNase H, 6.4 U/µL T7 RNA polymerase, and 0.42 µg/µL bovine serum albumin (BSA) (Promega and NEB). The enzyme mix is added after the 5-minute equilibration period, and the reaction is carried out for 90 min at 41°C. The reaction is stopped by freezing the samples.

Gel electrophoresis is the first method used to analyze results. Two assays, the RNA Nano 6000 and Small RNA Assay, are both used on the Bioanalyzer 2100 (Agilent), a microfluidic chip platform for electrophoresis. The RNA 6000 assay is used as a qualitative assessment to determine whether amplification occurred, while the Small RNA assay is used to more precisely evaluate the size of RNA and DNA fragments generated during the reaction. Manufacturer's (Agilent) directions are followed as in the protocol for each assay, except that the samples used for the Small RNA Assay are diluted 1:5 because the assay is extremely sensitive to salt concentrations. The results are visualized in both a standard gel plot as well as electropherogram plots, generated from the program provided by Agilent for the Bioanalyzer 2100.

Performing the Assay in a Fluorometer and Real-Time Detection

The concentrations for the reaction mix, nucleic acid mix, and enzyme mix are the same as above except that molecular beacon is exchanged for some of the water in the reaction mix to give a 100 nM beacon concentration. The volumes of the mixes are increased to 40 µL of reaction mix, 20 µL of nucleic acid mix, and 20 µL of enzyme mix. The reaction mix and nucleic acid mix are first heated to 65°C for 5 minutes and then cooled to 41°C for 5 minutes in a thermal cycler. During this period, a new, disposable cuvette (UVette, Eppendorf) is placed into a fluorometer with a temperature control for each reaction. The temperature is set to 41°C, so that the cuvette itself can be equilibrated to the reaction temperature as much as possible. At the end of this period, the enzyme mix is added to the reaction. The cuvette is then removed from the holder, and the total reaction mix is transferred into the cuvette, which is inspected to ensure that no bubbles appear. The cuvette is placed back into the fluorometer, and the reaction is run for 90 minutes.

The fluorometer is set to an absorbance of 494 nm and an emission of 518 nm. This data is the peak absorbance and emission for the fluorescent molecule, 6-FAM [58]. The fluorometer is set to acquire data every 5 minutes, starting at time 0 minutes. Five data points are taken for each time point. The data from the first time point, 0 minutes, is not presented in the final results as it was found that there is a spike in fluorescence between the first two time points, probably due to the mixes and cuvette equilibrating to 41°C. The reaction is run for 90 minutes. The data are plotted using a MATLAB script to average the 5 data points for each time point, remove the 0-minute time point, scale all time points to the 5-minute time point, and subtracting 1 so that the first time point is 0. Data are plotted as relative fluorescence against time.

10 *Quantification and Modeling of SMART amplification and real-time detection*

A model for the amplification step and real-time detection using molecular beacons was created using MATLAB (The Mathworks, Inc.) and Mathematica (Wolfram Research, Inc.) in order to iterate through the steps of the reaction to perform the calculations. The data is presented with the following notations:

15 Pr is Probe,

P1 is Primer 1,

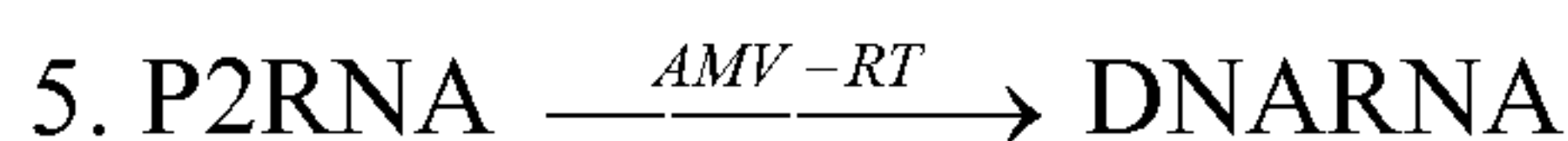
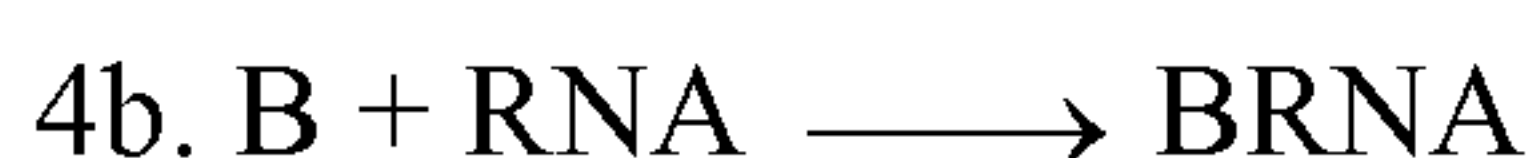
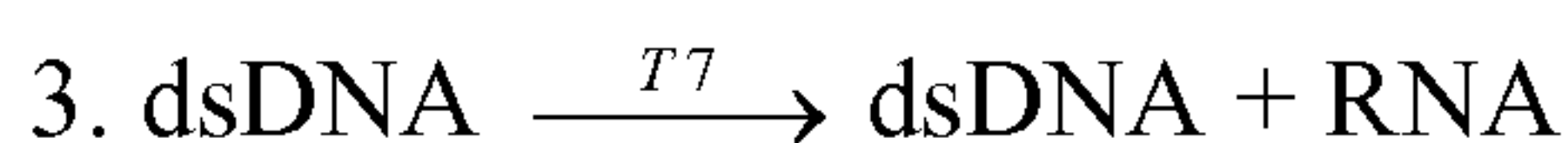
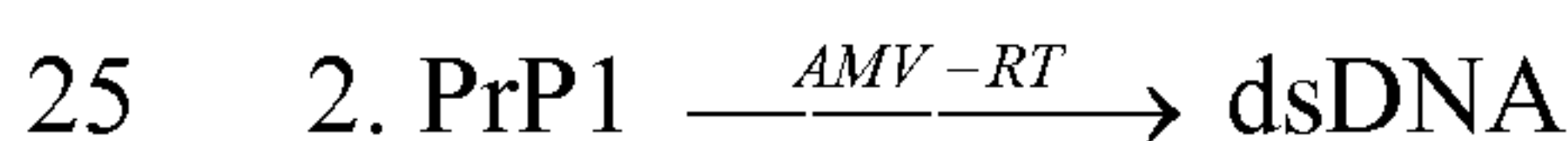
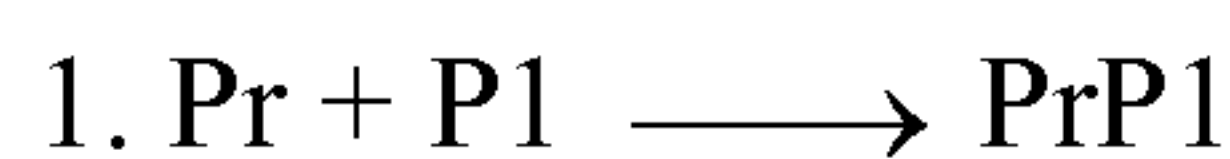
P2 is Primer 2,

B is beacon,

RNA is the RNA made, and

20 dsDNA is the double-stranded DNA that serves as the template for T7 RNA polymerase.

A complex of two of these species is denoted by placing the shorthand names together (i.e. PrP1 stands for Probe and Primer 1 bound together). The reaction was separated into 6 different steps:



All other events are considered to be negligible. In general, there were two types of steps: binding steps and enzymatic steps. The binding steps (1 and 4) can be described in similar ways. The equation for step 1 is shown below, where x is the change in concentration of Pr or P1 during a small time step, and k_1 is the equilibrium constant for step 1 (Eq. 1):

$$\frac{[PrP1] + x}{([Pr] - x)([P1] - x)} = k_1 \quad (1)$$

5 After finding x , the concentrations of the species are adjusted by the concentration x . Step 4 involves competitive binding of Primer 2 and Beacon to the RNA molecules. It differs from step 1 as two equations must be solved simultaneously. Equations 2 and 3 describe these binding events, where x is the decrease in concentration of Primer 2 or the increase of the complex Primer 2-RNA, and y is the decrease in concentration of Beacon or the increase of the complex
10 Beacon-RNA, and the sum of x and y represents the decrease in concentration of RNA.

$$\frac{[P2RNA] + x}{([P2] - x)([RNA] - x - y)} = k_{4a} \quad (2)$$

$$\frac{[BRNA] + x}{([B] - x)([RNA] - x - y)} = k_{4b} \quad (3)$$

The enzymatic steps are all described in similar ways as well, using the Michaelis-Menten equation. One assumption is that the two enzymes that polymerize nucleic acids, AMV-RT and T7 RNA polymerase, are saturated with NTPs through the reaction. Calculations show that this is true during the period of beacon binding. For rNTPs, a base count on the RNA
15 created in the reaction shows that U is the most used base. Since no RNA is in the sample in the beginning, the following is noted:

$$\frac{1RNA}{27UTP} \cdot 2mM UTP = 0.0741mM \cdot RNA = 74.1\mu M \cdot RNA \quad (4)$$

Thus, 74.1 μ M of RNA is made. This is on the order of 10^3 greater than the beacon concentration (50 nM), so rNTPs are not limiting. Again, without being A similar analysis can be done for dNTPs. For a conservative estimate, assume that the entire Probe begins as Primer 2,
20 and thus the double stranded DNA starts from Primer 2 and Primer 1. A base count shows that T is the limiting base for the reaction. Thus, a similar equation can be used as above:

$$1mM \cdot TTP \frac{1dsDNA}{31TTP} = 0.0323mM \cdot dsDNA = 32.3\mu M \cdot dsDNA \quad (5)$$

By this calculation a 32.3 μM of DNA is created. Since multiple strands of RNA can be made from one template of DNA, it is probable that rNTPs will be used in polymerization reactions before dNTPs, so dNTPs are not believed limiting in comparison to the beacon.

5 Since the enzymes are saturated with NTPs, the Michaelis-Menten equation collapses into the following form [59]:

$$V = \frac{V_{\max}[S]}{K_M + [S]} = \frac{k_{cat}[E][S]}{K_M + [S]} \quad (6)$$

where V_{\max} and K_M are the Michaelis-constants, k_{cat} is the general rate constant for a reaction or the limiting step of it, $[E]$ is the enzyme concentration, and $[S]$ is the substrate concentration. Given the assumption above, the substrate is treated as the nucleic acid to which the enzyme binds rather than the NTPs. For each iteration, V is multiplied by the iteration time step so that
10 the actual change in concentration of the substrate can be calculated for an iteration. From this, the concentration of the substrate and product involved in the enzymatic reaction is changed by the amount calculated for the iteration. Equation 6 is the general equation used for all of the enzyme steps (2, 3, 5 and 6).

The main assumption for the Michaelis-Menten equation is the steady-state assumption,
15 which states that $[ES]$, or the concentration of the enzyme bound to the substrate, does not change with time [60]. In the reaction as a whole, this is not true. Since the reaction is carried out iteratively, this statement can be treated as approximately true for each iteration, and thus the velocity of the reaction will change with time.

To solve the equation, k_{cat} , K_M , and $[E]$ are needed. For each enzyme, values were found
20 in literature for k_{cat} and K_M , and in some cases k_{cat} is found by using the relation that $k_{cat} = V_{\max}/[E]$. Since this reaction is different from other reaction conditions in literature, the final values used were estimated. For AMV-RT, it has been shown that the initiation step is the rate limiting step [61]. Therefore, the values for initiation were used. These values vary depending on the identity of the end base pair. For this reason an average was taken, with the value of k_{cat}
25 as $5.625 \times 10^5 \text{ s}^{-1}$ and K_M as $8.5 \times 10^{-8} \text{ M}$ [62]. To estimate k_{cat} for T7 RNA polymerase, Arnold *et al.* provided values for k_{eff} , which describes the amount of nucleotides transcribed per time while taking initiation, elongation, and termination into account. According to Arnold, the rate constant can vary between about 5 s^{-1} and 630 s^{-1} for the plasmids they used. The RNA created in the assay is much shorter than a plasmid. As such it is believed that initiation and termination

play a larger role in those examples. Considering that these are more likely to be the limiting factor, a small rate constant is chosen, 5 s^{-1} , which is divided by 71, the length of the RNA made, to retrieve the final k_{cat} , which describe the number of molecules made per second. Thus, the value chosen for k_{cat} was $7.04 * 10^{-2} \text{ s}^{-1}$, and K_M was used as in the paper, $6.3 * 10^{-9} \text{ M}$ [59]. For RNase H, many values were provided depending on the bases present. As with, AMV-RT, the average of these values were taken. The value for k_{cat} used was $1.6 * 10^{-1} \text{ s}^{-1}$ and the value for K_M was $7.32 * 10^{-8} \text{ M}$. The values for k_{cat} and K_M used in the model are summarized below (Table 5).

Table 1: Kinetic data used in the model for each enzyme in the SMART amplification reaction.

Enzyme	$k_{cat} (\text{s}^{-1})$	$K_M (\text{M})$
AMV-RT	$1.9369 * 10^{-2}$	$8.5 * 10^{-8}$
T7 RNA Polymerase	$7.04 * 10^{-2}$	$6.3 * 10^{-9}$
RNase H	$1.6 * 10^{-1}$	$7.32 * 10^{-8}$

Values for $[E]$ were calculated using the known number of Units/volume of each enzyme in the reaction mix according to the protocol along with specific activity and molecular weight (values were obtained from New England Biolabs) These values permitted conversion of Units/volume into concentrations.

Table 2: Data describing important parameters for each enzyme for the SMART amplification reaction.

Enzyme	Units/volume (U/L)	Specific Activity (U/mg)	Molecular Weight (Da) (g/mol)	$[E] (\text{M})$
AMV-RT	$3.25 * 10^5$	$4 * 10^4$	$1.58 * 10^5$	$5.14 * 10^{-8}$
T7 RNA Polymerase	$1.6 * 10^6$	$7.4 * 10^5$	$9.8 * 10^4$	$2.21 * 10^{-7}$
RNase H	$5 * 10^3$	$1.1 * 10^6$	$1.85 * 10^4$	$2.46 * 10^{-10}$

The final parameter to be chosen is the time step. Since the binding steps do not contain the value t for the length of an iteration of a time step, the time step represents approximately how long it takes for the binding events to reach equilibrium. For solution-phase kinetics, DNA with minimal secondary structure will nearly completely equilibrate by 60 seconds [63]. Thus, the time step chosen for the model is 60 seconds.

Results

The initial test checked if the amplification reaction was correctly designed and could be a viable option. In this experiment, Set 1 was used. A positive control was a reaction that contained all of the reagents along with 100 nM of SMART Probe in the final concentration. The other conditions involved removing one of the following: AMV-RT, T7 RNA polymerase, 5 Primer 1, and Primer 2. Gel electrophoresis using the RNA Nano 6000 assay was run, and the gel plot is shown Fig. 6.

The first lane in Fig. 6 shows the positive control. Two dark bands show that a high concentration of nucleic acid was present. Two peaks are shown. The RNA Nano 6000 assay did not separate small strands of nucleic acids, less than 100 bases. Removing either of the 10 enzymes or Primer 1 shows that the amplification does not occur when any of these reagents are missing. This agrees with the design of the assay, as AMV-RT was needed to make the template for RNA transcription. T7 RNA polymerase was the enzyme involved in transcription, and Primer 1 contained the promoter sequence for the T7 polymerase. Thus, absent any of these reagents, RNA was not polymerized. The final lane, which does not contain Primer 2 in the 15 reaction, shows that while amplification occurred, it did so at a diminished rate. This is shown as the bands were lighter, and fewer nucleic acids were made compared to the positive control. This also follows the design as removing Primer 2 did not allow for a cyclic, exponential reaction, but it still allowed for RNA transcription. It is important to note that 100 nM Probe was a relatively high concentration, and thus Primer 2 is still an important part of the reaction for 20 lower concentrations of Probe.

To optimize some of the conditions of the reaction, DMSO concentration and TrisHCl buffer pH were varied. For both experiments, Set 1 was used. DMSO is believed to reduce nonspecific interaction between different nucleic acids as well as disrupting secondary structure of the template and primers [52]. DMSO is also reported as difficult to work with as reagents 25 can precipitate out of solution. Since the length of the product has been decreased, it was hypothesized that a lower DMSO concentration could be a viable option. The reaction was performed with the normal 15% DMSO final concentration and with 5% DMSO final concentration in a thermal cycler and was analyzed with gel electrophoresis with the RNA Nano 6000 assay

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As shown in Fig. 7, three concentrations of probe, 100 nM, 10 nM, and 1 nM, were used for the different DMSO concentrations. The gel plot shows that the reaction was hindered by 5% DMSO. Throughout all of the experiments, even if some reagents precipitated out of solution at 15% DMSO, heating and mixing could put them back into solution and did not appear to affect the results of the reaction.

Initial tests used Tris-HCl pH 8.0, but others have reported using Tris-HCl pH 8.3 and 8.5 [44, 54]. As an experiment, Tris-HCl pH was also varied at pH 8.0 and pH 8.3. Three concentrations of probe, 100 nM, 10 nM, and 1 nM, and a negative control, 0 nM probe, were run at each pH level. The results of the gel plot are shown in Fig. 7. Lanes 1-4 correspond to Tris-HCl pH 8.0, and lanes 5-8 with pH 8.3.

The results as shown in Fig. 8 are that although the reaction continues to work at the higher pH level, the efficiency of the reaction drops due to the higher pH. Because of this, Tris-HCl pH 8.0 was used for all subsequent reactions to maintain efficiency.

Further studies were carried out to identify each individual strand as shown in Fig. 8 on a gel or electropherogram plot. For this purpose, the Small RNA Assay from Agilent was used in various experiments. Set 1 was used for this experiment. To identify known products, the individual DNA strands in the reaction, Probe, Primer 1, Primer 2, and combinations of the primers were added to a reaction mix without enzymes. In addition, the double stranded DNA expected from the reaction after the AMV-RT step was purchased and added into a mix without enzymes and with only the T7 RNA polymerase to identify the double stranded template DNA as well as the RNA produced in the reaction. Probe concentration was 10 nM in the reaction. The gel plot, Fig. 9, shows that the starting concentration of Probe is too low to be detected via gel electrophoresis. In addition, the combination of Probe and Primer 1 bound together show up where Primer 1 alone appears. Adding Primer 2 to this mix simply adds a band where expected for Primer 2, which seems to confirm that Primer 2 is not binding in significant amounts to either the Probe or Primer 1. Given the known lengths of these segments, the bands seem to come more quickly than the assay calculates via the ladder in the left most lane. This could be due to the high salt content of the reaction conditions. Furthermore, the cDNA band comes even quicker than expected. Thus, double-stranded species come quicker than expected, most likely due to

their minimal secondary structure, which would slow their migration through the gel. The final lane shows that multiple RNA products are actually made, as there are two strong bands that appear when T7 is simply added to cDNA. Abortive products are not uncommon with RNA polymerases, and thus this is not a cause for concern. The electropherogram of this data, Fig. 10, shows a clearer diagram of the last three lanes. In the data, the signal for the cDNA and T7 lane becomes jagged, rising a little over the baseline. This implies that other, smaller RNA or DNA products were also made during the reaction, likely as a result of incomplete transcription from a template.

The above data are shown in the electropherogram of Fig. 11. Without being bound by any particular theory it is believed that Fig. 11 establishes an increase in efficiency when RNase H is added despite a relatively high starting concentration of Probe and that RNase H facilitates the reaction. Known strands are marked on Fig 11. Fig. 11 confirms that the reaction makes the RNA strand that is designed to be made.

Fluorometer data was taken by using a series of 10:1 dilutions of SMART probe. Both sets were tested. Negative control was taken as 0 M SMART probe. The data for Set 1 is shown in Fig. 12. For the curves corresponding to starting concentrations of Probe of 1 nM, 100 pM, 10 pM, and 1 pM, the lines flatten at the top. This occurred because the detector of the fluorometer became saturated. Since relative fluorescence is plotted, though, and the background signal at 5 minutes varied, the curves peak at different points. Fig. 12 shows that decreases in starting concentration of Probe lead to increased time to achieve the same level of fluorescence. In the instant assay, the lower threshold for detection is about 100 fM Probe. Fig 12 also shows that a criterion for a positive signal is safely set at about three times the original background signal (corresponding with a relative fluorescence of 2). Fluorometric data is useful in both qualitative and quantitative measurement,

A microfluidic chip incorporating the present method allows for a cost-effective option as minimal reagents are used and facilitating uses in a point of care device. A chip design is presented in Fig 13. This example incorporates a section for the capture step in addition to the amplification step.

In Fig 13, gray areas in the schematic denote apparatuses for either heating at a constant temperature (two on the left) or for a detector (right). In this schematic, the left side will vary

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Claims:

1. A method of amplify Target RNA comprising the steps of
 - (i) introducing at least one Target RNA to a sample containing probe nucleotide
5 under hybridizing conditions ;
wherein said Target RNA comprises three regions,
said first region being a Hybrid Seq RC region; and,
a second region, being a Target RC regions contiguous with said first region; and
a third region being a Primer 2 region contiguous with said second region; and,
10 (ii) selectively amplifying the RNA of said Target RNA.
2. The method of Claim 1 further comprising detecting said amplified RNA.
3. The method of Claim 2 wherein said detecting is by the method of gel electrophoresis or
15 fluorescence
4. The method of Claim 3 wherein said detection by fluorescence is by molecular beacon.
5. The method of Claim 1 wherein said amplifying of step (ii) comprises transcribing
20 hybridized Target RNA into double-stranded DNA .
6. The method of Claim 5 wherein a promoter containing a T7 promoter sequence binds to said
probe RNA.
- 25 7. The method of Claim 5 wherein said transcribing is by means of a reverse transcriptase.
8. The method of Claim 7 wherein said reverse transcriptase is AMV-RT.
- 30 9. The method of Claim 5 further comprising transcribing said resulting double stranded
DNA into RNA.

10. The method of Claim 9 wherein said transcribing is by means of an RNA polymerase that catalyzes the formation of RNA in the 5'→3' direction polymerase forming a two strand, DNA-RNA hybrid.
- 5 11. The method of Claim 10 wherein said polymerase is a T7 RNA polymerase.
12. The method of Claim 10 wherein only said RNA strand of a DNA-RNA hybrid is degraded.
- 10 13. The method of Claim 12 wherein said degradation of said RNA strand is by RNase H.
14. The method of Claim 1 wherein said Hybrid Seq RC region, Target RC region and said Primer 2 region each comprise from about 8 to about 35 bases which number as to each region may be the same or different.
- 15 15. The method of Claim 1 further comprising binding a molecular beacon to the amplified RNA.
16. A method of detecting target nucleotide comprising
- 20 (i) exposing ProbeLeft nucleotides and ProbeRight nucleotides under hybridizing conditions to a Probe-specific ligase;
- (ii) permitting ligating of said ProbeLeft and ProbeRight nucleotides if they are adjacent to each other while said ProbeLeft and ProbeRight nucleotides are hybridized to a complementary nucleotide sequence; and,
- 25 (iii) detecting the presence or absence of said ligated ProbeLeft with ProbeRight nucleotide.

Fig. 1

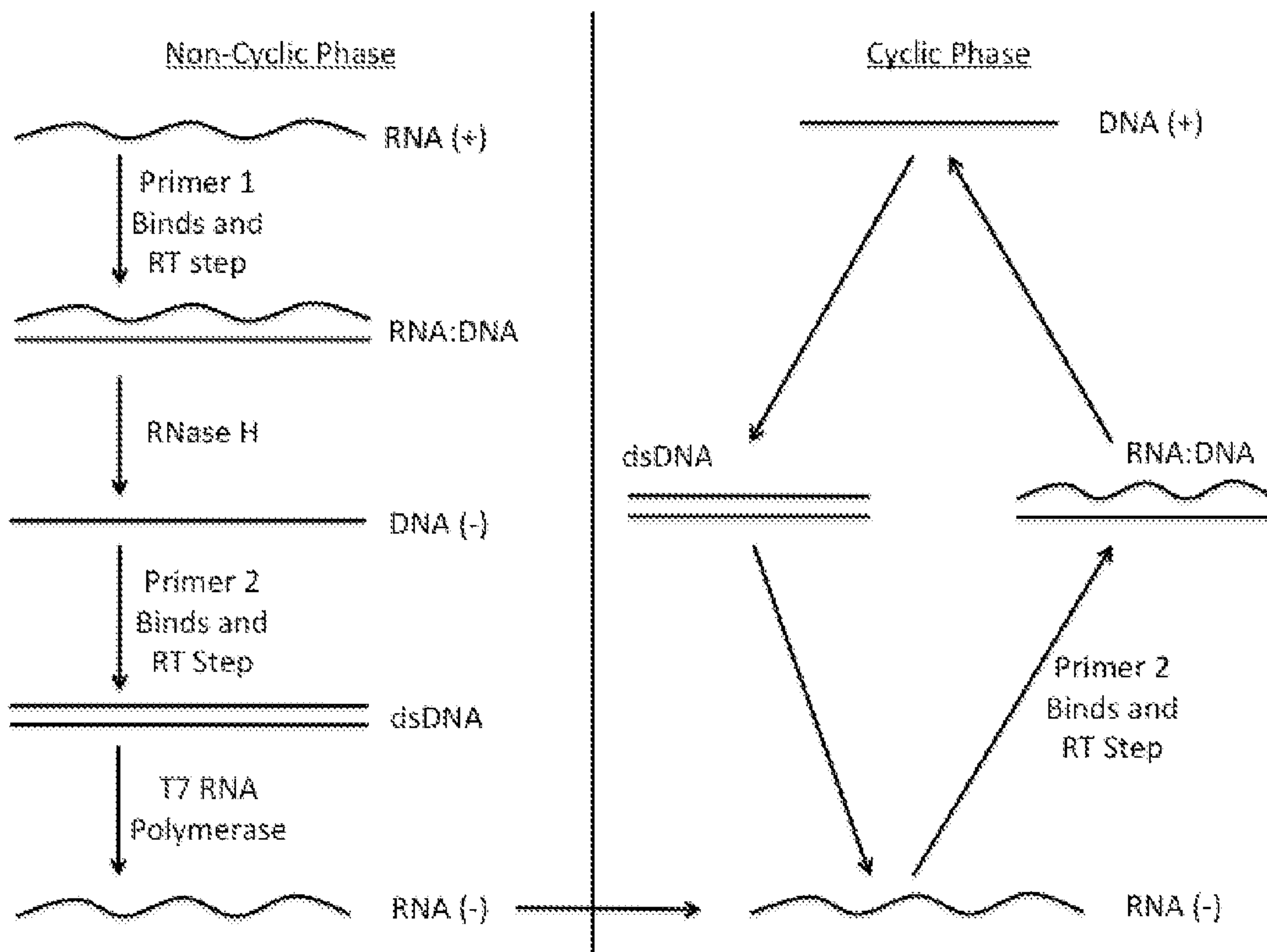


Fig. 2.

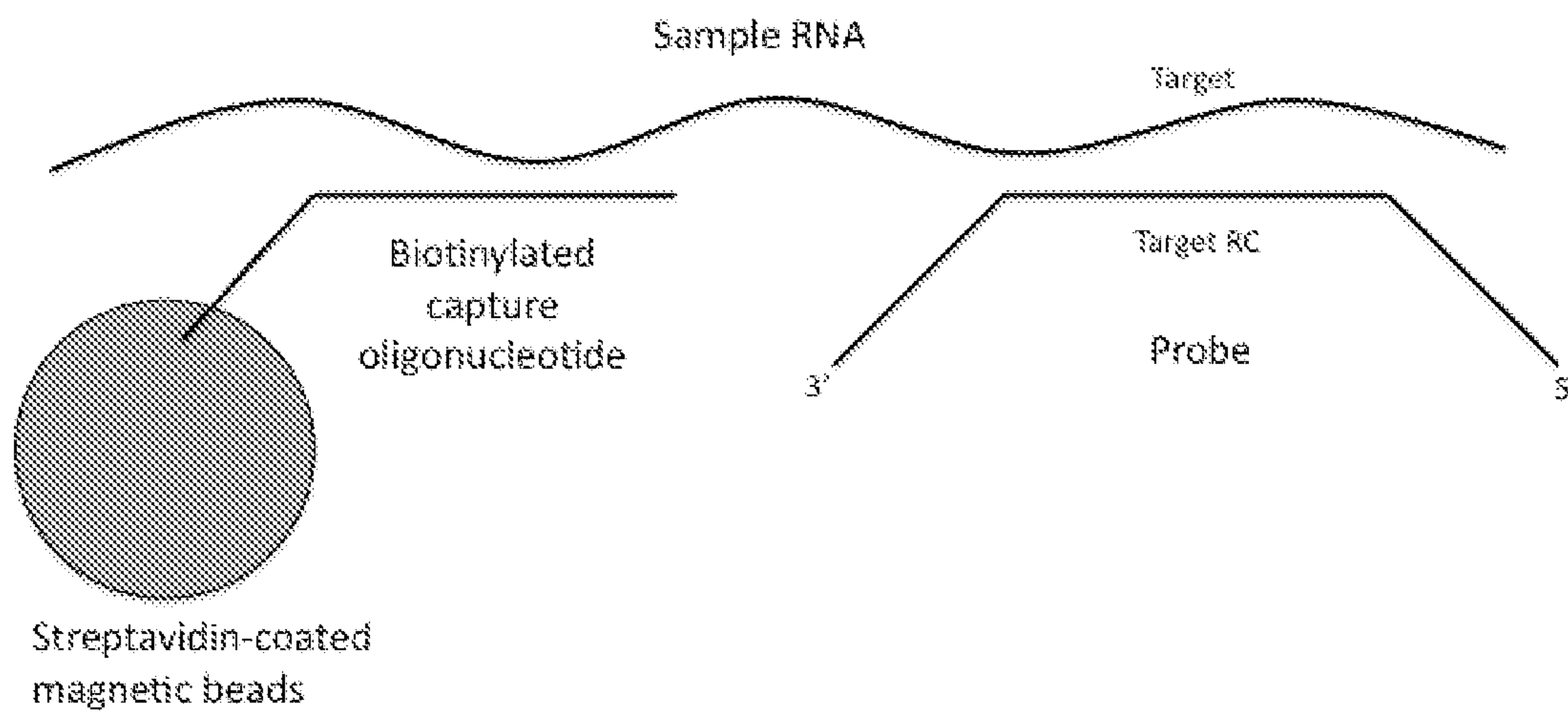


Fig. 2A

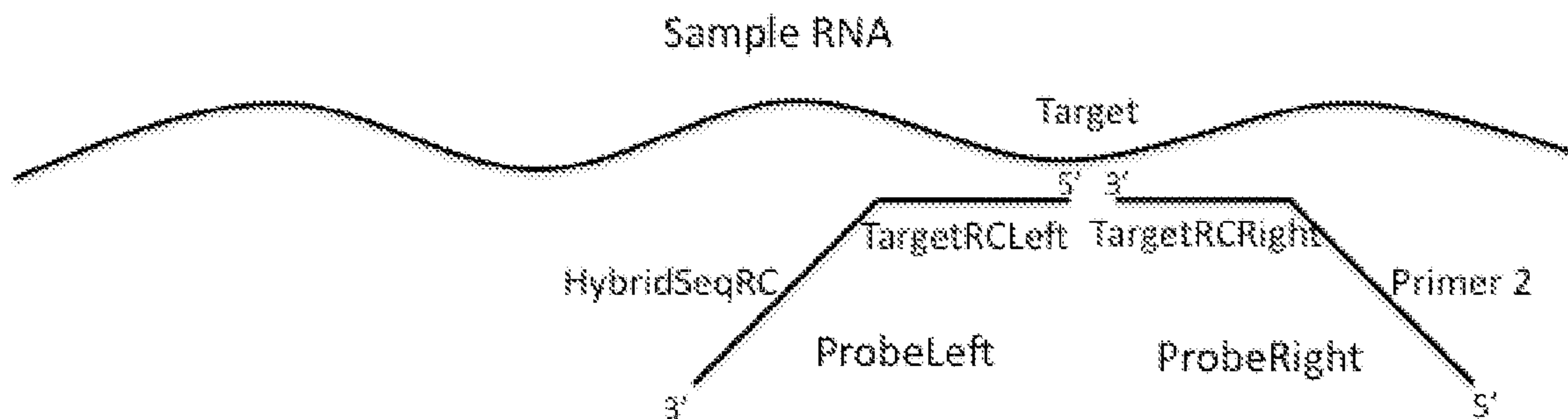


Fig. 3

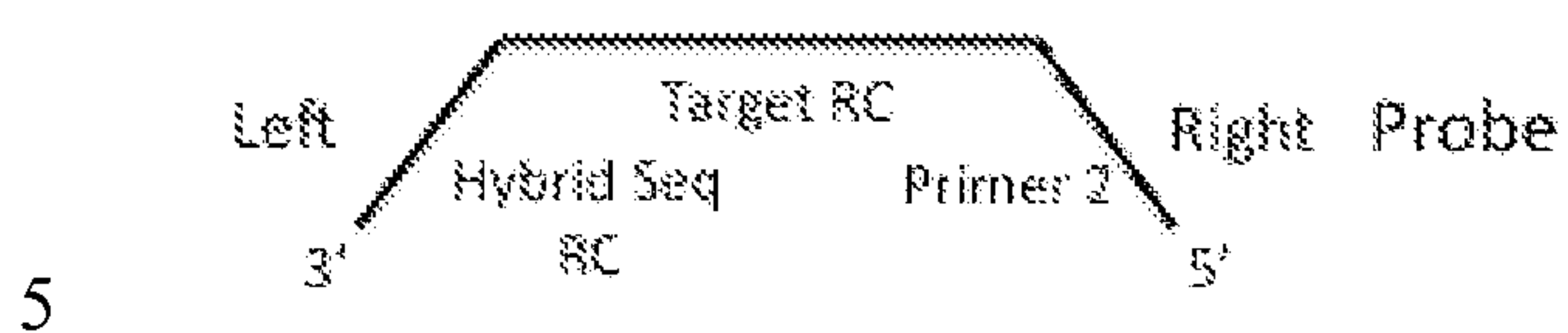


Fig. 4

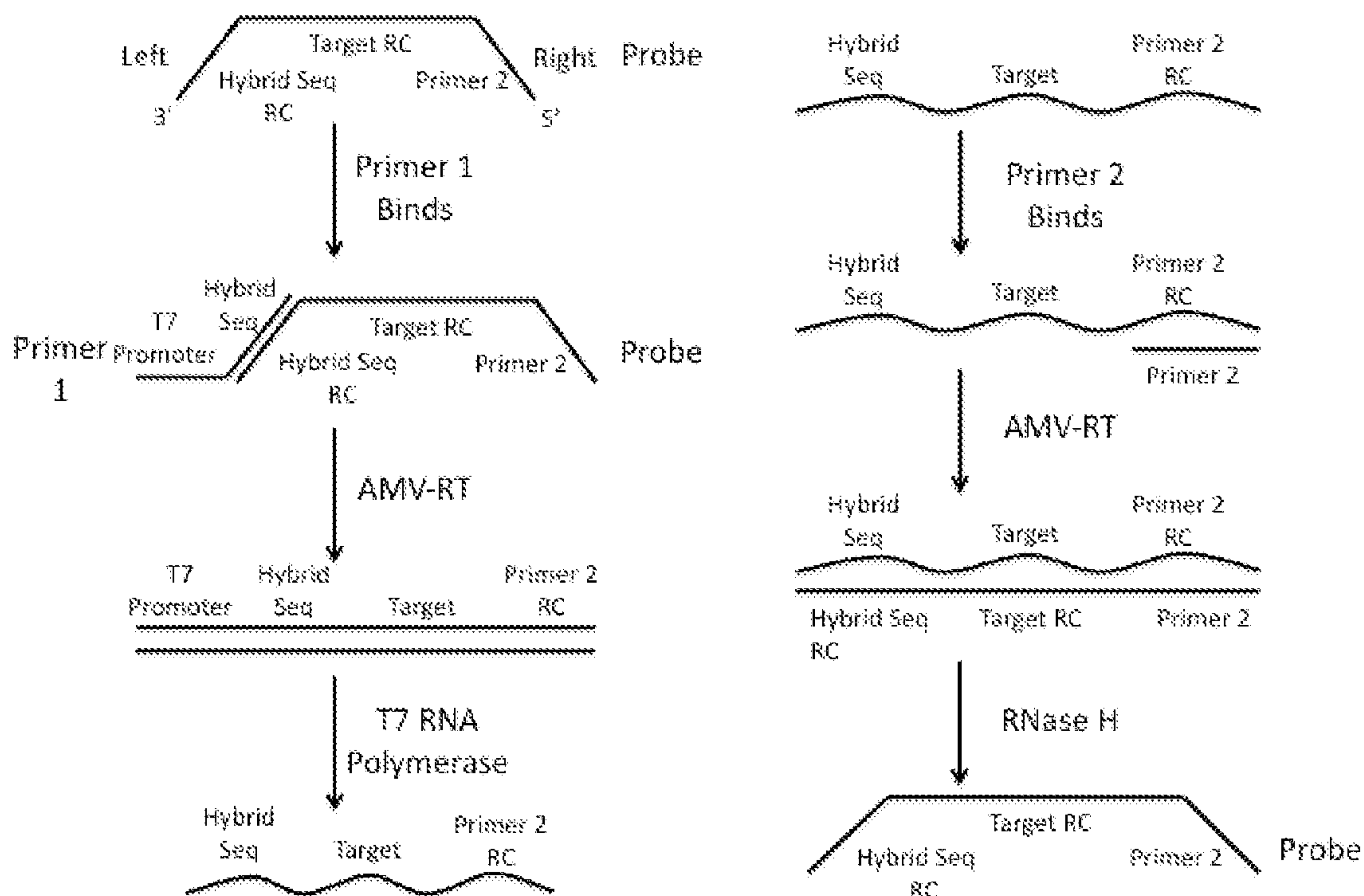


Fig. 5

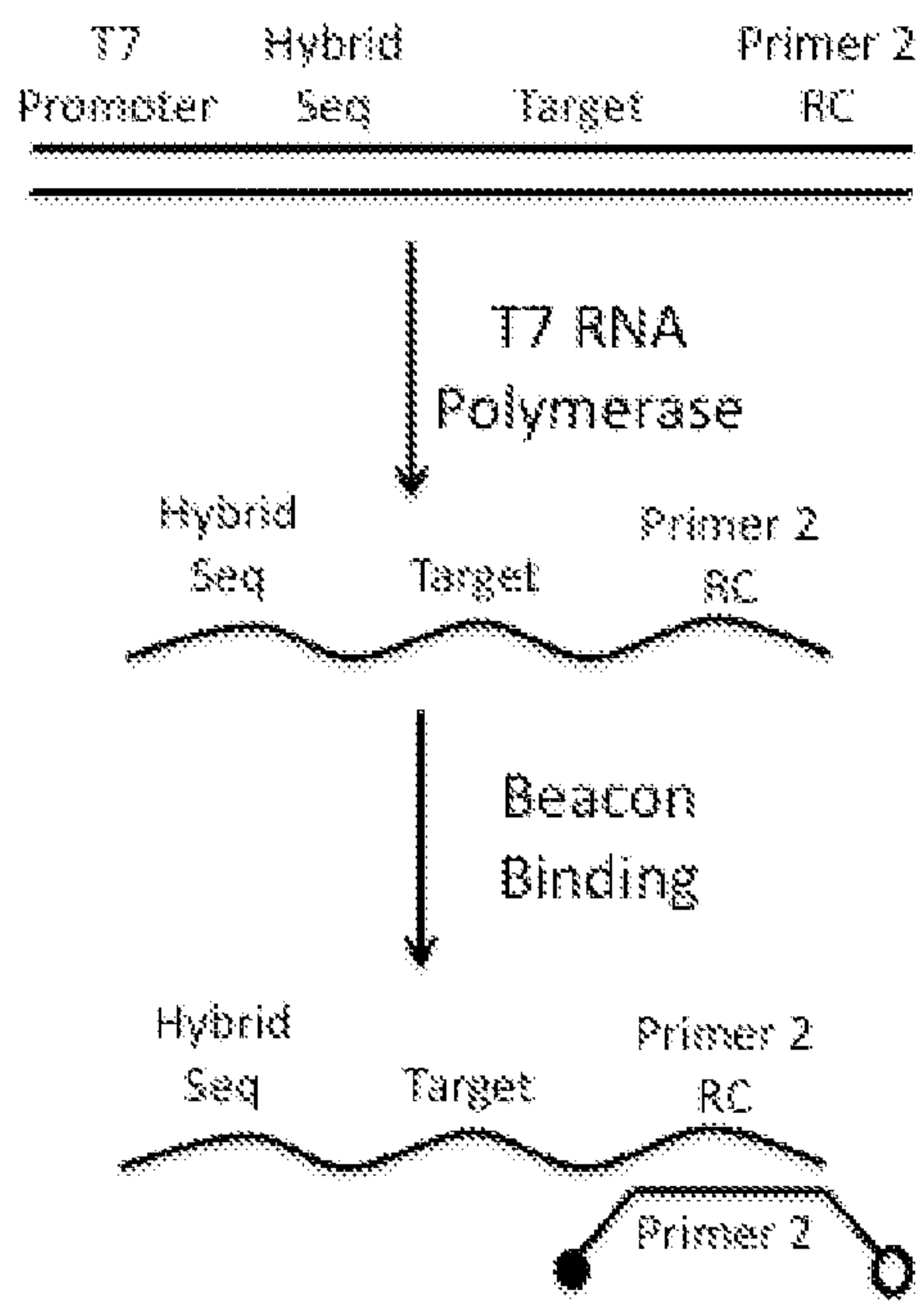


Fig. 6

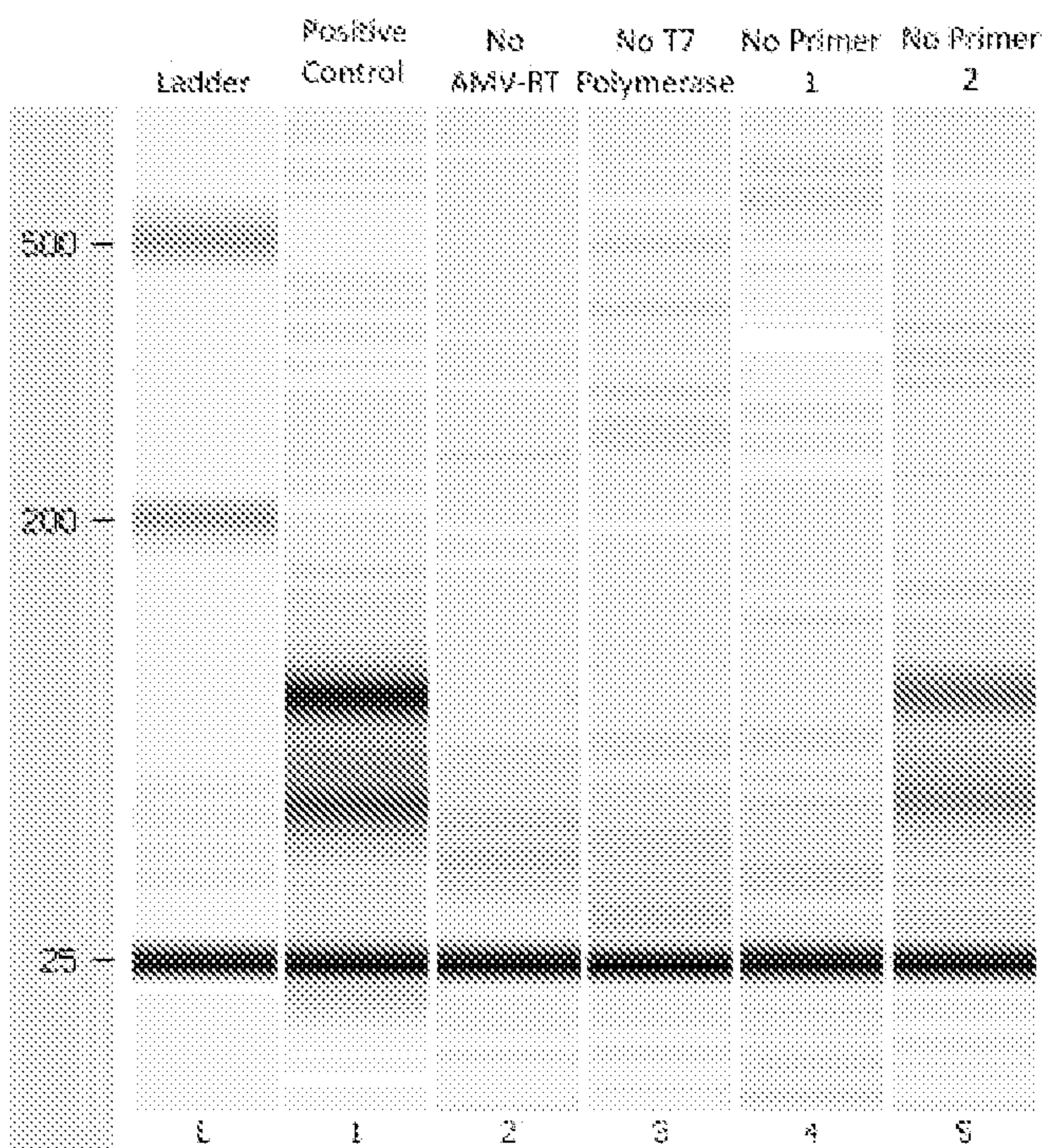


Fig. 7

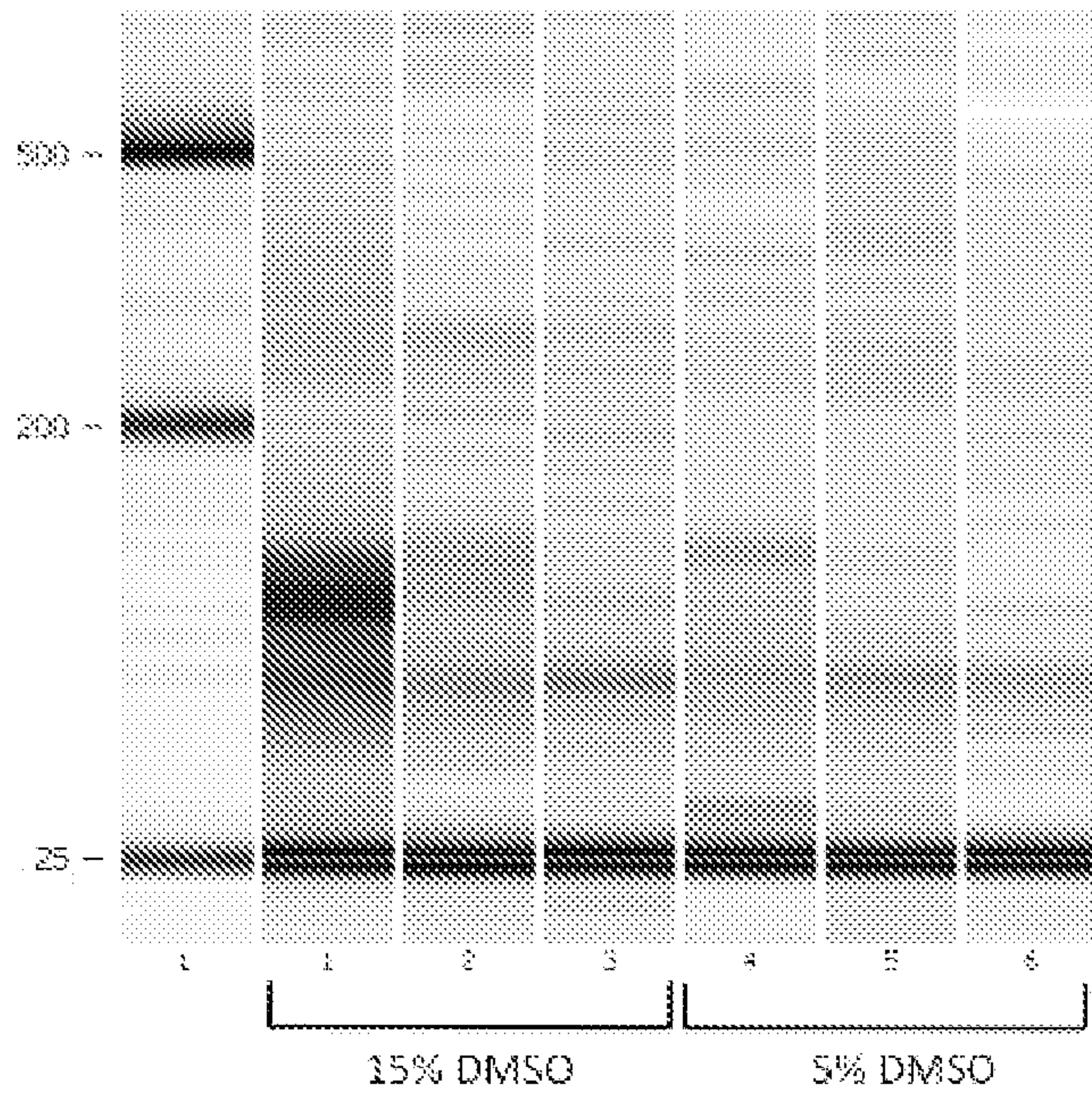


Fig. 8

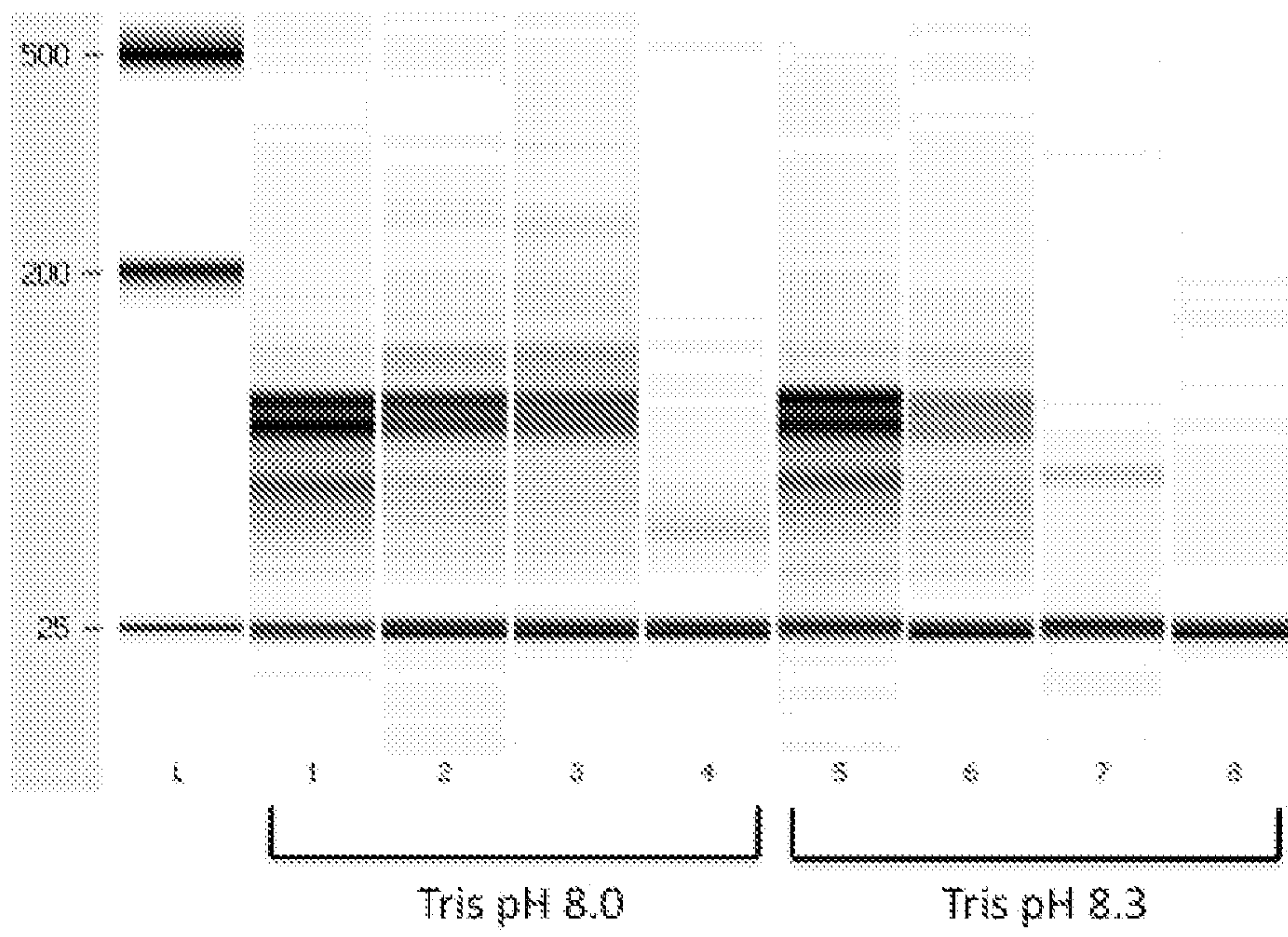


Fig. 9

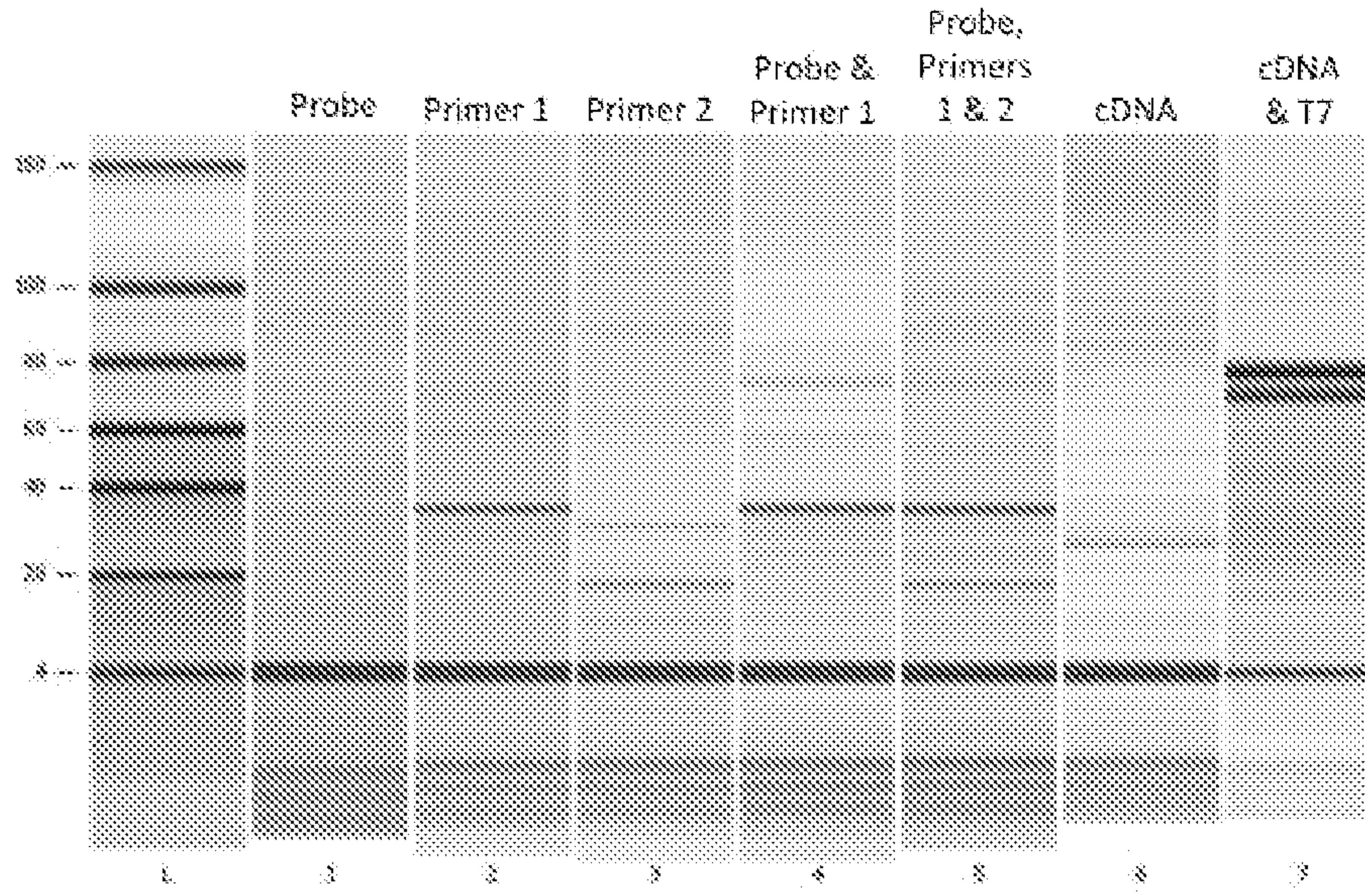


Fig. 10

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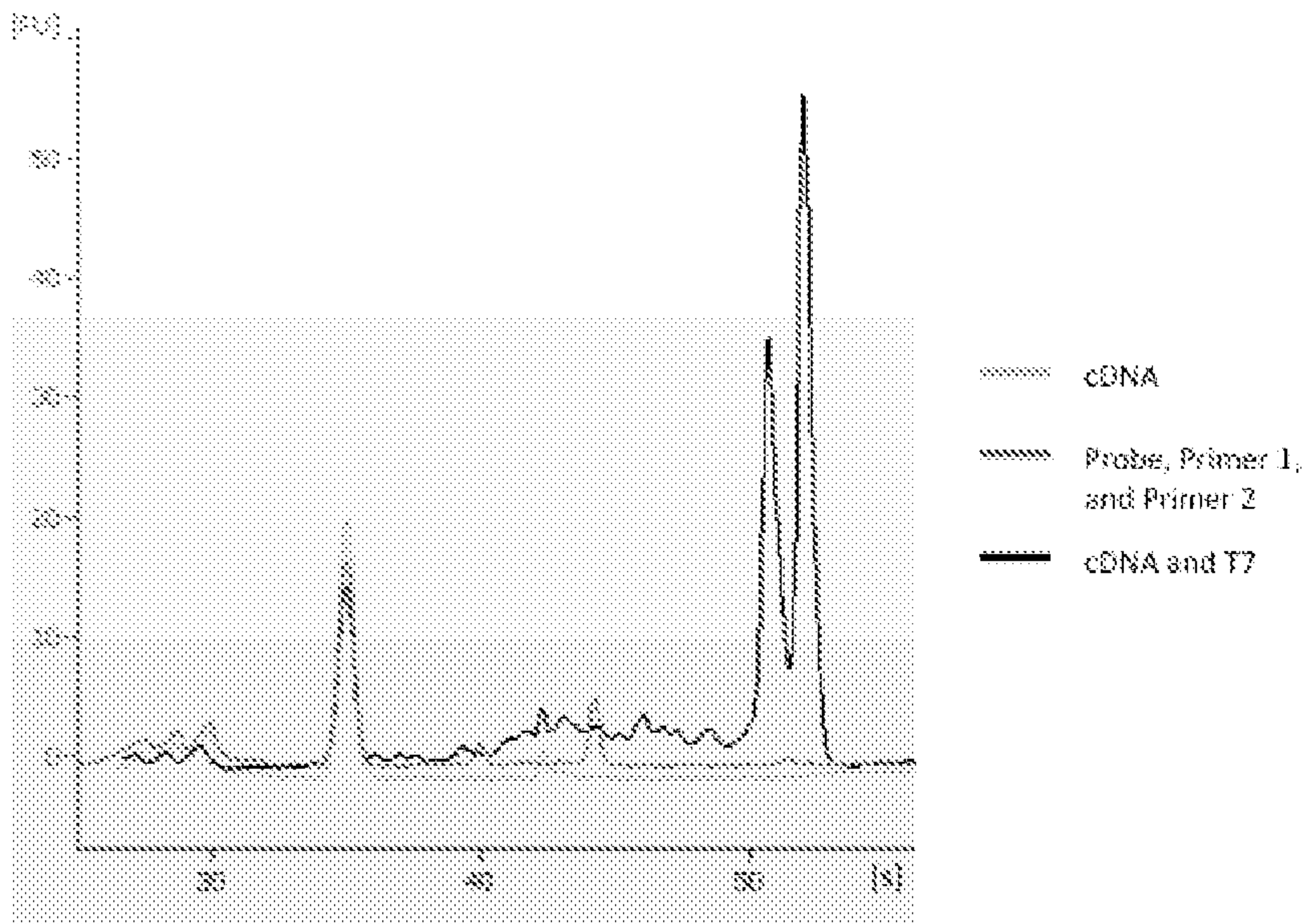


Fig. 11

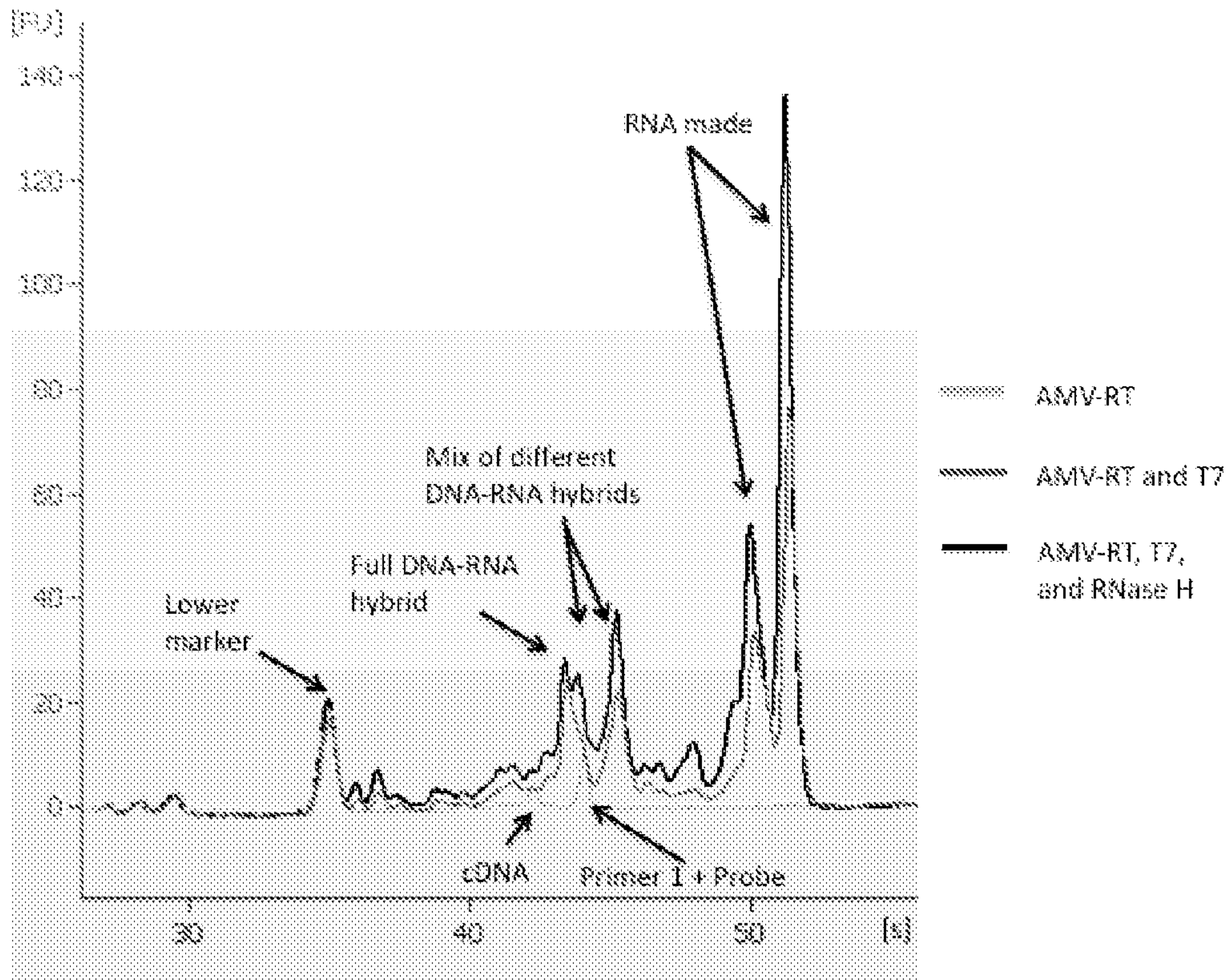


Fig. 12

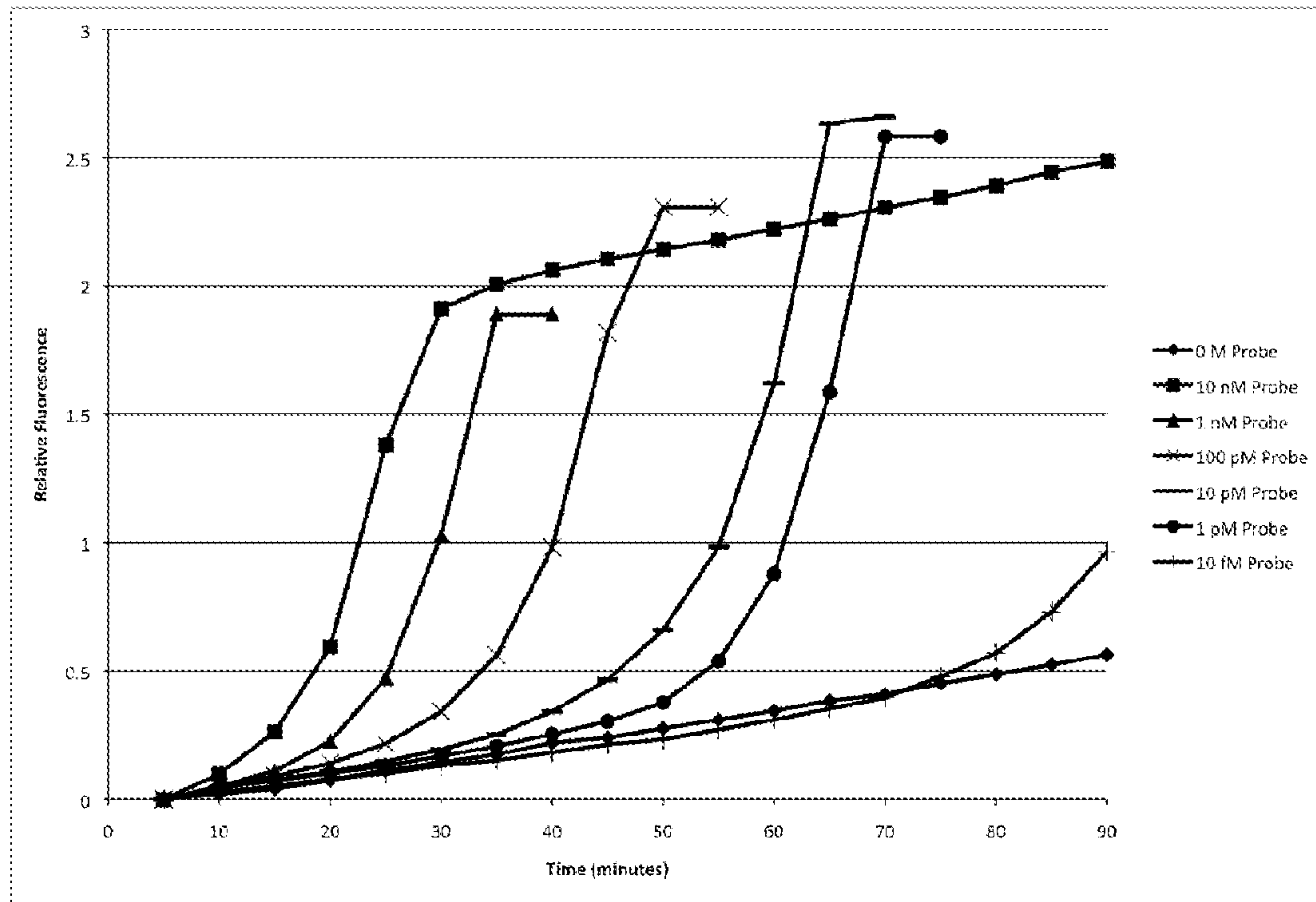


Fig. 13

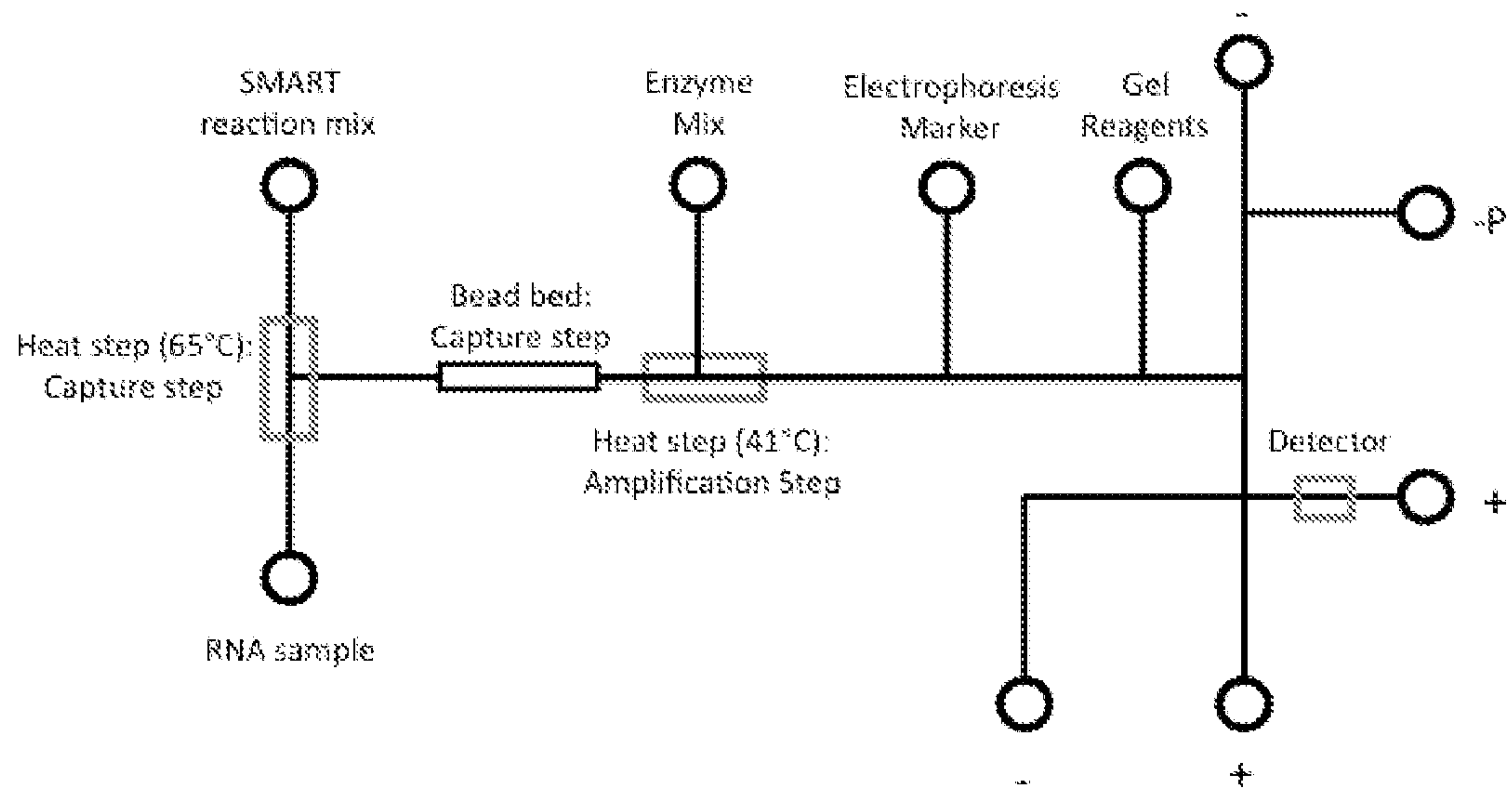


Fig. 1

