A disposable, integrated loop-mediated isothermal amplification cassette with thermally actuated valves

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Abstract An inexpensive, disposable, integrated, polymer-based cassette for loop-mediated isothermal amplification (LAMP) of target nucleic acids was designed, fabricated, and tested. The LAMP chamber was equipped with single-use, thermally actuated valves made with a composite consisting of a mixture of PDMS and expandable microspheres. The effect of the composite composition on its expansion was investigated, and the valve’s performance was evaluated. In its closed state, the valve can hold pressures as high as 200 kPa without any significant leakage. Both the LAMP chamber and the valves were actuated with thin film heaters. The utility of the cassette was demonstrated by carrying out LAMP of Escherichia coli DNA target and reverse transcribed loop mediated isothermal amplification (RT-LAMP) of RNA targets. The amplicons were detected in real time with a portable, compact detector. The system was capable of detecting as few as 10 target molecules per sample in well under 1 h. The portable, integrated cassette system described here is particularly suited for applications at the point of care and in resource-poor countries, where funds and trained personnel are in short supply.

Keywords Integrated microfluidic cassette • Loop-mediated isothermal amplification (LAMP) • Thermally actuated PDMS valves • Escherichia coli • Nucleic acid detection • Real-time fluorescence quantification • Expandable microsphere • Thermal actuation

1 Introduction

Nucleic acid amplification is an indispensable step in molecular diagnostics. Recently, there has been an increasing interest in point of care and home genetic diagnostic platforms using microfluidic chip technology (Yager et al. 2006; Easley et al. 2006; Chen et al. 2007; Liu et al. 2009). A great amount of effort has been invested in developing microfluidic chips for polymerase chain reaction (PCR) (Chen et al. 2004; Auroux et al. 2004; Zhang et al. 2006; Zhang and Xing 2007; Wang et al. 2006; House et al. 2010). Although PCR is a well established and widely used technology for DNA amplification, the need for thermal cycling complicates the use of PCR at point of care settings and increases the cost of PCR-based devices.

As an alternative to PCR, various isothermal nucleic-acid amplification methods have been proposed (Compton 1991; Walker et al. 1992; Piepenburg et al. 2006; Notomi et al. 2000). Isothermal nucleic-acid amplification technologies are more suitable for point of care diagnostics as they require less complex thermal control than PCR. Lutz et al. (2010) demonstrated a recombinase polymerase amplification (RPA)-based lab-on-a-foil for the detection of the antibiotic resistance gene mecA of Staphylococcus aureus. Gulliksen et al. (2005) and Dimov et al. (2008) reported, respectively, on using nucleic acid sequence-based amplification (NASBA) microfluidic chips for the detection of the human papilloma virus (HPV) and Escherichia coli bacteria. Sato et al. (2010) demonstrated a microbead-based, rolling circle amplification (RCA)
microchip for *Salmonella* detection. Fang et al. (2010) developed a microchip-based, loop-mediated isothermal amplification (LAMP) system for pseudorabies virus (PRV) detection. Ramalingam et al. (2009) reported combining helicase dependent amplification (HDA) with microchip technology for amplifying the BNI-1 fragment of SARS cDNA.

The devices reported above focus on the reaction chamber and do not integrate valves into the device. It is desirable to seal the reaction chamber to reduce the introduction of contaminants and to prevent evaporation as the nucleic acid amplification process takes place at an elevated temperature. Although valveless technologies have been developed to reduce evaporation (Zhang and Xing 2010), valves are still predominantly used for sealing in microfluidics (Chen et al. 2007; Wang et al. 2006; Das et al. 2007; Oh and Ahn 2006). Diverse, ingenious actuation mechanisms for microfluidic valves have been developed such as piezoelectric (Li et al. 2004; Shao et al. 2004), magnetic (Bae et al. 2002; Gaspar et al. 2008), thermal (Kim et al. 2004; Chen et al. 2008), pneumatic (Lagally et al. 2004; Lien et al. 2009), and hydraulic (Oh et al. 2005). Thermal actuation, which does not require moving components, is often adopted in microfluidic chips because of its simplicity, low cost, and reliability (Liu et al. 2004, 2002; Chen et al. 2005; Wang et al. 2005). However, thermally actuated valves, such as ice valves, require a continuous power supply during their entire actuation time, which is undesirable for a reactor operating for as long as an hour. Moreover, in many nucleic acid amplification reactors with real-time detection, in which the amplification products need not be further processed after detection, a single action “one-shot” valve would suffice. Recently, Stemme’s group reported on a simple, easy to fabricate, thermally responsive, single action valve cast with a mixture of PDMS and expandable Expancel® microspheres (Samel et al. 2007a, b).

In this paper, we report on a plastic, disposable cassette consisting of a reaction chamber for isothermal amplification of nucleic acids and two thermally actuated valves made with a PDMS–expandable microsphere composite. Prior to the amplification process, the valves were heated, expanded, and sealed the amplification reactor. The amplification process was monitored in real time with a minute detector. The paper describes the design and construction of the cassette and characterizes, for the first time, the expandability of the PDMS–expandable microsphere composite as a function of the PDMS’ composition and dopant concentration. The dopants were added to augment the thermal conductivity of the PDMS. Furthermore, we demonstrated the utility of the integrated device by amplifying, with LAMP, both DNA and RNA targets. As an example of a DNA target, we used the *E. coli* pathogen, which causes infections of extraintestinal sites such as the urinary tract and meninges. As an example of a RNA target, we used the RNA control included in the reverse transcription, loop-mediated, isothermal amplification (RT-LAMP) kit provided by the Eiken Chemical Co. In a separate work, we used the LAMP process to amplify and detect HIV RNA (Liu et al. 2011).

## 2 LAMP cassette and reagents

### 2.1 LAMP cassette

The 46 mm × 36 mm × 3.37 mm cassette consists of three layers of polycarbonate (McMaster Carr, Atlanta, GA, USA) (Fig. 1a): a top, 120 μm thick, polycarbonate film; a 3.0 mm thick cassette body; and a bottom, 250 μm thick polycarbonate film. The various layers were solvent-bonded with acetonitrile at room temperature. Figure 1a
and b are, respectively, an exploded view of the cassette and a photograph of the cassette’s body. The cassette’s body was fabricated using a precision, computer-controlled (CNC) milling machine (HAAS Automation Inc., Oxnard, CA) (Liu et al. 2009). The cassette has two functional domains: a LAMP/RT-LAMP incubation (reaction) chamber and two valve chambers. The reaction chamber is connected to the valve chambers with 500 μm wide and 250 μm deep conduits with rectangular cross-sections. Inlet and exit ports, each 0.8 mm in diameter, were drilled (with a tabletop milling machine, MF 70, Proxxon, Niersbach, Germany) in the side of the main body of the cassette (Fig. 1a). One valve chamber is connected to the inlet port, and the other valve chamber is connected to the exit port.

The reaction chamber is 10 mm in length, 2.0 mm in width, 1.0 mm in depth, and 20 μL in volume. The chamber is located at the bottom of the cassette’s main body, and it is separated from the heater by the 250 μm thick, bottom polycarbonate film. The inlet and outlet conduits of the reaction chamber are connected through vertical vias to conduits milled in the upper surface of the main body. To improve thermal isolation, the reaction chamber is surrounded by air-filled cavities.

Each valve chamber consists of a 4 mm diameter, 0.8 mm deep well to house a thermo-responsive, PDMS composite disk and a shallow 3 mm diameter, 150 μm deep compartment to facilitate fluid flow when the valve is open (Fig. 2). Figure 2a depicts the cross-section of the assembled valve (all the dimensions are given in mm). The valve structure will be described in detail in Sect. 2.2. The valves’ compartments were milled in the bottom surface of the main cassette body. Like the reaction chamber, the valves are surrounded by air-containing cavities to provide thermal isolation.

2.2 Thermo-responsive PDMS valves

The valve was formed with a composite of PDMS (Sylgard 184 Silicone Elastomer Kit, Dow Corning) and highly expandable microspheres (Expancel 031 DUX 40, Duluth, GA). Expancel microspheres consist of ~10 μm diameter spherical shells, made with vinylidene chloride, acrylonitrile, and methyl methacrylate, and filled with pressurized liquid hydrocarbon such as isobutane and isopentane. When the microsphere is heated above 80°C, the hydrocarbon undergoes a phase transition into a gas and inflates the shell. Due to the elevated temperature, the shell material is soft and undergoes plastic deformation. As a result, the sphere’s diameter increases approximately four-fold and its volume increases by a factor of ~60. This transition is essentially irreversible. Once the sphere is cooled, it remains in its expanded state (Roxhed et al. 2006; Griss et al. 2002).

To prepare the PDMS–expandable microsphere composite, we mixed the PDMS base, a curing agent, and expandable microspheres at predetermined ratios, and degassed the mixture in vacuum. In some of our experiments, we added silver powder (2–3.5 μm diameter particles with 99.9% purity, Sigma-Aldrich, St. Louis, MO) to the mixture to enhance the thermal conductivity of the composite. Various blend compositions were tested (Sect. 4.2). The suspension was carefully cast in a
polycarbonate mold consisting of 0.8 mm deep, 4 mm diameter milled wells (Fig. S1). Subsequently a flat, 5 mm thick polycarbonate slab was slowly placed over the PDMS–expandable microspheres suspension, allowing surface tension to pull the polycarbonate slab into intimate contact with the prepolymer without any bubble formation at the interface. To prevent the suspended microspheres from prematurely expanding, the curing of the PDMS–expandable microspheres composite was carried out at 60°C (i.e., below the phase transition temperature of the microspheres) for about 4 h.

Figure 2 depicts the normally open valve. Figure 2a is a cross-section of the valve in its open state. Figure 2b and c are, respectively, three-dimensional renderings of the open valve and closed valve. When the valve is in an open state (Fig. 2a, b), fluid is free to flow from conduit A to conduit B. When the valve is heated, the PDMS composite expands and permanently seals the passage between conduits A and B.

2.3 Reagents and protocol for DNA LAMP

On-chip LAMP was carried out in the 20 µL LAMP reaction chamber at 63°C. For a description of the LAMP process, see Notomi et al. (2000) and Nagamine et al. (2002). The LAMP template (plasmid DNA), extracted from pathogenic E. coli was obtained from urine samples, and the primers for E. coli were a gift from Professor Abhay Vats (Department of Pediatrics, Children’s Hospital of Pittsburgh, Pittsburgh, Pennsylvania, USA). The primers and their respective concentrations for E. coli DNA amplification were: outer primer F3 5’-GCCATCTCCTGATGACGC-3’ (0.2 µM), outer primer B3 5’-ATTTACCGAGCCGACACCG-3’ (0.2 µM), loop primer F loop 5’-CTTTGTAAACAACCTGATCGACATACTCAG-3’ (0.8 µM), loop primer B loop 5’-ATCAATCTGATATCCATGAAGG-3’ (0.8 µM), inner primer BIP 5’-CTGGGGCGAGTGTCGTGATTTGTAACAACCTGATCGACATACTCAG-3’ (0.8 µM), and inner primer B 5’-ATTATACTCCATATTTGTACATATTTGCCGACCCATATCCATATGTGGCT-3’ (1.6 µM) and inner primer BIP 5’-CATTTGTGACGCTGACGATC-3’ (1.6 µM) (Hill et al. 2008). The reaction mixture also contained 20 mM Tris–HCl (pH 8.8) (Fisher Scientific, Inc., PA), 10 mM KCl, 10 mM (NH4)2SO4, 2 mM MgSO4, 0.1% Triton X-100 (Sigma Chemical Co., MO), 0.8 M betaine (Sigma-Aldrich, St. Louis, MO), 8 U of Bst DNA polymerase (New England Biolabs, Inc., MA), 1.4 mM dNTPs (Promega, WI), 4.0 µM SYTO® 9 Green (Molecular Probes, Inc., Eugene, OR), and 5 µL of E. coli DNA at various concentrations. DNase/RNase-free water (Fisher Scientific, Inc., PA) was used throughout the experiments.

To confirm the real-time fluorescence detection, at the end of the experiment, we drilled into the reactor and removed the reaction products with a pipette. The LAMP reaction products were analyzed by gel electrophoresis. 3 µL of each LAMP-amplified product was loaded onto a lane of a 2.0% agarose gel. Electrophoresis of the amplified DNA fragment was carried out in Tris–Acetate–EDTA (TAE) buffer at a constant voltage of 114 V for 40 min. DNA marker VIII (Roche Diagnostic, Indianapolis, IN, USA) was used to determine the sizes of the amplified DNA bands. The gel was stained with ethidium bromide and was visualized with UV illumination.

2.4 Reagents and protocol for RNA LAMP

Lacking the necessary safety equipment to work with viruses, we carried out the RT-LAMP experiments with the Loopamp RNA amplification kit (Eiken, Japan). The primers’ composition and the amount of target analyte are considered proprietary information and are unavailable. The target analyte concentration is reported to be in the concentration range typical of clinical samples. Each RT-LAMP reaction was carried out with a 20 µL reaction volume containing 0.2 µM each of the outer primers F3 and B3, 0.8 µM each of loop primers F and B, 1.6 µM each of the primers FIP and BIP, 1.4 mM dNTPs, 0.8 M betaine, 0.1% Tween 20, 10 mM (NH4)2SO4, 8 mM MgSO4, 10 mM KCl, 20 mM Tris–HCl (pH 8.8), 8 U of Bst DNA polymerase, 0.625 units of AMV reverse transcriptase, 4.0 µM SYTO® 9 Green, and 2 µL of RNA template. The reverse transcription and amplification were carried out in a single step at 63°C. In a separate effort, Liu et al. (2011) amplified HIV RNA using our cassettes (without the thermally expandable valves) and the RT-LAMP process.

3 The integrated, real-time LAMP system

Figure 3 is a photograph of the experimental setup of the integrated LAMP cassette with the real-time fluorescence detection. The fluorescence excitation and detection were carried out with a portable optical detection system (Fluo Sens SD 003, Qiagen Lake Constance GmbH, Germany). The ESE optical detector (also shown in the inset in Fig. 3) uses 470 nm LED as the excitation light source and low-noise Si-photodiode for fluorescence detection.

In the experiment, the LAMP mixture was introduced into the LAMP cassette through the inlet port and the inlet, thermo-responsive, PDMS valve. The liquid filled the LAMP chamber, displacing the air through the outlet port. Once the liquid arrived at the outlet valve’s location, both the inlet and exit, thermo-responsive PDMS valves were heated to 80°C with a flexible, polimide-based thin film heater (Model HK5572R7.5L25A, Minco Products, Inc., Minneapolis, MN) located beneath the cassette next to the
valves’ positions (Fig. 4). Although in our experiments, the heater was powered with a DC power supply (Model 1611, B&K Precision Corporation, CA), it could be powered with a battery. When the thermo-responsive PDMS disks were heated to over 80°C, they expanded and closed the valves. Although the heaters can be readily patterned on the cassette, it is anticipated that, for economic reasons, it would be preferred to have the heaters part of a reusable reader/analyzer rather than part of the disposable cassette.

Once both the upstream and downstream valves were closed, the LAMP chamber’s temperature was raised to 63°C for 1 h. The heat was supplied with a thin film heater, similar to the one used to actuate the valves, positioned beneath the LAMP chamber (Fig. 4). In the experiments reported here, the chamber’s temperature was controlled in an open loop mode. To calibrate the device, we constructed a mock cassette, filled the LAMP chamber with water, and inserted into the chamber a type-K thermocouple (Omega Engr., Stamford, CT, USA). Each wire was 75 μm in diameter and the junction diameter was ~170 μm. The thermocouple reading was monitored with a HH506RA multilogger thermometer (Omega Engr., Stamford, CT, USA).

The intensity of the fluorescence signal emitted by the LAMP chamber was measured with the portable ESE optical detector. The detector was interfaced with a computer through a USB interface, and the graph of the fluorescence intensity as a function of time was displayed on the computer screen. In addition, images of the fluorescence of the LAMP chamber before and after the amplification process were obtained with an Olympus IX-71 inverted fluorescence microscope (Olympus America Inc., Melville, NY).

4 Results and discussion

4.1 Characterization of the thin film heating system

Figure 5 depicts the steady-state reactor’s temperature as a function of the heater’s power input. In the temperature range between 23°C and 95°C, the temperature depended nearly linearly on the heater’s power input. In the laboratory (room temperature of ~23°C), when power of 0.32 W (1.75 V) was applied to the heater, the reactor’s temperature was maintained at 63°C ± 0.2°C for the entire

![Fig. 5](image-url)
duration of the LAMP incubation process (1 h). For field operation, it would most likely be necessary to operate the heater in a closed loop control mode. The design and construction of such a control system would be relatively simple. As an interesting alternative, heat can be supplied by an exothermic chemical reaction without a need for any electrical power. For example, recently, Hatano et al. (2010) have successfully carried out LAMP amplification of *Bacillus anthracis* pag and capB gene fragments with commercially available heating packs of the type used to warm food products, albeit not in a microfluidic setting.

Although possible background fluorescence from the polyimide-based, thin film heater was of concern, we did not detect any significant emission from the heater that could interfere with our real-time LAMP detection system.

4.2 Thermo-responsive PDMS valve

The PDMS–Expancel microspheres composite is a convenient material for forming single-action, “one-shot,” valves for sealing the reaction chamber. When the microspheres are heated in air to above 80°C, they expand irreversibly with about a 60-fold increase in volume. One would expect, however, a much smaller increase in the volume of the PDMS–microspheres composite.

To evaluate the expandability of the composite for various ratios of PDMS base and curing agent, we cast 4 mm diameter, 0.8 mm thick disks of thermo-responsive PDMS composite using various ratios of PDMS base, curing agent, and microspheres. Figure 6 depicts the relative change in the volume of the composite as a function of composition (N = 3). Not surprisingly, as the concentration of microspheres increases so does the expandability of the thermo-responsive composite. Too large a concentration of microspheres, however, increased the apparent viscosity of the PDMS blend and adversely impacted its castability and compatibility with the mold.

Samel et al. (2007a) mentioned, in passing without any details, that the ratio of the PDMS base/curing agent had no effect on expandability. We found otherwise. When the volumetric ratio of the PDMS base to the curing agent was significantly altered, the expandability varied dramatically. As the relative amount of curing agent decreased, the elasticity and the expandability of the thermo-responsive composite increased. A composite consisting of 25% microspheres by volume and a PDMS-base/curing agent ratio of 20:1 expands nearly 550% upon heating. This is a much greater expandability than the previously reported 250% for a PDMS with base/curing agent ratio of 10:1 (Samel et al. 2007a).

In addition to expandability, the compliance of the PDMS composite is a major factor that affects valve performance. As we increased the concentration of the microspheres, we encountered difficulties in maintaining a hermetic seal. Thus, as a trade-off between expandability and compliance, we elected to make our valves with a 5% (by volume) concentration of microspheres in a PDMS matrix with a ratio of 20:1 of base to curing agent. The resulting material expanded about 150% ± 5% upon heating.

Although the time response of the valve was not a critical factor in our application, we nevertheless measured the closing time of the valve. We define the valve response time as the time interval from the instant when the power supply was turned on to the instant when the valve fully expanded as observed visually. Figure 7 depicts the response time of the thermo-responsive valve consisting of a 4 mm diameter, 0.8 mm thick disk (20:1 base PDMS-curing agent ratio and 5% microsphere concentration) as a function of the power input. The time response decreased as the power input increased and attained an asymptotic value of about 23 s when the power supply exceeded 2 W. The time response is, of course, a function of the disk’s size and the thermal capacitance of the substrate. Due to the relatively large size of our valve, its time response is relatively slow. The valve response time can be approximately related to the power input with an expression of the form $a\ln\left(\frac{bQ}{cQ+d}\right)$, where $Q$ is the power supplied to the heater and $a$, $b$, $c$, and $d$ are parameters that depend on the valve’s dimensions and thermophysical properties.

Since PDMS has a low thermal conductivity (≈0.15 W/m-K, Mark 1999), the valve is not heated uniformly. Microspheres next to the heater expand first in our thermoresponsive PDMS disk while microspheres located far from the heater expand only after a certain time delay. To examine the effect of valve thermal conductivity on the
valve’s performance, we varied the conductivity of the PDMS (20:1 base to curing agent ratio with 5% mass fraction of microspheres) by doping the PDMS with silver powder, and we monitored the temperatures of the valve’s heated surface and the valve’s distal surface as functions of time for different levels of silver doping. Figure 8 depicts the temperature of the valve’s heated surface (curves 1, 2, and 3) and the valve’s distal surface (curves 4, 5, 6) as functions of time in the absence of silver doping (curves 1 and 6), when the valve material is doped with a mass fraction of 21% silver (curves 2 and 5) and when the valve is doped with 41% of silver by mass (curves 3 and 4). At time $t = 0$, the heater power was set to 0.85 W. The inset depicts schematically the experimental set-up. As the silver content increases, the valve heated surface’s temperature decreases, the temperature of the valve distal surface increases, and the temperature difference between the heated surface and the distal surface decreases. All are indicators that the silver doping increases the effective thermal conductivity of the PDMS composite. We estimate (see Appendix) that the increase in the effective thermal conductivity is quite modest—about a factor of 1.8 in the presence of 21% silver mass fraction and a factor of four in the presence of 41% silver mass fraction. We did not use higher doping levels out of concern for the structural integrity of the valve.

Although the silver doping increased the valve’s effective thermal conductivity, it did not improve our valve’s performance. Since, in our case, the gap (Fig. 2) above the valve is very small, a relatively small expansion of the PDMS suffices to achieve a good seal. Thus, it was sufficient for just a fraction of the microspheres to expand in order to close the valve. In the absence of silver doping, the temperature of the heated surface of the valve increased faster and the valve provided a faster response time. Thus, in all the experiments described from here on we used undoped valves.

The holding pressure capability of the thermo-responsive PDMS valve was measured by applying back pressure and observing leakage through the valve. The valve successfully withstood pressures as high as 200 kPa without any visible leakage, which is also suitable for sealing a PCR amplification chamber (Wang et al. 2005).

In summary, the thermo-responsive PDMS valve is easy to construct, is reliable, and exhibits reproducible performance from one device to the other. Although, given the demands of our system, we used thermo-responsive PDMS valves with dimensions in the millimeter range, the valves’ size can be readily scaled down to the micrometer range with corresponding reductions in the response time and power consumption. Although not required in our application, the thermal conductivity of the valve can be controlled with appropriate doping.

4.3 Detection of DNA targets (E. coli)

*Escherichia coli* is the second major cause of neonatal meningitis, which is a major cause of mortality among newborns (Clarke 2001; Bonacorsi and Bingen 2005;
It is also associated with a high incidence of neurological sequelae. Thus, early detection of the presence of *E. coli* is highly desirable. Here, we use the *E. coli* as a model analyte to demonstrate the ability of our cassette to detect pathogen DNA. A dilution series of *E. coli* DNA samples were isothermally amplified using the protocol detailed in Sect. 2.3, and the fluorescent intensity was monitored as a function of time.

Figure 9a and b show, respectively, the fluorescence images of the *E. coli* sample (250 target copies) in the integrated cassette before and after LAMP amplification. Figure 9a shows that the fluorescent emission from the polycarbonate and from the heater was reasonably low. Figure 9b shows significant emission resulting from the dyed, amplified product. Figure 9c depicts the real-time, fluorescence intensity as a function of time when the initial target analyte consisted of 0 (negative control), 10, 25, 100, and 250 target copies. The fluorescent intensity of the negative control (no target) remains nearly level throughout the reaction, indicating negligible formation, if any, of primer-dimers and the absence of contamination. When the target was present, the signal intensity increased from a baseline of about 1000 relative fluorescent units (RFU) to a saturated level of about 3300 RFU. The threshold time (Tt) is defined as the time that elapses from the start of the heating until the fluorescent signal increases ~20% above the baseline level. Figure 9c indicates that Tt increases as the target molecule’s concentration, C, (DNA copy number/chamber) decreases. Figure 9c (inset) depicts Tt as a function of log(C). Tt decreases nearly linearly as a function of log(C) in the range of 10 < C < 250 copies per reaction chamber. The detection limit of 10 copies was comparable to prior reports on benchtop experiments (Hill et al. 2008).

Although it is quite possible that we could have obtained a lower detection limit, we did not attempt to carry out experiments with smaller numbers of target particles since our sample dilution method could not provide us with a sufficiently accurate estimate of the number of target molecules in the reaction mix at very low target concentrations. We are aware, however, of reports on single-molecule DNA amplification and detection (Zhang and Xing 2010).
To verify the real-time measurements, we subjected the LAMP products to gel electrophoresis. At the conclusion of the reaction, two holes were drilled into the cassette and 3 μL of the LAMP products was removed from each chamber and subjected to electrophoretic, size-based separation. Figure 9d shows the electropherograms of ethidium bromide, stained LAMP products in 2% agarose gel. Lanes 1, 2, 3, 4, and 5 correspond, respectively, to LAMP cassette products with 250, 100, 25, 10, and 0 (negative control) copies. The gel images feature separated bands, which is a characteristic ladder structure associated with LAMP products, indicating that the DNA was successfully amplified in our cassette. In contrast to a PCR product which consists of a single length amplicon and that exhibits a single band in the electropherogram, the LAMP amplicons are connected and the product molecules have different lengths. Therefore, the electropherograms exhibit the ladder pattern. For comparison and as a control, we carried out LAMP amplifications in a bench-top PCR thermocycler (Techne Incorporated, Princeton, NJ). The benchtop experimental results, provided in Fig. 2S, show similar results to the ones obtained with the cassette.

4.4 Detection of RNA targets

Since in many instances, such as in the case of HIV infection, one is interested in detecting RNA viruses at the point of care, we explored the feasibility of carrying out single-step, reverse transcription and amplification (RT-LAMP) in our LAMP cassette. Since our laboratory is not equipped to safely handle viruses, we used the Loopamp RNA amplification kit (Eiken Chemical Co. Ltd., Japan) as the model system. Unfortunately, Eiken Chemical Co. Ltd. does not provide the sequence of primers and the concentration of RNA template. Our experiments were therefore limited to comparing the RT-LAMP products produced in the cassette with the products of a commercial benchtop device.

Figure 10a depicts the real-time RT-LAMP fluorescent intensity as a function of time for a signal resulting from a negative control (no template) and a signal generated by the Loopamp RNA kit. When target is present, the signal rises from a baseline of about 970 RFU to a saturation level of about 3300 RFU.

To verify that the LAMP cassette, indeed, produced amplicons, we extracted a fraction of the products from the cassette and carried out electrophoretic separation in gel. At the end of the reaction, two holes were drilled into the cassette and 3 μL of the LAMP products was removed from the chamber and subjected to electrophoretic, size-based separation. Figure 10b shows electropherograms of ethidium bromide-stained LAMP products in the 2% agarose gel. Lane 1 corresponds to the product of the benchtop PCR cycler (positive control). Lanes 2 and 3 correspond, respectively, to positive control (with target) and negative control (no target) RT-LAMP, cassette products. The electropherograms show the characteristic ladder structure associated with LAMP products, indicating successful amplification in our cassette. Also, note the similarity between the benchtop (lane 1) and the cassette’s (lane 2) amplified products.
5 Conclusion and outlook

A new, single-use, low-cost, disposable, integrated LAMP/RT-LAMP, plastic cassette that utilizes thermo-responsive PDMS valves was developed, allowing isothermal nucleic acid amplification and on-chip, real-time monitoring of the amplification process via fluorescence detection. The device has the potential to be used as a component of a point of care device that can be operated by minimally trained personnel without any need for sophisticated laboratory equipment.

Although the idea of thermo-responsive valves, made with a PDMS-expandable microspheres mixture, was previously described in the literature (Samel et al. 2007a, b), our work demonstrates, for the first time, that the expandability of the thermo-responsive PDMS composite can be controlled by varying the volumetric ratio of the PDMS-base and curing agent and that the effective thermal conductivity of the valve material can be modified with appropriate doping. For a fixed concentration of expandable microspheres, the higher the PDMS base content, the higher the expandability. The paper also presents a new valve design that can withstand backpressures as high as 200 kPa without any observable leakage. The valves made with thermo-responsive PDMS composite are inexpensive, appropriate for disposable devices, easy to fabricate and integrate into functional microfluidic devices, robust, and reproducible. Finally, this is a first report on the integration of the expandable valve with an enzymatic amplification reactor.

Both the LAMP chamber and the valves are actuated (remotely) with thin film heaters. The heaters are part of the instrument that control the cassette and therefore do not add to the cost of the disposable cassette. The amplification results were monitored with a compact, portable ESE fluorescent detector. The use of real-time detection simplifies cassette design and eliminates the need to transfer the reaction products from the amplification reaction chamber into a separate detection chamber. The real-time detection also reduces the analysis time since the test can be stopped as soon as a detectable signal is obtained.

To demonstrate the system’s utility for the detection of DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, we carried out a series of experiments in which we detected *E. coli* DNA and RNA targets, and so on. The real-time detection also reduces the analysis time, and the desired incubation temperature of 63°C, move out of the way, and allow the hydration of the LAMP reagents. Another improvement may include equipping the cassette with a solid state membrane for the isolation and purification of nucleic acid targets (Liu et al. 2011). It would also be desirable to operate with microspheres that expand at temperatures below 60°C so that the same heat source that is used to incubate the isothermal amplification can also be used to actuate the valves.

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### Appendix

Although not central to the manuscript, we estimated the effect of the doping on the thermal conductivity of the PDMS disk. To this end, we model the disk with a one-dimensional, steady-state model. The steady state temperature distribution along the disk’s height is given by

\[
T(x) - T_\infty = \frac{q}{h} \left( 1 + B_i \left( 1 - \frac{x}{H} \right) \right),
\]

where \(T\) is the temperature at distance \(x\) above the heated surface; \(q\) is the heat flux at the heated surface; \(h\) is the heat transfer coefficient at \(x = H\); \(H\) is the height of the doped PDMS disk; \(B_i = \frac{\delta_i}{\delta_0}\) is the Biot number of the doped disk; and \(k_i\) is the thermal conductivity of the doped disk. The temperature ratio \(\theta_i = \frac{T(0)-T_\infty}{T(H)-T_\infty} = 1 + B_i\). The ratio of the thermal conductivities \(\frac{\theta_i}{\theta_0} = \frac{\delta_i}{\delta_0} = 1 + B_i\), where subscripts 0 and \(i\) denote, respectively, the undoped and doped disks.

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