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Reduction of background-triggered amplification in lesion-induced DNA amplification (LIDA)†

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Lesion-induced DNA amplification (LIDA) enables isothermal amplification of nucleic acids, and the only enzyme required is T4 DNA ligase. However, the application of LIDA for the amplification of trace amounts of nucleic acids has been hindered by the observed background-triggered amplification in the absence of the initial target due to a pseudo-blunt end ligation reaction of two of the primers. In this work, we have tested three approaches to minimize the background-triggered amplification: increasing and decreasing the concentration of salts such as NaCl and MgCl₂, respectively, and increasing the concentration of ATP. All these optimizations sharply decreased the background-triggered amplification. Employing the most favourable buffer condition of 2.5 mM MgCl₂ where the target-initiated amplification was least affected while reducing the background-triggered process enabled us to achieve a detection range of 14 nM–140 aM with an approximate limit of detection of 680 aM, which is five orders of magnitude more sensitive than using our standard amplification conditions. This optimization of the salt and co-factor concentrations to decrease the background and enhance the sensitivity of LIDA has demonstrated LIDA's potential for application in clinical diagnostics.

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Introduction

Nucleic acid amplification is an important method for detecting low-abundance nucleic acid sequences in biological samples associated with infectious pathogens or disease states.^{1,2} Polymerase chain reaction (PCR) is by far the most commonly used nucleic acid amplification method owing to its sensitivity.3 However, its dependence on expensive thermal cyclers for amplifying nucleic acids and its requirement for skilled operators make PCR less accessible for usage in resource-limited settings or in point-of-care testing. 4,5 The emergence of isothermal nucleic acid amplification techniques (INAATs) has improved the accessibility of NAATs in lowresource settings as these can be performed at a constant temperature using less equipment making them more convenient and cost-effective. 1,6,7 INAATs like loop-mediated isothermal amplification (LAMP),8,9 strand-displacement amplification (SDA), 10 and exponential-amplification reaction (EXPAR) 11 can amplify target nucleic acid sequences to produce detectable amplicons. For example, within one hour LAMP can detect $(\sim 10^2)$ copies per μ L of target, but it demands extensive primer design and high temperatures between 60 and 65 °C.9 Strand-

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displacement amplification (SDA) has potential for amplifying DNA with a nearly $\sim 10^7$ fold amplification factor under 2 hours,^{5,10} while the exponential amplification reaction (EXPAR) displays 10⁶-10⁸ amplification efficiency under 30 min. 11 However, the latter two methods employ two enzymes DNA polymerase and nicking endonuclease enzyme and are performed at 37 °C (ref. 10 and 12) and 60 °C, 11 respectively. Our group has developed an isothermal nucleic acid amplification process that uses only one enzyme based on a ligase chain reaction (LCR) rather than a polymerase chain reaction. 13,14 Traditionally, LCR is a highly sensitive exponential amplification technique that utilizes DNA ligase and needs a thermocycler for amplification. 15,16 Our approach to achieving DNA amplification isothermally involves introducing a destabilizing abasic lesion in one of the ligating primers that facilitates spontaneous turnover in the ligase chain reaction. This process is called lesion-induced DNA amplification (LIDA). 17 Our previous work has shown that LIDA can operate at room temperature, alleviating the need for temperature control. 17,18

In LIDA, one of the primers complementary to the target sequence, DNA-I, contains a destabilizing model abasic group (1',2'-dideoxy-5'-phosphate) at its terminus. The other primer is complementary to the target at the sequence adjacent to the destabilizing primer, leading to the formation of a nicked duplex (Fig. 1). The enzyme T4 DNA ligase can then catalyze the formation of a novel template through the ligation of these two primers (DNA-II, Fig. 1). Due to the presence of the abasic group on DNA-II, the original template DNA-I spontaneously

[†] Electronic supplementary information (ESI) available: DNA sequences, experimental details, graphical presentation and gel images. See DOI: https://doi.org/10.1039/d5an00047e

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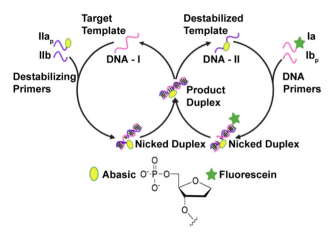


Fig. 1 Schematic of self-replication of DNA-I by lesion-induced DNA amplification (LIDA) from four primers (Ia, Ibp, IIap, and IIb). The IIap strand contains the destabilizing abasic group at the 5'-phosphate position. In typical LIDA, la is the limiting primer reagent, which is labeled with fluorescein (green star). The formation of DNA-I is measured by fluorescence imaging coupled with polyacrylamide gel electrophoresis.

dissociates. Subsequently, DNA-II functions as a template for hybridization with two additional primers that constitute the DNA-I sequence, one of which includes a fluorescent label. This results in the formation of another nicked duplex, which is subsequently ligated by T4 DNA ligase to form a fluorescently labeled copy of DNA-I. Once again, the duplex spontaneously dissociates due to the destabilization caused by the presence of the abasic group releasing DNA-I and DNA-II and allowing both to template more reactions. This cross-catalytic cycle leads to rapid, exponential self-replication of an 18 nt target and has been used to amplify a variety sequences. 14,17-19 By employing a serial-dilution strategy with low concentration of the primers in a cross-catalytic approach we were able to detect as low as 140 fM of target DNA. 14 In more recent work, lesion-induced DNA amplification has been utilized to detect RNA targets by incorporating an RNA-templated step where cDNA is generated, isothermally liberated and then subsequently amplified by LIDA. Using reverse transcription (RT-) LIDA with optimized primer concentrations we achieved a detection limit of 240 amol (32 pM) at 28 °C of target RNA spiked into samples of total cellular RNA.18

However, one major limitation of lesion-induced DNA amplification (LIDA) is the presence of background-triggered amplification that occurs in the absence of the target, which is observed in all the sequences that we successfully amplified by LIDA. 14,17-19 This background-triggered amplification leads to: (i) false positives by incorrectly indicating the presence of the target in the biological sample²⁰ and (ii) limited analytical sensitivity of the assay (too high of a limit of detection, LOD) leading to false negatives. 21 The background triggered reaction in LIDA results from T4 DNA ligase catalyzed ligation of the primers in the absence of the initial target to form DNA-II via a blunt-end like (pseudo blunt-end) ligation (Fig. 2)^{14,17,18} Recently, we determined that it is specifically due to the nontemplated formation of DNA-II (ligation of the 5'-phosphate

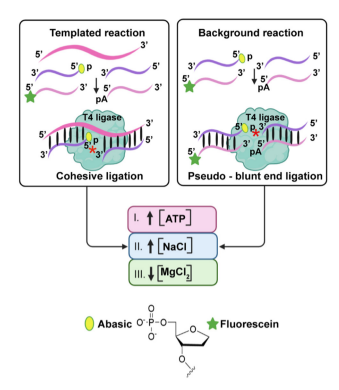


Fig. 2 Schematic illustration of the cohesive (templated reaction) and pseudo-blunt end ligations (background reaction) by the T4 DNA ligase involved in LIDA and the reported factors that help to reduce blunt end ligation, which we hypothesize would reduce background-triggered amplification in LIDA. *Shows the site of phosphodiester formation. Created in BioRender.com.

abasic primer) in the first cycle, which is a slow process but results in a template that can amplify by cross-catalysis.²²

Studies of T4 DNA ligase have shown that increasing the concentration of the enzyme co-factor ATP and decreasing the concentration of Mg²⁺ can inhibit the joining of blunt ends with only slight reduction in the rate of the templated ligation.²³ Other studies have also shown that increasing the concentration of monovalent salts like NaCl can inhibit both blunt end and cohesive ligation, albeit to different extents.²⁴ Therefore, we hypothesized that the factors that influence blunt end ligation by T4 DNA ligase would influence the pseudo-blunt end ligation that causes background-triggered LIDA (Fig. 2). Here we show how the background triggered process can be minimized by varying the concentration of ATP, NaCl, and Mg²⁺ leading to a reduction in the LOD of our standard LIDA process by five orders of magnitude. This enormous improvement in sensitivity opens the door for the use of LIDA in point-of-care testing.

Experimental section

Materials and instrumentation

Chemicals were purchased from Fisher Scientific or Sigma-Aldrich and used as received. T4 DNA ligase enzyme (2 000 000 cohesive end units per mL, catalog #M0202T) and ATP **Paper**

(10 mM, catalog #9804S) were purchased from New England Biolabs. For making 15% polyacrylamide denaturing gels, 40% acrylamide/bis-solution 19:1 was purchased from Bio-Rad (California, USA). The ligation master mix was prepared immediately before the experiment from separate solutions of TRIS-HCl (pH 7.5) (containing an appropriate amount of MgCl $_2$) and ATP and NaCl. Ultrapure water was obtained from a Milli-Q-Ultrapure Water System, which was autoclaved and used for the experiments.

The oligonucleotide synthesis was performed on an Applied Biosystems Model 392 DNA/RNA Synthesizer. A Torrey Pines Scientific Echotherm Chilling/Heating Plate (Model IC22) was used to maintain temperature for the experiments. 15% denaturing gels were made using 8 M urea. The gels were imaged using an Image Quant RT ECL Imager from GE Healthcare Life Science by UV transillumination. Igor Pro v7.08A (Wavemetrics, Oswego, OR) was used to fit the data with a sigmoidal growth function. We employed Axygen 0.6 mL Maxymum Recovery Snaplock Polypropylene Microcentrifuge tubes (MCT-060-L-C) for our experiments.

Oligonucleotide preparation and characterization

All the oligonucleotides listed in Table S1† were synthesized on an Applied Biosystems Model 392 automated solid-phase synthesizer using Glen Research reagents. All synthesized DNA was purified with Glen-Pak Purification cartridges (catalog # 60-5200-01) following the DMT-ON protocol and freeze-dried afterward.

Some of the DNA strands had special modifications with compounds such as 5'-phosphate, fluorescein, and the abasic group. The abasic group was incorporated using dSpacer CE phosphoramidite (catalog #10-1914-90) and fluorescein using 5'-fluorescein-dT phosphoramidite (catalog #10-1056-95). Solid phosphorylation reagent II (catalog #10-1902-90) was used to synthesize 5'-phosphate strands.

The synthesized strands were characterized by MALDI-TOF in a linear negative mode using a Voyager Elite time of flight-mass spectrometer (Applied Biosystems, Foster City, CA) following our earlier work.²² Purity was assessed by visualizing the oligonucleotides with Stains-All reagent after it was run on a 15% polyacrylamide denaturing gel and analyzed using the Image Quant RT ECL Imager from GE healthcare Life Science (with the fluorescein filter).

LIDA protocol

The isothermal self-replication experiments were performed as described in our earlier work 13,14 with some modifications. For 15 μL total reaction volume, we used 1 μL of T4 DNA ligase (2 000 000 CEU mL^{-1}). The final concentration of the three primers (Ibp, IIap, and IIb) was 2.8 μM and that of the limiting fluorescent primer (Ia) was 1.4 μM . The reaction mixtures were prepared by mixing four primers with the template (template-initiated reaction) and without the template (background-triggered reaction) to reach a volume of 10 μL . The DNA solution was incubated for 10 minutes at 30 °C. Each component of the ligation buffer and the enzyme were then added to different spots near the top of the Eppendorf tube containing the DNA.

The sample was then spun down and vortexed, thereby initiating the reaction. 1.5 µL aliquots of the reaction mixture were taken at various time points and placed in a microcentrifuge tube containing 2 µL of a mixture of bromophenol blue dye $(0.011 \text{ g mL}^{-1})$ and EDTA (0.5 M). The samples were analyzed using a 15% PAGE denaturing gel. The target concentration variation experiments were performed in the same way using the same primer ratios as mentioned above, but with various lower concentrations of the DNA target (14 nM, 140 pM, 14 pM, 14 fM, and 140 aM). A master mix containing each component of the ligation buffer and the enzyme was prepared and incubated for 10 minutes at 30 °C. To minimize any potential error or variation arising from different enzyme bottles, we combined 10 µL from each of the two T4 DNA ligase enzyme (2 000 000 CEU mL⁻¹) bottles. This pooled enzyme solution was then used to perform two replicate target concentration variation experiments to ensure consistency and reliability in the enzymatic activity across both experimental replicates.

Results and discussion

T4 DNA ligase catalyzes the formation of a phosphodiester bond between the 5'-phosphate and 3'-hydroxyl groups of two adjacent primers hybridized to a template strand.²⁵ This enzyme can also perform this bond sealing with blunt end duplex substrates. T4 DNA ligase requires divalent metal cations such as Mg²⁺ or Mn²⁺ and is dependent on ATP as an adenylate-donating co-factor that provides energy for ligation.²⁶ Ligation of nicked DNA involves three catalytic steps: (1) adenylation of the ligase at a lysine, which releases pyrophosphate (PPi) and results in the formation of a covalent ligase-adenosine monophosphate (AMP) complex; (2) transfer of AMP onto the 5'-phosphate of the nicked DNA, activating it for nucleophilic attack by the 3'-hydroxy group from the adjacent DNA strand; and (3) formation of the phosphodiester bond releasing the AMP. 26,27 For blunt-end ligation, the mechanism is virtually the same except that the enzyme can adenylate the 5'-phosphate on one duplex prior to the binding of the duplex containing the 3'-hydroxy group.²⁸

In the presence of the target, LIDA proceeds by cohesive ligation of the destabilizing primer strands hybridized to the target template to generate DNA-II (Fig. 2). In the absence of the target, however, ligation can occur because of the single-base overhang between the 5'-phosphate abasic primer and the 5'-phosphate adenosine primer, which also results in the formation of DNA-II. We refer to this mechanism of ligation to form DNA-II as a pseudo-blunt end ligation as there is no hydrogen-bonding in this single A:Abasic overhang (Fig. 2). We found that it occurs for LIDA-based amplification of all sequences that we have explored using T4 DNA ligase. ^{14,17,18}

Effect of increasing ATP on target-initiated and background-triggered LIDA

As the sensitivity of LIDA is impacted by the background amplification that arises from the *in situ* synthesis of a cata-

lytic amount of DNA-II by pseudo-blunt end ligation, our goal was to identify experimental conditions that decreased background-triggered amplification, thereby enhancing the sensitivity of LIDA. Studies have shown that an increase in ATP concentration decreases blunt-end ligation. Accordingly, we hypothesized that increasing the ATP concentration beyond our standard LIDA conditions (1 mM ATP) would adversely affect the background-triggered LIDA, leading to a shift to the right of the sigmoidal amplification profile corresponding to LIDA without the initial target.

The effect of increasing ATP concentration on LIDA in the absence and presence of the initial DNA-I target is presented in Fig. 3. For simplicity, we utilized standard primers that operate best at 30 °C where only one primer contains a destabilizing (abasic) group rather than our dual-destabilization primers that can be tuned to operate at lower temperatures.¹⁷ We also used a temperature controller to ensure that any changes in the rate of amplification could be attributed to changes in the buffer conditions. First, we evaluated the impact of ATP concentration on the target-initiated LIDA. Accordingly, a 14 nM DNA-I target was incubated with an excess of primers at varying ATP concentrations in the presence of T4 DNA ligase and the conversion of the fluorescent primer to a DNA-I product was monitored by gel electrophoresis. Increasing the ATP concentration from 1 mM to 7.5 mM did not have a significant effect on the amplification rate for the LIDA reactions initiated by 14 nM DNA-I (Fig. 3A). In contrast, the onset of amplification for the background-triggered LIDA in the absence of the initial DNA-I target slowed down with an increase in ATP concentration, supporting our hypothesis (Fig. 3B). The observed trends in the template-initiated and background-triggered LIDA are consistent with the observations by Ferretti and co-workers where they reported a significant decrease in the extent of blunt-end joining with an

increase in ATP concentration with only minimal reduction in the joining of the cohesive-ends by T4 DNA ligase.²³ However, at an ATP concentration of 10 mM, complete inhibition of LIDA was observed in both the absence and presence of the initial DNA-I target. Such a strong nonlinear effect from varying the ATP concentration as shown in Fig. 3A has been observed in T4DNA ligase catalyzed cohesive end ligation in the percent yield, 23 as well as in the enzyme turnover rate. 26 Moreover, in these reports, 10 mM ATP either completely shut down cohesive end ligation²³ or inhibited it significantly,²⁶ consistent with our observations. This hindering of ligation by excessive ATP has been attributed to the dissociation of the ligase from the DNA complex after the transfer of AMP. In the presence of excess ATP, the ligase can be readenylated prior to rebinding to the DNA complex disabling its ability to catalyze the ligation of the adenylated DNA intermediate (App DNA), leading to the accumulation of the latter. 23,26

Effect of increasing monovalent ion concentration on targetinitiated and background-triggered LIDA

Another approach to inhibit the activity of T4 DNA ligase in a manner that affects blunt end vs. cohesive ligation more is to introduce K⁺ or Na⁺ into the ligation mixture. Studies have shown that increasing monovalent ion concentrations to ionic strengths much higher than those provided by the presence of Mg²⁺ (vide infra) can inhibit the activity of the T4 DNA ligase albeit to varying extents on cohesive and blunt end substrates.²⁴ Therefore, we hypothesized that introducing NaCl would decrease the background-triggered amplification more so than the template-initiated LIDA reaction owing to its deleterious effect on the pseudo-blunt end ligation of the primers. LIDA was performed under our standard conditions with the addition of NaCl at various concentrations (Fig. 4).

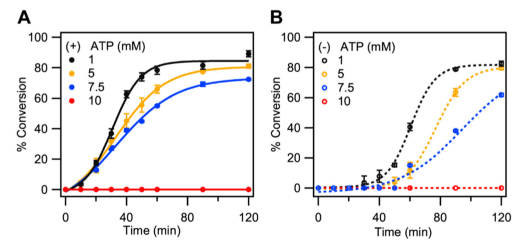


Fig. 3 Effect of increased ATP concentration on the kinetics of cross-catalytic formation of F-DNA-I initiated by (A) 14 nM and (B) 0 nM DNA-I in the presence of different ATP concentrations. % conversion refers to the amount of fluorescent primer converted to DNA-I. Standard experimental conditions: 14 nM DNA-I, 1.4 μ M Ia, 2.8 μ M IIa_P, 2.8 μ M IIb_P, 2.8 μ M IIb, 50 mM TRIS-HCl (pH 7.5), 10 mM MgCl₂, 1 mM ATP, 2000 CEU T4 DNA ligase, and 30 °C.



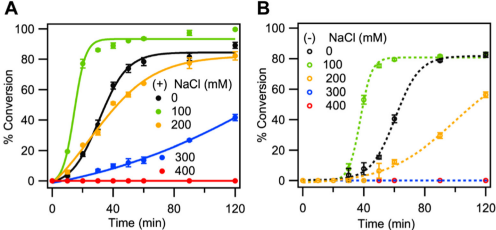


Fig. 4 Effect of NaCl concentration on the kinetics of LIDA initiated by (A) 14 nM and (B) 0 nM DNA-I using otherwise standard conditions. % conversion refers to the amount of fluorescent primer converted to DNA-I.

With 100 mM NaCl, we observed faster target initiated and background-triggered amplification compared to the reaction without any added NaCl. This increase in activity is attributed to the shielding by the monovalent sodium cations that stabilizes the hybridized DNA complexes via mitigation of the negative charge on the sugar-phosphate backbone.²⁹ However, T4 DNA ligase activity started to decline at salt concentrations above 200 mM, consistent with previous T4 DNA activity studies.30,31 We observed that at a 200 mM concentration both the target-initiated and background-triggered LIDA declined with a greater impact on the latter, suggesting that pseudoblunt end ligation was more compromised compared with cohesive ligation. This trend of slower amplification with increasing salt concentration continued at 300 mM, and by 400 mM no amplification was observed within two hours. The inhibitory effect of monovalent salts at higher concentration on T4 DNA ligase has been attributed to a decreased affinity of the enzyme-AMP towards the DNA resulting in a higher effective $K_{\rm m}$.^{30,31} This proposed decrease in ligase affinity for the DNA substrate is thought to stem from neutralization of the negative charges on the phosphate group that play an important role in binding of the DNA to the T4 ligase.^{29,30}

Effect of increasing divalent ion concentration on targetinitiated and background-triggered LIDA

In addition to ATP, the ligase requires a divalent metal cation such as Mg²⁺ or Mn²⁺. Studies have shown that a decrease in Mg²⁺ concentration results in the decreased joining or full inhibition of ligation of blunt ends, while only exhibiting a minimal decrease in the joining of the cohesive ends.²³ Accordingly, we hypothesized that decreasing the Mg²⁺ concentration below our standard LIDA conditions (10 mM MgCl₂) would adversely affect background-triggered LIDA. The effect of decreasing Mg²⁺ concentration on the target-initiated and background-triggered reactions is presented in Fig. 5.

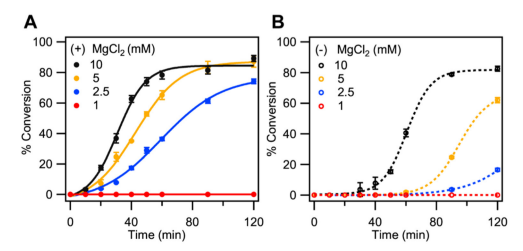


Fig. 5 Effect of decreasing MgCl₂ concentration on the kinetics of cross-catalytic formation of F-DNA-I initiated by (A) 14 nM and (B) 0 nM DNA-I under otherwise standard conditions. *% conversion* refers to the amount of fluorescent primer converted to DNA-I.

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Decreasing the Mg²⁺ concentration from 10 mM to 2.5 mM did not have a significant effect on the amplification rate for the target-initiated LIDA reactions (Fig. 5A) while a more significant effect was observed for the background-triggered process (Fig. 5B). A further decrease in Mg²⁺ concentration to 1.0 mM caused complete inhibition of both cohesive and blunt ends by the T4 DNA ligase. The observed trends in the templated and background reactions are consistent with the observations by Ferretti and co-workers, where they reported a significant decrease in the extent of blunt end joining with a decrease in Mg²⁺ concentration with only minimal reduction in the joining of the cohesive-ends by T4 DNA ligase.²³ The decrease in blunt end joining in the presence of low Mg2+ could be attributed to the abortive ligation that occurs at lower Mg2+ concentrations also leading to the accumulation of the App DNA. 26,32

Sensitivity of LIDA at 2.5 mM Mg²⁺

From the previous studies we found that the most optimal conditions for significantly mitigating the background-triggered process while still facilitating rapid LIDA in the presence of the initial target were by decreasing the concentration of MgCl₂ to 2.5 mM. To determine if these conditions with reduced Mg²⁺ that favor target-initiated LIDA improved the sensitivity of our assay, we performed LIDA at 2.5 mM MgCl₂ for a wide range of the initial DNA-I target covering eight orders of magnitude (Fig. 6A). Remarkably, by using this simple change in ligation mixture composition, we were able to distinguish as little as 140 aM of DNA-I versus the background-triggered process (Fig. 6A). In comparison, using the same primer concentrations and buffer but with 10 mM Mg²⁺ we can only distinguish down to 140 pM (Fig. S4†).

We next considered how to quantify the differences in kinetic traces as a function of the initial DNA-I target. The utilization of time-to-threshold is a common parameter in isothermal amplification assays to determine the presence of a target.33-35 Using a related approach reported in our previous work, 18 here we identified the point-of-inflection (POI) for each kinetic trace by fitting a sigmoidal curve to the data (Fig. 6A) and determining the difference in POI between the targetinitiated and background-triggered amplification measured with the same enzyme master mix (Fig. 6B, black trace). The average ΔPOI from two replicates versus the -logarithm of initial DNA concentration $(-\log C)$ gave a linear trend with a line-of-best fit of $\triangle POI = -9.47 \pm 1.41 \ (-\log C) + 157 \pm 17 \ (r^2$ value = 0.93) (Fig. 6B, black trace). From the line of best fit, we calculated the x-intercept to be 16.58 ± 1.41, which corresponds to the $-\log C$ value when ΔPOI is equal to 0. The red arrow over the black trace in Fig. 6B marks one σ greater than the x-intercept value ($-\log C$ of 15.17), which is equal to a concentration of 680 aM, and represents a conservative estimate of the limit of detection. However, we could discriminate as low as 140 aM ($-\log C$ of 15.85) initial DNA target from the background-triggered process based on a discernible ΔPOI for both replicates (Fig. 6B-black trace). Similar analysis for the targetdependent LIDA amplification using standard (10 mM) Mg²⁺ concentrations prevented us from discriminating the 14 pM

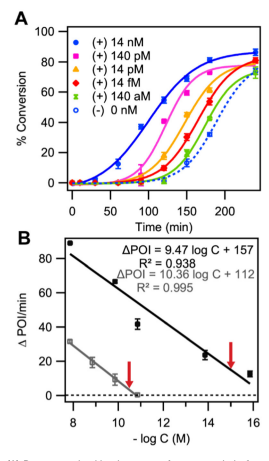


Fig. 6 (A) Representative kinetic traces of cross-catalytic formation of DNA-I at 30 °C initiated by different concentrations of the DNA-I target with 2.5 mM Mg²⁺ (B) The difference in point-of-inflection for kinetic traces with and without initial target DNA-I (Δ POI) as a function of the -log of the initial DNA-I concentration at both Mg²⁺ concentrations, standard 10 mM (grey) and optimized 2.5 mM (black) Mg²⁺. The red arrows indicate the approximate LOD for both Mg²⁺ concentrations, with grey and black lines representing 10 mM and 2.5 mM Mg²⁺ concentrations, respectively.

target from the background-triggered reaction, resulting in an approximate LOD of 55 pM (-log C of 10.26 Fig. 6B-grey trace). Thus, reducing the Mg²⁺ concentration led to an improvement of at least five orders of magnitude in sensitivity. These data reveal that optimizing the buffer conditions to suppress the background-triggered pseudo-blunt end process in LIDA significantly improves the assay sensitivity, underscoring the substantial impact of buffer conditions on the assay.

Conclusions

Lesion-induced DNA amplification (LIDA) represents a simple isothermal method for self-replicating 18-nucleotide DNA target sequences with an uncomplicated primer design that can be tuned to operate at room temperature making it well suited for point-of-care. Until now, this method has had a major drawback, which is the presence of a background-trig**Paper** Analyst

gered process that limited the sensitivity of LIDA. Here we explored conditions known to minimize the rate of blunt end ligation, examining the effects of mono (Na⁺) and divalent (Mg^{2+}) salt concentration as well as ATP concentration with the goal of delaying the onset of the LIDA background.

The results showed that increasing ATP concentration above 1 mM and NaCl concentration above 100 mM greatly diminished the background-triggered process. However, decreasing the concentration of MgCl2 had the most significant effect on the background-triggered process while still allowing for fast target-initiated amplification. Using the optimized conditions of 2.5 mM MgCl₂ enabled us to achieve a detection range of 14 nM to 140 aM, with a limit of detection of approximately 680 aM. This represents an improvement of five orders of magnitude in comparison with that of LIDA under our standard conditions. With this improvement, we have achieved detection down to clinically relevant concentrations (140 aM is 1200 copies in 15 μ L) paving the way to the application of LIDA to clinical diagnostics. Although the assay demonstrates high sensitivity, the longer reaction times remain a limitation. Future work will involve testing the optimized assay at room temperature with complex, real-world samples.

Author contributions

Anantha S. Ealeswarapu: conceptualization, methodology, investigation, resources, and writing - original draft. Nahida Akter: validation of experimental data. Julianne M. Gibbs: supervision, conceptualization, and writing - review & editing.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare no competing financial interest.

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References

- 1 Y. Zhao, F. Chen, Q. Li, L. Wang and C. Fan, Chem. Rev., 2015, 115, 12491-12545.
- 2 P. T. Monis and S. Giglio, Infect., Genet. Evol., 2006, 6, 2-12.
- 3 L. Garibyan and N. Avashia, J. Invest. Dermatol., 2013, 133, 1-4.
- 4 M. Fakruddin, K. S. B. Mannan, A. Chowdhury, R. M. Mazumdar, Md. N. Hossain, S. Islam and Md. A. Chowdhury, J. Pharm. BioAllied Sci., 2013, 5, 245-252.
- 5 P. Srivastava and D. Prasad, 3 Biotech, 2023, 13, 200.
- 6 P. Craw and W. Balachandran, Lab Chip, 2012, 12, 2469-
- 7 M. M. Islam and D. Koirala, Anal. Chim. Acta, 2022, 1209, 339338.
- 8 M. Soroka, B. Wasowicz and A. Rymaszewska, Cells, 2021, 10, 1931.
- 9 V. J. Gadkar, D. M. Goldfarb, S. Gantt and P. A. G. Tilley, Sci. Rep., 2018, 8, 5548.
- 10 G. T. Walker, M. S. Fraiser, J. L. Schram, M. C. Little, J. G. Nadeau and D. P. Malinowski, Nucleic Acids Res., 1992, 20, 1691-1696.
- 11 M. S. Reid, R. E. Paliwoda, H. Zhang and X. C. Le, Anal. Chem., 2018, 90, 11033-11039.
- 12 J. G. Nadeau, J. B. Pitner, C. P. Linn, J. L. Schram, C. H. Dean and C. M. Nycz, Anal. Biochem., 1999, 276, 177-187.
- 13 A. Kausar, R. D. McKay, J. Lam, R. S. Bhogal, A. Y. Tang and J. M. Gibbs-Davis, Angew. Chem., Int. Ed., 2011, 50, 8922-8926.
- 14 A. Kausar, C. J. Mitran, Y. Li and J. M. Gibbs-Davis, Angew. Chem., Int. Ed., 2013, 52, 10577-10581.
- 15 F. Barany, Genome Res., 1991, 1, 5-16.
- 16 Y. Li, X. Wang, M. Wang, M. Liu, H. Wang, W. Xia and L. Liu, Mol. Cell. Biochem., 2023, 478, 1621-1631.
- 17 B. S. Alladin-Mustan, C. J. Mitran and J. M. Gibbs-Davis, Chem. Commun., 2015, 51, 9101-9104.
- 18 B. S. Alladin-Mustan, Y. Liu, Y. Li, D. R. Q. De Almeida, J. Yuzik, C. F. Mendes and J. M. Gibbs, Anal. Chim. Acta, 2021, 1149, 238130.
- 19 A. Kausar, E. A. Osman, T. Gadzikwa and J. M. Gibbs-Davis, Analyst, 2016, 141, 4272-4277.
- 20 D. S. Mouliou and K. I. Gourgoulianis, Expert Rev. Respir. Med., 2021, 15, 993-1002.
- 21 J. Pugh, D. Wilkinson and J. Savulescu, J. Med. Ethics, 2022, 48, 329-333.
- 22 H. Park, S. Parshotam, S. C. Hales, A. K. Mittermaier and J. M. Gibbs, *Chem. - Eur. J.*, 2023, **29**, e202300080.
- 23 L. Ferretti and V. Sgaramella, Nucleic Acids Res., 1981, 9, 3695-3705.

- 24 K. Hayashi, M. Nakazawa, Y. Ishizaki and A. Obayashi, *Nucleic Acids Res.*, 1985, 13, 3261–3271.
- 25 I. R. Lehman, Science, 1974, 186, 790-797.
- 26 A. V. Cherepanov and S. De Vries, *Eur. J. Biochem.*, 2003, 270, 4315–4325.
- 27 A. V. Cherepanov and S. De Vries, *Eur. J. Biochem.*, 2002, **269**, 5993–5999.
- 28 R. Rossi, A. Montecucco, G. Ciarrocchi and G. Biamonti, *Nucleic Acids Res.*, 1997, **25**, 2106–2113.
- 29 S. El-Ashram, I. Al Nasr and X. Suo, *Biotechnol. Rep.*, 2016, **12**, 33–39.

- 30 A. J. Raae, R. K. Kleppe and K. Kleppe, *Eur. J. Biochem.*, 1975, **60**, 437–443.
- 31 D. Y. Wu and R. B. Wallace, Gene, 1989, 76, 245-254.
- 32 M. R. Taylor, J. A. Conrad, D. Wahl and P. J. O'Brien, *J. Biol. Chem.*, 2011, **286**, 23054–23062.
- 33 B. Tian, G. A. S. Minero, J. Fock, M. Dufva and M. F. Hansen, *Nucleic Acids Res.*, 2020, **48**, e30.
- 34 E. J. H. Wee and M. Trau, ACS Sens., 2016, 1, 670-675.
- 35 J. C. Rolando, E. Jue, N. G. Schoepp and R. F. Ismagilov, *Anal. Chem.*, 2019, **91**, 1034–1042.