Evaluation of a Loop-Mediated Isothermal Amplification Technique for the Rapid Visual Detection of *Hepatozoon canis* Infection

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**Abstract**

**Background** Laboratory diagnosis of *Hepatozoon canis* infection is tedious, especially in chronic and/or latent infections.

**Purpose** The study was planned to develop a simple read out loop mediated isothermal amplification (LAMP) assay targeting a partial 18S rRNA gene of *H. canis* with naked eye visualisation of LAMP products.

**Methods** A LAMP assay was employed to assess the DNA amplification by adding SYBR Green I dye for naked eye inspection of DNA accumulating in reaction tubes. Positive amplification was read through observation of change in colour of reaction mixture following addition of dye. The visual results were further verified with those of agarose gel electrophoresis. Genomic DNA of other haemoparasites of dog viz. *Babesia vogeli*, *B. gibsoni*, *Ehrlichia canis* and *Trypanosoma evansi* along with no-template control were used to determine the specificity of assay.

**Results** Among the 109 blood samples presented at Small Animal Clinics, Teaching Veterinary Clinical Complex, Guru Angad Dev Veterinary and Animal Sciences University, Ludhiana, Punjab (India) tested, 39 revealed colour change from orange to green indicating positive reaction while 70 were negative as revealed by no colour change. The results of visual inspection were comparable to those obtained by agarose gel electrophoresis. The LAMP primers specifically amplified *H. canis* DNA, whereas no amplification was detected in DNA samples of other haemoparasites and no-template control revealing specificity of the assay. The diagnostic sensitivity and specificity (95% CI) of visual LAMP assay with respect to microscopy in detection of *H. canis* varied from 100% (15.81–100.00%) and 65.42% (55.61–74.35%), respectively.

**Conclusion** The present investigation has developed a specific and rapid LAMP assay for the detection of *H. canis*, using SYBR Green I dye, which has practical applications for the screening of field samples.

**Keywords** *Hepatozoon canis* · LAMP assay · SYBR Green I dye · Visual inspection

**Introduction**

*Hepatozoon canis* is one of the most common tick transmitted apicomplexan protozoan parasite with worldwide distribution [1, 2]. The parasite is transmitted by ingestion of the brown dog tick, *Rhipicephalus sanguineus* sensu lato containing the mature oocysts [3]. Generally the infections are asymptomatic to mild, but may also become severe and fatal [4]. The clinical findings including anorexia, lethargy, fever, cachexia, pale mucous membranes and hind limb paralysis [5].

On diagnostic front, conventional parasitological examination viz. detection of ellipsoidal gamonts within the neutrophils or monocytes, in stained blood smears by microscopy and/or histopathological visualization of meronts or monozoic cysts in tissues are employed [6]. Serodiagnostic
tests, like enzyme linked immunosorbent assay and indirect fluorescent antibody test have also been utilized for detection of parasite specific antibodies [7, 8]. In recent times, nucleic acid based diagnostic techniques viz. polymerase chain reaction (PCR), real time-PCR assay and sequence analysis have been utilized worldwide for the diagnosis of H. canis infection and specification of the isolates, respectively [2, 9, 10] as they offer high levels of specificity and sensitivity in both, hosts (dogs) as well as vectors (ticks) [2, 11, 12].

The loop-mediated isothermal amplification (LAMP) method in the last decade has been used for early and sensitive diagnosis of various infectious agents [13–16]. The assay has advantages over other molecular tests in terms of simplicity, higher throughput amplification efficiency and ease of evaluation of results by visualization of the turbidity in positive reactions [17]. In our previous study, a LAMP assay was developed, for the first time, targeting the partial 18S rRNA gene of H. canis which revealed high levels of sensitivity, specificity and threshold detection over the traditional PCR assay and microscopy when applied on blood samples collected from dogs [16]. The present study was planned to develop a simple read out LAMP technology by naked eye visualization using SYBR Green I dye for interpretation of the LAMP results.

**Materials and Methods**

**Ethical Guidelines**

Approval and necessary guidelines of Institute Animal Ethics Committee (IAEC) was obtained GADVASU/2017/IAEC/39/12 vide Memo no. IAEC/2017/734-760 dated 20.03.2017 for conduction of the study.

**Samples**

The blood samples (n = 109) were collected randomly from the dogs presented in the Small Animal Clinics, Teaching Veterinary Clinical Complex, Guru Angad Dev Veterinary and Animal Sciences University, Ludhiana, Punjab (India). These animals were suspected for haemoprotozoan infections and blood collected for conduction of diagnostic tests was used for DNA isolation in the study.

**LAMP Assay**

The LAMP assay standardized in our previous study was employed on the isolated whole blood genomic DNA from the collected blood samples [16]. In brief, the reaction mixture consisted of 5 pmol of each outer primer (F3 and B3), 50 pmol of each inner primer (FIP and BIP), 0.6 M betaine (Sigma, USA), 6 mM MgSO$_4$ (New England Biolabs, UK), 1.6 mM dNTP mix (MBI Fermentas, USA), 1× Thermopol reaction buffer (20 mM Tris–HCl, 10 mM (NH$_4$)$_2$SO$_4$, 10 mM KCl, 2 mM MgSO$_4$, 0.1% Triton X-100) and 1 µL of target DNA. The reaction mixture was heated at 95 °C for 5 min, chilled on ice and subsequently 1 µL (8U) Bst DNA polymerase large fragment (New England Biolabs, UK) was added. The amplification reaction was carried out at 55 °C for 90 min and terminated by incubating at 80 °C for 2 min. Genomic DNA isolated from an infection free puppy and nuclease free water were used as negative and no-template control, respectively, while DNA isolated from a microscopically confirmed H. canis positive sample was used as positive control in all the runs.

**Naked Eye Visualization of LAMP Products**

The DNA amplification was assessed by adding 1.0 µL of the fluorescent intercalating SYBR Green I (Invitrogen) dye for naked eye inspection of DNA accumulating in the reaction tubes by visual fluorescence. The positive amplification was read through observation of change in colour of the reaction mixture following addition of dye to the tube. The results were further verified by electrophoresis of LAMP products in 2% ethidium bromide-stained agarose gel (Agarose Low EEO, SRL), and visualized using InGenius® Gel Documentation System (Syngene, UK). The specificity of the visual LAMP assay was determined by utilizing the genomic DNA extracted from whole blood of dogs infected with other haemoparasites viz. B. vogeli, B. gibsoni, E. canis and T. evansi along with no-template control. Further, the results of visual LAMP assay were compared with those of microscopy to determine the diagnostic sensitivity and specificity of the assay.

**Statistical Analysis**

The data obtained from visual LAMP assay and microscopy (gold standard test) were analyzed with the help of SAS software (SAS 9.3 Version). McNemar’s test $P$ value, specificity and sensitivity with 95% confidence interval (CI) were calculated.

**Results**

To evaluate the usefulness of the optimized LAMP assay as a rapid diagnostic tool, the whole blood genomic DNA extracted from the collected blood samples (n = 109) were tested. An observation of colour change following the addition of 1 µL of SYBR Green I dye was visually detected in the PCR tubes. In case of positive amplification, the original orange colour of the dye changed into green (n = 39) while in case of no amplification, the original orange colour of the
Dye was retained \((n=70)\), revealing 35.77\% positivity for *H. canis* infection (Fig. 1). It was observed that results of the agarose gel electrophoresis for the same samples were comparable with that of visual examination as revealed by characteristic ladder-like multiple bands in the positive samples (Fig. 2). The LAMP primers specifically amplified *H. canis* DNA (positive control), whereas no amplification was detected in DNA samples from dogs infected with *B. vogeli*, *B. gibsoni*, *E. canis*, *T. evansi*, and no-template control revealing specificity of the assay (Fig. 3).

The developed visual LAMP assay detected significantly higher number of samples positive for *H. canis* in comparison to microscopy \((\chi^2 \text{ value } = 35.027; \ P \text{ value } \leq 0.0001)\). The diagnostic sensitivity and specificity \((95\% \text{ CI})\) of visual LAMP assay with respect to microscopy in detection of *H. canis* varied from 100\% \((15.81–100.00\%)\) and 65.42\% \((55.61–74.35\%)\), respectively. In the present study the LAMP assay seemed to be more sensitive than microscopy as those samples of blood \((n=37)\) which were negative in microscopy were also amplified for the presence of parasite DNA.

**Discussion**

To facilitate the application of LAMP assay, monitoring of amplification can be carried out with naked eye inspection either in the form of visual turbidity or visual fluorescence [18]. However, the techniques relying on indirect detection methods, i.e. turbidity and colorimetry, may not be able to distinguish between real and false positives under some unexpected cases when non-specific amplification occurs [19]. In this regard, the monitoring may be done by use of fluorescent dyes viz. SYBR Green I. In the presence of sufficient amount of double stranded DNA, the color of SYBR

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**Fig. 1** Field application of visual LAMP assay for detection of *H. canis*. Positives reactions: tube no. 2, 3, and 12. Negative reactions: tube no. 1, 4–11, and 13.

**Fig. 2** Field application of LAMP assay for detection of *H. canis*. Lane M: Generuler™ 100 bp Ladder. Lane P: positive control. Lane 7: negative control. Lane 1–6, 8: field samples.

**Fig. 3** Specificity of visual LAMP assay for detection of *H. canis*. Tube 1: negative control. Tube 2: *H. canis* DNA. Tube 3: *B. vogeli* DNA. Tube 4: *B. gibsoni* DNA. Tube 5: *E. canis* DNA. Tube 6: *T. evansi* DNA.
Green I turns from orange to green which is apparent under natural as well as ultraviolet light [19, 20] making it ideal for easy monitoring through naked eye. Furthermore, the employment of SYBR Green I lead to increased sensitivity when compared with visual turbidity measurements as per Zhang et al. [20] and Soli et al. [21].

In practice, the visual inspection for amplification is performed through observation of color change by adding 1 μL of SYBR Green I to the tube and the result of the LAMP reaction can be monitored by the naked eye at the end point [20–23]. Although, the risk of contamination of amplicons may increase because of the need to open the reaction tubes to add the dye, however, opening the reaction tube after amplification with care to prevent carry-over contamination can be done to minimize the risk. On the other hand, agarose gel electrophoresis is a conventional method for monitoring the LAMP amplicons directly [13, 18, 19, 24], and even serves as a “gold standard” methodology in many situations. The results are interpreted as appearance of ladder-like banding in positive reaction tube under ultraviolet (UV) gel imaging analysis system. However, ethidium bromide, used in gel electrophoresis, may act as mutagen, carcinogen, or teratogen, though this depends on the organism exposed and the circumstances of exposure [25]. Additionally, the method possesses a risk of cross-contamination and also relatively long turnaround times [26]. Also, the requirement for electrophoresis apparatus and UV detection limits the suitability for field applications.

Conclusions

In the current investigation, a LAMP assay targeting the partial 18S rRNA gene of *H. canis* was evaluated for its application in field conditions, by monitoring the amplification with naked eye inspection in the form of visual fluorescence using the fluorescent intercalating dye SYBR Green I.

Therefore, it may be concluded that a simple and rapid LAMP assay with use of SYBR Green I dye eliminates the agarose gel electrophoresis used for documentation of results and thus is a rapid and time saving simple diagnostic technique for detection of *H. canis* infection in dogs. Furthermore, the LAMP assay can be employed as a simple read out technology (naked eye visualisation of LAMP products) in the field for routine examination, where sophisticated and high-end equipments are not feasible.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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