

TrueLAMP™ Instruction Manual

General Information

Description

TrueLAMP™ is a proprietary colorimetric LAMP reagent system consisting of 2× TrueLAMP buffer and TrueLAMP polymerase for rapid, high-specificity nucleic acid amplification. TrueLAMP chemistry suppresses primer-driven amplification in the absence of template, preventing nonspecific or false-positive reactions in no-template controls (NTCs). The liquid format provides flexibility for custom reaction design and scalable workflows while maintaining robust inhibition control.

Specification

- Product Name: TrueLAMP™
- Catalog #: TL0001S, TL0001M, TL0001L
- Contents:
 - TL0001S: 0.5 mL 2× TrueLAMP buffer and 5 µL TrueLAMP polymerase
 - TL0001M: 1.5 mL 2× TrueLAMP buffer and 15 µL TrueLAMP polymerase
 - TL0001L: 5 × (1.5 mL 2× TrueLAMP buffer and 15 µL TrueLAMP polymerase)

Storage

Store kit at –20 °C until use. Thaw 2x buffer at 22 °C for at least 30 min and vortex before use. After thawing, store 2x buffer at 22 °C tightly capped and use within a week or return it to –20 °C.

Terms & Conditions

- Product Usage: For in vitro laboratory research use only (RUO). Not for administration to humans or use in medical diagnosis.
- Warranties and Liabilities: Boltii Diagnostics Inc. accepts no responsibility and shall not be held liable for any loss, damage, or expenses—whether consequential or incidental—including damage to property, persons, or premises, arising from the use, results of use, or inability to use this product. Boltii Diagnostics Inc. makes no warranties, expressed or implied, including but not limited to warranties of merchantability or fitness for a particular purpose.

Patents, Trademarks & Copyrights

TrueLAMP™ is a trademark of Boltii Diagnostics Inc.

© 2026 Boltii Diagnostics Inc. All rights reserved.

TrueLAMP™ Assay Protocol

1. Prepare Reaction Mix

Prepare enough for 8 reactions at 10 µL each. Due to the small volumes, it's critical to use calibrated pipettes. Use immediately.

Component	1 Reaction (µL)	8 Reactions (µL)	+10% extra (µL)	Total (µL)
Nuclease-free water	3	24	2.4	26.4
10× LAMP primers	1	8	0.8	8.8
2× TrueLAMP buffer	5	40	4	44
TrueLAMP polymerase	0.05	0.4	0.04	0.44

2. Reaction Setup

- 1) Label each tube in an 8-strip 0.2 mL clear PCR tube set.
- 2) Add:
 - 1 µL nuclease-free water to NTC tube
 - 1 µL positive template (~500 copies, pre-validated) to PTC tube
 - 1 µL test sample to each test tube
- 3) Add 9 µL reaction mix to each tube.
- 4) Cap the tubes, vortex, and centrifuge the 8-strip briefly to remove bubbles.

3. Amplification

Incubate in submerged water bath or smart coffee mug or digital oven or thermocycler with heated lid:

- 60 °C for 10 min, then
- 65 °C for 50–80 min

4. Detection

Colorimetric Readout:

- Endpoint: photograph after incubation
- Semi-real-time: photograph every 15 min without moving camera

Quantify using smartphone colorimeter app:

- RGB mode → minimum green value
- CMYK mode → maximum magenta value

Expected Results:

NTC: red

PTC: yellow

Frequently Asked Questions

1. Should the kit be stored at -20 °C?

Store kit at -20 °C until use. Thaw 2x buffer at 22 °C for at least 30 min and vortex before use. After thawing, store 2x buffer at 22 °C tightly capped and use within a week or return it to -20 °C.

2. What incubator works best?

Uniform heating is critical to avoid evaporation and condensation in 10 µL reactions. Recommended: Fully submerged water bath (best for multiple reaction strips) or temperature-controlled smart coffee mug (POC) or thermocycler with heated lid. Convection oven may also work. Do not use heat block.

3. Why is my NTC positive?

1) Recalibrate pipettes. Incorrect volumes (< 4.75 µL buffer per 10 µL reaction) may reduce inhibition. 2) Verify incubation temperature. Nonspecific amplification can occur at ≤ 63 °C after ~30 min. 3) Increase buffer slightly (e.g., 5.5 µL 2x buffer per 10 µL reaction) for stubborn primer sets.

4. Why did my positive control fail?

Template concentration may be below LOD. Determine LOD for each primer set.

5. Can raw samples (e.g., saliva, plasma) be used?

Purified DNA/RNA is recommended. TrueLAMP is sensitive to pH and ionic strength. Water-suspended cells, bacteria or virions may be compatible after validation.

6. Can polymerase be premixed with 2x buffer for storage?

No. The polymerase and 2x buffer must be stored as separate components. Once combined, polymerase activity declines gradually as it binds to the inhibitor in the buffer, and this effect is irreversible. Prepare the master mix fresh and proceed to amplification within 30 minutes of assembly.

7. Why is a 60 °C pre-incubation needed and how long should reactions run?

An initial 58-61 °C 10 min pre-incubation followed by 65 °C for 50-80 min improves amplification consistency across DNA and RNA targets, primer sets, and incubation systems (oven, bath, thermocycler). Amplification is generally visible by 30 min and reaches maximum by 90 min. Strict endpoint timing is not required. Extended incubation does not typically alter endpoint color.

8. How should color be quantified?

Use a smartphone colorimeter app. Sample the region just below the liquid meniscus with the smallest aperture and record: minimum green (RGB mode) or maximum magenta (CMYK mode).

9. Do I need to determine differential time to threshold (DTT)?

No. Because NTCs remain reliably negative regardless of incubation duration, result interpretation does not depend on when the reaction is read. A simple visual or smartphone color comparison at any convenient endpoint is sufficient. This eliminates the need for DTT analysis that conventional LAMP requires to avoid false-positive interference.