

MBPCR125

Hi-PCR® Babesia bigemina SYBr PCR Kit

Description

Bovine babesiosis is a tick-borne disease of cattle caused by the protozoan parasites of the genus *Babesia* (Apicomplexa: Piroplasmorida) leading to significant economic losses in tropical and subtropical regions. More than 100 species of *Babesia* are recognized to infect domestic animals, wild animals, and humans worldwide. The principal species of *Babesia* that cause Bovine babesiosis are *Babesia bigemina*, *Babesia bovis* and *Babesia divergens*. *Babesia bigemina*, a major *Babesia* species responsible for bovine babesiosis is transmitted by *Rhipicephalus spp.* ticks in tropical and subtropical countries. Clinical manifestations of disease associated with *Bovine babesiosis* includes fever and intravascular hemolysis leading to progressive anemia, hemoglobinuria, and jaundice, and may result in death. The diagnostic methods of bovine babesiosis have relied mostly on clinical signs, microscopic examination of the parasites and serological testing for antibody detection. The advent of molecular diagnostic tests like polymerase chain reaction (PCR) has paved the way to efficient diagnosis than the conventional techniques. The Hi-PCR® *Babesia bigemina* SYBr PCR Kit is designed for specific and sensitive detection of *Babesia bigemina*.

NOTE: Hi-PCR® *Babesia bigemina* SYBr PCR Kit is for *in-vitro* use only.

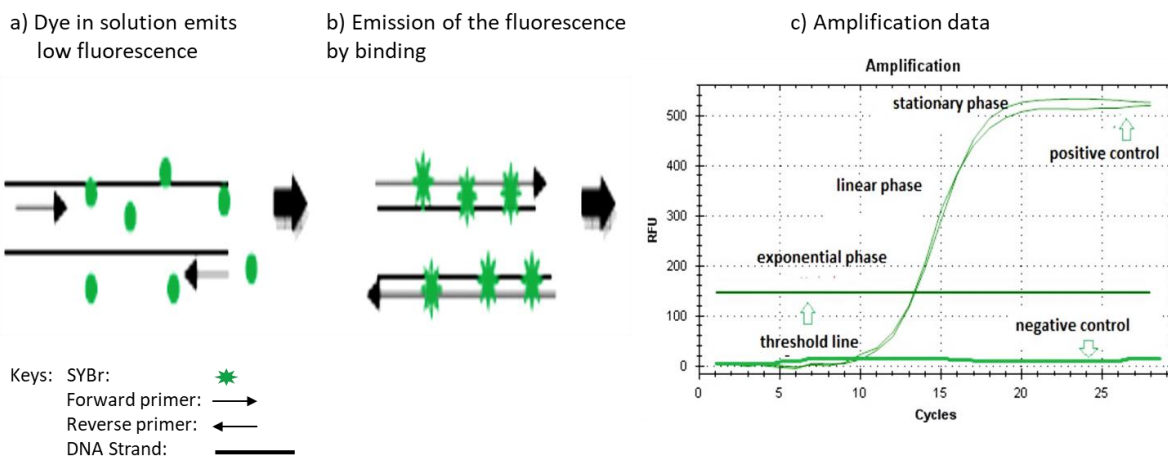
Intended Use

The Hi-PCR® *Babesia bigemina* SYBr PCR Kit is intended for use by qualified clinical laboratory personnel trained in the techniques of PCR and in vitro diagnostic procedures. The kit is recommended for sensitive and specific detection of *Babesia bigemina* in clinical samples.

Principle

Real-time PCR also called quantitative PCR (qPCR) or kinetic PCR, is a laboratory technique based on the principle of PCR. This technique is used to amplify and simultaneously quantitate a targeted DNA sequence. The presence of SYBr Green Dye, a dsDNA-binding dye, in the Hi-SYBr Master Mix allows for simplified assay design without the need for additional fluorescent probes and enables assay verification using a melt-curve analysis. Hi-PCR® *Babesia bigemina* SYBr PCR Kit is a qualitative Real-Time PCR kit that targets 18S rRNA gene of *Babesia bigemina* species with the amplification product size of 165 bp.

Diagrammatic representation of SYBr Green Chemistry in Real-Time PCR:



The SYBr Green dye cycles between an unbound (Denaturation step) and a bound (Annealing through Extension) state as the reaction progresses. Signal intensity increases as the quantity of amplicons increase in later cycles indicating amplification. During elongation, more and more dye molecules bind to the newly synthesized DNA.

If the reaction is monitored continuously, an increase in fluorescence is viewed in real-time. Upon denaturation of the DNA for the next heating cycle, the dye molecules are released and the fluorescence signal falls.

Positive control

A Positive control (PC) is a control reaction which contain the target DNA template that the PCR is designed to amplify. It is used to ensure proper and intended functioning of all the PCR reagents and is recommended to be used in every run to assess optimal assay performance.

Negative Control

A Negative Template Control (NTC) is essential to verify that the reagents, equipment, and environment used in the assay are free from contamination with target nucleic acid. In this control reaction, nuclease-free water is used as a template. It is recommended to include at least one negative template control reaction per run to ensure the reliability of the results.

Features

- Fast and simple
- Sensitive and specific results
- Guaranteed reproducible results
- Includes all reagents and controls

Sample Source: Blood samples

Storage and Shelf-life

The provided kit has a shelf-life of 12 months when stored between -10°C and -20°C. Repeated thawing and freezing of PCR reagents should be avoided as this may reduce the sensitivity. If reagents are to be used multiple times, we recommend storing reagents as aliquots to avoid repeated freeze and thaw. Degradation of sample DNA specimens can also reduce the sensitivity of the assay. HiMedia does not recommend using the kit after the expiry date stated on the pack.

Kit Contents

The provided PCR Kit contains:

Components	Product code	Reagents provided for (reactions)*	
		25R	50R
B. bigemina SYBr master mix	DS2685	270	540
<i>B. bigemina</i> SYBr Primer Mix	DS2686	27	54
<i>B. bigemina</i> Positive Control	DS0272B	70	140
Molecular Biology Grade water for PCR	ML065	70	140

*For a 25 µL PCR reaction

Materials needed but not provided:

Product name	Product Code
Real-Time PCR Instrument and equipment	
Thermal Cycler	LA948 / LA949 / LA950 / LA1006 / LA1015/ LA1059 / LA1060 / LA1066
TabSpin™ Microcentrifuge	LA1089/LA1090
Automated nucleic acid extraction system and materials	
Insta NX® Instrument - fully automated nucleic acid purification system utilizing the Innovative Super -S membrane column method	LA1056
Insta NX® Mag16, Insta NX® Mag16 ^{Plus}	LA1118, MBLA018
Insta NX® Mag32, Insta NX® Mag32 ^{Plus}	LA1096, MBLA019
Insta NX® Mag96	LA1097

Extraction Kits	
HiPurA® Pre-filled Clinical Multi-purpose Magnetic Nucleic Acid Purification kit (Cartridges)	MB583PC16200
HiPurA® Pre-filled Clinical Multi-purpose Magnetic Nucleic Acid Purification kit (Plates)	MB583MPF16200
HiPurA® DNA/ RNA Purification Kit	MB583
Tubes, plates and other consumables	
Varivol II Micropipettes (Capacity: 0.5 to 10 µL/10 to 100 µL/200 to 1000 µL)	LA611/LA614/LA615
µPet Autoclavable Micropipettes (Capacity: 0.5 - 10 µL/10 - 100 µL/20 - 200 µL/100 - 1000 µL)	LA955/LA958/LA959/LA960
Q4Pet Autoclavable Micropipette (Capacity: 0.5 to 10 µL/10 to 100 µL/100 - 1000 µL)	MBLA009/MBLA011/MBLA008
Barrier Tips, Maximum capacity 10 µL	LA749A
Barrier Tips, Maximum capacity 200 µL	LA751A
Barrier Tips, Maximum capacity 1000 µL	LA859A
8-strip tubes & optically clear flat caps for PCR	PR17, PR22, PR23
PCR Tubes, 0.1mL, 0.2 mL; PCR Plates	PW1255/PR2/PR3/PR19
Optical Sealing film	PR18

Specimen collection and Handling

Follow appropriate techniques for handling specimens; after use, contaminated materials must be sterilized by autoclaving before discarding. Standard precautions as per established guidelines should be followed while handling clinical specimens and items contaminated with blood and other body fluids. Safety guidelines may be referred to individual safety data sheets.

General Preparation Instructions

- Before use, a suitable amount of all PCR components should be completely thawed on ice (4°C).
- Perform the amplification reactions in a clean area.
- Use of aerosol barrier pipette tips is recommended to reduce contamination risks from extraneous DNA templates.
- Extract and store positive control material (if used) separately from all other reagents to avoid contamination and add it to the reaction mix in a separate area.
- Centrifuge the components briefly once thawed.

Protocol for PCR Master Mix Preparation

1. In the “Master mix Preparation” area, thaw all components from the kit on ice, mix by inverting the tubes and centrifuge the reagents for 5 seconds. Keep on ice for later use.
2. Based on the total number of specimens (including PC and NTC) to be tested (N), calculate the volume of the components to be added as **N X volume of “1R”**
3. Use 1.5 mL Nuclease free centrifuge tube(s) for the preparation of the PCR reaction mix. Refer the following table. After all the reagents are added, mix them thoroughly and centrifuge for 5 seconds.

Components	Product code	Volume for “1R” (one reaction)
Preparation of PCR Reaction Mix		
<i>B. bigemina</i> SYBr master mix	DS2685	10.0 µL
<i>B. bigemina</i> SYBr Primer Mix	DS2686	1.0 µL
Total PCR Reaction Mix	-	11.0 µL
Template addition		
Template (Extracted DNA)		14.0 µL
Total reaction volume	-	25.0 µL

4. Aliquot **11 µL** of **PCR reaction mix** into 0.1/0.2mL PCR tube/plate/strips, compatible to the PCR instrument to be used.
5. In the “Nucleic acid handling” area, add **14 µL** of extracted nucleic acid of test specimen into the plate/strip to respective wells.
6. For positive and negative control, template nucleic acid is replaced by Positive control mix and nuclease free water respectively. Refer the following table.

Set up of Positive controls (PC) for the PCR run		
Components	Product code	Volume for “1R” (one reaction)
PCR Reaction Mix	-	11.0 µL
<i>B. bigemina</i> Positive Control	DS0272B	14.0 µL
Total reaction volume	-	25.0 µL

Set up of Negative Template controls (NTC) for the PCR run		
Components	Product code	Volume for “1R” (one reaction)
PCR Reaction Mix	-	11.0 µL
Molecular Biology Grade water for PCR	ML065	14.0 µL
Total reaction volume	-	25.0 µL

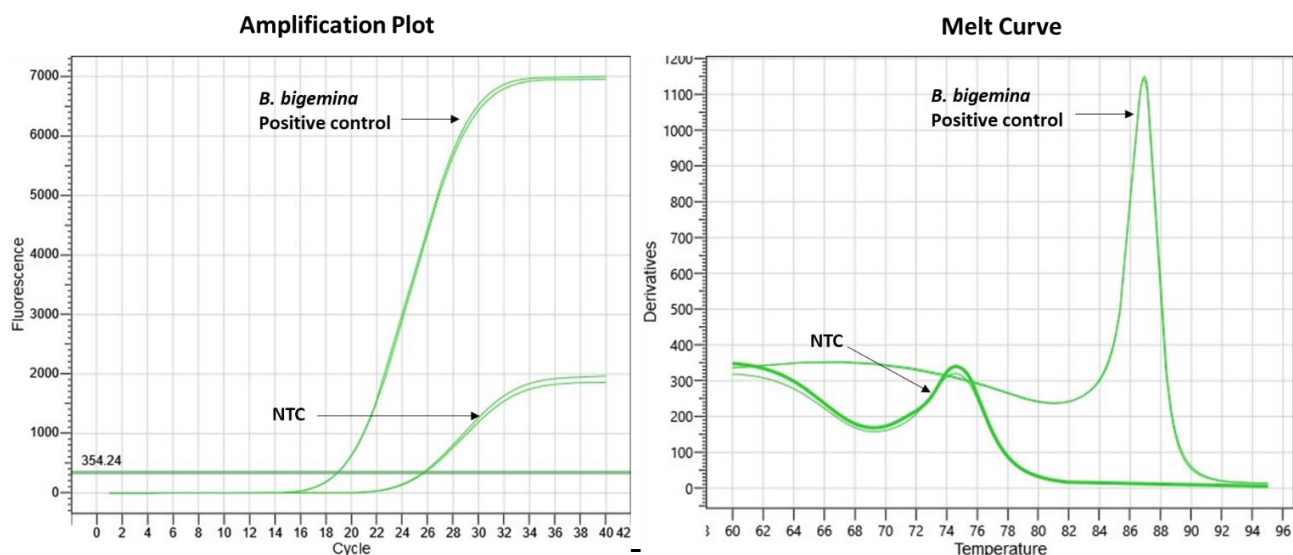
7. Tightly cap the tubes/strips or seal the plate using an optically clear adhesive film.
8. Centrifuge the tube briefly at 6000 rpm for about 10 seconds.
9. Place the tubes in Real-time PCR machine and set the recommended PCR program (mentioned below).

Recommended PCR program

1. Initial denaturation : 95°C for 5 minutes No. of cycles: 1
2. Denaturation : 95°C for 30 seconds
3. Annealing (Plate Read) : 67°C for 30 seconds } No. of cycles: 40
4. Melt Curve Analysis as per HiMedia’s Insta Q96 Real Time PCR Machine
 - a. 95°C : 15 seconds
 - b. 60°C : 1 minute
 - c. 95°C : 15 seconds
 - d. Increment : 0.5°C
 - e. On Hold : 10 seconds

NOTE: The user can also set up a melt curve program as per their existing PCR instrument.

Amplification data



Sr. No.	Sample	C _t value	T _m (°C)
1	<i>B. bigemina</i> Positive control (PC)	19.07	86.9
2	Negative Template control (NTC)	25.8	74.6

Image representing real-time amplification data of *Babesia bigemina* with Ct values (provided in table)

Data Interpretation

Melting Temperature (T _m)*	Result Interpretation
84.5°C-88.5°C	Positive for <i>Babesia bigemina</i>

* T_m values can slightly vary for different sample types. If the T_m values show significant variation from those mentioned in the above table, then the sample is considered to be negative for *Babesia bigemina*.

Warning and Precautions

Not for Medicinal Use. Read the procedure carefully before starting the experiment. Wear protective gloves/protective clothing/eye protection/face protection. Follow good clinical laboratory practices while handling clinical samples. Standard precautions should be followed as per established guidelines. Safety guidelines may be referred in safety data sheets of the product.

Performance and Evaluation

Each lot of Hi-PCR® *Babesia bigemina* SYBr PCR Kit is tested against predetermined specifications to ensure consistent product quality.

Quality Control

Each lot of HiMedia's Hi-PCR® *Babesia bigemina* SYBr PCR Kit is functionally tested in amplification assays. The Hi-PCR® *Babesia bigemina* SYBr PCR Kit provides reagents for controls - *B. bigemina* Positive Control (PC) and No Template Control (NTC) which are to be included in every run.

Troubleshooting Guide

Sr. No.	Problem	Cause	Solution
1.	No amplification	Degraded samples	1. Check the integrity of DNA using agarose gel electrophoresis. 2. Use freshly prepared DNA to ensure the availability of intact template sequence for efficient amplification.
		Error in protocol setup	Check the PCR conditions in the machine and make appropriate changes wherever required
2.	Variability between replicates	Error in reaction set-up	1. Verify that the correct reagent volumes, dilutions and storage conditions have been used. 2. Prepare large volume reaction mix, vortex thoroughly and aliquot appropriately into reaction tubes.
		Air bubbles in reaction mix	Briefly centrifuge reaction samples/plate prior to running on a PCR machine.
		Pipetting error	Replicates can show increased variation due to poor laboratory techniques or imprecise pipettes.
3.	Amplification in negative control	Reagents contaminated	1. Replace all critical solutions. 2. Repeat the analysis of all tests with fresh aliquots of critical reagents.

Safety Information

The Hi-PCR® Babesia bigemina SYBr PCR Kit is for laboratory use only, not for drug, household or other uses. Take appropriate laboratory safety measures and wear gloves when handling.




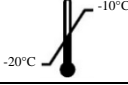




Disposal

The user must ensure safe disposal by autoclaving and/or incineration of used or unusable preparations of this product. Follow established laboratory procedures in disposing of infectious materials and material that comes into contact with clinical sample must be decontaminated and disposed of in accordance with current laboratory techniques.

Technical Assistance

At HiMedia, we pride ourselves on the quality and availability of our technical support. For any kind of technical assistance, mail at mb@himedialabs.com.

Symbols:

	Manufacturer		Do not use if package is damaged
	Batch code		Temperature limit
	Date of manufacture (YYYY-MM)		Consult instructions for use
	Use-by date (YYYY-MM)		Catalogue number

Identification No.: PIMBPCR125

Rev.No.:06

Date of Issue: 2025-10

Disclaimer :

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