

# ***PACKIT***

## ***Brucella abortus***

### **Reagent Set**

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**For *Brucella abortus* Detection**

# **User Manual**

**For Research Use Only**

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## **INTENDED USE**

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**POCKIT™** *Brucella abortus* Reagent Set uses insulated isothermal polymerase chain reaction (iiPCR) technology (Chang et al., 2012; Tsai et al., 2012) to detect the specific nucleic acid sequences of *Brucella abortus*. This reagent set is specially designed to be used on an iiPCR-compatible instrument, **POCKIT™** Nucleic Acid Analyzer. The intended users of the reagent set are veterinarians or technicians who have basic laboratory skills.

This reagent set is intended for research purpose and *in vitro* use only.

## **SUMMARY AND EXPLANATION**

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*Brucella abortus* (*B. abortus*) is a gram-negative bacterium that causes premature abortion of a cattle fetus. *B. abortus* is usually transmitted by contacting with the placenta, fetus, fetal fluids, and vaginal discharges from infected animals. This disease primarily affects cattle and causes abortion and infertility, but it can also be transmitted to humans through consumption of contaminated dairy products or direct contact with an infected animal (Corbel & Elberg et al, 2006). It is a highly contagious zoonosis pathogen and can lead to great economic loss especially in the dairy and agricultural industry.

## PRINCIPLES OF THE PROCEDURE

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The assay is based on iPCR for qualitative detection of *Brucella abortus*. Fluorogenic probe hydrolysis chemistry is used to generate fluorescent signal when a specific DNA sequence of *Brucella abortus* is amplified. The primers and probe target specific nucleic acid sequences between *alkB* gene and IS711 in *Brucella abortus*, and do not react with host genomic DNA and nucleic acids of other pathogens.

## PRODUCT DESCRIPTION

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### A. Materials Provided

#### 1) POCKIT™ *Brucella abortus* Reagent Set, 48 tests/package

Component	Contents or Purpose	Amount
Premix Pack	<ul style="list-style-type: none"> <li>■ <i>Brucella abortus</i> Premix (lyophilized pellet) containing dNTPs, primers, probe, and enzyme for amplification.</li> <li>■ Desiccating agent pack.</li> </ul>	6 bags (8 <i>Brucella abortus</i> Premix vials and 1 desiccating agent/bag)
Premix Buffer B	<ul style="list-style-type: none"> <li>■ Reaction buffer to re-dissolve the lyophilized pellet.</li> </ul>	2 vials (1.3 ml/vial)
P(+) Control	<ul style="list-style-type: none"> <li>■ Dried plasmid containing <i>Brucella abortus</i> partial sequence as positive control.</li> </ul>	1 vial
P(+) Control Buffer	<ul style="list-style-type: none"> <li>■ Reaction buffer to re-dissolve P(+) Control.</li> </ul>	1 vial (110 µl/vial)
User Manual		1 copy

- 2) **R-tube**, 48 tubes/pack

## **B. Materials and Equipment Required, but Not Provided**

- 1) **PetNAD™** Nucleic Acid Co-prep Kit or **taco™ mini** Automatic Nucleic Acid Extraction System (optional)
- 2) **POCKIT™** Nucleic Acid Analyzer: the iiPCR-compatible instrument.
- 3) **cube™** Mini-Centrifuge
- 4) Micropipette and tips

## **C. Storage and Stability**

- 1) The reagent set should be stored at 4°C and is stable until the expiration date stated on the label.
- 2) Store Premix vials in sealed Premix Pack to avoid hydration of lyophilized components.
- 3) Reconstituted P(+) Control is stable for 6 months at 4°C. Aliquot reconstituted P(+) Control to avoid degradation of nucleic acid.

## **PRECAUTIONS**

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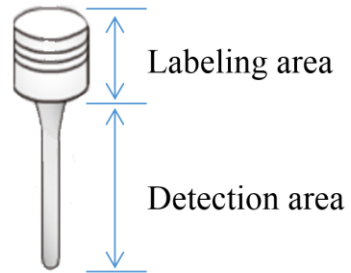
- A. Do not open R-tube(s) after reaction to prevent any carryover contamination.
- B. Perform extraction and amplification in two independent spaces to minimize contamination.
- C. Do not reuse R-tube and Premix.

D. Include the P(+) Control to:

- 1) Ensure **POCKIT™** Nucleic Acid Analyzer is working.
- 2) Ensure detection reagent performance after storage.

E. To get optimal fluorescence detection.

- 1) Wear powder-free gloves to handle R-tubes.
- 2) Do not label in the detection area of R-tube.



## LIMITATIONS

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- A.** The test should only be used for testing nucleic acid extracts. Do not add specimens directly into the Premix.
- B.** **PetNAD™** Nucleic Acid Co-prep Kit or **taco™ mini** Automatic Nucleic Acid Extraction System is recommended for nucleic acid extraction.
- C.** Any deviation from the recommended procedures may lead to sub-optimal results. Performance of the modified protocol should be validated by the users.
- D.** It is strongly recommended to use freshly prepared nucleic acids (within 1 hour after extraction) to achieve optimal results with **POCKIT™** *Brucella abortus* Reagent Set.

## SAMPLE TYPE

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This detection reagent is intended for analyzing nucleic acids extracted from whole blood, vaginal swab, and raw milk samples.

## OPERATION PROCEDURE

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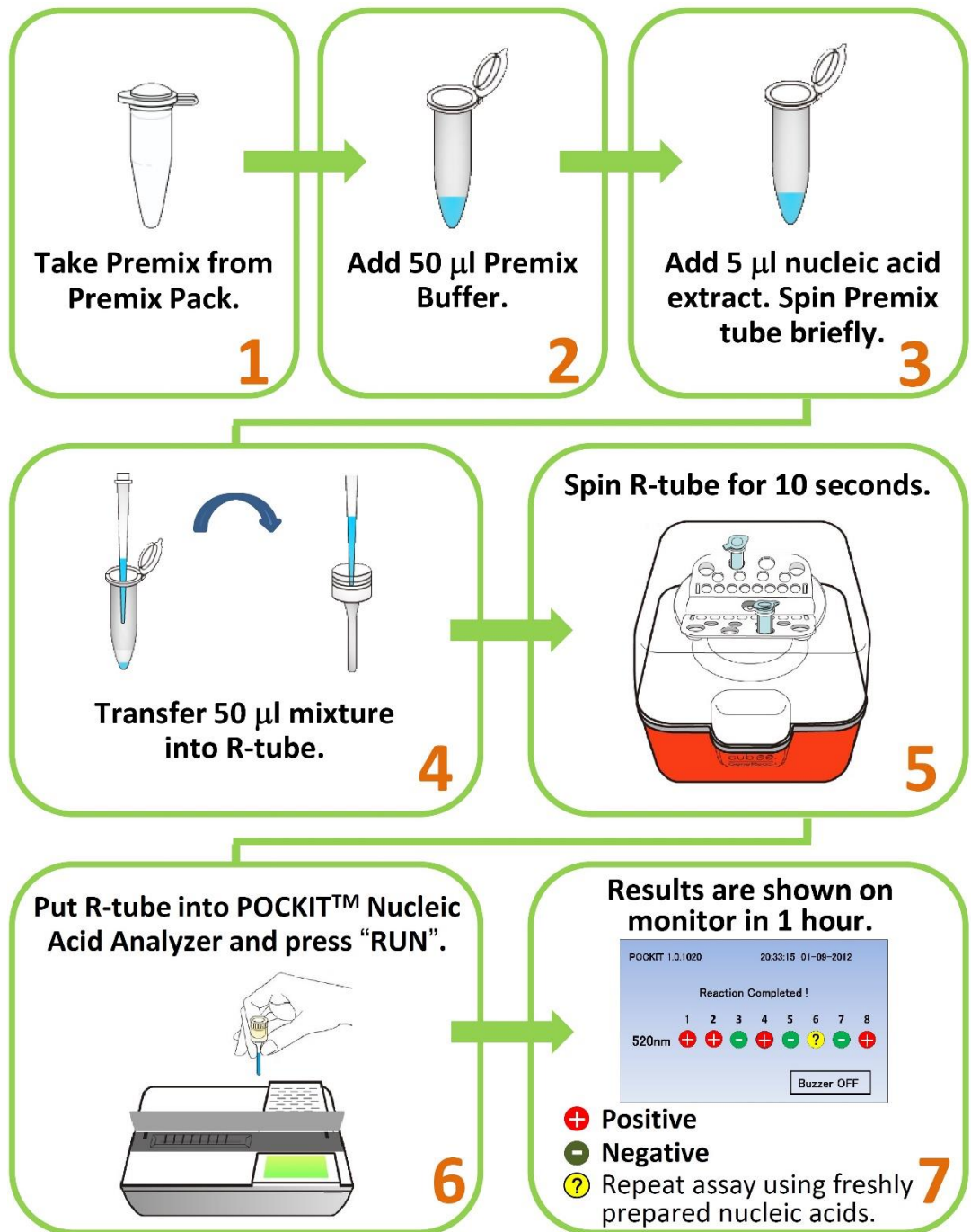
### A. Using POCKIT™ *Brucella abortus* Reagent Set

- **Note: Before preparing the reactions for iiPCR testing, turn on POCKIT™ Nucleic Acid Analyzer to initiate the calibration for the instrument. The device will complete self-test within 5 minutes. Please refer to the user manual of POCKIT™ Nucleic Acid Analyzer for further details.**
  - **Note: Before using for the first time, add 100 µl P(+) Control Buffer to P(+) Control. Store reconstituted P(+) Control at 4°C.**
- 1) Label R-tube(s) in the labeling area.
  - 2) Prepare one Premix for each sample. (Premix tubes are in Premix Pack. Each Premix Pack contains eight Premix tubes.)
    - **Note: When the pellet is not found at the bottom of the tube, spin tube briefly to bring it down.**
  - 3) Add 50 µl Premix Buffer B to each Premix tube.
  - 4) Add 5 µl nucleic acid extract or dissolved P(+) Control to each Premix tube. Spin Premix tubes briefly in a mini centrifuge (such as **cubee™** Mini-Centrifuge).



- 5) Transfer 50 µl Premix/sample mixture into R-tube.
- 6) Seal top of each R-tube with a cap. Make sure R-tube is capped tightly.
- 7) Place R-tube into the holder of **POCKIT™** Nucleic Acid Analyzer.
- 8) Spin tube/holder set briefly in **cubee™** Mini-Centrifuge to make sure all solution is collected at the bottom of R-tube.
  - **Note: Make sure there are no bubbles in the solution.**
  - **Note: Start reaction within 1 hour to reduce the risks of nucleic acid degradation and non-specific reaction.**
- 9) **POCKIT™** Nucleic Acid Analyzer reaction:
  - a) Select "520 nm".
  - b) When "System READY" is displayed, place the holder with R-tube(s) into the reaction chamber.
  - c) Tap cap of each R-tube to make sure the tube is positioned properly.
- 10) Close lid and press "Run" to start reaction program.
- 11) Test results are shown on the monitor after the reaction is complete.

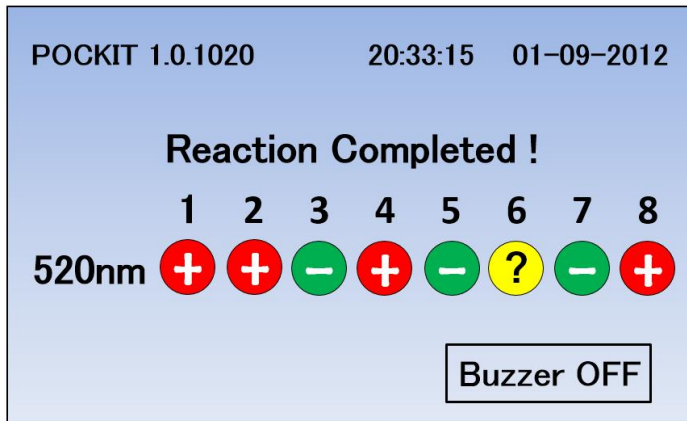
## B. Quick Guide



## DATA INTERPRETATION

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\*One example of results shown on the monitor.



520 nm	Interpretation
	<i>Brucella abortus</i> Positive
	<i>Brucella abortus</i> Negative
	Repeat reaction with freshly prepared nucleic acid.

## ANALYTICAL SENSITIVITY

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The detection limit of **POCKIT™** *Brucella abortus* Reagent Set is about 10 copies/ reaction.

## TROUBLESHOOTING

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Problems	Possible causes	Solutions
<b>False Positive</b>	1) Reuse of micro-centrifuge tubes, tips, R-tubes and Premix.	<ul style="list-style-type: none"> <li>■ Micro-centrifuge tubes, tips, R-tubes and Premix are for single-use only. Reusing these accessories would cause cross-contamination.</li> <li>■ Used micro-centrifuge tubes, tips, R-tubes and Premix should be collected and discarded according to local regulation. Do not place the waste close to the working area to prevent cross-contamination.</li> </ul>
	2) Contaminated micropipette.	<ul style="list-style-type: none"> <li>■ Use aerosol-free tips.</li> </ul>
	3) Contaminated reagent.	<ul style="list-style-type: none"> <li>■ Consult with a <b>GeneReach</b> technical support representative or local distributor.</li> </ul>
	4) Leakage or spill of reaction from R-tube into reaction chamber of POCKIT™ Nucleic Acid Analyzer.	<ul style="list-style-type: none"> <li>■ Consult with a <b>GeneReach</b> technical support representative or local distributor.</li> </ul>
	5) Contaminated working area.	<ul style="list-style-type: none"> <li>■ Consult with a <b>GeneReach</b> technical support representative on how to clean up working area.</li> </ul>

<b>Problems</b>	<b>Possible causes</b>	<b>Solutions</b>
<b>False Negative</b>	1) Nucleic acid extraction failed.	<ul style="list-style-type: none"> <li>■ Consult manual of Nucleic Acid Co-prep Set.</li> </ul>
	2) Poor nucleic acid quality.	<ul style="list-style-type: none"> <li>■ Check sample storage condition.</li> <li>■ Please refer to Troubleshooting section of Nucleic Acid Co-prep Set</li> </ul>
	3) PCR inhibition.	<ul style="list-style-type: none"> <li>■ Do not overload nucleic acid.</li> <li>■ Spike 5 µl nucleic acid sample into a positive control reaction for a parallel PCR reaction. Negative results indicate the presence of inhibitors in the nucleic acid. In that case, prepare another nucleic acid extract.</li> </ul>

## REFERENCE

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1. Chang, H. G., Tsai, Y., Tsai, C., Lin, C., Lee, P., Teng, P., et al. (2012). A thermally baffled device for highly stabilized convective PCR. *Biotechnology Journal*, 7(5), 662-666. doi: 10.1002/ biot.201100453
2. Corbel, M. J., Elberg, S. S., & Cosivi, O. (2006). *Brucellosis in humans and animals*. Hertfordshire: World Health Organization Press.
3. Tsai, Y., Wang, H. T., Chang, H. G., Tsai, C., Lin, C., Teng, P., et al. (2012). Development of TaqMan Probe-Based Insulated Isothermal PCR (iiPCR) for Sensitive and Specific On-Site Pathogen Detection. *PLoS ONE*, 7(9), e45278. doi:10.1371/journal.pone.0045278