BACTOTYPE® PCR Amplification Kit

Pasteurella multocida



Manual

Technology

The product group BACTOTYPE® PCR Amplification Kit comprises optimised systems for the identification of bacterial pathogens via polymerase chain reaction (PCR):

Highest sensitivity and specificity: All components are carefully developed and perfectly optimised

Easy and comfortable work: The reagents are ready-to-use and ready-to-load

Reliable results: The test kit contains an internal amplification control

The primer pair of the test kit is specific for two gene fragments of *Pasteurella multocida*. The PCR products are analysed on a conventional agarose gel. The ready-to-load reagents allow direct loading of the samples on the gel without addition of gel loading buffer. False negative results are minimized by using an internal amplification control. Manual work is reduced to a minimum. With exception of Taq DNA Polymerase, the test kit consists of all reagents necessary for the detection of the bacterial germs.

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Pathogenic Germ

Pasteurella multocida is part of the physiological bacterial flora of the upper respiratory tract of pigs. As secondary pathogen invading lungs injured by other bacteria or viruses *P. multocida* induces pneumonia. Subsets of *P. multocida* isolates are critical to induce the so called progressive atrophic rhinitis. The isolates synthesize a toxin encoded by the chromosomal *toxA* gene that serves as essential virulence factor. Nontoxigenic *P. multocida* isolates do not cause atrophic rhinitis. Culture, species identification, and toxin testing of *P. multocida* are time-consuming and costly. BACTOTYPE® PCR Amplification Kit *Pasteurella multocida* provides sensitive and specific results in about one day.

Ordering Information

BACTOTYPE® *Pasteurella multocida* 10 Reactions Cat. No. 04–109/10 BACTOTYPE® *Pasteurella multocida* 50 Reactions Cat. No. 04–109/50

Content

	10 reactions	50 reactions
PCR Mix (yellow cap) PCR buffer, dNTPs, primer mix (species-specific and amplification control), DNA for amplification control, NaN ₃	221 μL	935 μL
Magnesium Chloride (green cap) MgCl ₂ , gel loading buffer	80 μL	300 μL
Positive Control (red cap) non-infectious DNA with germ-specific sequences, NaN₃	25 μL	25 μL
Nuclease-free Water (blue cap)	1.0 mL	1.0 mL

Storage Conditions

Reagents of the test kit are stable for at least 12 months when stored at -20 °C in darkness. Repeated freezing and thawing should be avoided.

Solely, the use of this DNA Polymerase assures precise function of the BACTOTYPE® products. For ordering

Additionally Reagents

- Taq DNA Polymerase BACTOTYPE[®] test kits are validated with JumpStart™ Taq DNA Polymerase (Sigma-Aldrich, Cat. No. D4184).
 - and further information, please contact Sigma-Aldrich.

 Agarose and Ethidium bromide (for gel electrophoresis)
 - 100 bp DNA ladder (for gel electrophoresis)

Trademarks and Patents

- BACTOTYPE[®] is a registered trade mark of the Biotype AG.
- JumpStart™ is a registered trademark of Sigma-Aldrich.
- The PCR is under patent laws. Patentees are Hoffmann-La Roche Inc. and F. Hoffmann-La Roche (Roche).

Product Warranty

All contents of the BACTOTYPE® PCR Amplification Kit are subjected to extensive quality procedures by Labor Diagnostik Leipzig. In case of any problem, please report immediately.

Notes

Please, carry out precisely the instructions of the protocol!

In order to avoid any kind of contamination we recommend pipetting with filter tips. DNA preparation, amplification, and electrophoresis should be carried out in separate rooms. Take note of safety regulations for working in laboratories, and wear gloves.

The PCR Mix and the Positive Control contain sodium azide (NaN₃). Sodium azide is hazardous: Very toxic if swallowed, develops toxic gases when it gets in contact with acids (R28-32-50/53, S1/2-28-45-60-61). Ethidium bromide is hazardous, too. Please refer to the Material Safety Data Sheet (MSDS) and wear nitrile gloves.

MSDS of all BACTOTYPE® components are available from Labor Diagnostic Leipzig. For MSDS of additional reagents to be needed, please contact the corresponding manufactures.

DNA Isolation

Solely, ultrapure DNA implies the successful amplification. For DNA isolation of *Pasteurella multocida* from tissue samples or swabs the following test systems are recommended:

- NucleoSpin® Tissue Kit; Macherey-Nagel, Düren
- QIAamp[®] DNA Mini Kit; QIAGEN, Hilden

Please perform isolation according to the manufacturer's information.

Protocol

Preparation of the reagents: Before using the reagents of the test kit the first time, mix thoroughly and centrifuge the reagents for 5 sec at 3000 rpm. The blue colour in the magnesium chloride solution arises from the gel loading buffer and does not affect the PCR reaction. Please, avoid elongated centrifugation steps of the magnesium chloride solution. The gel loading buffer could be concentrated onto the bottom of the tube.

1. A) PCR Protocol for a Number of Samples different from the Packaging Unit (Multiplicator)

If there are less samples than the packaging unit, pipette as followed into a PCR reaction tube (also see pipetting scheme table 1):

17.0 µL PCR Mix

- + 4.5 µL Magnesium Chloride
- + 0.6 µL Nuclease-free Water
- + 0.4 μL Tag DNA Polymerase (here: JumpStartTM 2.5 U/μL)
- Pipet 22.5 μL of the master mix into a PCR reaction tube and add 2.5 μL of your DNA sample or control.
- **Attention:** Depending on the used Taq Polymerase, complete with Nuclease-free Water to a total volume of 25 μL per reaction.

1. B) PCR protocol for an Amount of Samples corresponding to Packaging Unit

If the number of samples corresponds to the packaging unit, we recommend the addition of Magnesium Chloride, Nuclease-free Water, and Taq DNA Polymerase directly to the PCR Mix.

For 10 samples (corresponding to the content of 10 reactions)

PCR Mix

- + 58.5 µL Magnesium Chloride
- + 7.8 μL Nuclease-free Water
- + 5.2 μL Taq DNA Polymerase (here: JumpStart™ 2.5 U/μL)

For 50 samples (corresponding to the content of 50 reactions)

PCR Mix

- + 247.5 μL Magnesium Chloride
- + 33 µL Nuclease-free Water
- + 22 μL Taq DNA Polymerase (here: JumpStart™ 2.5 U/μL)
- Pipet 22.5 μ L of the master mix into a PCR reaction tube and add 2.5 μ L of your DNA sample or control.

2. Positive Control

The test kit is provided with a ready-to-use Positive Control (red cap) of non-infectious DNA with germ-specific sequences. We strongly recommend performing a positive control for each run.

3. Negative Control

It is strongly recommended to perform a negative control (e. g. Nuclease-free Water) for each run.

Table 1: Pipetting Scheme

	Sample (x1)	Positive Control	PCR Control
PCR Mix	17.0 μL	17.0 μL	17.0 μL
Magnesium Chloride	4.5 μL	4.5 μL	4.5 μL
Sample DNA	2.5 μL	-	-
Positive Control	-	2.5 μL	-
Nuclease-free Water	0.6 μL	0.6 μL	3.1 μL
Taq DNA Polymerase (1 U)*	0.4 μL	0.4 μL	0.4 μL

^{*} Use 1 Unit for amplification (here: JumpStart™ 2.5 U/μL).

The volume of the DNA to be employed depends on its concentration and may be varied from 1 to 3 μ L. Adjust the volume with Nuclease-free Water to total reaction volume of 25 μ L.

PCR Amplification Parameters

We strongly recommend the use of Sigma JumpStart™ Taq DNA Polymerase.

Temperature	Time	
94°C	3 min	Hot Start*
94°C	30 s	
60°C RAMPING 1°C/s ^{**}	30 s	35 Cycles
72°C	30 s	
72°C	5 min	
10°C	∞	hold

^{*} For "hot start" Polymerase, please follow the activation time conditions specified by the manufacturer (for JumpStart™ Tag DNA Polymerase: 3 minutes).

4. Electrophoresis

PCR fragments are analysed on a 2% agarose gel and staining is done with ethidium bromide (0.5–1.0 μ g/mL) in TAE buffer.

Put 10 μ L of the PCR product per lane onto the gel (gel loading buffer already included). In order to compare the molecular size of the amplification products a size standard (100 bp DNA ladder) is recommended. Run the electrophoresis as described in the manufacturer's handbook.

The value is adjusted for most of the thermocyclers. Cyclers that exceed the ramping rate should be monitored and corrected as required.

5. Analysis and Interpretation of the Results

Size designation of the PCR fragments is done by comparison with the DNA size standard on a transilluminator. Photographic documentation is recommended.

Protect eyes and skin against UV-radiation!

Positive result: P. multocida strains encoding toxA show a toxin-specific band at 331 bp

in addition to the germ-specific band at 415 bp. If the strain does not encode ToxA, only the germ-specific band at 415 bp can be detected. The 838 bp (amplification control) band may be weak or may not appear at all due to low concentrations of bacterial DNA, whereas the germ-specific band is clearly visible at all. In this case the

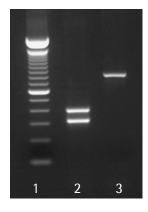
test is valid. The result is positive.

Negative result: If the 838 bp band (amplification control) appears only, germ detection is considered

to be negative.

Invalid result: If none of the bands mentioned above are detectable, the assay is invalid and has to

be repeated. Possibly, the PCR has been inhibited.



838 bp amplification control

415 bp germ-specific band 331 bp toxin-specific band

Lane 1: 100 bp DNA ladder (Invitrogen)

Lane 2: germ-specific (415 bp) and toxin-specific band (331 bp); sample is positive

Lane 3: only the amplification control is visible (838 bp); sample is negative