

## HANDBOOK

# Bio-T kit<sup>®</sup> *Actinobacillus pleuropneumoniae*

Cat. N° BIOTK099 - 50 reactions

**Detection of *Actinobacillus pleuropneumoniae* (APP)  
by real-time PCR (qPCR)  
with Endogenous internal positive control (IPC)**

## SWINE

### Sample types

- Deep nasopharyngeal swab
- Alveolar bronchial washing
- Organs (tonsils)
- Individual analysis or by pool up to 3 according to the matrix

### Recommended nucleic acids (NA) extractions

- Silica membrane columns extraction (e.g.: BioSella – BioExtract<sup>®</sup> Column Cat. N° BEC050 or BEC250)
- Magnetic beads extraction (e.g.: BioSella – BioExtract<sup>®</sup> SuperBall<sup>®</sup> Cat. N° BES384)

*Veterinary use only*



## DOCUMENTS MANAGEMENT

The Bio-T kit® *Actinobacillus pleuropneumoniae* has two technical handbooks:

- The extraction handbook for Bio-T kit® *Actinobacillus pleuropneumoniae*, displaying BioSella's recommended extraction protocols for each type of sample.
- The Bio-T kit® *Actinobacillus pleuropneumoniae* qPCR handbook, presenting the instruction information to perform the qPCR.

The last versions in use for each handbook are indicated on the certificate of analysis (CA) provided with the Bio-T kit® *Actinobacillus pleuropneumoniae*.

## MODIFICATIONS MANAGEMENT

BioSella indicates modifications done to this document by highlighting them using the rules presented in the Table below:

MODIFICATIONS MANAGEMENT			
Type of modification Highlighting color	Minor modifications	Type 1 Major modifications	Type 2 Major modifications
Impact on revision / version	Change of revision date No change of version	Change of revision date + change of version	Change of revision date + change of version
Examples of modifications	Corrections: typographical, grammatical or turns of phrase	EPC reference modification	Modification of Master Mix composition
	Addition of new sample type for extraction	Exogenous IPC reference modification	Modification of validated extraction protocol
	Addition of information giving more details or alternative protocol		
	Addition/Suppression of optional information		

## PRESENTATION

### Recommendations for sampling, shipping and storage of samples

Real-time PCR is a powerful technique allowing the detection of few amounts of pathogen genome. Genome can be rapidly degraded depending on the pathogen nature (bacteria / parasites, enveloped viruses...), the genome nature (DNA / RNA) and the sample type (presence of DNase / RNase). Thus, BioSellal recommends the following instructions to guarantee an optimal diagnosis.

#### Sampling

To prevent cross-contamination between samples leading to false positive results, it is mandatory to use disposable materials for single use and to avoid direct contact between specimens.

#### Shipping

It is recommended to ship soon as possible after sampling, under cover of positive cold.

#### Storage after reception

Recommended storage of samples at  $5^{\circ}\text{C} \pm 3$  for a maximum of 7 days and  $\leq -16^{\circ}\text{C}$  beyond.

### PIG Line

This kit belongs to the PIG line which gather a set of kits sharing common extraction and qPCR protocols. It is compatible with BioSellal's other kits belonging to the PIG and AVIAN lines. (information available on [www.biosellal.com](http://www.biosellal.com)).

## Description of the Bio-T kit® *Actinobacillus pleuropneumoniae*

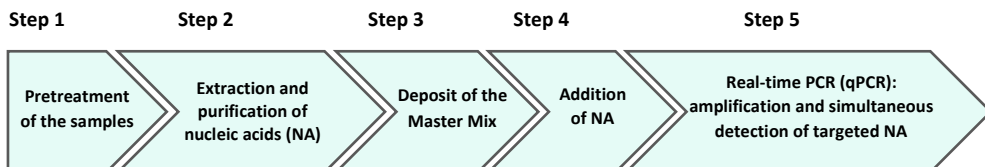
The **Bio-T kit® *Actinobacillus pleuropneumoniae*** (Cat. N° BIOTK099) contains a ready to use **PCR Master Mix** allowing the detection **in the same reaction well of**:

- ***Actinobacillus pleuropneumoniae* (APP)** with a 6-FAM labelling,
- **An Endogenous internal positive control IPC (beta actin)**, with a Cy5 labelling, to assess the presence of sufficient amount of host cells, sample integrity, nucleic acids extraction quality and absence of PCR inhibitors.

This kit, based on qualitative detection (detected or non detected) from deep nasopharyngeal swab, alveolar bronchial washing and organs samples (Individual analysis or by pool up to 3 according to the matrix), was developed and validated according to the **French regulatory standard NF U47-600-2 edited by AFNOR for the PCR part**.

Extraction protocols recommended by BioSellal are described in the **Bio-T kit® *Actinobacillus pleuropneumoniae* extraction handbook**.

## Description of the whole process



Extraction handbook of the Bio-T kit® <i>Actinobacillus pleuropneumoniae</i>		qPCR handbook of the Bio-T kit® <i>Actinobacillus pleuropneumoniae</i>		
Deep nasopharyngeal swab*	BioExtract® Column BioExtract® SuperBall®	Ready-to-use Master Mix MMAPP-A	Samples NC/NCS Process positive control EPC (EPCAPP-A)	Dyes: FAM/Cy5
Alveolar bronchial washing*				Passive reference: ROX
Organs*				Programs: PIG/AVIAN program ± RT Standard ramping

\* pretreatment mandatory

## Kit contents and storage

Table 1. Description of the kit contents				
Description	Reference	Volume/tube	Presentation	Storage
<b>Master Mix (MM)</b> Ready to use	MMAPP-A	750 µl	White cap tube Bag A	≤-16°C Protected from light, « MIX » Area
<b>External Positive Control (EPC)</b> Positive PCR control of APP	EPCAPP-A	110µl	Orange cap tube Bag B	≤-16°C « Addition of Nucleic acids » Area
<b>Water</b> RNase/DNase free	Aqua-A	1 ml	Blue cap tube Bag B	5°C ± 3 or ≤-16°C « Addition of Nucleic acids » Area

## List of consumables and reagents not included in kit

Table 2. Consumables and reagents not included in kit			
Consumables/ Reagents	Description	Provider	Cat. N°
<b>ATL Buffer</b>	Lysis Buffer	BioSellal	ATL19076
<b>BioExtract® Column</b>	DNA/RNA column extraction kit (50)	BioSellal	BEC050
<b>BioExtract® Column</b>	DNA/RNA column extraction kit (250)	BioSellal	BEC250
<b>BioExtract® SuperBall®</b>	DNA/RNA Magnetic beads extraction kit (4 x 96)	BioSellal	BES384

For consumables related to the thermal cycler, refer to the user manual of the device.

## Main critical points

- Wear appropriate personal protective equipment (lab coat, disposable gloves frequently changed).
- Work in dedicated and separate areas to avoid contamination: "Extraction" (unextracted samples storage, extraction equipment area), "Mix" (ready to use MM storage, qPCR plates preparation), "Nucleic acids Addition" (Nucleic Acids storage and addition of extracted nucleic acids and controls in the qPCR plate), "PCR" (final area containing the thermal cycler(s)).
- Use dedicated equipment for each working area (gloves, lab coat, pipettes, vortex, ...).
- Use filter tips.
- Before use, thaw all components at room temperature.
- Vortex and spin briefly (mini-centrifuge) all reagents before use.
- Avoid the repetition of freezing-thawing cycles for samples, lysates, extracted nucleic acids.
- **Pathogen's genome detected by the PIG line's kits can be DNA or RNA. Working with RNA is more demanding than working with DNA** (RNA instability and omnipresence of the RNases). For these reasons, special precautions must be taken:
  - o Always wear gloves, change them frequently, especially after contact with skin or work surfaces.
  - o Treat all surfaces and equipment with RNases inactivation agents (available commercially).
  - o When wearing gloves and after material decontamination, minimize the contact with surfaces and equipment in order to avoid the reintroduction of RNases.
  - o Use "RNase free" consumable.
  - o It is recommended to store the RNA at  $\leq 5^{\circ}\text{C} \pm 3$  during the manipulation and then freeze it as soon as possible, preferably at  $\leq -65^{\circ}\text{C}$  or by default at  $\leq -16^{\circ}\text{C}$ .
  - o Open and close tubes one by one in order to limit the opening times and avoid any contact with RNases present in the environment (skin, dust, working surfaces...).

# DETECTION OF APP BY qPCR WITH BIOTK099 KIT

## Global Procedure

### 1) Establish qPCR plate setup defining each sample position and including the following controls:

- **Negative Control Sample (NCS):** water (or PBS) replaces the sample from the first step of sample preparation.

This control is mandatory for each extraction series.

- **Negative Amplification Control (NC):** 5 µl of water RNase/DNase free (Aqua-A tube, **blue** cap) replaces sample Nucleic Acids extract on qPCR plate.

This control is recommended when using the kit for the first time or to verify the absence of Master Mix contamination.

- **External Positive Control of APP (EPC) :** Synthetic DNA (tube **EPCAPP-A**, **orange** cap), containing specific target of APP.

This control is mandatory.

**⚠ CAUTION:** *EPC tube handling represents nucleic acids contamination hazard, it is thus recommended to open and handle it in a restricted area, away from other PCR components and to take precautions to avoid cross-contamination with nucleic acids extracts during deposit on the qPCR plate.*

- If available, a **Process Positive Control (MRI)**, a weak positive sample is extracted in parallel with tested samples. After qPCR, MRI Ct values will be monitored on a Shewhart control card. Obtaining conform Ct values validates the whole process. In this case, the use of the EPC, provided with the kit, is not mandatory.

## 2) qPCR plate preparation

### In the “MIX” dedicated area

1. After thawing, vortex and rapid centrifugation, **transfer 15 µl Master Mix MMAPP-A (white cap)** in each well of interest (samples and controls).

### In the “Nucleic Acids addition” dedicated area

2. **Add 5 µl of extracted nucleic acids (or NCS, water, MRI, or EPC: EPCAPP-A orange cap tube)** in each well of interest. Make sure to pipet out in the bottom of the well, in the Master Mix, and to avoid the formation of bubbles.
3. **Seal the plate with an optically clear sealer or close the strip caps.**

### In the “PCR” amplification dedicated area

4. **Define the thermal cycler parameters** (see Table 3, Table 4, Table 5)
5. It is recommended to **spin the plate down prior to place it in the thermal cycler**, to prevent drops in the well pit walls.
6. Start the qPCR program. Approximate run time: 70 min.

## 3) Thermal cycler settings

This kit was developed and validated on AriaMx™ (Agilent Technologies, Fast ramping by default) and confirmed on ABI PRISM® 7500 Fast (Applied Biosystems) in standard ramping. It is compatible with all thermal cyclers with at least 6-FAM and Cy5 channels. For more information, contact our technical support.

Table 3. Thermal cycler configuration		
ABI PRISM® 7500 Fast		AriaMx™
<b>Mode</b>	Quantitation – Standard curve	Quantitative PCR, Fluorescence Probe
<b>Ramping</b>	Standard Ramping	Fast Ramping by default
<b>Passive Reference</b>	ROX	ROX



Table 4. Thermal cycler Settings			
Target	Detectors		Final Volume / well
	Reporter	Quencher	
APP	FAM	NFQ-MGB or None*	20 µl  = 15 µl Master Mix + 5 µl extracted nucleic acids or controls <sup>†</sup>
Endogenous IPC	Cy5	NFQ-MGB or None*	
To assign to samples and controls <sup>†</sup>			

\* Depends on the thermal cycler model. Do not hesitate to contact the BioSellal Technical Support (tech@biosellal.com)

† Controls are NC (water), NCS (extracted water), EPC and or extracted MRI.

Table 5. PIG/AVIAN Amplification program settings without RT <sup>†</sup>		
Standard ramping		
Cycles	Time	Temperature
1 cycle	5 min	95°C
40 cycles	10 sec	95°C
	45 sec	60°C
	+ data acquisition	

<sup>†</sup> optional step, in case of simultaneous detection of RNA genomes. Achieving a reverse-transcription (RT) step prior to PCR for the amplification of RNA genomes has no impact on the performances of the Bio-T kit® *Actinobacillus pleuropneumoniae* (see the summary of the validation file).

NB: This amplification program is compatible with all Bio-T kit® of the PIG and AVIAN LINES.

## RESULTS INTERPRETATION

To analyze and interpret the signals obtained by qPCR, the Threshold must be set up.

The threshold must be assigned carefully in order to obtain the most reproducible result between different manipulations according to the requirements defined in Annex C of the French Standard **NF U47-600 (part 1)**. A consistent set of positives controls, usually an In-house Reference Material (MRI) or the EPC, is used to set the threshold value above the baseline and in the exponential amplification phase of the plot.

The Threshold Cycle, named « Ct » or « Cq » (depending on thermal cyclers), corresponds to the intersection between the amplification curves and the threshold line. It allows the relative measurement of the concentration of the target in the PCR reaction when a calibrated extract is analyzed in the same series.

The qPCR series is validated if the controls (EPC, MRI, NCS and NC) present valid results, then the result of each sample can be interpreted.

## Main Scenarios

### Controls Reading

Table 6. PCR Controls results interpretation			
	Targets		Interpretation
	APP (FAM)	Endogenous IPC (Cy5)	
<b>NCS</b> Negative Control Sample  <b>MANDATORY</b>	Neg	Neg	Valid
	At least one of the two targets <b>Pos</b>		Contamination with a positive/negative sample during extraction step or during qPCR plate preparation.
<b>NC</b> Negative PCR Control  <b>OPTIONAL</b>	Neg	Neg	Valid
	At least one of the two targets <b>Pos</b>		Contamination with a positive/negative sample during extraction step or during qPCR plate preparation or Master Mix/water contamination
<b>EPC</b> APP PCR external positive control  <b>MANDATORY</b> <i>IN ABSENCE OF MRI</i>	Pos*	Neg	Valid
	Neg	Neg	Problem during qPCR plate preparation: Master Mix error? EPC omission?
	Pos*	Pos	Contamination with a sample during qPCR plate preparation?
<b>Sample process positive Control MRI</b>  <b>RECOMMENDED</b> <i>IF AVAILABLE</i>	Pos†	Pos‡	Valid
	Neg	Neg	Problem during qPCR plate preparation: Master Mix error? Nucleic acids extract omission or extract not in contact with Master Mix? Process drift: extraction and/or qPCR ? Degradation of the sample process positive control?

\* The Ct value obtained must be conform with the value indicated on the Certificate of Analysis (CA).

† The Ct value must be included within control card limits.

‡ The obtained Ct value depends on the thermal cycler, the sample type and the used extraction protocol. IPC Ct values for recommended extraction protocols are available upon request. BioSella recommends you determine your own maximal IPC Ct value depending on your own extraction method and thermal cycler.

#### Note:

Endogenous IPC targets a gene expressed by swine cells, thus it cannot be detected in NCS, NC and EPC.

However? Due to cross-reaction between swine beta actin and human beta actin, a slight signal can be observed for IPC in the controls, the Ct value of this signal must be higher than 35.

## Samples Reading

**Table 7. Different types of results obtained for the samples**

Targets		
APP (FAM)	Endogenous IPC (Cy5)	Interpretation
Neg	Pos*	Negative or Undetected
Pos		Positive or Detected
Pos	Neg or Ct>35	<b>Positive or Detected</b> Lack of host cells? Presence of inhibitors <sup>†</sup> ? Competition with the targets?
Neg	Neg or Ct>35	<b>Uninterpretable = Repeat the analysis</b> Problem during qPCR plate preparation: Master Mix error? Nucleic acids extract omission or extract not in contact with Master Mix? Presence of inhibitors <sup>†</sup> ? Nucleic acids degradation in the sample? Sampling problem: lack of cells? Extraction problem?

\*The obtained Ct value depends on the thermal cycler, the sample type and the used extraction protocol. IPC Ct values for recommended extraction protocols are available upon request. BioSella recommends you determine your own maximal IPC Ct value depending on your own extraction method and thermal cycler.

† In case of inhibition suspicion, 1) Repeat the qPCR with the dilution of extracted nucleic acids at 1/10 or 1/100 in the DNase/RNase free water.  
2) Restart the analysis from the extraction step.



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