

# Protocols for KingFisher\* instruments Using "ADIAMAG" extraction kit Ref. NADI003 (200 extractions) Ref. NADI003-XL (800 extractions)

\* If you use a different automat, please contact support.pcr@biox.com

# **Using "ADIAMAG"** extraction kit

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# Main change since previous version

N/A Not Applicable (first publication)
Correction Correction of document anomalies

Technical change Addition, revision and/or removal of information related to the product

Administrative Implementation of non-technical changes noticeable to the user

Note: minor typographical, grammar and formatting changes are not included in the revision historic

Release Date	Part Number	Change type	Change summary
2022/09	NEKF-24	Technical	Addition of protocols for the search of BVDv from humid TST-L from
		change	ALLFLEX ear notch (BAH) or conservation buffer of TST-L from
			ALLFLEX ear notch (BAL)
2022/11	NEKF-25	Technical	Addition of protocol for the search of <i>Mycobacterium avium</i> subsp
		change	paratuberculosis from faeces, tissue.
			Addition of protocol for search of Influenza virus and PRRS virus
			from oral fluids.
			Addition of environmental samples matrix for the search of
			Influenza Virus.
			Addition of a protocol for search of Bronchitis Virus.
2023/11	NEKF-26	Technical	Addition of protocols for the search of IHNV and VHSV from
		change	sperm/egg fish.
			Addition of protocols for the search of EHDV from blood and tissue.
			Addition of protocols for the search of BTV from tissue.
2025/01	NEKF-27	Technical	Addition of specific matrices for the search of African swine fever
		change	(environmental samples, blood on Whatman® 3 blotting paper,
			muscle exudate and semen).
			Addition of protocols for the search of <i>Lawsonia intracellularis</i> and
			Brachyspira.
			Addition of protocols for the search of <i>M. hyopneumoniae</i> from
			bacterial culture supernatants and FTA cards.
			Addition of protocols for the search of <i>Actinobacillus</i> pleuropneumoniae from supernatant bacterial culture and oral
			fluid.
			Addition of a protocol for the search of BVDV from ear notch
			combined with the ADIALYO BVDV Triplex amplification kit.
			Addition of a protocol for the search of PRRSV from semen.

## I. General information

## 1. Purpose of the kit

ADIAMAG is a nucleic acids DNA and RNA extraction kit based on nucleic acids adsorption on magnetic beads. The kit is adapted for KingFisher™ mL, DUO and 96/Flex instruments and contains all the buffers required to perform DNA/RNA extraction from different matrices.

If using a different automaton than those validated, please contact support.pcr@biox.com.

After samples lysis (performed separately from the instrument and using appropriate buffers), the supernatants can be handled with the same automated purification process using KingFisher $^{\text{TM}}$  mL, KingFisher DUO or KingFisher $^{\text{TM}}$  96/Flex instruments, irrespective of the pathogen and matrix used. The following table describes the protocol used with the different instruments.

Well/plate	Description
1	Nucleic acids release and binding to magnetic beads.
1	Beads collection by magnetic rods and transfer into the step 2.
2	Beads release. W3 buffer washing step.
2	Beads collection by magnetic rods and transfer into the step 3.
3	Beads release. W4 buffer washing step.
3	Beads collection by magnetic rods and transfer into the step 4.
4	Beads release. Ethanol 80 % washing step.
4	Beads collection by magnetic rods, air drying step followed by transfer into the step 5.
5	Nucleic acids elution step with E6 buffer.
3	Nucleic acid-free beads collection and transfer into the step 3.

Contact us to obtain the files for your KingFisher™ instruments.

## 2. Description of test

Bio-X Diagnostics validated the ADIAMAG kits with various matrices for the detection of different DNA/RNA pathogens in combination with the ADIAVET™ and ADIALYO™ range (please refer to the validation file for the kit concerned for validated matrices). The following table summarises the validated protocols.

											Sam	ples									
											Jan										
		EDTA blood	Sera, plasma	Ear notche	Skin biopsy	Tissue	Brain	Swab	Oral fluid	Faeces	Milk	tracheo –bronchial washing	Culture supernatant	Water/urine	Semen	Feather	FTA card	Fœtal gastric fluid	Environnemental sample	Muscle exudate	Blood on Whatman 3
Animal	Influenza virus					Х		Х	Х	Х		Х	Х			Х	Х		Х		
diseases	Chlamydia					Х		Х			Х							Х			
	BoHV-4							Х													
	Coxiella burnetii					Х		Х		Х	Х							Х			
	Leptospira					Х		Х						Х							
	Neospora caninum					Х	Х	Х													
	BVD virus	Х	Х	Х		Χ*					Х										
	Schmallenberg virus (SBV)	Х	Х			Х	Х														
Ruminant	Anaplasma phagocytophilum	х				Х		х													
diseases	Besnoitia besnoiti	Х			Х																
	Toxoplasma gondii					Χ	Χ	Х		Χ											
	Bluetongue virus (BTV)	Х				Х															
	Epizootic Haemorrhagic disease virus (EHDV)	х				Х															
	M. avium subsp. paratuberculosis					Х				Х									Х		
	Salmonella					Χ		Х										Х			
	Mycoplasma hyopneumoniae					Х		Х	Х			Х	Х				Х				
	Lawsonia intracellularis					х		Х	Х	Х									Х		
	Brachyspira					Х		Х		Х									Х		
	PRV Virus, Aujesky's disease					Х*	Х	Х					Х								
Swine diseases	PRRS virus	Х	Х			Х		Х	Х			Х			Х						
uiscuses	Actinobacillus pleuropneumoniae					Х		Х	Х				Х								
	Classical Swine Fever Virus (CSFV)	Х	Х			Х*															
	African Swine Fever Virus (ASFV)	Х	Х			Х*		Х							Х				Х	Х	х
	Porcine Circovirus (PCV2&PCV3)	Х	Х			Х															

<sup>\*</sup>If using the reference NADI003-XL, it is necessary to add the reference NADI004. For the pool size, refer to the user manual of the kit concerned.

			Samples																
		EDTA blood	Sera, plasma	Ear notch	Skin biopsy	Tissue	Brain	Swab	Oral fluid	Faeces	Milk	tracheo –bronchial washing	Culture supernatant	Water/urine	Sperm, egg, coelomic liquid	Feather	FTA card	Fœtal gastric fluid	Environnemental sample
Avian	Mycoplasma gallisepticum M. synoviae M. meleagridis M. iowea							Х									Х		
diseases	Ornithobacterium rhinotracheale							х											
	Marek virus					Х				Х						Х			Х
	Bronchitis Virus							Х											Х
Contagious equine metritis (CEM)	Taylorella equigenitalis, Taylorella asinigenitalis, Klebsiella pneumoniae Pseudomonas aeruginosa							X											
	Infectious hematopoietic necrosis virus (IHNV)					х							х		х				
Fish diseases	Viral hemorrhagic septicemia virus (VHSV)					х							Х		Х				
	Infectious pancreatis necrosis virus (IPNV)					X							Х		X				
Feline disease	Feline coronavirus (FIP)	х						х									_	_	

## II. Material & reagents

## 1. Composition of kit

The ADIAMAG kits contain the following buffers:

Designation of reagents	Type of reagent	NADI003 ADIAMAG 200 extractions	NADI003-XL ADIAMAG XL 800 extractions	NADI004 LB3 -125mL	Instruction
Lysis Buffer - LB1	Lysis buffer 1	1 x 20 mL	1 x 100 mL		Ready to use
Lysis Buffer - LB2	Lysis buffer 2	1 x 50 mL	2 x 125 mL		Ready to use
Lysis Buffer - LB3	Lysis buffer 3	1 x 125 mL		1x125 mL	Ready to use
ADIAMAG beads	Magnetics beads	2 x 1.5 mL	1 x 12 mL		Ready to use
Binding Buffer - B2	Binding buffer	1 x 180 mL	1 x 500 mL		Ready to use
Wash Buffer - W3	Wash Buffer	1 x 75 mL	1 x 300 mL		Ready to use
Wash Buffer - W4	Wash Buffer	1 x 75 mL	1 x 300 mL		Ready to use
Elution Buffer - E6	Elution Buffer	1 x 30 mL	1 x 125 mL		Ready to use
Proteinase K - PK	Enzyme	1 x 75 mg	4 x 75 mg		Lyophilized – to rehydrate
Proteinase Buffer - BPK	Rehydration buffer	1 x 8 mL	1 x 15 mL		Ready to use

## 2. Validity and storage

After reception, all the extraction reagents can be stored at room temperature ( $\pm$ 18 to  $\pm$ 25 °C) and are stable for up to 1 year and at the most until the shelf-life of the kit. Leave the flasks closed to prevent evaporation.

Prior to use, resuspend the lyophilized proteinase K using 2.6 mL of "Proteinase Buffer - BPK". The Proteinase K solution should be stored at < -15 °C.

In case the LB2 Lysis Buffer precipitates, preheat the solution at +70 °C until it clears. In case the LB1 and LB3 Lysis Buffer precipitate, preheat the solution at +30-40 °C until it clears.

## 3. Equipment required, but not supplied in the kit

Warning: The material should be Nuclease-free (e.g., autoclaved 25 minutes twice at +121 °C or once 60 minutes at +121 °C)

- Class II Microbiological Safety Cabinet.
- Centrifuge for microtubes and 20 mL tubes.
- Grinder (Mixer Mill or Fast Prep).
- Incubator, heating bath or block heater.
- Vortex
- 1 10  $\mu L$  pipette, 20 200  $\mu L$  pipette and 200 1 000  $\mu L$  pipette.
- Nuclease-free filter tips.
- Nuclease-free microtubes: 1.5 mL and 2 mL.
- Sterile tubes of 10 or 15 mL.
- Powder-free latex or nitrile gloves.
- Razor blades.
- 80 % ethanol solution.
- Sterile distilled water.
- Sterile saline water (NaCl 8.5 g/L)
- 1X PBS buffer (recommended composition, NaCl 150 mM, Na<sub>2</sub>HPO<sub>4</sub> 5 mM, KH<sub>2</sub>PO<sub>4</sub> 1.7 mM, without Ca<sup>2+</sup>, without K<sup>+</sup> another composition can be used after validation by the user).
- MEM medium + antibiotics (penicillin 100 IU/mL and streptomycin 100  $\mu g/mL$ ).

## Specific equipment for KingFisher instruments

## KingFisher 96/Flex

- DEEP WELL plates (Thermo scientific, 96 tests: ref. 10373480).
- ELUTION PLATES (Thermo scientific, 96 tests: ref. 10357939).
- TIPS (Thermo scientific, 96 tests: ref. 11744978).

## **KingFisher DUO**

- DEEP WELL plates (Thermo scientific, 96 tests: ref. 10373480).
- TIPS-12 (Thermo scientific, 600 tests: ref. 97003500).

## KingFisher mL

- KingFisher combi 240 (Thermo Scientific, 240 tests: ref: 97002141).

## Specific equipment for Mycobacterium avium subsp. paratuberculosis detection

- ADIAFILTER (Bio-X Diagnostics, 100 tests: ref. ADIFIL100).
- ADIAPREP (Bio-X Diagnostics, 100 tests: ref. ADPREP-200).
- Glass beads for grinding only for Mixer mill:
  - ADIAPURE™ ALIQUOTED GLASS BEADS (Bio-X Diagnostics, 480 tests: ref ADIADPBA1-480).
  - ADIAPURE™ GLASS BEADS RACKS 4x96 (Bio-X Diagnostics, 384 tests: ref ADPBIAR-4x96).
- Glass beads for grinding only for grinder Fast Prep or similar:
  - Lysing Matrix B tubes (MP Biomedical, 100 extractions: ref. 116911.100).

## Specific equipment for SBV virus or Influenza virus detection from tissue

- Lysing Matrix D tubes (MP Biomedicals, 100 extractions: ref. 116913.100) only for grinder Fast Prep.
- Metal beads 3 mm (e.g., Qiagen, 200 extractions: ref. 69997) only for grinder Mixer Mill.

## III. Use of samples and controls

## 1. Precautions

#### Caution:

Prepare buffers according to the §II.2.

Buffers can contain toxic substances, please consult the MSDS safety data sheet.

Store reagents at the recommended temperature.

Only appropriately trained personnel should perform this extraction. Ensure the micropipettes used are calibrated. The quality of the obtained results depends on rigorous respect of good laboratory practices.

PCR generates large amount of DNA. A few molecules of amplified products are sufficient to generate a positive result. Hence, PCR tubes should not be opened after amplification.

Samples for analysis should be handled and disposed of as biological waste. Take all measures of security and confinement required for the manipulation of the concerned biological agents.

Before starting the process, read the entire protocol and scrupulously respect it.

## 2. Storage of nucleic acid extracts

Extracted DNAs and can be stored at 4  $^{\circ}$ C for up to 24 hours after extraction. For long term storage, DNAs should be kept at < -15  $^{\circ}$ C.

Extracted RNAs are highly sensitive to temperature. Extraction should be performed at room temperature as fast as possible to avoid degradation. Crude extracts can be stored at 4 °C for a few hours after extraction. For long-term storage, RNAs should be stored at < -65 °C.

## 3. Controls preparation

Several controls should be included for each extraction.

The mix of the different controls included in the kits allows validation of all the steps (extraction and amplification) for all the samples.

- The endogenous or exogenous internal control included in the ADIAVET™ or ADIALYO™ kits allows validation of the extraction and amplification steps of each sample.
- The positive control included in the ADIAVET™ or ADIALYO™ kits allows validation of the specific target amplification.

Other controls should or must be added.

## A. Negative control of extraction (required)

To verify the absence of cross-contamination, at least one negative control has to be included per trial (e.g., AFNOR NF U47-600-1 guidelines recommends 1 negative control per 24 columns or 4 negative controls per trial of 96-wells plate). This control can be a negative sample or a buffer used for dilution.

## B. Positive control of extraction (recommended)

A positive control (including the specific pathogen) can be added in each trial. It can be a positive sample available in the laboratory or a negative sample spiked with the specific pathogen. This positive control will be detected close to the limit of detection of the method and allow comparison of the results obtained in different assays.

## IV. Preparation of the buffers and loading of the instruments

## 1. Preparation of the binding buffer

We recommend preparing the binding buffer just before use.

Carefully mix the "ADIAMAG Beads" solution before each use.

During buffer preparation, always add one more reaction than needed (to cover the loss of solution during multi-pipetting).

Mix (per reaction):

- 600 μL of Binding Buffer B2.
- 13 μL of ADIAMAG Beads.

## 2. Loading of the instrument

According to the KingFisher™ instrument used, the loading is carried out as follow:

- KingFisher™ mL: one tube comb per sample.
- KingFisher™ DUO: one 96-plate.
- KingFisher™ 96/Flex: one 96-plate per step.

Distribute the buffer in each well/line/plate as mentioned in the following table:

	Buffer to add	KingFisher™ mL	KingFisher™ DUO	KingFisher™ 96/Flex			
	600 μL binding buffer (600μL Binding Buffer B2 + 13 μL ADIAMAG beads)	Well 1	Line B	Plate 1			
	350 μL Wash Buffer W3	Well 2	Line C	Plate 2			
	350 μL Wash Buffer W4	Well 3	Line D	Plate 3			
Preparation of reagents	350 μL Ethanol 80 %	Well 4	Line E	Plate 4			
	60 μL or 100 μL* Elution Buffer E6	Well 5	Line F	Plate 5			
	Tip	On rail	Line A	Plate 6			
Loading of samples and run the instrument	and run the Place the comb or plates in the instrument						

<sup>\*</sup>For ADIAVET range: 100 µL for the influenza virus, PRV, Besnoitia besnoiti, Marek, Porcine Circovirus, *Mycobacterium avium* subsp *paratuberculosis* (with ADIAPREP protocol), Salmonella and fish diseases. 60 µL for other pathogens.

For ADIALYO range: 100 μL for all pathogens, except ADIALYO BTV/EHDV for which 60 or 100 μL can be used.

## 3. Instrument programs

All instrument programs listed below are compatible with the PCR kit range ADIAVET™ or ADIALYO™:

(Summarizing the standard and short programs for the different automates)

	Standard program (34 minutes)	Short program (21 minutes)
KingFisher™ 96/Flex	KF96V3-2	KF96V4-2
KingFisher™ DUO	KFDUOV3-1	KFDUOV4-1
KingFisher™ mL	KFMLV3	KFMLV4

The above programs, as well as programs for another instrument, are available on request (<a href="mailto:support.pcr@biox.com">support.pcr@biox.com</a>).

Warning, in the layout of our program, there may be a difference between real volume and programmed volume. This difference is validated.

## V. Samples preparation before transfer on the device

## 1. From blood/sera

	Virus: BVDV, BTV, SBV, EHDV, PRRSV, CSFV, FIP and ASFV	PCV2 and PCV3	A. phagocytophilum and Besnoitia besnoiti				
Sample preparation	Take <b>100 μL</b> of sample.	Take <b>100 μL</b> of sample.	Place 500 µL to 1 mL of bovine blood or 100 µL of equine blood in a microtube.  Add 1 mL of sterile distilled water.  Mix and incubate 10 minutes on ice.  Centrifuge 6 000 g/5 minutes.  Discard the supernatant.  Add 1 mL of sterile distilled water.  Centrifuge 6 000 g/5 minutes.  Discard the supernatant.				
	Add <b>250 μL</b> of <b>Lysis Buffer</b>	Add 250 μL of Lysis Buffer LB2 + 10 μL of Proteinase PK. <sup>1</sup>					
Lysis	Mix.	Mix. Incubate 15 minutes at room temperature.					
Loading of the instrument	Transfer 2	200 μL in the binding buffe	er (well 1/line B/plate 1). <sup>3</sup>				

<sup>&</sup>lt;sup>1</sup>A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for preparation, storage and use of the control.

## 2. From oral fluids

	L. intracellularis, A. pleuropneumoniae, M. hyopneumoniae, PRRS virus and Influenza virus
Sample preparation	Take <b>100 μL</b> of <b>sample</b> .
Lucio	Add <b>250 μL</b> of <b>Lysis Buffer LB2.</b> <sup>1</sup>
Lysis	Mix.
Loading of the instrument	Transfer <b>200 μL</b> in the <b>binding buffer</b> (well 1/line B/plate 1). <sup>2</sup>

 $<sup>^{1}</sup>$  Add **5 µL of EPC-Ext** provided in the kit of the pathogen of interest.

Refer to the corresponding package insert for preparation, storage and use of the control.

 $<sup>^2</sup>$  Add 5  $\mu\text{L}$  of EPC-Ext provided in the kit of the pathogen of interest.

<sup>&</sup>lt;sup>3</sup> For 96-plate lysis, the binding buffer can be added directly to the lysed sample.

<sup>&</sup>lt;sup>2</sup> For 96-plate lysis, the binding buffer can be added directly to the lysed sample.

## 3. From milk

	BVDV, C. burnetii and Chlamydia
Sample preparation	Take <b>100 μL</b> of <b>sample</b> .
	Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>Proteinase PK.</b> <sup>1,2</sup>
Lysis	Mix.
	Incubate 15 minutes at 56 °C.
Loading of the instrument	Transfer <b>the whole volume</b> in the <b>binding buffer</b> (well 1/line B/plate 1).

<sup>&</sup>lt;sup>1</sup>A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for preparation, storage and use of the control.

## 4. From culture supernatant

	PRV, IHNV, VHSV IPNV, M. hyopneumoniae, A. pleuropneumoniae and Influenza
Sample preparation	Take <b>100 μL</b> of <b>sample</b> .
	Add 100 μL of Lysis Buffer LB1 + 10 μL of Proteinase PK. <sup>1,2</sup>
Lysis	Mix.
	Incubate 15 minutes at room temperature.
Loading of the	Transfer the whole volume in the binding buffer.
instrument	(Well 1/line B/plate 1).

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for preparation, storage and use of the control.

## 5. From ear notch samples and humid TST-L from ALLFLEX ear notch (BAH)

	BVDV				
Sample	From 1 ear tissue sample, dry or humid.				
preparation	In the case of an ALLFLEX TST-L loop, remove the preservation liquid from the collection tube.				
	Add <b>300 μL</b> or <b>350 μL</b> of <b>Lysis Buffer LB3</b> . <sup>1,4</sup>				
Lysis	Mix.				
	Incubate 15 minutes at room temperature. <sup>2</sup>				
Loading of the instrument	Transfer 100 μL of individual or pooled <sup>3</sup> supernatant in the binding buffer (well 1/Line B/plate 1).				

<sup>&</sup>lt;sup>1</sup> Add **5 μL of EPC-Ext** provided in the kit of the pathogen of interest.

Refer to the corresponding package insert for preparation, storage and use of the control.  $^{1}$ 

<sup>&</sup>lt;sup>2</sup> Add **5 μL of EPC-Ext** provided in the kit of the pathogen of interest.

<sup>&</sup>lt;sup>2</sup> Add **5 µL of EPC-Ext** provided in the kit of the pathogen of interest.

<sup>&</sup>lt;sup>2</sup> For new analysis, each individual supernatant can be stored at +4 °C for 24 hours or at < -15 °C for long-term

 $<sup>^3</sup>$  Up to 25 samples can be pooled together. Mix 50  $\mu L$  of each sample and homogenize.

<sup>&</sup>lt;sup>4</sup> 300 μL if using the ADIALYO™ BVDV Triplex PCR kit or 350 μL if using the ADIAVET™ BVDV REAL TIME PCR kit.

## 6. From conservation buffer of TST-L from ALLFLEX ear notch (BAL)

In the case of ALLFLEX TST-L, eject the biopsy into the preservation buffer and adjust, if necessary, the volume to 250  $\mu$ L with ALLFLEX preservation buffer and incubate for 1 hour at room temperature. For extraction, use the buffer as matrix.

	BVDV				
Sample preparation	Transfer 100 $\mu$ L of conservation buffer or pool <sup>1</sup> of conservation buffer from ear notch sample.				
	Add 100 μL of LB1 Lysis Buffer + 10 μL of Proteinase PK. <sup>2</sup>				
Lysis	Mix.				
	Incubate 15 minutes at room temperature.				
Loading of the instrument	Transfer <b>the whole volume</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).				

 $<sup>^{1}</sup>$  Up to 25 samples can be pooled together. Mix 50  $\mu$ L of each sample and homogenize.

## 7. From skin biopsy

	Besnoitia besnoiti				
Sample preparation	Place <b>50 mg</b> of skin biopsy in a microtube.				
	Add 100 μL of Lysis Buffer LB1 + 10 μL of Proteinase PK + 100 μL PBS Buffer 1X (or sterile saline water) 1				
Lysis	Mix.				
	Incubate overnight at +56 °C.				
Loading of the instrument	Transfer <b>the whole volume</b> in <b>the binding buffer</b> (Well 1/Lane B/Plate 1).				

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

## 8. From trachea- bronchial washing

	M. hyopneumoniae, PRRS virus and Influenza virus				
Sample preparation	Transfer <b>1 mL</b> of <b>trachea-bronchial washing</b> in microtube.  Centrifuge 30 minutes at 10 000g.  Discard the supernatant.				
Lysis	Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>Proteinase PK</b> . <sup>1,2</sup> Mix and incubate 15 minutes at room temperature.				
Loading of the instrument	Transfer <b>the whole volume</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).				

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for preparation, storage and use of the control.

<sup>&</sup>lt;sup>2</sup> A pre-mix can be prepared just before use and added to each sample.

 $<sup>^2</sup>$ Add **5**  $\mu$ L of EPC-Ext provided in the kit of the pathogen of interest.

## 9. From water and urine

	Leptospira				
Sample Transfer 10 mL in a tube.  Centrifuge 30 minutes at 10 000 g or 10 minutes at 4 500 g.  Discard the supernatant. Add 1 mL PBS 1X. Mix.					
Transfer 100 μL in a microtube.  Lysis  Add 100 μL of Lysis Buffer LB1 + 10 μL of Proteinase PK.  Mix and incubate 15 minutes at +56 °C.					
Loading of the instrument	Transfer <b>the whole volume</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).				

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

## 10. From coelomic liquid

	IPNV				
Sample preparation	Take <b>100 μL</b> of <b>sample.</b>				
	Add 100 μL of Lysis Buffer LB1 + 10 μL of Proteinase PK + 5 μL of EPC-Ext. <sup>1,2</sup>				
Lysis	Mix.				
	Incubate 15 minutes at room temperature.				
Loading of the instrument	Transfer <b>the whole volume</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).				

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for the preparation, storage and use of the control.

## 11. From fœtal gastric fluid

	C. burnetii, Chlamydia, Salmonella				
Sample preparation	Take <b>100 μL</b> of <b>sample</b> .  (If the liquid is difficult to collect, dip a swab and treat it according to the swab protocol §15.A.f				
Lysis	Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>Proteinase PK.</b> <sup>1,2</sup> Mix and incubate 15 minutes at +56 °C.				
Loading of the instrument	Transfer <b>the whole volume</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).				

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for the preparation, storage and use of the control.

<sup>&</sup>lt;sup>2</sup> Add **5 µL of EPC-Ext** included in the kit.

 $<sup>^2</sup>$  Add 5  $\mu L$  of EPC-Ext included in the kit.

## 12. From feather

	Avian Influenza virus, Marek virus			
Sample	Cut the calamus of 1 to 5 feathers precociously to avoid any projections and place them in 2 mL of physiological saline.			
preparation	Take <b>100 μL</b> of <b>sample</b> .			
	Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>Proteinase PK.</b> <sup>1,2</sup>			
Lysis	Mix.			
	Incubate 15 minutes at room temperature.			
Loading of the	Transfer the <b>whole volume</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).			
instrument	Transfer the whole volume in the billiang burier (well 1/Line b) plate 1).			

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for the preparation, storage and use of the control.

## 13. From FTA Card

	Avian Influenza virus, Avian mycoplasmas, M. hyopneumoniae				
	Cut a 3mm² piece of the FTA Card.				
	Place it in a microtube.				
Lysis	Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>Proteinase PK.</b> <sup>1,2</sup>				
	Mix and incubate 15 minutes at room temperature.				
Loading of the instrument	Transfer the <b>whole volume</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).				

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for the preparation, storage and use of the control.

 $<sup>^2</sup>$  Add **5 \muL of EPC-Ext** included in the kit.

 $<sup>^2</sup>$  Add **5 \muL of EPC-Ext** included in the kit.

#### 14. From tissue or brain

## A. Sample preparation

# a) BVDV, CSFV, ASFV viruses (lymphoid tissues: spleen, ganglions, tonsil) and PRV virus (brain, lung) with Lysis buffer LB3.

Place 20 mg of sample in a microtube.

Continue according to the table below.

## b) BTV, EHDV, SBV (brain, spleen), influenza virus and Marek virus

Grind<sup>1 or 2</sup> **0.1** g of sample with **1 mL** of **PBS Buffer 1X** or sterile saline water.

Transfer **100 µL** of **sample** in a microtube.

Continue according to the table below.

## c) IHNV, VHSV and IPNV (spleen, anterior kidney and heart or brain)

Homogenize, by stomacher, mixer or mortar and pestle with sterile sand, re-suspend in the original transport medium with a ratio of 10 % w / v, centrifuge for 15 minutes at 4 000 g.

Grind <sup>1 or 2</sup> **0.1** g of sample with **1 mL** of **PBS Buffer 1X** or sterile saline water.

Transfer 100 µL of sample in a microtube.

Continue according to the table below.

# d) *M. hyopneumoniae*, PRRSV (lung), ASFV (lymphoid tissues: spleen, ganglions, tonsil) viruses, PCV2, PCV3, *L. intracellularis, Brachyspira*.

Place 20 mg of sample in a microtube.

NB: for the PRRS virus, pooled analysis (up to 3 samples) can be performed, n x 20 mg.

Add 1 mL of sterile saline water.

Grind (e.g., with a grinder like Mixer Mill, add a metal bead (3 mm) and grind 2 minutes at 30 Hz). Centrifuge 6 000 g/2 minutes.

Transfer 100  $\mu$ L of the supernatant of the ground sample in a microtube.

Continue according to the table below.

## e) Actinobacillus pleuropneumoniae

Vortex one biopsy in 1 mL of sterile saline water.

Place 20 µL to 100 µL of obtained liquid in a microtube.

Continue according to the table below.

# f) A. phagocytophilum, C. burnetii, Chlamydia, Leptospira., N. caninum, T. gondii and Salmonella (tissue e.g., cotyledon of placenta, foetal tissues)

The analysis from spleen is not recommended. Potential PCR inhibitors can disrupt the analysis. Rub within the tissue using dry swab.

Continue according to § 15.A.f.

## g) N. caninum and T. gondii (brain)

Mix one volume of brain and one volume of sterile saline water.

Vortex

Transfer 100 µL of the obtained liquid in a microtube.

Continue according to the table below.

<sup>&</sup>lt;sup>1</sup> For example with a grinder like Mixer Mill: add a metal bead (3 mm) and grind 2 minutes at 30 Hz then centrifuge 6 000 q/2 minutes.

<sup>&</sup>lt;sup>2</sup> For example with a grinder like Fast Prep: in a Lysing Matrix D tube, grind twice 20 seconds at 6m/sec with a 5 minutes break on ice between both. Then centrifuge 2 000 g/3 minutes.

<sup>&</sup>lt;sup>1</sup> For example with a grinder like Mixer Mill: add a metal bead (3 mm) and grind 2 minutes at 30 Hz then centrifuge 6 000 g/2 minutes.

<sup>&</sup>lt;sup>2</sup> For example with a grinder like Fast Prep: in a Lysing Matrix D tube, grind twice 20 seconds at 6m/sec with a 5 minutes break on ice between both. Then centrifuge 2 000 g/3 minutes.

## h) M. avium subsp. paratuberculosis

Place 20 mg of sample in a microtube.

Add 1 mL of sterile saline water.

Grind 10 minutes at 30 Hz (e.g., with a grinder like Mixer Mill) with a metal bead (3 mm) and 300 mg of glass beads (Bio-X Diagnostics, 480 tests: ref ADIADPBA1-480).

Centrifuge 3 000 g/5 minutes.

Transfer 100 μL of the supernatant of the ground sample in a microtube.

Continue according to the table below.

## B. Nucleic acids extraction and purification

	BVDV Virus	PRV, CSFV and ASFV viruses	M. hyopneumoniae, SBV, BTV, EHDV PRRSV, influenza, Marek, IHNV, VHSV, IPNV, ASFV viruses, PCV2 and PCV3	A. pleuropneumoniae, N. caninum, T. gondii, L. intracellularis and Brachyspira	M. avium subsp. paratuberculosis
Lysis	Add <b>350 μL</b> of <b>Lysis Buffer LB3.</b> <sup>4</sup> Mix.		Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>proteinase PK</b> . <sup>1,4</sup> Mix.		
Lysis	Incubate 15 minutes at room temperature.	Grind. <sup>2-3</sup>	Incubate 15 minutes at room temperature.	Incubate 15 minutes at +56 °C.	No incubation.
Loading of the instrument	Transfer <b>100 μL</b> <b>buffer</b> (well 1/L	_			

<sup>&</sup>lt;sup>1</sup>A pre-mix can be prepared just before use and added to each sample.

Refer to the corresponding package insert for the preparation, storage and use of the control.

<sup>&</sup>lt;sup>2</sup> For example with a grinder like Mixer Mill: add a metal bead (3 mm) and grind 2 minutes at 30 Hz then centrifuge 6 000 g/2 minutes.

<sup>&</sup>lt;sup>3</sup> For example with a grinder like Fast Prep: in a Lysing Matrix D tube, grind twice 20 seconds at 6m/sec with a 5 minutes break on ice between both. Then centrifuge 2 000 g/3 minutes.

 $<sup>^4</sup>$  Add **5 µL of EPC-Ext** included in the kit.

## 15. From swab

#### A. Sample preparation

## a) M. hyopneumoniae, PRRSV virus and PRV virus.

Add 2 mL of sterile saline water in the tube with the swab.

Vortex.

NB: for PRRSV virus, up to 3 swabs can be pooled together. Transfer the supernatant obtained in the tube with the next swab.

Press each swab to collect as much liquid as possible.

Transfer the liquid in a 2 mL-microtube.

Place 100 µL of the obtained liquid in a microtube.

Continue following the table below.

## b) FIP (feline coronavirus).

Add 2 mL of sterile saline water in the tube with the swab.

Vortex.

Press each swab to collect as much liquid as possible.

Transfer the liquid in a 2 mL-microtube.

Place 100 µL of the obtained liquid in a microtube.

Continue according to the table below.

## c) Avian Influenza Virus

Add 1 swab in a tube with 1 mL of MEM medium (+antibiotic if viral culture is needed) or sterile saline water.

Vortex.

Up to 10 swabs can be pooled together (mix 1 volume of each individual). Keep the individual mixtures at a temperature < -65 °C.

In case of pools, some weak positive samples can be not detected

Take 100 μL of the obtained liquid.

Or

Add 1 to 5 swabs into a tube with 2 mL of MEM medium (+antibiotic if viral culture is needed) or sterile saline water.

Vortex.

Up to 10 swabs can be pooled together (mix 1 volume of each individual).

Take 100 µL of the obtained liquid.

Continue according to the table below.

#### d) Swine Influenza Virus

Add 1 swab into a tube with 2 mL of MEM medium + antibiotic (to allow subsequent viral culture) or sterile saline water.

Vortex.

HO)

Take 100 µL of the obtained liquid.

Continue according to the table below.

## e) Actinobacillus pleuropneumoniae

Vortex one swab in 1 mL of sterile saline water.

Place 20  $\mu L$  to 100  $\mu L$  of the obtained liquid in a microtube.

Continue according to the table below.

## f) A. phagocytophilum, C. burnetii, Chlamydia, Leptospira., N. caninum, T. gondii, BoHV-

## 4, ASFV, Salmonella, L. intracellularis and Brachyspira

Vortex one swab in 1 mL of PBS Buffer 1X.

Place 100  $\mu$ L of the obtained liquid in a microtube.

Continue according to the table below.

g) Avian mycoplasmas, Ornithobacterium rhinotracheale and Infectious Bronchitis virus

Cut 1 to 3 swabs in 1 mL of sterile saline water.

Place 100  $\mu$ L of the obtained liquid in a microtube.

Continue according to the table below.

h) CEM: Taylorella equigenitalis, Taylorella asinigenitalis, Klebsiella pneumoniae and Pseudomonas aeruginosa

Vortex one swab in 500  $\mu L$  of PBS Buffer 1X.

Place 100  $\mu$ L of the obtained liquid in a microtube.

Continue according to the table below.

## B. Nucleic acids extraction and purification

	M. hyopneumoniae, FIP, PRRSV, PRV, influenza viruses and Infectious Bronchitis virus.	A. pleuropneumoniae, A. phagocytophilum, C. burnetii, Chlamydia, Leptospira., N. caninum, T. gondii, Salmonella, BoHV-4, Avian mycoplasma, Ornithobacterium rhinotracheale, CEMO, ASFV, L. intracellularis et Brachyspira			
Lysis	Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>proteinase PK</b> . <sup>1,2</sup> Mix.				
	Incubate 15 minutes at room temperature.	Incubate 15 minutes at +56 °C.			
Loading of the instrument	Transfer <b>the totality</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).				

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

 $<sup>^2\</sup>text{Add}$  5  $\mu\text{L}$  of EPC-Ext included in the kit of interest.

## 16. From faeces

## A. Sample preparation

## a) M. avium subsp. paratuberculosis

Note: Overnight rehydration at room temperature may be recommended for **goat and sheep faeces**. **Environmental samples** (for example soil scrapings from different breeding areas, ...) have to be treated as faecal samples. Dilute 3-10 g of sample in water.

Three ADIAMAG protocols are applicable:

## With ADIAPREP (ref. ADPREP-200)

Collect 1 spoon of faecal matter and transfer it into the ADIAPREP.

Vortex until a homogeneous suspension is obtained.

Transfer 1 mL into a clean tube, centrifuge 5 minutes at 3000 g and discard the supernatant.

Add 300 mg grinding beads and 500  $\mu L$  sterile deionized water to the pellet.

Grind for 5 minutes at 30 Hz on the Mixer Mill or 3 x 45 seconds on the Fast Prep/Ribolyser and centrifuge for 5 minutes at 3000 g.

Transfer 100 μL of sample in a microtube.

Continue according to the table below.

## • With ADIAFILTER (ref. ADIFIL100)

Collect 3 g (+/- 0.2) of faecal matter, vortex into 20 mL of sterile deionized water (or identical weight/volume ratio) until sedimentation for 10 to 20 minutes.

Place 10 mL of supernatant in the ADIAFILTER, centrifuge for 5 minutes at 3000 g and discard the supernatant and ADIAFILTER.

Add 500  $\mu$ L sterile deionized water to the pellet, vortex and transfer to a microtube containing 300 mg grinding beads.

Grind for 10 minutes at 30 Hz on Mixer Mill or 3 x 45 seconds on Fast Prep/Ribolyser and centrifuge for 5 minutes at 15 000 g.

Transfer 100  $\mu\text{L}$  of sample in a microtube.

Continue according to the table below.

## • Without prefiltration system

Collect 3 g (+/- 0.2) of faecal matter (or identical weight/volume ratio), vortex into 20 mL of sterile deionized water until sedimentation for 10 to 20 min.

Transfer 1 mL to a microtube containing 300 mg of grinding beads.

Grind for 10 minutes at 30 Hz on Mixer Mill or 3 x 45 seconds on Fast Prep/Ribolyser and centrifuge for 5 minutes at 15 000 g.

Transfer 100 µL of sample in a microtube.

Continue according to the table below.

## b) C. burnetii

## Add 5 mL of PBS Buffer 1X to 1 g of faeces.

Homogenize, e.g., with a vortex, for at least 15 seconds.

Centrifuge 3 000 g at 2 minutes.

Transfer 100 µL of sample in a microtube.

Continue according to the table below.

## c) T. gondii

Weigh 1 g of faeces in a previously labelled 10 mL or 15 mL sterile tube.

Add 10 mL of PBS Buffer 1X (This preparation is stable for 24 hours at room temperature).

Vortex until a homogenous solution is obtained.

Allow to settle 2 to 5 minutes.

Transfer **500 μL** of **supernatant** in a previously identified microtube.

Centrifuge at 3 000 g for 5 minutes. Discard the supernatant.

Homogenise the pellet with 1 mL of PBS Buffer 1X (This solution is stable for 24 hours at room temperature).

Transfer **500 μL** of **supernatant** in a tube containing 300 mg of glass beads.

Disrupt 10 minutes at 30 Hz with a Mixer Mill (or transfer the obtained solution in a Lysis Matrix B tube and disrupt 3 x 45 seconds at 4 m/sec with the Fast Prep).

Centrifuge at 15 000 g 5 minutes.

Transfer 100  $\mu L$  of supernatant in a microtube.

Continue according to the table below.

## d) Influenza virus, L. intracellularis and Brachyspira

Weigh 1 g of manure and add 5 mL of physiological water

Vortex until a homogenous solution is obtained.

Sediment 5 minutes.

Transfer **100 μL** of **supernatant** in a microtube.

Continue according to the table below.

## B. Nucleic acids extraction and purification

	M. avium subsp. paratuberculosis (with ADIAPREP)	M. avium subsp. paratuberculosis (with ADIAFILTER)  C. burnetii, T. gondii, L. intracellularis and Brachyspira	M. avium subsp. paratuberculosis (without prefiltration system) Influenza virus	
Lyse	Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>Proteinase PK + 5 μL of EPC-Ext</b> . <sup>1,2</sup> . Mix.			
	No incubation.	Incubate 15 minutes at +56 °C.	Incubate 15 minutes at room temperature.	
Chargement de l'automate	Transfer <b>the totality</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).			

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

<sup>&</sup>lt;sup>2</sup>Add **5 μL** of **EPC-Ext** included in the kit of interest.

## 17. From environmental sample (drag swab)

## A. Sample preparation

## a) M. avium subsp. paratuberculosis

Environmental samples (dung scrapings, holding area...) are treated as faeces. Continue to § 16.A.a).

# b) Influenza Virus, Marek virus, Infectious Bronchitis virus, ASFV, L. intracellularis and Brachyspira

Mix drag swab in a maximum volume of 40 mL of STP or PBS 1X Buffer. 70 mL for an overshoe. Centrifuge 1 mL of supernatant 10 to 15 minutes at 800 to 1 200 g.

Transfer 100  $\mu\text{L}$  of supernatant in a microtube.

Continue according to the table below.

## B. Nucleic acids extraction and purification

	Marek virus	Influenza virus Infectious Bronchitis virus, ASFV, L. intracellularis and Brachyspira
Lysis	Add 100 μL of Lysis Buffer LB1 + 10 μL of Proteinase PK + 5 μL of EPC-Ext. <sup>1,2</sup> Mix.	
	incubate 15 minutes <b>at +56 °C.</b>	incubate 15 minutes at room temperature.
Loading of the instrument	Transfer <b>the totality</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).	

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

<sup>&</sup>lt;sup>2</sup>Add **5 μL** of **EPC-Ext** included in the kit of interest.

## 18. From sperm or egg fish

## A. Sample preparation

## a) IHNV, VHSV, IPNV (liquid sperm)

Mix one volume of sperm and one volume of PBS 1X.

Vortex 20 seconds.

Centrifuge 2 minutes at 6 000 g.

Transfer 100 µL of the obtained liquid in a microtube.

Continue according to the table below.

## b) IHNV, VHSV, IPNV (coagulated sperm, egg)

Homogenize, by stomacher, mixer or mortar and pestle with sterile sand, re-suspend in the original transport medium with a ratio of 10 % w / v, centrifuge for 15 minutes at 4 000 g.

Grind <sup>1 or 2</sup> **0.1** g of sample with **1 mL** of **PBS Buffer 1X** or sterile saline water.

Transfer 100  $\mu$ L of sample in a microtube.

Continue according to the table below.

## B. Nucleic acids extraction and purification

	IHNV, VHSV, IPNV	
	Add 100 μL of Lysis Buffer LB1 + 10 μL of Proteinase PK. <sup>1,2</sup>	
Lysis	Mix.	
	Incubate 15 minutes <b>at room temperature.</b>	
Loading of the instrument	Transfer <b>the totality</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).	

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

<sup>&</sup>lt;sup>1</sup> For example with a grinder like Mixer Mill: add a metal bead (3 mm) and grind 2 minutes at 30 Hz then centrifuge 6 000 g/2 minutes.

<sup>&</sup>lt;sup>2</sup> For example with a grinder like Fast Prep: in a Lysing Matrix D tube, grind twice 20 seconds at 6m/sec with a 5 minutes break on ice between both. Then centrifuge 2 000 g/3 minutes.

<sup>&</sup>lt;sup>2</sup>Add **5 μL** of **EPC-Ext** included in the kit of interest.

## 19. From Whatman 3 blotting paper

	ASFV	
Sample preparation	Cut 5 pieces and place them in a microtube.  Add 1 mL PBS 1X. Mix and incubate 15 minutes at room temperature.  Transfer <b>100 μL</b> of sample in a microtube	
Lysis	Add 100 μL of Lysis Buffer LB1 + 10 μL of Proteinase PK. 1,2  Mix.  Incubate 15 minutes at room temperature.	
Loading of the instrument	Transfer <b>the totality</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).	

<sup>&</sup>lt;sup>1</sup>A pre-mix can be prepared just before use and added to each sample.

## 20. From muscle exsudate

	ASFV	
Sample preparation	Transfer <b>100 μL</b> of sample in a microtube	
Lysis	Add <b>100 μL</b> of <b>Lysis Buffer LB1 + 10 μL</b> of <b>Proteinase PK</b> . 1,2  Mix.  Incubate 15 minutes <b>at room temperature.</b>	
Loading of the instrument	Transfer <b>the totality</b> in the <b>binding buffer</b> (well 1/Line B/plate 1).	

<sup>&</sup>lt;sup>1</sup> A pre-mix can be prepared just before use and added to each sample.

## 21. From semen

	ASFV, PRRSV	
Sample preparation	Transfer <b>100 μL</b> of sample in a microtube	
Lysis	Add <b>250 μL</b> of <b>Lysis Buffer LB2</b> <sup>1</sup> Mix.	
Loading of the instrument	Transfer 200 μL in the <b>binding buffer</b> (well 1/Line B/plate 1).	

 $<sup>^{1}\</sup>text{Add}~\textbf{5}~\mu\text{L}~\text{of}~\textbf{EPC-Ext}$  included in the kit of interest.

 $<sup>^2</sup>$ Add **5**  $\mu$ L of **EPC-Ext** included in the kit of interest.

See the kit of interest user manual to prepare, store and use this control.

 $<sup>^2\</sup>mbox{Add}$  5  $\mu\mbox{L}$  of EPC-Ext included in the kit of interest.

See the kit of interest user manual to prepare, store and use this control.

# **VI.Amplification**

For the amplification of extracted nucleic acids, please refer to "Amplification" and "Interpretation of results" paragraphs of the ADIAVET™ or ADIALYO™ kit of interest user manual.

## VII. Index of symbols

Symbol	Meaning
REF	Catalog number
***	Manufacturer
<b>1</b>	Upper temperature limit
	Use by date
LOT	Batch code
[]i	Consult Instructions for Use
Σ	Contains sufficient for <n> tests</n>
类	Keep away from sunlight
VET	For veterinary in vitro use only

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