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Manual

MutaCLEAN® Mag RNA/DNA extraction kit

For extraction of nucleic acids using magnetic beads

Gültig ab/Valid from 2022-05-23

→ REF KG1023 KG1024 $\sum_{200/800}^{\Sigma}$ IVD CE



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1 INTENDED USE

MutaCLEAN® Mag RNA/DNA Kit is a magnetic bead-based extraction kit for nucleic acids (RNA and DNA) which may be used e.g. for subsequent real time PCR analysis. The kit is an accessory for an in vitro diagnostic device. Any in vitro diagnostic use of this sample preparation procedure in combination with a downstream nucleic acid test must be evaluated for the parameter in question. The nucleic acids can be extracted from clinical or environmental samples incl. food samples. The kit is intended to be used by professional users in a laboratory environment and the extraction can be performed manually or using an automated platform (e.g. KingFisher[™] Flex or Duo Prime, Microlab STAR, Freedom EVO, Biocomma M32, M96).

2 MODE OF ACTION

- a) Samples are lysed by incubation in working solution (binding buffer [P1] substituted with 2-propanol). Nucleic acids are bound to the magnetic beads.
- b) Bound nucleic acids are washed with Inhibitor removal buffer (P2) in order to remove PCR inhibitors from the sample.
- c) Bound nucleic acids are washed with Wash Buffer (P3) in order to purify them from salts, proteins and other cellular impurities.
- d) Purified nucleic acids are eluted from the magnetic beads with elution buffer (P4).

Purified nucleic acids can be used directly for downstream applications.

3 COMPONENTS

MutaCLEAN® Mag RNA/DNA (KG1023) is designed for 200 isolations.

MutaCLEAN® Mag RNA/DNA (KG1024) is designed for 800 isolations.

	Labelling	Content (KG1023)	Content (KG1024)
P1	Binding buffer	2 x 30 ml	2 x 120 ml or 1 x 240 ml
PA	Poly A/carrier RNA	1 x 4 mg	4 x 4 mg
P2	Inhibitor removal buffer	2 x 33 ml	2 x 132 ml or 1 x 264 ml
P3	Wash buffer	2 x 20 ml	4 x 40 ml or 2 x 80 ml
P4	Elution buffer	1 x 21 ml	1 x 84 ml
MB	MutaCLEAN [®] magnetic beads	4x 1.0 ml	1 x 16.0 ml

Table 1: Components of the MutaCLEAN® Mag RNA/DNA isolation kit.

Labelling		Content (KG1023)	Content (KG1024)
	Proteinase K*	1 x 200 mg	1 x 800 mg (respectively 4 x 200 mg)

* Proteinase K is provided separately along with the kit.

All solutions are clear and should not be used when precipitates have formed. Warm up the solutions at 18 to 25 $^{\circ}$ C or in a 37 $^{\circ}$ C water bath until the precipitates have dissolved.

4 EQUIPMENT AND REAGENTS TO BE SUPPLIED BY USER

Note: consumables not included in the kit are dependent of the mode of sample preparation, e.g. manual extraction or extraction using automated magnetic processors. Therefore, the customer needs to decide which consumables are necessary for the extraction process.

- · Laboratory equipment according to national safety instructions
- Nuclease-free 1.5 or 2.0 ml tubes
- Separation plate for magnetic beads separation, e.g. square-well Block (96-well block with 2.1 ml square-wells)
- · Elution plate for collecting purified nucleic acids
- Pipets with sterile pipet filter tips or tip combs (e.g. KingFisher 96tip comb for DW magnets)
- Tabletop microcentrifuge capable of 13,000 g centrifugal force
- absolute ethanol
- 2-propanol
- Thermoblock or laboratory furnace
- · Magnetic particle processor or magnetic separator

5 TRANSPORT, STORAGE AND STABILITY

The MutaCLEAN[®] Mag RNA/DNA Kits are shipped at ambient temperature. Kits must be stored at +18 to +25 °C. If properly stored, all kit components are stable until the date of expiry printed on the label.

Please note, that improper storage at +2 to +8 °C (refrigerator) or \leq -18 °C (freezer) will adversely impact nucleic acid purification when precipitates form in the solutions.

Reconstituted poly A/carrier RNA solution has to be aliquoted. Aliquots stored at \leq -18 °C are stable through date of expiry printed on kit label.

6 GENERAL NOTES ON THE TEST AND TEST PROCEDURE

- Read the Instruction for Use carefully before using the product.
- This instruction for use replaces the version dated 2021-01-05.
- This product has been produced and placed on the market in accordance with the regulation (EU) 2017/746 (IVDR).
- All serious incidents occurring in connection with the product must be reported to Immundiagnostik AG and (within the Union market) to the competent reporting authority of the respective member state.
- The MutaCLEAN[®] Mag RNA/DNA Kit must be utilised by qualified personnel only.
- Good Laboratory Practice (GLP) has to be applied.
- Clinical samples must always be regarded as potentially infectious material and all equipment used has to be treated as potentially contaminated.
- Binding buffer (P1) and inhibitor removal buffer (P2) contain guanidine hydrochloride which is an irritant. Always wear gloves and follow standard safety precautions.
- Do not let these buffers touch your skin, eyes, or mucous membranes. If contact does occur, wash the affected area immediately with large amounts of water; otherwise, the reagent may cause burns. If you spill the reagent, dilute the spill with water before wiping it up.
- Always wear gloves and follow standard safety precautions when handling these buffers.
- Do not pool reagents from different lots or from different bottles of the same lot. Immediately after usage, close all bottles in order to avoid leakage, varying buffer-concentrations or buffer conditions. After first opening, store all bottles in an upright position.
- Do not use a kit after its expiration date.
- Do not use any modified ethanol.

6.1 Waste Handling

• Dispose of unused reagents and waste should occur in accordance with country, federal state and local regulations. • Material Safety Data Sheets (MSDS) are available upon request from Immundiagnostik AG.

7 PREPARATION OF SOLUTIONS

Table 2: Preparation of MutaCLEAN® Mag RNA/DNA solutions.

Label	I	Storage and	
Label	KG1023	KG1024	stability
Poly A/ carrier RNA (PA)	Dissolve in 1 ml elution buffer and prepare aliquots.	Dissolve each vial in 1 ml elution buffer and prepare aliquots.	Store at ≤ -18°C, stable through date of expiry printed on kit label.
Binding buffer (P1)	Add 26 ml 2-propanol to each vial, mix well. Label and date bottle accordingly.	Filling 2 x 120 ml: Add 104 ml 2-propanol to each vial, mix well. Filling 1 x 240 ml: Add 208 ml 2-propanol to vial, mix well. Label and date bottle accordingly.	
Inhibitor removal buffer (P2)	Add 20 ml absolute ethanol to each vial, mix well. Label and date bottle accordingly.	Filling 2 x 132 ml: Add 80 ml ethanol to each vial, mix well. Filling 1 x 264 ml: Add 160 ml ethanol to vial, mix well Label and date bottle accordingly.	Store at 18 to 25 °C. Stable through the date of expiry printed on kit label.
Wash buffer (P3)	Add 80 ml absolute ethanol to each vial, mix well. Label and date bottle accordingly.	Filling 4 x 40 ml: Add 160 ml ethanol to each vial, mix well. Filling 2 x 80 ml: Add 320 ml ethanol to vial, mix well Label and date bottle accordingly.	

8 SAMPLE MATERIAL

Purification of nucleic acids from a wide range of sample material, such as the following:

- Human samples (EDTA blood, tissue, stool, urine, etc.)
- Veterinary samples (EDTA blood, tissue, raw milk, etc.)
- Insects and ticks
- Food samples (milk, drinking water)

- Environmental samples
- Plant material

Table 3: Volumes of binding buffer (P1) and pre-treatment for various sample matrices.

Sample material	Volume / Amount	Volume binding buffer	Pre-treatment of the sample
Stool, faeces	pea-size	500 µl	Prepare a suspension in 1.5 ml PCR- grade water. Vortex and briefly spin down sediments. Use 200 µl of the supernatant.
Swabs		500 µl	Add 500 µl PCR-grade water to a dry swab, suspend vigorously and use 200 µl of the suspension.
Liquid samples*	200 µl	500 µl	
Tissues	≤ 30 mg	500 µl	Homogenisation of tissue in 500 µl PCR-grade water. Spin down for 1 min at 8.000 <i>g</i> , use 200 µl of the supernatant.
Cells	≤ 2 x 10 ⁶	500 µl	Pellet up to 2 x 10 ⁶ cells. Resuspend pellet in 200 µl PCR-grade water. Homogenise and spin down for 1 min at 8 000 g. Use 200 µl of the supernatant.

* EDTA blood, serum, amniotic fluid, CSF, urine, water, milk etc.

9 HANDLING OF MAGNETIC BEADS

A homogeneous distribution of the magnetic beads to the individual wells of the separation plate is essential for a high well-to-well consistency. Therefore, before distributing the beads, make sure that beads are completely resuspended. Shake storage vial well or vortex briefly. Premixing magnetic beads with binding buffer allows easier homogenous distribution of the beads to the individual wells of the separation plate. During automation, a premix step before aspirating the beads / binding buffer mixture from the reservoir is recommended.

10 EXTRACTION OF NUCLEIC ACIDS

Before starting, prepare a working solution of the binding buffer (P1) supplemented with poly A/carrier RNA (PA) and proteinase K for at least one sample (N) more than required in order to compensate pipetting inaccuracies.

Samples containing precipitates must be centrifuged before purification!

Store eluted nucleic acid at ≤-18 °C for later analysis.

Table 4: Preparation of the working solution

Volume needed per sample	Mastermix working solution
500 μl binding buffer (P1)	500 μl x (N+1)
4 μl poly A/carrier RNA (PA)	4 μl x (N+1)
50 µl proteinase K [20 mg/ml]	50 μl x (N+1)

10.1 Protocol for manual use

This protocol is for manual use and serves as a guideline for adapting the kit to robotic instruments..

Step 1

- Add 20 µl MutaCLEAN[®] magnetic beads (MB) to a nuclease-free 2.0 ml microcentrifuge tube. Vortex MutaCLEAN[®] magnetic beads vigorously before pipetting.
- Add 550 µl working solution, freshly prepared, to each tube.
- Add 200 µl sample to each tube.
- Mix immediately.
- Perform incubation for 10 min at 60 °C.
- Following the incubation, centrifuge briefly to collect any sample from the lysis tube lid.

Step 2

• Separate the magnetic beads against the side of the tubes by placing the tubes on a magnetic separator. Wait at least 30 s until all the beads have been attracted to the magnets. Remove and discard supernatant by pipetting. Do not disturb the attracted beads while aspirating the supernatant.

Step 3

- Remove the tubes from the magnetic separator.
- Add 500 µl inhibitor removal buffer (P2) and resuspend the beads by shaking (optionally mix by pipetting up and down) until the beads are resuspended completely.

• Separate the magnetic beads against the side of the tubes by placing the tubes on the magnetic separator. Wait at least 30 s until all the beads have been attracted to the magnets. Remove and discard supernatant by pipetting. Do not disturb the attracted beads while aspirating the supernatant.

Step 4

- Remove the tubes from the magnetic separator.
- Add 450 µl Wash Buffer (P3) and resuspend the beads by shaking (optional mix by pipetting up and town) until the beads are resuspended completely.
- Separate the magnetic beads against the side of the tubes by placing the tubes on the magnetic separator. Wait at least 30 sec until all the beads have been attracted to the magnets. Remove and discard supernatant by pipetting. Do not disturb the attracted beads while aspirating the supernatant.

Step 5

• Repeat step 4.

Step 6

• Air-dry the magnetic bead pellet for 5-10 min at room temperature.

Step 7

- Remove the tubes from the magnetic separator.
- Add 100 µl elution buffer (P4) and resuspend the beads by shaking (optionally mix by pipetting up and down) until the beads are resuspended completely.
- Incubate for 10 min at room temperature.
- Separate the magnetic beads against the side of the tubes by placing the tubes on the magnetic separator. Wait at least 30s until all the beads have been attracted to the magnets.
- The supernatant contains purified nucleic acid.
- Transfer the supernatant to fresh nuclease-free tubes.

10.2 Protocol for KingFisher[™] Flex Magnetic Particle Processor

Protocols for other automated magnetic particle processors need to be adapted accordingly.

Step 1

- Add 20µl MutaCLEAN[®] magnetic beads (MB) to each well of an empty 96 deep-well block. Vortex MutaCLEAN[®] magnetic beads vigorously before pipetting.
- Add 550 µl working solution, freshly prepared, to each well.
- Add 200 µl sample to each well.

Step 2 - Prepare wash plates

- Add 500 μl inhibitor removal buffer (P2) to each well of an empty 96-well deep-well block.
- Add 450 µl Wash Buffer (P3) to each well of an empty 96-well deep-well block.
- Add 450 µl Wash Buffer (P3) to each well of a second empty 96-well deep-well block.

Step 3 - Prepare elution plate

• Add 100µl elution buffer (P4) to each well of an empty 96-well deep-well block.

Step 4 - Run purification protocol on instrument

- Insert plates as indicated on the KingFisher[™] Flex Magnetic Particle Processor.
- Method starts with a mixing step (combined lysis and binding step) after setting up the last plate to the instrument.

Step 5 – Remove elution plate

- The instrument stops after the final elution step. Follow the instructions on the instruments display and unload the plates from the instrument.
- The eluates contain purified nucleic acids.
- For storage purposes cover the elution plate with an adhesive foil.

For KingFisher[™] Flex Magnetic Particle Processor use the settings profile shown in Table 5 and Table 6.

Table 5: Reagent Information

Tip plate	Microtiter DW 96 plate	
Lysis/Binding		
Name	Well volume [µl]	Туре
Magnet Beads	20	Reagent
Working Solution	550	Reagent
Extraction Control	See Instruction Manual of the respective PCR Kit.	Reagent
Sample	200	Sample
Inhibitor Removal Buffer	Inhibitor Removal	
Name	Well volume [µl]	Туре
		_
Inhibitor Removal	500	Reagent
Inhibitor Removal 1st Wash Buffer	500 Inhibitor Removal	Reagent
		Reagent <i>Type</i>
1st Wash Buffer	Inhibitor Removal	
1st Wash Buffer Name	Inhibitor Removal <i>Well volume [µl]</i>	Туре
1st Wash Buffer <i>Name</i> Wash Buffer	Inhibitor Removal <i>Well volume [μl]</i> 450	Туре
1st Wash Buffer <i>Name</i> Wash Buffer 2nd Wash Buffer	Inhibitor Removal <i>Well volume [µl]</i> 450 2nd Wash Buffer	<i>Type</i> Reagent
1st Wash Buffer Name Wash Buffer 2nd Wash Buffer Name	Inhibitor Removal Well volume [μ]] 450 2nd Wash Buffer Well volume [μ]]	<i>Type</i> Reagent <i>Type</i>
1st Wash Buffer Name Wash Buffer 2nd Wash Buffer Name Wash Buffer	Inhibitor Removal Well volume [µl] 450 2nd Wash Buffer Well volume [µl] 450	<i>Type</i> Reagent <i>Type</i>

Table 6: Instrument Settings

Table 0. Instrument settings					
الاللكالير	Tip 1		96 DW tip comb		
	<u>م</u>	Pick-Up	Tip plate		
	Ċ	Binding	Lysis		
		Beginning of step Mixing / heating End of step	Pause Precollect Release beads Mixing time, speed Heating during mixing Heating temperature [°C] Postmix Collect count	No Yes 00:10:00, Bottom mix Yes 60 No 4 3	
	Å	Inhibitor Removal	Collect time [s]	3	
		Buffer Beginning of step	Precollect	No	
		Mixing / heating	Release time, speed Shake 1 time, speed Shake 2 time, speed Heating during mixing Postmix Collect count Collect time [s]	00:00:30, Medium 00:00:30, Bottom mix 00:00:30, Half mix No No 4 3	
	Å	1st Wash Buffer	1st Wash Buffer		
		Beginning of step Mixing / heating End of step	Precollect Release time, speed Shake 1 time, speed Shake 2 time, speed Heating during mixing Postmix Collect count Collect time [s]	No 00:00:30, Medium 00:00:30, Bottom mix 00:00:30, Half mix No No 3 2	
	ů	2nd Wash Buffer	2nd Wash Buffer		
		Beginning of step Mixing / heating End of step	Precollect Release time, speed Mixing time, speed Heating during mixing Postmix Collect count Collect time [s]	No 00:00:30, Medium 00:01:00, Bottom mix No 3 2	
	3333	Bead Drying			
			Dry time Tip position	00:05:00 Outside well / tube	
	3	Elution	Elution		
		Beginning of step Mixing / heating End of step	Precollect Release time, speed Mixing time, speed Heating temperature [°C] Preheat Postmix Collect count Collect time [s]	No 00:00:30, Fast 00:10:00, Slow 56 Yes No 5 4	
	\sim	Leave	Tip plate		

11 TROUBLESHOOTING

The following troubleshooting guide is included to help you with possible problems that may arise when isolating nucleic acid from different types of sample material. Especially when working with complex sample matrices such as fatty tissue, whole blood or highly contaminated environmental samples, preparation of samples can be crucial. For protocols on sample materials not covered in this manual or for further questions concerning nucleic acid extraction, please do not hesitate to contact our scientists on info@immundiagnostik.com.

Low nucleic acid yield	
Sample not sufficiently lysed	Supplement the working solution (Bind- ing Buffer + PolyA/Carrier RNA (PA) with 50 µl Proteinase K (20 mg/ml) per sample).
Incomplete Proteinase K digestion	Be sure to dissolve the lyophilized Protein- ase K completely, as follows:
	1. Pipette appropriate volume of PCR grade water to lyophilised Proteinase K in order to get a concentration of 20 mg/ ml (e.g. 2.5 ml PCR grade water to 50 mg Proteinase K).
	2. Close vial and invert until all the lyoph- ilisate (including any stuck to the lid) is completely dissolved. 3. Aliquot the rehy- drated enzyme, mark each aliquot with the date of reconstitution, and store at ≤-18°C. Rehydrated Proteinase K is stable for 12 months when stored properly.
Insufficient elution buffer volume	Bead pellet must be covered completely with elution buffer.
Aspiration of attracted bead pellet	Do not disturb the attracted beads while aspirating the supernatant, especially when the magnetic bead pellet is not vis- ible in the lysate.
Aspiration and loss of beads	Time for magnetic separation too short or aspiration speed too high.

Low nucleic acid yield	
Insufficient washing procedure	Use only the appropriate combinations of separator and plates. Make sure that beads are resuspended completely during the washing procedure. If shaking is not sufficient to resuspend the beads com- pletely mix by repeated pipetting up and down.
Carry-over of ethanol from wash buffers	Be sure to remove all of the ethanolic wash solution from the final wash, as re- sidual ethanol interferes with downstream applications.
Ethanol evaporation from wash buffers	Close buffer bottles tightly, avoid ethanol evaporation from buffer bottles as well as from buffer filled in reservoirs. Do not reuse buffers from buffer reservoirs.
Time for magnetic separation too short	Increase separation time to allow the beads to be completely attracted to the magnetic pins before aspirating any liquid from the well.
Aspiration speed too high (elution step)	High aspiration speed during elution step may cause bead carry-over. Reduce aspira- tion speed for elution.
Kit stored under non-optimal conditions.	Store kit at $+18$ to $+25$ °C at all times upon arrival.
Buffers or other reagents were ex-	Store all buffers at +18 to +25 °C.
posed to conditions that reduced their effectiveness	Close all reagent bottles tightly after each use to preserve pH and stability and to prevent contamination.
	Aliquot proteinase K and poly A/carrier RNA (PA) after reconstitution and store aliquots at \leq -18°C.

Low nucleic acid yield			
2-propanol not added to binding buffer (P1)	Add 2-propanol to the buffer before using. Mix the buffer well and store at +18 to 25 °C.		
	Always mark the buffer vial to indicate whether 2-propanol has been added or not.		
Ethanol not added to inhibitor removal buffer (P2) and/or wash	Add absolute ethanol to the buffers be- fore using.		
buffer (P3)	After adding ethanol, mix the buffers well and store at $+18$ to $25 ^{\circ}$ C.		
	Always mark the buffer vials to indicate whether ethanol has been added or not.		
Reagents and samples not com- pletely mixed	Always mix the sample tube well after ad- dition of each reagent.		
Impurities not completely removed	Perform a second wash step with wash buffer (P3) in order to completely remove salts, proteins and other residual impuri- ties from the bound nucleic acid.		

12 KIT PERFORMANCE

The scope of the validation was to show that the performance characteristics Muta-CLEAN® Mag RNA/DNA and if the method meets the requirements of the intended application, that is to extract DNA and RNA from various biological and environmental samples.

During validation, MutaCLEAN® Mag RNA/DNA was tested against other commercial extraction kits using standardized samples. Kits B to M mentioned in Tables 8 and 9 represent recommended kits of the manufacturers for the respective sample materials. The quality and quantity of extracted nucleic acids were determined using real time PCR and real time RT-PCR. The extractions of nucleic acids were performed according to the manufacturer's instructions.

12.1 Sample material

Table 7: Overview of the samples tested.

Sample	Pathogens detected	Genomic DNA detected
Avian faeces	Influenzaviruses	nd
Buccal swabs	Influenzaviruses, adenovirus, respiratory syncytial virus, <i>M. tuberculosis</i>	nd
Cerebrospinal fluid	Enteroviruses, tick-borne encephalitis virus	nd
Bacterial cultures	<i>E. coli,</i> Streptococci, <i>Legionella</i> , Mycobacteria incl. <i>Mycobacterium tuberculosis</i> complex, <i>Salmonella, Listeria, Campylobacter, Shigella</i>	nd
Bovine blood samples	Bovine viral diarrhea virus (BVD)	nd
Bovine brain samples	Schmallenberg virus	nd
Bovine faeces	Mycobacterium avium ssp. paratuberculosis	nd
Bovine raw milk	Mycobacterium avium ssp. paratuberculosis, E. coli, Streptococci, Yeast, Pseudomonas	nd
Bovine tissue samples	Coxiella burnetii (Q Fever)	yes
Drinking water	Legionella	nd
Human blood samples	Cytomegalovirus	yes
Human epithel	nd	yes*
Human hair with root	nd	yes*
Human muscle	nd	yes*
Human nails	nd	yes*
Human sperm	nd	yes*
Human sputum	nd	yes*
Human teeth	nd	yes*
Human urine samples	Cytomegalovirus	nd
Human stool samples	Norovirus, sapovirus, astrovirus, rotavirus, adenovirus, <i>Salmonella, E. coli</i>	nd
Ovine faeces	Mycobacterium avium ssp. paratuberculosis	nd

Sample	Pathogens detected	Genomic DNA detected
Tissue culture samples	Varicella zoster virus, cytomegalovirus, Epstein-Barr virus, Enteroviruses, Poliovi- ruses, Herpes simplex virus 1+2, Influenza- viruses, respiratory syncytial virus, rotavirus, adenovirus, <i>Babesia</i>	nd
Ticks	Tick-borne encephalitis virus, <i>Borrelia, Ehrli-</i> chia, Babesia	yes

* Samples were tested in a forensic lab.

The samples were either field samples, positive for pathogens (e.g. bovine feces and milk positive for *Mycobacterium avium ssp. paratuberculosis*, bovine ear notch samples positive for BVD, porcine saliva positive for PRRSV, ticks positive for *Borrelia* and TBEV, bovine tissue samples positive for *Coxiella burnetii*) or sample material was artificially spiked with pathogens or, in case of forensic samples, human genomic DNA should be isolated. If spiking was done, the sample materials were spiked with the respective pathogens, natively found in these materials in infected subjects (e.g. urine spiked with cytomegalovirus, buccal swabs spiked with influenzaviruses).

12.2 DNA extraction

The following table shows an overview of the performance of DNA extraction (genomic, bacterial, viral) using MutaCLEAN® Mag RNA/DNA (A) in comparison to competitors, indicated by characters in the first row of table 8.

The +/++/+++ indicate the DNA yield and outcome of the subsequently performed real time PCR for the respective pathogens mentioned in table 8 (Stratagene Mx3005P, Roche LightCycler 480II):

- + = Ct range >32
- ++ = Ct range 26 32
- +++ = Ct range <26
- na = not applicable
- nd = not done

Table 8: Comparison of DNA extraction efficiencies

Sample	Α	В	С	D	Ε	F	G	н	I.	К	L	М
Avian faeces	+++	na	na	na	na	na	na	na	na	na	na	na
Buccal swabs	+++	na	+++	nd	++	na	na	na	na	++	++	na
Cerebrospinal fluid	+++	na	+++	na	+++	na	na	na	na	na	++	++

Sample	Α	В	С	D	Ε	F	G	Н	I	K	L	М
Bacterial cultures	+++	na	++	na	na	na	na	na	na	na	++	na
Bovine blood samples	+++	na	++	++	na	++	na	na	na	na	++	na
Bovine brain samples	+++	na	na	++	na	na	na	++	na	na	na	na
Bovine feces	+++	na	++	na	na	na	++	na	na	na	na	na
Bovine raw milk	+++	na	na	na	na	na	na	na	na	na	na	na
Bovine tissue samples	+++	na	na	++	na	na	na	++	na	na	na	na
Drinking water	+++	na	+++	+++	na	+++	na	na	na	na	nd	na
Human blood samples	+++	na	+++	+++	na	+++	na	na	na	na	nd	+++
Human epithel	++	na	nd	nd	na	na	na	++	na	na	na	na
Human hair with root	++	na	nd	nd	na	na	na	+++	na	na	na	na
Human muscle	+++	na	nd	nd	na	na	na	+++	na	na	na	na
Human nails	+	na	nd	nd	na	na	na	+++	na	na	na	na
Human sperm	+++	na	nd	nd	na	na	na	++	na	na	na	na
Human sputum	+++	na	nd	nd	na	na	na	+++	na	na	na	na
Human teeth	+	na	nd	nd	na	na	na	+++	na	na	na	na
Human urine samples	+++	na	nd	na	++	na	na	na	na	na	na	na
Human stool samples	+++	+++	nd	na	na	na	na	na	na	na	na	na
Ovine faeces	++	na	na	na	na	na	na	na	na	na	na	na
Tissue culture samples	+++	na	na	nd	na	na	na	++	na	na	na	na
Ticks	+++	na	na	nd	na	na	na	++	na	na	na	na

The results shown in table 8 indicate that MutaCLEAN® Mag RNA/DNA can be used for the extraction of DNA from a variety of different sample materials. For the extraction of genomic DNA from human nails and teeth, a bead-beating step before using MutaCLEAN® Mag RNA/DNA is recommended. The results shown for these materials are without bead-beating prior to extraction. Furthermore, for the extraction of Mycobacteria DNA from feces and sputum, and RNA and DNA from ticks, a beadbeating or other mechanical disruption pre-extraction treatment is also highly recommended.

12.3 RNA extraction

The following table shows an overview of the performance of viral RNA extraction using MutaCLEAN[®] Mag RNA/DNA (A) in comparison to competitors, indicated by characters in the first row of the table.

The +/++/+++ indicate the RNA yield and outcome of the subsequently performed real time RT-PCR for the respective pathogens mentioned in Table 9 (Stratagene Mx3005P, Roche LightCycler 480II):

- + = Ct range >32
- ++ = Ct range 26 32
- +++ = Ct range <26
- na = not applicable
- nd = not done

Table 9: Comparison of RNA extraction efficiencies

Sample	Α	В	С	D	Ε	F	G	Н	I	Κ	L	М
Avian faeces	+++	+++	na									
Buccal swabs	+++	+++	na	++								
Cerebrospinal fluid	+++	+++	nd	na	na	na	na	na	na	++	na	++
Bovine blood	+++	++	++	na	na	na	na	na	na	++	na	++
Bovine brain samples	+++	++	na	na	na	na	na	na	++	na	na	na
Bovine faeces	+++	na	++	na								
Bovine raw milk	+++	na	na	na	na	na	na	na	na	na	na	na
Bovine tissue	+++	++	na									
Human urine	+++	na	nd	na								
Human stool	+++	+++	nd	na	na	na	++	na	na	na	na	na
Ovine faeces	++	na	na	na	na	na	na	na	na	na	na	na
Tissue culture	+++	+++	na	nd	na	na	na	na	++	na	na	na
Ticks	+++	na	na	nd	na	na	na	na	++	na	na	na

The results shown in table 9 indicate that MutaCLEAN® Mag RNA/DNA can be used for the extraction of RNA from a variety of different sample materials. For the extraction of RNA from ticks, a bead-beating or other mechanical disruption pre-extraction treatment is highly recommended.

13 ABBREVIATIONS AND SYMBOLS

DNA	Desoxyribonucleic acid		Catalog number / To be used with				
RNA	Ribonucleid acid	Σ Σ	Contains sufficient for <n> test</n>				
PCR	Polymerase chain reaction	ł	Limit of temperature				
RT	Reverse transcription	***	Manufacturer				
BINDING BUFFER P1	Binding buffer (P1)	><	Use by YYYY-MM-DD				
IR BUFFER P2	Inhibitor removal buffer (P2)	LOT	Batch code				
WASH BUFFER P3	Wash buffer (P3)	CONT	Content				
ELUTION BUFFER P4	Elution buffer (P4)	i	Consult instructions for use				
POLY A/CARRIER RNA PA	Poly A/carrier RNA (PA)	IVD	<i>In vitro</i> diagnostic medical device				
MAGNETIC BEADS MB	Magnetic beads	CE	European Conformity				
UDI	Unique Device Identification	Â	Attention				
(٢)	The product has been classified and marked in ac- cordance with EU Directives / Ordinance on Haz- ardous Materials.	Acute toxicity Skin irritation	; Category 4, H302 ; Category 4; H332 , Category 2; H315 Category. 2; H319				

14 LITERATURE

- 1. James H. Jorgensen , Michael A. Pfaller, Karen C. Carroll. Manual of Clinical Microbiology, 11th Edition, 2015.
- 2. Richard L. Hodinka, Benjamin Pinsky. Clinical Virology Manual, 5th Edition, 2016.