MACHEREY-NAGEL

NucleoMag[®] 96 Virus

Use of MACHEREY-NAGEL's NucleoMag® 96 Virus kit for the detection of Influenza virus A/H1N1



Abstract

The 2009 flu pandemic is a global outbreak of a new strain of influenza A virus subtype H1N1. The outbreak was first observed in Mexico, with evidence that there had been an ongoing epidemic for months before it was officially recognized as such. In the present study we describe the automated use of the MACHEREY-NAGEL NucleoMag[®] 96 Virus kit on the KingFisher[®] Flex magnetic separator for the isolation of viral nucleic acids and subsequent detection by PCR. For kit evaluation A/H1N1 samples from an interlaboratory test as well as clinical samples have been used.

Introduction

The so called "swine flue" is caused by the swine influenza virus (A/H1N1). This virus was discovered first in year 1930 in pig populations. The recently discovered subtype A of the H1N1 virus includes genes derived by reassortment from human, swine and avian viruses, which can be transmitted from human to human. As for all influenza caused infections a reliable and sensitive detection of the virus is an important prerequisite to identify suspicious patients and to support infection control.

For the detection of influenza viruses rapid tests using antibodies to detect virus proteins are available and commonly used. However, tests are often less sensitive in comparison to RT-PCR based methods and prone to false negative results. Therefore the RT-PCR based methods are considered as the gold standard.

Typically swabs or nasopharyngeal swab sample are taken from patients and used for viral RNA extraction. There are high demands for the RNA extraction methods with regard to sensitivity and reproducibility. Moreover, since usually a nigh number of samples have to be screened the extraction is typically automated with a high throughput.

In the current application note we describe the use of the MACHEREY-NAGEL magnetic beads based NucleoMag[®] Virus kit in combination with the Thermo King-Fisher[®] Flex magnetic particle separator. Swabs are washed and incubated for example in NaCl solution. The method starts with a lysis incubation of the swab wash solution and subsequently, after addition of binding buffer, the released nucleic acids are bound to magnetic particles. Following several washing steps the purified viral RNA

is eluted and can be used for RT-PCR downstream applications. The use of the KingFisher[®] Flex separation device allows the purification of up to 96 samples within approx 60 min.

Material and methods

The kit evaluation was done by Labcon OWL, Bad Salzuflen, Germany, with clinical samples and INSTAND interlaboratory test samples.

Samples that have been used in the study comprise:

- INSTAND round robin samples have been used throughout the validation. Samples are suitable for the detection of human Influenza A and B viruses as well as new Influenza A subtype H1N1 (swine flue).
- Samples are provided as lyophilized lysates of tissue culture media or allantoic liquid from infected and incubated hen's eggs.
- Samples have been extracted with the NucleoMag[®] 96 Virus kit automated on a KingFisher[®] Flex magnetic separator. After elution aliquots of the eluates have been used in conjunction with two different PCR system for the detection of Influenza A and A/H1N1.

In addition to the samples from the interlaboratory test swab samples from potential positive patients were subjected for viral RNA isolation.

The purification protocol used is summarized in Table 1.

Table 1: Purification protocol for viral RNA isolation using the NucleoMag[®] 96 Virus kit on the KingFisher[®] Flex instrument.

	KingFisher [®] Flex 96	
Plate 1	KF 96 Plate and KF 96 Tip comb	
Plate 2 (Thermo deep-well plate)	200 μL sample, 200 μL Lysis Buffer MV1, 10 μL Proteinase K, 4 μL carrier RNA	
	Mixing 15 min, fast, 56 °C	
	Add 600 μL Binding Buffer MV2 and 30 μL V-Beads	
	Mixing: 5 min, bottom mix, collect 3x	
Plate 3 (Thermo deep-well plate)	Wash Buffer MV3, 500 µL Mixing: 2 min, half mix, collect 3x	

	KingFisher [®] Flex 96
Plate 4	Wash Buffer MV4, 500 µL
(Thermo deep-weil plate)	Mixing: 2 min half mix, collect 3x
Plate 5 (Thermo deep-well plate)	Wash Buffer MV5, 550 µL
	90 sec, slow, beads are not resuspended in Wash Buffer MB5
Plate 5	Air-drying step, 3 min
(Thermo deep-well plate)	
Plate 6 KF96 plate	Elution Buffer MB6, 70 µL
	heating (preheat) 56 °C,
	Mixing: 5 min, slow, collect 3x
	release beads into plate 4

For downstream PCR analysis two different PCR systems have been used.

RT PCR amplification was performed according to the recommendations of the Robert Koch-Institute (RKI) or using a commercial detection kit (Astra Diagnostics, Germany).

Results

1) Method development using influenza INSTAND interlaboratory test samples

RT-PCR according to RKI recommendations (Lightcycler[®] 480) This RT-PCR system uses two independent mastermixes. One RT-PCR reaction detects Influenza A viruses, while the second PCR reaction is specific for subtype H1N1. Consequently, a sample which is positive for H1N1 should be positive in both PCR reactions. Influenza A subtypes different from H1N1 should yield only positive signals in the Influenza A reaction.

The results are summarized in Table 2.

Table 2: Results from INSTAND interlaboratory test samples. Real-Time PCR according to the recommendations of the RKI.

		In House PCR (TIB)	
RR-no.	sample specification In House PCR (TIB)	Influenza A	Influenza H ₁ N ₁
20088	Positiv for Influenza B (seasonal)	negativ	negativ
20089	Positiv for Influenza A/ H1N1 (swine lineage)	positiv	positiv
20090	Positiv for Influenza A/ H3N2 (seasonal)	positiv	negativ
20091	Negativ for Influenza A and B	negativ	negativ
20092	Positiv for avian Influenza A/H5N1 (seasonal)	positiv	negativ
20093	Positiv for Influenza A/ H1N1 (seasonal)	positiv	negativ
20094	Positiv for Influenza A/ H1N1 (swine lineage)	positiv	positiv
20095	Positiv for avian Influenza A/H5N1	positiv	negativ

RT-PCR kit from Astra Diagnistics, Hamburg, Germany (Influenza Screen & Type RT-PCR Kit 1.0)

This PCR system uses a mastermix for the detection of both Influenza A and subtype H1N1 (the primers bind to similar target sequences). Only one of the primers (either Influenza A or H1N1) bind to the target and only one signal is generated: one PCR reaction detects Influenza A viruses, while the second PCR reaction is specific for subtype H1N1. Consequently, a sample which is positive for H1N1 should be negative for Influenza A. Influenza A subtypes different from H1N1 should yield only positive signals in the Influenza A reaction.

The results are summarized in Table 3.

Table 3: Results from INSTAND interlaboratory test samples. Real-Time RT-PCR kit from Astra Diagnostics.

		Astra Kit	
RR-no.	sample specification	Influenza	Influenza
	In House PCR (TIB)	А	H ₁ N ₁
20088	Positiv for Influenza B (saisonal)	negativ	negativ
20089	Positiv for Influenza A/ H1N1 (swine lineage)	negativ	positiv
20090	Positiv for Influenza A/ H3N2 (saisonal)	positiv	negativ
20091	Negativ for Influenza A and B	negativ	negativ
20092	Positiv for avian Influenza A/H5N1 (saisonal)	positiv	negativ
20093	Positiv for Influenza A/ H1N1 (saisonal)	positiv	negativ
20094	Positiv for influenza A/ H1N1 (swine lineage)	negativ	positiv
20095	Positiv for avian Influenza A/H5N1	positiv	negativ

As can be seen from Table 2 and 3 with both PCR systems all samples from the interlaboratory test have been identified correctly. Neither false positives nor false negatives have been observed.

2) Testing of the method with swab samples from patients

The isolated viral RNA from swab samples was subjected to RT-PCR reactions specific for either Influenza A or H1N1. RT-PCR setup was performed according to the recommendations of the Robert Koch-Institute on a Roche Lightcycler[®] 480 instrument. Amplification plots are shown in figure 1 (influenza A) and figure 2 (H1N1). Calculated C_P values are summarized in Figure 3.



Figure 1: Amplification plots for Influenza A specific RT-PCR



Figure 2: Amplification plots for H1N1 specific RT-PCR



Figure 3: Summary for RT-PCR detection of H1N1 (upper figure) and Influenza A (lower figure). Amplification controls: black bars: negative controls; blue bars: positive controls.

All samples which were tested positive for the presence of the H1N1 virus (Fig. 3, top) were also tested positive for Influenza A (Fig. 3, bottom), clearly identifying the samples positive for the swine virus A/H1N1. All negative H1N1 also proved to be negative for Influenza A. In both RT-PCR systems the negative controls were always negative whereas positive controls were positive.

Summary

With both PCR systems all samples from the interlaboratory test have been identified correctly. Neither false positives nor false negatives have been observed. For swab samples consisten results were obtained. This supports the excellent quality of the eluted viral RNA and the unlimited use for downstream RT-PCR.

Acknowledgement

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Ordering information

Product	Specifications	Preps	REF
NucleoMag [®] 96 Virus	Kit based on magnetic bead technology for the isolation of viral RNA and DNA from serum or plasma.	1 x 96 / 4 x 96	744800.1 / .4
KingFisher [®] Accessory Kit A	Square-well Blocks, Deep-well Tip Combs, Elution Plates, for 4 x 96 NucleoMag [®] Virus preps using KingFisher [®] Flex platform	1 set	744950

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