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Rapid and generic identification of influenza A and other respiratory viruses with mass spectrometry



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The rapid identification of existing and emerging respiratory viruses is crucial in combating outbreaks and epidemics. Matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS) is a rapid and reliable identification method in bacterial diagnostics, but has not been used in virological diagnostics. Mass spectrometry systems have been investigated for the identification of respiratory viruses. However, sample preparation methods were laborious and time-consuming. In this study, a reliable and rapid sample preparation method was developed allowing identification of cultured respiratory viruses. Tenfold serial dilutions of ten cultures influenza A strains, mixed samples of influenza A virus with human metapneumovirus or respiratory syncytial virus, and reconstituted clinical samples were treated with the developed sample preparation method. Subsequently, peptides were subjected to MALDI-TOF MS and liquid chromatography tandem mass spectrometry (LC-MS/MS). The influenza A strains were identified to the subtype level within 3 h with MALDI-TOF MS and 6 h with LC-MS/MS, excluding the culturing time. The sensitivity of LC-MS/MS was higher compared to MALDI-TOF MS. In addition, LC-MS/MS was able to discriminate between two viruses in mixed samples and was able to identify virus from reconstituted clinical samples. The development of an improved and rapid sample preparation method allowed generic and rapid identification of cultured respiratory viruses by mass spectrometry.

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1. Introduction

Respiratory viruses are a major cause of infections, natural outbreaks, and epidemics. Approximately 200 million cases of viral community-acquired pneumonia occur every year: 100 million in children and 100 million in adults (Ruuskanen et al., 2011). A wide range of viruses, including current circulating subtypes of influenza A virus (influenza A(H1N1)pdm09 and influenza A(H3N2)), influenza B virus, respiratory syncytial virus (RSV), parainfluenza viruses, adenovirus, and rhinovirus, have been implicated in respiratory tract infections in past decades. In the last ten years, new viruses have emerged as severe acute respiratory syndrome (SARS) virus, Middle East respiratory syndrome (MERS), avian influenza A (H5N1) virus, human metapneumovirus (hMPV), coronaviruses

NL63 and HKU1, and human bocavirus. Rapid identification of existing and emerging respiratory viruses is of the utmost importance in combating outbreaks and epidemics, and starting early therapy and prophylaxis. Currently, cell culture, serology, and real-time reverse transcription-polymerase chain reaction (rRT-PCR) are used in virological diagnostics. However, there are a few drawbacks with these techniques: (i) serology and rRT-PCR-based assays are target directed and thus potentially miss non-selected or emerging pathogenic viruses (Binnicker et al., 2013; Yang et al., 2014); (ii) culturing is time consuming; (iii) serology-based assays are not applicable in the acute phase and/or have low sensitivity. Furthermore, multiple tests are needed to detect and subtype mixed infections.

Matrix-assisted laser desorption/ionization-time of flight mass spectrometry (MALDI-TOF MS) is a generic technique that can rapidly identify cultured microorganisms (Seng et al., 2009; van Veen et al., 2010). The analysis of bacteria by mass spectrometry (MS) has made great progress over the last two decades, and it is

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employed in many hospitals as a rapid and reliable alternative to traditional identification methods. Thus far, mass spectrometry-based methods have not been used for virological diagnostics. However, it has been shown that purified influenza virus particles could be identified with either MALDI-TOF MS or MALDI Fourier Transform Ion Cyclotron Resonance MS (MALDI-FT-ICR MS). Identification was based on the proteolytic digestion of concentrated and purified viral samples and on mass spectrometry analysis of the specific peptide mass profile. The sample preparation methods used in these studies included virus concentration and/or purification with either ultracentrifugation, differential centrifugation, precipitation, filtration, isolation of viral particles or protein(s) with an affinity capture immunoassay or gel electrophoresis (Downard et al., 2009; Schwahn et al., 2009a,b, 2010a,b; Chou et al., 2011; Jang et al., 2011; Nguyen and Downard, 2013; Fernandes and Downard, 2014). These sample preparation procedures were laborious and often time consuming (up to 24 h) and are therefore not applicable for high-throughput virological diagnostics.

The aim of this study was to develop a rapid, generic and robust sample preparation method for MALDI-TOF MS and LC-MS/MS that will enable reliable and fast identification of respiratory viruses. For this purpose, cultured influenza A virus strains and mixed samples of influenza A virus with hMPV or RSV were treated with the developed preparation method. Subsequently, obtained peptides were subjected to MALDI-TOF MS and LC-MS/MS for analyses. The identification of the respiratory viruses was based on peptide sequence differences in abundant viral proteins. To confirm correct identification of peptides by mass spectrometry, the amino acid sequences were compared to the corresponding DNA sequences, obtained by sequencing of the viruses. Finally, to determine the sensitivity of the in-house developed method the titers of cultured viruses were determined with rRT-PCR.

2. Materials and methods

2.1. Ethics statement

All of the clinical influenza strains, anonymized from routine diagnostics, originated from a collection at the University Medical Center Utrecht (UMCU). Collection of the samples and analysis of the isolated virus strains were approved by the local Medical Ethics Committee of the UMCU. The institutional review board (IRB) confirmed (protocol 12/320) that the viral strains were not regarded as patient-owned material; consequently, the use of these strains was not restricted by Dutch law ("Law Medical Scientific Research with People", WMO; art. 1b).

2.2. Viruses

Nasopharyngeal swabs and tracheal aspirates were collected between 2009 and 2011 from patients who were hospitalized with respiratory distress symptoms in the wards of the UMCU (Table S1). The collected samples were cultured immediately in either LLC-MK2 or HEp2 cell lines and were routinely checked for cytopathological effect (CPE) formation. The samples exhibiting specific CPE in 80% of the cells and testing positive for respiratory viruses by real-time PCR were harvested and stored at -80°C .

Next, eight influenza A H1N1 strains (designated here as BM1456, BM1457, and BM1480 through BM1485) and two influenza A H3N2 strains (designated here as BM1454 and BM1455) were cultured in LLC-MK2 cells in the presence of EMEM (Eagle's minimal essential medium) with trypsin at 37°C (Table S1). In addition, hMPV (designated here as BM1460) and RSV type A (designated here as BM1450) were cultured in LLC-MK2 cells in the presence of EMEM with trypsin at 37°C and in HEp2 cells in the presence of DMEM (Dulbecco's modified Eagle medium) with 5%

FBS (fetal bovine serum) at 33°C , respectively (Table S1). Subsequently, the culture supernatants were collected, and the cell debris was removed by centrifugation ($239 \times g$, 10 min). Aliquots were frozen at -80°C and were used without further purification steps.

2.3. Viral RNA extraction, cDNA synthesis, and rRT-PCR

To determine the sensitivity of the in-house developed sample preparation method the titers of cultured viruses were determined with rRT-PCR by using a standard curve. This standard curve was developed by counting virus particles using electron microscopy and by subsequently performing PCR.

Viral RNA extraction and PCR were performed according to previously described protocols (Tan et al., 2012). In short, viral genomic RNA was isolated using a MagnaPure LC total nucleic acid kit, according to the manufacturer's guidelines (Roche Diagnostics, Mannheim, Germany). Murine encephalomyocarditis virus was used as an internal control. Reverse transcription of the isolated viral RNA was performed using a MultiScribe reverse transcriptase kit and random hexamers (Applied Biosystems, Foster City, CA, USA), according to the manufacturer's guidelines. PCR primers and probes were designed on the basis of highly conserved genomic regions of the M1 gene for influenza A and the N gene for both hMPV and RSV A, and these primers and probes were used for the typing of the viral strains. The primers and probes used are listed in Table S2.

cDNA samples were analyzed in a $25 \mu\text{l}$ reaction mixture, containing $10 \mu\text{l}$ of cDNA, TaqMan universal PCR master mix (Applied Biosystems, ABI), primers and fluorogenic probes labeled with the 5' reporter dye 6-carboxy-fluorescein (FAM), and the 3' quencher dye 6-carboxy-tetramethyl-rhodamine (TAMRA). Amplification and detection were performed with an ABI 7500 system for 2 min at 50°C , 10 min at 95°C , and 45 cycles of 15 sec at 95°C and 1 min at 60°C . The samples were assessed for the presence of possible inhibitors of the amplification reaction using the indicated internal control, the signals of which had to range within a clear-cut interval.

2.4. Viral genomic cDNA synthesis and sequencing

To confirm the presence and amino acid sequence accuracy of the identified peptides, six of the clinical isolates were sequenced. The RSV strain (BM1450) with accession number JQ901450 was sequenced previously (Tan et al., 2012). For sequencing purposes, hMPV (BM1460) and influenza (BM1454, BM1456, BM1457, BM1480, and BM1483) PCR fragments were obtained by fractional amplification of MagnaPure LC genomic RNA isolates, using the Superscript III one-step RT-PCR System with Platinum Taq High Fidelity kit (Invitrogen) and a 9800 Fast thermal cycler (ABI), according to the manufacturers' protocols. Unlike hMPV and RSV, influenza virus contains a segmented genome consisting of 8 segments. All of the segments were completely amplified and purified from an agarose gel prior to fractional amplification, as previously described (Zhou et al., 2009). The PCR products were applied to a 1% agarose gel and were purified from the gel with a GeneJET Gel Extraction Kit (Thermo Fisher Scientific, Landsmeer, the Netherlands), according to the manufacturer's protocol. The isolated fragments were used for whole-genome sequencing.

The hMPV (BM1460) and influenza strains (BM1454, BM1456, BM1457, BM1480, and BM1483) listed in Table S1 were sequenced according to the whole-genome sequence protocol described recently, using the conventional Sanger technique (Tan et al., 2012). Fragments ranging between 650 and 1400 nucleotides were sequenced with an ABI 3730 48-capillary DNA analyzer, using Big-Dye Terminator 3.1 (ABI). The resulting sequence information was assembled into an hMPV whole-genome sequence through alignment with the corresponding reference sequence (GenBank: FJ168779; <http://www.ncbi.nlm.nih.gov/genbank/index.html>)

using Seqman software (DNASTAR lasergene 8); a similar approach was followed for the influenza strains. Table S3 provides an overview of the primers used for influenza, hMPV, and RSV sequencing. The DNA and amino acid sequences of the sequenced strains are provided in Supplementary Data.

2.5. Sample preparation of cultured viruses

The starting titers of the viral cultures are shown in Table S1. The starting titers for the influenza strains and RSV appeared roughly similar ($\sim 3 \times 10^{11}$ copies/ml), but the starting titer for hMPV was 100-fold lower. Tenfold serial dilutions for all of the strains (influenza A virus, RSV, and hMPV) were prepared in CyMol medium (Copan Italia, Brescia, Italy). CyMol is a collection, transport, and preservation medium for cells (cytology analysis), viruses, and nucleic acids (PCR-based assays) (Luinstra et al., 2011a). Subsequently, 200 μ l of each tenfold dilution step was treated according to the in-house developed sample pretreatment protocol, as follows. A 200 μ l aliquot was mixed with 25 μ l of a 0.5% stock solution of RapiGest (Waters Corporation, Milford, CT, USA) and was incubated for 7 min at 95 °C. Next, DL-dithiothreitol (DTT, Sigma–Aldrich) and acetonitrile (LC–MS Chromasolv, Sigma–Aldrich) were added to 5 mM and to 25% final concentration, respectively. Reduction and precipitation of the proteins were performed under controlled microwave radiation in a Rapid Enzymatic Digestion System (REDS; Hudson Surface Technology [HST] Inc., Old Tappan, NJ, USA) at 60 °C and 400 W for 10 min. The subsequent carbamidomethylation of cysteine residues was executed at 37 °C and 400 W for 5 min in REDS, after the addition of iodoacetamide (IAA; Sigma–Aldrich) to a 12.5 mM final concentration. Next, the proteinaceous particles were precipitated at 20,800 $\times g$ for 7 min, and the supernatants were discarded. The pellets were suspended in 40 μ l of 50 mM ammonium bicarbonate (Sigma–Aldrich) containing 4 μ g/ml modified and TPCK-treated trypsin from bovine pancreas (Sigma–Aldrich). Digestion was performed for 10 min in REDS at 37 °C and 400 W and was terminated by the addition of trifluoroacetic acid (TFA) to 1% final concentration. After incubation at 37 °C for 30 min, the RapiGest precipitates were removed by centrifugation, and the supernatants were filtered through a Microcon YM-10 filter (10 MWCO, Merck Millipore Ltd.; Co. Cork, Ireland). All of the samples were prepared at least twice.

For the analysis of mixtures of two different viral culture samples, each tenfold dilution of influenza A H3N2 (BM1454) or H1N1 (BM1456) was mixed with the corresponding dilution of the RSV (BM1450) or hMPV (BM1460) culture samples, respectively, followed by the exact sample preparation procedure described above. All the samples were prepared at least twice. To test whether human proteins present in clinical material interfere with the developed sample preparation method or identification by LC–MS/MS, reconstituted throat swabs were included. A swab that was rubbed around the tonsils of a volunteer was washed by vortexing the swab in CyMol medium. Subsequently, this so-called clinical sample was spiked with dilutions of cultured influenza A virus (BM1454, BM1456), RSV (BM1450), or hMPV (BM1460). The same sample preparation and LC–MS/MS analysis procedure as described above were performed on these reconstituted clinical samples. All tested samples were prepared at least twice.

2.6. MALDI-TOF MS analysis and database search

A 10 μ l aliquot of each sample was desalted on a C₁₈ ZipTip (Merck Millipore Ltd.), and the peptides were eluted with 1 μ l of 50% acetonitrile containing 10 mg/ml α -cyano-4-hydroxycinnamic acid (HCCA) and 2.5% trifluoroacetic acid (TFA). Elution was performed directly onto the MALDI target (MTP Polished steel, Bruker Daltonics, GmbH, Bremen, Germany), and the droplets were

air-dried. Spectra were recorded in reflector mode, using a MALDI-TOF MS instrument (Autoflex III, Bruker). Each spectrum was acquired by averaging 2000 laser shots (at 40% laser power) across a mass range of m/z 500–4000 followed by noise reduction. The instrument was mass calibrated with an external peptide standard (Peptide Calibration Standard, Bruker). The spectra were analyzed with FlexAnalysis and Biotoools (Bruker). Internal calibration was performed using identified peptides derived from viral nucleoprotein or trypsin. From the spectra, peak lists were generated and searched against the NCBI virus database (database collection of viruses: 2013.06.29) using MASCOT, version 2.2.04 (Matrix Science, London, UK) with a mass tolerance of 70 ppm (Perkins et al., 1999; Koenig et al., 2008). The carbamidomethylation of cysteine residues and oxidation of methionine were included in the calculations, and up to two missed cleavages were permitted. The MASCOT scoring system was used to determine protein expectation values that corresponded to the number of matches that were expected to occur by chance alone. The highest expectation value was considered the best match. All of the assays were performed at least twice.

In separate experiments, the peaks that were identified initially as nucleoprotein (NP) peptides of influenza A were subsequently confirmed by MALDI-TOF MS/MS. The MS/MS spectra were recorded in the reflector mode using LIFT technology (Bruker).

2.7. LC–MS/MS analysis and database search

The digests were analyzed by LC–MS/MS using a nano-Advance LC system (Bruker) coupled to a Q-TOF mass spectrometer (maXis impact, Bruker). A 5 μ l aliquot of each digested sample was injected onto a Magic C18AQ UHPLC NanoTrap column (C18, 100 μ m ID \times 100 mm, 5 μ m, 200 Å, Bruker) and was washed with loading solvent H₂O (0.2% formic acid) for 5 min at a flow rate of 4 μ l/min. Following valve switching, the peptides were separated on a Magic C18AQ analytical column (100 μ m ID \times 150 mm, C₁₈, 3 μ m, 200 Å, Bruker) at a constant flow of 800 nl/min. The peptide elution gradient was from 98% A (H₂O with 0.1% formic acid) to 45% B (CH₃CN) in 30 min, followed by an increase to 95% B for 5 min. Optima LC–MS H₂O (0.1% formic acid; Fisher Scientific) was used for the LC–MS/MS studies.

The nanoLC system was coupled to the mass spectrometer using a CaptiveSpray (Bruker) ionization source. The spray voltage was set at 1.4–1.6 kV, and the temperature of the heated capillary was set at 150 °C. The eluting peptides were analyzed using the data-dependent MS/MS mode. The ten most abundant ions (charge state 2⁺, 3⁺, and 4⁺) in an MS spectrum (400–1400 m/z) were selected for data-dependent MS/MS analysis by collision-induced dissociation, using nitrogen as the collision gas. The MS/MS scans were acquired over a mass range of 100–2000 m/z .

Peak lists were generated using DataAnalysis software, version 4.1 (Bruker), and the lists were exported as MASCOT Generic (MGF) files. These files were searched against the NCBI database (taxonomy: viruses) with the MASCOT search algorithm (MASCOT 2.2.04, Matrix Science). A mass tolerance of 50 ppm and an MS/MS tolerance of 25 mmu were used. Up to two missed cleavages for trypsin were allowed; carbamidomethylcysteine was selected for fixed modification and oxidation of methionine for variable modification. Only significant protein hits with at least two unique peptides with a score greater than 20 were selected.

3. Results

3.1. Sample preparation and identification of influenza A viruses by MALDI-TOF MS

Ten influenza A virus strains, isolated from hospitalized patients and characterized to the subtype level (eight H1N1 and two H3N2),

Table 1
Identified influenza A viruses based on nucleoprotein peptide detection by MALDI-TOF MS.

Strain ^a	Subtype	Id. score ^b	Seq. cov. ^c (%)	Identified as nucleoprotein of ^d (subtype/host)
BM1454	H3N2	90	30	Influenza A virus (H3N2/human)
BM1455	H3N2	108	35	Influenza A virus (H3N2/human)
BM1456	H1N1	91	37	Influenza A virus (H1N1/human)
BM1457	H1N1	89	31	Influenza A virus (H1N1/human)
BM1480	H1N1	95	41	Influenza A virus (H1N1/swine)
BM1481	H1N1	102	36	Influenza A virus (H1N1/human)
BM1482	H1N1	89	34	Influenza A virus (H1N1/human)
BM1483	H1N1	93	36	Influenza A virus (H1N1/swine)
BM1484	H1N1	95	32	Influenza A virus (H1N1/swine)
BM1485	H1N1	103	37	Influenza A virus (H1N1/human)

^a Viral strains were tested at approximately 1×10^9 genome copies in the total volume of each sample deposited onto the MALDI plate and analyzed.

^b Protein identification score given by MASCOT. A score ≥ 76 was considered significant identification.

^c Sequence coverage of nucleoprotein, as specified by MASCOT in percentage, based on the number of amino acids covered by mass values (peptides) matched with the identified protein.

^d The nucleoprotein that received the highest MASCOT score is shown. The influenza subtype should be considered as an indication because the identification was based on detection of nucleoprotein peptides.

were cultured in LLC-MK2 cells, resulting in viral titers ranging from 2.1 to 4.3×10^{11} genome copies/ml (Table S1). After the removal of cell debris, tenfold dilution series of the viral cultures were prepared in CyMol medium. Each dilution was treated according to the in-house developed sample preparation procedure. Subsequently, peptides were subjected to MALDI-TOF MS and were analyzed by MASCOT software. In two independent experiments, all of the tested influenza A strains (eight H1N1 and two H3N2) were correctly identified based on the recognition of peptides derived from nucleoprotein (Table 1 and Fig. 1). The ten influenza A strains were not only identified to their type, but also correctly identified to their subtype (Table 1). The entire procedure, *i.e.* the dilution series preparation followed by the treatment according to the in-house protocol, data analysis and the obtaining of results, was performed within 3 h for ten samples.

The identification limit at which the viral strains were correctly typed and subtyped appeared to be 1×10^9 viral genome copies in

the total sample volume deposited onto a MALDI plate for analysis (corresponding to a CT value of approximately 24; Table 1). At this viral titer, more than 20 mass peaks could be assigned with high significance to peptides derived from nucleoprotein (Table 2). The identity of approximately half of these peptide peaks was confirmed by MALDI-TOF MS/MS (data not shown). The intensities of the remaining peptide peaks assigned to nucleoprotein were too low for MS/MS analysis. The genomes of five influenza A strains, namely BM1454, BM1456, BM1457, BM1480, and BM1483, were sequenced, and the amino acid sequences of all of the identified peptides were found to be encoded in the corresponding genome sequences (Supplementary Data, genomic and amino acid sequences of sequenced strains).

Of the other proteins that constitute the viral proteome, only M1-derived peptides were identified occasionally (data not shown). The MS spectra also contained peaks from abundant non-viral proteins derived from cell lysates, such as actin and

Table 2
Peptides derived from influenza A nucleoprotein and detected by MALDI-TOF MS.

H1N1 strains	Observed [M+H] ⁺	H3N2 strains	Observed [M+H] ⁺
SALILR	672.4	SALILR	672.4
NLPFER	775.4	DS	2184.1
QDATEIR	832.4	QNATEIR	831.4
AMMDQVR	850.4	AMVDQVR	818.4
KTGGPIYR	891.5	KTGGPIYR	891.5
SRYWAIR	951.5	SGYWAIR	852.4
GKFTAAQR	1006.5	GKFTAAQR	1006.5
MVLSAFDER	1067.5	MVLSAFDER	1067.5
GVGTIAMELIR	1159.7	ND	–
LIQNSITIER	1186.7	DS	2207.2
MVLSAFDERR	1223.6	MVLSAFDERR	1223.6
MCSLMQGSTLPR	1380.6	MCSLMQGSTLPR	1380.6
ELILYDKEEIR	1420.7	ELVLYDKEEIR	1406.7
ATVMAAFSGNNEGR	1424.6	STIMAAFTGNTEGR	1455.7
ELILYDKEEIRR	1576.9	ELVLYDKEEIRR	1562.8
NPGNAEIEDLIFLAR	1671.9	NPGNAEIEDLIFLAR	1671.9
ASAGQISVQPTFSVQR	1675.8	ASAGQTSVQPTFSVQR	1663.9
MMESAKPEDLSFQGR	1725.8	MMEGAKPEEVSFR	1510.7
SQLVWMACHSAAFEDLR	2020.9	SQLVWMACHSAAFEDLR	2020.9
ESRNPNGAEIEDLIFLAR	2044.0	ESRNPNGAEIEDLIFLAR	2044.0
SYEQMETGGERQDATEIR	2099.9	SYEQMETGDRQNATEIR	2142.9
SCLPACVYGLAVASGHDFER	2209.0	ND	–
DS	775.4; 1424.6	NLPFEKSTIMAAFTGNTEGR	2184.1
DS	1186.7; 1067.5	LIQNSLTIEKMVLSAFDER	2207.2
LLQNSQVSLMRPNENPAHK	2275.2	LLQNSQIYSLIRPNENPAHK	2335.2
GVQIASNENVETMDSNTLELR	2320.2	GVQIASNENMDNMGSTLELR	2266.0

Peptides with a S/N of ≥ 6 are listed. These peptides were identified in tested viral strains at titers of 1×10^9 genome copies in the total volume of each sample, deposited onto a MALDI plate and analyzed.

ND, peptide not detected.

DS, detected sequence in the tested strains but in another peptide due to missed cleavage by trypsin.

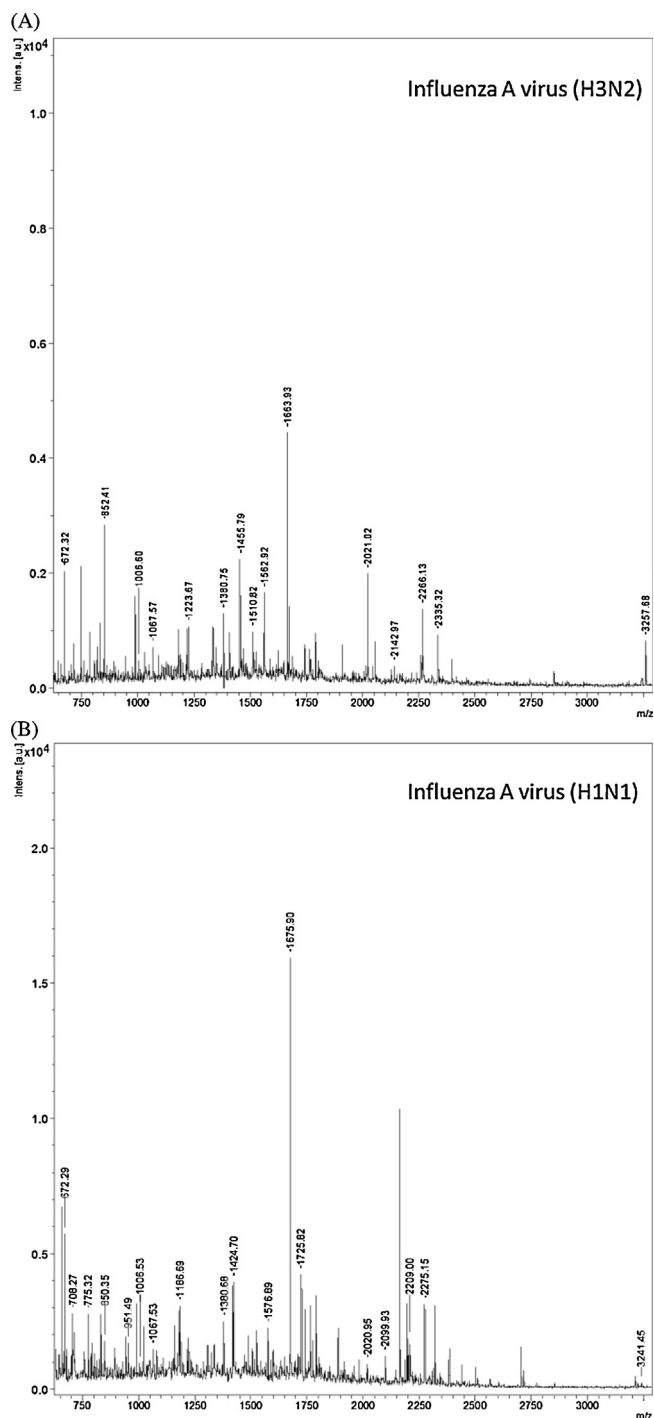


Fig. 1. Examples of MALDI-TOF mass spectra from whole influenza A virus tryptic digests of (A) H3N2 and (B) H1N1. The shown *m/z* ratios are examples of the detected peptides derived from influenza nucleoprotein.

trypsin auto-digestion fragments (shown as un-annotated peaks in Fig. 1).

3.2. Identification of influenza A viruses by LC–MS/MS

The same sample preparation procedure was followed as described above, after which the peptides were subjected to LC–MS/MS. Virus identification, including the sample preparation and analysis, was achieved in less than 6 h for ten samples. In two independent experiments, all ten influenza strains were correctly

identified as either H1N1 or H3N2 influenza A virus, with an identification limit of 7×10^6 genome copies in the total sample volume subjected to LC–MS/MS analysis (corresponding to a CT value of approximately 30–32; Table 3). At this viral titer, the influenza viruses were typed and subtyped based on the identification of peptides derived from the nucleoprotein protein, indicated in boldface in Table 4. The nucleoprotein-derived peptides that were identified repeatedly in at least 90% of the samples containing $\geq 7 \times 10^6$ viral genome copies are also shown in Table 4. In addition to the peptides derived from NP, the LC–MS/MS analysis identified peptides derived from M1, non-structural protein (NS1), HA, and NA (Table S4). The amino acid sequence coverage of these proteins (for each tenfold dilution of viral culture, an average value of two independent experiments was calculated) is shown in Table 3. Similar to the MALDI-TOF MS results, the identified amino acid sequences of the five sequenced strains (BM1454, BM1456, BM1457, BM1480, and BM1483) were in accordance with their respective genome sequences. The LC–MS/MS spectra also demonstrated proteins derived from cell lysates or culture additives, such as actin, lectin, histone, and enolase (data not shown). Moreover, most of the peptides detected with MALDI-TOF MS were also identified with LC–MS/MS (Tables 2 and 4).

3.3. Identification of respiratory viruses in mixed samples

Tenfold serial dilutions of strain BM1454 (H3N2) were mixed with the corresponding dilution factor of RSV (mixture A), resulting in an approximate 1:1 ratio of both strains. Similarly, in mixture B, tenfold serial dilutions of strain BM1456 (H1N1) were mixed with the corresponding dilution factor of hMPV, although the starting titer of hMPV was 100 times lower than those of influenza and RSV. The mixtures were treated according to the in-house developed sample preparation procedure, subjected to LC–MS/MS, and subsequently analyzed as described for the samples containing a single virus.

Using LC–MS/MS, both viruses in mixtures A and B were identified simultaneously based on the peptides derived from nucleoprotein proteins. The influenza strains were correctly typed and subtyped from sample dilutions containing approximately $\geq 5.3 \times 10^7$ genome copies in the total sample volume subjected to LC–MS/MS analysis. For influenza, peptides derived from M1, NS1, HA, and NA proteins were also identified (Table 5). For the hMPV and RSV strains, in addition to nucleoprotein-derived peptides, those of matrix protein (M), matrix protein 2–1 (M2-1), phosphoprotein (P), and fusion glycoprotein (F) were also identified (Tables 5, S5, and S6). The identification limit at which the RSV strain was identified correctly, was 3.6×10^7 genome copy equivalents in the total sample volume subjected to LC–MS/MS analysis (corresponding to a CT value of approximately 28; Tables 5 and S1). For hMPV, the identification limit was 1.1×10^7 genome copy equivalents in the total sample volume subjected to LC–MS/MS analysis (corresponding to a CT value of approximately 28; Tables 5 and S1). All of the identified peptide sequences of the tested influenza A, RSV, and hMPV strains were confirmed with their respective genome sequence data (Supplementary Data, genomic and amino acid sequences of sequenced strains).

3.4. Identification of reconstituted clinical samples by LC–MS/MS

Tenfold serial dilutions of influenza A H1N1 (BM1456), H3N2 (BM1454), RSV (BM1450), and hMPV (BM1460) were spiked in CyMol medium, containing material from a throat swab. These reconstituted clinical samples were treated according to the in-house developed sample preparation method, subjected to LC–MS/MS and analyzed as described above. All the spiked respiratory viruses were identified. In both influenza A and RSV samples,

Table 3
Influenza A viruses identified by LC–MS/MS.

Strains	Subtype	Titer ^a	Sequence coverage ^b (%)					Identified as ^c
			NP	M1	NS1	HA	NA	
BM1454-55	H3N2	10 ⁹	70	75	56	21	15	Influenza A H3N2
		10 ⁸	49	47	34	17	11	
		10 ⁷	25	18	0	2	0	
		10 ⁶	10	0	0	0	0	
BM1456-57 BM1480-85	H1N1	10 ⁹	72	71	61	18	5	Influenza A H1N1
		10 ⁸	56	49	39	10	4	
		10 ⁷	44	30	15	0	0	
		10 ⁶	14	0	0	0	0	

^a Tenfold dilution series of each tested influenza strain was prepared and analyzed by LC–MS/MS. Indicated are the total genome copies of the analyzed influenza virus subjected to LC–MS/MS.

^b Amino acid sequence coverage (%) of proteins determined by MASCOT, as based on identified peptides. Percentages are the averages of combined results for the H3N2 or H1N1 strain. Proteins were considered significantly identified when the MASCOT score was ≥ 50 and when a minimum of three peptides each with a score of ≥ 20 was identified. NP, nucleoprotein; M1, matrix protein; NS1, non-structural protein 1; HA, hemagglutinin; NA, neuraminidase;

^c Identification of type and subtype based on the highest MASCOT scores of nucleoprotein; thus, the given subtypes should be considered as indicative.

Table 4
Repeatedly LC–MS/MS-identified influenza A virus peptides derived from nucleoprotein at titers $\geq 7 \times 10^6$ genome copies.

H1N1 strains	Observed <i>m/z</i>	H3N2 strains	Observed <i>m/z</i>
FQTAAQR	820.4	FQTAAQR	820.4
LSDYDGR	824.4	ND	–
QDATEIR	831.4	QNATEIR	830.4
AMMDQVR	849.4	AMVDQVR	817.4
KTGGPIYR	890.5	KTGGPIYR	890.5
TRVAYER	893.5	TRSAYER	881.4
IDGKWMMR	904.5	RVDGKWMMR	1046.5
TGGPIYRR	918.5	TGGPIYRR	918.5
SRYWAIR	950.3	SGYWAIR	851.4
MCNILKKG	962.5	MCNILKKG	962.5
GKFQTAQR	1005.5	GKFQTAQR	1005.5
GVFELSDEK	1022.5	GVFELSDEK	1022.5
KTGGPIYRR	1046.6	DS	891.5
MVLSAFDER	1066.5	MVLSAFDER	1066.5
YLEEHPASGK	1129.5	YLEEHPASGK	1129.5
GVGTIAMELIR	1158.7	GIGTMVMELIR	1218.6
LIQNSITIER	1185.7	LIQNSITIER	1157.7
DS	849.4; 2044.0	AMVDQVRESR	1189.6
TSDMRTEVIR	1206.6	TSDMRAEIR	1190.6
DS	1022.5	GRGVFELSDEK	1235.6
MVLSAFDERR	1222.6	MVLSAFDERR	1222.6
SYEQMETGGER	1285.5	SYEQMETDGDR	1329.5
HSNLNDATYQR	1317.6	HSNLNDATYQR	1317.6
EGYSLVGIDPFK	1323.7	EGYSLVGIDPFK	1323.7
FYIQMCTELK	1331.6	FYIQMCTELK	1331.6
NKYLEEHPASGK	1371.7	NKYLEEHPASGK	1371.7
MCSLMQGSTLPR	1379.6	MCSLMQGSTLPR	1379.6
LLQNSQVSLMR	1386.8	DS	2334.2
ELILYDKEEIR	1419.7	ELVLYDKEEIR	1405.7
ATVMAAFSGNNEGR	1423.6	STIMAAFTGNTTEGR	1454.7
YLEEHPASGKDPK	1469.7	YLEEHPASGKDPK	1469.7
MCSLMQGSTLPRR	1535.7	MCSLMQGSTLPRR	1535.7
ELILYDKEEIRR	1575.9	ELVLYDKEEIRR	1561.8
YLEEHPASGKDPKK	1597.8	YLEEHPASGKDPKK	1597.8
NPGNAEIEDLIFLAR	1670.9	NPGNAEIEDLIFLAR	1670.9
ASAGQTSVQPTFSVQR	1674.8	ASAGQTSVQPTFSVQR	1662.9
NKYLEEHPASGKDPK	1711.9	NKYLEEHPASGKDPK	1711.9
MMESAKPEDLSFQGR	1724.8	MMEGAKPEEVSFR	1509.7
MCNILKKGKFTAAQR	1764.9	MCNILKKGKFTAAQR	1764.9
SQLVWVMACHSAAFEDLR	2019.9	SQLVWVMACHSAAFEDLR	2019.9
ESRNPNAEIEDLIFLAR	2043.0	ESRNPNAEIEDLIFLAR	2043.0
SYEQMETGGERQDATEIR	2098.9	SYEQMETDGDRQNATEIR	2141.9
SCLPACVYGLAVASGHDFER	2208.0	SCLPACAYGPAVSSGYDFEK	2177.9
RSYEQMETGGERQDATEIR	2255.0	ND	–
LLQNSQVSLMRPNENPAHK	2274.2	LLQNSQVSLMRPNENPAHK	2334.2
GVQIASNENVETMDSNTLELR	2319.2	GVQIASNENMDNMGSSTLELR	2265.0
QANNGEDATAGLTHIMIWHSNLNDATYQR	3240.5	QANNGEDATAGLTHIMIWHSNLNDATYQR	3256.5
DS	1323.7; 2208.0	SCLPACAYGPAVSSGYDFEKEGYSLVGIDPFK	3483.6

Identification data were extracted from MASCOT.

ND, peptide sequence not identified.

DS, identified sequence in another peptide.

The influenza viruses ($\geq 7 \times 10^6$ genome copies) were typed and subtyped based mainly on the identification of peptides derived from nucleoprotein (shown in bold).

Table 5

Identified proteins of two different respiratory viruses from mixed samples. Mixture A contained H3N2 (BM1454) and RSV (BM1450). Mixture B contained H1N1 (BM1456) and hMPV (BM1460).

Mix	Titer ^a	Id. virus ^b	Average sequence coverage of identified proteins ^c (%)								
			NP	M1	NS1	HA	NA	M2-1	F	P	
A	10 ⁹	Influenza A/H3N2	63	74	16	15	4				
		RSV	21	43				53	9	55	
	10 ⁸	Influenza A/H3N2	48	49	20	13	7				
		RSV	23	40				46	3	53	
	10 ⁷	Influenza A/H3N2	18	34	0	5	0				
RSV		14	0				23	0	0		
B	10 ⁹	Influenza A/H1N1	73	67	60	0	0				
		hMPV ^d	14	29				10	15	16	
	10 ⁸	Influenza A/H1N1	49	33	39	0	0				
		hMPV ^d	8	18				10	9	18	
	10 ⁷	Influenza A/H1N1	24	10	11	0	0				
hMPV ^d		0	0				0	0	0		

^a Total genome copy number of the tested influenza virus and RSV in a total sample volume subjected to LC–MS/MS analysis.

^b Identified virus.

^c An average sequence coverage (%) of the identified protein was calculated from two independent experiments. A protein was considered identified when the MASCOT score was ≥ 50 and when a minimum of three peptides with a score ≥ 20 was identified in the protein.

^d The starting concentration of hMPV was 100 times lower than that for influenza and RSV.

NP, nucleoprotein; M1, matrix protein; NS1, non-structural protein 1; HA, hemagglutinin; NA, neuraminidase; M2-1, matrix protein; F, fusion glycoprotein; P, phosphoprotein.

Table 6

Identification of viral proteins from reconstituted clinical material spiked with viruses.

Virus	Titer ^a	Average sequence coverage of identified proteins ^b (%)								
		NP	M1	NS1	HA	NA	PA	M2-1	F	P
H3N2	10 ^{9c}	69	71	32	13	12	0	ND	ND	ND
	10 ⁸	25	27	0	0	0	0	ND	ND	ND
	10 ⁷	4	4	0	0	0	0	ND	ND	ND
H1N1	10 ^{9c}	68	77	71	0	7	3	ND	ND	ND
	10 ⁸	24	14	9	0	0	0	ND	ND	ND
RSV	10 ^{9c}	54	54	27.3	ND	ND	ND	35	9	31
	10 ⁸	3	35	ND	ND	ND	ND	15	0	17
hMPV	10 ^{7c}	25	48	ND	ND	ND	ND	16	28	19
	10 ⁶	0	0	ND	ND	ND	ND	0	0	6 ^d
Neg. control	Throat swab	0	0	0	0	0	0	0	0	0

^a Total genome copy number of the tested influenza virus and RSV in a total sample volume subjected to LC–MS/MS analysis.

^b An average sequence coverage (%) of the identified protein was calculated from two independent experiments. A protein was considered identified when the MASCOT score was ≥ 50 and when a minimum of two peptides with a score ≥ 20 was identified in the protein. NP, nucleoprotein; M1, matrix protein; NS1, non-structural protein 1; HA, hemagglutinin; NA, neuraminidase; M2-1, matrix protein; F, fusion glycoprotein; P, phosphoprotein.

^c Positive control; isolation of the virus was without throat swab material.

^d Only in one of the two tested samples peptides were detected.

peptides derived from nucleoprotein, matrix protein M1, and non-structural protein NS1 were identified. In addition, in RSV samples also peptides derived from matrix protein M2-1 and phosphoprotein were identified. In one of the two tested hMPV samples, two peptides derived from phosphoprotein were identified (Table 6). The identification limit at which influenza A H3N2 was detected, was 1×10^7 genome copies in the total sample volume subjected to LC–MS/MS; for influenza A H1N1 and RSV this was approximately 1×10^8 genome copies; for hMPV 1×10^7 genome copies.

4. Discussion

The accurate and rapid identification of influenza and other respiratory viruses is important in combating outbreaks and epidemics, detecting newly emerging viruses, and initiating prophylaxis and early therapy. Antigen-detection and rRT-PCR tests are used most frequently for the detection of viruses in the acute phase. PCR-based methods can yield results within 3–8 h with high sensitivity and specificity (Centers for Disease Control and Prevention: Guidance for Clinicians on the Use of rRT-PCR and Other Molecular Assays for Diagnosis of Influenza Virus Infection, 2013), whereas the rapid antigen-detection tests can even yield results

within 15 min. However, both methods feature drawbacks. (i) They are target-directed and can potentially miss non-selected or emerging pathogenic viruses that are not covered by the PCR primer sets and antibodies used in the antigen-detection assays. For instance, it was shown that a new type of human papillomavirus (HPV) was not detected by a broad HPV-primer PCR system (Johansson et al., 2013) and newly emerging mutations in the conserved region of M1 prevented rRT-PCR detection of influenza A H1N1 and H3N2 from clinical specimens (Binnicker et al., 2013; Yang et al., 2014). (ii) Multiple tests (different PCRs or PCR followed by sequencing) are needed for (sub)typing and for detection of mixed infections. (iii) The sensitivity of rapid antigen-detection tests is low. The commonly used Rapid Influenza Diagnostic Test (RIDT), although fast, lacks accuracy and sensitivity (40–70%), resulting in a high percentage of false-negative results (Beck et al., 2012). Theoretically, mass spectrometry-based systems could overcome these drawbacks and they were already investigated for the identification of respiratory viruses. However, until now the used sample preparation methods were too laborious and time-consuming for mass spectrometry to be an identification method of clinical potential. Therefore, in this study a rapid, generic and robust sample preparation method for mass spectrometry-based virus identification was

developed. Subsequently, the sensitivity of the combined procedure of this sample preparation and mass spectrometry analysis was determined.

Tenfold serial dilutions of ten cultured influenza A strains of two different subtypes, mixed samples of influenza A virus with either hMPV or RSV, and reconstituted clinical samples were treated with the in-house developed sample preparation method without prior purification by laborious procedures, apart from standard removal of cell debris. Subsequently, peptides were subjected to MALDI-TOF MS and LC-MS/MS and identified by use of a web-based MASCOT search algorithm. To confirm correct identification, the amino acid sequences were compared to the corresponding DNA sequences, obtained by sequencing.

All of the influenza A strains were successfully identified to the subtype level by both mass spectrometry systems. Identification of viruses was achieved within 3 h with MALDI-TOF MS and within 6 h with LC-MS/MS, excluding the time necessary for culturing the viruses. Sensitivity of MALDI-TOF MS was lower (detection limit of 1×10^9 ; Table 1) compared to LC-MS/MS (detection limit of 7×10^6 ; Table 3); the latter corresponding to a CT value of 30–32.

Earlier studies have already shown that it is possible to identify highly pure and/or concentrated influenza A viruses with MALDI-TOF MS, MALDI-FT-ICR MS or LC-MS/MS (Schwahn et al., 2009a,b, 2010a,b; Chou et al., 2011; Jang et al., 2011; Nguyen and Downard, 2013; Fernandes and Downard, 2014; Li et al., 2014). However, the sample preparation methods used were time-consuming (up to 24 h) and laborious, and are therefore not suitable for rapid applications. In these studies, the same nucleoprotein-derived peptides were found among the identified viral peptides as detected here (Schwahn et al., 2009a,b; Li et al., 2014), which indicates that with the in-house developed sample preparation method, comparable results could be achieved in a much shorter time. Furthermore, in the developed sample preparation method, prior purification or extended concentration of cultured viruses, for example by applying ultracentrifugation, is not required for the identification of viral proteins in the presence of other viral or non-viral proteins. The sample preparation time was significantly reduced to approximately 1.5 h by the introduction of REDS for rapid reduction, alkylation, precipitation, and trypsin digestion. The tenfold serial dilutions of cultured viruses were prepared in CyMol, a collection, transport and preservation medium known for its compatibility with multiple diagnostic platforms including cytology and molecular diagnostics (Luinstra et al., 2011). Suspending viruses in CyMol significantly (approximately 100-fold) improved the identification limit compared to the other tested solutions, such as ammonium bicarbonate, NTE buffer (100 mM NaCl, 10 mM Tris-HCl [pH 7.5], 1 mM EDTA), PBS (phosphate buffered saline), and a universal transport medium (UTM-RT; data not shown).

Besides monocultures of influenza A strains, mixed samples of influenza A virus with either hMPV or RSV were subjected to MALDI-TOF MS and LC-MS/MS. Viruses were identified simultaneously and specifically by LC-MS/MS (Table 5), with a detection limit of 5.3×10^7 for influenza A (H1N1 and H3N2), 3.6×10^7 for RSV, and 1.1×10^7 genome copies for hMPV. Identification was not possible by MALDI-TOF MS due to its low sensitivity. In addition to monocultures and mixed samples, throat swabs spiked with influenza A, hMPV or RSV were subjected to LC-MS/MS. All the spiked viruses were identified, with detection limits of 10^7 , 10^8 , and 10^7 genome copies for influenza A H3N2, influenza A H1N1 and RSV, and hMPV, respectively. These results demonstrate that LC-MS/MS is not only able to identify monocultures of viruses, but can also discriminate between two viruses in the same sample and can identify a virus from a sample containing human proteins originating from a throat swab. Sensitivity of LC-MS/MS appeared to be lower when mixed samples or reconstituted samples were tested.

Viral loads in clinical samples vary depending on the type of respiratory material, subtype of the virus, severity and phase of infection, and host factors, such as immunity. In general, viral loads fall within the range of 10^4 – 10^6 genome copies/ml (Ngaosuwanuk et al., 2010; Lee et al., 2011). In addition, in a recent pan-European survey (Nuttall et al., 2011), the average CT values for influenza virus, RSV, and hMPV detected in patients were 29, 29, and 30, respectively (Coenjaerts, F.E.J., personal communication). Therefore, the ultimate identification limit of a mass spectrometry method should reach 10^4 genome copies/ml for direct clinical application. The sensitivity of the developed sample preparation method in combination with LC-MS/MS is 100-fold lower and therefore not good enough for direct application in virological diagnostics. For detection of viruses from clinical samples, viral culture is needed before samples can be prepared and analyzed. Because culturing is required, viable virus has to be recovered and successfully grown. For application in virological diagnostics, sensitivity should be increased by developing techniques that enable fast and efficient virus concentration and purification from respiratory samples. Furthermore, mass spectrometry systems should be improved to increase sensitivity. In the meanwhile, keeping the drawbacks into mind this in-house developed sample preparation method in combination with LC-MS/MS could be experimentally used for patients with suspicion of a respiratory infection where routine diagnostics are negative.

The identification of the influenza A strains, RSV and hMPV was based on detection of peptides derived from different proteins. All these identified peptides were found to be encoded in the corresponding genome sequences. MALDI-TOF MS and LC-MS/MS both detected nucleoprotein-derived peptides. In addition, LC-MS/MS-identified peptides derived from other viral proteins, namely M1, NS1, HA, and NA proteins of influenza viruses and M, M2-1, P, and F proteins of RSV and hMPV. The identification of peptides from different proteins not only accomplishes identification but also potentially delivers information about virulence, resistance to antiviral medications, and/or transmission efficiency. For instance, an amino acid substitution in nucleoprotein (M136L) was shown to enhance human-to-human transmission (Zhou et al., 1999). Interestingly, one of the identified nucleoprotein-derived peptides, *i.e.*, QANNGEDATAGLTHIMIWHNSLNLDATYQR, contained this substitution. This peptide was found in all of the tested influenza strains (Table 4), suggesting that these strains might exhibit increased human-to-human transmission efficiency.

In conclusion, mass spectrometry systems have been investigated for the identification of highly pure and/or concentrated influenza A viruses. However, used sample preparation methods were laborious and time-consuming. The developed sample preparation method in combination with LC-MS/MS allowed rapid and reliable identification of cultured respiratory viruses and viruses spiked in throat swabs. To introduce this LC-MS/MS based identification method in virological diagnostics, sensitivity must be improved up to 100-fold and clinical samples have to be tested. The combination of an improved mass spectrometry technique and/or clinical sample preparation method allowing direct identification of viruses from clinical samples will be an important new diagnostic tool in clinical virology.

Conflict of interest

The authors declare that they have no competing interests.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.jviromet.2014.11.014>.

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