

# Evaluation of a Multiplex One-Step TaqMan Real-Time Reverse Transcription-PCR Assays for the Detection of H5N1 Avian Influenza Viruses in Clinical Specimens

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## Introduction

The highly pathogenic influenza A virus subtype H5N1 is an emerging avian influenza virus that continues to pose a significant threat to human health. Highly pathogenic avian influenza (HPAI) is very contagious among wild and domestic birds, and can be deadly to them, especially domestic poultry [1-3]. Outbreaks of avian epizootic in the world after first appearing in 2003 have shown that HPAI A (H5N1) could cross the species barrier to infect humans with high mortalities. Since 2003, 650 human infections with HPAI have been reported to the World Health Organization (WHO) by 15 countries. About 60% of these people died [4,5].

Since October 2005, H5N1 subtype influenza A virus has been circulating in the industrial and domestic poultry population in Crimea Autonomous Republic and other districts of Ukraine. During this outbreak of H5N1 avian epizootic, virus killed or led to the destruction of 238 000 poultry. Thus, HPAI A (H5N1) represents a serious problem to public health. Sensitive and robust surveillance measures are required to detect any evidence that the virus has acquired the ability to transmit between humans and emerge as the next pandemic strain.

Diagnosis of avian influenza can be made by complex of methods, including clinical signs, serologic methods and direct virus detection methods. Currently, methods for detection of H5N1 virus include virus isolation, serologic tests and molecular detection of viral nucleic acids [6]. Virus isolation method is time consuming, technically difficult to perform, has a high false-negative rate and therefore it isn't suitable for routine diagnostics [7]. Commercially available antigen detection tests are based on either immunofluorescence or enzyme immunoassay (EIA) methods and detect all subtypes of influenza A virus. They have low sensitivity for the detection of avian influenza H5N1 in specimens of humans and thereby have limited utility for detection of HPAI A(H5N1) [4]. At the same time serological tests for the detection of virus specific antibodies (hemagglutination inhibition test, EIA, and virus neutralization test) are available. However due to slow seroconversion and technical limitations these assays are impractical for the routine diagnostic testing of clinical samples and are more useful for research studies [4-7].

Molecular diagnostics are based on amplifying nucleic acid to high levels to allow easy identification of the sample. There are several different types of molecular diagnostic tests. Reverse transcription-polymerase chain reaction (RT-PCR) and real-time polymerase chain reaction (RRT-PCR) are the most commonly used tests for rapid detection of influenza viral RNA in clinical and laboratory specimens

[8]. The analytical sensitivity of the RRT-PCR and the one-step RT-PCR, were 10,000-fold and 100-fold greater than the commercial Antigen Capture EIA in detection of H5N1, respectively [9].

The objective of this study was to develop and evaluate rapid TaqMan RRT-PCR for the detection and subtyping of H5N1-specific influenza A viruses. Samples collected during the 2005 H5N1 birds epizootic, H1N12009 human pandemic and experimental H5N1 human infection were used to assess the relative sensitivity and specificity of the designed TaqMan RRT-PCR.

## Materials and methods

### Virus strains and avian specimen

Avian influenza A (H5N1) virus isolate (LA-NK-21205) was isolated in Crimea AR in 2005 during epizootic outbreaks among the industrial and domestic poultry. It was kindly provided for our study by National Centre of microbial strains (Kyiv, Ukraine).

Human influenza virus reference strains A/FM1/47 (H1N1), A/Panama/2007/99 (H3N2), A/New Caledonia/20/99 (H1N1), B/Hong Kong/330/01 were kindly provided for our study by Svitlana L. Rybalko and WHO Collaborating Center for Influenza (CDC, USA).

Cloaca swab specimens of birds from Odessa zoo park (eagle, pelican, hawk and peacock) and throat swab samples from died horses were collected during the respiratory disease outbreak in 2005. These samples were kindly provided for our study by the Institute of veterinary medicine UAAN (Kyiv, Ukraine).

Human respiratory samples (nasopharyngeal swabs and aspirates, sputum) from viral influenza (VI) patients (n=10) were kindly provided by Irina G. Kostenko (the Main Military Clinical Hospital of Ukraine). The clinical material was validated by Seeplex® Influenza A(H1N1pandemic) RT-PCR assay (Seegene Inc., South Korea) and TaqMan Influenza A (H1N1) Assay Sets [10].

In addition, five respiratory samples (nasopharyngeal swabs and sputum) collected from uninfected patients were contaminated by LA-NK-21205 Crimean virus isolate to simulate respiratory clinical H5 samples due to the limited availability of H5 clinical samples from human cases in Ukraine.

### Viral RNA extraction

Viral RNA was extracted from 140 µl of avian specimens or standard virus samples using NucleoSpin RNA Virus Kit (Macherey-Nagel

GmbH, Germany) based on spin column extraction technology (according to the manufacturer's instructions). It was used 5 µl of control RNA (MS2-phage) as Internal Control (IC) during the RNA purification stage. All procedures were carried out under the containment conditions of a biosafety level 3 with additional safety precautions.

### Nucleotide sequence analysis

Reaction mixture contained 13 µl of purified DNA, 1 µl of primer (20 pmol), 4 µl RR mix of BigDye Terminator v3.1 Cycle Sequencing Kit (ABI, Foster City, CA), and 2 µl of 5x sequencing buffer, according to the manufacturer's instructions. The PCR was performed as follows: denaturation for 1 min at 94°C, followed by 25 cycles of PCR amplification, with each cycle consisting of 10 s of denaturation at 96°C, 5 s of annealing at 50°C, and 4 min of elongation at 60°C. The PCR products were purified with a Sephadex G-50 column and subjected to capillary electrophoresis at 3100-Avant genetic Analyzer (ABI, Foster City, CA). The sequence data were analyzed by Nucleotide BLAST Software [11], using the NCBI database [12].

### Primers and probe design

Nucleotide sequences (N=30) of the matrix (M1), hemagglutinin (HA) and neuraminidase (NA) genes were taken from the Influenza Virus Resource of the NCBI database. The M1, H5, N1 genes A(H5N1) positive during of 2004-2012 period subjected to multiple alignments using the BioEdit Sequence Alignment Editor Software version-7.0- (<http://www.mbio.ncsu.edu/BioEdit/bioedit.html>) [13]. The conserved regions were used for design of specific primers and TaqMan fluorescent probes with Primer Express 2.0 Program (Applied Biosystems, Cheshire, United Kingdom). The TaqMan-probes were labeled with the fluorescent reporters FAM, HEX/JOE at the 5'-end and non-fluorescent quencher BHQ1, BHQ2 at the 3'-end. Primers and TaqMan fluorescent probes were synthesized by Invitrogen (Invitrogen, Germany) and Sintol, Evrogen (). The amplified fragments have length of 193 bp, 190 bp and 130 bp.

### Internal quality control (IC-control)

MS2 bacteriophage was used to control nucleic acids extraction, as described by Dreier et al. [14]. Specific primers and TaqMan fluorescent probe were designed for simultaneous detection of MS2 bacteriophage genomic RNA and M-gene RNA of influenza HPAI A (H5N1) viruses. TaqMan-probe was labeled with the fluorescent reporter HEX at the 5'- end and non-fluorescent quencher BHQ1 at the 3'-end. The amplified fragment has length of 57 bp. We used method of limited titration of internal control (MS2-phage) dilution

to evaluate the effectiveness of virus RNA isolation from different clinical materials.

### Engineering of positive control plasmids for HPAI A(H5N1) detection

Synthetic positive controls, which represent fragments of M, H5 and/or N1 genes were generated by PCR and cloned into pGEM3Zf(+) vector (Promega). All plasmids were subjected to nucleotide sequencing to ensure the correct target sequences by sequencing with BigDye Terminator v3.1 Cycle Sequencing Kit (ABI, Foster City, CA). Identity of cloned fragments and corresponding virus sequences of target genes were confirmed by multiple alignment using the BioEdit Sequence Alignment Editor Software. Then sequences of engineering fragments M1, H5, N1 genes and corresponding genes A(H5N1) positive during (N>100) were analyzed by Nucleotide BLAST Software. The recombinant plasmids were used for amplification as external standards. Standard curve was created with 10-fold serial dilutions of transcribed RNA that contains from 10<sup>7</sup> to 10<sup>1</sup> copies of target sequences per reaction.

### PCR optimization

Real-Time RT-PCR was carried out in a 25 µl mixture containing 5 µl RNA, 12.5 µl 2X TaqMan One Step RT PCR Master Mix (Applied Biosystems, USA), 0.75 µl 40X MultiScribe and RNase inhibitor mixture, 0.25 µl forward primer, 0.25 µl reverse primer and 0.125 µl probe using a fluorometric PCR thermocycler (ABI PRIZM 7000).

The fixed amount of template in reaction was used to optimize the concentration of primers and probes titration method. The annealing temperature was estimated by the amplification of positive samples.

The following criteria were used for determination the optimal amplification conditions: minimal value of threshold cycle (Ct) at maximal value of fluorescence above the background signal (ΔRn). Optimization of temperature regime was carried out to adopt test kit to ABI PRIZM 7000 (Applied Biosystems, USA) and RotorGene 3000/6000 (Corbett Research, Australia).

## Results

### Nucleotide sequence analysis

Nucleotide sequences (N=30) of MA, HA, NA genes isolated from HPAI A(H5N1) positive samples during 2004-2006 period downloaded from NCBI database were aligned ( Table 1).

No.	Virus nomenclature	Host	Year
1	A/human/Vietnam/CL01/2004(H5N1)	human	2004
2	A/human/Vietnam/CL02/2004(H5N1)	human	2004
3	A/human/Vietnam/CL17/2004(H5N1)	human	2004
4	A/human/Vietnam/CL20/2004(H5N1)	human	2004
5	A/human/Vietnam/CL26/2004(H5N1)	human	2004
6	A/human/Vietnam/CL36/2004(H5N1)	human	2004

7	A/human/Vietnam/CL115/2005(H5N1)	human	2005
8	A/human/Vietnam/CL2009/2005(H5N1)	human	2005
9	A/human/Vietnam/CL105/2005(H5N1)	human	2005
10	A/Ck/Viet Nam/C57/2004(H5N1)	birds	2004
11	A/Ck/Viet Nam/36/2004(H5N1)	birds	2004
12	A/Ck/Viet Nam/37/2004(H5N1)	birds	2004
13	A/Ck/Viet Nam/38/2004(H5N1)	birds	2004
14	A/Ck/Viet Nam/39/2004(H5N1)	birds	2004
15	A/duck/Kurgan/08/2005(H5N1)	birds	2005
16	A/goose/Krasnoozerka/627/2005(H5N1)	birds	2005
17	A/chicken/Tambov/570-2/05(H5N1)	birds	2005
18	A/wild duck/Omsk/103-01/05(H5N1)	birds	2005
19	A/goose/Novosibirsk/4/2005(H5N1)	birds	2005
20	A/swan/Astrakhan/Russia/Nov-2/2005(H5N1)	birds	2005
21	A/chicken/Crimea/08/2005(H5N1)	birds	2005
22	A/chicken/Crimea/04/2005(H5N1)	birds	2005
23	A/goose/Crimea/615/05(H5N1)	birds	2005
24	A/Egypt/2782-NAMRU3/2006(H5N1)	birds	2006
25	A/chicken/Nigeria/641/2006(H5N1)	birds	2006
26	A/chicken/Sudan/2115-9/2006(H5N1)	birds	2006
27	A/chicken/Krasnodar/199/06(H5N1)	birds	2006
28	A/common bussard/Bavaria/2/2006(H5N1)	birds	2006
29	A/chicken/Volgograd/236/06(H5N1)	birds	2006
30	A/domestic goose/Iraq/812/2006(H5N1)	birds	2006

**Table 1:** Viruses selected from NCBI database for alignment.

We compared human and avian H5N1 HPAI viruses' nucleotide sequences, which were isolated in different countries: Vietnam, Russia, Iraq, Nigeria, Sudan and three Crimean virus isolates from Ukraine. Conservative regions (99, 8-100% homology) of 130-200 nt in length were selected. These regions were used to design specific primers and probes for detection of HPAI A(H5N1) by multiplex Real-Time RT-PCR protocol. BLAST analysis of nucleotide sequences of MA, HA and NA genes of HPAI A(H5N1) strains isolated during 2007-2013 within selected conservative regions revealed 97-99% of nucleotide homology.

### Multiplex Real-Time RT-PCR protocol

In this study we developed a two-stage multiplex RRT-PCR based on the one-step MA, H5 and N1 genes detection. The first stage should distinguish influenza A and B. Then, positive samples were tested by RRT-PCR in one-tube targeting H5- and N1-specific sequences.

### Design of primers and probes for TaqMan Real-Time RT-PCR

Using Primer Express V.2 (Applied Biosystems) we designed primers and TaqMan-probes for detection of selected conservative regions of MA, HA and NA genes of HPAI A(H5N1) and for internal control (MS2-fage RNA) detection. Specific primers and probes targeting the M1 gene of influenza A were selected for typing of influenza A virus from influenza B and C and other respiratory viruses. The specific primers and probes targeting the H5 and N1 genes were selected for subtyping of HPAI A(H5N1) from influenza A(H1,H3,H7) viruses.

BLAST (Basic Local Alignment Search Tool) analysis of designed primers and probes for M, H5 and N1 genes conferred that they have high homology (>98%) with the corresponding nucleotide sequences of more than 120 genomes of HPAI A(H5N1) strains isolated from birds in Asia, Europe, Russia in 2004 and in Ukraine and Crimea AR in 2005.

### Optimization of the multiplex assay

Primers titration from 100 to 700 nM as well as probes titration from 100 to 300 nM indicated an optimal primers concentration of 500 nM and an optimal probes concentration of 250 nM for all four assays in the multiplex real-time RT-PCR. Application of higher or lower concentrations hadn't influence on the sensitivity of the multiplex assay significantly (results aren't shown). The optimized multiplex real-time RRT-PCR assay has a 25 µl PCR reaction volume.

The lowest Ct value and highest Δ Rn were observed in the amplification with an annealing temperature of 55°C (at the 1-st step of assay) and of 54°C (at the 2-nd step of assay). Optimization of reaction protocol have been done for Applied Biosystems(ABI PRIZM 7000/7500) and for Corbett Research(Rotor Gene 3000/6000) instruments.

### Specificity of TaqMan multiplex RRT-PCR assay

Specificity of multiplex assay was recently evaluated by cross-reactivity with different genotype influenza viruses (H3N8, H7N7) and other avian diseases viruses (Rhinotracheitis virus, Newcastle Disease Virus, Variola avium virus, Chickens Egg Drop Syndrome virus) [15]. In this study we investigated specificity of the Multiplex H5N1 TaqMan RRT-PCR assay by analyzing avian clinical samples from Odessa zoo park, reference CDC human influenza samples and clinical human samples (H1N1) and simulated clinical human samples (H5N1).

As it shown in Table 2, positive results at 1<sup>st</sup> and 2<sup>nd</sup> stage were obtained only for specimen of peacock died of avian influenza in Odessa zoo park and for cultural sample of Crimean isolate LA-NK-21205 of avian influenza virus A (H5N1), which were used as a positive control.

№	Clinical materials	Obtained results by Multiplex Real-Time RT-PCR protocol			
		1 <sup>st</sup> stage		2 <sup>nd</sup> stage	
		IC (internal control) (Ct)	Influenza viruses A (M-gene) (Ct)	Hemagglutinine HA5 (Ct)	Neuraminidase NA1 (Ct)
1	Eagle cloak swab	+ (24,29)	-	-	-
2	Hawk cloak swab	+ (24,9)	-	-	-
3	Pelican cloak swab	+ (25,69)	-	-	-
4	Sample of the 1 <sup>st</sup> passage in the allantoic cavities of embryonated hen's eggs (isolated from died horse)	+ (23,6)	-	-	-
5	Cultural avian influenza virus A (H5N1) sample (isolated from died peacock in Odessa zoo park)	+ (20,37)	+ (16,62)	+ (19,38)	+ (19,51)
6	Cultural sample of Crimean isolate LA-NK-21205 of avian influenza virus A (H5N1)	+ (25,39)	+ (13,43)	+ (20,11)	+ (19,3)
7	Samples (n=10) of the non-infected allantoic cavities of embryonated hen's eggs	+ (28,69)	-	-	-

Remarks: + positive result or detecting sample fluorescence signal; negative result or no detecting sample fluorescence signal; (Ct) – threshold cycle value.

**Table 2:** Testing of specimens from Odessa zoo park by Multiplex H5N1 TaqMan RRT-PCR protocol.

The concentration of cultural sample of Crimean isolate LA-NK-21205 was 105 EID50. So, the concentration of cultural avian influenza virus A (H5N1) sample isolated from died peacock in Odessa Zoo Park was the same. Fluorescence increasing was not detectable in the samples of ten non-infected allantoic fluides of embryonated hen's eggs and clinical avian specimens from Odessa zoo

park (eagle, hawk, pelican cloak swabs and sample, isolated from died horse).

No cross-detection was observed with CDC human influenza B viruses' sample and influenza viruses A(H1N1), A(H3N2) by targeting the H5-gene. The data are represented in Table 3.

№	CDC influenza samples	Obtained results by Multiplex Real-Time RT-PCR protocol			
		1 <sup>st</sup> stage		2 <sup>nd</sup> stage	
		IC (internal control) (Ct)	Influenza viruses A (M-gene) (Ct)	Hemagglutinine HA5 (Ct)	Neuraminidase NA1 (Ct)
1	Influenza B virus: B/HongKong/330/01	+ (24,53)	-	-	-

2	Influenza virus A(H1N1) : A/FM1/47	+ (23,9)	+ (20,6)	-	+ (19,6)
3	Influenza virus A (H1N1): A/New Caledonia/20/99	+ (25,19)	+ (24,56)	-	+ (21,3)
4	Influenza virus A(H3N2): A/Panama/2007/99	+ (23,36)	+ (25,69)	-	-

Remark: positive result or detecting sample fluorescence signal; negative result or no detecting sample fluorescence signal; (Ct) – threshold cycle value.

**Table 3:** Testing of CDC influenza samples by Multiplex H5N1 TaqMan RRT-PCR protocol.

Although the pandemic outbreak of influenza A/H1N1/2009 declared finished, this virus continues to circulate and remains a challenge for patient management. Respiratory human clinical samples (n=10) collected during an H1N1 pandemic influenza in in 2009 were also analyzed. All these samples had previously been demonstrated to contain influenza A H1N1pandem 2009 viral RNA (tested by Seplex®

Influenza A (H1N1pandemic), manufactured by Seegene Inc, South Korea) and then were verified by H1N1pandemic2009 primers and probes, recommended by WHO protocol TaqMan Influenza A (H1N1) Assay Sets [16]. Following designed multiplex H5N1 real-time RT-PCR analysis, no cross-detection were observed with all tested human respiratory samples (influenza A(H1N1) pandemic2009).

Recombinant (target) plasmids	No. of positive samples/no. of samples tested/25 µl reaction mixture									Analytical sensitivity/per 25 µl reaction mixture
	10 <sup>7</sup>	10 <sup>6</sup>	10 <sup>5</sup>	10 <sup>4</sup>	10 <sup>3</sup>	10 <sup>2</sup>	10	1		
Ten-fold dilutions										
pIHC-11(H5-gene)	3/3	3/3	3/3	3/3	3/3	3/3	3/3	3/3	1/3	10
pINC-8(N1-gene)	3/3	3/3	3/3	3/3	3/3	3/3	3/3	3/3	1/3	10
pIMC-3(MA-gene)	3/3	3/3	3/3	3/3	3/3	3/3	3/3	3/3	2/3	10

**Table 4:** Determination of detection limits and the amplification efficiency of Multiplex H5N1 Real-Time RT-PCR assay.

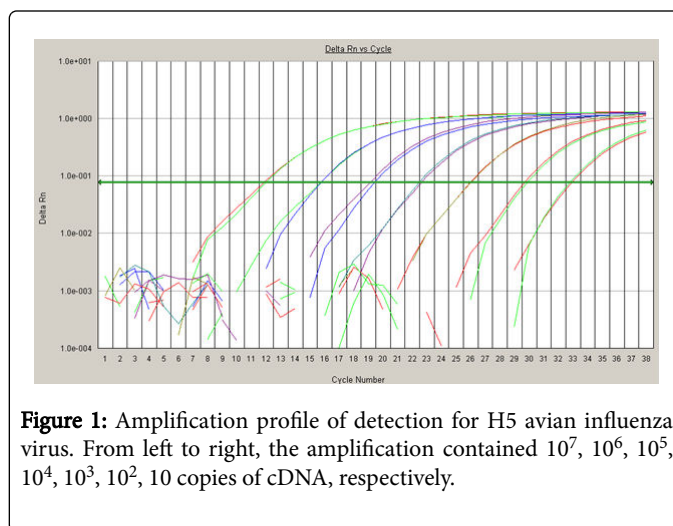
Due to the limited availability of H5 clinical samples from human cases in Ukraine, we contaminated five human respiratory samples (nasopharyngeal swabs and sputum) from not infected human by LA-NK-21205 Crimean virus isolate. All of five simulated HPAI (H5N1) specimens, which tested by the H5N1 TaqMan RRT-PCR, were shown positive results, contained influenza A H5 viral RNA.

**Analytical sensitivity (or detection limit) measuring of the assay**

Ten-fold dilutions of AIV H5N1 recombinant plasmids pIMC-3(M-gene), pIHC-11(H5-gene) и pINC-8(N1-gene) were used for the determination of detection limits and the amplification efficiency of the assay (Table 4). Samples were tested in triplicate for each dilutio

For determination of the sensitivity of RRT-PCR and creation the standard curve for quantitative analysis and estimation of the linear range, the 10-fold serial dilutions (1-10<sup>7</sup> copies) of plasmids were used as standards (Figures 1 and 2).

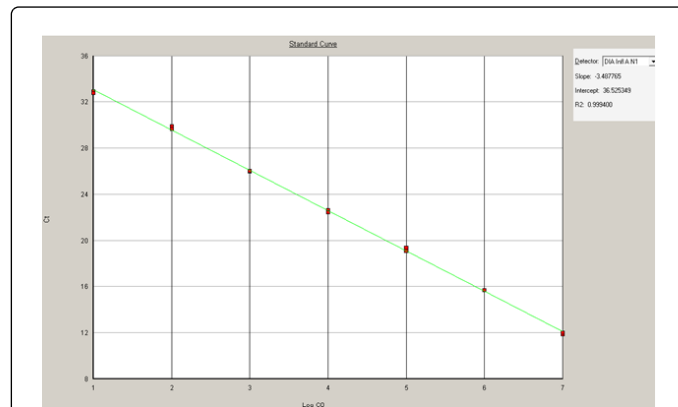
The limit of detection was determined as 10 copies in reaction with the Ct value 33 (for H5). The linear range for detection was detected as 10 to 10<sup>7</sup> copies in reaction. By using the Ct slope method [17,18] and slope values of -3.48 with pIHC-11(H5-gene), -3.45 with pINC-8(N1-gene), and -3.32 for pIMC-3(MA-gene), the amplification efficiency values of the RRT-PCRs were calculated (98, 99, and 100%, respectively).



**Discussion**

A number of scientific reports about the use of rapid detection tests for type A influenza virus in poultry have been published [19-25] pointing out the importance of having such tests available. Information about the application of these assays using human samples or samples collected during an epizootic from naturally infected birds isn't published.

In Ukraine H5N1 infections aren't endemic, so diagnostic tests for H5N1 avian influenza determination could be performed for detection of influenza virus among tourists returning from H5N1-affected regions and among potentially exposed poultry workers. That's why, the actual question is the necessity of robust effective HPAI A (H5N1) diagnostics in human and birds populations.



**Figure 2:** Reactions with copy numbers of positive plasmid (H5) from  $10^1$  to  $10^7$  were used for creating the standard curve. The standard curve was plotted by the log concentration of copy numbers against Ct values ( $Y = -3.487X + 36.52$ ,  $R^2 = 0.999$ ).

The H5N1 RRT-PCR was designed as a two-stage, TaqMan-based multiplex real-time assay to improve efficient processing and reduce the risk of carryover contamination. It proposes to distinguish influenza virus A from influenza virus B and other pathogens at the 1<sup>st</sup> stage of assay, and to subtype sequence by H5 and N1 assay in one tube at the 2<sup>nd</sup> stage. Less than 4 h is enough to obtain complete results for the clinical samples.

Primers and TaqMan-probes that fit the criteria for suitable real-time PCR primers were selected from highly conserved regions of the H5, N1 and M- gene of H5N1 HPA influenza viruses. Our investigation of MA, HA and NA genes of HPAI A(H5N1) strains isolated in period of 2007-2012 with selected conservative regions showed 97-99% of nucleotide homology. Therefore, mutations appeared after 2006 year hadn't affected the scope of the annealed primers and probes. We suggest, that designed diagnostic primers and probes are useful for identification of current A (H5N1) strains.

Despite the selection of conserved regions, in order to ensure the amplification of target viral RNA from the HA gene of influenza viruses, two degenerate bases were incorporated into the forward H5 primer and one was incorporated into the H5-probe sequence.

The H5N1 assay was designed for use on a TaqMan sequence detection platform (Applied Biosystems); however, the test will be useful in a range of laboratories since the assay is transferable to other real-time platforms, such as Corbett Rotor-Gene.

For current WHO recommendations [10] all diagnostics test protocols should be validated to ensure adequate specificity and sensitivity. In order to make sure that designed H5N1 Primers and TaqMan-probes are highly sensitive for the determination of H5N1 infection in different clinical sample types we analyzed human and birds clinical samples. The influenza A H5N1 virus Crimean isolate LA-NK-21205 was selected as the positive control material for the RRT-PCR assay in birds, since this strain is representative of H5N1

HPA strains, circulated in poultry in Crimea AR (2005). It was shown that the Multiplex H5N1 RRT-PCR is specific for the detection of influenza A H5N1 HPA viruses, amplifying viral RNA from cloac swab, allantoic fluids of embryonated hen's eggs, cultural samples.

It wasn't possible to establish the sensitivity of the Multiplex H5N1 RRT-PCR for human clinical sample types, such as secretions from the upper human respiratory tract (sputum, nasopharyngeal swabs and aspirates) because of the limited availability of H5N1 clinical samples from human cases in Ukraine. In this study we tested five simulated respiratory clinical H5N1 human samples. The study demonstrated that the H5N1 RRT-PCR assay is applicable for human samples testing.

Non-reactivity of the H5N1 RRT-PCR assay with other influenza A virus subtypes (seasonal H1N1: A/FM1/47, A/New Caledonia/20/99, H3N2: A/Panama/2007/99), influenza B virus subtype and pandemic 2009 H1N1 was demonstrated on four reference and ten human clinical samples respectively. The results confirmed the high specificity of the assay.

We analyzed the effectiveness of viral RNA extraction from different clinical materials by using limitation titration method of internal control (MS2-phage) dilutions (data not shown), which amplified together with M-gene of influenza viruses RNA. Our study demonstrated that viral RNA extraction technology by spin column is suitable for analyzing human samples and birds specimens. Thus, the addition of an internal control (IC) could further improve the assay as a diagnostic tool, as each sample could be tested for the quality of the nucleic acid extraction.

The limit of detection of H5N1 RRT-PCR test was determined as 10 copies in reaction and the linear range for detection was detected as  $10^1$  to  $10^7$  copies in reaction by titration of AIV H5N1 recombinant plasmids methods in this study. This is confirmed by our recent research [26] by titration of cultural sample of Crimean isolate LA-NK-21205, when analytical sensitivity was detected as  $10^2$  to  $10^3$  copies in clinical sample.

## Conclusion

The Multiplex H5N1 real-time RT-PCR is reproducible and cost-effective assay because it is rapid, specific, sensitive. Therefore, it is applicable for diagnostics and monitoring of influenza virus infection in patients with respiratory symptoms. Also, the designed test-kit is an effective tool for the epizootic monitoring of avian H5N1 HPA influenza viruses in birds.

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