

## Subtyping Animal Influenza Virus with General Multiplex RT-PCR and Liquichip High Throughput (GMPLex)\*

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**Abstract:** This study developed a multiplex RT-PCR integrated with luminex technology to rapidly subtype simultaneously multiple influenza viruses. Primers and probes were designed to amplify NS and M genes of influenza A viruses HA gene of H1, H3, H5, H7, H9 subtypes, and NA gene of the N1 and N2 subtypes. Universal super primers were introduced to establish a multiplex RT-PCR (GM RT-PCR). It included three stages of RT-PCR amplification, and then the RT-PCR products were further tested by LiquiChip probe, combined to give an influenza virus (IV) rapid high throughput subtyping test, designated as GMPLex. The IV GMPLex rapid high throughput subtyping test presents the following features: high throughput, able to determine the subtypes of 9 target genes in H1, H3, H5, H7, H9, N1, and N2 subtypes of the influenza A virus at one time; rapid, completing the influenza subtyping within 6 hours; high specificity, ensured the specificity of the different subtypes by using two nested degenerate primers and one probe, no cross reaction occurring between the subtypes, no non-specific reactions with other pathogens and high sensitivity. When used separately to detect the product of single GM RT-PCR for single H5 or N1 gene, the GMPLex test showed a sensitivity of  $10^{-5}$  (= 280ELD<sub>50</sub>) for both tests and the Luminex qualitative ratio results were 3.08 and 3.12, respectively. When used to detect the product of GM RT-PCR for H5N1 strain at the same time, both showed a sensitivity of  $10^{-4}$  (=2800 ELD<sub>50</sub>). The GMPLex rapid high throughput subtyping test can satisfy the needs of influenza rapid testing.

**Key words:** Influenza Virus; General multiplex RT-PCR; luminex assay; Subtyping; HA and NA genes

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Influenza A Virus (IV) belongs to the family *Orthomyxoviridae*<sup>[9]</sup>. Other members of the family include influenza viruses type B and C, which infect only humans. The virus has an envelope with a host-derived lipid bilayer and is covered with about 500 projecting glycoprotein spikes with hemagglutinin (HA) and neuraminidase (NA) activities. There are 16 subtypes of HA and 9 subtypes of NA.

Avian influenza is a significant threat to the poultry industry. Highly pathogenic AIVs (HPAI) belonging to H5 and H7 subtypes have caused respiratory disease with nearly 100% mortality in poultry<sup>[16]</sup>. Accidental genetic reassortments of RNA segments from different subtypes of avian, swine and human IVs are capable of inducing a global pandemic<sup>[16]</sup>. An HPAI H5N1 was reported to be transmitted from birds to pigs and humans leading to significant mortality<sup>[3]</sup>. The virus was spread by wild free flying birds which increase the threat of a pandemic.

The prerequisite for controlling the disease is rapid and accurate identification of this virus and subtyping. The method recommended for definitive antigenic subtyping of influenza A viruses by the World Health Organization (WHO) Expert Committee involves the use of highly specific antiserum, prepared in an animal giving minimum nonspecific reactions, directed against the H and N subtypes. The agar gel immunodiffusion (AGID) test and enzyme-linked immunosorbent assays (ELISAs) remain the diagnostic assay most utilized for detection of AIV antibodies in commercial poultry worldwide and are considered the “standard” by the World Organization for Animal Health<sup>[11]</sup>. AGID test detects antibodies to two influenza virus proteins, NP and M1, which are highly conserved and type specific. There is a

sensitive and specific ELISA that demonstrates nucleoprotein of type A influenza virus using a monoclonal antibody against type A influenza nucleoprotein. Although these assays detect antibodies against all AIV subtypes, it does not distinguish subtypes amongst them. Reverse transcriptase polymerase chain reaction (RT-PCR) and real time RT-PCR (rRT-PCR) have been developed as rapid detection tests for AIVs<sup>[2, 7, 8, 13]</sup>. However, the single RT-PCR only recognizes one specific subtype of AIV gene. Multiplex RT-PCR could only detect limited subtypes of AIV<sup>[12]</sup>. Studies have demonstrated that using multiplex assay for typing and subtyping of IVs<sup>[9]</sup> However, the numbers of target genes are limited and no more than five. To distinguish the most common subtypes in animals, a GM RT-PCR integrated with luminex technology (GMPLex) was developed for simultaneously differentiating of 7 subtypes of avian influenza A viruses.

## MATERIALS AND METHODS

### Virus strain and major reagents

Inactivated Avian Influenza A virus/Chicken/HK/HI/1997(H5N1), A/Chicken/ HK /921/1997(H9N2), A/PFV/Restock/1/1934(H7N1)A/PFV/Restock/1/1934 (H7N1), A/Swine/SZ/111/2005(H1N1), A/Swine/SZ /321/2005(H3N2) were obtained from Harbin Veterinary Research Institute, Chinese Academy of Agricultural Sciences. Qiagen one-step RNA PCR kit, DNA fragment purification kit, QIAamp viral RNA Mini-prep kit, Qiagen one-step RT-PCR kit and EZ1 virus mini kit 2.0 were used. Luminex carboxy bead from Luminex (Valencia, CA), Luminex calibration bead mix and Luminex control bead kit from Qiagen (Valencia, CA).

**Design of the probe and primer**

To design a set of primers and probes specific to each single HA and NA subtype, we compared nucleotide sequences of all subtypes of HA genes retrieved from the Genbank. Sequences from regions that were conserved in only a single subtype were chosen for the primer design. A total of 9 sets of primers and probes were designed by this strategy for GM RT-PCR amplification (Table 1). Briefly for H5, for example, probes were selected from highly conserved regions of target genes specific for HA gene of H5 influenza viruses, by the sequence data available in the Genbank. The sequence data were generated using the sequence analysis of the influenza database at <http://www.ncbi.nlm.nih.gov/genomes/FLU/Database/select.cgi?go=1>. AIV HA gene of H5 subtype from different years was selected, but mainly after 2004. As well as different countries mainly from

Asia, different hosts but mainly avian, especially chicken and duck, Nine hundred twenty six strains were selected and analyzed, and made multiple alignment with software DNASTar 5.06(DNASTAR, Inc., Madison, WI). It selected 20 most conservative oligonucleotides in the HA gene of H5 subtype as detection probe for Luminex (Fig.1). The Tm of probe was kept around 56~58 °C and labeled 5' with -NH<sub>2</sub>.

We designed nested degenerated primers separately before and after probes. There was a pair of universal primer tagged designed primers, and the 5' of downstream reverse primer was labeled with biotin. To amplify HA gene of all the H5 strain, the 3' of each primer should have at least 5 conservative bases. All the above primers and probes had been tested by a software PrimerExpress3.0 (Applied Biosystems Inc. Foster City, CA).

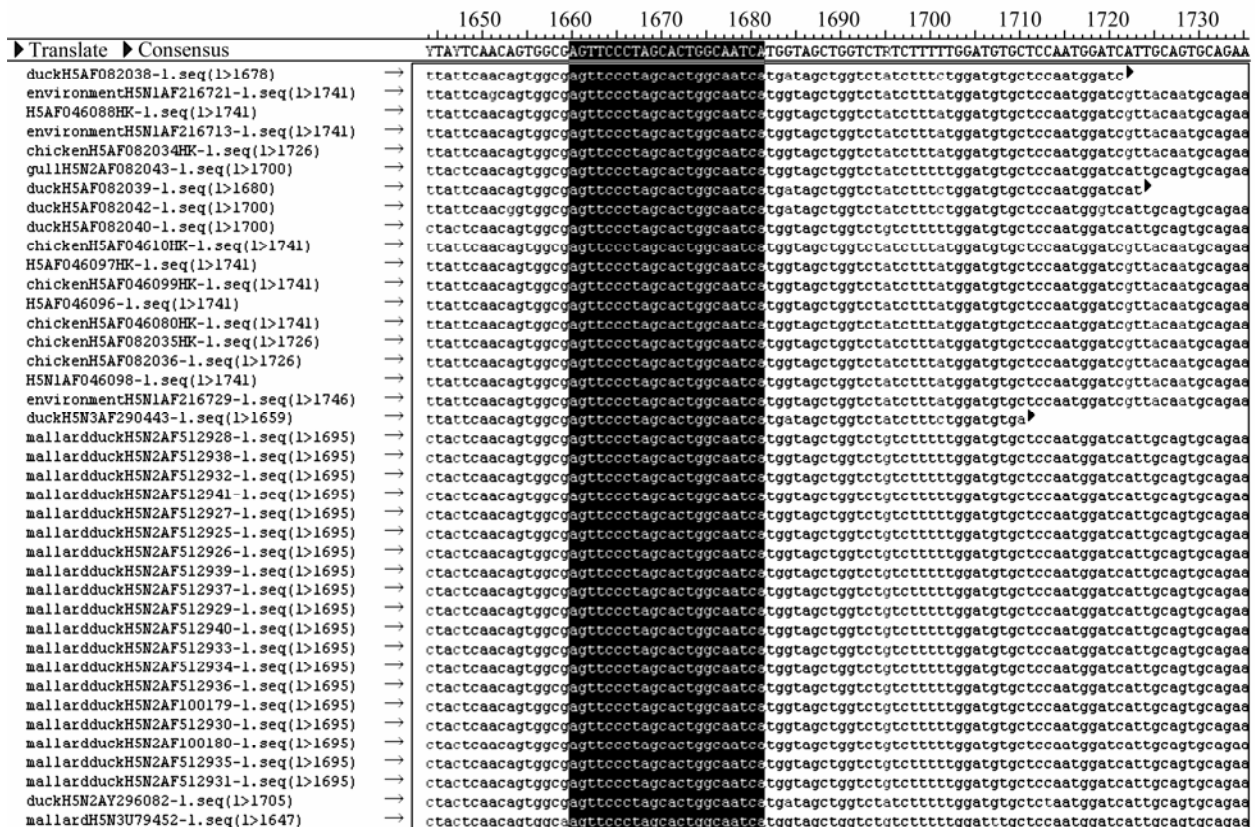


Fig. 1. Location of the probe in HA gene of AIV H5 subtype.

Table 1. Primers and Probes of GMPLex for each subtype of influenza virus

Name	Target gene	Primer name	Sequence	Expected PCR product(bp)
Influenza virus (IV)	M	IVMprobe	ATCATTGGGATCTTGCACTTGA	174
		Ri	CCCTCTTTTCAAACCGTATTTAA	
		IVMRi	TTCTTTGCGTTATGTCTCTGCCCTCTTTTCAAACCGTATTTAA	
		IVMRo	AGGCACTCCTTCCGTAGAAG	
		Fi	AAATGCAGCGATTCAAGTGA	
		IVMFi	CAGGCCACGTTTTGTTCATGCAAATGCAGCGATTCAAGTGA	
Influenza virus (IV)	NS2	IVNSprobe	TCTACAGAGATTTCGCTTGG	145
		IVNS Ri	TTCTTTGCGTTATGTCTCTGGTTCTCGCCATTTTCCGTTTC	
		IVNSRo	AAWYTAGATGCAAATTCTGCA	
		IVNSFi	CAGGCCACGTTTTGTTCATGCTTGAATGGAATGATAACACAG	
		IVNSFo	TGCAATTGGGGTCCCTCATCGG	
AIV H5	HA	RAIVH5Probe	AGTTCCTTAGCACTGGCAAT	185
		Ri	ATCCATTGGAGCACATCCA	
		RAIVH5Ri	TTCTTGCGTTATGTCTGCTGATCCATTGGAGCACATCCA	
		AIVH5Ro	AAWYTAGATGCAAATTCTGCA	
		Fi	CAAGRCTAAACAGRGAGGAAATA	
		RAIVH5Fi	CAGGCCACGTTTGTTCATGCCAAGRCTAAACAGRGAGGAAATA	
		AIVH5Fo	TGACTATCCRCARTAYTCAGAAGA	
AIV H9	HA	AIVH9Probe	CTCCACACAGAGCACAATGG	218
		Ri	TCGACGATGTAGGACCATTCT	
		AIVH9Ri	TTCTTTGCGTTATGTCTCTGTGACGATGTAGGACCATTCT	
		AIVH9Ro	CCRGGGTAACACATTTCCATT	
		Fi	MCAATGTTCTGTGACACAT	
		AIVH9Fi	CAGGCCACGTTTTGTTCATGCMCAATGTTCTGTGACACAT	
		AIVH9Fo	GCTACCAATCAACAACTCCAC	
AIV H7X	HA	AIVH7XProbe	TTGGTTTAGCTTCGGGGC	122
		Fi	GCAGYGGCTACAAAGATGTGA	
		AIVH7XFi	CAGGCCACGTTTTGTTCATGCGCAGYGGCTACAAAGATGTGA	
		AIVH7XFo	RATACAGATTGAYCCAGTCA	
		Ri	AGACAAGGCCATTGCAA	
		AIVH7XRi	TTCTTTGCGTTATGTCTCTGAGACAAGGCCATTGCAA	
AIV H72	HA	AIVH72Probe	GCAGTTTGAGCTGATAGACAATGA	160
		Fi	ATCACAGGCAAATTGAATCGT	
		AIVH72Fi	CAGGCCACGTTTTGTTCATGCATCACAGGCAAATTGAATCGT	
		AIVH72Fo	TGCAGCTGACTACAAAAGCAC	
		Ri	TTGCGTCTCGTGTCCAATT	
		AIVH72Ri	TTCTTTGCGTTATGTCTCTGTTGCGTCTCGTGTCCAATT	
Influenza virus	HA	IVH1N2Probe	GGTTGATGATGGTTTTCTGGAT	198
		IVH1N1Probe	AGTTGATGATGGATTTCTGGAC	
		Fi	TGAAAAGATGAACACACAATTCA	
		IVH1Fi	CAGGCCACGTTTTGTTCATGCTGAAAAGATGAACACACAATTCA	
		IVH1Fo	AGAGCACACAGAATGCCATT	
		Ri	TCCAAAGTTCTTTTCATTTTCCA	
		IVH1Ri	TTCTTTGCGTTATGTCTCTGTCCAAAGTTCTTTTCATTTTCCA	
		IVH1Ro	TTTGACGCTTTCCATGCA	
Influenza virus H3(redesign)	HA	RIVH3Probe	TGGATTTCTTTGCCATATCAT	145
		Fi	AGCTGAAGTCAGGATACAAAGA	
		RIVH3Fi	CAGGCCACGTTTTGTTCATGCAGCTGAAGTCAGGATACAAAGA	
		RIVH3Fo	TTCAAATATACCACAAATGTGA	
		Ri	TTTTGGCAGGCCACAT	
		RIVH3Ri	TTCTTTGCGTTATGTCTCTGTTTTGGCAGGCCACAT	
RIVH3Ro	ATGTTGCACCTAATGTTGCC			

Table 1 (continued)

Influenza virus N1	NA	IVN1Probe	TGGTCTTGCCAGACGGTGCTG	126
		Fi	GGAGCAGCATATCTTTTTGTGG	
		IVN1Fi	CAGGCCACGTTTTGTTCATGCCGAGCAGCATATCTTTTTGTGG	
		IVN1Fo	TAAGACCTGTTTCTGGGTTGAG	
		Ri	TGTCAATGGTGAATGGCAAC	
		IVN1Ri	TTCTTGCCTTATGTCTCTGTGTCAATGGTGAATGGCAAC	
		IVN1Ro	AACAAGGAGTTTTTGAACAACAACTACT	
Influenza virus N2	NA	RIVN2Probe	AGGCTCATGGCCTGATGG	131
		RFi	CAGTATTGTTGTGTTTTGTGG	
		RIVN2Fi	CAGGCCACGTTTTGTTCATGCCAGTATTGTTGTGTTTTGTGG	
		RIVN2Fo	GGAAACTGAAGTCTKGTGGAC	
		RIVN2Ri	CTTATATAGGCATGAGATTGAT	
		RIVN2Ri	TTCTTGCCTTATGTCTCTGCTTATATAGGCATGAGATTGAT	
		RIVN2Ro	TTTTTCTAAAATTGCGAAAG	

Using the same procedure described above, we designed corresponding nested-degenerated primer and probe targeted highly conserved fragments for NS gene and M gene against all subtype IV, HA gene against H1, H3, H7 and H9 subtype, and NA gene against N1 and N2 subtype (Table 1) manually.

#### RNA isolation

Viral RNA was extracted from the supernatant of the allantoic fluids from different birds by using QIAamp RNA extraction kit (Valencia, CA) according to the manufacturer's instruction. The RNA was eluted from the QIASpin columns in a final volume of 50  $\mu$ L of elution buffer and immediately stored at -70  $^{\circ}$ C until used.

#### Development of a GM RT-PCR for the detection of single subtype of IV

The General multiplex RT-PCR (GM RT-PCR) was carried out in a reaction mixture (50  $\mu$ L volume) containing 1  $\times$  RT-PCR reaction buffer (Qiagen), 0.4 mmol/L each of four dNTPs (Qiagen), 2.5 mmol/L MgCl<sub>2</sub> (Qiagen), 2  $\mu$ L one step RT-PCR enzyme mix (Qiagen), 0.6~2.4  $\mu$ mol/L of super primer mixture, 0.05~0.2  $\mu$ mol/L single primer, 10 U RNase inhibitor (Qiagen), 2  $\mu$ L of RNA template. The GM RT-PCR conditions were step 1, reverse-transcription, 50  $^{\circ}$ C

for 35 min; step 2, Inactivate reverse-transcriptase and hot-start, 95  $^{\circ}$ C for 15 min; step 3, enrich, fifteen cycles of 94  $^{\circ}$ C 30 s, 52  $^{\circ}$ C 1 min and 30 s, and 72  $^{\circ}$ C for 1 min; step 4, adding tail, 6 cycles of 94  $^{\circ}$ C 30 s, 70  $^{\circ}$ C 1 min and 30 s; step 5, amplification, 35 cycles of 94  $^{\circ}$ C 30 s, 55  $^{\circ}$ C 30 s, and 72  $^{\circ}$ C for 30 s, followed by a final 72  $^{\circ}$ C for 10 min were done. The products were subjected to electrophoresis on a 1.5% agarose gel and visualized under UV.

#### Development of a GM RT-PCR for the detection of multiple subtypes of AIV

*Development of a GM RT-PCR for the detection of two subtype IVs*

2  $\mu$ L of viral RNA from each strain, with any two combination mixtures from the following strains, A/Chicken/HK/HI/1997(H5N1), A/Chicken/HK/921/1997(H9N2), A/PFV/Restock/1/1934(H7N1), A/Swine/SZ/111/2005(H1N1), A/Swine/SZ/321/2005(H3N2), GM RT-PCR was then done by the protocol described in 2.2.4.

*Development of a GMT RT-PCR for the detection of H5, H7 and H9 subtype*

Viral RNA of 2 $\mu$ L from each of the A/Chicken/HK/HI/1997(H5N1), A/Chicken/HK/921/1997(H9N2), A/PFV/Restock/1/1934(H7N1), with combination

mixtures, then the GM RT-PCR followed the protocol described in 2.2.4.

*Development of a GMT RT-PCR for the detection of five subtypes of IV*

Viral RNA of 2µL from each of the above 5 strains, with combination mixture, set the total volume as 50 µL and then GM RT-PCR followed the protocol described in 2.2.4.

**Development of a GMPLex to detect AIV**

*Binding the oligonucleotide probe to Luminex carboxylated fluorescent coding microspheres by one-step EDC coupling reaction*

According to the sequence of fluorescent coding microspheres suggested by manufacture, 10 of them were selected, which had the lowest fluorescent signal misreading probability, binding with the following probe according to table 2, NS gene, M gene of IV, HA gene of AIV subtype H1, H3, H5, H7 and H9, NA gene of N1 and N2. The couple reaction of fluorescent code microsphere to probes of AIV was done according to manufacturer’s instruction. Briefly, all reagents were recovered to room temperature (RT), probes diluted to 0.2 mmol/L with 0.1 mol/L MES solution (pH 4.5). All the following operations were done in a dark room. The fluorescent coding microspheres storage solution were votexed evenly, then 200 µL solution ( $5 \times 10^6$ ) was aspirated to 1.5 mL eppendorf tube, centrifuged at 10 000 r/min for 5 min,

supernatant discarded, added to 50 µL 0.1 mol/L MES solution (pH4.5), vortexed and added to the appropriate amount of 0.2 nmol probes. The solution was mixed and 10 µL of fresh made EDC solution (10 mg/mL) was added and mixed and incubated for 30 min. This step was repeated, 1 mL Tween -20(0.02% w/v) was added to wash the microsphere, centrifuged at 10 000 r/min for 10 min, supernatant discarded, 1.0 mL SDS(0.1% w/v) added, centrifuged at 1 000 r/min for another 10 min to precipitate the microsphere. The appropriate amount of 0.1 mol/L MES (pH 4.5) was added. The microsphere was counted by using blood cell counting assay, the concentration of the microsphere calculated and then diluted the microsphere concentration to 5 000 microspheres per 1 µL with MES solution, and this solution stored at 2~8 °C.

**Development of a GMPLex to detect single subtype of IV**

To confirm whether the various IVs subtype probes coupled microsphere could hybridize with the corresponding product of GM RT-PCR, GM RT-PCR products were taken from IVs subtypes of A/Chicken/HK/HI/1997(H5N1), A/Chicken/HK/921/ 1997(H9N2), A/PFV/Restock/1/1934(H7N1), A/Swine/ SZ/111/2005 (H1N1), A/Swine/SZ/321/2005(H3N2), hybridized with the corresponding probe coupled microspheres according to the following protocol. All reagents were recovered to room temperature (RT) and all the

Table 2. Microspheres Code and its corresponding AIV probes

Number	1	2	3	4	5
Microspheres	33	34	35	36	37
Probe	IV M probe	IV NS probe	RAIV H5 Probe	AIV H9 Probe	AIV H7x Probe
Target	A subtype	A subtype	H5 AIV	H9 AIV	H7 AIV
Number	6	7	8	9	10
Microspheres	38	42	43	44	45
Probe	AIV H72 Probe	IV H1 Probe	RIV H3 Probe	IV N1 Probe	RIV N2 Probe
Target	H7 AIV	H1 IV	H3 IV	N1 IV	N2 IV

following operations were done in a dark room. First 50  $\mu\text{L}$  of fresh made GM RT-PCR product was purified by using Qiagen DNA Fragment Purification Kit (Valencia, CA), diluted the purified PCR product to 36  $\mu\text{L}$ . the probe coupled microspheres prepared in step 2.3.1 were taken with 10  $\mu\text{L}$  respectively and further diluted with 1.5 $\times$ TMAC hybridization solution to 200 microspheres per 1  $\mu\text{L}$ . then probe coupled microspheres working solution were added to the 96 U microplate and set 17  $\mu\text{L}$  of TE solution (pH 8.0) as blank control. purified and diluted GM RT-PCR product with 5  $\mu\text{L}$  were added to the wells, the microplate coated with the film and the reaction in a PCR apparatus to hybridize was performed. The reaction condition was set as 95  $^{\circ}\text{C}$  5min, 52  $^{\circ}\text{C}$  15 min, then centrifuged at 10 000 r/min for 3 min to collect the microspheres. Fresh Reporter Mix was done by diluting streptavidin-R- phycoerythrin to 10  $\mu\text{g}/\text{mL}$  in 1 $\times$ TMAC Hybridization Solution. Reporter Mix of 25  $\mu\text{L}$  was added to each well and gently mix by pipetting several times. The reaction plate was incubated at the hybridization temperature for 5 min. Total 50  $\mu\text{L}$  at hybridization temperature were done on the Luminex analyzer according to the system manual for analysis.

According to the determination criteria recommended by Luminex, if the microspheres of each fluorescent is no less than 20 and the fluorescent number of the blank control is not more than 3 000, the results can be standardized. Luminex qualitative ratio result (LQRR) equal to Median fluorescence intensity (MFI) divided by MFI of blank control, that is  $\text{LQRR} = \text{MFI}_S / \text{MFI}_B$ . The  $\text{LQRR} \geq 3$ , is positive; if the LQRR is between 2 and 3, is suspicious; and  $\text{LQRR} \leq 2$ , is negative.

### **Development of a GMPLex to detect multiple subtypes of IV**

Ten fluorescence coded microsphere with 100  $\mu\text{L}$  of respectively were diluted with 1.5 TMAC hybridization solutions to 200 microspheres per 1  $\mu\text{L}$ . It was vortexed and made into a mixture and other steps were the same as the protocols described in 2.3.2. The set machine could detect 10 fluorescence coded microspheres simultaneously, determination criteria was the same as the methods described in 2.3.2. In order to verify whether GMPLex detection results were identical with subtypes of different AIV, the above 5 different AIV GM RT-PCR products were taken respectively, then 2 or 3 combinations of the GM RT-PCR products were used, and 5 of them were hybridized with the 10 fluorescence coded microspheres mixtures, following the conditions described in 2.3.2.

### **Sensitivity and specificity of GMPLex high throughput test**

The preparation of RNA transcripts of different subtypes of IV for GMPLex was conducted with the stain of A/Chicken/HK/HI/1997(H5N1). The sensitivity of the GMPLex was evaluated with different (50% embryo lethal dose,  $\text{ELD}_{50}$ ) ranging from 1 to  $10^9$  with ten-fold dilutions<sup>[14]</sup>. The assay of each IV dilution was conducted in triplicates. The specificity of the GMPLex was examined by using RNA extracted from 83 viral samples delivered from Hongkong; all of the samples were investigated with virus isolation and the GMPLex respectively.

## **RESULTS**

### **GM RT-PCR for single subtype of influenza virus**

The nucleotide sequences of NP gene and M gene were highly conserved in all subtypes of influenza

Table 3. Result of GM RT-PCR for multiplex strains mixture of influenza virus

Virus mixture	H7x	N1	N2	NS	H3	M	H5	H1	H9	Fragment shown in gel
H5N1+ H7NX+H9N2	+	+	+	+	-	+	+	-	+	3
H5N1+H7NX+H9N2+ H1N1+H3N2	+	+	+	+	+	+	+	+	+	5

viruses. Five virus strains were all amplified with 174 bp fragment targeted M gene and 145 bp fragment targeted NS1 (data not shown). For a single subtype, only one of the GM RT-PCR reactions are the product of expected size. The size of PCR product ranged from 126 bp to 245 bp, depending on the HA and NA subtype. The specificity of GM RT-PCR indicated it was positive for H5 gene and N1 gene of A/Chicken/HK/HI/1997(H5N1), H1 gene and N1 gene of A/Swine/SZ/111/2005(H1N1), H3 gene, N2 gene of A/Swine/SZ/321/2005(H3N2), H7 gene of A/PFV/Restock/1/1934(H7N1), H9 gene of A/Chicken/HK/921/1997(H9N2)(Fig. 2,Table 1).

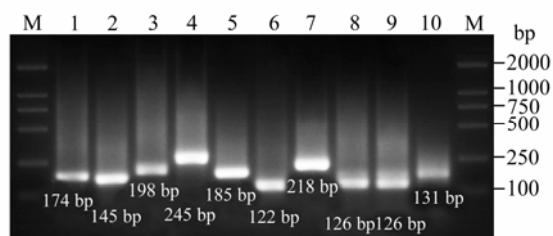


Fig.2. Gel electrophoresis result of GM RT-PCR for single subtype of influenza virus. M: Marker DL 2000; 1: M gene of A/Chicken/Hongkong/SZ-HI/1997(H5N1); 2: NS gene of A/Chicken/Hongkong/SZ-HI/1997(H5N1), 3: HA gene of H1 subtype(A/Swine/Shenzhen/111/2005(H1N1)); 4: HA gene of H3 subtype(A/Swine/Shenzhen/321/2005(H3N2)); 5: HA gene of H5(A/Chicken/Hongkong/SZ-HI/1997(H5N1)); 6: HA gene of H7(A/PFV/Restock/1/1934(H7N1)); 7: HA gene of H9(A/Chicken/Shenzhen/921/1997 (H9N2)); NA gene of N1(A/Chicken/Hongkong/SZ-HI/1997 (H5N1)); NA gene of N1(A/Swine/Shenzhen/111/2005 (H1N1)); NA gene of N2(A/Swine/Shenzhen/321/2005(H3N2)).

**GM RT-PCR for multiple strains mixture of IV**

GM RT-PCR could amplify expected products from mixture of multiplex Influenza A virus, as A/Chicken/HK/HI/1997(H5N1), A/Chicken/HK/921/1997(H9N2), A/PFV/Restock/1/1934(H7N1), A/Swine/SZ/111/2005(H1N1), A/Swine/SZ /321/2005(H3N2)) (Table 3). The result of amplification was a smear because of strategy of asymmol/Letricol RT-PCR for 9 genes (Fig. 3).

**GMPLex for single subtype of influenza virus**

MFI of GM RT-PCR product ranged from 81~18826, according the guideline of Luminex, the LQRR result of H72 was 1 and the result was negative, indicating that there was no specific hybridization between microsphere #38 and H7x GM RT-PCR products. Other probes were positive, the value of LQRR was between 3~27(Table 4).

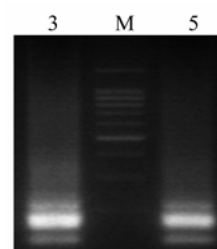


Fig. 3. Gel electrophoresis of GM RT-PCR for Multiplex strains mixture of influenza virus. M: 100 bp DNA Ladder Marker (Qiagen, Valencia, CA);Lane 3: A/Chicken/Hongkong/SZ-HI/1997(H5N1) + A/Chicken/Shenzhen/921/1997(H9N2) + A/PFV/Restock/1/1934(H7N1) ;Lane 5: A/Chicken/Hongkong/SZ-HI/1997(H5N1) + A/Chicken/Shenzhen/921/1997(H9N2) + A/PFV/Restock/1/1934(H7N1) + A/Swine/Shenzhen/111/2005 (H1N1) +A/Swine/Shenzhen/321/2005(H3N2).



Table 4. MFI value of GMPLex for single subtype

Code	33	34	35	36	37	38	42	43	44	45
Probe	IVM	IVNS	RAIVH5	AIVH9	AIVH7X	AIVH72	IVH1	RIVH3	IVN1	RIVN2
MFI <sub>B</sub>	180	621	80	38	149	73	694	185	50	26
MFI <sub>S</sub>	976	2003	608	599	608	81	18826	542	609	833
Cutoff	540	1863	240	114	447	219	2082	555	149	78
LQRR	5	3	8	16	4	1	27	3	12	32
<b>Result</b>	+	+	+	+	+	-	+	+	+	+

Table 5. MFI value of GMPLex for multiplex strain mixture

Code	33	34	35	37	38	36	42	43	44	45
probe	IVM	IVNS	RAIVH5	AIVH7X	AIVH72	AIVH9	IVH1	RIVH3	IVN1	RIVN2
MFI <sub>B</sub>	1835	2839	143	647	216	322	1245	583	124	172
MFI <sub>H5N1+H7NX+H9N2</sub>	13658	11377	12643	14729		16400			12390	8470
LQRR <sub>H5N1+H7NX+H9N2</sub>	7.4	4.0	88.4	22.8		50.9			99.9	49.2
MFI <sub>H5N1+H7NX+H9N2+H1N1+H3N2</sub>	14615	8479	9683	13225		12801	15239	14644	10798	6848
LQRR <sub>H5N1+H7NX+H9N2+H1N1+H3N2</sub>	8.0	3.0	67.7	20.4		39.8	12.2	25.1	87.1	39.8

### GMPLex for multiple strains mixture of influenza virus

Using mixture of GM RT-PCR product of either 3 mixtures of different subtype strains or 5 mixtures of subtypes, the probe mixture was hybridized. MFI of the product was between 6848~16400 and the MFI of blank control was lower than 3 000. Probe mixture had specific reaction with GM RT-PCR product and LQRR was 3~99.9. (Table 5)

### Sensitivity of the GMPLex

The sensitivity of the GMPLex was determined by testing a serial dilution of allantoic fluid of the A/Chicken/HK/HI/1997 (H5N1). GMPLex could detect H5 fragment or N1 fragment amplified by single pairs of primers GM RT-PCR in  $10^{-5}$  (equivalent to 280 ELD<sub>50</sub>), which was tenfold less compared with the sensitivity of real time RT-PCR of the national standard in  $10^{-5}$  (equivalent to 28 ELD<sub>50</sub>). H5 and N1

were subtype together in 2800 ELD<sub>50</sub> (Table 6). When Avian Influenza virus A/Chicken/HK/HI/1997(H5N1), A/Chicken/HK/921/1997(H9N2), A/PFV/Restock/1/1934 (H7N1) mixed together, its detection sensitivity was 2800 ELD<sub>50</sub>, 28000 ELD<sub>50</sub> and 280 ELD<sub>50</sub> respectively.

### Specificity of the GMPLex

The specificity of GMPLex assay was evaluated with the main infectious respiratory diseases including Newcastle diseases virus(NDV) (LASOTA), infectious bronchitis virus(IBV) (4/91), infectious laryngotracheitis virus(ILTV) (isolation strain), infectious bursal disease virus(IBDV)(GX strain), marker's disease virus(MDV) (WH strain) and main bacteria infectious diseases including *E.coli* O157, fowl cholera and *Salmonella pullorum*. There was no cross reactions, the MFI was lower than 3000 and the LQRR value was lower than 0.88 (below the 1). Therefore, the assay constructed was specific.

Table 6. Comparison of GMPLex for single type and single strain

		Single subtype		Single strain			
	Code	35	44	35	44	Code	
	Sample/probe	RAIVH5	IVN1	RAIVH5	IVN1	Sample/probe	
1	LQRR <sub>10</sub> <sup>-1</sup>	7.32	10.80	7.1	9.81	LQRR <sub>10</sub> <sup>-1</sup>	1
2	LQRR <sub>10</sub> <sup>-2</sup>	6.91	8.72	6.38	8.16	LQRR <sub>10</sub> <sup>-2</sup>	2
3	LQRR <sub>10</sub> <sup>-3</sup>	4.26	6.41	3.13	4.06	LQRR <sub>10</sub> <sup>-3</sup>	3
4	LQRR <sub>10</sub> <sup>-4</sup>	2.13	3.21	2.09	2.11	LQRR <sub>10</sub> <sup>-4</sup>	4
5	LQRR <sub>10</sub> <sup>-5</sup>	2.08	2.18	1.90	1.93	LQRR <sub>10</sub> <sup>-5</sup>	5
6	LQRR <sub>10</sub> <sup>-6</sup>	1.92	1.96	<2	<2	LQRR <sub>10</sub> <sup>-6</sup>	6
7	LQRR <sub>10</sub> <sup>-7</sup>	<2	<2	<2	<2	LQRR <sub>10</sub> <sup>-7</sup>	7
8	LQRR <sub>10</sub> <sup>-8</sup>	<2	<2	<2	<2	LQRR <sub>10</sub> <sup>-8</sup>	8
9	LQRR <sub>10</sub> <sup>-9</sup>	<2	<2	<2	<2	LQRR <sub>10</sub> <sup>-9</sup>	9

**Evaluation of the GMPLex assay using clinical samples**

To evaluate the clinical sensitivity of the GMPLex method, a total of 83 samples derived from vaccines, detection antigens and separations, which were sequenced, were re-tested by GMPLex. The results indicated that the sensitivity and specificity were identical with virus isolation (Table 7).

**DISCUSSION**

Avian influenza (AI) is a highly contagious disease in poultry and outbreaks can have dramatic economic and health implications. The rapid spread of highly pathogenic H5N1 AIV throughout Asia, Europe, and Africa and its zoonotic potential pose for both public health and the economic integrity of the poultry

industry<sup>[1,4]</sup>. Thus, there is a critical need for comprehensive and sensitive assays for AIV diagnosis in poultry, especially to distinguish the outbreak subtypes. More importantly, the unique characteristics of distinct AIV subtypes complicate widespread vaccination and prevention strategies, further increasing the importance of effective surveillance.

It is not only needed to detect its IVs positive, but also it must subtype the isolates, so as to adopt corresponding measures to control the disease. Current subtyping needs to detect virus with each subtype anti-serum separately. There were reports that some groups adopted multiplex RT-PCR to identify subtype simultaneously, but due to the technical shortcoming of multiplex RT-PCR, the number it could detect was limited<sup>[12]</sup>. This study established a completely new General Multiplex RT-PCR (GM RT-PCR) test method, resolving the chip testing “bottle neck” of having to perform many individual PCR amplifications. The influenza virus subtyping GM RT-PCR test and the LiquiChip test methods were combined to give an IV GMPLex subtyping test method. The GMPLex subtyping test can satisfy the needs of influenza rapid high throughput testing at ports of entry and exit, but also has erected a platform

Table 7. Comparison of virus isolation and GMPLex for detection of IV in 83 clinical samples

Sample subtype*	Virus Isolation	GMPLex
H5N1	15	15
H7	0	0
H9N2	27	27
Negative	41	41
Total	83	83

\*: No. of positive samples for assay

for more novel IV rapid throughput tests. We constructed a platform of molecular differential diagnostic (MDD) assay that could identify, differentiate, and pinpoint the offending pathogen associated with a clinical syndrome. Compared with the routine diagnostic methods of RT-PCR or rRT-PCR, the GMPLex assay had more high throughput and better improved platform.

The application of a multiplex RT-PCR assay (an RT-PCR assay for the simultaneous detection of different viruses in a sample), offers a significant time and cost-saving advantage, especially when large numbers of samples are analyzed [12]. The principal challenge in developing a multiplex PCR system is choosing the right primers to overcome primer-dimer formation. Therefore, the oligonucleotide primers selected for the amplification of IV nucleic acids were analyzed to ensure that they not only meet the essential criteria for optimal PCR primers [5], but also could be used together in a multiplex reaction under identical amplification conditions. The primers were designed with regard to similar length (20- or 21-mers), similar GC content (Table 1), similar annealing temperature, and as little primer-dimer formation as possible between all six primers. An annealing temperature of 55 °C was evaluated to give maximum product yields and specificity. In addition, "hot-start" PCR conditions were used to minimize primer-dimer formation. Another prerequisite for a multiplex PCR assay is that the PCR products must have different sizes to be clearly identified and differentiated after gel electrophoresis. The 10 pair of primers when used together in the multiplex reaction, amplified only specific products of the expected sizes ranging from 122~198 bp, respectively (Table 1), which could easily be distinguished by agarose gel

electrophoresis (Fig. 2).

To achieve rapid subtype identification of influenza virus, universal primers were added to the novel GM RT-PCR test so that one PCR would be able to simultaneously amplify many target fragments and, combined with the luminex high throughput test, an influenza GMPLex rapid and high throughput subtype identification test was developed, which attained the target of rapid, accurate and IV subtype identification. General Multiplex RT-PCR integrated with Luminex (GMPLex) aimed at the establishment of a rapid method for the screening or detection of not only one subtype of AIV, but the relatively multiple subtypes of AIV in all type A virus, as well as successful detection determined on the primer used for GM RT-PCR discussed above, also depended on the probes used in liquid chip test. The probes were designed with regard to similar length (18- or 25-nt),  $T_m$  was 5°C higher than primer's in order to assure hybridization prior to the primers. The LiquiChip system is a bead-based platform that offers the potential to rapidly assay up to 100 different analytes in a single sample; LiquiChip assays are based on xMAP technology and involve the interaction of immobilized, bead-bound capture molecules with a reaction partner (analyte) in solution. A reporter molecule, specific for the analyte, is used to quantify the interaction [6,10,15]. Validation of the specificity of the GMPLex revealed there was no cross-reaction with other avian viruses (data not shown), including NDV, IBV, IBDV, Duck hepatitis virus (DHV), avian enterovirus (AEV) and host-derived RNA.

The GMPLex assay specifically detected the targeted viruses, as it was demonstrated by sequencing of the amplification products. Because the assay

detected 280 ELD<sub>50</sub> ~2800 ELD<sub>50</sub>, theoretically it could identify even one single infected chicken within a pool of several hundreds of chicken swab samples. The results of this study indicate that the GMPLex described in this paper is a specific, sensitive and reliable tool for the simultaneous diagnosis of multiple subtypes of AIV, as it was confirmed by virus isolation. Because sensitivity and specificity of the system was very similar to that of the monospecific assays, the GMPLex is a cost-saving alternative to single PCRs in routine diagnostic submissions or surveys.

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