The Development of Pathogenicity of Avian Influenza Virus Isolated from Indonesia

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Abstract
Highly pathogenic avian influenza outbreak in Indonesia has been reported in various poultry due to H5N1 subtype. The presence of multiple basic amino acids within the cleavage site of HA glycoprotein has been identified to be associated with the pathogenicity of avian influenza virus. The study was retrospective study which was designed to characterize the cleavage site and fusion site region of haemagglutinin gene of AIV isolated from various poultry in 2003 to 2013. Isolation, Identification and propagation were carried out to collect viral stock. For virus detection, reverse transcriptase PCR (RT-PCR) method on H5 and N1 gene fragment was performed. All of RT-PCR HA gene positive products were sequenced for further nucleotide analysis and to determine the nucleotide composition at the targeted fragment. The results are all AIV isolates were identified as H5N1 subtype. The sequence analyses revealed some motives of basic amino acid motive that were classified as highly pathogenic avian influenza virus. Further analyses on fusion domain of all AIV isolated during the period 2003 to 2013 showed conserved amino acid.

Keywords: avian influenza, haemagglutinin, cleavage site, basic amino acid, fusion site

Introduction
Highly pathogenic avian influenza (AI) in Indonesia has been reported first time as causative agent of mortality in commercial farm in West Java and Central Java, in since 2003. The outbreak was continued to spread to other provinces and all provinces have reported the disease, except Gorontalo and Maluku provinces (Komnas FBPI, 2008). Unfortunately in 2011, AI outbreak has been reported in Gorontalo (http://ditjenmak.deptan.co.id), where previously known as free AI cases and the infection still existing sporadically throughout Indonesia. So far it has been known that the infection is caused by H5N1 subtype (Asmara et al., 2005; Dharmayanti et al., 2005-a; Mahardika 2005; Wibowo et al., 2006; Wibowo et al., 2007). The virus is highly pathogenic for poultry and it causes high economic loss impact, owing to high mortality of birds, birds eradication cost, and poultry product export-import restriction.

Avian influenza virus is a member of the family Orthomyxoviridae, genus Influenza A virus. Viral genome is segmented and consist of single stranded-RNA, negative sense, and has eight segments gene which responsible for internal and surface protein. Surface protein consists of Haemagglutinin (H/HA), neuraminidase (N/NA) and matrix (M2). Internal proteins are nucleoprotein (NP), polymerase complex (PB1, PB2, and PA), matrix (M1), and nonstructural protein (NS1 and NS2). Influenza A virus is classified into type A, B, and C based on matrix or
nucleoprotein antigenic character. Influenza virus A are further divided into subtype based on antigenicity character of HA and NA proteins (Cox and Kawaoka, 1998). To date, 16 HA and 9 NA subtype have been recognized (Fourchier et al., 2005).

Haemagglutinin (HA) is glycoprotein projection viral and formed as a homotrimer comprising three identical subunits that contain two polypeptide chains called HA-1 and HA-2. This polypeptide is linked in each subunit by single disulphide bond (McCauley and Mahy, 1983). It is encoded by the fourth largest RNA segment and widely distributed on the surface of virion reaching approximately 80%. This Haemagglutinin is a principal antigen which plays vital role for infection. In the process of infection, HA will undergo post translational cleavage by certain protease enzyme into HA1 and HA2 sub-unit(Cox and Kawaoka, 1998). The HA1 is receptor binding protein as the major target of immune responses, whereas HA2 is an anchor protein of the envelope, mediating fusion of the envelope to the cellular endosomal membrane (Suzuki and Nei, 2002).

Recent studies have revealed an understanding of pathogenicity at molecular level. It has been shown to be associated with the presence of multiple basic amino acids at the cleavage site of the HA glycoprotein (Senne et al., 1996). Cleavage site (CS) region of HPAI viruses differ from those of a virulent influenza A viruses by the virtue of possessing multiple basic amino acid of the carboxyl terminus of HA1, especially arginine and lysine. This feature permits cellular protease such as furine-like endoprotease to recognize multiple basic amino acids, which in turn cleaves the HA and renders the virus infection. Such a kind of condition permits the virus to spread in a variety of organs, leading to systemic infection. Whereas most a virulent strains have a single arginine or monobasic cleavage site that will be cleaved only by trypsin secreted from cells in the respiratory and intestinal tract, so the virus only produced localized infection (Harimoto and Kawaoka, 2001; Ito et al., 2001).

Field examination focusing on clinical signs of infected poultry in Indonesia indicated some variations in macroscopic lesions. Formerly, the infected bird exhibited characteristic symptoms, e.g.: high mortality rate, hemorrhages on the shank and hock joint, congestion and cyanosis on the comb and wattle, which can be followed by edema gelatinous. The development of clinical signs, started in 2005, today in some cases of AI infection could not always indicate specific symptoms although the circulating virus remain highly pathogenic H5N1. So far, there has not been any evidence that another H5N1 subtype and pathotype infection occurs in poultry. This research was aimed to characterize the fragment of HA gene of avian influenza virus responsible for pathogenicity marker of AIV, isolated from various poultry since 2003 to 2013 would be elucidated by amplifying and sequencing of this regions.

Materials and Methods
The study was retrospective study which was designed to characterize the cleavage site and fusion site of HA gene fragment of AIV isolated from 2003 to 2013. Isolation, identification, and molecular characterization was done in the Laboratory of Microbiology, Gadjah Mada University and Biotechnology Center, Disease Investigation Center, Lampung. Sequencing was performed in the laboratory which equipped with sequencing facilities in PT. Charoen Pockphan Indonesia, Jakarta and Centre for Veterinary Biologics (Pusat Veterinaria Pharma), Surabaya.

Materials
Field poultry samples suspected to AIV infection from several outbreak places in Java Island and Lampung province from 2003 to 2013 have been collected. Those samples were collected from AI cases with and without specific symptoms for AIV infection. Some samples stocked as organ samples need for further process to viral isolation and propagation.
Methods
Isolation and Propagation

Lung samples collected from suspected poultry have been prepared for further processed and then inoculated into allantoic sac of SPF embryonated chicken egg of 11 days old. Cropped viral in allantoic fluid was tested on its ability to agglutinate chicken red blood cell or haemagglutination (HA) test. HA positive indicated the growth of both AIV and NDV. Positive HA test isolate then continued to haemagglutination inhibition (HI) test using specific antibody to AIV H5N1 subtype to confirm the presence of AIV. Isolation and identification were based on the procedures of World Health Organization on avian influenza diagnostic and surveillance (WHO, 2002). Infected allantoic fluid samples were stored for further processed of molecular characterization.

Viral Sub-typing Identification
RNA extraction.

Viral RNA was extracted from allantoic fluid using RNA extraction kit Invitrogen Pure Link™ Micro to Midi 50xRxn Total RNA Purification System (Catalog number: 12183-018) according to the manufacturer’s protocol. To purify total RNA from 0.2 ml allantoic fluid, in a 1.5 ml RNAse-free micro centrifuge tube was added 0.2 ml of RNA lyses solution contain 1% (v/v) 2-mercaptoethanol. The mixture was vortexed thoroughly to disrupt and lyse blood cells, and centrifuged at 12,000xg for 2 minutes at room temperature. Supernatant was transferred to clean 1.5 ml RNA-se free micro centrifuge tube and 200 μl of 100% ethanol was added. Any precipitate was dispersed by vortexing or pipetting up and down several times. The sample was transferred to the RNA spin cartridge, centrifuged at 12,000xg for 15 seconds at room temperature. The flow-through was discarded and 700 μl of Wash Buffer I to the spin cartridge was added and centrifuged at 12,000xg for 15 seconds at room temperature. The spin cartridge was placed into a clean RNA Wash Tube and 500 μl of Wash Buffer II with ethanol was added to the spin cartridge then centrifuged at 12,000xg for 15 seconds at room temperature. The flow-through was discarded. This step was repeated once. The spin cartridge was centrifuged at 12,000 x g for 1 minute at room temperature to dry the membrane, and the cartridge was moved into an RNA Recovery Tube. Sixty μl of RNase-free water was added to elute the RNA and incubated for 1 minute at room temperature then centrifuged for 2 minutes at 12,000xg at room temperature. This step was repeated, and then the cartridge was discarded and elutes was stored at -4° C.

Thermocycling

One step Reverse transcriptase-PCR (RT-PCR) was carried out in GeneAmp ® PCR System 2400 machine and performed using Invitrogen SuperScript™ III One-Step RT-PCR System. Primers designed for this research were H5 and N1 and listed in the Table 1.

Primers 1 and 2 used to amplify fragment HA and NA gene of AIV isolated from 2003 to 2008, meanwhile primers 3 and 2 used to amplify fragment HA and NA gene of AIV obtained from 2009 to 2013. However, primer 3 from AAHL. (2010) also be used to amplify some AI viruses isolated up and down several times. The sample was transferred to the RNA spin cartridge, centrifuged at 12,000xg for 15 seconds at room temperature. The flow-through was discarded and 700 μl of Wash Buffer I to the spin cartridge was added and centrifuged at 12,000xg for 15 seconds at room temperature. The spin cartridge was placed into a clean RNA Wash Tube and 500 μl of Wash Buffer II with ethanol was added to the spin cartridge then centrifuged at 12,000xg for 15 seconds at room temperature. The flow-through was discarded. This step was repeated once. The spin cartridge was centrifuged at 12,000 x g for 1 minute at room temperature to dry the membrane, and the cartridge was moved into an RNA Recovery Tube. Sixty μl of RNase-free water was added to elute the RNA and incubated for 1 minute at room temperature then centrifuged for 2 minutes at 12,000xg at room temperature. This step was repeated, and then the cartridge was discarded and elutes was stored at -4° C.

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<table>
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<th>Table 1. Primer used in this study.</th>
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in 2008. Cycling condition for H5, HA.30, and NA1 were initially done with a reverse transcription step at 50°C for 30 minutes and a hot start step at 94°C for 5 minutes. For H5 amplification was performed as follow: denaturation (94°C for 30 seconds), annealing (50°C for 1 minute) and extension (68°C for 45 seconds), and ended by final extension at 68°C for 5 minutes. For HA.30 amplification was performed as follow: denaturation (94°C for 30 seconds), annealing (50°C for 40 second), extension (68°C for 40 seconds), and ended by final extension at 68°C for 5 minutes. Amplification of NA gene using NA-1 primer was done as follow: denaturation (95°C for 30 seconds), annealing (55°C for 45 seconds), extension (72°C for 60 seconds), and ended by final extension at 72°C for 5 minutes. Total cycles performed in PCR session was 40 times.

Electrophoresis of PCR Product

PCR products were electrophoresis in 1.5% Agarose gel in 1x TBE. Electrophoresis tank used MSMDIDUO, Cleaver Scientific Ltd. Gel was running at 100V for 40 minutes, and read under UV transilluminator at 302 nm wave length to determine the size of PCR product fragment.

Sequencing and sequence analysis

PCR products of HA gene were sent to laboratory which has sequencing facilities, and prior to be sequenced the PCR product should be purified. Primers used to sequence the targeted HA fragment genes were the same with previous primers to amplify. Sequences yielded were analyzed by MEGA version 5.0, include sequence editing, multiple alignment, and deductive amino acid prediction (Tamura et al., 2007).

Result and Discussion

Molecular identification

Propagation and identification of all samples showed positive for AIV H5N1 and negative for Newcastle disease virus (NDV), which were characterized using serum anti specific for both AIV and NDV. Molecular identification in this study was based on the amplification of H5 fragment gene at position 775 to 1021 with product size at 246 bp (Ito et al., 2001) particularly for AIV isolated from 2003 to 2008 (Figure 1), meanwhile AIV obtained from 2009 to 2013 used the primer which was targeted of HA gene at position 657 to 1361 with product size at 705 bp (AAHL., 2010) (Figure 2). Molecular identification of H5 gene according to Wibowo et al. (2007) reported that the same primer designed has been used successfully to characterize fragment HA gene of some virus isolated from 2003 to 2005. Molecular identification has been conducted by Asmara et al. (2005) for AIV.

Figure 1. Electrophoresis profile of HA-5 gene fragment indicated DNA band of 246 bp product size. All samples in lane 11sb, 15 sb, 16 sb, 20 sb, 25 sb, 30 sb and 31 sb were positive result.

Figure 2. Electrophoresis profile of HA-5 gene fragment using AAHL primer indicated DNA band of 705 bp product size. Samples in lane 1,2,3,4,6,7,8,9,10, and 11 were positive, but lane 5 was negative.
isolated from various birds sample taken from Java in the early AIV outbreak. He used H5 primer designed by Spackman et al. (2002) to amplify fragment HA gene. The primer was actually designed for AIV American isolates but it could work for Indonesian isolates. The H5 identification has been reported by Dharmayanti et al. (2005-a and 2005-b) which used primer designed by Lee et al. (2001) with the amplification target at 545 bp. Meanwhile Wibowo et al. (2007), were using WHO primer designed with 600 bp product size. They could identify some AIV isolated from various regions in Indonesia. WHO primer that once suggest by World Health Organization (WHO) also broadly used to VAI detection, particularly in the early AIV outbreak in Indonesia. Primers designed by Australian Animal Health Laboratory already validated and used in all diseases investigation centre throughout Indonesia.

Amplification of NA-1 gene fragment of AIV isolated from 2003 to 2013 was based on primer which were designed by Payungporn et al. (2004) at position 458 to 589 with product size at 131 bp (Figure 3). The N1 sub typing using the same primer designed has been reported by Mahardika (2005) and Wibowo et al. (2007) particularly to detect some cases of AIV infection in an early outbreak in Indonesia. They successfully performed the amplification by using the primer.

**Analysis of cleavage site area.**

Further characterization of AIV at the pathogenicity marker, was based on amplification of HA gene fragment of cleavage site area. It is believed that amino acid configuration at cleavage site region would determine pathotype of AIV. The presence of polybasic amino acids in that region is important as a pathotypic marker of AIV (Bank and Plowright, 2003). Our result of pathogenicity study at molecular level indicated that some AI viruses obtained from West Java, Central Java, Yogyakarta Special Province, East Java, and Lampung Province which were isolated from various species of birds since 2003 to 2013 could be classified into five motives of basic amino acids. Detail of multiple alignment of this study could be found in Table 2. The first motive was –PQERRRKKRR/GLFGAIAGFIE- which could be observed in the early outbreak during the year of 2003 to 2005. The CS feature was the same with the ancestral virus Goose/Guangdong/1/1996. The second motive was –PQRE-RRKKRR/GLFGAIAGFIE- which indicated a deletion of amino acid at upstream -6 and was found in AI viruses isolated in the year 2003, 2004, and 2007. The third motive was –PQRESRRKKRR/GLFGAIAGFIE- and could be found in AIV isolated from 2006 to 2011 which were predominantly showed a mutation upstream -6 of arginine into serine. The forth motive was –PQRESRRRRKRR/GLFGAIAGFIE- which indicated a mutation at upstream -2 lysine into arginine. The last identified motive was –PQRE-RRRKRR/GLFGAIAGFIE- which showed not only deletion at position upstream -6 but also a mutation at upstream-3 lysine into arginine, which was found in AIV obtained in 2012 and 2013 and isolated from waterfowl.

According to Senne et al., (1996), minimal motive of amino acid concept is B X B R//, which B is basic amino acid; X is a non-basic amino acid and R is arginine that is related to restriction area (sign //). Meanwhile Harimoto and Kawaoka (1994) reported two kinds of structural features which are
Table 2. Multiple alignment of cleavage site area at position upstream -16 until downstream +14. Sign / indicated cleavage site position.

<table>
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Note: Bold type indicate investigated AIV isolates
critical for determining HA cleavability by protease. A series of amino acids in
the cleavage site area and the presence or absence of carbohydrate chain in the near vicinity will determine cleavability of HA
gene. In order HA to be cleaved, at least six basic amino acids have to be present at the
 cleavage site if carbohydrate site chain is
 nearby, otherwise only for basic amino acid
 is needed. Following in amino acid level
conversion, all AIV isolates in this research
have the same motive to fulfill the minimal
concept of HPAI virus according to Senne
et al. (1996); Harimoto and Kawaoka (1994), and
still consistent with few kinds of studies that
were reported by another researcher.

The feature of -PQRERRRKKR/ GLFGAIAGFIE- at the cleavage site area in our
result is the same to the most Indonesian AI
viruses in the early of outbreak (Dharmayanti et al., 2005-b; Dharmayanti and Indriani,
2007; and Smith et al., 2006). Compared with
that of H5N1 from Indonesia, Thailand, and
Vietnam seem to be identical and consistent with -PQRERRRKKR/GLF- but differ from
Guandong isolate (A/China/GD01/2006) which lost a basic amino acid (Lys) at position
-2 and showed a Gln to Leu substitution at
position -8 at cleavage site area (Zhou et al., 2007). One of cleavage site motive
in our result is identical with the previous
study that 11 isolates avian viruses from
Vietnam, including A/Dk/Vietnam/568/05, has arginine deletion (PQRE-RRRKR// GLF).It is different from another virus (A/Ck/Vietnam/147/04) that has arginine to
isoleucine substitution at position -5 from the
cleavage site area (Smith et al., 2006). Another
motive which could be reported from our
study was -PQRSRRRKR/GLFGAIAGFIE-
The feature has substitution amino acid at
position upstream -2 lysine into arginine
and a bit similar with other two viruses from
Indonesia (Ck/KP/BBVet-XII-1/04) and (Ck/KP/BBVet-XII-2/04) motive that
have been reported by Smith et al., (2006) which indicated lysine deletion at position
upstream -2 (PQRERRRK-R/GLF-).

Predominantly cleavage site motive
could be found in AIV obtained from the
year 2006 to 2011 and isolated from various poultry. The motive was -PQRESRRRKR/
GLFGAIAGFIE- with a mutation a serine at position upstream -6 into arginine.
According to Dharmayanti and Darminto
(2009), a new motive of basic amino acid
area was discovering at AIV isolated from
poultry in 2006, in West Java, Indonesia. They reported a mutation on upstream-6
from arginine to serine at cleavage site
position within HA gene. The mutation had
never been found in previous isolates, before
2006. Our result emphasized that such kind
mutation could be found not only in AIV
isolated from West Java but also in other
regions in Indonesia.

The new motive which could be
reported from our study was PQREE-RRRKR/
GLFGAIAGFIE- which showed not only
deletion at position upstream -6 but also a
mutation at upstream-3 lysine into arginine.
The motive could be found in the AIV
obtained during 2012 and 2013 and isolated
from waterfowl. Our result is differ with the
previous report in Indonesia that cleavage
site motive of AIV isolated from duck was
-PQRRERRRKR/GLF-, (Wibawa et al., 2012).
The CS feature is also differ to AIV belong
to clade 2.3.2 which were reported for the
first time in China 2009 (Li et al., 2011), in
India (Nagarajan et al., 2012), in Bulgaria and
Rumania (Reid et al., 2011).

Most AIV have arginine on terminal
carboxyl HA-1 at upstream-1 position,
whereas upstream-9 and -10 is the proximal
end that according to another similar study
this position always placed by amino acid
 glutamine at position-9 and proline at
position-10 and these position have so far
been considered as a conserved region
(Dharmayanti et al., 2005-b; Dharmayanti and
Indriani, 2007; Dharmayanti and Darminto,
2009; Susanti et al., 2007; Senne et al., 1996;
and Zhou et al., 1999). Our study showed
the same result with the previous report that
those position are conserved, which can be
observed not only in AIV isolated in the early outbreak but also in current outbreak.

**Analysis of fusion site**

According to Cross et al. (2009), fusion peptide was considered as HA fragment adjacent to cleavage site and has 23 amino acid residues at N-terminal of HA2. Fusion peptide influenza A virus, among 16 VA1 subtype were considered highly conserved and lies at position downstream +1 to +11 or a half of N-HA2 (Nobusawa et al., 1991; Perdue et al., 2003). Fusion domain also rich with hydrophilic amino acid, such as: tryptophan (F), valine, leucine, isoleucine, and alanine. This domain was characterized by glycine residue with certain interval. However, this feature is not indicate specific motif, but it may related to structural fusion peptide (Steinhauer et al., 1995; Cross et al., 2009). Fusion domain analysis of HA2 fragment of AIV H5N1 subtype at glycine residue, is generally consistent at position +1, +4, and +8. According to both Russand Engelman (2000) and Kleiger et al. (2001) reported that glycine residue which has GXXXG or GXXG sequences are considered to have important role in the helix stability and interaction between helices.

Amino acid analysis at position downstream +1 to +11 indicated that all AIV isolates showed highly conserved with the motif -/GLFGAIAGFIE- (Table 2). Our result confirmed previous report of Dharmayanti et al. (2005-b), Dharmayanti and Indriani (2007), Dharmayanti and Darminto (2009), and Susanti et al. (2007) which reported that this motif is highly conserved at HA gene fragment of AIV isolated in Indonesia. Our result also in accordance with previous report of Nobusawa et al. (1991), Senne et al. (1996), Ito et al. (2001), Perdue et al. (2003), Zhou et al. (1999), Zhou et al. (2007). However, an AIV H5N1 subtype has been reported showed an amino acid mutation at position of +8 of fusion peptide domain of HA2 (Susanti et al., 2007).

Amino acid position at downstream +1 and +2 in this study so far remains glycine and leucine. Both are aliphatic amino acid. This result was supported by Dharmayanti and Darminto (2009), Dharmayanti et al. (2005-b), Dharmayanti and Indriani (2007), Susanti et al. (2007), Ito et al. (2001), Senne et al. (1996), Zhou et al. (1999), and Zhou et al. (2007). According to Steinhauer et al. (1995), that fusion peptide HA at position +1 glycine may substituted by alanine, but the mutation will lead HA stability decreased due to an increase of fusion pH. However, +1 glycine cannot be substituted by serine, histidine, leucine, isoleucine, and phenylalanine. The mutation will interfere HA folding and fusion process. Further analysis indicated that glycine residue has a role of structural of virus stability.

**Conclusion**

Based on the data in our study, all AIV isolated 2003 to 2013 could be identified as H5N1 subtype. Molecular characterization at pathogenicity marker of HA gene, revealed some motives at the cleavage site position, that were:-PQRERRKKR/GLFGAIAGFIE-, -PQRE-RRKRR/GLFGAIAGFIE-, -PQRESRRKKR/GLFGAIAGFIE-, -PQRESRRKRR/GLFGAIAGFIE-, and -PQRE-RRRKR/GLFGAIAGFIE-. All the features are classified as highly pathogenic avian influenza virus. Fusigenic domain of all AIV isolated during the period 2003 to 2013 showed conserved amino acid, with the motive is -/GLFGAIAGFIE-.

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