Bioinformatics study of complete amino acid sequences of neuraminidase (NA) antigen of H1N1 influenza viruses from 2006 to 2013 in Iran

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ABSTRACT

Introduction: Influenza is a contagious acute viral disease of the respiratory tract that causes fever, headache, muscle aches and cough. One of the unique features of influenza virus is antigenic variation in viral protein neuraminidase (NA) which causes emergence of new virus variants. NA is responsible for the release and spread of progeny virions. Due to the continuous changes of NA genes, vaccine strains must be re-selected annually. Methods: Complete NA amino acid sequences of 97 strains circulating from 2006 to 2013 in Iran were downloaded from NCBI. The sequences were edited and classified by the year of isolation and their diversity and important changes as well as changes in the predicted ligand binding sites and their resistance to anti-NA drugs, were analyzed. Bioinformatics software such as MEGA6.0, BioEdit, DNAsisMAX and DNASTar were used for the sequence alignments and phylogenetic analyses. Web-based analysis such as SWISS-MODEL, Phyre2 and 3DLigandSite were used for evaluation of the second and third protein structures and prediction of the ligand binding sites. Results: The results showed that 2009 could be considered as an important transition year which caused to classify the isolates into two different distinct groups. This shows the importance of changes made during possible mutations in the genomic structure of the virus which have made it antigenically different from the previous years. Anti-NA drug resistance was observed in 2009. This pandemic strain has become dominant in the following years and is used as a standard vaccine strain from 2010 onwards. Conclusion: The results obtained in this study can aid in better understanding of the antigenic evolution of H1N1 influenza viruses and can potentially accelerate the selection of the vaccine strains.

KEYWORDS: H1N1 influenza virus, Neuraminidase, Antigenic drift, Sequence analysis, Anti-NA drug resistance.

INTRODUCTION

Influenza is a contagious acute viral disease of the respiratory tract that causes fever, headache, muscle aches, sweating, runny nose, sore throat and cough. It is caused by infection with type A or B influenza virus almost every year [1]. Influenza A viruses are enveloped viruses which belong to the Orthomyxoviridae family with segmented genome made of eight single-stranded negative RNA segments from 890 to 2,341 nucleotides that encode 10 and in some cases, 11 proteins [2]. The classification and nomenclature of influenza A viruses are based on the type of hemagglutinin (HA) and neuraminidase (NA) glycoproteins which are presented on the surface membrane [3]. To date, 16 serologically-distinct HA types and 9 different NA types have been described. Recently, a new HA subtype (H17) has been identified in Bats in South America [4]. NA and HA are two main external transmembrane glycoproteins, extended from the surface of the virus envelope.

These two antigens are the most variable antigens that cause protective immunity. HA is the receptor binding glycoprotein which mediates membrane fusion process [5]. NA is responsible for assisting in the virus entry and the release and spread of the progeny virions [6-8]. In fact, NA is the component of the virus membrane with enzymatic activity which is involved in the spread of influenza infection from cell to cell [9, 10]. At the late stages of the viral life cycle, NA removes sialic acids from the virus progeny and the infected cells, thus prevents the virus aggregation and promotes its detachment and release [11].

Antigenic variation is a significant feature in influenza [1]. In other words, changes in proteins of virus specially (i.e. HA and NA) would cause variants that are new in the population. Every year or every few years, occurrences of point mutations that are minor antigenic changes leads to the antigenic drift. The antigenic shifts are major antigenic changes, leading to completely new virus variants that might have a pandemic potential. Vaccination and antiviral drugs are main methods for prevention and treatment of influenza disease. Continuous changes of HA and NA genes (antigenic changes) have made vaccine strains selection challenging and they must be selected.
annually, based on the currently circulating strains. Antiviral drugs, especially NA inhibitors, are the valuable addition to the options used to control the influenza infection [12, 13].

Collecting, reporting and analysis of epidemiologic data obtained from inspection of antigenic changes in each year is an important procedure that global influenza surveillance program has emphasized on it and could be useful in selection of the vaccine strains [14, 15]. The recognition of the importance of anti-NA antibodies in immunity to influenza has led to studies of small molecules that can block the functions of NA, with a view to the antiviral drug development [16]. In the case of NA, the enzyme inhibitors; Zanamivir (GG167; Relenza, GlaxoSmithKline, Research Triangle Park, NC, USA) and Oseltamivir (GS4104; Tamiflu; Genentech, South San Francisco, California and Chugai Pharmaceutical Co., Japan) [17] are already licensed as anti-influenza drugs and are widely marketed for therapeutic and prophylactic uses [18-21]. They act specifically against all subtypes of influenza A and influenza B NAs that prevent the virus from being released from the infected cells.

Although the emergence of antiviral resistance is generally associated with drug treatments, due to the drug pressure, it may emerge coincidentally during a virus evolution when mutations are either neutral or confer an evolutionary advantage on the (epidemic) virus variant and are maintained in the absence of the drug pressure. The emergence and spread of anti-NA drug resistant viruses that should be monitored, emphasizes the importance of developing other effective agents against different viral targets and application of antiviral combinations to reduce the emergence of the resistance [22].

In this study which was conducted for the first time in Iran, the complete amino acid sequences of NA antigen of type A H1N1 influenza viruses available in NCBI GenBank from 2006 to 2013 in Iran were inspected to define the prevalence library and to study the trend of changes in NA sequences while determining their relation with the changes of the virus using bioinformatics methods. The relations of gene alterations and the tertiary structure (3D model) with the relevant regions in bioinformatics methods. The relations of gene alterations and the tertiary structure (3D model) with the relevant regions in bioinformatics methods. The relations of gene alterations and the tertiary structure (3D model) with the relevant regions in bioinformatics methods. The relations of gene alterations and the tertiary structure (3D model) with the relevant regions in bioinformatics methods.

The remarkable point is that, the first group, in terms of amino acid sequences and gene positions, is similar to the sequences of 2006-2008 isolates and in some cases is entirely identical while the second group is similar to the sequences of 2010-2013 isolates and in some cases is entirely identical. This similarity, in terms of alignment analyses of amino acid sequences and the structure of phylogenetic tree between the two different groups in 2009, shows the importance of changes in the genomic structure of the virus in this year which were possibly due to mutations in the genome and the occurrences of frame shift or gene shuffling, causing different sequences and a new group. This new group became dominant in the following years, with significant amino acid changes in NA genome that resulted in isolates with different clinical properties. It was also found that, in 2009, the distinct sequence which had the highest similarity (approximately 99%) with the other sequences of this year (ADJ18168) had also the highest similarity with the sequences of the following years isolates (i.e. 2010-2013) and the distinct sequence which had the lowest similarity (approximately 81%) with the other sequences of 2009 (ACY01439) had the highest similarity with the sequences of the previous year’s isolates (i.e. 2006-2008) (Table 1). This can explain the domination of 2009 pandemic strain from 2010 onwards.

RESULTS

By investigating the results of sequence alignments and phylogenetic trees and percent similarity/divergence matrices and predicted secondary and tertiary structures (3D structures) of the ligand binding sites of different sequences of each year (data not shown), it was found that from 2006 to 2008, the sequences are identical and there is no change in the ligand binding sites, or the type and number of the ligands.

In 2009, with respect to the occurrence of 2009 H1N1pandemic, by studying the available similarities and differences in alignments of the sequences (Fig.1) and by investigating the phylogenetic tree of the 2009 isolates (Fig. 2), it was found that influenza isolates of Iran, are located in two entirely distinct and separated branches and form two different groups in terms of amino acid properties.

MATERIALS and METHODS

Complete amino acid sequences (complete CDS) of NA antigen of human H1N1 influenza virus available in NCBI GenBank (http://www.ncbi.nlm.nih.gov) from 2006 to 2013 in Iran were downloaded and saved in FASTA format for creation of required data bank. All downloaded sequences were edited using DNASIS MAX 3.0 software (Hitachi Pharmacia, Hitachi Software Engineering Company, Yokohama, Japan) to have proper and identical sequences in terms of name and format for comparison and performing of the intended analyses. Totally, 97 sequences from 16 provinces of Iran were downloaded, edited and classified based on the year of isolation. The sequences were aligned based on ClustalW method using BioEdit 7.1 program [23]. The Molecular Evolutionary Genetic Analysis (MEGA 6.0) software (http://www.megasoftware.net) was used to analyze the sequence alignments and the evolutionary distances between the strains were computed with the same software using the maximum composite likelihood method for evaluation and exact comparison of the sequences, determination of conserved and variable regions and similarities and differences [24, 25]. Megaline–Dnastar software (Expert Analysis Software for PC, Inc., USA) was used for construction of phylogenetic trees by UPGMA (unweighted pair group method with arithmetic means) method.

Using the percent similarity/divergence matrix created by BioEdit program in each year, the sequences with the highest and lowest similarity with other sequences of that year were selected as distinct sequences for comparison and analysis. Then evaluation of secondary and tertiary protein structure and prediction/determination of the ligand binding sites, type and number of the ligand(s) of distinct sequences of each year were done using following web-based analysis software: SWISS-MODEL (http://swissmodel.expasy.org) [26], Phyre2 (http://www.sbg.bio.ic.ac.uk/phyre2) [27] and 3DLigandSite (http://www.sbg.bio.ic.ac.uk/3dligandsite) [28].
Fig. 1. Sequence alignment of 2009 influenza virus isolates in Iran.
Fig. 2. Phylogenetic tree of the Neuraminidase (NA) gene by UPGMA method. Evolutionary relationships of individual coding regions from influenza A viruses sampled in Iran during 2009. Length of each branch pair represents the evolutionary distance between sequence pairs. Scale indicates the number of substitution. MegalignTM constructed phylogram (DNAStar, Madison, WI).

Table 1. Percentage similarity/divergence of 2009 isolates. The triple asterisks indicate when rows and columns refer to the same isolate. A diagonal line formed by the triple asterisks divide the table into two triangles, above and under it. The percentage similarity values are indicated in the upper triangle and the percent divergence values are indicated in the lower triangle.
By predicting and analyzing the secondary and tertiary structures in the amino acid sequences of 2009 and comparison of these structures and analysis and comparison of regions and amino acid sequences in the binding ligands and active sites of the protein in molecular interactions between the ligand and the receptor, the differences between these two groups in amino acid sequence analysis (bifurcation of phylogenetic tree) of 2009 isolates became more clear. It was found that in the ligand binding site 1, amino acid positions 83-84-235-411, in two distinct sequences of 2009 were changed from Val-Thr-Asn-Glu with 15 ligands to Val-Lys-Asn-Glu with 17 ligands (Fig. 3).

Since 2010 to 2013, sequences are identical, there is no change in their ligand binding sites and the type and number of the ligands (data not shown). This result shows that the predominant 2009 pandemic strain is also dominant in these years.

In general, studying changes and mutations in conserved positions and regions, especially those located in active/catalytic sites or the areas associated with anti-NA drug resistance, are very important because these regions usually have functional or structural values in the protein sequence and can be used to be developed as targets for the new drugs, the infection control, the treatments and design of new vaccines.

In the next phase of this study, the sequences of different isolates were studied in terms of their position and the number of glycosylation sites at Asn-X-Ser/Thr motif, where X can be any amino acid except proline [29]. These sites play an important role in some features of the protein such as folding, stability, antigenicity and immunogenicity [30]. The sequences of 2006-2008 isolates have 9 conserved glycosylation sites at positions: 44, 58, 63, 70, 88, 146, 235, 434 and 455. The sequences of 2009-2013 isolates have 8 glycosylation sites at: positions 50, 58, 63, 68, 88, 146, 235 and 386 except for some cases in 2009 (creation of glycosylation site at positions 50, 68, 386 and deletions at positions 44, 434 and 455) and 2010 (the creation of glycosylation site at position 42) that some changes were detectable in different positions (Table 2). This implies a change in the number and location of the glycosylation sites of 2009 pandemic strain which became dominant in the next years.

One of the important conserved regions in NA is 150 loop (cavity) which includes residues Gly147 to Asp151 surrounding the active site of NA and functions as a laterally gate for the drug binding [31, 32]. It was found that this region was conserved from 2006 to 2008. In 2009, the mutation occurred in Val149Ile which became dominant from 2010 to 2013. Residues within the active site are highly conserved among all of the NA subtypes, including 8 charged and polar residues (Arg118, Asp151, Arg152, Arg225, Glu277, Arg293, Arg368 and Tyr402). They interact with the substrate directly at the catalytic site. The structure of catalytic site is stabilized and supported by the conserved framework residues (Glu119, Arg156, Trp179, Ser180, Asp/Asn199, Ile223, Glu228, His275, Glu278, Asn295 and Glu425) [33, 34]. It was found that the catalytic site residues were conserved from 2006 to 2013 and the framework residues were also conserved except for some cases in 2009 that H275Y mutation, which is the most common molecular marker of NA drug resistance, has occurred.

Besides monitoring of the naturally occurring changes in NA and its respective genes, monitoring of mutations especially those associated with resistance to the NA inhibitors is necessary. Antigenic changes and mutations in the gene of NA protein often cause resistance to the available anti-influenza drugs in influenza viruses. Zanamivir and Oseltamivir that inhibit the function of NA are used as antiviral drugs for the prevention and treatment of influenza [35]. As mentioned above, H275Y mutation is the most prevalent molecular marker of drug resistance to Oseltamivir that occurs in or near the active site of NA and decrease the sensitivity of NA and virus to this drug [36]. We found that this mutation was occurred in 3 cases in 2009. Other mutations associated with the resistance to anti-NA drugs were also inspected; however none of them was observed (Table 3).

Molecular and phylogenetic analyses were also performed to reveal the relationship of changes of each year’s isolates to the vaccine strains (Table 4). It was found that all changes and the obtained results of each year were in compliance with and were observed in the standard vaccine strain of that year and the seasonal isolates turned out to be closely related to the corresponding vaccine strains. This means that studying and identifying the antigenic changes of the virus in each year can be used for the prediction of changes in future and designing new vaccines or anti-influenza drugs.

Fig. 3. Predicted 3D structure of ligand binding site of 2009 distinct sequences. 3DLigandsite visualization of the protein structure with the residues which form part of the binding site are colored blue. The ligands that form the cluster used for the prediction are also displayed as wireframes.
DISCUSSION

The selection of influenza vaccine strains and the development of new vaccines have been always a demanding task because of the antigenic changes of influenza viruses. Due to continuous and high mutation rate in NA, studying the changes in important sites of NA in the available sequences of different virus strains in a specified time period using conventional bioinformatics methods, can provide useful and important information about the trend of changes for the virologists and the public health authorities. Such insights can help to monitor the emergence of new influenza variants and to detect new antigenic changes occurring in NA protein which are important for updating the vaccine composition.

In this study, by computational investigation of sequences isolated between 2006 to 2013 from Iran and the inspection of the amount of phylogenetic similarity amongst them and by determination of the binding sites and relevant ligands of NA antigen, it was found that the amounts of similarities in sequence alignment analyses in 2006-2008 in one side and in 2010-2013 from the other side were significant while in 2009, a continuous shuffling, leading to different sequences and a new group.

The results of our study are consistent with the results of others in terms of changes in number and location of glycosylation sites, mutations/conservation in important locations such as 150 loop, catalytic sites (NA active sites), framework residues and NA drug resistance were observed in 2009 strains. Therefore, it can be assumed this new group and its relevant mutations became dominant and stable in the following years. The 2009 H1N1 pandemic strain is used as a standard vaccine strain from 2010 onwards and mutations associated with anti-NA drug resistance were observed in 2009 strains.

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In conclusion, our findings suggest that the 2009 H1N1 pandemic strain might continue to mutate in its NA gene and create new dominant strains with the potential to cause pandemics in the future. Therefore, conduction of this study was important to assess the extent of diversity of the circulating strain.

was an important transition year which almost classified the isolates into two groups, namely the 2006-2008 and the 2010-2013 isolates while each group had similar changes/mutations. The 2009 phylogenetic tree was also divided into two distinct branches (due to the occurrence of pandemic) with two groups of isolates. One group was similar to 2006-2008 isolates in terms of amino acid properties and changes/mutations and the other group was similar to 2010-2013 isolates. This shows the importance of changes in the genomic structure of the virus in 2009 which were made possibly during mutations in the genome and likely by the occurrence of frame shifts or gene shuffling, leading to different sequences and a new group.

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strains and to analyze the amino acid changes and the tertiary structures which can change the function of the virus and its virulence or pathogenicity. This also emphasized the importance of continuous monitoring of influenza A H1N1 virus strains (especially the A (H1N1) pdm09 strain) in different geographical areas in order to get better selection of the vaccine strain for the upcoming seasons. To the best of our knowledge, this study was done for the first time in Iran and the obtained results can be potentially useful for the epidemiological studies, as well as for better understanding of the antigenic evolution of H1N1 influenza viruses and the prediction of changes in the future that together can help to select the proper vaccine strains or to design new effective anti-influenza drugs.

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

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