Extraction and purification of the H9N2 virus nucleoprotein: A simple and practical method

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Abstract

Background: Avian Influenza disease annually entails a significant economic loss to the poultry industry around the world. Influenza virus is a polymorphic virus of the orthomyxoviridae family (single-stranded RNA genome), and nucleoprotein (NP) is the structural and internal protein of the virus. The aim of the work was to purify nucleoprotein for further investigations with a simple, low-cost, fast and practical method.

Methods: In this study, H9N2 influenza virus was isolated in specific pathogen-free embryonated chicken eggs by allantoically inoculating 103 to 105 egg-infective doses (EID50) for 9 to 11 days, purified by 10% (W/V) polyethylene glycol (PEG) 6000 with a sucrose gradient of 60% to 30%. The influenza virus proteins were collected and prepared as fractions by preparative electrophoresis. Finally, the purified NP was subjected to sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and Western blot procedures.

Results: The protein analysis with SDS-PAGE and silver nitrate staining indicated that the desired samples contained purified nucleoprotein and lacked other viral proteins. The results of the investigation of lyophilized fractions containing nucleoprotein on the SDS-PAGE revealed the absence of viral RNA in nucleoprotein and its high purity.

Conclusion: According to this study, purified nucleoprotein can be used to produce nucleoprotein vaccines, as well as to study structural, molecular and diagnostic and therapeutic materials.

Keywords: Avian influenza virus, H9N2, Nucleoprotein, Extraction, Purification

Introduction

Avian Influenza is a highly contagious disease that causes symptoms in respiratory, digestive and nervous systems in a wide range of birds and can be transmitted to mammals, including humans (1). Dramatic changes reassortment between human and avian viruses are due to an-

What is “already known” in this topic:
The nucleoprotein H9N2 influenza virus functions as a significant adaptor molecule between the virus and host cell processes. The purified nucleoprotein is used to detect new virus species caused by interbreeding and their resultant disease and to provide a vaccine for avian influenza for prevention and control programs of influenza in the industry.

What this article adds:
The aim of this study was to purify virus NP in order to further project with a simple, low-cost, fast and practical method. In the current study, firstly the H9N2 virus identified and the locations of the viral proteins were determined. Subsequently the nucleoprotein was purified with adequate and accurate information about the viral proteins.
Purification of the H9N2 virus nucleoprotein

nual epidemics and pandemics (2). To date, the most appropriate method for fighting avian influenza is control and eradication. An effective human influenza vaccine development is a critical component to control and prevent an influenza pandemic (3). Influenza virus is an enveloped, single-stranded negative-sense RNA segmented virus (3). Based on the two proteins on the surface of the virus; hemagglutinin (H), and neuraminidase (N), influenza A viruses are divided into 15H and 9N subtypes (4). The proteins play key roles in infecting a host cell. In addition, inside the envelope are segmented single-stranded RNA and nucleoprotein (NP) (5). Avian influenza (AI) H9N2 subtype is generally reported as the primary isolate from turkey flocks in Wisconsin in America in 1966, then the virus has been reported and circulated among various countries (6). Although H9N2 is a low-pathogenic virus, the disease has been observed in all types of poultry in the field (7). Because of the wide range of the virus and the great losses it caused, initially a local vaccination program was implemented, but mass vaccination was quickly authorized (8). The virus has spread to other avian species such as chickens (breeders, broilers, and layers) and ostriches (8), it has been panzootic in Eurasia and is now the most dominant subtype of influenza virus in poultry industry (9). The viruses were classified as low pathogenic on the basis of experimental inoculation of chickens by different routes, but under field conditions, they have proved to cause a severe respiratory syndrome in birds, with heavy reductions in egg production and high mortality (5, 7). The H9N2 outbreak has been reported in the world from 1944 to 1996 in Europe, Africa, and Asia. Subsequently, in 1997, severe H9N2 cases were reported in many Asian countries (10). All viruses with negative-sense RNA genomes encode an ssRNA binding nucleoprotein (NP) (11). The NP is one of the key determinants of species specificity, RNA encapsulation, transcription, replication, and packaging (11). The NPs have interaction with different molecules such as; PB1, PB2, M1, Importin α, F-actin, CRM1/exportin-1 and BAT1/UAP56 (11). The NP Functions as a key adaptor molecule between the virus and host cell processes (9, 12). The purified nucleoprotein is used to detect new virus species caused by interbreeding and their resultant disease and to provide a vaccine for avian influenza for prevention and control programs of influenza in the industry (13). The aim of this study was to purify virus NP in order to further project with a simple, low-cost, fast and practical method.

Methods

Viral propagation

In this experimental study, viral propagation was performed by the method described (14). Briefly, the isolates of avian influenza A/Chicken/Iran/259/2014/H9N2 were propagated in specific-pathogen-free chicken embryonated eggs by allantoically inoculating 10³ to 10⁵ egg-infective doses (EID₅₀) for 9 to 11 days. Eggs were incubated at 37°C with relative humidity of 55% and were candled daily for 7 days to embryo viability. Allantoic fluid from embryos that died after 24 h was collected aseptically and tested for hemagglutinating virus by the hemagglutination (HA) test. If no deaths occurred in the eggs after 6 days, all of the eggs were opened aseptically. The allantoic fluids were pooled and inoculated into 5 more embryonated eggs. If no deaths occurred after 7 days in the eggs of the second passage, all the eggs were opened, and the allantoic fluid of each egg was tested for hemagglutinating activity. If deaths occurred in the eggs and the material had a positive HA test, samples were considered positive, and this result was confirmed with the hemagglutinating inhibition (HI) test. Allantoic fluids from embryos showing early mortality after inoculation were tested for bacterial contamination according to routine procedures, and subsequently the infected fluids which contained between 32×10⁵ EID 50 of virus per ml were harvested and pooled. Reference stocks were lyophilized and working stocks were stored in ampoules at -20°C (15). However, HA and HI tests were performed by standard methods (15).

Extraction and Purification of nucleoprotein

The mixture was incubated for 1 hour at 4°C and clarified by centrifugation at a speed of 3500 rpm and 4°C for 15 minutes. The supernatant was collected, and PEG 5% (w/v) was added to it. Then, clarified by centrifugation at a speed of 9000 rpm and 4°C for 60 minutes, precipitate in TNB buffer was created as a homogenous suspension. Finally, a homogenous suspension was prepared for dialysis to remove PEG from the suspension. The HA test was performed on the sample after dialysis. Then, the dialyzed fluid was ultra-centrifuged at a speed of 20000 rpm and 4°C for 2 h. The resulting precipitate was converted into homogeneous suspension in 100 ml of TNE buffer. The HA test was also carried out at this stage. The purification of virus was performed by sucrose density-gradient centrifugation method (16). Finally, the resulting virus samples were lyophilized.

Protein Assay

For this purpose, Bradford method was performed to determine the protein concentration of the sample according to the previously described method with some modifications (17). In this method, duplicate volumes of bovine serum albumin (BSA) (10-100 mg/ml) was used to prepare the calibration graph. The basic Laemmli SDS-PAGE procedure was used to separate the protein of interest (18). Afterward, one of the gels was stained with Coomassie blue G-250 (Merck, Germany) and the other gel with silver nitrate. Also, native and functional biological molecules were purified by preparative electrophoresis (model 491 Prep Cell, Bio-Rad, CA), using reduced and non-reduced methods (19). Finally, to determine protein fractions the optical density was read at 280 nm using a spectrophotometer. The reduced 12% SDS-PAGE was used to analyze the integrity of the purified influenza virus proteins and the detection of fractions containing nucleoprotein. In brief, 200 μl of each protein fraction [with a high absorption] was mixed with 30 μl of 2-mercaptoethanol and incubated for 4 h at 37°C. The protein fractions [with high absorption] were dissolved in non-reducing SDS-PAGE loading buffer (without 2-mercaptoethanol) and stained with silver nitrate (20). The protein concentration was assessed with the Bradford method. Finally, in order to confirm H9N2 presence, the purified

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sample was loaded and resolved by SDS-PAGE and proteins were transported to polyvinylidene difluoride (PVDF) membranes. Membrane blocking was done using 5% bovine serum albumin (BSA), overnight. Primary specific antibody versus H9N2 was used at a 1:1000 dilution, and secondary antibody was used at a 1:10000 dilution (HRP-conjugated). Immunoreactive bands visualized with chemiluminescence HRP substrate (Abcam).

Results

Protein assay before SDS-PAGE
A protein assay was performed for influenza virus, and at this stage, BSA was used as a standard in volumes of 100, 300, 500 and 1000 mg/ml, and the results indicated that the volume of the viral solution was appropriate. According to Table 1, the highest and lowest amounts of protein were the sample F act 3 and the sample F act 5 with 387.07 mg/ml and 212.5 mg/ml, respectively.

SDS-PAGE after protein assay
Gels were prepared with concentrations of 2.5, 5 and 10 μg of each sample. According to our results (Fig. 1), the molecular weight of the nucleoprotein was estimated between 45 and 66.2 kDa.

Preparative electrophoresis results
The absorption of the fractions was obtained from the purified nucleoprotein at 280 nm (UV). Graphs 1 and 2 were plotted from the 121 fractions of the reduced and, 123 fractions of the non-reduced methods. Our results showed that 26 fractions from the reduced and 42 fractions from the non-reduced methods had a positive absorption, and these fractions contained viral NP.

SDS-PAGE and Western blotting after protein purification
SDS-PAGE was performed for the presence of NP after purification with preparative electrophoresis method (Figs. 2 and 3). According to the molecular weight of NP, fractions 2 and 3 (reduced purification) contained viral NP, but in fractions with non-reduced purification, no band was detected, and no viral protein was isolated. The NP band (H9N2) was verified again by immunoblotting assay (Fig. 4).

Discussion
The purpose of this work was to prepare and purify the influenza virus nucleoprotein in a simple, low-cost, applied and rapid method. In the current study, firstly the H9N2 virus identified and the locations of the viral proteins were determined. Subsequently the nucleoprotein was purified with adequate and accurate information about the viral proteins. The use of a live virus for propagation is necessary, but epidemiological hygiene needs to be considered, and this issue is more important in influenza viruses due to their genetic rearrangement and transmission potential to humans. However, the advantages of using the inactive virus are due to the lack of infectious potential and long-term maintenance and the absence of contamination with other microorganisms. The virus from dialysis was lyophilized to facilitate further protein purifications. By SDS-PAGE analyses, the virus protein bands were examined, and it was determined that a nucleoprotein with a molecular weight ~ 54 kDa is clearly identifiable [between bands of 45 to 65 kDa of ladder], and two bands (top and down), which are related to HA protein, can be identified and separated. However, to ensure that the NP protein can be observed in low amounts of protein and the diluted sample, gels are stained with silver nitrate, which is about 10 to 100 times more accurate than the Coomassie blue. For purity and the presence of the desired protein, we examined the molecular weight and position of the nucleoprotein in terms of movement and detachment in the polyacrylamide gel; samples were run on a preparative electrophoresis machine. The Prep Cell 491 has a special conctructional complexity and requires the

Table 1. Results of Protein assay

<table>
<thead>
<tr>
<th>Sample name</th>
<th>OD 595nm</th>
<th>Protein mg/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>St 100</td>
<td>0.08</td>
<td>100</td>
</tr>
<tr>
<td>St 300</td>
<td>0.31</td>
<td>300</td>
</tr>
<tr>
<td>St 500</td>
<td>0.53</td>
<td>500</td>
</tr>
<tr>
<td>St 1000</td>
<td>0.97</td>
<td>1000</td>
</tr>
<tr>
<td>F act 1</td>
<td>0.22</td>
<td>212.9</td>
</tr>
<tr>
<td>F act 2</td>
<td>0.32</td>
<td>309.67</td>
</tr>
<tr>
<td>F act 3</td>
<td>0.40</td>
<td>387.07</td>
</tr>
<tr>
<td>F act 4</td>
<td>0.27</td>
<td>261.29</td>
</tr>
<tr>
<td>F act 5</td>
<td>0.17</td>
<td>212.5</td>
</tr>
</tbody>
</table>

Fig. 1. SDS PAGE gel for absorption (reduction method) protein fractions stained silver nitrate
1. F act 4 (10mg), 2. F act 4 (5mg), 3. F act 4 (2.5mg), 4. F act 5 (10mg), 5. F act 5 (5mg), 6. F act 5 (2.5mg)

Fig. 2. SDS PAGE gel for adsorption (reduction method) protein fractions
1. Prestaining Marker, 2. Virus sample before purification, 3. Absorption fraction containing nucleoprotein, 4. Absorption fraction containing nucleoprotein
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Graph 1. Absorption of 121 fractions from (reduced method) at 280 nm

Graph 2. Absorption of 123 fractions from (non-reduced method) at 280 nm

precision and observance of the basic principles of assembling its parts. The components should be tightly assembled and the gel preparation, spraying, fractionation and droplet adjustment steps should be done correctly, and since the application and operation of the device is long, it is necessary to schedule proper testing. The accuracy of the purification of the influenza virus proteins is very important due to the small size and the number of proteins and types of proteins. By adjusting the fractionation device, the volume of 8 ml and duration of 20 minutes for each fraction, this step lasted 48 h. Initially, the fractions contain a protein-free stain which should be discarded. After preparing the fractions and determining their absorbance at 280 nm, the precise and pure presence of the nucleoprotein in fractions 2 and 3 was observed after complete removal of color, and this result was confirmed by SDS-PAGE, but in the non-reducing procedure, no purification was performed, and no band was found in SDS-PAGE. To increase the volume of pure nucleoprotein fractions, fractions containing the nucleoprotein were lyophilized. After lyophilization of the fractions containing the nucleoprotein, protein assay, SDS-PAGE and staining with silver nitrate were performed again. The significant points about the electrophoresis of the influenza virus are the importance of the preparation stages and purification of the virus and the electrophoretic conditions affecting the pattern of electrophoresis of viral proteins. So that with the slightest change, it is possible to observe a change in the electrophoretic pattern of the virus. The lack of similarity of the electrophoretic pattern of various influenza viruses can be attributed to this cause. To ensure the absence of the influenza virus genome in purified nucleoprotein, (because the nucleoprotein and genome of the virus are in the structure of the virus and the nucleoprotein is the viral genome coverage, and together with the RNA, the RNP forms), the resulting samples were electrophoresed.

To check the accuracy of the results, the influenza virus before and after lyophilization, and a purified nucleoprotein after lyophilization were tested. The results of electrophoresis indicated no genomic band in the nucleoprotein sample. Thus, during testing, it has been proven that with the help of the Prep Cell 491, we will be able to completely purify the H9N2 avian influenza virus and save time and cost. Purified nucleoprotein can be used to provide nucleoprotein anti-sera against influenza viruses in future researches. Significant points in this study are the high speed and precision of the test results, which can be expressed as a superior advantage over the success of the research. Studies have been conducted on nucleoprotein and different methods used for purification of nucleoprotein. Harmon et al., investigated an immunoassay for serological diagnosis of influenza type using a recombinant DNA producing nucleoprotein antigen and monoclonal human antibody IgG (21). They used immunoaffinity chromatography method for NP purification. This method is time-consuming and

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costly, and although the appropriate amount of nucleoprotein was purified, supplementary experiments and a large number of viral samples are required. In the present study, with the least number of the viral samples, nucleoprotein can be obtained. Li et al., (2014), isolated and purified the NP by SDS-PAGE for NP purification and then mass spectrometry was performed (22). In the SDS-PAGE electrophoresis method, viral proteins are isolated but not completely purified, and when cutting from the gel, there is a possibility of error. On the other hand, in this method, the accuracy and skill of the researcher are of great importance, and the cost of mass spectrometry is significant for confirmation of the protein.

Conclusion

Considering the results of our study, the purified nucleoprotein was obtained very dynamically, with high speed, precision, and purity. Purified nucleoprotein can be used to produce nucleoprotein vaccines, as well as to study structural, molecular and diagnostic and therapeutic materials.

Acknowledgments

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Conflict of Interests

The authors declare that they have no competing interests.

References