Expression of Insoluble Influenza Neuraminidase Type 1 (NA1) Protein in Tobacco

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ABSTRACT

The avian influenza virus, particularly H5N1 strain, is highly virulent to poultry and mankind. Several expression systems, like yeast, baculovirus and mammalian cells, have been adopted to produce vaccine candidate for this lethal disease. The present research aimed at developing a recombinant vaccine candidate, neuraminidase type 1 (NA1), for the Malaysia isolate of H5N1 in Nicotiana benthamiana. The NA1 gene was fused directly in-frame in cowpea mosaic virus (CPMV)-based pEAQ-HT vector with C-terminal polyhistidine-tag incorporated to ease the subsequent purification step. The expression of the NA1 gene in tobacco was confirmed at RNA and protein levels at 6 days post-infiltration (Dpi). From the insoluble fraction of the protein, a recombinant glycosylated NA1 protein with a molecular weight of ~56 kDa was immunogenically detected by a specific anti-NA polyclonal antibody. We report for the first time the insolubility of the plant-made NA1 protein where a native sequence was used for its expression. This study signifies the necessity of the use of optimised sequences for expression work and provides great opportunity for the exploration of plant-manufactured NA1 protein as vaccine candidate.

Keywords: Avian Influenza, H5N1, Neuraminidase, Plant virus-based expression system, Recombinant protein

INTRODUCTION

Highly pathogenic avian influenza (AI), H5N1, in poultry has fuelled intense media coverage and health concerns due to the severe outbreaks in Asia, Africa and Europe. This deadly pandemic is still showing a great potential of spreading. The development of vaccine candidates for AI is utmost crucial and could serve as the best strategy for managing the disease. Two of the currently licensed vaccines against influenza are conventional inactivated virus vaccine (CIV) and live-attenuated vaccine (LAV) with an objective of complete infection inhibition. These vaccines are dominated by the anti-haemagglutinin (HA) immune response, and hence, the effectiveness is restricted to virus strains with closely matched HA. They are susceptible to protect failures in cases where antigenic drift or shift occurred on HA [1]. Therefore, antiviral drugs like Tamiflu® (oseltamivir) and Relenza® (zanamivir) have been used widely to treat patients with early infection [2]. These drugs inhibit neuraminidase (NA) main function, which is to facilitate the release of the progeny virions from infected cells. By this, the virus spread is restricted and disease onset is suppressed. However, virus strain with NA-inhibitor resistance has been observed and identified [3]. Therefore, it seems the vaccine candidate targeting NA is still handy in this case.

For a better protection against heterovariant influenza viral infection, vaccines containing equal levels of immunogenic HA and NA antigens are required. Inclusion of NA offers a better prospect in this case due to its slower rate of antigenic evolution [1]. Instead of infection inhibition, NA presents an alternative strategy called infection-permissive immunization. Previous studies have described that CIV and LAV supplemented with purified NA protein or recombinant NA protein yielded equivalent level of both anti-HA and anti-NA antibody titres [4,5]. The viral replication was significantly
reduced in response to heterotypic virus challenges. Recombinant NA protein is favourable due to its production simplicity as compared to the laborious purification process of native NA protein from influenza virion. Plant system has been used to produce several recombinant proteins due to its great advantages over other competing systems. Plant systems offer opportunities to produce vaccines at low cost, it is easy to scale-up, and has low risk of contamination with animal pathogens. Plant-based vaccine also provides a new approach of oral delivery [6], provided these immunogenic peptides are expressed in an edible crop plant, such as corn [7]. Hence, downstream purification cost or injection-related hazard can be eliminated. In this study, we aimed to engineer a recombinant NA1 protein from a Malaysian isolate and express it in tobacco using a viral vector. We describe the construction of the recombinant vector and the expression of NA1 gene at both RNA and protein levels.

MATERIALS AND METHODS

NA1 Sequence and Plant Viral Vectors

The NA1 gene of a local isolate Avian Influenza (A) virus strain A/chicken/Malaysia/5744/2004(H5N1) was kindly supplied by Professor Abdul Rahman Omar from the Institute of Bioscience, Universiti Putra Malaysia, Malaysia. Native NA1 sequence was used in the construction of recombinant vector. CPMV-based plant viral expression vectors, pEAQ-HT and pEAQ-HT-GFP were a kind gift from Professor George Lomonossoff (John Innes Centre, UK).

Construction of Recombinant Vector and Its Characterisation

Full-length of the neuraminidase type 1 (NA1) (450 amino acids) was used for PCR-based cloning. A set of primers incorporated with restriction sites, namely AgeI (Forward NeuF-AgeI: GCGACCGGTATGAATCCAAATAGAAAGATTA) and SmaI (Reverse NeuR-SmaI: GTCCCGGGCTGTCAATGTGAATGCTG) was used for PCR amplification. PCR reaction was prepared by adding 1 X of PCR buffer S (160 mM (NH4)2SO4, 500 mM Tris-HCl, 17.5 mM MgCl2 and 0.1 % Triton X-100), 0.4 mM of dNTP, 0.5 μM of each primer, 20 ng template, 1 U of Taq polymerase and made up to 20 μl with sterile deionised water. The tubes were heated at 94 °C for 4 min and then subjected to twenty five amplification cycles with 45 sec at 94 °C for denaturation, 45 sec at 50 °C for annealing and 1 min at 72 °C for extension. Another 5 min at 72 °C was added as the final extension step. The PCR products were then electrophoresed at 80 Volt for 40 min in 0.8 % (w/v) of agarose gel. The PCR products were purified using MEGAquick-spin™ PCR and Agarose gel DNA extraction system (Intron Biotechnology, Korea) according to manufacturer’s instructions. For ligation purpose, the NA1 gene and pEAQ-HT vector were digested by the same restriction enzymes (RE), AgeI and SmaI, to form sticky ends at both cutting sides. The resulting recombinant construct was named as pEAQ-HT::NA1 (Figure 1) and verified by RE digestion. AgeI was used to linearize the recombinant vector. Reaction mixture was prepared by adding 1 X of buffer V2 buffer (10 mM Tris-HCl pH 7.5 at 30 °C, 10 mM MgCl2, 50 mM NaCl and 0.1 mg/ml BSA), 2 U of AgeI (Vivantis, Malaysia), 100 ng of pEAQ-HT::NA1 plasmid and made up to 20 μl with molecular grade distilled water. AgeI and XbaI were used to release the NA1 gene from the pEAQ-HT vector. Reaction mixture was prepared by adding 1 X of V2 buffer, 2 U of AgeI (Vivantis, Malaysia), 2 U of XbaI (Promega, USA) and 200 ng of pEAQ-HT::NA1 vector and made up to 20 μl with molecular grade distilled water. The reaction mixtures were incubated at 37 °C for 2 h. After that, the digested product was electrophoresed in a 1.0 % (w/v) agarose gel. The plasmids were subsequently sent for nucleotide sequencing (Macrogen Inc., Korea).

Agroinfiltration Procedure

Empty pEAQ-HT, recombinant pEAQ-HT::NA1 and pEAQ-HT-GFP were transformed into A. tumefaciens LBA4404 using electroporation method. The resultant transformants were thereafter named as Ag/pEAQ-HT, Ag/pEAQ-HT::NA1 and Ag/pEAQ-HT-GFP. The cultures were grown to stable phase in Luria-Bertani medium supplemented with 50 μg/ml rifampicin and kanamycin at 28 °C in shaking condition of 200 rpm. The cells were collected by centrifugation at 6,000 rpm for 10 min at 4 °C and resuspended in infiltration buffer (10 mM 2-(N-morpholino)ethanesulphonic acid (MES), pH 5.5, 10 mM MgCl2, 0.1 mM acetylsyringone) to an OD600 of 0.3-0.4. After 2 h of incubation period at room temperature, the suspensions were infiltrated into fully expanded N. benthamiana leaves using a syringe without a needle. After infiltration, the plant was further grown under
mRNA Assessment of pEAQ::NA1 Infiltrated Plant Samples

Leaf samples from both pEAQ-HT::NA1-infiltrated and mock-infiltrated plants were harvested on six days post-infiltration (Dpi) for total RNA extraction procedure by using TRIzol® reagent (Invitrogen, USA) following manufacturer’s protocol. Reverse transcription (RT) was conducted on the RNAs extracted by using Viva 2-steps RT-PCR kit (Vivantis, Malaysia). First-strand cDNA synthesis reaction mixture was prepared by adding 2 µg of total RNA, 2 µM gene-specific primer (NeuR-SmaI), 0.5 mM dNTP mix and made up to 10 µl with DEPC-treated water. The mixture was incubated at 65 °C for 5 min and then placed on ice for 2 min. A 10 µl mixture of 1 X RT buffer (500 mM Tris-HCl (pH 8.3 at 25 ºC), 750 mM KCl, 30 mM MgCl2 and 100 mM DTT) and 100 U M-MuLV Reverse Transcriptase was added to each individual tube. The mixture was incubated at 42 ºC for 60 min. Then, the reaction was terminated at 85 ºC for 5 min and chilled on ice. PCR reaction mixture and its program were similar to previous section as described. RNA extracted was also subjected to PCR without first strand cDNA synthesis as a negative control for tracking of DNA contamination. The PCR products (20 µl) were then electrophoresed at 80 Volt for 40 min in 1.0 % (w/v) agarose gel.

Protein Extraction and SDS-PAGE

Electrophoresis

Proteins were extracted from the Ag/pEAQ-HT::GFP infiltrated leaves after 3, 6, 9 and 12 Dpi and Ag/pEAQ-HT::NA1 infiltrated leaves after 6 Dpi. Total soluble proteins were extracted according to protocol as described by [8]. Inoculated leaf tissues were frozen in liquid nitrogen and ground to powder using mortar and pestle. Then, the samples were suspended in 2 volumes of soluble protein buffer (50 mM Tris-HCl pH 7.25, 150 mM NaCl, 0.1 % (v/v) Triton X-100, 2 mM DTT and 1/100 protease inhibitor cocktail) and centrifuged at 13,000 rpm for 10 min at 4 ºC. Supernatants were collected and stored at 4 ºC. The pellets of Ag/pEAQ-HT::NA1-infiltrated samples were kept for insoluble proteins extraction.

One millilitre of pre-cooled methanol (with 1/100 protease inhibitor cocktail) was added into each tube. The tubes were vortexed for 30 sec, incubated at -20ºC for 5 min and centrifuged for 5 min at 13,000 rpm at 4ºC. The supernatant was removed with a pipettor and the washing steps were repeated once for methanol and once for acetone. The pellets were brief-dried at room temperature and re-suspended in 1 volume of insoluble protein buffer (7 M urea, 2 M thiourea, 40 mM Tris HCl pH 7.5, 10 mM DTT and 1/100 protease inhibitor cocktail). The tubes were incubated at 23 ºC for 15 min under gentle agitation. Then, the samples were centrifuged at 13,000 rpm for 30 min at 18 ºC and supernatants
were collected. The proteins extracted were then electrophoresed at 120 Volt for 90 min in 12 % SDS-PAGE gel.

**Protein Purification by Immobilized Metal Affinity Chromatography (IMAC) Resin**

Profinity Ni-charged IMAC resin (BioRad, USA) was suspended thoroughly and 200 µl was transferred to Micro Bio-spin column™ (BioRad, USA). Storage solution was removed by centrifugation (1,000 rpm) for 10 sec and the column was washed by 5 volumes of distilled water. The resin was pre-equilibrated by 5 volumes of binding buffer (50 mM NaH₂PO₄, 300 mM NaCl, 5 mM imidazole and 8 M urea, pH 8.0) prior to adding 500 µl of Ag/pEAQ-HT::NA1 infiltrated insoluble protein sample. The sample and resin were mixed by pipetting for 5 times. The unbound protein was removed by centrifugation and the column was washed by 5 volumes of washing buffer (50 mM NaH₂PO₄, 300 mM NaCl, 20 mM imidazole and 8 M urea, pH 8.0). The column was transferred into a new microcentrifuge tube and protein was eluted by 150 µl of elution buffer (50 mM NaH₂PO₄, 300 mM NaCl, 250 mM imidazole and 8 M urea, pH 8.0).

**Western Blotting Analysis**

Western blotting analysis was carried out on total crude protein extracts and IMAC-purified protein extracts. The extracts separated on 12 % SDS-PAGE gel were then transferred onto nitrocellulose membrane (Invitrogen, USA). Primary rabbit polyclonal antibodies against GFP (Abcam, UK) (1:10,000 dilution), primary rabbit polyclonal IgG antibody rose against avian influenza A neuraminidase (Abcam, UK) (1:10,000 dilution) and horseradish peroxidase-conjugated anti-rabbit IgG as the secondary antibody (Zymed, USA) (1:10,000 dilution) were used in the detection procedure. The membrane was blocked with 5 % (w/v) non-fat dried milk in 1 X PBS-T buffer (137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 2 mM KH₂PO₄ and 0.05 % (v/v) Tween-20) together with primary antibody for 2 h at room temperature. Then, the membrane was washed three times with PBS-T buffer and incubated with secondary antibody for 1.5 h at room temperature. After washing three times in PBS-T buffer, the membrane blotted with GFP protein was incubated with 3 ml of 3,3′,5,5′-Tetramethylbenzidine (TMB) substrate until development of signal at desired level. Image was captured using GS800 calibrated imaging Densitometer (BioRad, USA). For membrane blotted with NA1 protein, SuperSignal West Dura Chemiluminescent substrate solution (Pierce, USA) was spread evenly on the membrane for 5 min and image was captured by CCD camera (Alpha-Innotech Corporation).

## RESULTS AND DISCUSSION

**Results**

**Construction of pEAQ-HT::NA1 and Its Characterisation**

The full-length NA1 gene of Malaysian strain H5N1 with a molecular size of 1.4 kbp was isolated by PCR and its sequence was confirmed by nucleotide sequencing. The NA1 gene was inserted successfully into pEAQ-HT vector via AgeI and SmaI sites with His-tag at C-terminal. Following the gene cloning, the recombinant vector, pEAQ-HT::NA1 was successfully characterized and identified by PCR and RE digestion. The RE profile of different enzymatic digestion verifying the desired pEAQ-HT::NA1 is shown in Figure 2. AgeI produced the linearized pEAQ-HT::NA1 (~12 kbp) and NA1 gene with His-tag was cleaved out from the pEAQ-HT vector when digested with AgeI and Xhol. The fragments obtained were approximately 1.4 kbp and 10 kbp in molecular sizes as expected for NA1 gene and pEAQ-HT vector, respectively. Following cloning process, nucleotide sequencing analysis showed an in-frame fusion of the NA1 gene with His-tag of pEAQ-HT vector.

![Figure 2. Restriction profile of recombinant pEAQ-HT::NA1. Lane M: Vivantis 1 kb ladder; lane 1 AgeI-digested; lane 2 AgeI- and Xhol-digested. Approximately 1.4 kbp of NA1 fragment is observed after AgeI and Xhol digestion.](image-url)

**Physical Observation of Plants Post-agroinfiltration**

When infiltration was conducted with low OD600 (0.3-0.4) of *A. tumefaciens*, at 6 Dpi,
mock-infiltrated (Figure 3a) and Ag/pEAQ-HT-GFP-infiltrated (Figure 3c) plant leaves showed minimum symptoms, while the Ag/pEAQ-HT::NA1-infiltrated zones bleached (Figure 3b).

Expression of NA1 Gene in N. benthamiana

Following a kinetic expression assessment of pEAQ-HT-GFP on N. benthamiana leaves at protein level, specific and distinct GFP protein bands (~27 kDa) were detected in soluble fraction of proteins at 3, 6, 9, 12 Dpi from Ag/pEAQ-HT-GFP-infiltrated leaves using primary rabbit polyclonal anti-GFP antibody (Figure 4). The expression reached the maximal level within 6 to 9 Dpi and diminished slowly from day 12. No band was detected for mock-infiltrated protein samples.

Expression of NA1 Gene in N. benthamiana at RNA and Protein Levels

The intact total RNA of infiltrated tobacco leaves was successfully extracted and subjected to RT-PCR amplification. Using the specific primers (NeuF-AgeI and NeuR-Smal), a specific and district band (~1.4 kbp) of the full-length NA1 was successfully amplified from cDNA sample of Ag/pEAQ-HT::NA1-infiltrated plant on 6 Dpi as shown in Figure 5 (Lane 2). In contrast, no amplification was obtained from cDNA sample of mock-infiltrated plant (Figure 5, Lane 1) and RNA sample of Ag/pEAQ-HT::NA1-infiltrated plant (Figure 5, Lane 3).

Using the protein extraction method as described above, the insoluble fraction of protein containing the NA1 protein was recognized by polyclonal anti-NA antibody in Western blotting analysis. In contrast, negative outcome was obtained from the detection of recombinant NA1 protein in soluble fraction. The insoluble NA1 protein band with a molecular size of ~56 kDa was detected in a sample of Ag/pEAQ-HT::NA1-infiltrated plant (Figure 6, Lane 2) but not in that of mock-infiltrated plant (Figure 6, Lane 1). Ni-charged IMAC resin purification was done on the crude protein extracted and similar band size was detected (Figure 6, Lane 4) with unspecific background reduced to minimum.

Discussion

The present study was aimed at developing a recombinant vaccine candidate, NA1, for the Malaysian isolate of AI virus strain H5N1 in a plant system, tobacco. This was commenced by inserting the NA1 gene into CPMV-based plant expression vector, pEAQ-HT, to obtain recombinant pEAQ-HT::NA1. The expression profile of NA1 protein in this particular expression system was firstly determined by using a reporter, GFP. Based on the period attaining the most optimal expression level of
GFP, the NA1 expression in tobacco was then studied at both RNA and protein levels.

In the present study, a relatively low OD600 (0.3-0.4) was used for the delivery of recombinant vectors. This was considered necessary given the hypersensitive response of plant leaves to A. tumefaciens infection as reported [9, 10]. This is in agreement with several reports in which A. tumefaciens OD600 used ranged from 0.2 to 0.5 [11, 12, 13]. There are reports, however, which have clearly showed the use of higher A. tumefaciens density with OD600 range of 1.0-1.2 for plant infiltration [14, 15, 16]. The effects of A. tumefaciens density have in fact been well-studied for stable Agrobacterium-mediated transformation system in term of expression efficiency and plant tissues hypersensitivity response [9, 17], but not the detail mechanism. For agroinfiltration transient approach, A. tumefaciens density has been reported as an independent factor for GFP expression [18]. N. tabacum responses to this plant pathogen had been studied [10] and three of the major responses recorded from the infiltrated zones were the induction of pathogenesis-related gene expression, chlorosis and inhibition of leaf expansion. In addition, Gilis and friends [19] also suggested that plant leaves responded differently to different constructs targeting their recombinant proteins in different cell compartments.

Based on the expression profile of GFP, protein expression levels at 6 and 9 Dpi were comparable. Hence, we decided to harvest and analyse the infiltrated plants at 6 Dpi for NA1 expression. This expression profile was in agreement with the previous research [13, 18, 20]. Sheludko and colleagues [20] found that GFP accumulation achieved the maximal level at 3 Dpi and remained stable up to 8 Dpi. Similar tendency was mentioned by another report, which explained that within 4-5 Dpi, the intensity of GUS staining reached the peak level and decreased gradually after that time [13]. Phenomenon of the decreasing expression level can be explained by triggering of RNA silencing mechanism which stops the foreign gene expression [21, 22].

The overall results of RT-PCR and Western blotting analyses confirmed the successful transcription and translation of NA1 gene in infiltrated leaf sample. In order to eliminate the possibility of plasmid DNA contamination during extraction, RNA sample without RT step had been recruited in PCR amplification procedure. Based on the results obtained, the generation of NA1 mRNA in infiltrated leaf sample was proven and the production of NA1 protein was expected. Following the detection of ~56 kDa protein from Ag/pEAQ-HT::NA1-infiltrated leaves, but not in that of mock-infiltrated leaves, we herewith claimed that the plant-based full-length NA1 protein had been produced successfully in N. benthamiana. This indicates the native NA1 gene without sequence modification can be expressed in tobacco leaves using CPMV system, but unlikely in the insoluble state. The expected size of full-length native NA1 (450 amino acids) is ~49 kDa, and thus the observed band of ~56 kDa protein signified a glycosylated NA1. Similar size of recombinant NA was obtained by Schmidt and colleagues [23] who used baculovirus system. In their study, with the use of mass spectrometry, MALDI-ToF, they estimated the molecular weight of the recombinant NA protein fragments to be 54.6 kDa and 58.5 kDa. On the other hand, Yongkiettrakul and colleagues [24] reported that, in yeast system, the predominant hyperglycosylated recombinant NA has a size of ~72 kDa with ~60 kDa protein as an intermediate product of glycosylation.

In Western blotting analysis, unspecific background generated from insoluble fraction of protein is most likely due to the low stringent immunogenicity of polyclonal antibody which corresponds only to 15 amino acids at the C-terminal of NA. To minimise this predicament, the cloning method of NA1 gene into pEAQ-HT vector has allowed the generation of C-terminal His-tag to facilitate the purification of recombinant proteins by Ni-charged IMAC resin.
This purification step of the total insoluble protein had increased the specificity of detection by which the similar band size of ~56 kDa (Figure 6, Lane 4) was resulted with unspecific background reduced to minimum. IMAC is a widely used purification technique for recombinant His-tagged proteins. IMAC resin comprises of iminodiacetic acid charged with transition metals such as Cu\(^{2+}\), Ni\(^{2+}\), Co\(^{2+}\) or Zn\(^{2+}\) and offers a high selectivity of binding to His-tagged proteins [25]. Mohanty and Miener [26] reported His-tag length plays a significant role in relation to protein yield, but its insertion position does not. However, for native protein with signal sequence, in general, His-tag in N-terminal is impractical as it will be cleaved off after transporting the protein to its targeted location. A few related studies have revealed its feasibility in purifying recombinant proteins [27, 28] which can be further employed in subsequent investigations such as immunogenicity assessment. Nevertheless, this purification step in this case has further enhanced the identity of plant-made NA1 protein.

Expression of influenza NA has been well studied in mammalian cell [29], yeast [24] and baculovirus systems [4, 5, 23]. In plant, different types of AI NA were expressed in N. benthamiana, i.e. NA type 2 from influenza A/Wyoming/3/03 (H3N2) [30], NA1 from influenza A/Hatay/2004 (H5N1) [31] and A/Vietnam/04 (H5N1), reported as personal communication in a review article by Musiychuk et al. [32]. These three studies reported the expression of soluble form of recombinant NA protein. In the present study we obtained insoluble recombinant NA1, apparently contradicting these reports. The NA1 sequence that we used was native, not optimised based on plant codon usage. The differences in synonymous codon usage between heterologous expression system and the natural host were believed to be significant. Codon-usage variations among different species are frequently related to the failures in heterologous recombinant protein expression, such as lack of expression and truncated or non-functional proteins production [33]. However, in present study, insolubility of recombinant protein was discovered from native sequence expression, which is seldom reported in plant system. Improper folding and protein insolubility could be due to biased codon usage as translational pause occurs in the presence of slow codons [34].

Many studies have shown that optimisation or harmonisation of codon usage according to expression host increased protein yield considerably [35, 36, 37, 38]. Among these, report by Angov’s group [35] was the most impressive as the expression of sequence harmonised genes exceeded that of native genes by 4- to 100-folds. Another potential protective surface glycoprotein of AI virus, HA, has also been well investigated in heterologous expression studies. HA type 3 and 5 were expressed successfully in tobacco either by transient [28, 30, 39, 40, 41] or stable [41] transformations. Most of the studies reported gene sequence optimisation before subjection to heterologous expression in plant system. The antigenicity of these plant-produced HAs has been confirmed in animal models [28, 39, 40]. Apparently, gene sequence optimisation is crucial for expressing foreign genes in plant system.

However, codon optimisation is not always obligatory for some cases. There are a few exceptional cases in which codon optimisation based on host system does not necessarily give positive effect on expression level. Generally, a higher codon adaptation index (CAI) is generated by codon optimised sequence and protein expression level is believed to be high. However, the A+T content are consequently increases with the increase of CAI for a foreign gene. This might contribute to the instability of mRNA and hence reduces protein expression level [42, 43]. This is supported by Laguia-Becher et al. [44], who demonstrated that the higher the A+T content in codon optimised gene, the lower the expression obtained. The effect of codon optimisation on Toxoplasma gondii surface antigen 1 (SAG1) gene was studied in tobacco system. SAG1 sequence was optimised based on plant codon usage, CAI value and the A+T content were both increased in this case. Results showed that the protein expression level was significantly lowered for optimised variant as compared to native gene. However, advance studies should be carried out in order to comprehend further. Besides, Maclean and colleagues [15] found that human codon-optimised human papillomavirus type 16 (HPV-16) L1 gene resulted to the highest accumulation of the protein in tobacco system as compared to plant codon-optimised and native sequences. In this report, they demonstrated that the plant codon-optimised sequence did not generate detectable protein while native sequence caused the second highest accumulation of the L1 protein. In this case, the A+T content cannot be responsible as native sequence was holding a higher value. The reason behind the poor expression of plant codon-optimised gene
remains unknown and requires further investigations.

Therefore, the native sequence of NA1 was applied in present study instead of codon optimised gene, as its outcome was thought to be unpredictable and worthwhile to be studied. The findings of the present study are the first of its kind to report the insoluble state of expression of a native NA1 sequence in tobacco system. Nonetheless, further modification on the insoluble NA1 is absolutely necessary in order to make the downstream assays such as protein activity and immunogenicity possible. For instance, there are various in vitro refolding processes reported to be effective in renaturing the insoluble recombinant proteins into bioactive form [45, 46]. This is commonly done for *E. coli* recombinant proteins as this system tends to produce recombinant proteins as inclusion body.

**CONCLUSION**

In conclusion, the native NA1 sequence has been successfully expressed in an insoluble form in tobacco. This study might also signify the necessity of the use of optimised sequences for expression work. Further investigations, like protein characterisations and codon optimisation need to be conducted in order to make a comparison with the present study and produce NA1 protein that is suitable for subsequent immunogenicity trial. Nevertheless, this study provides a great opportunity for the exploration of plant-manufactured NA1 protein as vaccine candidate. In future, other palatable plant species like alfalfa or lettuce could serve as a target system as combined feed/vaccine for poultry to combat AI disease.

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**REFERENCES**

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