Recombinant Subunit Rotavirus Trivalent Vaccine Candidate: Physicochemical Comparisons and Stability Evaluations of Three Protein Antigens

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Abstract

Although live attenuated Rotavirus (RV) vaccines are available globally to provide protection against enteric RV disease, efficacy is substantially lower in low- to middle-income settings leading to interest in alternative vaccines. One promising candidate is a trivalent nonreplicating RV vaccine, comprising 3 truncated RV VP8 subunit proteins fused to the P2 CD4\(^\text{\textsuperscript{+}}\) epitope from tetanus toxin (P2-VP8-P\([4/6/8]\)) to the P2-VP8-P\([4/6/8]\) displayed similar physical stability profiles as function of pH and temperature while P2-VP8-P\([8]\) was relatively more stable. Forced degradation studies revealed similar chemical stability profiles with Met\(^\text{1}\) most susceptible to oxidation, the single Cys residue (at position 173/172) forming intermolecular disulfide bonds (P2-VP8-P\([6]\) was most susceptible), and Asn\(^\text{7}\) undergoing the highest levels of deamidation. These results are visualized in a structural model of the nonreplicating RV antigens. The establishment of key structural attributes of each antigen, along with corresponding stability-indicating methods, have been applied to vaccine formulation development efforts (see companion paper), and will be utilized in future analytical comparability assessments.

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Introduction

Rotavirus (RV) is a leading cause of acute diarrhea and gastroenteritis among infants and young children across the world with approximately 215,000 children under 5 years of age dying from RV infection each year. Most deaths occur in developing countries. Furthermore, millions of children worldwide require home care, ER visits, and hospitalization, which contribute significantly to the disease burden. Improvements in sanitation, personal hygiene, or food quality reduce but do not eliminate the risk of this infection. Therefore, vaccination is the most cost-effective strategy to control the burden of RV-related illness.

Currently, 4 WHO prequalified, live attenuated oral RV vaccines (RotaTeq®, Rotarix®, Rotavac®, Rotasil®) are commercially available and combined cover more than 100 countries. In general, widely used RV vaccines (RotaTeq® and Rotarix®) show good efficacy (>85%) in developed countries, however, efficacy is reduced (40%-60%) in the low-income countries where the need is most. Although the causes for their reduced efficacy are unknown, and are an active area of investigation, contributing factors possibly include lower viral titer (transplacentally acquired RV antibodies, components of breast milk and stomach acid) and/or impaired immune response (malnutrition, interfering microbes, and other coinfections). From limited available data, lower efficacy in
certain subpopulations of the developing world of other live, attenuated oral vaccines has also been observed against enteric pathogens such as poliovirus and Vibrio cholera. Thus, there is interest in developing recombinant subunit protein, injectable RV vaccine candidates to address some of the current deficiencies of live attenuated oral vaccination.

To this end, a candidate recombinant protein—based, parenterally administered RV vaccine is under development (non-replicating RV vaccine [NRRV]). In a phase 1 clinical trial conducted in infants and toddlers in South Africa, a monovalent NRRV vaccine containing aluminum-adjuvanted P[8] antigen was shown to be well-tolerated and immunogenic, thus demonstrating a promising approach to develop a new vaccine against RV. Moreover, a trivalent version of the aluminum-adjuvanted NRRV vaccine is currently being evaluated in infants and toddlers in early stage clinical trials in South Africa. The trivalent NRRV vaccine candidate is composed of 3 recombinant protein antigens designed to protect against the most prevalent RV strains in the developing world (P[4], P[6], and P[8]). Each of these 3 antigens is a recombinant fusion protein based on the RV surface protein VP4. Upon host infection, VP4 is cleaved by proteases in the intestinal lumen into VP5* and VP8* leading to production of neutralizing antibodies. Wen et al. at the NIH successfully expressed ∆VP8*, a truncated, soluble version of the VP8* protein, in E. coli while preserving its key epitopes. The tetanus toxoid CD4+ T cell epitope (P2) was added on the N-terminus via a GSFGS linker to create a fusion protein which enhanced the immunogenicity of ∆VP8* as demonstrated in guinea pigs. In terms of nomenclature, the 3 recombinant protein antigens are P2-VP8-P[4], P2-VP8-P[6], and P2-VP8-P[8], where P2 stands for the carrier protein, VP8 refers to the ∆VP8* part of VP8* protein, and P[4], P[6], and P[8] represent the RV strain DS-1 (G2P[4]), 1076 (G2P[6]), and Wa (G1P[8]), respectively. For simplicity, the protein antigens are referred to as P[4], P[6], or P[8], respectively, in this work (see main text for further discussion).

Successful development and eventual commercialization of this recombinant subunit RV vaccine candidate will not only depend on clinical safety and efficacy results, but also the ability to produce the vaccine at low cost for use in the developing world. To this end, demonstrating a product candidate retains its critical quality attributes (CQAs, i.e., key aspects of structural integrity, stability, and immunological potency) as the manufacturing process is changed and scaled-up to ensure low-cost production is a key aspect to its development. To ensure CQAs are retained pre-versus post-process change, it is important to develop analytical assays that are robust and sensitive to detect changes in the physicochemical and immunological properties of the protein antigen. Thus, it is essential to develop a battery of analytical assays capable of monitoring structural and functional equivalence during comparability assessments. In addition, these analytical characterization methods can be applied to the formulation development to ensure vaccine potency (and physicochemical stability) of the 3 NRRV antigens across the vaccine’s shelf life.

The work presented in here, along with an accompanying companion paper, describes physicochemical characterization, forced degradation, and formulation development studies of the 3 NRRV recombinant fusion proteins as bulk vaccine antigens. In this work, a wide variety of physicochemical characterization techniques were used to characterize and compare the primary and higher-order structures, post-translational modifications, conformational stability, and aggregation propensity of each of the 3 NRRV antigens. In addition, forced degradation studies were performed to compare physical stability profiles as a function of pH and temperature as well as to elucidate chemical degradation pathways to identify chemically labile residues (i.e., “weak spots”). Analytical tools were developed to monitor or quantify degradative changes and the most informative assays for each structural attribute will be applied to formulation development and future comparability assessments. In a companion paper, the 3 NRRV antigens are further characterized in terms of formation of aggregates and particles (e.g., counting, sizing, and physicochemical attributes) during agitation and freeze-thaw stresses. In addition, formulation development studies are described to identify candidate formulations to minimize protein aggregation and particle formation during frozen storage of these NRRV antigens as a bulk drug substance (see companion paper by Agarwal et al. 27).

Materials and Methods

The P[4] and P[6] antigens were produced and purified from E. coli at Walter Reed Army Institute of Research, MD, and provided in 0.5 mM sodium phosphate, 150 mM NaCl, pH 7.2. The P[8] antigen was produced and purified from E. coli by Blue Sky Bio-Services, MA, and provided in 600 mM ammonium sulfate, 50 mM Tris, pH 7.5 buffer. Unless otherwise noted, each of the antigens was dialyzed overnight at 4°C in 10 mM sodium phosphate, 150 mM NaCl, pH 7.2 buffer which is referred as formulation buffer in the text hereafter. Sodium phosphate dibasic heptahydrate, and sodium chloride were purchased from Thermo Fisher Scientific (Waltham, MA). All other buffer reagents and chemicals including sodium phosphate monobasic monohydrate, citric acid, and ammonium bicarbonate were purchased from Sigma-Aldrich (St. Louis, MO) and were of analytical grade or higher unless noted otherwise. The extinction coefficient of each NRRV antigen was calculated from the primary sequence using online ExPaSy tool (Swiss Institute of Bioinformatics), resulting in values of 1.653 mg/mL−1 cm−1, 1.708 mg/mL−1 cm−1, and 1.733 mg/mL−1 cm−1 for P[4], P[6], and P[8] antigen, respectively, to determine protein concentration. It should be pointed out that the protein samples used in this study are bulk and by inference do not represent the aluminum-adjuvanted drug product vaccine.

Descriptions of many of the analytical methods used in this work have been published previously, and detailed descriptions for all methods in this work are provided in the Supplemental Methods section. These methods include intact mass spectrometry (MS), LC-MS peptide mapping, Fourier transform infrared (FTIR) spectroscopy, far-UV circular dichroism (CD) spectroscopy, fluorescence spectroscopy, SDS-PAGE, size-exclusion chromatography, sedimentation velocity analytical ultracentrifugation, reversed-phase (RP) ultra-high pressure chromatography, hydrophobic interaction chromatography (HIC), differential scanning calorimetry, extrinsic fluorescence spectroscopy (DSF), and OD₅₅₀ (turbidity) analysis. Experimental details for physical stability assessments using radar plots, chemical stability studies under forced degradation conditions, and structural modeling using I-TASSER are also described in the Supplemental Methods section.

Results

Physicochemical Characterization and Comparisons of the Three NRRV Antigens

The primary structure of the 3 fusion proteins was initially assessed by intact mass analysis. As shown in Figure 1, panels A, B, and C, predominantly a single peak was observed for each antigen and the average molecular weight results (20,517.6 ± 0, 20,732.0 ± 0, 20,433.7 ± 0.1 Da for P[4], P[6], and P[8], respectively, n = 3) correspond to the expected native protein mass based on the amino acid sequence plus an additional +132 Da (similar results were obtained for non-reduced samples, see Supplemental Fig. S1A). As
confirmed by LC-MS peptide mapping results described below, an additional Met residue is present on the N-terminus of these antigens, as would be expected for a foreign protein expressed in E. coli.

LC-MS peptide map analysis was conducted to confirm each antigen’s primary sequence and identify any posttranslational modifications. Each antigen displayed a unique peptide elution profile as shown in Figure 1 panels D, E, and F, demonstrating the ability to distinguish the 3 proteins as a potential identity test. Using intact (MS1) and fragmentation (MS2) mass analysis, 44 peptides were identified in the P[4] digest, 50 peptides in P[6], and 37 peptides in P[8], which covered 100% of each of the protein’s primary sequence (Supplemental Fig. S1B). Moreover, the peptide mapping results suggested deamidation at several Asn residues in P[4] and P[8], and at 1 Asn residue in P[6]. For P[4], Asn7, Asn47, and Asn179 were ~3%-4% deamidated. For P[6], Asn7 was ~2%-3% deamidated. For P[8], Asn7 was ~20%, Asn184 ~6%, and Asn178 ~3% deamidated. Thus, the peptide mapping method was shown to be useful not only as an overall fingerprint analysis for the structural integrity and identity of each NRRV antigen, but also as charge heterogeneity test for Asn deamidation (also see chemical stability section below).

The overall higher-order structure (HOS) of the 3 proteins was assessed by a combination of different biophysical tools. The secondary structure composition by FTIR by monitoring the absorbance in the amide I region (1600-1700 cm⁻¹). All 3 antigens were primarily β-sheet in structure with main peak at ~1642, ~1639, and ~1640 cm⁻¹ for P[4], P[6], and P[8], respectively, as shown in Figure 2, panels A, B, and C. The secondary structure composition by Fourier self-deconvolution (see Fig. 2d) correlated very well with the secondary structure assignments from X-ray crystal structures of P[4] and P[8] antigens in the literature. Far-UV circular dichroism spectroscopy was used to confirm the findings from FTIR analysis and a peak minimum around 215-216 nm was observed for each antigen indicating β-sheet structure in solution (Fig. 2e). The overall tertiary structure was compared using intrinsic tryptophan fluorescence emission spectrum from 305 to 405 nm. Peak position or λmax for P[4], P[6], and P[8] was 334 ± 0, 333.7 ± 0.6, and 335.3 ± 0.6, respectively (Fig. 2f). This result suggests the average Trp residue for each of the 3 antigens is located in a similar environment (each NRRV antigen has 4 Trp residues).

A combination of sedimentation velocity analytical ultracentrifugation and size exclusion-high performance liquid chromatography (SE-HPLC) were used to probe the size of the 3 protein antigens and to assess the presence of soluble aggregates (monomer, dimer, etc.). Each antigen had monomeric composition and about 1% higher molecular weight (HMW) species based on the peak areas from sedimentation coefficient (s) distribution (Fig. 3a). Similar s value of 2.09 ± 0.01 was recorded for P[4] and P[8], whereas for P[6], s value was 2.03 ± 0.01 suggesting smaller
hydrodynamic size for P[6] compared to P[4] and P[8] (which show similar size). These observations were supported by SE-HPLC as an orthogonal tool. The retention time for P[4] and P[8] was 16.28 ± 0.01 and 16.24 ± 0.01 min, respectively, whereas P[6] eluted at 17.08 ± 0.06 min consistent with a smaller hydrodynamic size for P[6] (Fig. 3b). As expected based on the size results from intact mass analysis (see aforementioned), P[6] antigen band migrated at a somewhat higher molecular weight on SDS-PAGE gels as compared to P[4] and P[8] (which were at similar mass level, Fig. 3c).

From HIC analysis (Fig. 3d), all 3 antigens appeared heterogeneous when eluted from the phenyl column with the major species accounting for >92% for each protein. Of the 3 antigens, P[4] was most homogeneous at 98.3 ± 0.3%, followed by P[6] at 92.8 ± 0.4% and P[8] at 92.3 ± 0.3% (n = 3). P[6] was most hydrophobic of the 3 antigens as it eluted at a relatively later retention time as compared to P[4] or P[8] (Fig. 3d). Similarly, through reversed-phase high pressure chromatography analysis (RP-HPLC), the major peak accounted for >93% of each protein. Of the 3 antigens, P[4] was most homogeneous at ~100 ± 0% followed by P[6] at 96.5 ± 0.2% and P[8] at 93.4 ± 0.1% (n = 3). P[6] eluted later than P[4] or P[8] as shown in Figure 3e. These RP-HPLC results are consistent with the HIC results (Fig. 3d). No difference in the total area was observed with and without the column in HIC and RP-HPLC assays indicating excellent recovery (i.e., no notable levels of protein were lost because of column adsorption).

**Physical Stability Profiles and Comparisons of the Three NRRV Antigens as a Function of Temperature and pH**

The HOS and conformational stability of each NRRV antigen was measured and compared as a function of temperature (10°C to 90°C) in the formulation buffer. Secondary and tertiary structure stability was evaluated by monitoring CD molar ellipticity at 216 nm and intrinsic Trp fluorescence emission peak intensity, respectively. As shown in Figures 4a and 4b, a single transition event was noted for each antigen and P[8] showed highest onset (T onset) and apparent melting temperature (Tm) as shown in the bar graphs on right. This result suggests that the HOS of the P[8] antigen is the most stable of the 3 antigens against thermal stress. Similar rank order of conformational stability was observed between the 3 antigens with P[8] being most stable (and P[4] was slightly more stable than P[6]) as measured by differential scanning calorimetry and DSF (Figs. 4c and 4d), respectively.

Two light scattering methods (SLS and OD350) were used to assess thermally induced aggregation. Results indicate P[8] is most stable (highest T onset value) as shown in the bar graphs on right panels of Figures 4e and 4f. The P[4] and P[6] antigens showed similar thermal induced aggregation profiles. It is interesting to note the substantially lower T onset values (by ~20°C) by SLS as compared to OD350 method for each of the 3 antigens. These differences in SLS versus OD350 results likely reflect the higher sensitivity of SLS toward formation of smaller aggregates compared to OD350 method which is likely more sensitive to the formation of larger aggregates/particles in solution.

Physical stability of each antigen as a function of both solution pH (from 3.0 to 8.0) and temperature (from 10°C to 90°C) was then monitored to measure changes in secondary structure, tertiary structure, and aggregation behavior. Overall, a pH-dependent destabilization was observed in secondary structure for each antigen (Supplemental Figs. S2a–S2c). For tertiary structure analysis, intrinsic Trp fluorescence mean spectral center of mass peak position showed some subtle changes in the thermal melt profiles, but in general, was not sensitive enough to monitor any potential differences between the antigens (Supplemental Figs. S2d–S2i). By contrast, DSF showed a clear pH-dependent destabilization with
lower pH being less stable as the temperature was increased (Supplemental Figs. S2j–S2l). Aggregation propensity versus solution pH was assessed by monitoring the intensity of total scattered light at 295 nm at 90°C. P[8] showed highest intensity values at pH 6.0 and pH 7.0 before precipitation occurred (Supplemental Fig. 2o). In summary, all 3 antigens showed a pH-dependent destabilization at lower pH values (with P[6] being least stable at pH 4.0 and pH 5.0).

To better visualize and compare the entire biophysical stability data sets of each antigen as a function of pH and temperature, a radar chart, data visualization analysis was utilized (Fig. 5). Each radar chart has 5 axes corresponding to the 5 biophysical measurements and the data are mapped to a pentagon such that smaller area of polygon indicate native-like state of protein and relatively larger area represents structural alterations. For each antigen, 4 distinct regions were observed (regions I, II, III/IV, V). Region I corresponds to a native-like structure of the antigen, region II represents partially structurally altered state, region III denotes low pH structurally altered state, region IV shows an aggregation prone region identified specifically for P[8] antigen, and region V represents more extensively aggregated and structurally altered state. Data for each technique can be read in the radar charts by following a particular axis. For instance, SLS signal is mapped to axis “a” (Fig. 5, radar chart key) and if we follow axis “a” for P[8] antigen in radar charts I to V; I—low signal indicate native-like or monomeric state, II—low to medium signal indicate low levels of aggregation, IV—high signal that is significant aggregation, V—low signal due to precipitation of aggregated protein. If the total relative area of native-like state (i.e., region I) of each antigen is compared; P[8] was most stable with an area of 39%, followed by P[4] with 33%, and P[6] was least stable (27%). In addition, a specific aggregation prone region (region IV) was identified for P[8], suggesting it might be prone to aggregation in that range of pH and temperature (note that P[4] and P[6] are also structurally altered and/or aggregated in region IV).

**Chemical Stability Profiles and Comparisons of the Three NRRV Antigens**

After subjecting the 3 NRRV antigens to elevated pH and temperature stress (pH 9.0, 25°C for 6 days), an increasing trend in molecular mass was observed for each protein by intact mass analysis compared to pH 7.2, 4°C control samples (Fig. 6a). Peptide mapping analysis was conducted to further probe the nature of the mass increase. Representative data for P[4] are shown in Figure 6b and similar data for P[6] and P[8] antigens are shown in Supplemental Figures 3a and 3b, respectively. Under no stress...
Figure 4. Higher-order structure (HOS) stability and aggregation propensity of the NRRV antigens in 10 mM PBS, pH 7.2, as a function of thermal stress (10°C–90°C). (a) Far-UV CD normalized mean residue ellipticity at 216 nm, (b) intrinsic tryptophan fluorescence normalized MSM peak intensity, (c) representative DSC thermograms of each antigen, (d) extrinsic ANS fluorescence MSM peak intensity, (e) static light scattering intensity at 295 nm, and (f) OD_{350} values for P[4] (black), P[6] (red), and P[8] (blue) antigens. Bar graphs on the right side of each panel show thermal onset (Tonset) and thermal melting temperature (Tm) values for each antigen. Error bars represent 1SD from triplicate measurements. DSC, differential scanning calorimetry; MSM, mean spectral center of mass.
conditions (pH 7.2, 4°C), deamidation was observed at Asn7 in each antigen and at each of the Asn-Gly (NG) sites (P[4] and P[8] antigens have 2 NG sites, whereas, P[6] has none). On incubation at pH 9.0 25°C for 6 days, P[4] antigen showed increased levels of deamidation at Asn7 and Asn-Gly residues (Asn47 and Asn179) as shown in Figure 6c. For P[6] antigen, Asn7 showed increased levels of deamidation under the same stress conditions. The P[8] antigen also showed increased levels of Asn7 deamidation and an increasing trend of deamidation was observed for the Asn-Gly residues (Asn46 and Asn178) under similar stress, although it was not statistically significant under these conditions (more notable levels of Asn46 and Asn178 deamidation were observed in P[8] when subjected to more aggressive stress condition of pH 9.0 40°C for 6 days; however, P[4] and P[6] antigens could not be tested for deamidation under this condition due to aggregation; data not shown. Overall, forced deamidation studies demonstrated the susceptibility of Asn7, Asn47, and Asn179 in P[4], Asn7 in P[6], and Asn7, Asn46, and Asn178 in P[8] toward deamidation under basic conditions at elevated temperature with the Asn7 residue being the most labile across all 3 antigens.

SDS-PAGE analysis (under non-reducing conditions) of the same elevated pH and temperature stressed P[6] samples showed an additional band corresponding to dimeric species which was absent or less abundant under reducing condition as shown in Figure 7a. This result suggests the dimeric species were linked with intermolecular disulfide bond because each NRRV antigen has single Cys residue. A faint dimeric band was also present under non-reducing condition for P[6] pH 9.0, 4°C stressed samples.

Figure 5. Biophysical stability profile from radar chart analysis of each NRRV antigen versus temperature across the pH range of 3.0 to 8.0. Sample buffer is 20 mM citrate phosphate buffer containing 150 mM NaCl at indicated pH value. (a) P[4], (b) P[6], and (c) P[8] show radar charts generated from multivariate stability data sets shown in Supplemental Figure S2. Radar chart key is provided in the bottom panel (see text for further explanation).
ultra-high pressure chromatography analysis showed an additional peak (~34.5 min) eluting at later retention time than the monomer peak (~30 min) as shown in Figure 7c. This additional peak was identified as P[6] dimer species by MS1 analysis (right panel, Fig. 7c) which correlates with SDS-PAGE results. However, the dimeric species were not observed by intact mass analysis presumably due to their low abundance and a single peak corresponding to monomeric protein was observed in control and stressed samples (Fig. 7b). For the P[4] antigen, a very faint HMW smear was observed at pH 9.0, 25°C (but not at pH 9.0, 4°C) under non-reducing conditions in SDS-PAGE analysis as shown in Supplemental Figure S4a. The P[4] dimer species also eluted at later retention time on reversed-phase chromatography as confirmed by the MS1 analysis of the eluting peaks (see Supplemental Fig. S4c). For the P[8] antigen, no additional HMW species was observed under the tested conditions suggesting higher stability of this antigen to non-native disulfide formation as compared to P[4] and P[6] (see Supplemental Fig. S5). Overall, the chemical stability profile demonstrates that P[6] is more susceptible to this non-native disulfide bond reaction compared to P[4] and P[8] antigens.
In terms of oxidative stress reactions, the 3 NRRV antigens were subjected to different amounts of H$_2$O$_2$. Figures 8a and 8b show representative data for P[6] antigen, a prominent peak for native species and low level of oxidized species were observed in the control sample (without H$_2$O$_2$) as measured by intact mass spectrometry and LC-MS peptide mapping, respectively. With the increasing amount of H$_2$O$_2$ the relative abundance of native species decreased, and the number and abundance of oxidized species increased. Peptide mapping analysis was conducted to identify the amino acids undergoing oxidation and also to quantify the oxidation relative to the control sample. As shown in Figure 8c, for P[6] antigen which has 2 Met residues, Met$^1$ was most susceptible to oxidation followed by Met$^{100}$. Similar observations were made for the 2 Met residues in P[8] antigen (Met$^1$ more susceptible than Met$^{99}$), as shown in Figure 8c and Supplemental Figure S8. The P[4] antigen has 3 Met residues and their susceptibility to oxidation can be rank ordered as Met$^1$ > Met$^{100}$ > Met$^{124}$ (Fig. 8c). Oxidized Met$^{100}$ in P[4], P[6], and Met$^{99}$ in P[8] were only detectable when samples were incubated with $>$0.1% H$_2$O$_2$ ($>$1000 ppm) as described in Figure 8c. Similarly, Met$^{124}$ in P[4] was oxidized only when exposed to $>$0.25% H$_2$O$_2$ (data not shown). No detectable change in the elution profile of digested peptides was observed until exposure to 0.1% H$_2$O$_2$, at which point the elution profile was altered and sequence coverage was reduced from 100% to ~80% for all 3 antigens (Supplemental Figs. S6-S8). In addition, the elution profiles of each sample (stressed or control) of a particular antigen were similar on a reversed-phase column, in which the primary peak eluted at ~9.5 min (P[4]), ~9.7 min (P[6]), and ~9.5 min (P[8])
as shown in Figure 8d and Supplemental Figure S9. In addition, the area of the peak corresponding to native species did not change substantially between samples (Fig. 8d) indicating no measurable loss of protein. In summary, intact protein mass measurement and peptide mapping by LC-MS methods are assays of choice to detect and monitor oxidation. The Met1 residue in each NRRV protein antigen was most susceptible to H₂O₂ induced oxidation, and oxidation was also observed at other Met residues under more aggressive stress conditions (≥1000 ppm H₂O₂).

**Discussion**

In this work, the primary and higher-order structures, post-translational modifications, and product-based impurities (e.g., charge heterogeneity, aggregates) of the 3 recombinant NRRV protein antigens were measured and compared using a wide variety of physicochemical characterization tools. In addition, degradation pathways of each of these proteins were elucidated via forced degradation studies to identify “weak spots” in terms of physical and chemical stability profiles. Although the potential impact of physicochemical structural alterations on immunogenicity is unknown at this time, developing such stability-indicating analytical tools and applying the structural knowledge gained in this work will be useful to (1) set critical manufacturing process parameters to ensure
consistency, (2) monitor key structural attributes during comparability assessments, and (3) develop stable formulations for the bulk drug substance and adjuvanted final drug product. The pharmaceutical stability profiles encountered during manufacturing and storage are not only dependent on such intrinsic properties (e.g., primary sequence/post-translational modifications,33 conformational stability,34 solubility35), but also extrinsic stress factors (e.g., storage temperature, freeze-thaw, agitation).36,37

Figure 9. Structural modeling of the 3 NRRV protein antigens. (a) Schematic description of the protein composition and nomenclature of each NRRV antigen, and (b) 3D structure prediction of each NRRV antigen using I-TASSER modeling; ΔVP8* protein is shown in gray, GSGSG linker in cyan, and P2 epitope in black. Residues susceptible to chemical degradation under stressed conditions are highlighted (Cys—red, Asn—green, Met—magenta).

Primary Structure, Post-Translational Modifications, and Chemical Stability Profile

An additional Met residue identified at N-terminus of each antigen was not unexpected because these antigens were expressed in E. coli and the efficiency of methionine aminopeptidase, enzyme responsible for excision of N-terminal methionine during translation, is about 30%-60% depending on the host.38 Residue next to the N-terminal Met for each NRRV antigen is Glu with a bulky side chain which could also hinder the efficiency of methionine aminopeptidase by steric effects as noted in the literature.39,40 Peptides generated after chymotrypsin digestion during peptide mapping analysis of each antigen showed distinct and very reproducible elution profile on a RP column which could be used as an identity test during future development of these antigens.

Oxidation via reactive oxygen species is a commonly observed chemical degradation pathway for protein biotherapeutics and vaccine antigens which can affect product quality, stability, and raises immunogenicity concerns.41,42 Peroxides are an example of
reactive oxygen species which can come in contact with vaccine drug products through various means such as the use of vaporized \( \text{H}_2\text{O}_2 \) as a sterilizing agent, as contaminants in formulation excipients such as polysorbates, and potentially even as trace quantities present in water depending on its source.\(^{43-46}\) Under forced degradation conditions (100-5000 ppm \( \text{H}_2\text{O}_2 \)), the \( \text{Met}^1 \) residue was most sensitive to oxidation in each of the 3 NRRV antigens. Interestingly, the \( \text{Met}^{100} \) or \( \text{Met}^{99} \) residue in each antigen, and \( \text{Met}^{126} \) in \( P[4] \) antigen were less prone to \( \text{H}_2\text{O}_2 \)-induced oxidation compared to \( \text{Met}^1 \) (which could be due to their reduced solvent accessibility within the three dimensional (3D) structure of the protein; see below).

Forced deamidation studies confirmed that \( \text{Asn}^7 \) in each NRRV antigen is susceptible to deamidation under conditions of basic pH and elevated temperature. \( \text{Asn}^7 \) in these proteins is followed by Ser residue which is the second most susceptible site for deamidation after NG sequence.\(^{36}\) The \( P[4] \) and \( P[8] \) antigens each have 2 NG sites (\( P[6] \) has none) in their primary sequence, and the \( \text{Asn} \) residues at these sites also showed increasing trend in deamidation levels for stressed samples (Fig. 7c). It is known that Asn deamidation in proteins and peptides depends on a combination of factors including their primary sequence, 3D structure (i.e., flexibility and solvent accessibility of these sites) and solution conditions (i.e., pH, buffer type, temperature).\(^{47,48}\) Overall, the peptide mapping method was able to detect and quantify the Asn deamidation in each NRRV antigen and thus will be a valuable tool during future formulation development and analytical comparability assessments.

The labile \( \text{Met}^1 \) and \( \text{Asn}^7 \) residues are both located in the P2 epitope region of these antigens which is highly flexible and unstructured as predicted by I-TASSER modeling (Fig. 9b) and some preliminary hydrogen deuterium exchange mass spectrometry studies (data not shown). Higher flexibility of this P2 region would explain the higher propensity of \( \text{Met}^1 \) to oxidation and \( \text{Asn}^7 \) to deamidation due to their relatively higher solvent accessibility compared to other Met and Asn residues which could be buried inside the protein structure.

In addition, the 1 free Cys residue in each NRRV antigen is also a “weak spot” in each antigen. Under stressed conditions of elevated pH and temperature, we observed reducible dimeric or multimeric species which were linked with intermolecular disulfide bonds. The \( P[6] \) antigen showed the highest susceptibility to this degradative reaction, followed by \( P[4] \), whereas dimeric or multimeric species were not observed for \( P[8] \) under the tested conditions. This result is in agreement with the physical stability data which showed higher conformational stability of \( P[8] \) antigen suggesting some sort of conformational alteration is needed for the Cys residue to get more soluble exposed allowing it to form non-native intermolecular disulfide bonds. Susceptibility of \( \text{Cys}^{173} \) in \( P[4] \) and \( P[6] \) to non-native disulfide formation could be due to lower conformational stability of the helical domain containing this Cys residue, thus exposing the free Cys to solvent and promoting the degradative reaction. Although the \( P[8] \) antigen was more stable to this degradative reaction under these conditions, disulfide linked covalent aggregates were formed with all 3 NRRV antigens when subjected to agitation stress (see companion paper by Agarwal et al.\(^{37}\)).

### Ongoing and Future Work

In terms of ongoing and future work, both physical and chemical degradation processes elucidated in this work (leading to structural alterations, chemical changes, or aggregate formation) could be detrimental to the development of NRRV antigens as a candidate vaccine without appropriate formulation development. The physicochemical characterization tools developed in this work were applied to monitor and characterize protein aggregation and particle formation during agitation and freeze-thaw stresses, and to develop candidate formulations to minimize their occurrence during storage and processing of these NRRV antigens as frozen liquid bulk drug substances (see companion paper, Agarwal et al.).\(^{37}\) It is important to note that these assays can also be adopted and modified to characterize and assess the physicochemical stability of the final vaccine drug product (a trivalent mixture of the 3 NRRV antigens formulated with an aluminum adjuvant) during storage as well as any potential effects of chemical alterations of the NRRV antigens on their interaction with aluminum adjuvants (work in progress). However, it is essential as part of future work to better correlate these physicochemical changes with in vitro potency using immunochemical (binding) assays (e.g., ELISA, biolayer interferometry, or surface plasmon resonance) utilizing antibody reagents which are specific to each antigen and bind to neutralizing
epitope(s). Eventually the results from physiochemical and in vitro potency assays need to be correlated with in vivo immunogenicity (animal studies) to determine their true impact on biological potency of the vaccine. For example, the chemical variants of NRRV antigens generated during forced degradation studies (e.g., oxidation, deamidation, and/or intermolecular disulfide bond formation) may or may not form during real-time storage, and similarly, may or may not impact their antigenicity and/or immunogenicity. Finally, the physiochemical analytical tools described in this work can be used during future comparability assessments of different lots of each of the NRRV antigens made from scaled-up manufacturing processes. It is not uncommon to introduce changes in the manufacturing processes during scale-up or switching to a different manufacturing site to keep the cost of vaccine production low, which is a key focus for the success of this subunit RV vaccine candidate for use in the developing world.

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