Chemical Reactivity of Brome Mosaic Virus Capsid Protein

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Viral particles are biological machines that have evolved to package, protect, and deliver the viral genome into the host via regulated conformational changes of virions. We have developed a procedure to modify lysine residues with S-methylthioacetimidate across the pH range from 5.5 to 8.5. Lysine residues that are not completely modified are involved in tertiary or quaternary structural interactions, and their extent of modification can be quantified as a function of pH. This procedure was applied to the pH-dependent structural transitions of brome mosaic virus (BMV). As the reaction pH increases from 5.5 to 8.5, the average number of modified lysine residues in the BMV capsid protein increases from 6 to 12, correlating well with the known pH-dependent swelling behavior of BMV virions. The extent of reaction of each of the capsid protein’s lysine residues has been quantified at eight pH values using coupled liquid chromatography–tandem mass spectrometry. Each lysine can be assigned to one of three structural classes identified by inspection of the BMV virion crystal structure. Several lysine residues display reactivity that indicates their involvement in dynamic interactions that are not obvious in the crystal structure. The influence of several capsid protein mutants on the pH-dependent structural transition of BMV has also been investigated. Mutant H75Q exhibits an altered swelling transition accompanying solution pH increases. The H75Q capsids show increased reactivity at lysine residues 64 and 130, residues distal from the dimer interface occupied by H75, across the entire pH range.

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Introduction

Viral particles are biological machines whose structure has evolved to package, protect, and deliver the viral genome into the host.¹,² The close integration of virion structure and function underscores the need to study virion architecture and the structural transitions that regulate the viral life cycle.³⁻⁸ There is also considerable recent interest in studying virion structure to advance the nanoengineering of viral particles for the delivery of drugs and molecular therapeutics.⁹⁻¹³
electron microscopic image reconstruction. Virions also show rapid fluctuations around an equilibrium structure that have been described as “breathing” vibrational modes. Conformational changes on both time scales can be studied using labeling techniques such as hydrogen–deuterium exchange (HDX) and covalent modification. Covalent modification techniques can either decrease the masses of proteins, as is seen with limited proteolysis to remove flexible “loops and fringes” of a protein, or increase protein masses, as in selective modification of specific amino acid side chains. In contrast to HDX, covalent modifications are not labile on the time scale of the analysis, thus minimizing back exchange of the label and reducing complexities in data interpretation. Side-chain reactivity can be affected by secondary, tertiary, and quaternary structure, and the patterns of reactivity displayed by a protein can be used to infer structural features and conformational changes. An additional advantage of covalent modification procedures is that hydrolytic enzymes and side-chain modifying reagents have well-characterized residue selectivities that allow additional sequence information to be derived from mass spectrometric analysis of modified proteins or peptides.

Thioimidates modify amino groups according to Fig. 1a, resulting in the mass increases summarized in Fig. 1b. We have used thioimidate reagents such as S-methylthioacetimidate (SMTA) to probe the structure of soluble proteins and the proteins of the bacterial ribosome. Lysines that are protected from SMTA modification are buried at the interface of protein–protein or protein–nucleic acid interactions or are involved in ionic or hydrogen-bonding interactions with other amino acid residues. The average extent of modification of proteins shows excellent agreement with predictions based on crystal structures, indicating minimal effects of native biomolecular structure.

Brome mosaic virus (BMV) was chosen to demonstrate the application of SMTA labeling to a dynamic macromolecular complex because of its simplicity (180 copies of a single capsid protein), the availability of a high-resolution crystal structure of the virion, and its high sequence homology (70% sequence identity) to cowpea chlorotic mottle virus (CCMV). High sequence homology to CCMV allows us to use crystallographic and electron microscopic structural models of CCMV virions to interpret our results from BMV. Both viruses undergo a pH-dependent swelling transition that results in a 10% increase in the virion’s 28-nm diameter as the solution pH increases from 5.0 to 8.0, with a transition midpoint at pH 6.5. The swelling transition was first characterized by analytical ultracentrifugation. Subsequent research described the sensitivity of the virion’s RNA and protein components to nuclease and protease digestion as a function of pH and the role of divalent cation binding to virions. More recently, the CCMV swelling transition was the subject of a molecular modeling study that compared in silico structural parameters to previous in vitro results. The generally accepted mechanism of bromovirus swelling involves the loss of divalent metal ions ligated by clusters of glutamic acid residues and consequent structural expansion driven by charge–charge repulsion. Tama and Brooks’ molecular dynamics study indicated that the pK of CCMV residue E81 (homologous to BMV E80) was elevated in the virion’s compact form relative to more expanded swelling intermediates (pK = 7.2–8.5 in the compact virion versus pK = 4.0–4.5 in the most expanded intermediate), implicating this residue as a keystone in the structural transition. Assuming a typical pK of 10.5, the BMV capsid protein’s 12 lysines are unlikely to be directly involved in the transition. However, their distribution on the exterior and interior surfaces of the virion and at the interfaces between capsid protein monomers makes their SMTA reactivity a useful readout of virion quaternary structure as a function of pH. The present report demonstrates that the chemical reactivity of lysine residues at the interface of capsid subunits is an accurate monitor of the pH-dependent structural transition of BMV virions. To further illustrate the utility of SMTA modification, we applied our labeling procedure to mutant virion that has an altered pH-dependent swelling transition. The results demonstrate that a residue at a protein–protein interface in a virion’s structure can influence the conformations of distal residues or the global flexibility of the entire virion.

![Fig. 1.](image-url) (a) Modification of protein or peptide amino groups by the thioimidate reagents described in the text. (b) The structural features of each reagent and the mass increase per modified amino group.
Results

BMV virions demonstrate pH-dependent SMTA reactivity

Thioimidate reagents have a lower tendency to engage in unproductive side reactions compared to oxygen-containing analogs such as O-methylacetimidate.\textsuperscript{49–51} However, thioimidates can still be hydrolyzed to unreactive amide and thiol products. Protonation of the imine (pK\textsubscript{a} \textapprox 7.0) increases the rate of hydrolysis,\textsuperscript{52–54} which would compete directly with the rate of lysine modification and decrease the SMTA concentration. Consequently, the observed pH dependence of the extent of lysine modification would contain contributions from both protein structure and SMTA hydrolysis. In order to control for the effect of hydrolysis of SMTA, conditions that allowed complete modification of protein amino groups across the pH range from 5.4 to 8.5 were developed. The 10 lysines and amino terminus of bovine pancreatic RNase A all appear to be surface accessible,\textsuperscript{54} and the protein exists predominately as a monomer in solution at the concentrations used for our experiments (2.5 μM, 0.03 g/L).\textsuperscript{55,56} When RNase A is subjected to 10 cycles of modification between pH 5.7 and 8.4, the result is nearly complete modification of all 11 of the protein’s amino groups across the entire pH range (Fig. S1 and Fig. 2a, open circles). The deconvoluted spectrum of RNase A reacted with SMTA at pH 5.4 (Fig. S1a) shows a distribution of labeled forms of the protein ranging from 7 to 11 amidinated lysine residues. At pH 5.7,

![Fig. 2.](image)

(a) The pH dependence of the extent of amidination of RNase A and the capsid protein of the BMV virion. The values on the ordinate axis are intensity weighted averages of the extent of modification. Error bars are the standard deviations of three determinations. (b) Apparent melting points [T\textsubscript{m(app)}] of WT BMV virions as a function of pH for three heating rates. Each point is the average of three determinations. (c) Melting points of unmodified (filled circles) and SMTA-modified (open squares) BMV virions as a function of pH. (d) Melting points of unmodified (open circles) and SMTA-modified (filled squares) BMV virions as a function of urea concentration. (e) Electron micrographs of unmodified and SMTA-modified virions. Virions labeled “acetate, pH 5.4” and “Tris, pH 8.6” were subjected to 10 cycles of buffer exchange with no SMTA. Virions labeled “SMTA, acetate” or “SMTA, Tris” were subjected to 10 cycles of SMTA modification as described in the text.
the distribution has converged to either 10 or 11 amidinated lysines (Fig. S1b). At pH values higher than 5.7, complete modification was observed (Fig. S1c and d). After 10 cycles of modification, the only indication of pH dependence for RNase A’s SMTA reactivity is at low pH values: as the pH decreases from 5.6 to 5.4, the average extent of modification decreases by one lysine residue (Fig. 2a, open circles). This decrease is most likely the effect of the pH decrease by one lysine residue (Fig. S2 and Fig. 2a, filled triangles). Significantly, as the pH increases from 5.4 to 8.5, an inflection point appears in the data at around pH 6.5. These changes in the SMTA reactivity of capsid protein lysines as a function of pH suggest that lysine modification by SMTA can be used to probe the relative solvent accessibility of the capsid protein structure during the pH-dependent swelling transition as long as chemical modification of the lysines does not significantly alter fundamental properties of the capsid protein.

Virion structure is preserved after SMTA modification

Differential scanning fluorimetry (DSF), a method to examine the thermal denaturation profile of proteins, and electron microscopy results demonstrate that the structure of BMV virions is essentially unchanged by amidination. Wild-type (WT) virions exhibit a sharp decrease in stability as a function of increasing pH, with an inflection point at pH ~6.5 (Fig. 2b). When the Tm(app) curve of unmodified virions (Fig. 2c, closed circles) is compared to SMTA-modified virions (Fig. 2c, open squares), differences between the two curves are minimal. First-derivative plots from raw melting point data for unmodified and SMTA-derivatized virions are presented in Supplemental Figs. S3 and S4, respectively. Modified virions exhibit a 2 °C decrease in Tm(app) at low pH that is likely due to a general loosening of the virion structure, a phenomenon that has been observed in other chemically modified viral capsids. A modest 2 °C increase in the melting point for SMTA-modified virions at high pH values was also observed, likely due to the more intense positive charge conferred by the amidinated lysine’s elevated pKa (estimated to be between 12.5 and 13.5), which could increase virion stability by enhancing either protein–protein or protein–RNA interactions. To compare RNA binding by the WT and SMTA-modified BMV capsid proteins, we performed a fluorescence titration of a well-characterized ligand, fluorescein-labeled B-Box stem loop RNA. Unmodified or amidated capsid protein gives nearly identical protein–RNA dissociation constants (0.13 µM for unmodified versus 0.11 µM for amidated capsid protein), as shown in Fig. S5. These results suggest that the stabilizing effect of amidination is due to enhanced protein–protein interactions. Previous fluorescence anisotropy titration experiments determined a Kd value for B Box binding to WT capsid protein of 449 nM. However, our Kd determination of the unmodified and modified capsid protein was made in parallel, and the small difference suggests that the stabilizing effect of amidination is due to enhanced protein–protein interactions. To further demonstrate the minimal structural effect of SMTA modification, the Tm(app) of virions was determined as a function of urea concentration. The nearly identical decreases of unmodified and SMTA-derivatized virions’ Tm(app) (Fig. 2d, open circles and filled squares, respectively) to increasing urea concentrations again demonstrate that the structural effects of lysine amidination are minimal. To put the changes observed in these analyses in context, virions containing capsid protein mutants with single amino acid substitutions of key residues or empty virus-like particles result in 5 to 15 °C decreases in BMV virion Tm(app) values (Fig. S6 and Ref. 63). The series of electron micrographs in Fig. 2e demonstrate that there are no gross structural differences between virions taken directly from SAMA storage buffer, virions that were repeatedly diluted and concentrated in either acetate or Tris buffer, or virions reacted with SMTA at pH 5.4 or 8.6.

SMTA reactivity of lysine residues in the BMV virion

Modified virions were proteolytically digested and the resulting peptide mixtures were analyzed by LC–MS/MS in order to map the locations of SMTA-modified lysine residues in BMV virions. Extracted ion chromatograms for identified peptides were integrated to yield intensities for each peptide in a digest sample, and the resulting data were used to determine the extent of labeling of each lysine within the capsid protein as a function of pH. Data on the extent of amidination of BMV capsid protein lysines as a function of pH are shown in Fig. S8. Amidination of lysine residues eliminates tryptic digest sites in a protein, resulting in digest mixtures containing progressively larger peptides. Manual integration of the total ion chromatogram (TIC) data is a partial remedy, as discussed below, but inevitably information on some portions of the protein is lost. Digestion of modified BMV virions with chymotrypsin is a better solution, and these data are discussed below. Results from the analysis of tryptic and chymotryptic digests are presented in Table 1. Peptides from all enzymatic digestions were
propionamidinated to maintain uniform ionization efficiency. Propionamidinated lysines have a 14-Da higher mass than acetylated lysines, allowing unambiguous identification of residues protected from SMTA modification in the native virion (Fig. 1b). Although the peptide acetyl-STSSTGKMTR with an amidination at K8 was detected irregularly, quantitatively reliable labeling data could not be obtained for K8, and results of this residue’s reactivity are not included in this work. Figures S9 and S10 in the supplemental material illustrate the distribution of tryptic and chymotryptic cleavage sites in the BMV capsid protein sequence, respectively. Partial lists of tryptic and chymotryptic peptide sequences are tabulated in Tables S4 and S5, respectively, along with the peptides’ predicted mass-to-charge ratios.

We identify four classes of lysines, based on inspection of the crystal structure and patterns of SMTA labeling: highly reactive surface residues, residues that are exposed on either the exterior or the interior surface of the virions, and residues that are sequestered at the interfaces of tertiary or quaternary structural features.

Table 1. Extent of amidination of lysines of the BMV capsid protein

<table>
<thead>
<tr>
<th>Lysine</th>
<th>Classificationa</th>
<th>% Amidinated lysine (standard deviation)</th>
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<tbody>
<tr>
<td></td>
<td>pH 5.4</td>
<td>pH 5.6</td>
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<td></td>
<td>pH 6.0</td>
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<td>pH 6.7</td>
<td>pH 7.2</td>
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<td>pH 8.0</td>
<td>pH 8.5</td>
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<tr>
<td>Tryptic data</td>
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<tr>
<td>41</td>
<td>Internal surface</td>
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<tr>
<td>44</td>
<td>Internal surface</td>
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<td>53</td>
<td>Interface</td>
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<td>64</td>
<td>External surface</td>
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<tr>
<td>81</td>
<td>Interface</td>
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</tr>
<tr>
<td>83</td>
<td>External surface</td>
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</tr>
<tr>
<td>86</td>
<td>Internal surface</td>
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<tr>
<td>105</td>
<td>External surface</td>
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<tr>
<td>111</td>
<td>External surface</td>
<td>—</td>
</tr>
<tr>
<td>130</td>
<td>Interface</td>
<td>—</td>
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<tr>
<td>165b</td>
<td>External surface</td>
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<td></td>
<td>Chymotryptic data</td>
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<td>41</td>
<td>Internal surface</td>
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<td>44</td>
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<td>Interface</td>
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<tr>
<td>165b</td>
<td>External surface</td>
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The extent of SMTA modification is quantified as a percentage of peptides containing the indicated lysine with an amidination out of all identified peptides containing that lysine. Mean values and standard deviations (in parentheses) are the averages of three to five determinations.

a Classification as “Surface” or “Interface” residues was accomplished using VIPERdb. Surface residues were further classified as “External” or “Internal” by inspection of a reconstructed capsid structure.

b Peptides containing lysine 8 were not observed reliably in either tryptic or chymotryptic digests. Peptides containing K165 were not observed in tryptic digests. See the text for discussion.

Highly reactive surface-exposed lysines of the BMV virion

Data for K105, K111, and K165 introduce the format used in all subsequent figures (Fig. 3). The locations of these three residues are depicted in an isolated capsid protein monomer and on the external surface of a hexamer of capsid proteins (Fig. 3a and b, respectively). The color used to depict each residue in the structure is also used to plot that residue’s extent of labeling data (Fig. 3c). The extent of modification of these three lysine residues is 80% or greater at all pH values, demonstrating that our procedure compensates adequately for SMTA hydrolysis at low pH. The 10% decrease in the reactivity of these residues at the lowest pH values serves as an estimate of the contribution of SMTA hydrolysis to any observed decreases in lysine reactivity at low pH.

There are few tryptic peptides that contain residue K165. Even in unmodified capsid protein, the smallest tryptic peptide containing K165 has a mass of 2420.8 Da (G145–K165); amidination of K165 increases the size of the smallest possible tryptic peptide to 5772.8 Da (G145–R189). Chymotryptic digestion pro-
duces more easily separated and identified peptides such as A158–L171 (1389.8 Da). Chymotryptic digest data in Table 1 indicate that the modification of K165 is nearly complete at all pH values. As a result, K165 is included in this class of lysine residues. The reactivities of K105, K111, and K165 imply that the capsid protein’s other nine lysines are responsible for the pH dependence of SMTA reactivity.

Surface-exposed exterior lysines of the BMV virion

K64 and K83 are on the exterior of the BMV capsid but are located near the interface of capsid subunits (Fig. 3d and e). At pH values below 6.0, chymotryptic digest data show that less than 40% of the K64 and K83 residues were modified by SMTA (Fig. 3f). At pH 6.0, however, approximately 80% of the K64 and K83 residues were modified. These results demonstrate that some residues on the exterior of the BMV particle are partially protected at low pH but become more solvent accessible as the pH increases. The observation that the ~30% and ~50% increases in SMTA reactivity for K64 and K83 occur before the pH transition midpoint of pH 6.5 suggests that the associated structural changes are sequential and affect the lysines exposed on the external surface of the virion before either the interfacial or the interior surface lysine residues.

Surface-exposed interior lysines of the BMV virion

Lysines K41, K44, and K86 are found on the interior surface of the virion (Fig. 4a and b). In the crystal structure of BMV, these lysines exist in a region of low electron density above the shell of encapsulated RNA. Chymotryptic digest data demonstrate that K41 is easily modified even at pH 5.4, indicating that SMTA can access the virion’s interior even in its low pH compact form (Fig. 4c). Approximately 50% of K44 positions and nearly 80% of K86 are less reactive at pH values below the major structural transition midpoint of 6.5, indicating that these lysines are partially protected by structural features of the compact virion (Fig. 4c and Table 1). Residue K44’s SMTA reactivity shows a 40% increase below the swelling transition midpoint of pH 6.5, suggesting that this position, adjacent to a pore on the pentamer axis, becomes more accessible before the main expansion of the virion. The reactivity of K41 and K44 is consistent with the currently accepted model of viral particles as dynamic structures whose multiple “breathing modes” allow solvent and small molecules access to the particle core without disrupting its structure. Partial tryptic digest experiments using BMV demonstrated that the N-terminal region containing K41 and K44 is flexible and can access bulk solution.
even in intact virions. The reactivity of K86 increases more slowly, consistent with its position close to the interface between subunits and only reaches 90% reactivity at pH 7.0. The large standard error in the extent of modification for K86 at intermediate pH values is due to digest-to-digest variability in chymotrypsin activity. Cleavage at M69 or L73 and L85 provides peptides that report reliably on the labeling states of K81 and K83, resulting in the low standard deviations seen for these residues (Figs. 5c and 3c, respectively). However, the peptides containing K86 are formed by cleavage at L85 and one of six succeeding sites (L91, L92, W93, L94, L96, or L97), effectively diluting the signals that report on K86’s extent of labeling among a collection of seven peptides.

**SMTA labeling of interfacial lysines of the BMV virion**

K53, K81, and K130 occupy positions on capsid protein surfaces that are buried by quaternary protein–protein interactions (Fig. 5a and b). K81 shows a dramatic increase in reactivity, starting at less than 10% modified below the transition midpoint pH and increasing to almost complete modification around pH 6.5, tracking the pH dependence of the overall virion swelling almost exactly. The interpretation of the data for residues K53 and K130 is more complex. SMTA modification of K53 increases steadily through the midpoint for the main structural transition but never reaches 100% at pH 8.5 (Fig. 5c), suggesting that it is exposed to solution late in the swelling transition and still somewhat protected from SMTA even in the expanded form of the virion. K130’s 40% reactivity at pH 5.4 and 5.6 shows it to be the most accessible interfacial residue in the compact virion. Modification of K130 shows a constant increase to complete reactivity over the pH range from 5.4 to 8.5. These results demonstrate that the interfacial residues make the most dramatic transition from a sequestered environment at low pH to full solvent exposure at high pH.

**Characterization of mutant virions**

Lysine modifications report on the responses of residues in three structural regions of the BMV virion. However, the high pK₅ of lysine side chains...
**(ε-amino pKₐ of 10.5)** suggests that they probably do not participate directly in the BMV virions' swelling transition. Histidine residues, with pKₐ values of ~6.4, are better candidates for sensing solution pH or triggering the pH-dependent conformational change of BMV virions. The BMV capsid protein contains three histidines, at positions 75, 170, and 175 (Fig. 6a). Perhaps relevant to the swelling transitions, all three histidines are located at the interface of dimers of the capsid subunits (Fig. 6b). Analysis of the conservation of the three residues revealed that, among members of the *Bromoviridae*,

![Partial sequence alignment](image)

Fig. 6. (a) Partial sequence alignment of histidine residues in the capsid protein sequences from members of the *Bromoviridae*. Conservative sequence substitutions are highlighted in red; others are highlighted in blue. Sequence numbering is for the BMV capsid protein. Additional abbreviations are as follows: BBMV, broad bean mottle virus; SBLV, Spring beauty latent virus; CYBV, Cassia yellow blotch virus; MYFV, Melandrium yellow fleck virus. (b) The location of residues H75, H170, and H175 at the interface of a capsid protein dimer. The two capsid protein subunits are colored white and gray. (c) Transmission electron micrographs of virions isolated from tobacco leaves transfected with the H75Q or H170Q mutants. (d) Northern blot analysis of the RNAs extracted from WT and mutant virions. The RNAs were detected with a riboprobe that recognizes the nearly identical 3' untranslated regions of the BMV RNAs as described by Hema et al. Each sample was from an independent preparation of virion from *N. benthamiana* plants. R1-R3: BMV genomic RNAs 1–3; R4: subgenomic RNA 4. (e) Virion melting points determined by DSF as a function of pH for WT BMV, H75Q, and H170Q containing mutant virions. Each value represents the mean T_m(app) of three determinations. Error bars for the standard deviation of each measurement are smaller than the symbols used for each data point. (f) WT and mutant virion melting points as a function of urea concentration. Solution pH was as indicated in the figure legend. Each value is the mean of three measurements, and standard deviations are smaller than the size of the symbols used for each data point.
H175 is absolutely conserved, H170 is conserved in all but one member, and H75 is the least conserved (Fig. 6a). To investigate the structural role of these residues, glutamine substitutions of each histidine were produced to generate viruses H75Q, H170Q, and H175Q.

Virion production was quantified by Bradford protein assays of the final virion isolates using bovine serum albumin (BSA) as a standard. Mutants H75Q, H170Q, and H175Q produced virions at approximately 50%, 10%, and less than 1% of WT, respectively. Isolated virions were also examined by electron microscopy. The yields of mutant H175Q precluded further investigation. Mutants H75Q and H170Q produced viral particles whose diameter and morphology are similar to that of WT (Fig. 6c). The RNAs packaged in WT and the two remaining mutants were extracted from CsCl-purified virions and electrophoresed on an agarose gel (Fig. 6d). Mutant H75Q has a distribution of genomic RNA comparable to WT, while the H170Q virions show decreased packaging of BMV RNA3 and RNA4. This aberrant RNA packaging pattern suggests that H170 exerts a direct or indirect influence on interactions between the BMV capsid and RNA3 or RNA4.

The DSF assay was used to characterize the stability of H75Q and H170Q virions as a function of pH (Fig. 6e). Mutant H170Q exhibits an inflection point at ~pH 6.5 similar to that of the WT, with a dramatic 30 °C decrease in the $T_{\text{m(app)}}$ above the transition midpoint. Mutant H75Q exhibits stability comparable to the WT both above and below the swelling transition. However, H75Q’s transition midpoint occurs at pH 6.2, suggesting that the presence of imidazole in H75Q is a significant factor in the virions’ response to changing pH. WT and H75Q virions show nearly identical decreases in their denaturation temperatures as a function of increasing urea concentration at pH 5.3 (Fig. 6f). The H170Q mutant has a marginally lower $T_{\text{m(app)}}$ at low urea concentrations and significantly higher $T_{\text{m(app)}}$ at high urea concentrations. These changes should be considered in the context of the 15 to 20 °C changes in the $T_{\text{m(app)}}$ for the WT virions when the pH is increased from 5.3 to 6.3 (Fig. 6f, filled black squares versus filled black circles). Furthermore, the $T_{\text{m(app)}}$ values for SMTA-modified H75Q and H170Q virions are nearly identical with the WT and their unmodified counterparts, lending further support to the conclusion that SMTA modification

![Fig. 7.](image-url)
does not alter or destabilize virion structure. Denaturation of the compact, low pH form of the virions is less likely to reflect contributions from molecular interactions that affect the swelling transition but would be expected to contain contributions from intramolecular interactions that destabilize the capsid protein's structure.

The three histidines at the interface of BMV capsid subunits have a range of effects on virion production, RNA encapsidation, and pH-dependent conformational changes. The H75Q mutant's structural integrity and the perturbation of its response to pH prompted us to examine the SMTA labeling of its lysines as a function of pH. To decrease virion heterogeneity, we made H75Q in a form that packaged only RNA3 and RNA4 (referred to as R3/4H75Q). Virions composed of WT capsid protein and containing only RNA3 and RNA4 (referred to as R3/4) were prepared for comparison to the R3/4H75Q particles. Both R3/4 and R3/4H75Q virions were purified at pH 5.5 and were indistinguishable from each other in electron micrographs (data not shown).

SMTA labeling of mutant virions

Both R3/4 and R3/4H75Q virions were subjected to the SMTA modification procedure, enzymatic digestion, and LC–MS/MS analysis. The WT R3/4 virions give results that are nearly identical with virions containing a natural distribution of all three BMV genomic RNAs (Fig. S10a and Table S6). The major differences are that R3/4 exhibited lower average standard deviations (5.6% and 5.8% for the tryptic and the chymotryptic digest data), compared to natural, heterogeneous virions (9.5% and 9.2% for tryptic and chymotryptic data) (Table S6 and Table 1, respectively). In contrast, R3/4H75Q revealed some significant changes in the pH-dependent reactivity of lysine residues (Fig. 7a and b). Although K64 was inaccessible to modification until the solution pH was above 6.0 in R3/4 virions, it was nearly completely modified even at pH values below 6.0 in R3/4H75Q particles (Fig. 7a). The twofold increase in interfacial residue K130's SMTA reactivity at pH values below the transition midpoint (Fig. 7b) indicates that the effect of the H75Q mutation can be detected using SMTA modification and that this mutation has a detectable structural influence on residues distal from the dimer interface occupied by H75.

Discussion

Chemical labeling of lysines followed by mass spectrometric analysis of peptides was previously used to probe protein–nucleic acid interactions in bacterial ribosomes. In order to extend this technique to different solution conditions, a procedure for the amidination of lysine residues that mitigated the effect of changes in solution pH on labeling efficiency was optimized using RNase A. When subjected to 10 cycles of SMTA modification, the SMTA reactivity of monomeric RNase A is nearly pH independent (Fig. S1 and Fig. 2a). In contrast, only half of the BMV capsid protein's lysine residues are modified at low pH, and as the reaction pH increases, the broad distribution of modified states converges to a single mass representing 12 completely amidinated lysines (Fig. S2 and Fig. 2a). These results demonstrate that protein–protein interactions in the BMV capsid can preclude the SMTA modification of the 12 lysines that are on the external surface, the internal surface, and the interfacial regions of the virion. Furthermore, the degree of modification of the lysines as a function of pH will provide insights to the conformational dynamics of the BMV virion. The details of the extent of modification were examined using proteolytic digestion and LC–MS/MS analysis.

Lysine modification and pH-dependent conformational changes of the BMV virion

There are four interesting exceptions to expected labeling behavior among the surface residues. At pH values above the pH 6.5 midpoint for the swelling transition, residues K44, K64, K83, and K86 are uniformly modified. As the pH decreases to values below 6.5, their reactivity is significantly attenuated, suggesting that the environments occupied by K44, K64, K83, and K86 become more sterically restricted in the compact, low pH virion structure. These observations are consistent with Wang et al.'s experiments on HDX of backbone amides in BMV capsid protein. They found that the backbone amide protons of six residues in the capsid protein sequence containing residues 1–47 and four residues in the range from 70 to 90 decreased their rates of HDX when the solution pH was lowered from 7.30 to 5.43, indicating a decrease in solvent exposure or conformational flexibility in the regions of the molecule containing K44, K83, and K86. A single backbone amide in a peptide containing residues 62–69 also decreased its rate of exchange, but the HDX exchange data did not provide residue-level resolution, preventing this change from being correlated directly to K64. Formaldehyde cross-linking of BMV RNA and capsid protein recovered peptides containing residue K44, indicating that this residue is capable of interacting with the viral RNA. Crystal structures of the closely related CCMV virion showed distinct bulges in the RNA-associated electron density directly below the lysine homologous to BMV K86, but the structure of the RNA could not be resolved due to crystallographic disorder. The combination of observed proximity
and pH-dependent differential protection from SMTA modification suggests that the interaction of K44 and K86 with the phosphodiester backbone of the virion's RNA changes as the virus swells.

Figure 7a and b shows the effect of the H75Q mutation on the SMTA labeling of K64 and K130. The H75Q mutant is of special interest since it encapsidates RNA similarly to WT BMV and has a urea denaturation profile comparable to WT virions at pH 5.3. The fact that the swelling transition of the H75Q mutant was altered suggests that the imidazole side chain of H75 either contributes to or is affected by the swelling transition. Capsid protein dimers such as the one depicted in Fig. 6b have contact surface areas between 1244 and 1589 Å² and are thought to be the basic assembly unit of the virion, leading to the expectation that any disruption of the interfacial region would have measurable effects on the stability of the virion. Comparing the locations of H75 in Fig. 6b with the location of K64 or K130 in Fig. 3b or Fig. 5b, it seems unlikely that direct interactions take place between these residues. When the low and high pH forms of the CCMV virion were compared, the largest radial increases in the low pH extent of labeling of K64 and K130 (Fig. 7a and b) indicate greater conformational flexibility in the virion structure due to the loss of a hydrogen bond donated by the H75 δ-N in the protonated imidazole. Interestingly, asparagine or aspartate residues are found in four of the six bromovirus sequences compared in Fig. 6a. The localized molecular interactions responsible for the differential reactivity of K64 and K130 (Fig. 7a and b) indicate greater conformational flexibility in the virion structure due to the loss of a hydrogen bond donated by the H75 δ-N in the protonated imidazole. Interestingly, asparagine or aspartate residues are found in four of the six bromovirus sequences compared in Fig. 6a. The localized molecular interactions responsible for the differential reactivity of K64 and K130 (Fig. 7a and b) indicate greater conformational flexibility in the virion structure due to the loss of a hydrogen bond donated by the H75 δ-N in the protonated imidazole.

Characteristics of the SMTA modification procedure

In addition to the information on the swelling transition of the BMV virion, this study should provide guidance to analysis of other proteins whose activities are affected by pH. These include endosome-resident proteins such as mammalian-binding proteins and Toll-like receptors. With regard to the activities of the BMV protein, we observed that SMTA modification had only minimal effects on the quaternary structures of the BMV virions or RNA binding by the capsid protein. These observations are consistent with SMTA modifications having only minimal effects on the structure of prokaryotic ribosomes.

An issue with the analysis of amidinated samples is trypsin inability to cleave at SMTA-modified lysines. Elimination of tryptic digest sites creates errors and inconsistencies in the data for three reasons. First, large peptides generated from samples modified at high pH do not always elute effectively from our C18 columns, resulting in the outright loss of information for some lysine residues. A dramatic example can be seen by comparing the K81 labeling data from R3/4 virions generated from tryptic and chymotryptic digests (Fig. 5c and f). As the pH increases in the tryptic digest data, K81's extent of modification approaches 80% and then drops suddenly to ~40% (Fig. 5c). The chymotryptic data for K81 show a smooth increase to ~90% modification at pH 8.5 (Fig. 5f). This difference occurs because the number of tryptic peptides that report on the labeling of K81 decreases as K53, K64, K83, and K86 are more completely amidinated at higher pH, eliminating these lysines as tryptic cleavage sites. Conversely, the chymotryptic data are based on a uniform collection of peptides at every pH value. Second, the progressively larger tryptic peptides that do elute into the mass spectrometer support higher charge states. Although the Orbitrap mass analyzer can accurately measure the precursor masses of such peptides, the MS/MS spectra produced in the linear ion trap contain +3 and +4 fragment ions that are not recognized by automated database searches. Moreover, if more than one lysine in such a peptide is differentially modified, assignment of the position of the amidination will be ambiguous, increasing the errors in the extent of modification data. Finally, the current version of the ProtParser software was designed for use in bottom-up proteomics experiments and does not parse peptides with charge states >+4. The +5, +6, or +7 charge states of the
large peptides generated in tryptic digests of extensively modified BMV display higher intensities than the +3 or +4 charge states. A remedy for this problem is inspection of the Mascot output to identify large peptides, followed by manual extraction and integration of XICs for their higher charge states. Tryptic digest data presented in Table 1 and Table S6, and in Figs. S4a-c and S5a-c have been corrected this way.

Summary and conclusions

Our data provide a dynamic view of the BMV virion structure in solution to augment static crystallographic or cryoelectron microscopic data. The reactivity of some “surface-exposed” lysine residues was essentially pH independent, as expected from their classification. However, other surface lysine residues showed pH-dependent changes in SMTA reactivity, specifically protection at lower pH. These results demonstrate the ability of SMTA modification to provide experimental evidence of interactions between amino acid residues that otherwise must be inferred from structural data or that might be overlooked.

Materials and Methods

Chemicals

Water was purified by a Barnstead Thermolyne Nanopure system. Ammonium chloride, magnesium chloride hexahydrate, and methyamine as a 40% (v/v) solution in water were supplied by Aldrich. HPLC-grade acetonitrile, 2-propanol, methanol, trifluoroacetic acid, and formic acid were products of J. T. Baker. To determine the approximate pH of reaction mixtures, EMD Chemicals (Hepes, free acid), leucine enkephalin, 4-(2-hydroxyethyl)-piperazine-1-ethanesulfonic acid (Hepes, free base), dithiothreitol, and 1,3-propanediol (Tris, free base) were products of J. T. Baker. Dye Reagent, Bio-Rad, Hercules, CA) using BSA as a standard.

Preparation of thioimidate reagents

SMTA was synthesized according to Thumm’s method as described by Beardsley and Reilly. The product form of this reagent is the hydroiodide salt of SMTP. S-Ethylthiopropionimidate (SETP) was synthesized using an adaptation of the Pinner synthesis of diethylsulfothioimidate reported recently by Lauber and Reilly. Propionitrile (0.02 mol, 1.4 mL) was dissolved in methylene chloride (4.3 mL, dried over molecular sieves) in a round-bottom flask. A 10-fold molar excess of ethanethiol (0.20 mol, 14.8 mL) was added, and the solution was immersed in an ice bath and stirred while bubbling HCl gas into the solution for 1 h. During this interval, the reaction flask was vented into a solution of 1 M sodium hydroxide. After charging the reaction flask with HCl, the flask was stoppered and placed in an airtight bag (to minimize release of ethanethiol) and placed in a cold room at 4 °C. After overnight incubation, the hydrochloride salt of the product was precipitated by adding diethyl ether and placing the reaction mixture into a −20 °C freezer. After 1 h at −20 °C, 2.26 g of white solid were collected by filtration through Whatman’s #1 paper, a 73% yield of the clear and colorless SETP hydrochloride.

The final products of all syntheses were stored in vacuo over Drierite pellets until used.

Production of BMV virions and capsid protein

Agrobacterium tumefaciens was used to launch BMV infection in the geranium tobacco, Nicotiana benthamiana, as described previously. To produce BMV virions containing RNA3 and its subgenomic copy RNA4 encapsidated by WT capsid protein (referred to as R3/4 BMV), we infiltrated N. benthamiana plants with a mixture of Agrobacterium cells engineered to express the BMV 1a and 2a replication proteins and the replication-competent RNA3 that also directs transcription of RNA4. Three separate cDNAs for BMV RNA3 containing mutations of histidines 75, 170, or 175 glutamines were made by PCR-mediated site-directed mutagenesis using the QuikChange kit and protocols recommended by the manufacturer (Stratagene, Torrey Pines, CA). Primers for each mutant are available upon request. WT virions and virions expressing each of the three histidine mutants were all purified from N. benthamiana leaves according to previously published procedures. Virion yields were quantified by Bradford dye binding assays (Quick Start Bradford Dye Reagent, Bio-Rad, Hercules, CA) using BSA as a standard.

Dissociated capsid protein samples were prepared from WT virions by dialysis in Slide-A-Lyzer dialysis cassettes (Thermo-Fierce, Rockford, IL) against a buffer containing 50 mM Tris, pH 7.5, 500 mM CaCl2, and 1 mM DTT for 24 h at 4 °C. Following centrifugation at 97,000 g for 1 h to remove the precipitated RNAs, the supernatant was dialyzed against 50 mM Tris, pH 7.5, 300 mM NaCl, and 1 mM DTT for 24 h. The final dialyze had an A260/A280 ratio <0.65, and the protein concentration was determined by Bradford assay using BSA as a standard.

Modification of viral particles with SMTA

The buffers used were 500-mM stock solutions of acetic acid (pH 5.4 and 5.8), the free acid forms of Mes (pH 6.0 and 6.5) and Heps (free pH 6.7 and 7.2), and Tris free base
were removed by diluting the reaction to 500 μM ammonium hydroxide. Stock solution pH was determined with a Corning Scholar 425 pH meter equipped with a glass pH electrode. The buffer stocks were used to prepare solutions of SMTA and to prepare more dilute 100-mM reaction buffer solutions. When modifying BMV virions at varying pH values, an IQ400 ion-selective field effect transistor (Hach Company, Loveland, CO) was used to measure the initial pH of the dilute buffer solution, the pH of the dilute buffer solution after addition of an aliquot of SMTA-containing solution, and the pH of the reaction solution after 1 h. Addition of the SMTA solution to the dilute buffer solution typically caused a 0.1–0.2 unit decrease in pH, but the pH stayed constant after 1 h. Tris-buffered solutions showed large decreases in pH after SMTA addition (from 8.7 to 8.0 and from 10.6 to 8.5). The reported reaction pH values were measured after the initial addition of SMTA to the dilute reaction buffer solutions.

The SMTA protein modification procedure was developed using bovine pancreatic RNase A before application to virions. When BMV virions were modified, 160 pmol of BMV protein were diluted into a total volume of 100 μL in an Amicon Microcon centrifugal concentrator with a 10,000-Da molecular mass cutoff filter (protein concentration: 1.6 μM or 0.03 g/L). All SMTA reactions and subsequent sample cleanup were performed in the same filter cartridge at room temperature (22–24 °C). Reaction solutions contained a 100-mM concentration of one of the buffers described above and 20 mM MgCl2. The solution was centrifuged for 16 min and diluted with 80 μL of the reaction buffer, and then a 20-μL aliquot of 200 mM SMTA in concentrated buffer solution was added. The BMV capsid protein contains 12 lysines and an acetylated N-terminus; thus, the SMTA:protein mole ratio was 25,370:1 and the SMTA:amino group mole ratio was 2100:1. This process was repeated 10 times to ensure that any observed protection of lysine amino groups from SMTA reaction was due only to tertiary and quaternary structure rather than competing hydrolytic degradation of the SMTA. After 10 cycles of modification, residual reaction products were removed by diluting the reaction to 500 μL with 100 mM ammonium bicarbonate containing 50 mM DTT to reduce any mixed disulfide bonds formed by reaction with methanethiol produced by amidination of lysines in proteins.31 Residual DTT and ammonium bicarbonate were removed by diluting the protein twice to 500 μL with 0.1% (v/v) formic acid in water and concentrating it to a final volume of 50 μL. Samples were then transferred to septum-capped vials for whole protein LC–MS analysis.

Whole protein LC–MS

The average extent of SMTA modification of RNase A or BMV capsid protein was determined with a Micromass QTOF Micro spectrometer. Sample handling and gradient development were provided by a Waters Alliance 2795 chromatograph using the gradient of Table S1. Protein samples in 0.1% (v/v) formic acid in water from the previous section’s modification procedure were concentrated and desalted by injecting 30 to 40 pmol of protein onto a reversed-phase capillary column (Upchurch Scientific Blue PEEK, 10 cm × 0.0254 cm, packed with Phenomenex Jupiter C4 material). Raw profile mode spectra were summed and extracted from TICs manually, and the masses of SMTA-modified forms of the proteins were determined by deconvolution of the raw spectra using MaxEnt 1 (Micromass). The resulting whole protein spectra were centroided and exported as text files containing mass/intensity pairs for calculation of the intensity weighted average extent of modification using Microsoft Excel 2007.

Differential scanning fluorimetry

The effect of SMTA modification on BMV structural integrity was examined using DSF. After 10 cycles of modification with SMTA in Tris buffer at pH 8.0, aliquots containing 5 μg of BMV capsid protein were diluted into 30 μL of reaction buffer solutions and then transferred to a 96-well PCR plate. A 1-μL aliquot of a 200× stock solution of SYPRO Orange dye in dimethyl sulfoxide (Sigma) was added to each well.38 Samples of virions containing unmodified protein were diluted and loaded into a second set of wells at the same concentrations of BMV virions and SYPRO Orange as above. The temperature gradient and fluorescence were monitored with an Agilent Stratagene Mx3005P (Agilent, Santa Clara, CA). The samples were equilibrated at 25 °C for 5 min, and then their temperature was increased to a final value of 95 °C in 0.5 °C increments with a ramp time of 30 s. Fluorescence intensity was monitored at 570 nm. Raw data were processed with Microsoft Excel 2007.

WT mutant virions were diluted into buffers containing 0, 2, 4, and 6 M urea to further characterize the effects of the H75Q and H170Q mutations on capsid protein and virion structure. Quantities of protein, SYPRO Orange, and the temperature gradient program controlling the Agilent Stratagene Mx3005P were identical with the details above.

Fluorescence polarization titration

Samples contained 0.25 μM fluorescein-labeled B Box stem loop (Integrated DNA Technologies, Coralville, IA) in 50 mM Tris, pH 7.5, and 50 mM NaCl.67 Concentrations of unmodified or SMTA-modified BMV capsid protein ranged from 2 pM to 7 μM. Fluorescence polarization measurements were made using a PanVera Beacon 2000 fluorescence polarization system (PanVera Corp., Madison, WI) equipped with 488-nm excitation and 535-nm emission filters. Prior to data collection, samples were incubated at 37 °C for 15 min, and the 37 °C temperature was maintained by the instrument. Dissociation constants for the RNA–protein complex were extracted by fitting the raw data against a noncooperative saturation binding curve with OriginPro 8.6 (OriginLab, Northampton, MA).

Electron microscopy sample preparation and image collection

BMV virions were added to glow-discharged copper-coated grids at a final concentration of 0.01 mg/mL and stained with 1% (w/v) depleted uranyl acetate. Electron
micrographs with a 60,000 × magnification were recorded using a JEOL 1010 transmission electron microscope operating at 80 kV. 

**Enzymatic digests and peptide LC–MS/MS**

Labeled BMV capsid proteins were exchanged into 20 mM ammonium bicarbonate or 20 mM Tris for trypsin or chymotrypsin digestion, respectively. The protease was added to a final concentration of 4% or 3.3% (w/w) of the capsid protein, and enzymatic digestion was allowed to proceed under the conditions described in Table S2. Peptides produced by enzymatic digest were reacted with SMTP or SETP to ensure uniform ionization efficiency and to mass tag lysine residues that were protected from SMTA modification in the native virions. Because thioimidate modification of free amino groups proceeds rapidly at pH values above 7.5, enzymatic digestions were terminated by the addition of an aliquot of 200 mM SMTP or SETP stock solution in 250 mM Tris free base to a final concentration of 100 mM reagent. The propionamidination reaction was incubated at 25 °C for 1 h and terminated by the addition of 10% trifluoroacetic acid in water. Doubly labeled peptide samples were purified using C18 spin columns (Pierce Biotechnology). Eluted peptides were concentrated in vacuo and then resuspended in 25 μL of 0.1% aqueous formic acid. Peptide samples were analyzed by LC–MS/MS on a 75-μm i.d. fused silica column packed with C18 material (Magic C18, Michrom Bioresources, Auburn, CA), using a Dionex Ultimate 3000 to supply the gradient shown in Table S3. Eluting peptides were detected with a Thermo LTQ-Orbitrap hybrid mass spectrometer. Peptide masses were measured to a specified resolution of 30,000 in the Orbitrap cell while the instrument’s linear quadrupole ion trap was used to produce and measure CID MS/MS spectra of the five most intense peptides detected in a given precursor spectrum.

**Bioinformatics and quantitation of labeling results**

The extent of SMTA modification of each BMV capsid protein lysine was quantified by integration of TIC data using programs from the National Center for Glycomics and Glycoproteomics ProteinQuant Suite. Lists of precursor ions and associated fragment ions were extracted from raw TIC data files into Mascot Generic Format using the program TurboRAW2MGF. Peptides were identified by Mascot searches against either the SWISSPROT database filtered for “Viruses” or a database containing only BMV protein sequences. Modifications used as search parameters included acetylation of the protein’s N-terminus, amidation of lysine, propionamidination of lysine or a peptide’s N-terminus, oxidation of methionine, and deamidation of asparagine or glutamine. Mascot results were parsed with ProtParser allowing for peptide charge of +1 to +4, the minimum MOWSE score for all peptides of 1 and 400, respectively, a minimum intensity for all peptides being 1, and with up to 5 allowed missed cleavages and no exclusion of doubled tryptic cleavage sites. Redundant appearances by the same peptide sequence were combined under the appearance with the highest MOWSE score.

Peptides were quantified using ProteinQuant 2.6. Parameters for peptide signal integration include the initial retention time for a peptide from the ProtParser results, a maximum chromatographic peak width of 1.0 min, a search window for peak reassignment of 1.5 min, beginning and ending edge definitions based on intensity alone, a baseline intensity calculated from the first 10 min of the chromatogram, and normalization based only on the total intensity of the identified peptides. ProteinQuant output files were converted to Microsoft Excel 2007 spreadsheets to tabulate each appearance by the 12 lysines from the BMV capsid protein.

Crystal structures of BMV and RNase A were obtained from the Protein Data Bank‡ and visualized using PyMOL v. 0.99 (DeLano Scientific) or UCSF Chimera. The classification of lysine residues in the BMV virion as surface or interfacial used the Amino Acid Information utility available at VIPERdb§.

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**Acknowledgements**

W.E.R. would like to thank Dr. Stella Aniagyei and Professor Bogdan Dragnea for an introduction to BMV preparation and handling. This work was supported by National Institutes of Health grant 1R01AI090280 to C.K. and National Science Foundation grants CHE-1012855 and CHE-0832651 to J.P.R.

**Supplementary Data**

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.jmb.2012.06.031

**References**


‡ [www.pdb.org](http://www.pdb.org)

‡ [www.pymol.org](http://www.pymol.org)

§ [http://viperdb.scripps.edu](http://viperdb.scripps.edu)


