NOTES

Disulfide Bonds and Membrane Topology of the Vaccinia Virus A17L Envelope Protein

TATIANA BETAKOVA† AND BERNARD MOSS*

Laboratory of Viral Diseases, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland 20892-0445

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The envelope protein encoded by the vaccinia virus A17L open reading frame is essential for virion assembly. Our mutagenesis studies indicated that cysteines 101 and 121 form an intramolecular disulfide bond and that cysteine 178 forms an intermolecular disulfide linking two A17L molecules. This arrangement of disulfide bonds has important implications for the topology of the A17L protein and supports a two-transmembrane model in which cysteines 101 and 121 are intramemnlal and cysteine 178 is cytoplasmic. The structure of the A17L protein, however, was not dependent on these disulfide bonds, as a recombinant vaccinia virus with all three cysteine codons mutated to serines retained infectivity.

Crescent-shaped membranes, comprising the earliest defined vaccinia virus structures in the cytoplasm of infected cells, develop into spherical, immature virus particles and subsequently into dense, brick-shaped, infectious intracellular mature virions (IMV) (4). Some IMV are then wrapped by modified trans-Golgi membranes, transported to the periphery, and released from the cell (8, 19). Electron micrographs suggested that the crescent membranes, and their precursors which accumulate in the presence of the drug rifampin, are composed of single lipid bilayers unconnected to cellular organelles (3, 7, 9). Other studies, however, suggested that these viral structures are comprised of two closely apposed membranes that are derived from the cellular intermediate compartment connecting the endoplasmic reticulum to the Golgi network (11, 18, 21). Further biochemical, morphological, and genetic studies are needed to understand the biogenesis of IMV membranes and to resolve conflicting data. IMV contain at least 11 virus-encoded membrane proteins, of which A14L, A17L, and D13L are needed for assembly of the spicule-covered crescents (14, 17, 26, 27). Of these, the A17L protein is the best characterized. Disulfide-linked A17L protein dimers have been reported to form a complex with dimers of the A14L protein (16) and trimers of the A27L protein (15). The A17L protein is posttranslationally cleaved near the N and C termini (2, 15, 22) within AGX consensus motifs previously identified as cleavage sites for other vaccinia virus structural proteins (25). The C-terminal truncation of the A17L protein is dependent on the F10L kinase (2), which is also required for envelope formation (23, 24). The A17L (2, 5) and A14L (2) proteins are phosphorylated directly or indirectly by the F10L kinase, providing a role in morphogenesis for the latter enzyme.

The topology of the A17L protein determines the intracellular sites of protein interactions, cleavage, phosphorylation, and disulfide bond formation. Krijnse-Locker et al. (11) demonstrated that the N and C termini of the A17L protein are cytoplasmically oriented, consistent with an even number of membrane-spanning segments. Although the A17L protein has four hydrophobic domains, recent topological studies of Betakova et al. (1) supported a two-transmembrane model. In the present study, we mutagenized the three cysteines of the A17L protein and found evidence for a previously unrecognized intramolecular disulfide bond between cysteines 101 and 121 which is compatible with a two-transmembrane model (11). The formation of disulfide-bonded A17L dimers was dependent on cysteine 178, which is in the C-terminal cytoplasmic domain of the A17L protein. Nevertheless, the structure of the A17L protein was not dependent on these disulfide bonds, since a recombinant virus in which all three cysteine codons were mutated to serines retained infectivity.

Evidence for intra- and intermolecular disulfide bonds. Purified virions were treated with sodium dodecyl sulfate (SDS) and 2-mercaptoethanol (2-ME) and analyzed by SDS–12.5% polyacrylamide gel electrophoresis (PAGE) and Western blotting with an antibody to the mature N terminus of the A17L protein (2). As previously shown (2), the predominant band was a C-terminally truncated monomer of about 23 kDa (Fig. 1A, lane 6). In addition, a minor dimer band was detected, in agreement with observations of Krijnse-Locker and Griffiths (12), suggesting difficulty in completely reducing the intramolecular disulfide bond. When the reducing agent was omitted, there was relatively more dimer and the major dimer and monomer bands migrated more rapidly than the reduced forms, of which only trace amounts could be detected (Fig. 1A, lane 4), suggesting a more compact structure due to an intramolecular disulfide bond. Iodoacetamide or N-ethylmaleimide (NEM) is frequently used to alkylate free sulfhydryl groups and prevent artificial postlysis disulfide bond formation. Addition of 50 mM iodoacetamide prior to virus disruption had no major effect on the relative amounts of the unreduced monomeric and dimeric forms of the A17L protein from IMV membranes (Fig. 1A, lane 5). In other experiments, cells were lysed in the presence of 50 mM iodoacetamide and 20 mM NEM and the alkylating agents were maintained during virus
purification through a sucrose cushion. Under these conditions, the dimer and monomer bands were also similar in intensity (data not shown). Although the effect was barely evident, iodoacetamide caused the monomer species to migrate slightly faster than the form that received neither iodoacetamide nor 2-ME (Fig. 1A, compare lanes 5 and 4), presumably due to alkylation of the same cysteine that can form an intermolecular disulfide bond.

Lysates of vaccinia virus-infected BSC-1 cells were analyzed in a manner similar to that used for purified virions. The polypeptide pattern, however, was more complex due to mixtures of N- and C-terminally cleaved and uncleaved forms. Under favorable electrophoretic conditions, the C-terminally truncated form of the A17L monomer was resolved from its uncleaved precursor (2), as shown in Fig. 1A, lane 3. The incompletely reduced disulfide-bonded dimer was also resolved as faint C-terminally truncated and precursor species (Fig. 1A, lane 3). Without 2-ME, the dimer bands were more prominent and both the dimer and monomer bands migrated more rapidly than the reduced forms, consistent with the presence of an intramolecular disulfide bond (Fig. 1A, lane 1). Iodoacetamide had only a slight effect on the mobility of the unreduced monomers (Fig. 1A, lane 2), suggesting that the intramolecular disulfide bond had been formed before cell lysis. In contrast, there was much less of the dimeric forms of A17L in the presence of iodoacetamide (Fig. 1A, lane 2) than in its absence (Fig. 1A, lane 1), consistent with postlysis dimer formation. In some experiments, however, iodoacetamide had little effect on the amount of dimer, corresponding to data obtained by Krijnse-Locker and Griffiths (12) using NEM as the alkylating agent. One explanation is that some of the A17L protein exists in a noncovalently linked dimeric form with closely apposed cysteines so that disulfide bond formation can compete with alkylation.

Effects of cysteine-to-serine mutations on disulfide bond formation. The A17L open reading frame (ORF) contains cysteines at positions 101, 121, and 178. To determine which ones are involved in disulfide bond formation, we needed a way of expressing mutated proteins during a virus infection. The A17L ORF with its late promoter was inserted into a plasmid, and the cysteines were individually mutated to serines using the QuickChange Site-Directed Mutagenesis Kit (Stratagene) and appropriate primers. For expression, the plasmids were transfected into cells that were subsequently infected with a conditionally lethal vaccinia virus mutant, vA17LΔS, with a stringently repressed A17L gene (26). In the absence of isopropyl-β-D-thiogalactopyranoside (IPTG), the only detectable A17L protein was derived from the transfected plasmids but all other viral proteins were expressed from the viral genome. The 2-ME-induced decrease in mobility of the monomeric species detected with the wild-type A17L protein (Fig. 1A) was not observed when either cysteine 101 or 121 was changed to serine, although a dimeric species was still formed (Fig. 1B, lanes 1 to 6). However, no dimeric form was detected when cysteine 178 was changed to serine but the 2-ME-induced shift of the monomeric species occurred (Fig. 1B, lanes 7 to 9). These data suggested that the intramolecular disulfide bond of the wild-type A17L protein forms between cysteines 101 and 121 and that cysteine 178 is required for intermolecular disulfide bond formation.

Complementation of plaque formation by transfected A17L genes. We used the transfection-infection protocol described above to determine whether the mutated A17L genes are functional for virus assembly and infectivity. Repression of A17L gene expression results in a severe reduction of vA17LΔS plaque size and number (26). Plaque formation was enhanced in the absence of IPTG, however, by transfection of a plasmid containing the A17L gene regulated by its own late promoter (Fig. 2A, A17L). Remarkably, plaque formation was also enhanced by transfection of the A17L gene with mutations of...
cysteine 101 (Fig. 2A, C101S), 121 (Fig. 2A, C121S), or 178 (Fig. 2A, C178S). Furthermore, the A17L gene with mutations of two cysteines (Fig. 2A, C121, 178S) or three cysteines (Fig. 2A, C101, 121, 178S) also rescued plaques. By contrast, enhanced plaque formation did not occur when a stop codon was formed by introducing an A residue at position 309 to give a C-terminal truncation (Fig. 2A, A17L-309A) or when eight amino acids between the second and third hydrophobic domains of A17L were replaced with a similar-length influenza virus hemagglutinin epitope tag (1) (Fig. 2A, TAG). These results indicated that neither the intramolecular nor the intermolecular disulfide bonds of the A17L protein are needed for virus assembly and spread, whereas certain other mutations produce nonfunctional A17L proteins.

Isolation and characterization of mutant vaccinia viruses with cysteine-to-serine mutations. Following the protocol in the previous section, BSC-1 cells were transfected with the plasmid containing all three cysteine mutations and infected with vA17LΔ5. Viruses in plaques that formed in the absence of IPTG were subjected to repeated rounds of plaque purification, and 30 were amplified and screened by SDS-PAGE under reducing conditions and Western blotting. The majority of isolated viruses made the wild-type A17L protein. Three viruses that made A17L proteins with or without IPTG that migrated faster than the wild type in the presence of 2-ME were further characterized. One of these formed disulfide-linked dimers, and two did not. PCR analysis indicated that each virus had a single copy of the A17L gene that had replaced the inducible copy, and sequencing indicated that one (vA17LC101,121S) had serine substitutions of the first two cysteines but retained the cysteine at position 178, whereas in the other two (vA17LC101,121,178S), which did not form A17L protein dimers, serines replaced all three cysteines. The yields of the A17L cysteine mutants in both the cell lysates and medium were similar to those of WR and vA17LΔ5 at 37.5°C in the presence of IPTG (Fig. 3). Thus, the three cysteines of the A17L ORF were not required for replication of vaccinia virus.

When the A17L protein of purified vA17LC101,121S virions was examined, the disulfide-linked dimer was present but the 2-ME-induced mobility change of the monomer was not observed (Fig. 4A), consistent with the roles of cysteines 101 and 121 in intramolecular disulfide bond formation. As predicted, the A17L protein of purified vA17LC101,121,178S virions lacked both inter- and intramolecular disulfide bonds (Fig. 4A). The absence of the intermolecular disulfide bond correlated with the ability to efficiently extract the A17L protein from purified vA17LC101,121,178S virions in the absence of a reducing agent (Fig. 4B). In contrast, the extraction of the A17L protein from vA17LΔ5 (made in the presence of IPTG), WR, or vA17LC101,121S virions was enhanced by a reducing agent (Fig. 4B). The extraction of other membrane proteins in the absence of a reducing agent, detected by polyclonal antiserum to vac-
cinia virus, was not enhanced by the mutations of the A17L protein (data not shown), suggesting that the general integrity of the membrane had not been radically altered. In addition, preincubation of purified viruses at elevated temperatures prior to plaque formation did not reveal enhanced thermal sensitivity of the mutants (data not shown).

Conclusions. The present study confirmed previous reports regarding the existence of disulfide-bonded dimers of the A17L protein, provided evidence for an intramolecular disulfide bridge, and demonstrated that cysteine 178 is required for the latter. Disulfide bonds usually form within the endoplasmic reticulum of mammalian cells or the periplasm of prokaryotic cells, which provide a favorable redox potential, as well as molecular chaperones that assist in folding (6, 10, 13). It seems likely, therefore, that the disulfide bonds of the A17L protein would form in the endoplasmic reticulum. In this regard, a recent topological analysis of the A17L protein provided evidence for two membrane-spanning domains and placed cysteines 101 and 121 in an intraluminal location (11), consistent with stoichiometric disulfide bond formation. In contrast, a suggested model with four membrane-spanning domains (11) would place cysteines 101 and 121 in different compartments and is incompatible with the data presented here. Cysteine 178 is located in the cytoplasmic tail of the A17L protein, accounting for the variable amounts of disulfide-bonded molecules reported previously (12). It seems likely, however, that cysteine 178 residues of noncovalently linked A17L dimers are closely apposed, allowing some cytoplasmic disulfide bond formation and further disulfide bond formation when virions are released by cell lysis.

The A17L protein is required for assembly of vaccinia virions and is conserved in other vertebrate poxviruses. Of the three cysteines, the two that form the intramolecular disulfide bond are more highly conserved, as they are present in the orthologous protein of molluscum contagiosum virus, a distantly related member of the poxvirus family (20). Based on the conservation of the two cysteines, we anticipated that they might be required for proper folding or function of the A17L protein. To test this hypothesis, a recombinant vaccinia virus with all three cysteines of the A17L protein mutated to serines was constructed. The infectivity of this mutant in cell culture established that the disulfide bonds of the A17L protein are not needed for virion assembly or spread. Although the disulfide bonds help to retain the A17L protein in the viral membrane, as shown by the need for reducing agents to extract the wild type but not the mutated polypeptide, the mutant virions were not noticeably less heat stable than wild-type virions. Our inability to demonstrate an appreciable in vitro effect of mutating cysteines, however, does not preclude advantages of disulfide bond formation with regard to the assembly, structure, or stability of vaccinia virions in vivo.

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REFERENCES