**Vaccinia virus B5 protein affects the glycosylation, localization and stability of the A34 protein**

**Adrien Breiman and Geoffrey L. Smith**

Department of Virology, Faculty of Medicine, Imperial College London, St Mary's Campus, Norfolk Place, London W2 1PG, UK

Vaccinia virus has two infectious forms, the intracellular mature virus, which has a single envelope, and the extracellular enveloped virus (EEV), which is surrounded by two lipid bilayers. The outer membrane of the EEV contains at least six viral proteins. Among them A34, a type II membrane glycoprotein, and B5, a type I membrane glycoprotein, form a complex and are involved in processes such as morphogenesis and EEV entry. A34 is required for normal incorporation of B5 into the EEV membrane. Here, we used a virus lacking B5 and viruses with mutations in the B5 membrane-proximal stalk region and looked at the effect of those modifications on A34.

Data presented show that B5 is required for the correct glycosylation, trafficking and stability of A34, emphasizing the complex interactions and mutual dependence of these vaccinia EEV proteins.

**Vaccinia virus** (VACV) is the prototypical member of the genus *Orthopoxvirus* of the *Poxviridae*. It replicates in the cytosol and produces multiple types of infectious virions (Smith et al., 2002; Condit et al., 2006; Roberts & Smith, 2008). The first infectious progeny is the intracellular mature virus (IMV), which is surrounded by a single-lipid envelope (Dales & Siminovitch, 1961; Hollinshead et al., 1999) and remains in the cell until cell lysis. However, some IMV are transported via microtubules to the early endosomes or trans-Golgi network where they are wrapped by two cellular membranes containing several VACV proteins. The resulting intracellular enveloped virus (IEV) is then transported on microtubules to the cell surface where the outer membrane fuses with the plasma membrane to externalize a double-enveloped virus by exocytosis. This virion is called cell-associated enveloped virus (CEV) if it remains on the cell surface, or extracellular enveloped virus (EEV) if it is released from the cell. The CEV/EEV outer membrane contains at least six viral proteins: A33 (Roper et al., 1996), A34 (Duncan & Smith, 1992), A56 (Shida, 1986), B5 (Engelstad et al., 1992; Wolfe et al., 1993), F13 (Blasco & Moss, 1991) and K2 (Turner & Moyer, 2006; Wagenaar & Moss, 2007). A34 is a type II transmembrane protein with different glycoforms between 23 and 28 kDa and its extracellular part contains a C-type lectin-like domain (Duncan & Smith, 1992). A K151D point mutation in the VACV strain Western Reserve (WR) A34, which is present naturally in the VACV International Health Department (IHD)-J strain, caused an increase in EEV release (Blasco et al., 1993). Similarly, deletion of the A34R gene (vA34R) from VACV WR caused a 25-fold increase in EEV, but such EEV had a fivefold reduction in specific infectivity (McIntosh & Smith, 1996). Deletion or suppression of the A34R gene caused a small plaque phenotype (Duncan & Smith, 1992; McIntosh & Smith, 1996), inability to form actin tails (Wolfe et al., 1997; Sanderson et al., 1998) and severe attenuation (McIntosh & Smith, 1996).

B5 is a 42 kDa type I transmembrane glycoprotein (Engelstad et al., 1992; Isaacs et al., 1992) with an extracellular domain composed of four short consensus repeats (SCRs) characteristic of complement control proteins (Takahashi-Nishimaki et al., 1991), although there is no evidence that B5 regulates complement activity. After the SCRs B5 has an acidic stalk region (ST) before the transmembrane domain (TM) and a short cytoplasmic tail (CT). Both the SCRs and CT are dispensable for targeting B5 to the EEV membrane (Herrera et al., 1998; Lorenzo et al., 1998; Mathew et al., 1998), although the latter affects its transport to the cell surface (Mathew et al., 2001) and recycling via endosomes (Ward & Moss, 2000). B5 is needed for IMV wrapping to form EEV (Engelstad & Smith, 1993; Wolfe et al., 1993). B5 and A34 interact (Rottger et al., 1999; Earley et al., 2008; Perdiguerio et al., 2008; Roberts et al., 2009) and in the absence of A34, the amount of B5 incorporated in EEV is decreased markedly (Earley et al., 2008; Perdiguerio et al., 2008; Roberts et al., 2009). B5 and A34 each affect the glycosaminoglycan (GAG)-dependent rupture of the EEV outer membrane during EEV entry (Law et al., 2006; Roberts et al., 2009).

Although B5 expressed on its own displays a cellular localization profile very similar to the one observed in the context of viral infection (Katz et al., 1997; Lorenzo et al., 2000), this is not the case for A34. In infected cells, A34 is found at the Golgi, on the cell surface and in CEV/EEV. In
contrast, when it is expressed alone it accumulates in the perinuclear region and does not go to the plasma membrane (Lorenzo et al., 2000). In addition, attempts to express A34 on its own from classical eukaryotic expression vectors (pcDNA3, pCI) and in several recombinant expression systems yielded poor levels of expression. For example, in a system where soluble forms of EEV proteins A56, B5 and A33 were expressed in CHO cells and secreted into the medium, the yield obtained for A34 was about 20-fold lower than for B5 (Law et al., 2005; Putz et al., 2005; M. Law unpublished data). Here, we present data showing that in the absence of B5, the level of A34 is markedly decreased, most probably because of misfolding and consequential degradation.

In a recent study on EEV entry, we generated several VACV mutants with alterations in the B5 stalk acidic residues (Roberts et al., 2009), for the structure of these mutants see Fig. 1a. Using those viruses, we analysed lysates from infected RK13 cells by immunoblotting with mouse monoclonal antibodies (mAbs) against B5 (36-6; Roberts et al., 2009), A34 (34-1; Roberts et al., 2009) and the IMV protein D8 (AB1.1; Parkinson & Smith, 1994) as an infection control (Fig. 1b). Mouse mAbs against the A34 and B5 proteins were produced by immunization of mice with purified recombinant protein expressed from mammalian cells (Law et al., 2005; Putz et al., 2005). As noted previously, these mutations affected the electrophoretic mobility of B5 (Roberts et al., 2009). In addition, this analysis showed that in the absence of B5, the amount of A34 in the infected cells was reduced considerably, and with some of the B5 mutants the glycosylation profile of A34 was different. When B5 was deleted, one distinct band at about 20 kDa (A34*1) was observed instead of the 23–28 kDa bands made by wild-type (WT) virus. In contrast, deletion of all the B5 SCRs (vST23-35ala) had no effect and A34 retained the WT profile (note the remaining B5 fragment was not visible due to its small size). However, substitution of the acidic residues of the stalk with alanines (vST2-35ala) had the same effect as deleting B5. Moreover, substitution of the five acidic residues closest to the membrane (vST23-35ala) also led to the A34* profile. Interestingly, vST2-16 and vST28-35 showed a mixed profile with the A34* band evident together with higher molecular mass forms (Fig. 1b). Since vST23-35ala and vST28-35ala differ only at residue 23 and display distinct A34 glycosylation profiles, we wondered if that amino acid could by itself influence A34 glycosylation. To address this, a recombinant VACV in which the aspartic acid 23 of the stalk region was mutated to alanine (vST23ala) was constructed by using transient dominant selection as described previously (Roberts et al., 2009). When tested by immunoblotting as above, vST23ala showed the same A34 profile as the WT virus and therefore, mutating the aspartic acid 23 is not sufficient to alter A34 glycosylation. To determine more precisely the nature of the A34 20 kDa isoform, we used drugs that affect glycosylation: namely tunicamycin, an inhibitor of N-acetylglucosamine transferase and kifunensin, an inhibitor of α-mannosidase I (Fig. 2a). RK13 cells were infected with WR or vAB5R at 5 p.f.u. per cell for 90 min and then incubated overnight in Dulbecco’s modified Eagle's medium containing 2.5 % fetal bovine serum with or without 1 μM tunicamycin or 5 μM kifunensin. Cell lysates were then prepared and analysed by immunoblotting (Fig. 2b). Treatment of WR-infected cells with tunicamycin produced a single A34 band, corresponding to the unglycosylated polypeptide (A34ug). The A34ug is

![Fig. 1](image-url)
predicted to be 19.6 kDa (Duncan & Smith, 1992), and we observed a slightly smaller band of about 17 kDa. In vΔB5R-infected cells, A34 is slightly larger than A34ug and was still reduced in size in the presence of tunicamycin, indicating that A34* is a partially glycosylated form. Notably, levels of B5 and A34 both decreased in the presence of tunicamycin, indicating that glycosylation is required for stability of these proteins. In the presence of kifunensin, the A34* pattern was observed in WR-infected cells, suggesting that A34* represents an intermediate with nine mannose residues (Man9), before trimming by the α-mannosidases.

Another interesting observation was that the amount of A34 in the absence of B5 was increased by kifunensin treatment. This is in agreement with reports showing that processing by α-mannosidases acts as a signal to target misfolded proteins for proteasomal degradation and that inhibition of these enzymes by kifunensin treatment leads to accumulation of misfolded (Man9)-glycoproteins (Olivari & Molinari, 2007). To address this further, the stability of A34 with time was investigated (Fig. 2c). RK13 cells were infected with WR or vΔB5R as before and cell lysates were prepared at 4, 8, 12 and 24 h p.i. and analysed by immunoblotting. Up to 8 h p.i., A34 was easily detected in cells infected by either virus, although it was less abundant in vΔB5R-infected cells, but thereafter A34 declined substantially in vΔB5R-infected cells and was barely visible at 24 h. This suggests that synthesis and accumulation of A34 starts normally without B5, but as the rate of synthesis decreases later during infection, the level of A34 declines. Collectively, those data suggest that in the absence of B5, A34 is misfolded and degraded.

Next, the effect of A34 glycosylation status on the subcellular localization was investigated. BSC-1 cells were infected with viruses at 2 p.f.u. per cell for 8 h, fixed with PBS-4% paraformaldehyde (PFA) for 10 min on ice and then in PBS-8% PFA for 20 min at room temperature. Fixed cells were permeabilized with 0.2% Triton X-100 and incubated with anti-A34 mAb and a rabbit anti-protein disulphide isomerase Ab (anti-PDI; Abcam) to stain the endoplasmic reticulum (Fig. 3). Consistent with Fig. 2(c), significant levels of A34 were present in both WR- and vΔB5R-infected cells at 8 h p.i. In WR-infected cells, the anti-A34 mAb labelled the Golgi as well as punctate structures corresponding to virions in the periphery, as described previously (Lorenzo et al., 2000), but no significant co-localization with PDI was observed (Fig. 3a). In contrast, in vΔB5R-infected cells A34 was

Fig. 2. In the absence of B5, A34 has aberrant glycosylation and is targeted for degradation. (a) Simplified diagram of the glycosylation pathway showing where tunicamycin and kifunensin act. Glc, Glucose; GlcNAc, N-acetylglucosamine; Man, mannose. (b) RK13 cells were infected with WR and vΔB5R and incubated with or without tunicamycin (Tunic.) or kifunensin (Kifu.). At 16 h p.i., cell lysates were prepared and analysed by immunoblotting with anti-B5, anti-D8 and anti-A34 mAbs. Another image of the bottom section of the membrane is shown for clarity. A34ug and B5ug, unglycosylated A34 and unglycosylated B5. (c) A34 is synthesized in the absence of B5, but is degraded over time. RK13 cells were infected with WR or vΔB5R and harvested at the indicated times p.i. Cell lysates were prepared and analysed by immunoblotting as in (b). The positions of molecular mass markers are shown on the left in kDa.
present throughout the cell in a reticular pattern colocalizing with PDI (Fig. 3a), similar to that seen when A34 was expressed from a Semliki Forest virus vector (Lorenzo et al., 2000). No staining of VACV particles was observed, and this may be explained by the wrapping defect of vAB5R (Engelstad & Smith, 1993; Wolffe et al., 1993). Cells infected with vAB5R, vST23-35ala and vST28-35ala all showed a significant amount of A34 in the ER and nuclear envelope, but vST23-35ala and vST28-35ala also showed some staining of viral particles (Fig. 3b; enlargement of the punctate staining representing virions is shown for the vST28-35 image). Cells infected with vST23-35ala generally had fewer particles than WR- or even vST28-35ala-infected cells. This is consistent with the fact that vST23-35 produced much less EEV than WR, whereas EEV production by vST28-35ala was only slightly reduced (Roberts et al., 2009). Overall, these data show that the A34* band in SDS-PAGE correlates with the presence of A34 in the ER. This is consistent with A34* being a (Man9)-A34 that accumulates in the ER.

Data presented here indicate that in the absence of B5, or in presence of some mutated forms of B5, A34 is not correctly folded and accumulates in the ER as a partially glycosylated intermediate. Ultimately, at least in the case of vAB5R, this would lead to proteasomal degradation. A hypothesis to explain these data would be that an interaction of the negatively charged acidic residues in the B5 stalk region with positive charges of A34 might help A34 to acquire the correct conformation. Alternatively, the B5 stalk could play a role in the trafficking of the B5/A34 complex.

Taken together with the previous data showing that A34 is required for proper incorporation of B5 in the EEV membrane, this shows that there is a complex mutual interaction between these two VACV proteins making it difficult to unravel their respective roles in virus wrapping, egress and re-entry due to their inter-dependence.

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