Vaccinia virus protein C16 acts intracellularly to modulate the host response and promote virulence

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The vaccinia virus (VACV) strain Western Reserve C16 protein has been characterized and its effects on virus replication and virulence have been determined. The C16L gene is present in the inverted terminal repeat and so is one of the few VACV genes that are diploid. The C16 protein is highly conserved between different VACV strains, and also in the orthopoxviruses variola virus, ectromelia virus, horsepox virus and cowpox virus. C16 is a 37.5 kDa protein, which is expressed early during infection and localizes to the cell nucleus and cytoplasm of infected and transfected cells. The loss of the C16L gene had no effect on virus growth kinetics but did reduce plaque size slightly. Furthermore, the virulence of a virus lacking C16L (vΔC16) was reduced in a murine intranasal model compared with control viruses and there were reduced virus titres from 4 days post-infection. In the absence of C16, the recruitment of inflammatory cells in the lung and bronchoalveolar lavage was increased early after infection (day 3) and more CD4+ and CD8+ T cells expressed the CD69 activation marker. Conversely, late after infection with vΔC16 (day 10) there were fewer T cells remaining, indicating more rapid clearance of infection. Collectively, these data indicate that C16 diminishes the immune response and is an intracellular immunomodulator.

INTRODUCTION

Vaccinia virus (VACV) is the prototypical member of the genus Orthopoxvirus (OPV) of the Poxviridae and is famous as the live vaccine used to eradicate smallpox, an extinct human disease caused by variola virus (VARV) (Fenner et al., 1988). Following the eradication of smallpox, VACV continues to be studied intensively because of its development as a vaccine vector (Panicali et al., 1983; Smith et al., 1983), and because of its fascinating interactions with the host cell and immune system (Seet et al., 2003; Haga & Bowie, 2005). In this study, we have characterized the C16L gene of VACV strain Western Reserve (WR) and the encoded protein.

The VACV WR C16L gene is located in the inverted terminal repeat (ITR) at both ends of the VACV genome. Many genes within and adjacent to the ITRs encode proteins that modulate the host immune response or interactions with the host cell. For example, the gene upstream of C16L encodes the VACV growth factor (Twardzik et al., 1985; Buller et al., 1988), which promotes cell growth, and the gene two genes downstream encodes the interleukin (IL)-18-binding protein (Smith et al., 2000; Symons et al., 2002; Reading & Smith, 2003). The C16L gene is predicted to encode a 331 aa protein with a mass of 37.5 kDa (www.poxvirus.org). There are highly conserved orthologues of C16 in several other OPVs, suggesting an important function, and bioinformatic analysis identified a conserved 6 aa sequence at the C terminus of C16 that is present in the same region of the IL-1 receptor antagonist (IL-1ra) protein (Kluczyk et al., 2002). This 6 aa sequence is also conserved in the VACV strain Copenhagen orthologue of C16 (termed C16) (Goebel et al., 1990), and in orthologues in other OPVs (www.poxvirus.org). This sequence is critical for the ability of IL-1ra to antagonize signalling from the IL-1 receptor (IL-1R) and prompted the proposal that the VACV protein might act as a secreted viral IL-1ra and inhibit signalling from the IL-1R by receptor blockade (Kluczyk et al., 2004).

The goals of this study were to characterize the C16 protein, determine its location and investigate if it affects virus replication or virulence. A VACV deletion mutant lacking both copies of the C16L gene (vΔC16) and a revertant virus (vC16Rev) in which the C16L gene was reinserted into the deletion mutant were constructed. Data
presented show that C16 is non-essential for virus replication, but promotes VACV virulence in a murine intranasal (i.n.) model and delays the infiltration and activation of cells in the infected lungs. As such, it represents an intracellular virulence factor that functions as an immunomodulator.

**METHODS**

**Cell culture.** BS-C-1, TK−143 and HeLa cells were grown at 37 °C in a 5% CO₂ atmosphere in Dulbecco’s modified Eagle's medium supplemented with 10% fetal bovine serum (FBS; Gibco). The source of VACV strain WR was described previously (Alcami & Smith, 1992).

**Construction of plasmid vectors.** A cassette containing the *Escherichia coli* guanylylphosphoribosyl transferase (*Ecogpt*) gene fused in-frame with the enhanced green fluorescent protein (*Egfp*) gene and downstream of synthetic early/late VACV promoter was assembled as follows. The *Ecogpt* gene was amplified by PCR with primers AF1 (5’-AGTCTAGATCCGGATCCACCTGCGG-3’) and AF2 (5’-GGGCTAGACGCGTCCGATGGGAGGTGCGGGG-3’) and pGpt9/14 as template (Boyle & Coupar, 1988). The *Egfp* gene was amplified by PCR using primers AF3 (5’-GGTCGCTCAGATCCGCTAGCGCTACCG-3’) and AF4 (5’-AGTCGAATTCATAGCGAAAAATACATCGTCACCTGGGAC-3’) gene and pEGFPC2 (BD Biosciences) as template. These PCR products were ligated together using a synthetic early/late promoter wherein both copies of the synthetic early/late VACV promoter was assembled as follows. The resultant plasmid, pET28a-C16L-CHis, was termed pAFE. Recombinant C16 protein was expressed from *E. coli* strain BL21 according to the vector manufacturer's instructions (Novagen) and was purified by nickel chelate and ion-exchange chromatography as described for the A41 protein (Ng et al., 2001). The purified protein (100–200 μg with an estimated purity of 90%) was injected into New Zealand White rabbits to produce anti-C16 serum (Harlan Sera Laboratories).

**Immunoblotting.** Cells were infected with the indicated viruses at 10 p.f.u. per cell and cell lysates and supernatants were prepared as described previously (Bartlett et al., 2002). Antibodies used included rabbit polyclonal sera against VACV proteins C16 (1:2000) (see above), A41 (1:1000) (Ng et al., 2001) and A56 (1:1000), or mouse monoclonal antibody AB1.1 against D8 (1:1000) (Parkinson & Smith, 1994). Secondary antibodies and detection systems were as described previously (Bartlett et al., 2002).

**Cell fractionation.** BS-C-1 cells were infected at 0.1 p.f.u. per cell for 48 h, scraped from the dish and collected by centrifugation (800 g, 5 min). The cell pellet was resuspended in hypotonic buffer (10 mM Tris/HCl, pH 9.0) and disrupted by Dounce homogenization. The nuclei were removed by centrifugation (300 g, 5 min) and the cytoplasmic fraction (supernatant) was removed. Nuclei were washed four times in 10 mM Tris/HCl, pH 9 before incubation in RIPA buffer for 10 min. The mixture was centrifuged (1500 g, 10 min) and the supernatant retained as the nuclear fraction.

**Immunofluorescence.** HeLa cells were grown on sterilized glass coverslips (borosilicate glass; BDH) in six-well plates and infected at 5 p.f.u. per cell. Where indicated, 1 ng leptomycin B (Sigma-Aldrich) ml⁻¹ was added to the cells for 4 h (Wolff et al., 1997). Cells were washed, fixed, permeabilised and blocked as described previously (Chen et al., 2006). Cells were stained with anti-C16 (1:100) or anti-p65 (1:50; Santa Cruz) at room temperature for 1 h, followed by secondary antibody staining and mounting as described previously (Chen et al., 2006).

**Virus growth curves.** Monolayers of BS-C-1 cells were infected with either 10 or 0.01 p.f.u. per cell for measurement of one-step or multi-step growth kinetics, respectively, as described previously (Chen et al., 2006).

**Murine i.n. and intradermal models of infection.** Female BALB/c mice (n=5, 6–8 weeks old) were infected i.n. with 5×10³ p.f.u. and fluorescent microscopy and, after further rounds of plaque purification on BS-C-1 cells, were analysed by PCR to ensure replacement of both copies of the C16L gene with the Ecogpt/Egfp cassette. The resultant plaque-purified virus was called vAC16/Ecogpt/Egfp.
monitored as described previously (Williamson et al., 1990; Alcamí & Smith, 1992). Female C57Bl/6 mice (n=5, 6–8 weeks old) were inoculated intradermally (i.d.) in the ear pinnae with 1 × 10⁶ p.f.u. as described previously (Tscharke & Smith, 1999; Tscharke et al., 2002).

**Analysis of cell populations of infected organs.** Mice were infected i.n. with 1 × 10⁶ p.f.u. and at the indicated time post-infection (p.i.) were sacrificed, and the bronchial alveolar lavage (BAL) and lung tissue were processed as described previously (Clark et al., 2006; Jacobs et al., 2006). Live cells in BAL or lung samples were counted, washed with FACS buffer (0.1 % BSA, 0.1 % NaN₃ in PBS), blocked and stained with appropriate combinations of fluorescent isothiocyanate-, phycoerythrin- or tricolour-labelled antibodies. These were grouped into anti-CD25, anti-CD69, anti-CD3, anti-CD8, anti-CD4 for T lymphocytes, anti-Ly6G on neutrophils, anti-DX5 on natural killer cells, anti-F480 on macrophages and the relevant isotype controls (BD biosciences) as described previously (Tscharke & Smith, 1999; Tscharke et al., 2002). The presence of cell-surface markers was determined on a FACScan flow cytometer with CellQuest software (BD Biosciences). A lymphocyte gate was used to analyse data from at least 20 000 events.

**Murine vaccination and challenge analysis.** Female BALB/c mice (n=5, 6–8 weeks old) were vaccinated i.d. in the ear pinnae with 1 × 10⁶ p.f.u. of the indicated virus, and, 28 days later, were challenged i.n. with 5 × 10⁴ p.f.u. of VACV WR and monitored as described previously (Clark et al., 2006).

**Statistical analysis.** Student’s t-test (two tailed, unpaired) was used to examine the significance of raw data.

**RESULTS**

**Construction of C16L deletion mutant and revertant viruses**

To study the function of the C16 protein, a VACV strain WR mutant lacking both copies of the C16L gene, vAC16, was constructed (Methods) from plaque-purified VACV WR (vC16). The isolation of this virus demonstrated that the C16L gene was non-essential for virus replication. A revertant virus in which the C16L gene was reinserted into both ITRs, vC16Rev, was also constructed. PCR using primers for the C16L gene locus confirmed the presence of the C16L gene in vC16 and vC16Rev, and its absence in vAC16 (Supplementary Fig. S1 available in JGV Online). Analysis of genomic DNA by restriction enzyme digestion and agarose gel electrophoresis showed that the only discernible change between the viruses was caused by alteration to the C16L loci (data not shown).

**Analysis of C16 expression**

Immunoblot analysis using anti-C16 (Methods) identified 25 and 37 kDa proteins in extracts of cells infected with vC16 but not with vAC16 (Fig. 1a) or in mock-infected cells (Fig. 2a). These proteins were found in cell lysates and not in the concentrated culture supernatant, whereas the secreted protein A41 (Ng et al., 2001) was present in the culture medium. Notably, the 25 kDa C16 protein was present only late during infection and its formation was ablated in the presence of AraC.

To determine when C16 is expressed during infection, cells were infected in the presence or absence of cytosine arabinoside (AraC), an inhibitor of viral DNA replication and therefore late protein expression, and extracts of cells were analysed by immunoblotting (Fig. 1b). The 37 kDa C16 protein was detected from 2 h p.i., and at all times thereafter. Notably, it was expressed in the presence of AraC, indicating expression early during infection. In contrast, AraC blocked the expression of A56, a protein expressed predominantly late during infection (Brown et al., 1991). Notably, the 25 kDa C16 protein was present only late during infection and its formation was ablated in the presence of AraC.

The intracellular localization of the C16 protein was examined by biochemical fractionation of cells followed by immunoblotting (Fig. 2a) and immunofluorescence (Fig. 2b). Immunoblotting showed that cellular proteins tubulin and lamin A+C were present in the cytoplasmic or nuclear fraction, respectively, as expected. In contrast, the 37 kDa C16 protein was in both nuclear and cytoplasmic fractions after infection with vC16 and vC16Rev, but, even after longer exposure of the film, the 25 kDa C16 protein localized exclusively to the cytoplasm. For comparison, the VACV protein D8 was only in the cytoplasm.
Immunofluorescence also showed that C16 localized to the cytoplasm and nucleus [Fig. 2b(ii)]. There was minimal background immunofluorescence in vC16-infected cells (data not shown). The partial nuclear localization was investigated further using leptomycin B, an inhibitor of active nuclear export. In the presence of leptomycin B, the great majority of C16 and p65 were present in the nucleus [panels (v) and (vi)], suggesting active transport rather than diffusion through the nuclear pores (Fig. 2b). This localization pattern and active transport were similar in C16-transfected cells (data not shown).

**Computational analysis of the C16L gene**

The VACV WR C16L gene is present in both ITRs (GenBank accession nos YP_232892 and YP_233091) and is predicted to encode a protein without a transmembrane domain or signal peptide (www.poxvirus.org). Computational analysis found no cellular proteins with similarity to the VACV WR C16 protein, with the exception of the C-terminal peptide in the IL-1ra protein. However, C16 is highly conserved in several OPVs including variola virus (VARV), cowpox virus (CPXV), ectromelia virus (ECTV) and horsepox virus (HSPV). C16 is diploid in some strains of VACV, but not in other OPVs. In monkeypox virus (MPXV), camelpox virus (CMLV) and taterapox virus (TATV) the reading frame of the C16L orthologue is disrupted by mutation into shorter fragments.

The phylogenetic relationships of these proteins are shown in a rooted tree (Fig. 3a) produced from the aligned amino acid sequences. Group I proteins share 95–99 % amino acid identity and these proteins all have the C-terminal VTRFYF sequence, which is present in the IL-1ra protein (Kluczyk et al., 2004).

Another group of more distantly related OPV proteins, typified by protein C4 from VACV WR, was identified and is shown as group II (Fig. 3a). The C4 protein family is also conserved in several OPVs (VACV, VARV, CMLV, TATV, MPXV and CPXV) and these have 40–44 % amino acid identity to VACV WR C16. The degree of similarity between C16 and C4 is greatest in the C-terminal 100 aa, where these proteins share 55 % identity. Within the C4 group the proteins are highly conserved (95–99 % identity). Notably, C4 is encoded by CMLV, TATV and MPXV viruses that lack C16. The close similarity between the C4 and C16 proteins within OPVs suggest that they are likely to have been generated by a gene duplication event followed by diversification. The third group (III) of proteins (Fig. 3a) comprises more distantly related proteins from other chordopoxviruses including goatpox virus (GPXV), lumpy skin disease virus (LSDV), sheeppox virus (all genus Capripoxvirus), fowlpox virus (genus Avipoxvirus), Yaba monkey tumor virus (genus Yatapoxvirus) and deerpox virus (DPXV) (unclassified). These proteins have 26–37 % amino acid identity (49–61 % similarity) to C16, and are closer to C16 than C4. Within the groups II and III proteins, the IL-1ra-like sequence, VTRFYF, is partially conserved with the consensus sequence VT(R/K)-Y-

The expression of C16 by 14 VACV strains, two strains of CPXV and CMLV strain CMS was investigated by immunoblotting (Fig. 3b). Protein(s) recognized by the anti-C16 serum were expressed by all 14 VACV strains and both strains of CPXV, but not by CMLV or vC16. The latter observation demonstrated that the anti-C16 antibody does not recognize the related C4 protein encoded by VACV strain.
Infection with most viruses (but not VACV strains Dairen, Copenhagen and Tashkent) produced proteins of 37 and 25 kDa that were recognized by the anti-C16 antibody. These are both encoded by the C16L gene because they were absent in cells infected by vD16 and were expressed by vC16 and vC16Rev controls. The smaller polypeptide might be derived from the 37 kDa protein by proteolytic cleavage or (less likely) it might be generated by different translation initiation and its significance is unknown.

**C16 is non-essential in cell culture**

The isolation of vΔC16 demonstrated that C16 is not essential for VACV replication. To determine if deletion of...
C16L caused alterations in virus replication kinetics or spread, the growth of vAC16 was compared with wild-type and revertant controls. Plaques formed by vAC16 were slightly smaller ($P<0.05$) than controls in BS-C-1 and RK-13 cells (Fig. 4a). To investigate if the reduced plaque size was due to reduced virus titres or reduced virus spread, the replication kinetics were analysed after low (0.01) or high (10) m.o.i. However, no difference was observed between vAC16 and controls (Fig. 4b and data not shown). Virus plaque formation is influenced by the production of actin tails beneath cell-associated enveloped virus particles on the cell surface (Smith et al., 2002, 2003), and so the formation of actin tails by vAC16 was analysed by confocal microscopy. However, no difference between vAC16 and controls was observed (data not shown) and so the reason for the small difference in plaque size remains unknown.

### C16 affects virus virulence in the murine i.n. model

The virulence of vAC16 was compared with control viruses in two murine models. In an i.d. model, no significant difference in lesion size or cell recruitment was observed in animals infected with vAC16 at $10^4$ p.f.u. compared to control viruses (Supplementary Fig. S2 available in JGV Online and unpublished data). However, in an i.n. model, vAC16 induced significantly ($P<0.05$) less weight loss and milder signs of illness compared with controls (Fig. 5a and b) between days 6 and 11. Moreover, although all viruses replicated initially to similar titres (day 2) there was significantly ($P<0.05$) less infectious virus in lungs infected by vAC16 from day 4 p.i. onwards (Fig. 5c and d). This accelerated virus clearance after infection with vAC16 compared with controls suggested a more effective antiviral host response, and therefore the cellular inflammatory response in lungs was analysed by flow cytometry.

### C16 affects immune cell recruitment in the murine i.n. model

Cells in infected lungs and BAL were extracted, quantified by trypan blue exclusion and identified by flow cytometry (Fig. 6a). This revealed a statistically ($P<0.05$) higher number of cells in vAC16-infected BALs at day 3 p.i., compared with controls. By day 7, the cell numbers had equalized between the viruses, whereas at day 10, there were fewer lymphocytes present in the BAL of mice infected with vAC16. To test whether this early increase in lymphoid cells was due to an enhanced recruitment of a particular lymphoid subset, the percentage of macrophages (Fig. 6b), neutrophils (Fig. 6c), natural killer cells and CD4$^+$ and CD8$^+$ T lymphocytes (data not shown) present was analysed. This showed a slightly greater number of all cell types examined in the vAC16-infected BAL at day 3, but there was no dramatic increase in any lymphoid subset.

Next, the recruitment and activation of T cells (CD3$^+$) in the lungs was investigated by staining for CD3, CD4, CD8 and CD69 (Fig. 6d–f). Natural killer cell (DX5$^+$) recruitment in the lungs was also analysed and showed no significant variation between vAC16 and control viruses (Supplementary Fig. S3 available in JGV Online). T-cell analysis showed that at day 3 more of the CD4$^+$ and CD8$^+$ T cells, recruited to the site of infection with vAC16, were activated (CD69$^+$) compared with controls. By day 7, there was no difference in the number of activated T lymphocytes after infection with the different viruses. However, by day 10 there was a reduction in CD4$^+$ and CD8$^+$ T cells present.
after infection with vΔC16. One interpretation of these data is that the host response to infection with vΔC16 is accelerated compared with controls, so that early p.i. there are more cells in vΔC16-infected tissues and these show a greater degree of activation, whereas at late times there are fewer cells because the infection is being cleared sooner. By inference, the C16 protein may function to diminish the innate response to infection.

Vaccination and challenge with vΔC16

To characterize further the immune response to infection with vΔC16, the effectiveness of this virus as a vaccine was compared with that of the control viruses (Fig. 7). Mice were vaccinated i.d. with the indicated viruses and were challenged with VACV WR 28 days later. The challenge dose was 500 LD₅₀ for BALB/c mice of this age and hence even vaccinated mice began to lose weight rapidly. Weight loss reached its maximum at day 3 post-challenge when the average weight loss of the three immunized groups (vC16, vΔC16 and vC16Rev) was 20–21%. However, under the conditions tested there was no difference in the protection afforded by immunization with vΔC16 compared to the control viruses.

DISCUSSION

A characterization of the C16 protein from VACV strain WR and the effects of this protein on virus replication, virulence and immunogenicity are reported. Bioinformatic analysis indicated that very closely related proteins (95–99% amino acid identity) are encoded by five OPV species and that the gene is present, but disrupted, in three others. In addition, C16 is related to another family of OPV proteins exemplified by C4 from VACV WR, and a more distantly related group from other chordopoxviruses. C16 was expressed by all VACV and CPXV strains tested and was synthesized early during infection. This early expression profile is consistent with the analysis of the C16L promoter (data not shown) and a genome-wide transcriptome analysis that detected C16 mRNA by 1 h p.i. (Assarsson et al., 2008). C16 is shuttled actively between the cytoplasm and nucleus but lacks a recognizable nuclear localization signal and nuclear export signal, suggesting it might be transported in a complex with another protein. The partial nuclear localization of C16 is interesting because it is one of only a few nuclear proteins encoded by VACV, which replicates in the cytoplasm. Another example is the E3 protein (Yuwen et al., 1993) that binds dsRNA (Chang et al., 1992) and contributes to virulence (Brandt & Jacobs, 2001). In contrast, a group of other VACV proteins that affect transcription of host nuclear factor kappa B (NF-κB)-responsive genes function from within the cytoplasm (Bowie et al., 2000; Bartlett et al., 2002; Shisler & Jin, 2004; Chen et al., 2006, 2008). There are also reports of endogenous nuclear proteins being recruited to the cytoplasm during VACV infection (Oh & Broyles, 2005).

The intracellular location of the C16 protein makes its postulated function as an IL-1ra-like protein that mediates extracellular blockade of signalling from the IL-1R (Kluczyk et al., 2002, 2004) unlikely. C16 would
only be in a position to modulate signalling from the IL-1R after release due to lysis of infected cells, and while this might happen very late during infection, such a function is unlikely to be its primary role given its early expression. However, there are also intracellular (ic) IL-1ra proteins, produced by differential splicing (Butcher et al., 1994; Malyak et al., 1998a, b) and whose functions remain poorly defined. These icIL-1ra isoforms were reported to downregulate the IL-1 pathway (Watson et al., 1995; Arend et al., 1998; Banda et al., 2005). Moreover, the precursor form of IL-1α, pro-IL-1α, can localize to (Wessendorf et al., 1993) and function within the nucleus (Cheng et al., 2008) and icIL-1ra may regulate these functions. It is therefore more likely that C16 functions by mimicking icIL-1ra. Interestingly, the deerpox virus protein 054 shares 89% amino acid identity with the secreted IL-1ra protein from Bos Taurus (Afonso et al., 2005) and is substantially more likely to mimic extracellular IL-1ra activity than C16, but does not have a counterpart in VACV or other OPVs.

VACV already targets the IL-1 pathway by several mechanisms, indicating the importance of IL-1 against poxvirus infections. VACV restricts the formation of IL-1β in infected cells by expressing a caspase 1 inhibitor (Dobbelstein & Shenk, 1996; Kettle et al., 1997) and blocks the function of IL-1β systemically by the expression of a soluble IL-1βR (Alcamí & Smith, 1992; Spriggs et al., 1992; Alcamí & Smith, 1996). Furthermore, VACV expresses intracellular proteins A46 (Bowie et al., 2000; Stack et al., 2005), A52 (Bowie et al., 2000; Harte et al., 2003; Graham et al., 2008), N1 (Bartlett et al., 2002; DiPerna et al., 2004; Cooray et al., 2007; Graham et al., 2008) and B14 (Chen et al., 2006, 2008) that inhibit IL-1x- or IL-1β-induced signalling from the IL-1R that would otherwise activate NF-kB.

A virus deletion mutant, vΔC16, formed a slightly smaller plaque when compared with wild-type and revertant controls, but replicated normally in the cell lines analysed and in vivo to reach equivalent titres at 2 days p.i. However, there was no defect in virus-induced actin tail formation, a pre-requisite for efficient cell-to-cell spread. Hence, the basis for this small
in vivo inflammatory response produced a slightly small plaque in cell culture and an altered explanation that B14 interacts with the IκB kinase complex and inhibits NF-κB activation (Chen et al., 2008). Most VACV immunomodulatory proteins, like B14, are expressed early during infection and the early expression of C16 would be consistent with such a role.

The deletion of C16 from the VACV WR genome reduced virus virulence in a murine i.n. model (systemic infection) but not in an i.d. (local infection) model. This attenuation was characterized by significant reductions in virus-induced weight loss, signs of illness and virus titres. Although, vΔC16 virus titres in infected lungs were equivalent to control virus-infected lungs at 2 days p.i., at later times the virus titres were considerably reduced in vΔC16-infected tissues. This indicated that, in the absence of C16, the virus can replicate efficiently in vivo, but, in the absence of C16, virus clearance is accelerated.

Investigation of this attenuated phenotype by analysis of lymphoid cells within infected tissue indicated a difference in the number and properties of the infiltrating cells and the kinetics of their recruitment. At 3 days p.i., the number of infiltrating cells was greater after infection with vΔC16, and more CD4+ and CD8+ T cells were activated (CD69+). By day 7, the number of cells present and the proportion that were activated were equivalent for all viruses. However, by day 10 the total number of infiltrating cells and the number of CD4+ and CD8+ T cells were reduced after infection with vΔC16, consistent with more rapid resolution of the infection. The accelerated recruitment and activation of T cells early after infection fits with the diminished virus titres observed from day 4 p.i., and with the reduced weight loss and signs of illness. The recruitment of leukocytes to sites of inflammation is dependent on the expression of chemokines and cytokines. It follows that C16 is either directly or indirectly influencing the expression or function of such inflammatory mediators. The intracellular location of C16 suggests that it might mediate such an effect by modulating intracellular signalling pathways and this possibility is currently under investigation.

Despite the reduction of virulence of vΔC16 in the i.n. model and the difference in T-cell recruitment and activation, vΔC16 was as effective as wild-type and revertant controls in inducing protective immunity against subsequent virus challenge with an otherwise lethal dose of VACV strain WR. The vΔC16 phenotype (reduced virulence yet undiminished immunogenicity) make VACV strains lacking the C16L gene attractive vaccine candidates.

In summary, C16 is an intracellular virulence factor, which modulates the host response to infection. Investigation of its role in virus virulence should provide further insight into host–virus interactions and viral immunomodulation.

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