Antibody limits in vivo murid herpesvirus-4 replication by IgG Fc receptor-dependent functions

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Antibody is an important antiviral defence. However, it is considered to do little against human gamma-herpesviruses, which establish predominantly latent infections regulated by T cells. One limitation on analysing these infections has been that latency is already well-established at clinical presentation; early infection may still be accessible to antibody. Here, using murid herpesvirus-4 (MuHV-4), we tested the impact of adoptively transferred antibody on early gamma-herpesvirus infection. Immune sera and neutralizing and non-neutralizing monoclonal antibodies (mAbs) all reduced acute lytic MuHV-4 replication. The reductions, even by neutralizing mAbs, were largely or completely dependent on host IgG Fc receptors. Therefore, passive antibody can blunt acute gamma-herpesvirus lytic infection, and does this principally by IgG Fc-dependent functions rather than by neutralization.

INTRODUCTION

Antibody provides an important defence against many epidemic viruses. Its protective efficacy correlates with the neutralization of cell-free virions (Zinkernagel & Hengartner, 2006). This is reflected epidemiologically in virus strains that differ mainly where neutralizing antibodies bind. Whether antibody is also effective against persistent pathogens such as gamma-herpesviruses is less clear. The archetypal gamma-herpesvirus, Epstein–Barr virus (EBV), causes disease mainly when latent; inhibiting its lytic replication during acute infectious mononucleosis makes little difference to latent viral loads (Yao et al., 1989); also, long-term latent infection appears to be controlled by T cells (Rickinson & Moss, 1997). Therefore, antibody would seem to play little part in host defence. However, EBV infection takes at least 1 month to become symptomatic (Hoagland, 1964), by which time latency is already well-established. The early events in infection remain poorly defined. Substantial CD8+ T-cell responses to viral lytic antigens at clinical presentation (Callan et al., 1996) suggest that significant lytic replication occurs early on, and vaccination with a viral lytic antigen can alleviate the symptoms of primary infection (Sokal et al., 2007). Therefore, antibody might still be useful therapeutically in acute settings such as organ transplantation into EBV-naive recipients, where primary EBV infection confers a poor outcome (Walker et al., 1995).

Genetic immunodeficiencies can provide clues as to what controls human viruses (Carneiro-Sampaio & Coutinho, 2007). However, B cell-deficient humans lack the main EBV latency reservoir and consequently fail to support normal persistence (Faulkner et al., 1999). The contribution that antibody makes to host defence has therefore been hard to define. The failure of EBV to establish a significant experimental infection of non-primates further limits the opportunities for direct analysis. We and others (Virgin & Speck, 1999; Nash et al., 2001; Stevenson & Efstathiou, 2005) have therefore developed murid herpesvirus-4 (MuHV-4) as a small-animal model of gamma-herpesvirus pathogenesis. MuHV-4 is genetically closer to Kaposi’s sarcoma-associated herpesvirus than to EBV, but its tropism and pathogenesis are similar to those of EBV, and it currently provides the most tractable means to establish how antibody might interact with a gamma-herpesvirus in vivo.

T-cell depletion causes disease in B cell-deficient (Stewart et al., 1998) but not immunocompetent (Stevenson et al., 1999) MuHV-4 carriers, implying that antibody helps to control established infection. In support of this idea, adoptively transferred immune sera reduced MuHV-4 replication in B cell-deficient (Gangappa et al., 2002) and T cell-depleted, CD28-deficient (Kim et al., 2002) mice, and a rabbit serum raised against the MuHV-4 gp70 complement control protein reduced MuHV-4 replication in B cell-deficient mice (Gangappa et al., 2002). However, human gamma-herpesviruses cause disease more by latency-associated tumours rather than by chronic lytic replication, so the main opportunity for antibody to protect against them is likely to be early on. The impact of antibody on early MuHV-4 replication has not been tested.

The mechanism of antibody-mediated protection was not addressed in the MuHV-4 studies listed above, but was
speculated to be neutralization in one study (Kim et al., 2002) and correlated with in vitro virus neutralization in the other (Gangappa et al., 2002). However, gamma-herpesvirus neutralization is not well understood. For example, EBV can be neutralized in vitro (Thorley-Lawson & Geilinger, 1980), but EBV carriers continue to transmit infection with little evidence of viral antigenic variation, despite making virus-specific antibodies. Immune sera can block MuHV-4 infection of fibroblasts in vitro (Stevenson & Doherty, 1998), but they block in vitro Immune sera can block MuHV-4 infection of fibroblasts in vivo host entry poorly (Gillet et al., 2007a) and tend to enhance MuHV-4 infection of IgG Fc receptor-positive (FcR⁺) cells such as macrophages (Rosa et al., 2007). Thus, whilst immune sera can block virions binding to fibroblasts (Gill et al., 2006), virion membrane fusion seems to remain intact. Poor in vivo neutralization could consequently reflect the availability of additional binding routes that are harder for antibody to block. mAbs to gH–gL and gB can block infection post-binding and achieve a more universal neutralization (Gill et al., 2006; Gillet et al., 2006), but gH–gL and gB are poorly immunogenic, so this activity of immune sera is weak (Gillet et al., 2007b). The virion fusion machinery – gH and gB – is also protected by glycosylation (Gillet & Stevenson, 2007a) and by gB and gH–gL associating on the virion surface (Gillet & Stevenson, 2007a; b; Gillet et al., 2009). The gB–gH–gL complex appears to unfold only after a post-endocytic dissociation of gL (Gillet et al., 2008a, b), so extracellular gB and gH can remain protected. Another consideration is that in vivo gamma-herpesvirus lytic spread may involve direct cell–cell contact more than cell-free virion release, and thereby resist neutralization as described for other herpesviruses (Hooks et al., 1976).

We asked here whether immune sera or monoclonal antibodies (mAbs) could reduce the primary lytic replication of MuHV-4 in immunocompetent hosts and, if so, how they might act. Our aim was to define a basis for antibody-based interventions in primary gamma-herpesvirus infections.

**METHODS**

**Mice.** BALB/c, C57BL/6, 129Sv (Harlan UK) and 129Sv x B6 FcRγ²−/− mice (Taconic Europe) were bred in the Cambridge University Department of Pathology animal unit and infected with MuHV-4 when 6–12 weeks old. Intranasal infections were performed in a volume of 30 μl, given under general anaesthesia. For luciferase imaging, mice were injected intraperitoneally with luciferin (2 mg per mouse), anaesthetized with isoflurane, then scanned with an IVIS Lumina (Caliper Life Sciences). Quantitative comparisons used the total radiance (photons s⁻¹ cm⁻²) over each region of interest, relative to a negative-control region. All experiments conformed to local animal-ethics regulations and to Home Office Project Licence 80/1992.

**Sera and mAbs.** Immune sera were collected at least 3 months post-infection and pooled from 10–12 mice each. Hybridomas were derived from MuHV-4-immune mice by fusion with NSO myeloma cells (Kohler & Milstein, 1975), except for mAb DW-6F6, which was derived from mice infected with influenza A/PR/8/34 and is specific for the influenza H1 haemagglutinin. Antibodies were concentrated from hybridoma supernatants by ammonium sulfate precipitation, dialysed against PBS, isotyped by ELISA (Sigma) and quantified by Mancini assay (Mancini et al., 1965). Those used for in vivo protection are listed in Table 1.

**Cells and viruses.** BHK-21 fibroblasts (ATCC CCL-10) and NMuMG epithelial cells (CRL-1636) were grown in Dulbecco’s modified Eagle’s medium (DMEM) with 2 mM glutamine, 100 U penicillin ml⁻¹, 100 μg streptomycin ml⁻¹ (all from Sigma) and 5 % fetal calf serum (PAA Laboratories). Hybridomas were grown in RPMI medium (Sigma), supplemented as for DMEM. All viruses were derived from a MuHV-4 bacterial artificial chromosome-cloned genome (Adler et al., 2000). The wild type and its luciferase-expressing derivative (Milho et al., 2009) were grown and titrated in BHK-21 cells (de Lima et al., 2004). Infected cells were freeze–thawed

**Table 1. Summary of mAbs used in vivo**

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Target</th>
<th>Isotype</th>
<th>Neutralization*</th>
<th>In vivo titre†</th>
</tr>
</thead>
<tbody>
<tr>
<td>LT-6E8</td>
<td>gp70 (SCR1 + 2)</td>
<td>IgG₂b</td>
<td>−</td>
<td>+</td>
</tr>
<tr>
<td>58-16D2</td>
<td>gp70 (SCR4)</td>
<td>IgG₂a</td>
<td>−</td>
<td>+</td>
</tr>
<tr>
<td>3F7</td>
<td>gN</td>
<td>IgG₂a</td>
<td>−</td>
<td>+</td>
</tr>
<tr>
<td>SC-9A5</td>
<td>gB</td>
<td>IgG₃</td>
<td>+ +</td>
<td>+</td>
</tr>
<tr>
<td>MG-2C10</td>
<td>gB</td>
<td>IgM</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>BN-3A4</td>
<td>gp150</td>
<td>IgG₁</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
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<td>IgG₂a</td>
<td>−</td>
<td>−</td>
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<tr>
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<td>gp150</td>
<td>IgG₂a</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>T1A1</td>
<td>gp150</td>
<td>IgG₂a</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>230-5B2</td>
<td>gH–gL</td>
<td>IgG₂a</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>T2C12</td>
<td>gH–gL</td>
<td>IgG₂a</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>230-4A2</td>
<td>gH–gL</td>
<td>IgG₂a</td>
<td>+</td>
<td>−</td>
</tr>
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</table>

*Plaque reduction on BHK-21 cells, scored from negative (−) to strong (++)
†Reduction in lung titre by plaque assay, scored from negative (−) to strong (++)
and virions concentrated by high-speed centrifugation (38 000 g, 90 min). Cell debris was then removed by low-speed centrifugation (1000 g, 10 min).

**Virus titres and neutralization assays.** MuHV-4 was titrated by plaque assay on BHK-21 cells (de Lima et al., 2004). Infectious virus in ex vivo tissues was measured by plaque assay of freeze–thawed tissue homogenates. To determine virus titre in noses, we removed a block of tissue bounded by the cartilaginous tip of the nose anteriorly, the orbits posteriorly, the zygomatic arches laterally, the palate ventrally and the nasal bones dorsally. Cell monolayers were incubated with virus (2 h, 37 °C), overlaid with 0.3% carboxymethylcellulose and, 4 days later, fixed (4% formaldehyde) and stained (0.1% toluidine blue) for plaque counting. Titres were compared statistically between experimental groups by unpaired, two-tailed t-tests, assuming equal variance. The calculations were performed in Excel. P values of <0.05 were considered statistically significant.

**Flow cytometry.** MuHV-4-infected cells (2 p.f.u. per cell, 18 h) were trypsinized, washed in PBS and incubated (1 h, 4 °C) with MuHV-4 glycoprotein-specific mAbs, followed by fluorescein-conjugated rabbit anti-mouse IgG pAb (Dako Cytomation). The cells were washed twice in PBS after each incubation and analysed on a FACSCalibur (BD Biosciences).

**Immunoblotting.** N-Linked glycans were removed with protein N-glycanase F; O-linked glycans were removed with sialidase A + O-glycanase according to the manufacturer’s instructions (Prozyme). Deglycosylated and intact samples were denatured (95 °C, 5 min) in Laemmli’s buffer (60 mM Tris-Cl (pH 6.8); 3% (v/v) SDS; 5% (v/v) 2-mercaptoethanol; 10% (v/v) glycerol; 0.002% (v/v) bromophenol blue), resolved by SDS-PAGE and transferred to PVDF membranes. The membranes were blocked with 10% non-fat milk and probed with MuHV-4-specific mAbs plus horseradish peroxidase-conjugated rabbit anti-mouse IgG pAb (Dako Cytomation). Membranes were washed four times in PBS/0.1% Tween 20 after each step. Bound antibody was visualized by ECL substrate development (APBiotec).

**RESULTS**

**Immune sera reduce MuHV-4 replication in naive hosts**

Adoptively transferred immune sera can reduce MuHV-4 replication in persistently infected mice that lack normal antibody production (Gangappa et al., 2002; Kim et al., 2002). We wanted to know whether immune sera could also reduce acute viral replication in naive, immunocompetent hosts. We tested this by giving MuHV-4-immune sera intraperitoneally to naive mice at the same time as intranasal MuHV-4 virions (Fig. 1). Significant reductions in virus titres were observed in both C57BL/6 (Fig. 1 a, b) and BALB/c (Fig. 1c) mice and across multiple time points. Naive sera had no effect (Fig. 1c).

We used luciferase-positive MuHV-4 (Milho et al., 2009) as an alternative means of tracking in vivo lytic infection (Fig. 2a, b). Immune sera again reduced significantly the extent of acute host colonization. This was evident both in lungs and in noses. Plaque assays (Fig. 2c) confirmed a

![](image)

**Fig. 1.** Protection against MuHV-4 replication by immune sera. (a) C57BL/6 mice were given 3×10⁴ p.f.u. MuHV-4 intranasally (i.n.) and at the same time treated or not with 200 µl immune serum intraperitoneally (i.p.). Infectious virus in lungs was measured by plaque assay 3 days later. Each point shows the titre for one mouse; x shows mean values. Immune serum reduced lung titres significantly (P<0.0005 by two-tailed Student’s t-test). (b) C57BL/6 mice were infected and treated or not with immune serum as in (a). The effects of immune serum on day 3 and day 6 titres were then compared. The reduction in titre was significant at day 6 (P<0.0001), but not at day 3 (P=0.06). (c) BALB/c mice were infected and given immune serum as in (a), or naive serum as an additional control. Virus titres in lungs were determined after 5 days. Immune serum reduced titres significantly compared with either virus only or naive serum (P<0.01). Naive serum had no effect compared with virus only (P=0.5).
reduction in nose infection by immune serum. In other experiments, we obtained similar data for noses and lungs, but nose titres were generally more variable between individual mice.

Non-neutralizing mAbs reduce MuHV-4 replication in naive hosts

Since the complexity of immune sera makes mechanisms of protection difficult to establish, we used mAbs to explore further the interaction between antibody and MuHV-4. Table 1 summarizes the results. Fig. 3 shows neutralization assays. In Fig. 4(a), mAb LT-6E8 reduced MuHV-4 replication compared with a control mAb (influenza haemagglutinin-specific). LT-6E8 recognizes gp70, the main product of MuHV-4 ORF4 (Gillet et al., 2007c) and the target used by Gangappa et al. (2002) for protection by immune serum. mAb 58-16D2, which recognizes a different gp70 domain (Gillet et al., 2007c), was also effective (Fig. 4b). As expected, neither LT-6E8 nor 58-16D2 affected the replication of MuHV-4 lacking ORF4 (Fig. 4b).

The amounts of mAb used depended on what each hybridoma produced. In general, we used the maximum amount possible, so as not to miss a therapeutic effect. We also ensured that at least 200 μg of any ineffective mAb was given, so as to be sure that ineffectiveness did not reflect
low antibody concentration. Titration of LT-6E8 (Fig. 4c), a standard effective mAb, showed that even 20 μg antibody reduced MuHV-4 replication significantly.

LT-6E8 blocks heparan sulfate binding by gp70, but does not neutralize (Fig. 3), because virions can also bind to heparan sulfate via gH–gL. mAb 230-4A2 blocks heparan sulfate binding by gH–gL and so neutralizes synergistically with LT-6E8 (Gillet et al., 2008c). However, LT-6E8 and 230-4A2 gave no synergistic reduction in in vitro virus titres (Fig. 4d). mAb MG-2C10 (Gillet et al., 2006), a neutralizing, gB-specific IgM (Fig. 3), had no effect on in vitro virus replication (Fig. 4e). mAbs 3F7 (gN-specific, non-neutralizing) (May et al., 2005a) and SC-9A5 (gB-specific, neutralizing) both reduced in vivo lytic replication, whilst mAb BN-3A4 (gp150-specific, non-neutralizing) did not (Fig. 4f). Therefore, no correlation, either positive or negative, was evident between an antibody’s capacity to neutralize cell-free virions in vitro and its capacity to reduce virus replication in vivo.

Reduced lytic infection by antibody depends on host IgG Fc receptors

The antiviral effect of IgG occurs principally by Fab-dependent virion neutralization and by Fc-dependent recruitment of other immune effectors. Recruiting cellular effectors requires signalling through the common IgG Fc receptor gamma chain (FcyRγ) (Nimmerjahn & Ravetch, 2008). We therefore explored the mechanism of antibody-mediated protection by infecting FcγRγ−−− mice. We used mice that also lacked the FcgRII inhibitory receptor (FcγRγ−−−/FcgRII−−−), so as to reduce any excess immunosuppressive effect of IgG binding to cells lacking activating Fc receptors. mAbs 3F7 and LT-6E8 both failed to reduce virus titres significantly in the FcγR− mice, whereas they were effective in FcγR+ 129Sv controls (Fig. 5a). We obtained an equivalent result (Fig. 5b) with C57BL/6 controls (the FcγR− mice were 129Sv × C57BL/6) with mAb 58-16D2 (Fig. 5c), and with immune sera using both 129Sv and C57BL/6 controls (Fig. 5c, d). Therefore protection by both non-neutralizing mAbs and immune sera depended on IgG Fc receptors.

Neutralization has at most a minor role in antibody-mediated MuHV-4 control

As mAb SC-9A5 is an IgG3, it should not engage strongly with IgG Fc receptors (Nimmerjahn & Ravetch, 2008). Therefore, as a neutralizing mAb (Fig. 3), it might be expected to reduce virus titres equally in FcγR− and FcγR+ mice. In one experiment (Fig. 6a), SC-9A5 reduced the virus titres of FcγR− mice, but not to a significant level; in a repeat experiment (Fig. 6b), the reduction was significant, although there was also a significant reduction by mAb 3F7 (P<0.02), something not seen in other experiments (e.g. Figs 5a, 6a). In both experiments, SC-9A5 gave a substantially greater reduction in FcγR+ than FcγR− mice (4.6-fold rather than 1.9-fold in Fig. 6a; 17-fold rather than 3.7-fold in Fig. 6b).

We examined further a possible contribution of neutralization to in vivo titre reductions with the gH–gL-specific neutralizing mAbs T2C12 and 230-5B2 (Fig. 6c). Again, there were substantial reductions in FcγR+ mouse titres, and only small reductions that did not reach statistical significance in FcγR− mouse titres. Therefore, even neutralizing mAbs acted in vivo largely through IgG Fc receptor engagement.

gp150 is not an effective target for MuHV-4 control by mAbs

gp150 is expressed abundantly on MuHV-4-infected fibroblasts and epithelial cells. Therefore, it was surprising that the gp150-specific mAbs BN-3A4 (Fig. 4f), T7F5 (Fig. 6a) and 150-5A10 (Fig. 7a) all failed to reduce virus titres. These mAbs all recognize epitopes contained within gp150 aa 108–151 (Gillet et al., 2007b). mAb T1A1, which
Fig. 4. Reduction in in vivo MuHV-4 infectivity by glycoprotein-specific mAbs. (a) C57BL/6 mice were infected i.n. with MuHV-4 (3×10⁴ p.f.u.) and at the same time given antibody i.p. (500 μg). Infectious virus titres in lungs were then determined by plaque assay. Each point shows the titre for one mouse; x shows mean values. The gp70-specific mAb LT-6E8 reduced virus titres significantly at day 5 (P<0.005 by Student's two-tailed t-test) compared with the influenza haemagglutinin (flu HA)-specific control, although not at day 3 (P=0.1). (b) In an equivalent experiment, mAb LT-6E8 did not affect the replication of gp70⁻ MuHV-4 (P=0.2). mAb 58-16D2 (50 μg), which recognizes a different domain of gp70, also significantly reduced the 5 day post-infection titres of wild-type (wt; P<0.00002) but not gp70⁻ (P=0.8) MuHV-4. (c) BALB/c mice were infected i.n. with MuHV-4 (3×10⁴ p.f.u.) and at the same time given LT-6E8 i.p. in differing amounts. Infectious virus titres in lungs were determined by plaque assay 5 days later. Each point shows the titre for one mouse; x shows mean values. All LT-6E8 amounts reduced virus titres compared with the no-antibody control (P<0.003; 500 μg was marginally more effective than 56 or 19 μg (P<0.04), but no more effective than 167 μg (P=0.3). (d) BALB/c mice were infected i.n. with MuHV-4 (3×10⁴ p.f.u.) and at the same time given mAb LT-6E8 (500 μg) or mAb 230-4A2 (500 μg) i.p., both together or no antibody. Infectious virus titres in lungs were determined by plaque assay 5 days later. All mAb treatments reduced virus titres significantly compared with the no-antibody control (P<0.02). Both LT-6E8 and 230-4A2 together were more effective than LT-6E8 alone (P<0.03), but not more effective than 230-4A2 alone (P=0.8). (e) C57BL/6 mice were infected i.n. as in (a), and the effect of immune serum (200 μl) was compared with that of the neutralizing gB-specific IgM mAb MG-2C10 (200 μg). Immune serum reduced virus titres significantly (P<0.001), whereas MG-2C10 had no effect (P=0.7). (f) Mice were infected i.n. as in (a) and at the same time given antibody or not i.p. (results of two separate experiments are shown). Infectious virus in lungs was titrated by plaque assay 5 days later. mAbs 3F7 (IgG2a, anti-gN, non-neutralizing, 40 μg) and SC-9A5 (IgG2a, anti-gB, neutralizing, 500 μg) reduced virus titres (P<0.002), whereas mAb BN-3A4 (IgG1, anti-gp150, non-neutralizing, 700 μg) did not (P=0.9).
recognizes an epitope in the region 151–270, also gave no reduction (data not shown).

gp150 is predicted to be heavily O-glycosylated (Fig. 7b). All of the gp150-specific mAbs used recognize glycan-independent epitopes, as they recognize gp150 fragments expressed in Escherichia coli (Gillet et al., 2007b and data not shown), but glycans could potentially hide their epitopes. We have found previously (Gillet & Stevenson, 2007a) that O-glycosylation inhibits gB recognition by mAb MG-2C10. This manifests as virions from NMuMG epithelial cells resisting recognition by MG-2C10 unless O-
glycans are removed, whereas virions from BHK-21 fibroblasts are recognized with or without deglycosylation. (Cells of epithelial origin presumably add more extensive, mucin-like O-glycans.) Flow cytometry of MuHV-4-infected NMuMG cells showed less staining than infected BHK-21 cells for all gp150-specific mAbs (Fig. 7c). There was also less gp150 detected by immunoblot of NMuMG cell-derived compared with BHK-21 cell-derived virions (Fig. 7d). The mAbs used recognize no products of the corresponding gp150+ or gp70− viruses; the bands below 70 kDa in Fig. 7(e) are all alternative glycoforms of gp70 (Gillet et al., 2007c). Reduced gp150 detection on NMuMG cell-derived virions could reflect a lower gp150 content. However, these virions do not show the typical gp150-deficient phenotype (de Lima et al., 2004) of heparan sulfate-independent infection (data not shown). Moreover,

**Fig. 6.** Antibody-mediated protection by neutralizing mAbs is also largely IgG Fc receptor-dependent. (a) C57BL/6 (IgG FcR+/+) or FcR−/−;RNL−/− (IgG FcR−/−) mice were infected with MuHV-4 i.n. (3×10⁴ p.f.u.) and given antibody or not i.p. Five days later, infectious virus in lungs was titrated by plaque assay. Each point shows the titre for one mouse; x shows mean values. IgG FcR+/+ mice showed a significant reduction in virus replication by mAbs SC-9A5 (500 µg) and 3F7 (40 µg) (P<0.001 by Student’s two-tailed t-test), but not by mAb T7F5 (200 µg) (P=0.2). None of the mAbs reduced virus replication significantly in IgG FcR−/− mice. (b) A repeat experiment again showed a significant reduction in IgG FcR+/+ lung titres by mAbs 3F7 (40 µg) and SC-9A5 (500 µg) (P<0.0003). This time the reduction in IgG FcR−/− lung titres by SC-9A5, whilst lower than that of IgG FcR+/+ (3.7-fold rather than 17-fold), was also significant (P<0.0003), as was that by 3F7 (P<0.02). (c) IgG FcR+/+ and IgG FcR−/− mice were further compared for virus titre reductions by the neutralizing gH−gL-specific mAbs T2C12 (500 µg) and 230-5B2 (200 µg), using the same infection protocol as in (a). Both mAbs reduced lung virus titres significantly in IgG FcR+/+ (P<0.005) but not IgG FcR−/− (P>0.13) mice.
gp150 recognition was not reduced uniformly: recognition by T1A1 was relatively preserved, whereas recognition by T7F5 was reduced markedly. Therefore, it seemed more likely that gp150 recognition was being masked.

gp150 has at least some some O-linked glycans, because it is immunoprecipitated readily by jacalin (May et al., 2005b). Removing O-linked glycans did not increase convincingly the detection of NMuMG-cell-derived gp150 (Fig. 7e). However, the detection of BHK-21 cell-derived gp150 decreased markedly, presumably because gp150 became unstable (or was otherwise lost) when its O-linked glycans were removed. (Note that the actual molecular mass of gp150 is very difficult to infer from SDS-PAGE: O-glycan digestion retards its migration even further.) The greater equivalence of NMuMG and BHK-21...

**Fig. 7.** O-Glycosylation may make gp150 a poor therapeutic antibody target. (a) C57BL/6 (IgG FcR+/+) or FcR−/−FcγRII−/+ (IgG FcR−/−) mice were infected i.n. with MuHV-4 (3×10⁴ p.f.u.) and given antibody or not i.p. Five days later, infectious virus in lungs was titrated by plaque assay. Each point shows the titre for one mouse; × shows mean values. IgG FcR+/+ mice showed a significant reduction in virus replication by mAb LT-6E8 (500 μg) (P<0.002), but not by mAb 150-5A10 (200 μg) (P=0.9), using Student's two-tailed t-test. Neither mAb reduced virus replication significantly in IgG FcR−/− mice. (b) Predicted O-glycosylation sites in gp150 (Julenius et al., 2005). The predicted signal sequence (residues 1–22) and transmembrane domain (residues 459–481) are also shown. (c) Flow-cytometric analysis of BHK-21 cells (solid lines) and NMuMG cells (dashed lines) infected with MuHV-4 (2 p.f.u. per cell, 18 h). Control=secondary antibody only. mAbs 8C1 (anti-gH) and BN-1A7 (anti-gB) provide controls for the level of infection. None of the mAbs stained uninfected cells. Staining with mAb 150-5A10 was equivalent to that of BN-5A4. (d) Immunoblot of virions derived from NMuMG cells (NMu) or BHK-21 cells (BHK). Each blot shows two independently grown virus stocks from each cell type. mAb 150-7D1 recognizes the ORF17 capsid component and provides a loading control. (e) Virions derived from NMuMG cells or BHK-21 cells were untreated (nil) or digested with PNGase F to remove N-linked glycans (N−) or digested with sialidase + O-glycanase to remove O-linked glycans (O−). Samples were then immunoblotted for gp150. mAb 150-7D1 (ORF17) provides a loading control; mAb 58-16D2 demonstrates O-linked glycan removal from gp70 in the same samples.
gp150 signals after O-linked glycan removal suggested that glycosylation substantially modified recognition by gp150-specific mAbs. Therefore, differential glycosylation could explain the inefficacy of gp150-specific mAbs in reducing in vivo virus titres.

**DISCUSSION**

 Passive antibody, given promptly, attenuates primary alphaherpesvirus infections (Whitley, 1994; Ogilvie, 1998). Gamma-herpesviruses are generally viewed as less accessible to antibody-based interventions, as they colonize their hosts largely by latency-associated lymphoproliferation. We show here that immune sera and mAbs can nonetheless reduce MuHV-4 acute lytic replication and that they act predominantly by IgG Fc receptor engagement.

Antibody can protect mice against herpes simplex virus (HSV) both by neutralization and by IgG Fc receptor-dependent functions (McKennald, 1985; Ishizaka et al., 1995). Early studies mainly compared neutralizing and non-neutralizing mAbs. Whilst antibodies capable of neutralization need not act by neutralization, the finding that glycoprotein C was a viable target for antibody in vivo and not for neutralization (Balachandran et al., 1982; Kümel et al., 1985) argued that, here at least, protection involved host Fc receptors. Protection by immune serum was subsequently shown to be IgG Fc receptor-dependent (Chu et al., 2008). The difficulty with interpreting such results is that the HSV gE–gI binds human but not mouse IgG Fc (Johansson et al., 1985). In contrast, MuHV-4 naturally infects mice (Kozuch et al., 1993) and gammaherpesviruses are not known to express IgG Fc receptors. Our findings with MuHV-4 – i.e. that non-neutralization epitopes provided effective antibody targets in vivo and that protection by immune serum was largely IgG Fc receptor-dependent – therefore suggested that antibodies generally act via Fc receptors in host defence against herpesviruses.

One difference between MuHV-4 and HSV is that the HSV gD provides a good neutralization target (Minson et al., 1986); the analogous (although non-homologous) MuHV-4 gp150 does not, and seems to function instead in antibody evasion. gp150 is abundant on infected-cell surfaces, but gp150-specific mAbs failed to reduce in vivo virus replication. Some uncertainty remains over gp150 processing: the equivalent gene of most other gammaherpesviruses can be spliced (Beisel et al., 1985) to exclude some of the region equivalent to the immunogenic ‘stalk’ of gp150 (Gillett et al., 2007b). However, we have found no evidence of gp150 splicing by RT-PCR, either in vitro or in vivo (data not shown). A mAb recognizing the C-terminal part of the gp150 extracellular domain identifies additional, faster-migrating forms on SDS-PAGE, suggesting the possibility of post-translational cleavage (data not shown), but we have not detected corresponding N-terminal fragments in infected-cell supernatants, so these may simply be different glycoforms. Also, gp150 cleavage alone would not explain gp150-specific mAbs being ineffective, as gp70 is shed abundantly into infected-cell supernatants (Kapadia et al., 1999; Gillet et al., 2007c) yet remained a good antibody target. Differential gp150 glycosylation seems more likely to limit the impact of gp150-specific mAbs.

The importance of IgG Fc receptors in limiting in vivo MuHV-4 replication was consistent with MuHV-4 neutralization being much more difficult than in vitro assays of fibroblast infection would suggest (Gillett et al., 2007a). One consideration is virions blocked for fibroblast infection by immune sera using IgG Fc receptors to productively infect myeloid cells (Smith et al., 2007). However, there was no evidence for neutralization being more effective in FcRγ−/− mice. Therefore, a more important consideration is probably the in vivo accessibility of virions to antibody. gp48, which promotes direct MuHV-4 cell–cell spread via actin polymerization (Gill et al., 2008), contributes more to in vivo virus replication (May et al., 2005c) than does gp150, which functions in virion release (de Lima et al., 2004), or gL, which functions in virion cell binding (Gillett et al., 2007d). Similarly, HSV cell–cell spread via gE contributes significantly to dissemination in vivo but not in vitro (Dingwell et al., 1994), whilst cell binding by gp50 is important for pseudorabies virus dissemination in vitro, yet is largely dispensable in vivo (Peeters et al., 1993). Thus, herpesviruses seem generally to remain cell-associated in vivo, affording limited opportunities for neutralization. As the main function of cell-free virions is genome transmission between hosts, their release need not extend beyond apical epithelial cells. The difficulty of herpesvirus neutralization may limit what passive antibody can achieve – certainly the inhibition of MuHV-4 replication achieved here was relatively modest. Further study is required to determine whether this reflects a specific mechanism of viral evasion.

Which IgG Fc receptor-dependent effector mechanism limits MuHV-4 replication remains unclear. Antibody-dependent cytotoxicity seems the most likely. Depleting natural killer (NK) cells with mAb PK136 (anti-NK1.1) had no effect on antibody-mediated protection (data not shown). However, macrophages, monocytes or granulocytes may be more important mediators of such cytotoxicity in mice (Nimmerjahn & Ravetch, 2008). Another (non-exclusive) possibility is enhanced virion uptake by dendritic cells improving T-cell priming. Thus, the failure of gp150-specific antibodies to reduce viral titres could have reflected that their opsonization allows particularly efficient dendritic-cell infection after uptake, and therefore disruption of antigen presentation (Smith et al., 2007; Gillet et al., 2007b), as opposed to virion uptake without infection, which should allow efficient antigen presentation. The in vivo fate of opsonized virions and infected cells clearly requires further study. The present data indicate that it is this, rather than neutralization, that chiefly determines the therapeutic effect of antiviral antibodies.
Might passive antibody also be effective against EBV? Whilst unlikely to prevent infection, it could potentially be helpful acutely in immunosuppressed patients exposed to EBV by reducing the extent of lytic viral spread. Antibody might also be a useful adjunct to latency-directed therapies such as adoptive T-cell transfer. Most considerations of anti-EBV antibody have focused on gp350 because anti-gp350 mAbs can block B-cell infection in vitro (Thorley-Lawson & Geilinger, 1980). The ineffectiveness of mAbs against the MuHV-4 gp150 suggested that other targets might be better in vivo.

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