Granulocyte-macrophage colony-stimulating factor does not increase the potency or efficacy of a foot-and-mouth disease virus subunit vaccine

Luizinho Caron2, Mario C.S. Brum2, Mauro P. Moraes2,3, William T. Golde2, Clarice Weis Arns4 and Marvin J. Grubman2*

ABSTRACT.- Caron L., Brum M.C.S., Moraes M.P., Golde W.T., Arns C.W. & Grubman M.J. 2005. [Granulocyte-macrophage colony-stimulating factor does not increase the potency or efficacy of a foot-and-mouth disease virus subunit vaccine.] Pesquisa Veterinária Brasileira 25(3):150-158. USDA, ARS, PIADC-FMD Research Unit, PO.Box 848, Greenport, NY 11944 0848, USA. E-mail: mgrubman@piadc.ars.usda.gov

Foot-and-mouth disease (FMD) is one of the most feared diseases of livestock worldwide. Vaccination has been a very effective weapon in controlling the disease, however a number of concerns with the current vaccine including the inability of approved diagnostic tests to reliably distinguish vaccinated from infected animals and the need for high containment facilities for vaccine production, have limited its use during outbreaks in countries previously free of the disease. A number of FMD vaccine candidates have been tested and a replication-defective human adenovirus type 5 (Ad5) vector containing the FMDV capsid (P1-2A) and 3C protease coding regions has been shown to completely protect pigs against challenge with the homologous virus (FMDV A12 and A24). An Ad5-P1-2A+3C vaccine for FMDV O1 Campos (Ad5-O1C), however, only induced a low FMDV-specific neutralizing antibody response in swine potency tests. Granulocyte-macrophage colony-stimulating factor (GM-CSF) has been successfully used to stimulate the immune response in vaccine formulations against a number of diseases, including HIV, hepatitis C and B. To attempt to improve the FMDV-specific immune response induced by Ad5-O1C, we inoculated swine with Ad5-O1C and an Ad5 vector containing the gene for porcine GM-CSF (pGM-CSF). However, in the conditions used in this trial, pGM-CSF did not improve the immune response to Ad5-O1C and adversely affected the level of protection of swine challenged with homologous FMDV.


RESUMO.- [Fator Estimulante de Colônias de Granulócitos e Macrófagos (GM-CSF) não aumenta a eficácia ou potência da vacina de subunidades da Febre Aftosa em suínos.] A febre aftosa é uma das doenças mais temidas nos rebanhos em todo o mundo. A vacinação tem sido uma arma eficiente no controle da doença, no entanto há preocupações com as vacinas atualmente utilizadas incluindo a necessidade de instalações de alta segurança para a produção dessas vacinas e a falta de um teste de diagnóstico aprovado que faça distinção precisa entre animais vacinados dos infectados. Várias vacinas têm sido testadas contra a febre aftosa e uma dessas utiliza como vetor um vírus defectivo para replicação, derivado do adenovírus humano tipo 5 (Ad5), o qual contém as proteínas que compõe a cápside do vírus da febre aftosa (P1-2A) e a protease 3C, protegendo completamente suínos contra o desafio de uma cepa homóloga (A12 e A24). Uma vacina com o Ad5-P1-2A+3C proveniente da cepa O1 Campos (Ad5-O1C), no entanto, somente induziu um baixo título de anticorpos neutralizantes específicos em testes de potência vacinal em suínos. O fator estimulante de colônias de granulócitos e macrófagos (GM-CSF) tem sido utilizado com sucesso na formulação de vacinas para estimular a resposta imune contra inúmeras doenças, incluindo HIV, Hepatite C e B. Na tentativa de melhorar a resposta imune específica contra
a febre aftosa induzida pelo Ad5-O1C, suínos foram vacinados com Ad5-O1C juntamente com Ad5-GM-CSF porcino. Entretanto, as condições utilizadas nesse teste, o GM-CSF suíno não melhorou a resposta imune do Ad5-O1C e adversamente afetou o nível de proteção de suínos desafiados com o vírus homólogo da febre aftosa.

TERMS OF INDEXAÇÃO: Febre Aftosa Virus O1 Campos, Adenovirus, Fator Estimulante de Colônias de Granulócitos e Macrófagos.

INTRODUCTION

Foot-and-mouth disease (FMD) is a highly contagious, acute disease of domestic and wild cloven-hoofed animals (Callis & McKercher 1986, Donaldson et al. 1986). The etiologic agent, FMD virus (FMDV), is a member of the Aphthovirus genus Picornaviridae family (Murphy et al. 1999) and contains a single-stranded, positive-sense RNA genome of approximately 8.5kb. The virus consists of seven serotypes (O, A, C, SAT 1-3 and Asia) and multiple subtypes within each serotype.

FMD outbreaks can result in significant economic losses in disease-free countries due to direct and indirect costs, most of which are related to international trade embargoes of animals and animal products. Recent outbreaks in Taiwan and the United Kingdom have highlighted the major risk of introduction and rapid spread of FMD in a susceptible population (Mathews 2001). Currently, conventional binary ethyleinimine (BEI) inactivated vaccines emulsified with adjuvant have been widely used in effective control and eradication programs around the world (Brownlie 2001). However, there are no approved diagnostic tests that can reproducibly differentiate vaccinated from infected/convalescent animals. Furthermore, conventional vaccines require growth and inactivation of live virus in containment facilities and introduce the potential for escape of live virus or incomplete inactivation (Barteling & Vreeswijk 1991). As a result some FMD-free countries have not considered vaccination as part of their disease control strategy.

Several approaches are being used to develop alternative FMD vaccines that address these concerns, including construction of modified live-virus (Mason et al. 1997, Almeida et al. 1998), biosynthetic proteins (Kleid et al. 1981, McKercher et al. 1985), synthetic peptides (Bittle et al. 1982, Clark et al. 1983, Brown 1988, Blanco et al. 2001), naked DNA vectors (Ward et al. 1997, Chinsangaram et al. 1998, Wong et al. 2000, Benvenisti et al. 2001, Cedillo-Barrón et al. 2001), and recombinant viruses (Sanz-Parra et al. 1999a, b, Mayr et al. 1999, 2001, Berinstein et al. 2000, Moraes et al. 2002, Wu et al. 2003b). However, most of these approaches have either been unsuccessful in both swine and cattle or require multiple inoculations to induce protection.

Human adenovirus (Ad5) has been used as a vector for FMD vaccines encoding the capsid (P1-2A) (Sanz-Parra et al. 1999a, b) or P1-2A and 3C protease coding regions of FMDV (Mayr et al. 1999, 2001, Moraes et al. 2002, Wu et al. 2003b). Inoculation with one dose of Ad5 containing the serotype A24 Cruzeiro P1-2A and A12 3C coding regions (Ad5-A24) protected swine from direct inoculation challenge with homologous virus (Moraes et al. 2002). More recently we have attempted to develop Ad5 vaccine vectors for other FMDV serotypes, in particular isolates of serotype O that are currently causing disease outbreaks throughout the world. We have constructed Ad5 and Ad2 vectors containing the P1-2A region of O1 Campos and the 3C of A12 (Ad5-O1C) and found that swine inoculated with these vectors developed a lower FMDV-specific neutralizing antibody response than Ad5-A24 inoculated animals (unpublished data). Likewise, swine inoculated with an Ad5-FMD bivalent vaccine containing the P1-2A regions from A24 Cruzeiro and O1C developed a higher neutralizing antibody response against A24 than against O1 (Wu et al., 2003b). It has been established that serotype O antigen induces a lower immune response as compared to serotype A antigen (Pay & Hingley 1987, Doel et al. 1994). Because of this commercial vaccines usually contain about 4-5 times more O1 inactivated virus (146S antigen) than A 146S antigen.

The use of cytokines to modulate responses against immunization with DNA and recombinant virus vectors is being actively investigated. Granulocyte-macrophage colony-stimulating factor (GM-CSF), a hematopoietic growth factor, has been widely used as a molecular adjuvant to induce immunity. GM-CSF activates neutrophils, macrophages, dendritic cells (DCs), and other mononuclear cells, and also stimulates progenitor/stem cells to mature and migrate from the bone marrow to the peripheral circulation (Kusakabe et al 2000). Intramuscular (IM) inoculation of the GM-CSF gene together with plasmids carrying viral genes, such as those encoding the glycoprotein of rabies virus and VP1 of encephalomyocarditis virus, increased antigen-specific immune responses and protective immunity in mice as compared to inoculation with the plasmid containing only the viral gene (Xiang & Ertl 1995, Sin et al. 1997). Somasundaram et al. (1999) found an adjuvant effect of porcine GM-CSF (pGM-CSF) on a DNA vaccine containing the gD and gB glycoprotein genes of Aujeszky’s disease virus (PRV) in pigs. This effect was characterized by an early appearance of anti-PRV IgG2, a significantly enhanced anti-PRV IgG1 and IgG2 antibody response, a significantly decreased and shortened period of virus shedding in nasal swabs and improved protection against viral challenge. In contrast, co-administration of porcine IFN-γ or IL-2 had no adjuvant effect.

Lee et al. (1998), observed that both humoral and cellular immune responses to hepatitis C virus (HCV) envelope proteins were augmented in rats by the co-delivery of the GM-CSF gene. Moreover, inoculation of bicistronic plasmids elicited higher levels of antibody and lymphoproliferative responses than did the co-inoculation of two independent expression plasmids that encoded the GM-CSF gene and each HCV envelope gene. Similarly Barouch et al. (2002) demonstrated that a bicistronic plasmid containing GM-CSF and HIV gp120 elicited a dramatic augmentation of vaccine-elicted CD4+ T cell responses in mice as compared to gp120 alone or individual plasmids containing each gene. These results suggest that the local concentration of GM-CSF may be a critical factor that contributes to the enhancement of the immune response to the co-expressed antigens. It has also been demonstrated that the time of administration of GM-CSF with respect to the DNA antigen vaccine can profoundly influence the nature of the Th1/Th2 balance of an antigen-specific immune response (Kusakabe et al. 2000).
We have tested pGM-CSF as an adjuvant for our Ad5-O1C vaccine and have delivered this cytokine with a recombinant Ad5 vector. We have previously utilized the Ad5 vector system to effectively deliver various cytokines including type I interferons (IFN-α/β) to swine (Chinsangaram et al. 2003, Moraes et al. 2003; Wu et al., 2003a) and cattle (Wu et al., 2003a). Swine inoculated with both Ad5-pGMCSF and Ad5-O1C, however, did not develop an enhanced immune response against O1C as compared to Ad5-O1C inoculated animals and administration of pGM-CSF had an adverse effect on the degree of protection afforded by the Ad5-O1C vaccine.

**MATERIALS AND METHODS**

**Virus and cells**
All adenovirus clones were generated, grown, and titered in human 293 cells (Graham et al. 1977, Graham & Prevec 1991), between passages 20 and 30. Plaque reduction neutralization assays with FMDV were performed in baby hamster kidney 21 cells (BHK-21) between passages 62 and 70. TF-1 cells, obtained from the American Type Culture Collection, are an indicator cell clone that is dependent on human GM-CSF or IL-3 for growth. These cells were used to analyze the biological activity of pGM-CSF obtained from the supernatants of IB-RS-2 cells (Instituto Biológico-Rim Suíno 2) infected with Ad5-pGMCSF, as pGM-CSF supports growth of this cell line as well.

**Construction of recombinant adenoviruses Ad5-pGMCSF and Ad5-O1C**

The pGM-CSF gene, containing the signal sequence, was obtained from plasmid p3Cia-pGM-CSF kindly provided by Dr. Steve Martin, Pharmacia-Upjohn. This plasmid was digested with HindIII and EcoRI, ligated to similarly digested pBluescript II KS (Stratagene, La Jolla, CA) and chemically transformed into Top 10 competent cells (Invitrogen, Carlsbad, CA) to generate the plasmid, pKSII-pGMCSF. The coding sequence of pGM-CSF was removed from pKSII-pGMCSF by digestion with ClaI and XbaI and ligated into similarly digested pAd5-Blue (Moraes et al. 2001) to generate the infectious clone pCRM8 (Sá-Carvalho et al., 1997) a chimeric FMDV clone containing the P1-A2 coding region of FMDV O1C in the background of FMDV A2, was digested with Ncol, Apal to remove the O1C P1-A2 coding region. Plasmid p12X3C (Mayr et al., 1999), which contains only the complete P1-A2 and 3C coding regions and partial 2B and 3B coding regions of FMDV A12, was digested with Ncol together with the same enzymes to remove the A12 P1-A2 coding region and ligated to p12X3C. This plasmid was ligated with BglII and XbaI to remove the A12 P1-A2 and 3C coding regions and ligated to similarly digested Ad5 transfer vector pShuttle (He et al., 1998), pShuttle-P1-A2(O1C)X3C was digested with Clal and XbaI and the fragment containing the FMDV coding regions was ligated to similarly digested pAd5-Blue to generate pAd5-O1C. Recombinant virus Ad5-O1C was produced by transfection of 293 with Pacl digested pAd5-O1C and purified virus prepared as described above. In both the Ad5-pGMCSF and Ad5-O1C viruses the inserted coding regions are under the control of the cytomegalovirus (CMV) immediate early promoter (Moraes et al., 2001).

**Expression of pGM-CSF and O1C capsid proteins synthesized in Ad5-pGMCSF or Ad5-O1C infected cells**

IB-RS-2 cells were grown in 6 well plates and infected with Ad5-pGMCSF or Ad5-O1C at a multiplicity of infection (moi) of 20. At 5 or 23 h postinfection (hpi), the cells were preincubated for 1 h in methionine-free culture medium in the absence or presence of 5 μg/ml tunicamycin, followed by radiolabeling for 1 h with [35S]methionine in methionine-free medium in the absence or presence of tunicamycin. The supernatants were harvested, frozen at -70°C and the cells were washed twice with PBS, lysed, centrifuged and the cytoplasmic extracts frozen at -70°C. Samples were immunoprecipitated with polyclonal goat anti-pGM-CSF antibody (R&D Systems, Minneapolis, MN) or various FMDV-specific antiseras as described in the captions to Figs. 1 and 3 and the immune complexes were precipitated with Streptococcus aureus protein G (Calbiochem, San Diego, CA) and analyzed by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) on a 15% gel. The gels were dried and exposed to Kodak XBR-1® X-ray film.

**Virus and cells**
The pGM-CSF gene, containing the signal sequence, was obtained from plasmid p3Cia-pGM-CSF kindly provided by Dr. Steve Martin, Pharmacia-Upjohn. This plasmid was digested with HindIII and EcoRI, ligated to similarly digested pBluescript II KS (Stratagene, La Jolla, CA) and chemically transformed into Top 10 competent cells (Invitrogen, Carlsbad, CA) to generate the plasmid, pKSII-pGMCSF. The coding sequence of pGM-CSF was removed from pKSII-pGMCSF by digestion with ClaI and XbaI and ligated into similarly digested pAd5-Blue (Moraes et al. 2001) to generate the infectious clone pCRM8 (Sá-Carvalho et al., 1997) a chimeric FMDV clone containing the P1-A2 coding region of FMDV O1C in the background of FMDV A12, was digested with Ncol, Apal to remove the O1C P1-A2 coding region. Plasmid p12X3C (Mayr et al., 1999), which contains only the complete P1-A2 and 3C coding regions and partial 2B and 3B coding regions of FMDV A12, was digested with Ncol together with the same enzymes to remove the A12 P1-A2 coding region and ligated to p12X3C. This plasmid was ligated with BglII and XbaI to remove the A12 P1-A2 and 3C coding regions and ligated to similarly digested Ad5 transfer vector pShuttle (He et al., 1998), pShuttle-P1-A2(O1C)X3C was digested with Clal and XbaI and the fragment containing the FMDV coding regions was ligated to similarly digested pAd5-Blue to generate pAd5-O1C. Recombinant virus Ad5-O1C was produced by transfection of 293 with Pacl digested pAd5-O1C and purified virus prepared as described above. In both the Ad5-pGMCSF and Ad5-O1C viruses the inserted coding regions are under the control of the cytomegalovirus (CMV) immediate early promoter (Moraes et al., 2001).

**Expression of pGM-CSF and O1C capsid proteins synthesized in Ad5-pGMCSF or Ad5-O1C infected cells**

IB-RS-2 cells were grown in 6 well plates and infected with Ad5-pGMCSF or Ad5-O1C at a multiplicity of infection (moi) of 20. At 5 or 23 h postinfection (hpi), the cells were preincubated for 1 h in methionine-free culture medium in the absence or presence of 5 μg/ml tunicamycin, followed by radiolabeling for 1 h with [35S]methionine in methionine-free medium in the absence or presence of tunicamycin. The supernatants were harvested, frozen at -70°C and the cells were washed twice with PBS, lysed, centrifuged and the cytoplasmic extracts frozen at -70°C. Samples were immunoprecipitated with polyclonal goat anti-pGM-CSF antibody (R&D Systems, Minneapolis, MN) or various FMDV-specific antiseras as described in the captions to Figs. 1 and 3 and the immune complexes were precipitated with Streptococcus aureus protein G (Calbiochem, San Diego, CA) and analyzed by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) on a 15% gel. The gels were dried and exposed to Kodak XBR-1® X-ray film.

**Biological activity of pGM-CSF synthesized in Ad5-pGMCSF infected cells**

Biological activity was measured using a Cell Proliferation ELISA System RPN 250 (BIOTRAK® Amersham Pharmacia Biotech, Piscataway, NJ). A standard curve was generated by growth of TF-1 cells, in 96-well tissue culture plates, in the presence of dilutions of recombinant pGM-CSF (R&D Systems) (Kitamura et al., 1989). Dilutions of supernatant fluids from Ad5-pGMCSF infected IB-RS-2 cells were added, in duplicate, to a 96-well plate containing TF-1 cells. After 5 days incubation, bromodeoxyuridine (BrdU) was added and cells labeled for approximately 20 h. Cell proliferation was measured following the manufacturers recommended protocol (Bautista et al. 2002).

**Animals**

Eighteen out-bred Yorkshire gilts approximately 7-8 weeks old, weighting between 35-40 lbs, were divided into five groups and each group was housed in a separate room in the high-containment facilities at the PIADC. All animals were handled humanely according to protocols approved by the Animal Care and Use Committee (ACUC) of PIADC. The animals were observed for one week prior to the start of the experiment. The vaccine was administered by IM inoculation in the neck with 2 ml of various doses of Ad5 vectors in PBS (Table 1). All animals were challenged 21 days post vaccination (dpv) and 35 dpv the remaining pigs were humanely euthanised.

**Serology and virus isolation**

Serum samples were obtained at 0 and 4 days after...
inoculation and weekly thereafter. Sera were tested for the presence of neutralizing antibodies against FMDV O1C, in a plaque reduction neutralization (PRN) assay. Neutralization titers were reported as the serum dilution yielding a 70% reduction in the number of plaques (PRN70) (Moraes et al. 2002).

Heparinized blood was obtained prior to challenge and daily for 7 days postchallenge (dpc) and examined for the presence of virus by a standard plaque assay in BHK-21 cells. Nasal secretions were obtained on the same days as heparinized blood, inoculated into IB-RS-2 cells, and observed for 48 h for cytopathic effects. Negative samples were frozen and a second passage performed. For positive samples, titration was performed from the original sample by a plaque assay on BHK-21 cells.

**Challenge**

The 18 animals were challenged with homologous FMDV O1C 21 days after receiving the respective vaccine combinations (Table 1). FMDV O1C was provided by the Instituto Pan-Americano de Febre Aftosa, Rio de Janeiro (PANAFTOSA). This virus was found at the site of inoculation; the score listed was determined at 9 dpc or the day of death; a Four days post-vaccination.

**RESULTS**

**Ad5-pGMCSF and Ad5-O1C expression**

IB-RS-2 cells were infected with Ad5-pGMCSF and at 6 or 24 hpi were radiolabeled with $^{[35S]}$methionine in the absence or presence of 5 µg/ml tunicamycin. The cell culture supernatants were examined by SDS-PAGE. As can be seen in Fig. 1 (lane 2) multiple bands are present in the supernatant at 24 hpi, while in the supernatant from cells treated with tunicamycin a single band is present (lane 3). This band is immunoprecipitated by polyclonal serum that was produced against E. coli-expressed pGM-CSF (lane 6). These results are in agreement with published sequence data indicating that mature pGM-CSF has three potential N-linked glycosylation sites (Inumaru & Takamatsu 1995). The inefficient immunoprecipitation of glycosylated pGM-CSF (lane 5) may be the result of interference with the antigen-antibody reaction by the extensive glycosylation of this protein. We further analyzed production of pGM-CSF in IB-RS-2 cells by Western blot analysis and show glycosylated pGM-CSF is efficiently detected, suggesting that denatured protein is a better antigen for this polyclonal antibody (compare Fig. 2, lanes 6 and 7 to Fig. 1, lane 5).

To examine the expression of O1C capsid proteins by Ad5-O1C, IB-RS-2 infected cells were radiolabeled, cell lysates prepared at 5-7 hpi, immunoprecipitated with various FMDV-specific antibodies, and analyzed by SDS-PAGE (Fig. 3). We and others have previously shown that the A12 3C protease can efficiently process the P1-2A precursor from A24, O1C, and O1 Taiwan/99 (Sa-Carvalho et al. 1997, Almeida et al. 1998, Moraes et al. 2002, Wu et al. 2003b unpublished). Reaction with bovine convalescent serum that was produced against E. coli-expressed pGM-CSF (lane 5) may be the result of interference with the antigen-antibody reaction by the extensive glycosylation of this protein. We further analyzed production of pGM-CSF in IB-RS-2 cells by Western blot analysis and show glycosylated pGM-CSF is efficiently detected, suggesting that denatured protein is a better antigen for this polyclonal antibody (compare Fig. 2, lanes 6 and 7 to Fig. 1, lane 5).

<table>
<thead>
<tr>
<th>Animal Group</th>
<th>Virus constructs</th>
<th>Dose (PFU/animal)</th>
<th>Clinical scores</th>
<th>Weeks post-vaccination (PRN 70)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2107 1</td>
<td>Ad5-Blue</td>
<td>1X10⁹</td>
<td>16</td>
<td>&lt;8 &lt;8 &lt;8 &lt;8 &lt;8 Dead</td>
</tr>
<tr>
<td>2108 1</td>
<td>Ad5-Blue</td>
<td>18</td>
<td>&lt;8 &lt;8 &lt;8 &lt;8 &lt;8 Dead</td>
<td></td>
</tr>
<tr>
<td>2109 1</td>
<td>Ad5-Blue</td>
<td>18</td>
<td>&lt;8 &lt;8 &lt;8 &lt;8 &lt;8 Dead</td>
<td></td>
</tr>
<tr>
<td>2110 2</td>
<td>Ad5-pGMCSF</td>
<td>17</td>
<td>&lt;8 &lt;8 &lt;8 &lt;8 &lt;8 160 1000</td>
<td></td>
</tr>
<tr>
<td>2111 2</td>
<td>Ad5-pGMCSF</td>
<td>15</td>
<td>&lt;8 &lt;8 &lt;8 &lt;8 &lt;8 Dead</td>
<td></td>
</tr>
<tr>
<td>2112 2</td>
<td>Ad5-pGMCSF</td>
<td>17</td>
<td>&lt;8 &lt;8 &lt;8 &lt;8 &lt;8 640 1000</td>
<td></td>
</tr>
<tr>
<td>2104 3</td>
<td>Ad5-O1C</td>
<td>5X10⁹</td>
<td>11</td>
<td>&lt;8 &lt;8 16 16 &lt;16 2560 2000</td>
</tr>
<tr>
<td>2113 3</td>
<td>Ad5-O1C</td>
<td>8</td>
<td>&lt;8 &lt;8 &lt;8 &lt;8 &lt;8 4000 1000</td>
<td></td>
</tr>
<tr>
<td>2114 3</td>
<td>Ad5-O1C</td>
<td>11</td>
<td>&lt;8 &lt;8 32 64 32 4000 2000</td>
<td></td>
</tr>
<tr>
<td>2115 3</td>
<td>Ad5-O1C</td>
<td>8</td>
<td>&lt;8 &lt;8 8 16 16 2560 2000</td>
<td></td>
</tr>
<tr>
<td>2105 4</td>
<td>Ad5-O1C + Ad5-pGMCSF</td>
<td>5X10⁹ + 1X10⁸</td>
<td>9</td>
<td>&lt;8 16 32 16 8000 2000</td>
</tr>
<tr>
<td>2116 4</td>
<td>Ad5-O1C + Ad5-pGMCSF</td>
<td>12</td>
<td>&lt;8 8 16 16 &lt;16 4000 2000</td>
<td></td>
</tr>
<tr>
<td>2117 4</td>
<td>Ad5-O1C + Ad5-pGMCSF</td>
<td>10</td>
<td>&lt;8 16 64 16 &lt;16 8000 2000</td>
<td></td>
</tr>
<tr>
<td>2118 4</td>
<td>Ad5-O1C + Ad5-pGMCSF</td>
<td>15</td>
<td>&lt;8 8 16 16 &lt;16 2560 4000</td>
<td></td>
</tr>
<tr>
<td>2106 5</td>
<td>Ad5-O1C + Ad5-pGMCSF</td>
<td>12</td>
<td>&lt;8 32 32 16 4000 4000</td>
<td></td>
</tr>
<tr>
<td>2119 5</td>
<td>Ad5-O1C + Ad5-pGMCSF</td>
<td>11</td>
<td>&lt;8 32 32 16 4000 4000</td>
<td></td>
</tr>
<tr>
<td>2120 5</td>
<td>Ad5-O1C + Ad5-pGMCSF</td>
<td>9</td>
<td>&lt;8 32 32 16 &lt;16 1240 4000</td>
<td></td>
</tr>
<tr>
<td>2121 5</td>
<td>Ad5-O1C + Ad5-pGMCSF</td>
<td>13</td>
<td>&lt;8 8 16 16 &lt;16 4000 2000</td>
<td></td>
</tr>
</tbody>
</table>

a Serum dilution that reduces the number of plaques by 70%; b Clinical scores were determined by the number of toes presenting FMD compatible lesions plus the presence of lesions in the snout and mouth, with a maximum score of 18; we did not count a lesion found at the site of inoculation; the score listed was determined at 9 dpc or the day of death; c Four days post-vaccination.

The biological activity of pGM-CSF obtained from supernatant fluids of Ad5-pGMCSF infected IB-RS-2 cells was measured in vitro by a TF-1 cell proliferation assay. As shown in Fig. 4, Ad5 expressed pGM-CSF has biological activity as demonstrated by the support of TF-1 cell growth, while supernatant fluids from a control Ad5 lacking the pGM-CSF gene, Ad5-Blue, had no biological activity.

Immune response prior to challenge

To examine the potency and efficacy of Ad5-O1C in susceptible animals and determine if pGM-CSF can enhance the immune response of the FMDV antigens, 3 groups of swine, 4 animals per group, were inoculated IM with Ad5-O1C at 5x10^8 pfu/ml in the absence (Group 3) or presence of 1x10^8 or 1x10^9 pfu/animal Ad5-pGMCSF (Groups 4 and 5, Table 1). Control groups, 3 animals per group, were inoculated with an Ad5 vector, Ad5-Blue (Group 1), or with Ad5-pGMCSF alone (Group 2).

After vaccination no adverse side effects, such as fever, apathy or inflammation at the inoculation site, were observed in any of the pigs. The FMDV-specific neutralizing antibody titers of the Ad5-O1C inoculated animals were detectable by 4-7 dpi, but were low and were not increased by co-administration of Ad5-pGMCSF (Table 1, Fig. 5). As expected the animals in the control groups did not develop an FMDV-specific neutralizing antibody titer.

Clinical response after challenge

All animals were challenged 21 dpv. After challenge, swine in the control groups (1 and 2) developed a fever (temperature of 40°C or higher for 2 or more consecutive days) by 1-2 dpc and vesicular lesions by 1 dpc. All of these animals had extensive disease by 2-3 dpc (Fig. 6). Between 3-5 dpc all animals in control group 1, that received the Ad5-Blue vector, died and on day 3 one animal in group 2, that received Ad5-pGMCSF, also died. Upon necropsy, these four animals all had extensive signs of heart necrosis (tiger heart) which was confirmed by histopathological examination. These cardiac complications are typical causes of death as a result of FMDV infection in young swine. The two control animals that survived each lost at least one hoof by one week postchallenge.

Nine of twelve of the vaccinated pigs (Groups 3, 4 and 5) developed fever by 2-5 dpc, while 1 pig in group 3 and two in group 4 never had fever. All vaccinated swine developed signs of infection, but the onset of disease was delayed 1-2 days compared to the controls, disease severity was reduced, and no animals in these groups died or lost their hoofs (Fig. 6). The group that received only Ad5-O1C (Group 3) had a slightly reduced clinical
score as compared to the groups that received both Ad5-O1C and Ad5-pGMCSF (Groups 4 and 5).

**Serological response after challenge**

The two surviving animals in the control groups became FMD positive by 7 dpc and all animals in the FMD vaccinated groups developed a substantial increase in FMDV-specific neutralizing antibody response after challenge (Table 1, Fig. 5).

In both control groups viremia was detected at 1 dpc and reached a peak of 1.5-3.5x10⁶ pfu/ml at 2 dpc (Fig. 7A). In the
Luizinho Caron et al.

FMD vaccinated groups, viremia was first detected in groups 4 and 5 (vaccine + pGMCSF) at 2-3 dpc and by 3-4 dpc in group 3 (vaccine alone). One pig in group 3 never developed viremia or fever and a second animal only had a low level of viremia for 1 day (Fig. 7A). Peak viremia was at 3 dpc in the dual-inoculated groups and was approximately 1,000-fold lower as compared to the controls (Fig. 7A). In the Ad5-O1C vaccinated animals, peak viremia was at 4 dpc and was approximately 10,000-fold lower than in the control pigs.

Virus present in nasal swabs reached a peak in both control groups by 2-3 dpc and at 4 dpc in all three FMD vaccinated groups (Fig. 7B). Group 3, vaccinated with only Ad5-O1C, had the lowest levels of virus shedding.

**DISCUSSION**

We have previously demonstrated that a single IM inoculation of Ad5-FMDV vectors containing the P1-2A coding regions from A12 (Mayr et al. 2001) or A24 (Moraes et al. 2002) and the A12 3C protease coding region can induce significant FMDV-specific neutralizing antibody titers in swine and protect most or all of the animals when challenged either by contact or direct inoculation. However, inoculation with an Ad5 vector containing the O1C capsid coding region or an Ad5 vector containing the O1C capsid coding region in a bicistronic expression cassette induces a significantly lower FMDV-specific neutralizing antibody response in swine (unpublished observations; Wu et al. 2003b). Similar results have been reported with the conventional inactivated FMD vaccine and therefore to be an efficacious vaccine, manufacturers include a higher dose of type O 146S antigen than type A in vaccine formulations (Pay & Hingley 1987, Doel et al. 1994).

To enhance the immune response induced by Ad5-O1C, we tested a vaccination regimen that included co-administration of pGM-CSF, a cytokine that has been shown to enhance the immune response of animals to some vaccines (Xiang & Ertl 1995, Lee et al. 1998, Somasudaran et al. 1999, Cedillo-Barrón et al. 2001, Barouch et al. 2002). We constructed an Ad5-pGMCSF virus that expressed biologically active pGM-CSF. Swine inoculated with Ad5-O1C and either a low or high dose of Ad5-pGMCSF developed a detectable FMDV-specific neutralizing antibody response prior to the animals administered only Ad5-O1C, i.e., by 4 days, and the response was higher at 1 week postinoculation. However, by 2 weeks postinoculation and at the time of challenge the neutralizing antibody response of these 3 groups was equivalent (Fig. 5) and was considerably lower than the response to one inoculation of Ad5-A12 or Ad5-A24 (Mayr et al. 2001, Moraes et al. 2002). Thus, the co-administration of Ad5-pGMCSF did not enhance the long-term FMDV-specific neutralizing antibody response.

After challenge the control groups (Groups 1 and 2) rapidly developed significant disease and all 3 animals in Group 1 and one of 3 animals in Group 2 died. The histopathology findings upon necropsy of the four animals that died after challenge revealed that they all had severe vesicular lesions on the mouth and all feet, resembling typical FMD lesions, and moderate to severe myocardial necrosis resulting in heart failure, which was most likely the result of FMDV infection. In previous experiments with FMDV serotype A, control groups of swine rapidly developed significant disease after direct inoculation challenge, but did not die if they were healthy prior to the start of the experiment (Moraes et al. 2002, Chinsangaram et al. 2003, Moraes et al. 2003), suggesting that the challenge virus used in the present studies was either too virulent or the dose used was excessive. In subsequent titration of this virus, 1/100th the dose used in this study still resulted in the death of 1 of 2 naïve animals and the animal that survived had severe disease including loss of hoofs. In either case the “over-challenge”, in this experiment, masked the protective effects of the Ad5-O1C vaccine, since all vaccinated animals developed vesicular lesions and 3 of 4 animals had viremia. However, vaccination with Ad5-O1C clearly delayed and reduced the severity of disease. None of the animals inoculated with only Ad5-O1C died or lost their hoofs, the appearance of vesicular lesions was delayed for 1-
2 days as compared to the control groups and was less severe. Furthermore, the peak of viremia was delayed 2 days, one animal never developed viremia, and virus titers were approximately 10,000-fold lower than in the controls, and the peak of virus shedding was delayed 2 days.

The addition of pGM-CSF did not enhance the efficacy of the Ad5-O1C vaccine. Although none of the animals given Ad5-pGM-CSF died or lost their hoofs, all developed vesicular lesions that were somewhat more severe than the Ad5-O1C vaccinated animals. Furthermore, while viremia was delayed and reduced as compared to the control groups all of these effects can be attributed to the vaccine alone, as the Ad5-O1C inoculated animals had even lower levels of viremia.

We have subsequently repeated this experiment, and included a second route of inoculation, and obtained very similar results. Swine inoculated with Ad5-O1C IM with a needle or subcutaneously/IM by needleless delivery and challenged had no detectable viremia and clinical disease was delayed and less severe than control animals (data not shown). However, the addition of Ad5-pGM-CSF reduced the level of protection afforded by the vaccine alone.

Cedillo-Barrón et al. (2001) demonstrated that the addition of plasmid encoded pGM-CSF along with a DNA based FMDV empty capsid vaccine resulted in a statistically significant increase in the antibody levels against FMDV and somewhat improved the protection of vaccinated swine as compared to swine only given the empty capsid vaccine. However, in these studies the swine were challenged after 3 DNA inoculations and the increase in FMDV-specific antibody levels was only detected after the second inoculation. Additional studies examining the adjuvant effect of GM-CSF for other antigens have suggested that the presence of this cytokine at the site of vaccine inoculation is important (Wang et al. 2002) and that plasmids co-expressing antigen and GM-CSF are more effective than co-administration of plasmids separately expressing each antigen (Lee et al. 1998, Barouch et al. 2002). The addition of pGM-CSF did not enhance the efficacy of the Ad5-O1C vaccine alone.

Efficacy of a foot-and-mouth disease virus subunit vaccine

2 CONCLUSION

In previous work, we have demonstrated that swine given one dose of an Ad5-O1C vector developed only low FMDV-specific neutralizing antibody response and did develop clinical disease after challenge, disease signs and viremia were considerably reduced as compared to control groups. While the addition of Ad5-pGM-CSF induced a more rapid neutralizing antibody response, it adversely affected the level of protection afforded by the Ad5-O1C vaccine alone.

Acknowledgements - We are thankful to Marla Koster, Dr Elida Bautista and Dr. Vladimir G. Andreyev for their technical assistance, Dr. Peter W. Mason for providing the FMDV O1 Campos challenge virus and plasmid pCRM8, the Plum Island animal caretakers for their assistance with the swine, and the USDA-ARS for financial support.

REFERENCES


