In Vivo Expression of and Cell-Mediated Immune Responses to the Plasmid-Encoded Virulence-Associated Proteins of *Rhodococcus equi* in Foals

Stephanie Jacks,1 Steeve Giguère,1,* and John F. Prescott2

Department of Large Animal Clinical Sciences, College of Veterinary Medicine, University of Florida, Gainesville, Florida 32610,1

and Department of Pathobiology, Ontario Veterinary College, University of Guelph, Guelph, Ontario N1G 2W1, Canada2

Received 30 November 2006/Returned for modification 19 December 2006/Accepted 5 February 2007

*Rhodococcus equi* is a facultative intracellular pathogen that causes pneumonia in foals but does not induce disease in adult horses. Virulence of *R. equi* depends on the presence of a large plasmid, which encodes a family of seven virulence-associated proteins (VapA and VapC to VapH). Eradication of *R. equi* from the lungs depends on gamma interferon (IFN-γ) production by T lymphocytes. The objectives of the present study were to determine the relative in vivo expression of the vap genes of *R. equi* in the lungs of infected foals, to determine the recall response of bronchial lymph node (BLN) lymphocytes from foals and adult horses to each of the Vap proteins, and to compare the cytokine profiles of proliferating lymphocytes between foals and adult horses. *vapA, vapD*, and *vapG* were preferentially expressed in the lungs of infected foals, and expression of these genes in the lungs was significantly (*P < 0.05*) higher than that achieved during in vitro growth. VapA and VapC induced the strongest lymphoproliferative responses for foals and adult horses. There was no significant difference in recall lymphoproliferative responses or IFN-γ mRNA expression by bronchial lymph node lymphocytes between foals and adults. In contrast, interleukin 4 (IL-4) expression was significantly higher for foals than for adults for each of the Vap proteins. The ratio of IFN-γ to IL-4 was significantly higher for foals than for adult horses for most Vap proteins. Therefore, foals are immunocompetent and are capable of mounting lymphoproliferative responses of the same magnitude and cytokine phenotype as those of adult horses.

*Rhodococcus equi*, a gram-positive facultative intracellular pathogen, is one of the most important causes of pneumonia in foals between ages 3 weeks and 5 months. *R. equi* has also emerged as a significant opportunistic pathogen in immunosuppressed people, especially those infected with the human immunodeficiency virus (3, 9, 14). In foals, the course of the disease is insidious and pathology is often extensive by the time the disease is diagnosed. Unlike environmental *R. equi* isolates from pneumonic foals typically contain an 80- to 90-kb plasmid. Plasmid-cured derivatives of virulent *R. equi* strains lose their ability to replicate and survive in macrophages (12). Plasmid-cured derivatives also fail to induce pneumonia and are completely cleared from the lungs of foals, confirming the absolute necessity of the large plasmid for the virulence of *R. equi* (12, 37).

A 27.5-kb region of the virulence plasmid bears the hallmark of a pathogenicity island and contains the genes for a family of seven closely related virulence-associated (Vap) proteins, designated VapA and VapC to VapH (35). Although a recent study has proposed the designation vapI for another gene of the pathogenicity island, vapI is not functional (31). In a recent study, an *R. equi* mutant lacking a 7.9-kb DNA region spanning five vap genes (*vapA, -C, -D, -E, and -F*) was attenuated for virulence in mice and failed to replicate in macrophages (20). Only complementation with *vapA* could restore full virulence, whereas complementation with *vapC, vapD, or vapE* could not (20). All *vap* genes are expressed during in vitro growth and are upregulated when *R. equi* is grown in equine macrophage monolayers (32).

Because of the facultative intracellular nature of *R. equi*, cell-mediated immune mechanisms are thought to be of major importance in resistance. Most knowledge of cell-mediated immunity to *R. equi* infections comes from studying infection of mice. In mice, functional T lymphocytes are required for the clearance of virulent *R. equi*. Although both CD4+ and CD8+ T cells contribute to the host defense against *R. equi* in mice, CD4+ T lymphocytes play the major role and are required for complete pulmonary clearance (21). Studies with mice have also clearly shown that a Th1 response, characterized by gamma interferon (IFN-γ) induction, is sufficient to effect complete pulmonary clearance of *R. equi*, whereas a Th2 response, characterized by interleukin 4 (IL-4) induction, is detrimental (22, 23). How these findings in mice relate to the foal remains to be determined. As opposed to foals, adult horses completely clear an intrabronchial challenge with virulent *R. equi* and do not develop clinical signs. Clearance of *R. equi* in adult horses is associated with a significant increase in bronchoalveolar lavage fluid CD4+ and CD8+ lymphocytes, lymphoproliferative responses to *R. equi* antigens, including VapA, development of *R. equi*-specific cytotoxic T lymphocytes, and IFN-γ induction (15–17, 24, 30). A few studies have examined antibody responses of foals and adult horses to the
Vap proteins of R. equi (18, 24). However, cell-mediated immune responses and cytokine profiles of foals and adult horses for each of the Vap proteins have never been evaluated. Knowledge of gene products preferentially induced during infection in the natural host would provide important insight into the pathogenesis of the disease and may prove important for vaccine development. Despite the documented importance of some of the vap genes in the virulence of R. equi, relative expression of these genes during active infection in foals has not been studied.

In regard to the above, the objectives of the present study were to determine the relative in vivo expression of the functional vap genes of R. equi in the lungs of infected foals, to determine the recall lymphoproliferative responses of foals to each of the functional Vap proteins and compare them to those of resistant adult horses, and to determine the cytokine profiles of proliferating lymphocytes for both foals and adult horses.

MATERIALS AND METHODS

Animals and intrabronchial challenge. Five foals between 7 and 10 days of age and five adult horses between 2 and 12 years of age were used in this study. Adequate transfer of passive immunity was confirmed in foals at 12 to 24 h of age by measurement of the plasma immunoglobulin G concentration using a commercial immunoassay (DVM Stat; Corporation for advanced Applications, Newburg, WI). Foals together with their dams were moved to individual stalls in an isolation facility the day after birth. Adult horses were moved to the isolation facility at least 2 days prior to the beginning of the study. Prior to initiation of the study, all animals were determined to be healthy on the basis of a thorough physical examination performed by two veterinarians. Animals were supplied with water and feed ad libitum, and their body weight was measured daily. None of the animals showed signs of previous R. equi infection.

For determination of in vitro expression of the vap genes, in vitro RNA was obtained from in vitro-grown bacteria. The vap genes were induced for 12 h. This time selection was based on a time-response curve of VapA and ConA, respectively. The cells were stimulated at 37°C for 72 h in 6% CO2. Twelve hours before the end of the assay, 20 µl of alamar blue (Accumed International, Inc., Westlake, OH) was added to each of the plates. Plates were read on a fluorometer (Synergy HT; BioTek Instruments, Inc., Winooski, VT) using an excitation wavelength of 530 nm and an emission wavelength of 590 nm. The change in fluorescence was calculated as the mean for the stimulated plates minus the mean for the cells without antigen or mitogen (blank).

BLN cells used for quantification of mRNA expression were prepared exactly as described above with the exception that the cells were stimulated with each of the Vap antigens for 12 h. This time selection was based on a time-response curve for IFN-γ and IL-4 mRNA expression.

RNA isolation from BLN cells. RNA treatment of bacterial RNA samples, and cDNA synthesis. Isolation of total RNA from BLN cells was performed using the QIAGEN RNeasy kit (QIAGEN, Inc., Valencia, CA) according to the manufacturer’s instructions. The RNA concentration was determined by measuring the optical density at 260 nm. All RNA samples were treated with amplification-grade DNase I (Gibco BRL, Rockville, MD) to remove any trace of genomic DNA contamination. Briefly, 1 U of DNase I and 1 µl of 10× DNase I reaction buffer were added to each sample. The mixture was incubated for 10 min at room temperature and then inactivated by addition of 1 µl of 25 mM EDTA and heating at 65°C for 10 min. cDNA was synthesized with the Advantage RT-for-PCR kit (Clontech, Palo Alto, CA) by using the protocol of the manufacturer. Briefly, 1 µg of total RNA was mixed with 1 µl of oligo(dT)12 primer (20 U) and heated at 70°C for 2 min. After cooling to room temperature, the following reagents were added: 5 µl of 5× first-strand buffer, 1 µl of 10 mM dNTPs, 50 µg of Escherichia coli RNase (10 mM each), 0.5 µl of RNase inhibitor (40 U/µl), and 1 µl of Moloney murine leukemia virus reverse transcriptase (200 U/µl). The mixture was incubated at 42°C for 1 h, heated at 94°C for 5 min, diluted to a final volume of 100 µl, and stored at −70°C until used for PCR analysis.

Quantification of cytokine mRNA. Gene-specific primers and internal oligonucleotide probes for equine gyceraldehyde-3-phosphate dehydrogenase, IL-4, and IFN-γ have been described previously (11). The internal probes were labeled at the 5′ end with the reporter dye 6-carboxyfluorescein and at the 3′ end with the quencher dye 6-carboxytetramethylrhodamine. Amplification of 2 µl of cDNA was performed in a 25-µl PCR mixture containing 900 nM of each primer, 250 nM of TaqMan probe, and 12 µl of TaqMan Universal PCR Mastermix (Applied Biosystems, Foster City, CA). Amplification and detection were performed using the ABI Prism 7900 sequence detector system (Applied Biosystems) with initial incubation steps at 50°C for 2 min and 95°C for 10 min, followed by 40 cycles of 95°C for 15 s and 60°C for 1 min. Serial dilutions of cDNA from 24-h concanavalin A-stimulated equine blood mononuclear cells were used to generate a standard curve for relative quantification of each gene of interest. Each sample was assayed in triplicate, and the mean value was used for comparison. Samples without cDNA were included in the amplification reactions to determine background fluorescence and check for contamination. To account for variation in the amount and quality of starting material, values obtained for IFN-γ and IL-4 transcripts were normalized by dividing by the gyceraldehyde-3-phosphate dehydrogenase transcript level for the same sample. The sample with the lowest level of gene expression was designated 1, and relative quantification between samples was reported as the n-fold difference relative to cytokine mRNA expression in that sample.

Bacterial RNA isolation, DNase treatment of bacterial RNA samples, and cDNA synthesis. Bacterial RNA extraction for determination of in vivo expression of the vap genes was performed on lung tissue from the five infected foals. For determination of in vitro expression of the vap genes, one colony of R. equi ATCC 3701 was plated in five separate tubes, each containing 5 ml of brain heart infusion broth with 1% yeast extract. The suspensions were grown in a shaking incubator at 37°C for 18 h. A 125-mm3 section of lung tissue and pellets from in vivo-grown bacteria were independently pulverized in cold Trizol reagents (Invitrogen Corporation, Carlsbad, CA) using glass tissue grinders. Total RNA was extracted using the Trizol reagents by following the instructions from the manufacturer. RNA from each source was further purified independently by passing the resulting sample through RNeasy MiniElute spin columns (QIAGEN, Scientific and Technical Publishing Company).
TABLE 1. Oligonucleotide primer and probe sequences for amplification of vap genes from R. equi

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer or probe</th>
<th>Sequence (5’ to 3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>vapA</td>
<td>Forward</td>
<td>CATCAAACCTTCTGATAGCAGTTA</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>GACGCACCACAGTATAGAATCCT</td>
</tr>
<tr>
<td></td>
<td>Probe</td>
<td>TCTCAGGCGCATATCGAGGCT</td>
</tr>
<tr>
<td>vapC</td>
<td>Forward</td>
<td>GGGTACGCGTGGTGGG</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>CCGTATACTGTGTCGCGGA</td>
</tr>
<tr>
<td></td>
<td>Probe</td>
<td>TCGGAGTTCTCCGAGCAGGAC</td>
</tr>
<tr>
<td>vapD</td>
<td>Forward</td>
<td>CAGAGCTTTTGGCG</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>CATIGAAGGCGGTGGCATC</td>
</tr>
<tr>
<td></td>
<td>Probe</td>
<td>TTATTCAGGTCTTGGCTCAGGCT</td>
</tr>
<tr>
<td>vapE</td>
<td>Forward</td>
<td>TGAGTACAAAGCTTGCTGGCC</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>ACCTGCGCAGCAAACC</td>
</tr>
<tr>
<td></td>
<td>Probe</td>
<td>TACTGAAACTAATATCAAG</td>
</tr>
<tr>
<td>vapF</td>
<td>Forward</td>
<td>CATCACTGCTGGGAAAGTACT</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>CCCATGAAAGCAGGATTTG</td>
</tr>
<tr>
<td></td>
<td>Probe</td>
<td>CGGCAATCAATAACATGGCAGGAC</td>
</tr>
<tr>
<td>vapG</td>
<td>Forward</td>
<td>CACTGCAAACCCCGGGA</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>GCCGCGCAGAAAGCCTG</td>
</tr>
<tr>
<td></td>
<td>Probe</td>
<td>TCCGAAATCCGCGAGCAGA</td>
</tr>
<tr>
<td>vapH</td>
<td>Forward</td>
<td>GTCAACTTTCCTGACAGGTCA</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>ACAGGGACTCTCCCTCCTCA</td>
</tr>
<tr>
<td></td>
<td>Probe</td>
<td>CGGGCAATCCTCGGCCATAGC</td>
</tr>
</tbody>
</table>

Inc.) with on-column DNase treatment according to the manufacturer’s instructions. To deplete equine RNA and further enrich bacterial RNA, RNA samples from lung tissue were additionally subjected to the MICROBEnrich kit (Ambion, Inc., Austin, TX). Then, in an attempt to completely eliminate genomic DNA contamination, each RNA sample underwent an additional treatment with amplification-grade DNase I (Gibco BRL) before reverse transcription. Synthesis of cDNA was performed as described above except that random hexamer priming (20 μM) was used. Duplicate RNA samples were subjected to the same reverse transcription protocol minus the reverse transcriptase to confirm the absence of genomic DNA contamination or give a baseline used to subtract traces of remaining genomic DNA contamination.

Quantification of R. equi vap mRNA. Gene-specific primers and internal oligonucleotide probes for vapA, vapC, vapD, vapE, vapF, vapG, and vapH were selected based on the plasmid DNA sequence of R. equi ATCC 33701 using Primer Express software (Table 1). Primer and probe sequences for 16S rRNA, selected based on the plasmid DNA sequence of R. equi strain 33701, the plasmid-cured derivative of ATCC 33701, was used to generate a standard curve for relative quantification of the vap genes. Genomic DNA from R. equi strain 33701, the plasmid-cured derivative of ATCC 33701, was used to generate a standard curve for relative quantification of 16S rRNA. Amplification and detection were performed as described above under “Quantification of cytokine mRNA.” Calculated amplification efficiencies for primer/probe assays ranged between 89.0% and 99.9%. Coefficients of correlation (r) of the standard curves ranged between 0.981 and 0.985. To account for variation in the amount and quality of starting material, values obtained for vap transcripts were normalized by dividing by the 16S rRNA transcript level for the same sample. The sample with the lowest level of gene expression was designated 1, and relative quantification between samples was reported as the n-fold difference relative to mRNA expression in that sample. In rare instances when traces of genomic DNA were detected, the concentration of genomic DNA was subtracted from that for gene expression.

Statistical analyses. Statistical analyses were performed by using SigmaStat (version 3.0; SPSS, Inc., Chicago, IL). Differences in lymphocyte proliferation and cytokine expression between foals and adult horses, as well as differences in cytokine expression between Vap proteins for a given group (foals or adults), as well as differences in the relative expression levels of each vap gene within a given sampling site (in vivo or in vitro), were assessed using the Friedman repeated measure analysis of variance on ranks. When appropriate, multiple pairwise comparisons were done using the Student-Newman-Keuls test. Significance was set at a P value of <0.05.

RESULTS

Disease process and macroscopic findings. All animals maintained normal vital signs during the study and showed no evidence of disease. All five infected foals had macroscopic pulmonary lesions that consisted of mild to moderate areas of consolidation in the ventral lung lobes, along with multiple small nodular lesions up to 1 cm in diameter. The mean number of R. equi bacteria (± standard deviation [SD]) was 9.63 × 10^8 ± 1.21 × 10^7 (range, 6.23 × 10^8 to 2.53 × 10^7) CFU/g of lung tissue. The lungs of adult horses were free from lesions, and bacterial culture was negative.

In vivo expression of R. equi vap genes. Real-time PCR was used to quantify the relative in vivo expression of the vap genes in the lung tissue of infected foals, and the expression was compared to that of in vitro-grown bacteria. Expression levels of vapA, vapD, and vapG were significantly higher during in vivo growth than during in vitro growth (Fig. 1). Expression of vapG in lung tissue was significantly higher than that of all other vap genes. Expression of vapA and vapD in lung tissue was significantly higher than that of vapC, vapE, vapF, and vapH (Fig. 1). The expression of the vap genes during in vitro growth followed a similar pattern, with significantly greater expression of vapA, vapD, and vapG (Fig. 1).

Vap-specific proliferative responses and cytokine expression. To determine whether differences in lymphoproliferative responses and cytokine profiles may be at the origin of the peculiar susceptibility of foals to infection with R. equi, recall responses of susceptible, infected foals was compared to those of R. equi-resistant adult horses. Proliferation of BLN mononuclear cells following stimulation with each recombinant Vap protein was measured. There were no significant differences in lymphoproliferative responses to each of the Vap proteins expression between in vivo and in vitro expression of the plasmid-borne vap genes of R. equi. In vivo expression was determined in lung tissue from five foals experimentally infected with R. equi. In vitro expression was determined during exponential growth of R. equi in broth. Results are presented as means ± SD. Different letters (a, b, and c) indicate a significant difference in expression in vivo between vap genes. Different numbers (1, 2, and 3) indicate a significant difference in expression in vitro between vap genes. * Significant difference in expression of a given vap gene between in vivo and in vitro growth (P < 0.05).

FIG. 1. Comparison of in vivo and in vitro expression of the plasmid-borne vap genes of R. equi. In vivo expression was determined in lung tissue from five foals experimentally infected with R. equi. In vitro expression was determined during exponential growth of R. equi in broth. Results are presented as means ± SD. Different letters (a, b, and c) indicate a significant difference in expression in vivo between vap genes. Different numbers (1, 2, and 3) indicate a significant difference in expression in vitro between vap genes. * Significant difference in expression of a given vap gene between in vivo and in vitro growth (P < 0.05).
between infected foals and adult horses (Fig. 2). For foals, lymphoproliferative responses to VapA were significantly greater than those to all of the other Vap proteins, and lymphoproliferative responses to VapC were significantly greater than those to all Vap proteins except VapA (Fig. 2). In adult horses, lymphoproliferative responses to VapA and VapC were significantly greater than those to the other Vap proteins (Fig. 2).

Induction of IFN-γ and IL-4 in proliferating cells was quantified by real-time reverse transcription-PCR. There was no significant difference in IFN-γ expression by BLN cells between foals and adult horses for any of the Vap proteins (Fig. 3A). In contrast, IL-4 mRNA expression for each of the Vap proteins was significantly greater for adult horses than for foals (Fig. 3B). The ratio of IFN-γ to IL-4 was significantly higher with foals than with adult horses for VapA, VapC, VapD, VapF, and VapH (Fig. 3C).

**DISCUSSION**

Despite a central role for cell-mediated immune responses in protection against *R. equi*, most studies have focused on antibody responses against the Vap proteins. This is the first study to examine the expression of *vap* genes in the lungs of *R. equi*-infected foals and to measure lymphoproliferative responses and the cytokine profiles of infected foals and adult horses for each of the functional Vap proteins. It complements an earlier study by Hooper-McGrevy et al. (18) that examined the serum immunoglobulin G subisotype responses to each of the Vap proteins for foals and for adult horses, as well as a study evaluating the expression of the pathogenicity island genes in equine macrophages (32).

VapA is expressed on the bacterial surface, and its expression is temperature regulated, occurring between 34 and 41°C (34). In contrast, VapC, -D, and -E are secreted proteins concomitantly regulated by temperature with VapA, being produced at 37°C but not at 30°C (7). The cellular locations of VapF, VapG, and VapH are unknown. In one study, all *vap* genes were upregulated when *R. equi* was grown in equine macrophage monolayers compared to expression during in vitro growth at 30°C (32). In the same study, *vapA* and *vapC*...
were the most highly upregulated of the vap genes (32). In the present study, only vapA, vapD, and vapG were expressed at significantly higher levels in vivo than during in vitro growth. The discrepancy between findings of these studies may result from the fact that culture in macrophage monolayers does not accurately reflect all the conditions experienced by the pathogen in the lungs of foals. In addition, relative expression of the vap genes may depend on the stage of infection. The present study evaluated expression in the lungs of foals 2 weeks after infection, whereas the macrophage study evaluated expression 4 h after the monolayers were infected with R. equi. Finally, differences in temperatures for in vitro growth between the two studies may at least partially explain some of the inconsistencies. The macrophage study grew R. equi in vitro at 30°C, a temperature known to decrease expression of the Vap proteins (7, 32). In contrast, the present study used the more physiological temperature of 37°C to maximize vap gene expression during in vitro growth, hence minimizing the likelihood of identifying differences due only to dissimilar growth conditions.

The exact role of each Vap protein remains to be determined, but the proteins are probably involved in preventing acidification and late endosomal maturation of R. equi containing vacuoles within macrophages (10, 36). Regulation of expression of the vap genes is complex and depends on at least five environmental signals, temperature, pH, oxidative stress, magnesium, and iron (4, 5, 32, 34). The vapA and vapD genes are the major acid-inducible determinants carried by the virulence plasmid (4). In another study, vapA and vapG were predominantly induced by H2O2 treatment (5). These findings have led to the hypothesis that VapA, VapD, and VapG may play a dominant role in protection against macrophage-related stresses (5). That vapA, vapD, and vapG were expressed at significantly higher levels than the other vap genes in the lungs of infected foals in the present study is consistent with this hypothesis. These three genes were also significantly more expressed in vivo than in vitro.

Several lines of evidence suggest that immune responses against VapA and potentially other Vap proteins confer a role in protection (17, 24). A recent study with mice has shown that DNA immunization with vapA protects against R. equi infection and that the immunoglobulin G subtype response is consistent with a Th1-based immune response (13). A similar DNA vaccine containing the vapA gene has been shown to induce strong cell-mediated immune responses in adult horses (25). In foals, production of antibody to VapA and VapC but not of that to other Vap proteins increases following natural exposure to R. equi (18). The present study shows that VapA and VapC are also the proteins inducing the strongest lymphoproliferative recall responses following experimental challenge for both foals and adult horses. A prior study had demonstrated strong lymphoproliferative recall responses to VapA in association with clearance of R. equi in adult horses (24). The present study confirmed this finding and also demonstrated that lymphocytes from foals and from adult horses proliferate similarly in response to each Vap protein.

The documented importance of IFN-γ in protection against infection with R. equi (22) and the recognized Th2 bias in immune responses of neonates from many species (1) have led to the widespread hypothesis that a similar Th2 bias may be at the basis of the peculiar susceptibility of foals to infection by R. equi. The recent finding that young foals are deficient in their ability to produce IFN-γ in response to mitogens led to the conclusion that neonatal foals may also be predisposed to develop a Th2-like immune response (6). In one study, pulmonary lymphocytes from adult horses collected 7 days following challenge with R. equi expressed predominantly IFN-γ, but they also expressed IL-4 mRNA in response to in vitro stimulation with VapA (24). Similar results were obtained following stimulation of BLN lymphocytes with VapA in the present study. In addition, the present study extends these findings to the other Vap proteins of R. equi. The present study also demonstrates that BLN lymphocytes from foals can express IFN-γ in response to stimulation with each Vap protein just as well as those from immune adult horses. In fact, as a result of the significantly lower level of IL-4 expression for foals than for adult horses, foals showed a significantly higher IFN-γ/IL-4 ratio than adult horses in response to stimulation with most Vap proteins. The findings in the present study, therefore, do not support the conclusion that neonatal foals are predisposed to develop a Th2-like immune response. The reason for the peculiar susceptibility of foals to R. equi must be more complex and also must take into account the unique association of the VapA-containing virulence plasmid with disease in foals.

This strong predominating IFN-γ response in a neonate is not unprecedented. Many studies have shown that despite an apparent Th2 bias, neonates can mount strong Th1 responses to some antigens (1, 27). For example, Mycobacterium bovis BCG, a pathogen closely related to R. equi, triggers a Th1 response in human neonates of a magnitude similar to that seen when it is given later in life (26). Virulent R. equi is widespread in the environment of horse breeding farms (28, 33), yet development of clinical disease is the exception rather than the rule. The fact that oral administration of live virulent R. equi to newborn foals confers almost complete protection against subsequent heavy intrabronchial challenge (8, 19) and the fact that most foals on farms where it is endemic do not develop disease or develop subclinical disease and eventually clear the infection are consistent with the results of the present study, which demonstrate that foals develop adequate cell-mediated immune responses to key antigens of R. equi.

In conclusion, the data presented here indicate that vapA, vapD, and vapG are the most biologically relevant vap genes because they are preferentially induced during infection in the natural host. All the Vap proteins are immunogenic, with VapA and VapC providing the strongest lymphoproliferation stimulus. The peculiar susceptibility of foals to infection by R. equi cannot be explained by a failure to mount Th1 immune-reactivity to the Vap proteins. Further work is required to identify the fundamental host basis of the susceptibility of foals to R. equi pneumonia.

ACKNOWLEDGMENTS

This project was supported by the Morris Animal Foundation and by the Florida Department of Regulation Pari-Mutual Wagering Trust Fund. We thank the Florida Thoroughbred Breeders’ and Owners’ Association for support of the equine research breeding herd.

REFERENCES


