Immunoglobulin G Subisotype Responses of Pneumonic and Healthy, Exposed Foals and Adult Horses to *Rhodococcus equi* Virulence-Associated Proteins

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*Rhodococcus equi* causes severe pyogranulomatous pneumonia in foals and in immunocompromised humans. Replication of virulent isolates within macrophages correlates with the presence of a large plasmid which encodes a family of seven virulence-associated proteins (VapA and VapC to VapH), whose functions are unknown. Although cell-mediated immunity is thought to be crucial in eliminating *R. equi* infection, antibody partially protects foals. The antibody response to both VapA and VapC was similar in six adult horses and six naturally exposed but healthy foals, as well as in eight foals with *R. equi* pneumonia. The immunoglobulin G (IgG) subisotype response of pneumonic foals to Vap proteins was significantly IgGb biased and also had a trend toward higher IgGT association compared to the isotype association of antibody in adult horses and healthy exposed foals. This suggests that in horses, IgGb and IgGT are Th2 isotypes and IgGa is a Th1 isotype. Furthermore, it suggests that foals which develop *R. equi* pneumonia have a Th2-biased, ineffective immune response whereas foals which become immune develop a Th1-biased immune response. Pneumonic foals had significantly more antibody to VapD and VapE than did healthy exposed foals. This may indicate a difference in the expression of these two Vap proteins during persistent infection. Alternatively, in pneumonic foals the deviation of the immune response toward VapD and VapE may reflect a bias unfavorable to *R. equi* resistance. These data indicate possible age-related differences in the equine immune response affecting Th1-Th2 bias as well as antibody specificity bias, which together favor the susceptibility of foals to *R. equi* pneumonia.

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*Rhodococcus equi* causes pyogranulomatous bronchopneumonia in foals younger than 4 months and induces significant economic losses on endemically infected horse-breeding farms (29). This gram-positive, facultatively intracellular bacterium is an opportunistic pathogen in immunocompromised humans, such as those infected with human immunodeficiency virus (2, 9, 13). Foal-virulent *R. equi* strains possess an 81-kb plasmid and express VapA, a plasmid-encoded, surface-expressed lipoprotein (38, 40–42). VapA is a member of a family of seven Vap proteins (VapA and VapC to VapH), which have homology in their C-terminal halves and are encoded within a 27-kb pathogenicity island on the large plasmid (37). The function of these proteins is not known.

Protection of foals against *R. equi* appears to rely on cooperation and interdependency between the antibody-mediated and cell-mediated immune response. Antibody appears to contribute to protection since the period of maximum susceptibility coincides with declining levels of maternally derived antibody (29). In addition, antibody opsonizes *R. equi* for uptake and killing by macrophages and neutrophils in vitro (45, 47, 48) and may facilitate the killing of *R. equi* in vivo. Vap-specific antibodies protect immunosuppressed mice since purified immunoglobulin G (IgG) from APTX (a VapA-enriched antigen)-vaccinated horses protected against intraperitoneal challenge with *R. equi* whereas nonimmune equine IgG failed to provide protection (10).

Pulmonary clearance of virulent *R. equi* in mice requires functional T lymphocytes (3, 46). Both CD4+ and CD8+ T cells apparently contribute to protection (26, 33), and CD4+ cells are necessary for complete pulmonary clearance of *R. equi* in mice (16). Mice in which a Th2 cytokine response was induced by administration of monoclonal antibodies against gamma interferon (IFN-γ) prior to experimental infection with virulent *R. equi* failed to clear the bacteria and developed pulmonary granulomas (17). In contrast, immunocompetent BALB/c mice developed a Th1 cytokine response and cleared the infection (17). Adoptive transfer of *R. equi*-specific Th1 or Th2 cell lines in mice supported the conclusion that a Th2 response is detrimental whereas a Th1 response is beneficial in clearance of *R. equi* (18). Virulent *R. equi* can modulate the cytokine response in foals, down-regulating IFN-γ mRNA expression in CD4+ T cells and up-regulating lung interleukin-10 expression (12). These cytokines may influence the Th1-Th2 balance of the immune response in foals.

Although antibody provides partial protection against *R. equi* pneumonia (14), the role of the Ig isotype has not been described. Prescott et al. determined that the antibody response of young foals to the APTX antigen with aluminium hydroxide adjuvant induced a more IgGb- and IgGT-biased subisotype response than did natural infection, which induced an IgGa-dominant response (30). Vaccination with this antigen and adjuvant exacerbated disease in the foals after natural challenge with *R. equi* (30). An aluminium hydroxide-based influenza vaccine also induced an IgGT-biased response in horses, with some evidence of an IgGc response. These vacci-
nated horses were not resistant to infection even though they had an anamnestic IgG response (25). In contrast, natural infection with influenza virus induced virus-specific IgG and IgGb (25).

In mice and humans, the antibody isotype reflects the Th1-Th2 bias of the immune response. A similar bias may occur in horses. The IgG subclass profile of the antibody response associated with protective immunity to *R. equi* infection has not been described. The study reported here addresses the hypothesis that resistance or susceptibility to *R. equi* pneumonia in foals is associated with distinct IgG subclass-related antibody responses to the seven virulence-related Vap proteins and that in pneumonic foals the profile reflects a Th2-biased response whereas in healthy foals and adults the profile reflects a Th1-biased response.

**Materials and Methods**

**Experimental design.** Serum was collected biweekly from clinically normal pony foals (*n = 6*) kept on pasture at a University of Guelph research farm that had a history of *R. equi* infections in foals. The serum sample in which the peak antibody response to VapA was observed over the first 6 months of a foal’s life was used to determine the isotype profile of the anti-Vap response compared to sera obtained from a group of clinically normal, unrelated adult horses (*n = 6*) at the same farm and from a third group of foals with clinical *R. equi* pneumonia (*n = 8*). The clinical case samples were obtained from client animals at the University of Florida and the Ontario Veterinary College. The horses were categorized according to defined resistance or susceptibility status to provide a framework on which to make correlations based on specificity and isotype relatedness of serum anti-Vap antibody. The three groups represent (i) a clinically normal, healthy *R. equi*-exposed foal group; (ii) a clinically normal and healthy adult group (and since adult horses do not become sick with *R. equi* pneumonia, presumably an immune group); and (iii) a susceptible group of foals that had developed *R. equi* pneumonia and were therefore assumed to be nonimmune. The use of the horses for this research complied with all relevant national guidelines and University of Guelph Animal Care Committee policies.

**Vap protein purification.** Vap proteins were obtained as glutathione S-transferase fusion proteins. Each Vap gene was amplified from the *R. equi* strain 103†-plasmid by PCR using primers described in Table 1. The primers contained a 5′-extension encoding restriction enzyme recognition sites for BamHI or EcoRI. Amplified products were digested with BamHI and EcoRI, and the digestion products were ligated to a similarly digested plasmid vector (pGEX-2T; Amersham Pharmacia Biotech, Baie d’Urfé, Quebec, Canada). The resulting plasmids were transformed into *Escherichia coli* XL1Blue (Stratagene, La Jolla, Calif.). These plasmids allowed in vitro production of recombinant Vap proteins fused to glutathione S-transferase. Fusion proteins were purified, using glutathione-Sepharose 4B beads (Amersham Pharmacia Biotech), and Vap proteins were cleaved from glutathione S-transferase using thrombin (Amersham Pharmacia Biotech). Protein production was assessed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis, Western blot analysis, and protein assay (Pierce, Rockford, Ill.).

Although cloning and protein production were successful for each Vap protein, purification using the glutathione-Sepharose 4B beads was not always achieved because of the inability of the fusion protein to bind to the beads. In these cases, a protein agglutination protocol was employed to concentrate the Vap protein. Briefly, a 100-ml overnight culture, which had reached an optical density at 600 nm (OD<sub>600</sub>) of 0.6, was stimulated with 5 mM isopropyl-β-D-thiogalactopyranoside (IPTG) for 3 h. The culture was then centrifuged at 1,500 × g for 15 min at 4°C, and the supernatant was decanted. The pellet was resuspended in 4 ml of 25% sucrose–50 mM Tris (pH 8.0) and frozen at −70°C. The suspension was then thawed quickly in a 37°C water bath, and 1 ml of a lysosome solution (10 mg of lysosome [Sigma-Aldrich, Oakville, Ontario, Canada] per ml of 250 mM Tris [pH 8.0]) was added. This mixture was incubated on ice for 45 min, and 30 ml of a 20% sucrose solution were added. The mixture was then centrifuged at 15,000 × g for 15 min at 4°C, and the supernatant was decanted. The pellet was resuspended in 500 μl of sterile water and analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis, Western blot analysis, and protein assay.

**Serological testing.** Equine sera were absorbed for *E. coli*-specific antibodies because Vap proteins produced by agglutination contained trace amounts of *E. coli* antigens. Briefly, *E. coli* XLI1Blue cells were grown on tryptic soy agar plates at 37°C for 24 h. The bacteria were washed from the plates with 5 ml of phosphate-buffered saline, using a glass rod. The OD of the bacterial suspension was measured and adjusted with phosphate-buffered saline to an OD600 of 2.0. The suspension was subjected to three rounds of 3 min of sonication and freeze-thawing and cultured to confirm killing. The suspension was then added in a 1:1 ratio to each serum sample and incubated at 4°C for 18 h with gentle rocking (Lab-line 3-D bidirectional rotator; Fisher Scientific, Nepean, Ontario, Canada). The sera were then centrifuged (14,000 × g for 30 min) to remove debris.

Titers of serum antibodies to each of the Vap proteins were determined by enzyme-linked immunosorbent assay (ELISA). The natural log of the dilution which produced an OD double that of the negative control was considered the titer of that sample. Each sample was run in duplicate, and the average titer of the group (adults, clinically ill foals, and clinically normal foals) was used in a comparison of the total antibody response (expressed as a ratio over the total antibody response to VapA) and the IgG subtype response to each of the Vap proteins. A ratio was used for the total antibody comparison to control for possible variation associated with differences in antibody avidity. It was not used for the comparison of isotype response because this would have removed most of the data. Ninety-six-well polystyrene plates (Immunolon 1; Dynatech Laborato-

**TABLE 1. Primers used for PCR of each of the seven members of the Vap family**

| Vap   | Primer 1                                  | Primer 2                                  |
|-------|-------------------------------------------|-------------------------------------------|
| VapA  | 5'-GGCGGATCCACATTTGCGACGTTTCT-3'          | 5'-CATGAATTCCTAGGCGGTGCGA-3'              |
| VapC  | 5'-GGCGGATCCAGAATGTTGTCGTCCTGTCGTC-3'     | 5'-CATGAATTCGGGAGGTGTTACCTTCCACGGC-3'     |
| VapD  | 5'-GGCGGATCCGCAGAGAGAccAAAGAATGTTT-3'     | 5'-CATGATTCTTGTTCTCCACGGCAGGCA-3'         |
| VapE  | 5'-GGCGGATCCAGAATGTTGTCGTCCTGTCGTC-3'     | 5'-CATGAATTCCTAGGCGGTGCGA-3'              |
| VapF  | 5'-GCGGATCCACCGACGTTTCTCTATCAAGCCA-3'     | 5'-CATGAATTCCTGTCGTCCTGGCAATC-3'          |
| VapG  | 5'-GGCAATTCAGAGAGGATCTGCGATTTGCTG-3'      | 5'-CATGAATTCCTGTCGTCCTGGCAATC-3'          |
| VapH  | 5'-CATGATTCTCGTGAACGGGCTGAAGCTC-3'        | 5'-CATGAATTCCTGTCGTCCTGGCAATC-3'          |

**Analysis.** Two-tailed Student’s t tests and LSD random permutation tests (analysis of variance) in SAS (SAS/STAT user’s guide version 6, 1990; SAS Institute Inc., Cary, N.C.) were used to analyze the data and to determine the significance of differences between groups (*P < 0.05*). The first test was used to test for significant differences in the actual titers of each group, and the second test was used to test for significant differences between the groups for each
RESULTS

Antibody response to each of the Vap proteins. Antibody to VapA and VapC increased following natural exposure to R. equi (Fig. 1). In contrast, antibody to VapF, VapG, and VapH did not change throughout the 6-month observation period and antibody to VapE was not detected. The level of anti-VapD antibody rose slightly in some foals a few weeks after the peak of the response to VapA and VapC. Figure 1 data from one foal are representative of the pattern of the response observed in six foals kept at pasture, which remained apparently healthy, although the response pattern shifted along the x-axis between foals, so that the day on which the anti-VapA response peaked differed between foals. For this reason, data for foals were not combined.

There was a marked antibody response to VapA and VapC by all groups in the study. The antibody response relative to anti-VapA was significantly lower in both foal groups than in the immune adult horses for anti-VapD, anti-VapE, anti-VapF, and anti-VapH (Fig. 2). The pneumonic foals had significantly more antibody to VapD and VapE than did the normal foals, which as a group had no antibody response to VapE and only a low response to VapD by one foal.

Subisotype of the IgG response. The antibody response to each of the proteins by the three groups of animals (pneumonic foals, clinically normal foals, and immune adults) was further studied by investigating the subisotype (IgGT, IgGb, IgGc, and IgGa) of IgG and the differences were analyzed by Student’s t test. With the exception of three pneumonic and two clinically normal foals responding to VapA, an IgGc subisotype response was not detected to any of the Vap proteins. The IgGT response of pneumonic foals was significantly greater than that of the immune adult group to VapA, VapC, and VapG (Fig. 3). The adults produced an IgGa- versus an IgGT- or IgGb-biased response against VapA, VapC, VapF, and VapH (Fig. 3). Although the pneumonic foals also produced IgGa against VapA and VapC, this group had an IgGb-dominant response to all the Vap family members (Fig. 3). The IgGb response of ill foals was significantly greater (P ≤ 0.05 by Student’s t test) than that of immune adults for VapA, VapC, VapD, VapE, and VapG and significantly greater than the normal foal group for VapC, VapE, and VapG (Fig. 3).

The mean IgGa-to-IgGb and IgGa-to-IgGT ratios against all of the Vap family members were consistently higher in the adults than in the pneumonic foals (Table 2) and were usually higher in adults than in healthy foals. When the immune adult group was compared to the pneumonic-foal group, the IgGa-to-IgGb ratio was as much as three times higher and the ratio difference was significant for VapA, VapC, and VapF. The IgGa-to-IgGb ratios in the healthy foals tended to also be higher than in the pneumonic foals, as did the IgGa-to-IgGT ratio. When the overall IgGb response to all the Vap proteins was compared between the horse groups by analysis of variance, the IgGb response was significantly (P ≤ 0.05) greater in the pneumonic-foal group than in both the clinically normal group and the adult group.

DISCUSSION

Antibody function differs by immunoglobulin subisotype (7, 22). The Ig subisotype bias of the response to infectious organisms reflects a finely regulated multifactorial system that...
steers the response to provide an environment appropriate for control of the particular organism (15). In effect, the isotype profile of the immune response reflects the overall type of response (Th1 or Th2) induced by the pathogen or vaccine (32). This division of the CD4+ T helper cell population into two complementary groups has been extensively investigated by using humans and mice (32). The relationship of the bias of the response (Th1/Th2) to the degree of resistance to specific pathogens or to the response to vaccination has become an important area of research since it may be possible to increase resistance to disease by steering the immune response (11).

Assuming that the Th1 bias is protective in equine infections by the intracellular pathogen *R. equi* as it is in mice, the results reported here support the conclusion that IgG antibodies reflect a Th2-like response since a significantly greater IgGT response to *R. equi* antigens VapA, VapC and VapG was observed in foals that had developed *R. equi* pneumonia than in the immune adult group. However, this difference was not noted between the two foal groups. The significantly greater overall amount of IgGb anti-*R. equi* antibody in the sera of pneumatic foals compared to adult and healthy foals suggests that the ratio of IgGa to IgGb is important in determining resistance, since the amounts of IgGa were similar in clinically normal foals, pneumatic foals, and adults (Fig. 3). These results suggest that a higher IgGa-to-IgGb anti-*R. equi* ratio (Table 2) reflects a Th1-biased immune response and greater protection from *R. equi* infection. This is an important finding, since a fundamental question regarding *R. equi* pneumonia in foals is why foals are particularly susceptible to disease since this infection is almost unique to foals (as well as occurring in immunodeficient individuals of other species). The present results support the hypothesis that foals develop *R. equi* pneumonia because an inappropriate Th1-dominant response to infection develops whereas foals and adults which develop a Th1 response become immune (and remain so as adults). It has been suggested that the neonatal period in general is marked by a high susceptibility to infections and that although neonatal T cells are immunocompetent, their differentiation is biased toward a Th2 profile under neutral conditions (19). Work with rodents and humans indicates that this susceptibility may be a result of a combination of factors including a greater costimulatory requirement of neonatal T cells than of adult T cells (1), differences in antigen handling by neonatal B cells and low major histocompatibility complex-peptide density which favors priming of Th2-type CD4 cells (23), the lack of anatomical structures (germinal centers) required for lymphoid cell maturation (8, 24), and an immature phenotype of neonatal B cells in comparison to adult B cells (21). In addition to these possible mechanisms which may or may not operate to produce a relative Th2 bias in foals, the *R. equi* virulence plasmid may drive a Th2-biased immune response. For example, Giguère et al. (12) reported that foals with severe pneumonia caused by virulent *R. equi* strains differed significantly from foals infected with avirulent *R. equi* strains in that the former had reduced amounts of IFN-γ mRNA in bronchial lymph node CD4+ T cells as well as enhanced quantities of interleukin-10 mRNA in the lungs. Since the presence of the virulence plasmid carrying *R. equi* Vap genes was the only difference between the virulent and avirulent *R. equi* strains used in the experimental infections, this suggests that an important function of the virulence plasmid involves driving an ineffective Th2 immune response so that foals develop pneumatic disease rather than clearing infection. One factor determining whether foals become immune (Th1 response) or develop disease (Th2 response) in response to virulent *R. equi* may be the dose of virulent bacteria initiating the infection. By analogy, the dose of *Mycobacterium tuberculosis* BCG has been shown experimentally to determine whether a Th1 or a Th2 response developed, with relatively low doses leading to an almost exclusively cell-mediated, Th1 response (28). It may also be predicted that, as in mycobacterial infections, individuals vary in their ability to mount Th1 and Th2 immune responses (36, 44).

Aluminum hydroxide induces a Th2-biased response in...
mice, humans, and other species, which is reflected in the Ig subisotype of the antibodies produced (5, 6, 43). It was previously demonstrated that vaccination of foals with APTX antigen in aluminum hydroxide was associated with exacerbation of R. equi pneumonia and induced a greater IgGb and IgGT response than did natural infection, which induced an IgGa-biased response (30). Serum antibody responses following vaccination of horses with an aluminum hydroxide-based influ-
TABLE 2. IgGa-to-IgGb and IgGa-to-IgGT ratios for mean IgGa-, IgGb-, and IgGT-associated antibody to each of the \textit{R. equi} Vap proteins for each horse group

| Group and ratio | Value of ratio for: |
|-----------------|---------------------|
|                 | VapA    | VapC    | VapD    | VapE    | VapF    | VapG    | VapH    |
| IgGa/IgGb       |         |         |         |         |         |         |         |
| Healthy adults  | 1.89 ± 0.61 | 1.76 ± 0.94 | 0.56 ± 0.77 | 3.75 ± 0.44 | 2.62 ± 0.44 |         |         |
| Pneumonic foals | 1.01 ± 1.75 | 0.50 ± 1.66 | 0.37 ± 0.85 | 0.53 ± 1.31 | 0.35 ± 0.91 | 0.35 ± 0.94 | 0.50 ± 1.24 |
| Healthy foals   | 1.30 ± 0.94 | 1.56 ± 0.94 | 0.00 ± 0.00 | 0.62 ± 0.91 | 0.67 ± 0.94 | 0.47 ± 1.36 |         |
| Significancea   | ac      | ab      | a       | b       |         |         |         |
| IgGa/IgGT       |         |         |         |         |         |         |         |
| Healthy adults  | 2.57 ± 0.88 | —       | 0.56 ± 0.77 | 3.75 ± 0.44 | 2.62 ± 0.44 |         |         |
| Pneumonic foals | 0.95 ± 1.07 | 1.10 ± 0.72 | —       | —       | 0.62 ± 0.78 |         |         |
| Healthy foals   | 1.35 ± 0.50 | 5.49 ± 0.98 | —       | —       | 1.25 ± 0.79 | 4.90 ± 1.64 |         |
| Significanceb   | ac      | b       |         |         |         |         |         |

* Dashes indicate an inability to calculate the ratio due to the absence of a response and hence a zero denominator.

* Significant differences (P ≤ 0.05 by Student’s t test) between the adult and pneumonic foal group are indicated by a, those between the pneumonic and healthy foal groups are indicated by b, and those between the adult and healthy foal group are indicated by c.

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