**Induction of Protective Immunity in Rabbits by Coadministration of Inactivated *Pasteurella multocida* Toxin and Potassium Thiocyanate Extract**

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*Pasteurella multocida* is a bacterial pathogen that causes rhinitis (snuffles), pneumonia, otitis media, septicemia, metritis, and death in domestic rabbits. Currently, there are no effective vaccines to prevent infection by this organism. Subcutaneous (s.c.) immunization with either exotoxin or thiocyanate extracts of *P. multocida* induces partial protection in rabbits. Since disease begins at mucosal sites, induction of local immunity may be important in preventing systemic disease. Little is known concerning the efficacy of intranasal (i.n.) administration of these antigens in inducing protective mucosal immunity to *P. multocida* in rabbits. The purpose of this study was twofold: (i) to investigate the effectiveness of vaccination with purified *P. multocida* toxin (PMT) and a potassium thiocyanate extract of *P. multocida* (CN) in combination and (ii) to evaluate the efficacy of administration of these antigens i.n. versus s.c. Forty-eight rabbits were randomly divided into eight different treatment groups. Rabbits received either one or both antigens by either s.c. or i.n. administration. Following vaccination, each group received an i.n. challenge of *P. multocida*. Rabbits vaccinated with both antigens i.n. or s.c. had a 100% survival rate, few or no bacteria in the liver and lungs, high serum immunoglobulin G (IgG) and IgM antibody titers, and significant numbers of IgG antibody-secreting cells (ASC) in the spleen and tracheobronchial lymph node. Rabbits vaccinated i.n. had significant nasal and bronchoalveolar lavage IgA antibody levels. Rabbits vaccinated with only one antigen, either PMT or CN, had lower antibody titers, moderate to severe liver and lung infections, and fewer ASC compared to rabbits receiving both antigens. Rabbits in the control groups had moderate to severe liver and lung infections. This study indicates that i.n. immunization with both PMT and CN induces an effective response against homologous *P. multocida* challenge.

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upper respiratory tract, the mucosal immune response is likely to be an important defense mechanism. Secretory IgA (sIgA) antibodies are abundant in mucosal secretions and function to inhibit microbial adherence to epithelial cells (22). sIgA is preferentially induced following mucosal immunization; thus, the production of sIgA following i.n. vaccination should help prevent bacterial colonization and subsequent infection. The objective of this research was twofold: (i) to determine if coadministration of CN and PMT offers better protection against Pasteurellosis in New Zealand White male rabbits than either one given alone and (ii) to evaluate the efficacy of i.n. versus s.c. administration in stimulating protective immunity.

MATERIALS AND METHODS

Experimental animals. Forty-eight New Zealand White male rabbits (Oryctolagus cuniculus) (Hazelton Research Products, Inc., Kalamazoo, Mich.) weighing 2 to 2.5 kg were used. The colony from which the rabbits were obtained was Pasteurella free. Rabbits were placed in individual stainless steel cages upon arrival and allowed to acclimate to their environment for 5 days. Commercial feed (Purina Lab Rabbit Chow 5321; PMI Inc., Richmond, Ind.) and tap water were supplied ad libitum. The use of rabbits in this study was authorized by the Purdue University Animal Care and Use Committee.

Inactivation and preparation of CN for inoculation. CN (1 mg/ml) was toxic at concentrations of 40 mg/ml when assayed with embryonic bovine lung cells and CN (Oxford Laboratories, Worthington, Minn.) was toxic at 1 mg/ml when assayed with the dengue dengue dengue dengue dengue dengue dengue dengue dengue dengue dengue von der Lippe isolate (29). Each dose of CN, whether administered i.n. or s.c., was prepared as a 1:1,000 dilution of 1% NaHCO3 and 50 mM KSCN (Fisher Scientific Co., Pittsburgh, Pa.) and by use of a DNA molecular probe for the Pasteurella multocida isolate (29). Briefly, P. multocida was grown to confluence on 5% horse blood agar (Becton Dickinson, Cockeysville, Md.) in a 37°C CO2 incubator for 24 h. After 24 h of incubation, 6 ml of 10% nalidixic acid (Norit X, Minco) and grown overnight at 37°C. The numbers of CFUs of P. multocida were counted on blood agar plates. To remove the supernatant, and the remaining pellet was washed three times with PBS and resuspended by vortexing in 1 ml of PBS.

Inactivation of CN was confirmed with the API 20E system (Biomerieux, St. Louis, Mo.). The identity of organisms recovered from cultures was confirmed with the API 20E system (Biomerieux, St. Louis, Mo.).

Pathologic and histologic evaluations. At necropsy, each lung lobe was scored for percentage of pneumonia lesions. The lungs were fixed in 10% buffered formalin. Sections of liver and lungs were taken for histologic examination. Blood and nasal wash samples were collected on days 0, 14, 28, and 52 by tilting a rabbit's head to one side, injecting 1 ml of saline gently into the upturned naris with a feeding needle, and allowing the washings to drip from the contralateral naris into a petri dish. This process was repeated for the other side. Washings were pooled and centrifuged at 200 x g for 10 min to remove cellular debris and the supernatant was placed in a tube and frozen at –20°C until analyzed. Blood and nasal wash samples were taken (i) prior to inoculation (day 0), (ii) 1 week following each inoculation (days 14, 28, and 42), and (iii) 3 days following challenge. Thighs with vidin-4 CFU/ml P. multocida (day 52). Eyes of all rabbits were necropsied on day 53.

Preparation and confirmation of challenge dose. Bacteria were diluted to an absorbance value equal to 0.01 CFU/ml on a standard curve and then diluted to the challenge dose on the day of challenge. Counting dilutions was performed by using triprep-10 concentrators (Millipore Corp., Bedford, Mass.) with a nominal pore-size filter (Amicon, Inc., Beverly, Mass.). The protein concentration of CN was determined by use of a bichinchoninic acid assay (Bio-Rad Laboratories, Hercules, Calif.) to bovine serum albumin (BSA) as the protein standard. In general, seven bacterial plates grown to confluence will yield 1 ml of CN at a concentration of 1 mg/ml, and in this study 70 plates were used.

Inactivation and preparation of PMT for inoculation. The day of inoculation, each 1-ml dose of PMT, whether administered i.n. or s.c., was prepared separately for each rabbit. One hundred microliters (5 μg/ml) of purified PMT (generous gift of Richard Richler, Ames, Iowa) was placed in a 1.5-ml microcentrifuge tube and inactivated in a 60°C water bath for 30 min. For i.n. administration of vaccines containing PMT, 1-ml doses contained 5 μg of inactivated PMT and 20 μg of CT (1 mg/ml) in sterile saline. For i.n. administration of vaccines containing PMT, 1 ml of PMT (50 μg) was inactivated, mixed with 22.5 μl of 0.01 M NaHCO3 and 0.2 M AIBO3 at pH 8.5, and vortexed. The mixture was microcentrifuged for 5 min (13,000 x g) to remove the supernatant. The pellet was washed three times with PBS, and the final pellet was brought to 1 ml with PBS. The remaining pellet was washed three times with PBS and resuspended by vortexing in 1 ml of PBS.

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absorbed with a 1:1,000 dilution of inactivated CN. They were then incubated overnight at 4°C.

On the following day, plates were washed three times with PBS-Tween, and rabbit anti-goat IgG (heavy and light chains) conjugated to alkaline phosphatase (Southern Biotechnologies) was diluted to 1:1,000 with PBS-Tween and absorbed with inactivated CN. Seventy-five microliters of conjugated antibody was added to each well. Plates were incubated for 3 h at room temperature and washed three times with PBS-Tween. Substrate was prepared by diluting 1 part 3% (wt/vol) agarose (ultrapure, high melting temperature; Gibco BRL, Grand Island, N.Y.) with 4 parts 5-bromo-4-chloro-3-indolylphosphate (5-BCIP) (Sigma). The agarose was first melted in a boiling water bath and then mixed with 5-BCIP. The mixture was kept at 40°C by placing the tube containing the mixture in a beaker containing hot water. One hundred microliters of the warm mixture was added to each well. Plates were incubated for 10 min at room temperature and placed in a humid chamber overnight at 4°C. Spots were counted with a Photo-Zoom inverted microscope (Cambridge Instruments), and the number of spots per six wells was averaged and recorded as the mean number of antibody-excreting cells (ASC) per 106 cells.

ELISA. Enzyme immunoassay-radioimmunoassay flat-bottom high-binding 96-well plates were coated with 100 μl of inactivated CN (2.5 μg/ml) per well as described above for the ELISpot assay. This quantity of antigen was chosen because it gave optimal enzyme-linked immunosorbent assay (ELISA) readouts for rabbit serum samples. On the following day, plates were washed and blocked with 0.01% BSA as described above. Plates were washed three times with PBS-Tween, serum samples were serially diluted from 1:100 to 1:6,400 and assayed in duplicate.

Each ELISA plate included six negative control wells and six positive control wells. Negative controls consisted of pooled sera obtained from six unimmunized rabbits that were used at a dilution of 1:120 with PBS-Tween (50 μl/well). For positive controls, sera obtained from six rabbits with high anti-CN titers were pooled and diluted to 1:100 with PBS-Tween.

Plates were incubated for 2 h at 37°C and washed three times with PBS-Tween. Goat anti-rabbit IgG, IgA, and IgM sera were diluted 1:20,000 with PBS-Tween and absorbed with inactivated CN at a 1:1,000 dilution. Fifty microliters of each isotype was added to each well of the plate and incubated overnight at 4°C. Plates were washed three times with PBS-Tween, and 50 μl of rabbit anti-goat IgG (heavy and light chains) conjugated to alkaline phosphatase, diluted 1:1,000 in PBS-Tween, and absorbed with inactivated CN was added to each well. Plates were incubated for 3 h at room temperature and washed three times with PBS-Tween, and 50 μl of the substrate p-nitrophenyl phosphate (Sigma) (1 mg/ml) was added to each well and incubated for 15 min at 25°C.

The absorbance of each well was recorded at 405 nm with a 96-well microplate reader (Molecular Devices, Menlo Park, Calif.). Results for serum were reported as end-point antibody titers. The titer was designated the reciprocal of the last end-point dilution. Each BAL and nasal wash sample was assayed in triplicate.

Statistical analysis. All results were analyzed with Statistica for Windows (StatSoft, Tulsa, Okla.). Multivariate analysis of variance (MANOVA) was used to analyze between- and within-group interactions for serum, nasal wash, BAL, and ELISpot results, with a P value of <0.05 being considered significant. Within-group interaction took into consideration the repetitive measure (antibodies) for each treatment group at each sampling period. Analysis of variance (ANOVA) was used to analyze between-group interaction for severity of pneumonia and CFU of P. multocida recovered from liver and lungs, with a P value of <0.05 being considered significant.

RESULTS

Survival rate after challenge with P. multocida. Following challenge, 100% of rabbits in the inPMT, scPMT, inCN, inPMTCN, and scPMTCN groups survived. However, only 50% of rabbits in the scCN group, 33% of rabbits in the inCo group, and 0% of rabbits in the scCo group survived to day 53. All rabbits in the scCo group had severe pneumonia and purulent discharge from the nostrils and eyes, while rabbits in the inCo group displayed signs of lethargy, anorexia, and discharge from the nostrils. With the exception of rabbits in the scCN group, rabbits immunized with either PMT or CN survived challenge and did not appear to develop nasal or ocular discharge.

P. multocida cultured from the liver and lungs of challenged rabbits. The numbers of CFUs of P. multocida cultured from the liver and lungs of each rabbit were converted to log10 values (Tables 1 and 2). Rabbits in the inPMT group and the scPMT-CN group had no P. multocida cultured from the liver. Furthermore, the two PMT- and CN-vaccinated groups had significantly fewer CFU recovered from the liver than the scCo group [P < 0.03].

In contrast to rabbits in the inPMT-CN and scPMT-CN groups, which had the lowest numbers of CFUs of P. multocida recovered from the lungs (Table 2), rabbits in the scCN, inCN, inCo, and scCo groups had the largest amounts of P. multocida cultured from the lungs (P < 0.02).

P. multocida-positive cultures from the bullae and nasal pharynx following challenge. P. multocida was cultured from the bullae of none of the rabbits in the scPMT-CN group, 16% in the inPMT-CN group, 40% in the inCo group, 50% in the inCN, scCN, and inPMT groups, and 60% in the scPMT group. When the nasal pharynx was cultured, the inPMT-CN group had the lowest culture-positive rate (33%), followed by the inCo group (80%), the inPMT and scPMT-CN groups (83%), and the scPMT, inCN, and scCN groups (100%). No data were collected for the scCo group, since all of the rabbits died 3 days postchallenge. In general, rabbits which were vaccinated with both antigens had the lowest culture-positive rates.

Summary of lung lesions. Lung lesion scores were ranked according to severity. The inPMT-CN group had the lowest severity score (0.1), followed by the scPMT-CN group (0.2), the
scPMT group (0.5), the inCN group (2.8), the inPM group (3.6), the inCo group (4.4), the scCN group (10.3), and the scCo group (17.5). Rabbits in the scCN and scCo groups had significantly higher lesion severity scores (*P*, 0.02) than those in all the other groups.

Serum end-point antibody titers. Following the second vaccination period, rabbits receiving CN, either alone or in combination with PMT, had an increased IgG level (Fig. 1), and most rabbits receiving CN had an increase followed by a decrease in the IgM level (data not shown). No group had a significant increase in serum IgA antibody titers over time (data not shown). Control rabbits or rabbits receiving PMT alone had little or no increase in anti-CN antibody titers following each subsequent immunization. Following challenge with live *P. multocida*, rabbits receiving any form of CN had an increase in the IgG level (Fig. 1). Furthermore, rabbits which received any form of CN had an average twofold-higher IgG level than control rabbits or rabbits receiving PMT alone (*P*, 0.05).

Percent positive nasal wash antibody titers. Preimmunization titers were not significantly different between groups for any isotype. The inPMTCN group had the greatest IgA response of all groups both after the first inoculation and following challenge (Fig. 2). The greatest increase in IgG antibody levels was for rabbits vaccinated i.n. or s.c. with CN or PMTCN (Fig. 3), while only rabbits receiving both antigens had a significant increase in IgM antibody levels (*P* < 0.04) (data not shown).

Throughout the study, the inPMTCN group consistently had higher IgA and IgG antibody titers than all the other groups, except for the third immunization, after which the inCN group had the highest IgA and IgG antibody titers (Fig. 2 and 3). Following challenge with *P. multocida*, the inPMTCN group had the highest IgA and IgG antibody titers. In general, rabbits which received any form of CN had higher IgA and IgG antibody titers than rabbits in the two PMT and control groups (*P*, 0.05).

BAL isotype-specific antibody titers. Rabbits receiving CN or PMTCN i.n. or i.m. had the highest levels of IgG antibodies in BAL (*P* < 0.04) (Fig. 4). The inCN and inPMTCN groups had a significant (*P* < 0.05) increase in IgA antibody levels, approximately three times higher than in the other groups. The inPM, inCN, inPMTCN, and scPMTCN groups had the highest levels of IgM antibodies.

### TABLE 2. Mean *log*<sub>10</sub> CFU of *P. multocida* per g of tissue recovered from the lungs of challenged rabbits<sup>a</sup>

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>Mean <em>log</em>&lt;sub&gt;10&lt;/sub&gt; CFU/g of lung</th>
<th>Statistical significance (<em>P</em>) of difference between groups</th>
</tr>
</thead>
<tbody>
<tr>
<td>inPMT</td>
<td>2.8</td>
<td>0.01</td>
</tr>
<tr>
<td>scPMT</td>
<td>2.9</td>
<td>0.01</td>
</tr>
<tr>
<td>inCN</td>
<td>3.4</td>
<td>0.01</td>
</tr>
<tr>
<td>scCN</td>
<td>3.9</td>
<td>0.01</td>
</tr>
<tr>
<td>inPMTCN</td>
<td>0</td>
<td>0.01</td>
</tr>
<tr>
<td>scPMTCN</td>
<td>0.7</td>
<td>0.03</td>
</tr>
<tr>
<td>inCo</td>
<td>3.4</td>
<td>0.01</td>
</tr>
<tr>
<td>scCo</td>
<td>5.9</td>
<td>0.01</td>
</tr>
</tbody>
</table>

<sup>a</sup> See the legend to Fig. 1 for details. ANOVA was used to analyze between-group interactions, with a *P* value of <0.05 being considered significant. Individual *P* values are shown.

![Fig. 1. Mean *log*<sub>2</sub> serum anti-CN IgG antibody titers. Rabbits were vaccinated (on days 7, 21, and 35, except for groups receiving PMT, which were vaccinated on days 7 and 35) either i.n. or s.c. with either PMT or CN alone or in combination with each other (PMTCN) or with a control (Co) (aluminum hydroxide for s.c. administrations and saline for i.n. administrations). Serum samples were collected prior to vaccination, 1 week after each vaccination, and 3 days following challenge. Rabbits were challenged i.n. with virulent *P. multocida* on day 49. Samples were analyzed by an ELISA as described in Materials and Methods. Between- and within-group interactions were analyzed by MANOVA, with a *P* value of <0.05 being considered statistically significant. Lowercase letters signify comparisons made between groups, with different letters signifying statistical significance between groups at each time period. Vacc, vaccination.]
ELISpot assay results. The numbers of spots detected in the ELISpot assay were averaged and recorded as the number of ASC per 10^6 tissue cells. No significant amounts of ASC were produced in the MLN or tonsils of any group for any isotype. However, differences were detected for the spleen and TLN. Rabbits receiving both antigens had significantly more anti-CN IgG ASC in the spleen than rabbits in all the other groups (Table 3). The number of anti-CN IgG ASC produced in the TLN by the scCN group was larger than that in any other group (Table 4). Furthermore, the inPMT, inCN, inPMTCN, and scPMTCN groups had a significantly larger number of anti-CN IgG ASC in the TLN than the scPMT group. The inPMTCN group was the only group to produce a significant amount of anti-CN IgM ASC in the spleen (127 ASC/10^6 spleen cells; P < 0.01), 3 to 12 times more than any other group (data not shown).

DISCUSSION

One of the most difficult aspects of pasteurellosis and other P. multocida-related infections is that they can be caused by various toxigenic and nontoxigenic serotypes of the bacterium. Most infections are caused by type A P. multocida, but type D is also a prominent serotype responsible for the development of atrophic rhinitis in pigs and rabbits (13). Due to the ability of various serotypes to induce disease, vaccination becomes difficult and immunity against one serotype does not necessarily result in cross-protection against a heterologous serotype. However, this is the first study to demonstrate that coadministration of PMT and CN antigens stimulates homologous protective immunity to pasteurellosis in rabbits. Rabbits which were vaccinated with both antigens, whether by i.n. or s.c. inoculation, had the best protection, as indicated by high sur-
vival rates, few or no *P. multocida* organisms cultured from the middle ear, liver, or lungs, and minimal histologic lung lesions.

Protection cannot be directly linked to any one isotype or antibody response, since several groups had similar serum, nasal wash, or BAL titers. This finding suggests a possible role for cell-mediated immunity. Enhanced alveolar macrophage or T-cell responses are important for the clearance of bacterial infections (7, 26), and this study supports previous findings which demonstrated a direct correlation between decreased bacterial load and enhanced survival of challenged animals (7).

Rabbits vaccinated with PMT alone, whether s.c. or i.n., still showed partial protection against challenge, as indicated by low lung lesion scores and high survival rates, supporting the results of previous studies (28). However, these groups had more bacteria cultured from their tissues than those which received both antigens, indicating that PMT and CN together induce greater protective immunity.

In general, rabbits which were immunized, regardless of antigen or route of administration, had a 100% survival rate following challenge with homologous *P. multocida*, with the exception of the scCN group, which had only a 50% survival rate. It is surprising to find that the scCN group had such a low survival rate and severe lung lesions, given that the rabbits in this group had high antibody titers. However, samples were analyzed for the whole isotype present and not for the specific subclass of an isotype produced in response to the CN antigen. Therefore, even though the rabbits in the scCN group had high serum IgG antibody titers, they may not have mounted the appropriate subclass of IgG to opsonize *P. multocida* efficiently. Poor opsonization could lead to severe systemic infections by the bacterium. Lack of effective cell-mediated immunity could also be a factor. Previous studies have shown that the administration of antigen stimulates macrophage activity and results in greater protection against challenge by *P. aeruginosa* in rats (7).

Rabbits in the scCo group had the largest amount of *P. multocida* cultured from the liver and lungs. They also had one of the most severe lung lesion scores and a 0% survival rate, while the inCo group had an 83% survival rate. The high survival rate of the inCo group suggests that these rabbits were partially protected. The reason for this result is not obvious, but it is possible that i.n. inoculation with a feeding needle results in minor tissue damage and the release of cytokines that provide partial, nonspecific protection. Another possibility is that a few rabbits in the inCo group were exposed to PMT via a gastric feeding needle which was used to immunize the in-PMT rabbits. However, none of the control animals had significant levels of CN-specific antibodies in serum, nasal wash, and BAL.

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>Mean no. of anti-CN IgG ASC per 10^6 spleen cells</th>
<th>Statistical significance (P) of difference between groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>inPMT</td>
<td>scPMT</td>
</tr>
<tr>
<td>inPMT</td>
<td>59</td>
<td></td>
</tr>
<tr>
<td>scPMT</td>
<td>19</td>
<td></td>
</tr>
<tr>
<td>inCN</td>
<td>79</td>
<td></td>
</tr>
<tr>
<td>scCN</td>
<td>155</td>
<td>0.01</td>
</tr>
<tr>
<td>inPMTCN</td>
<td>196</td>
<td>0.01</td>
</tr>
<tr>
<td>scPMTCN</td>
<td>164</td>
<td>0.01</td>
</tr>
<tr>
<td>inCo</td>
<td>75</td>
<td></td>
</tr>
<tr>
<td>scCo</td>
<td>NA</td>
<td></td>
</tr>
</tbody>
</table>

*See the legend to Fig. 1 for details. MANOVA was used to analyze between- and within-group interactions, with a P value of <0.05 being considered statistically significant. Individual P values for between-group comparisons are shown. NA, rabbits did not survive to day of necropsy, and no data were available.*

FIG. 4. Mean percent positive BAL anti-CN IgA, IgG, and IgM antibody titers. See the legend to Fig. 1 for a description of the study. Four days following challenge with live *P. multocida*, BAL was collected for each rabbit. ELISA was used to analyze lavage samples, and MANOVA was used to determine between- and within-group interactions, with a P value of <0.05 being considered statistically significant.
or BAL samples. Even though survivors in the inCo group had some anti-CN IgG ASC in the spleen and TLN, they were ill at the time of necropsy, displaying signs of lethargy, anorexia, and nasal discharge. Nonspecific immune features were most likely responsible for the decrease in bacterial numbers and the increase in the survival rate of the inCo group.

High levels of IgA antibodies were present in nasal secretions of the inPMTCN group and in BAL secretions of the inCN and inPMTCN groups. This finding confirms those of numerous studies showing that mucosal but not s.c. inoculation results in the production of IgA antibodies (6, 29). i.n. immunization causes an increase in IgA antibody levels in the nasally associated lymphoid tissue (NALT) in the upper respiratory tract (32) as well as in the bronchially associated lymphoid tissue in the lower respiratory tract (1, 2, 15). These two inductive sites allow for isotype switching and differentiation of B cells to IgA-secreting cells. NALT plays an important role in that it has the capacity to aid in B-cell maturation and differentiation as well as to maintain immune memory (32). Furthermore, the presence of IgA antibodies has been shown to be important for blocking bacterial attachment and antigen uptake across mucosal membranes (22). If the bacteria can be sequestered and destroyed at the site of infection, then further systemic damage and possible death can be prevented. However, the presence of IgA alone does not correlate highly with protection, as both i.n. and s.c. immunizations induced protective immunity in rabbits.

Even though i.n. immunization with PMTCN is an effective way to control infection, the method of vaccine delivery is not necessarily practical, especially when vaccinating a large number of rabbits. However, the efficacy of mucosal vaccination suggests that it may eventually be possible to deliver these antigens by alternative routes, such as orally, to induce mucosal immunity in respiratory tracts of rabbits (29). Furthermore, oral delivery of a vaccine could provide a safe, practical, and effective means of protecting rabbits without concern for the development of injection site reactions or discomfort from i.n. administration.

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REFERENCES


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**TABLE 4. Mean number of anti-CN IgG ASC per 10⁶ TLN cells produced in response to challenge with virulent P. multocida**

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>Mean no. of IgG ASC/10⁶ TLN cells</th>
<th>Statistical significance (P) of difference between groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>inPMT</td>
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<td>inPMT</td>
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<td>scPMT</td>
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<td>inCN</td>
<td>89</td>
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<td>scCN</td>
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<td>inPMTCN</td>
<td>107</td>
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<td>scPMTCN</td>
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</tr>
<tr>
<td>inCo</td>
<td>72</td>
<td>NA</td>
</tr>
<tr>
<td>scCo</td>
<td>NA</td>
<td></td>
</tr>
</tbody>
</table>

*See the legend to Fig. 1 for details. MANOVA was used to analyze between- and within-group interactions, with a P value of <0.05 being considered significant. Individual P values for between-group comparisons are shown. NA, rabbits did not survive to day of necropsy, and no data were available.*

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