Adoptive Transfer of Lymphocytes Sensitized to the Major Surface Glycoprotein of *Pneumocystis carinii* Confers Protection in the Rat

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Abstract

*Pneumocystis carinii* is a major opportunistic pathogen and a leading cause of morbidity in patients with AIDS. CD4+ cells have been shown to be important in host defenses against *P. carinii*, but the antigen(s) involved with this response have not been identified. We undertook the present study to determine whether the major surface glycoprotein (MSG) of *P. carinii* contains epitopes that can elicit a protective cellular immune response. Spleen cells and purified CD4+ cells isolated from Lewis rats, pulsed 1–4 d with MSG, and injected into corticosteroid-treated Lewis rats with pneumocystosis resulted in significant reduction in the *P. carinii* burden, as judged by organism quantitation and lung histology. The protective response demonstrated by the donor cells was dependent on previous exposure to *P. carinii*, cell concentration, and time of incubation with MSG. In addition, reconstitution with MSG-specific CD4+ cells resulted in an early hyperinflammatory response within the lungs of these animals with a high percentage of mortality. Thus, in this model, MSG can elicit an immune response mediated by CD4+ cells, which has a harmful as well as helpful effect on the host, and these responses occur despite the presence of corticosteroids. (*J. Clin. Invest. 1995. 95:2587–2593*) Key words: AIDS • immunity • T cells • pneumonia • opportunistic infection

Introduction

*Pneumocystis carinii* is a major opportunistic pathogen and a leading cause of morbidity in patients with AIDS. Impaired cell-mediated immunity is considered to be the major host predisposing factor for *P. carinii* pneumonia (1). Major underlying conditions in *P. carinii* patients include AIDS, malignancies, protein malnutrition, and immunosuppressive drugs (particularly corticosteroids). In AIDS patients, the risk of developing *P. carinii* pneumonia rises with the progressive decline in the number and function of CD4+ T cells (2). In previous animal studies, Furuta (3) found that the transfer of T cells to *P. carinii*-infected nude mice enhanced clearance of pneumocystosis. Other studies have shown that *P. carinii*-infected severe combined immunodeficiency (SCID)1 mice reconstituted with spleen cells isolated from immunocompetent mice were able to resolve the infection (4). Further work using SCID mice demonstrated that reconstitution with CD4+ cells resulted in complete clearance of *P. carinii* from the lung (5). These data suggest the importance of adoptive transfer of immunocompetent cells in the clearance of *P. carinii* pneumonia. The role, if any, that specific *P. carinii* antigens play in this process has not yet been investigated.

Much of the immunological work on *P. carinii* has focused on a major surface glycoprotein referred to as MSG or gpA. MSG, which actually represents a family of related proteins encoded by unique genes (6–9), is highly immunogenic and elicits both humoral and cellular immune responses (10–14). It has been demonstrated that MSG is a specific T cell antigen and significantly increases the percentage of CD4+ T cells during the proliferative response (14). MSG also elicits the secretion of cytokines, suggesting the importance of this antigen in host T cell recognition of *P. carinii* by recruitment of inflammatory cells and cytokine production.

In the present study, we have tested the hypothesis that MSG contains epitopes which can elicit a protective cellular immune response. Since MSG has been shown to stimulate an immune response in exposed subjects, protective epitopes contained in this antigen might become attractive as vaccine targets. The secondary aim of this study was to determine whether the corticosteroid-treated rat could be used to analyze this immune recognition and clearance of *P. carinii* from the lungs.

Methods

Animals. Male Lewis rats (viral antibody–positive) were bred and raised in a conventional colony with food and water ad lib. at the Veterans Affairs Medical Center, Cincinnati, OH with ample environmental exposure to *P. carinii*. At 6–8 wk of age, 150–200 g, *P. carinii* infection was induced by subcutaneous injections of 4 mg/0.4 ml Depo-Medrol (The Upjohn Co., Kalamazoo, MI) every seventh day. Ampicillin (1 mg/ml) was given in the water to control secondary bacterial infections. After 7 wk of immunosuppression, animals received the donor cells and continued on corticosteroid treatment for 10 wk. At this time, animals were killed by an overdose of CO2 and the lungs removed as described below. In each experiment, at least 10 animals were in each treatment group. Effectiveness of the adoptive transfer was based on histologic and quantitative analysis of the severity of *P. carinii* infection rather than on survival because some rats died from causes unrelated to *P. carinii* (e.g., other opportunistic infections, injection procedure). Only animals that survived 10 d after adoptive transfer were included in the data analysis.

Antigen preparation. After 8–10 wk of immunosuppression, *P. carinii* organisms were isolated from the lungs of corticosteroid-treated

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1. Abbreviations used in this paper: MSG, major surface glycoprotein; SCID, severe combined immunodeficiency.

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Sprague–Dawley rats as described (14). Briefly, lungs were removed, minced, and ground through a 60-mesh wire screen in PBS. Erythrocytes were lysed by exposure to 0.85% ammonium chloride, centrifuged, and the pellet washed twice with PBS. Samples of homogenates were streaked on Mueller–Hinton and Sabouraud dextrose agar plates for detection of any bacterial or fungal contaminants. Specimens with no detectable contamination were pooled and digested with Zymolase 100 T (ICN Biomedicals Inc., Costa Mesa, CA) for 30 min at room temperature. The treated organisms were then fractionated by differential centrifugation. MSG was purified from the 100,000 g supernatant by HPLC. MSG was isolated on the basis of size by injecting 100,000 g supernatants onto a Macroscope GPC 150 column (Alltech Associates Inc., Deerfield, IL) under isocratic conditions in 0.1 M KH2PO4 (pH 7.0), 0.2 M NaCl as previously described (14). MSG migrated as a band with a molecular weight of 120 kD on SDS-PAGE under reducing conditions. The purity of the preparation was shown by the following: Coomassie blue staining; reactivity with MSG-specific monoclonal antibodies; and by the absence of endotoxin, as verified by the Limulus amebocyte assay (<0.125 U/ml; Whittaker M. A. Bioproducts,Walkersville, MD).

Donor cell culture and transfer. In each adoptive transfer experiment, donor spleen cells were isolated from two groups of six adult Lewis rats. The first group of animals was born and raised in a protected environment free from exposure to common viruses and P. carinii. The second age-matched group consisted of animals raised in a conventional colony providing sufficient environmental exposure to P. carinii. In all cases, the presence or absence of antibodies to MSG was confirmed by Western blot analysis of sera. Spleens were obtained from both sets of animals and cells dispersed into PBS by passage through a stainless steel screen. Erythrocytes were lysed by incubation in 0.85% ammonium chloride, the cells washed twice in PBS, and then plated in RPMI 1640 containing 2 mM l-glutamine, 100 U of penicillin/ml, 100 μg of streptomycin/ml, 5 × 10−3 M β-mercaptoethanol, and 10% heat-inactivated FCS. Cells were cultured for 1–7 d in the presence of MSG (10 μg/ml) or ovalbumin (10 μg/ml) to act as a nonspecific protein control. After culture, the cells were harvested and washed by centrifugation. Additionally, in one experiment, a standard proliferation assay was performed on the donor cells as previously described (14). Viability was determined by trypsin blue dye exclusion (>96%). Cells were injected intravenously in 200–μl vol into the tail vein of recipient Lewis rats that had undergone immunosuppression for 7 wk. In each experiment, all recipient animals received donor cells from the same spleenocyte preparation with the intravenous injections administered 3–5 d apart, depending on the length of incubation time of the donor cells. In all cases, recipient animals continued oral corticosteroid treatment 10 wk after adoptive transfer. Control animals received an injection of an equal volume of pyrogen-free normal saline.

Separation and FACS® analysis of cell populations. In one set of experiments, donor spleen cells were separated into distinct CD4+ and CD8+ cell populations before adoptive transfer. Cells were harvested following a 4-d incubation and separated using Cellset–plus Rat CD4+ or CD8+ affinity chromatography (BioTex Laboratories, Edmonton, Canada) according to the manufacturer’s instructions. The spleen cell suspensions were precipitated with either monoclonal mouse anti–rat CD8+ (for CD4+ purification) or monochlonal mouse anti–rat CD4+ (for CD8+ purification) on ice for 1 h. Cells were washed twice in pyrogen-free PBS and cell concentration adjusted to 5 × 105 cells/ml before loading onto the column. Purified populations of CD4+ or CD8+ cells were eluted in the flow-through from the columns. Samples from each purification were analyzed for purity before adoptive transfer.

Sample purity and percentage of subpopulations of T cells after incubation for 1–4 d with and without MSG were analyzed by FACS® analysis (Becton Dickinson and Co., Mountain View, CA). Briefly, 106 cells in 50 μl were incubated with a fluorescein-conjugated mouse anti–rat CD8+ mAb (PharMingen, San Diego, CA) and/or phycoerythrin-conjugated mouse anti–rat CD4+ mAb (PharMingen) at 4°C for 20 min, washed three times in cold HBSS, and analyzed by flow microfluorometry as previously described (15). Purity of each subset was >92%; however, the remaining 8% showed no detectable contamination by the alternate subset of T cells.

Evaluation of P. carinii. Assessment of P. carinii in the lungs was based on histologic and quantitative procedures previously reported in detail (16). A highly significant correlation was found by both methods of assessment. At death or time of killing, the lungs were removed and weighed. The left lung of each animal was infused with a 4% formaldehyde solution until fully expanded and fixed further in formaldehyde for several days before conventional paraffin embedding. Three horizontal 0.5-μm sections of the lung were stained with hematoxylin and eosin and Grocott-Gomori methenamine-silver. The lung sections were coded and read blindly. The following scoring system was used to evaluate P. carinii based on alveolar involvement: 0, no infection; 0.5, minimal (<1% alveoli); 1, light (1–25% alveoli); 2, moderate (26–50% alveoli); 3, severe (51–75% alveoli); and 4, very severe (>75% alveoli).

The right lung was used for quantitation of P. carinii cysts and nuclei as described by Cushion and Walzer (17). Lung pieces were minced and then homogenized by passage through a stainless steel 60 mesh screen. The homogenate was centrifuged at 1,000 g at 4°C, for 15 min. The supernatant was decanted and erythrocytes lysed by incubation in 0.85% NH4Cl. The organisms were then washed twice in PBS and slides were prepared for enumeration. Stains used to identify P. carinii were cresyl ech violet (CEV), which selectively stains the cell wall of the cyst, and Diff-Quik (DQ), which stains the nuclei of cysts, trophozoites, and intermediate forms. Three 10-μl drops were placed on a glass slide and stained with either CEV or DQ. Each drop covered ~1 cm2. 10 oil immersion fields in each drop were randomly scanned. The lower limit of detection by this evaluation method is 1.1 × 103 organisms per lung. Cyst counts were performed routinely; Diff-Quik counts were performed in selected instances. Specimens to be studied were coded and read in a blind manner.

Statistical analysis. Data are reported as the log transformation of the geometric mean±SD. Scalar comparisons were made by ANOVA using the Newman–Keuls multiple range test or Student’s t test. Ordinal comparisons were made by the Mann–Whitney or the Wilcoxin-paired sample test as appropriate by INSTAT (GraphPad Software for Science, San Diego, CA). Significance was accepted when the P value < 0.05.

Results
Adaptive transfer of P. carinii–exposed or unexposed lymphocytes. In the first experiment, we compared the effectiveness of administering donor spleen cells from Lewis rats with exposure to P. carinii; donor spleen cells from unexposed (naive) rats; or saline in the clearance of P. carinii pneumonia among recipient animals (Fig. 1). The log10 mean P. carinii cyst count among rats administered saline (9.29±0.35/lung) was very similar to the cyst counts for all groups of recipients administered splenocytes from naive animals (ranging from 8.92±0.69/lung to 9.08±0.25/lung). By contrast, a significant (P < 0.05) reduction in cyst counts was achieved in recipients administered splenocytes from rats which had been exposed to P. carinii; these groups included fresh cells and cells incubated for 24 h in the presence or absence of MSG or ovalbumin (ranging from 8.04±0.55/lung to 8.46±0.67/lung). Among each of these groups, a significantly lower (P < 0.05) log10 mean cyst count was achieved with the administration of exposed splenocytes than with the unexposed splenocytes. When the recipients administered the exposed spleen cells were compared, the greatest reduction was obtained with cells incubated with MSG (8.04±0.55/lung). However, the differences among these groups were not statistically significant.
Incubation of the splenocytes from exposed rats for 7 d in culture produced results similar to those obtained with cells incubated for 4 d. A highly significant ($P < 0.001$) reduction in *P. carinii* cyst counts was achieved with spleen cells pulsed with MSG.

To investigate the properties of these donor cells before adoptive transfer, aliquots of the whole spleen cell suspensions from both naive and exposed animals were examined for their in vitro proliferative response to MSG and their cellular composition. Peak proliferative response to MSG by *P. carinii*-exposed T cells occurs after a 4-d incubation with the antigen (14). As shown in Table I, only spleen cells from exposed animals displayed a significant proliferative response ($39,600\pm 6,200$ mean cpm, $P < 0.05$) to MSG compared to the values obtained from media alone. All four of the experimental groups displayed normal frequencies of CD4$^+$ and CD8$^+$ populations after 1 d of incubation; yet, by day four of incubation, these groups had elevated levels of CD4$^+$ cells. This change was most marked with exposed CD4$^+$ cells pulsed with MSG which increased from 46.2$\%$ at 1 d to 72.3$\%$ at 4 d. In contrast to the CD4$^+$ cells, there was no difference in the frequency of CD8$^+$ cells in any of the groups despite time of incubation.

**Adoptive transfer of lymphocyte subsets.** Based on the previous observations, to determine if a specific T cell subset provided the reduction of *P. carinii* burden observed, the donor cells were separated into CD4$^+$ and CD8$^+$ populations after a 4-d incubation in the presence or absence of MSG. As shown in Fig. 3, rats administered whole spleen cells after the 4-d incubation had cyst counts similar to those previously observed. There was no significant difference in the log$_{10}$ mean cyst counts among rats administered saline (9.2$\pm 0.13$/lung) and those recipients administered naive donor splenocytes pulsed for 4 d with MSG (8.76$\pm 0.4$/lung). Animals administered exposed splenocytes had significantly ($P < 0.05$) lower mean cyst counts (8.07$\pm 0.52$/lung), while the most significant ($P < 0.001$) reduction was observed in the recipients administered exposed splenocytes pulsed with MSG (7.17$\pm 0.36$/lung). In contrast to the whole spleen cells, a significant reduction in the log$_{10}$ mean cyst count was observed in all recipient animals receiving CD4$^+$ donor cells. These groups included unexposed CD4$^+$ cells pulsed for 4 d with MSG (8.2$\pm 0.63$/lung, $P < 0.05$), exposed CD4$^+$ cells (6.99$\pm 0.58$/lung, $P < 0.001$), and exposed CD4$^+$ cells pulsed for 4 d with MSG (5.93$\pm 0.77$/lung, $P < 0.001$). These data suggest that CD4$^+$ cells, regardless of origin or sensitization, are important in protection. However, an ~10-fold difference existed between each of the treatment groups, with MSG-pulsed CD4$^+$ cells isolated from exposed animals being the most effective in reducing the log$_{10}$ mean of *P. carinii* cysts. After reconstitution with CD8$^+$ cells, a significant ($P < 0.05$) reduction in the mean cyst count was observed only in the recipient rats administered exposed CD8$^+$ cells (8.29$\pm 0.27$/lung). No further reduction in the log$_{10}$ mean cyst count by the CD8$^+$ cells was obtained after incubation with MSG (8.32$\pm 0.12$/lung).

**Histological analysis.** Evaluation of the magnitude of *P. carinii* pneumonia and the effects of the adoptive transfer of the different donor cell groups by histological scoring correlated well with the data obtained by organism quantitation (Table II). The hematoxylin and cosin–stained sections of the lungs from animals receiving saline or naive donor cells revealed the typical foamy intraalveolar exudate with only a mild infiltration.
of alveolar macrophages. Examination of alveoli by silver staining revealed that a high percentage of the alveoli were involved in the infection (3.4±0.6) with cysts closely aggregated and clustered in the center of the exudate in these groups. The lungs from animals receiving exposed donor cells pulsed for 4 and 7 d exhibited significantly (P < 0.05) lower histological scores of 2.2±0.6 and 2.4±0.4, respectively. The results of these cell transfers were also seen in changes in the lung weight to body weight ratio, which has previously been shown to correlate well with histologic score and *P. carinii* organism burden (16).

**Mortality and the hyperinflammatory response with CD4+ cells.** Earlier reports have shown that reconstitution of SCID mice with CD4+ cells from a *P. carinii*-immunized or environmentally exposed donor results in an early hyperinflammatory reaction which leads to an increase in mortality (5). This extreme hyperinflammatory response was characterized by an increase in lung weight. In the present study, adoptive transfer of exposed CD4+ cells pulsed with MSG resulted in the lowest histologic score (1.6±0.5), but the highest lung weight to body weight ratio (18±2) of all rat groups (Table II). The results suggested that the CD4+ cells were effective in clearing *P. carinii* but caused a hyperinflammatory response which was responsible for the premature death of five rats. This conclusion was supported by histologic analysis which showed heavy infil-

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**Table I. Properties of Lewis Rat Donor Cells before Adoptive Transfer**

<table>
<thead>
<tr>
<th></th>
<th>[H]thymidine uptake</th>
<th>Positive CD4⁺</th>
<th>Positive CD8⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>cpm × 10⁻³</td>
<td>%</td>
<td>%</td>
</tr>
<tr>
<td>Experiment groups</td>
<td></td>
<td>1 d</td>
<td>4 d</td>
</tr>
<tr>
<td>Naive</td>
<td>1.5±0.3</td>
<td>39±2</td>
<td>48±5</td>
</tr>
<tr>
<td>Exposed</td>
<td>1.6±0.2</td>
<td>41±3</td>
<td>54±4</td>
</tr>
<tr>
<td>Naive</td>
<td>2.5±0.7</td>
<td>43±1</td>
<td>62±5</td>
</tr>
<tr>
<td>Exposed</td>
<td>7.9±1.7</td>
<td>46±4</td>
<td>72±3</td>
</tr>
</tbody>
</table>

* Whole spleen cell suspensions from naive and *P. carinii*-exposed Lewis rats were incubated in RPMI 1640 complete with and without MSG (10 μg/ml) for 1 or 4 d. After this incubation, the cells were pulsed overnight with 1 μCi/well [H]thymidine before harvest on glass fiber filter strips. Samples were then counted by liquid scintillation counting. Data are expressed as mean cpm±SD of triplicate cultures of a representative experiment.

1 Whole spleen cell suspensions from naive or *P. carinii*-exposed Lewis rats were incubated in RPMI 1640 complete with and without MSG (10 μg/ml) MSG for 1 or 4 d. Surviving cells were isolated over Ficoll-Hypaque and stained with Fluorescein-conjugated mouse anti- rat CD8 and phycoerythrin-conjugated mouse anti-rat CD4. Cells were analyzed by flow microfluorometry in a fluorescence-activated cell sorter.

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**Figure 2.** The effects of *P. carinii*-exposed or nonexposed donor cell incubation time with MSG on the clearance of pneumocystosis in recipient Lewis rats. Four groups of immunosuppressed Lewis animals were reconstituted at week seven with 10⁶ donor cells from either *P. carinii*-exposed or nonexposed Lewis rats. One group received nonexposed donor cells cultured for 1, 4, or 7 d; one group received exposed donor cells cultured for 1, 4, or 7 d; one group received nonexposed donor cells pulsed for 1, 4, or 7 d with MSG; one group received exposed donor cells pulsed for 1, 4, or 7 d with MSG; and one group received only saline. Raw numbers for quantitation of *P. carinii* cysts were obtained by staining lung homogenates with cresyl echt violet. Counts are reported as log transformation of the geometric mean±SD. *P < 0.05 compared to saline injection control. **P < 0.001 compared to saline injection control.
trates with mononuclear cells and alveolar macrophages within the lungs. By contrast, the three rats which survived this episode had a lung weight to body weight ratio of 9±2 and the lungs, which were cleared of *P. carinii*, had normal histology.

**Discussion**

Results described in this paper represent the first demonstration that MSG contains protective T cell epitopes. This protective response observed after adoptive transfer of the donor cells was dependent on previous exposure to *P. carinii* and time of incubation with MSG. The effect was specific for MSG, as demonstrated by the lack of clearance in the lungs of animals receiving donor cells incubated with an equivalent concentration of ovalbumin.

The role of MSG in recognition of and immune response to *P. carinii* has yet to be fully elucidated. Gigliotti et al. (18), using immunoelectron microscopy, demonstrated that MSG is distributed on the surface of all developmental stages of *P. carinii*. Through molecular studies of rat and ferret *P. carinii*, it has been demonstrated that MSG represents a group of proteins encoded by multiple genes (6–8, 19). Numerous unique MSG cDNAs have been identified, establishing that multiple mRNAs are transcribed within a population of organisms (6–9). These data raise the possibility that antigenic variation may occur in MSG as a means of evading the host immune defenses. The present report has shown that native MSG can elicit a protective immune response. Further studies with different recombinant MSG preparations are needed to identify conserved, protective epitopes before consideration can be given to MSG as a potential vaccine candidate.

Protection against *P. carinii* in the present study was achieved with the adoptive transfer of sensitized whole spleen cell suspensions and purified CD4+ cells, regardless of their origin or previous sensitization. Several methods have been utilized in vivo to demonstrate the critical role CD4+ cells have in resistance to *P. carinii* pneumonia. Shellito et al., (20) utilized CD4+ -depleted mice which, after intratracheal inoculation, developed a vigorous *P. carinii* infection that was cleared after repletion of these cells. Using the SCID mouse model, Hammsen and Stankiewicz (4) demonstrated that reconstitution of these mice with immunocompetent spleen cells resulted in clearance of *P. carinii*. In the same study, treatment of the reconstituted animals with an anti-CD4+ monoclonal antibody resulted in the inability to resolve the infection. Similarly, Roth and Sidman (5), also using the SCID model, showed that adoptive transfer of CD4+ cells was sufficient for resistance; yet, this reconstitution caused an intense hyperinflammatory response, resulting in high mortality in the recipient mice.

The present study was performed using the standard corticosteroid-treated rat model of pneumocystosis. We were reluctant to attempt the adoptive transfer experiments using this model because of the broad effects of steroids on the host immune system. These actions include inhibition of all lymphoid tissues (21); suppression of T lymphocyte function (22), macrophage differentiation and function (23), natural killer activity (24), and antibody responses to *P. carinii* (25); and peripheral blood lymphopenia, which occurs through a process of cell redistribu-

![Figure 3. The effects of purified lymphocyte subsets on the clearance of pneumocystosis in recipient Lewis rats. Immunosuppressed Lewis rats were reconstituted at week seven with donor cells from *P. carinii-*exposed or nonexposed Lewis rats. One group received 10⁶ naive donor cells pulsed for 4 d with MSG; one group received 10⁶ exposed donor cells cultured for 4 d; one group received 10⁶ exposed donor cells pulsed for 4 d with MSG; two groups received either 10⁶ purified nonexposed CD8+ or CD4+ cells pulsed for 4 d with MSG; two groups received either 10⁶ purified exposed CD8+ or CD4+ cultured for 4 d; two groups received either 10⁶ purified exposed CD8+ or CD4+ cells pulsed for 4 d with MSG; and one group received only saline. Raw numbers for quantitation of *P. carinii* cysts were obtained by staining lung homogenates with cresyl echt violet. Cyst counts are reported as log transformation of the geometric mean±SD. *P < 0.05 compared to saline injection control. **P < 0.001 compared to saline injection control.](image-url)
Table II. Histological Score, Lung wt to Body Weight Ratio, and Percent Mortality of Lewis Rats after Adoptive Transfer of MSG-pulsed T Cells

<table>
<thead>
<tr>
<th>Experimental group*</th>
<th>No. of animals</th>
<th>Histological score</th>
<th>Lung wt to body wt</th>
<th>Premature death</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saline</td>
<td>14</td>
<td>3.4±0.6</td>
<td>17±3</td>
<td>4/14</td>
</tr>
<tr>
<td>1 d pulse</td>
<td>10</td>
<td>2.8±0.8</td>
<td>10±4**</td>
<td>0/10</td>
</tr>
<tr>
<td>4 d pulse</td>
<td>10</td>
<td>2.2±0.6**</td>
<td>9±3**</td>
<td>0/10</td>
</tr>
<tr>
<td>7 d pulse</td>
<td>9</td>
<td>2.4±0.4**</td>
<td>11±2**</td>
<td>1/9</td>
</tr>
<tr>
<td>CD8+</td>
<td>8</td>
<td>2.8±0.4</td>
<td>11±3**</td>
<td>2/8</td>
</tr>
<tr>
<td>CD4+</td>
<td>8</td>
<td>1.6±0.5**</td>
<td>18±2</td>
<td>5/8</td>
</tr>
</tbody>
</table>

* Five groups of immunosuppressed Lewis rats were reconstituted at week seven with donor spleen cells from P. carinii–exposed Lewis rats. Groups received 10^6 cells pulsed for 1, 4, or 7 d with MSG. Two groups received either 10^6 purified CD8+ or CD4+ cells pulsed for 4 d with MSG. One group received only saline. 4 Only animals which survived 10 d after adoptive transfer were included in the histologic and quantitative analysis for severity of P. carinii infection. 6 Histological score was determined by analysis of hematoxylin and cosin and Grocott Gomori methenamine silver-stained sections of the lungs. Scoring for alveoli involvement: 0, no infection; 1, light (1–25% alveoli); 2, moderate (25–50% alveoli); 3, severe (51–75% alveoli); and 4, very severe (76–100% alveoli). Values represent mean±SD of all lungs scored in the experiments. ** Values represent mean ratio × 10^3±SD. 1 Numbers represent animals that died between 10 d after adoptive transfer and week 10 of the experiment. ** Significantly different compared to saline, P < 0.05.

tion rather than cell destruction with the primary site of influx being the bone marrow (15, 26, 27). Since at least a portion of the lymphocytes in the lung are derived from peripheral blood (28), we feared that corticosteroids might impede recruitment of lymphocytes from the bone marrow to the lung. However, studies demonstrating the enhanced ability of T cell clones to survive and function in immunocompromised hosts suggested that steroid-treated rats might provide a permissive environment for adoptive transfer experiments (29, 30). Perhaps the intravenous injection of activated lymphocytes in our rats resulted in immediate recruitment of the donor cells to the lung, where they were sequestered at the site of infection.

The treatment of P. carinii pneumonia with drugs or by immunological reconstitution has led to some unexpected results. In both our study and a previous study using the SCID mouse model (5), reconstitution with CD4+ cells alone produced an extreme hyperinflammatory response resulting in deleterious effects to the host. This response occurred in our model despite the presence of steroids. In contrast, administration of corticosteroids to AIDS patients with pneumocystis prevwnt the deterioration in blood oxygenation which occurs soon after beginning anti-P. carinii therapy; the hypothesis here is that the drug-induced damage to P. carinii exacerbates the lung’s inflammatory response (31–34). It, thus, appears that the pathophysiologic mechanisms underlying the clinical deterioration in AIDS patients and rodents are different. P. carinii pneumonia causes significant changes in the lung surfactant system characterized by a fall in phospholipids and an increase in surfactant protein A (35). Since corticosteroids are known to increase surfactant phospholipids and cause other changes in the alveolar microenvironment (36), it is possible that the beneficial effects of steroids in AIDS patients with P. carinii pneumonia are exerted through the surfactant system rather than through the host inflammatory response.

Although the data presented here indicate that CD4+ cells play a critical role in the immune response to P. carinii, the possible role for CD8+ cells has not been elucidated. CD8+ cells offer some resistance to pneumocystosis when adoptive transfer (Fig. 3). In contrast to our findings, adoptive transfer of purified CD8+ cells into SCID mice was ineffective in clearance (5). Previous studies using the corticosteroid-treated rat and the CD4+–depleted mouse demonstrated a large influx of CD8+ cells into the lungs of P. carinii–infected animals (15, 37). Since CD8+ cells are recruited to the lung in response to P. carinii infection, perhaps this subset plays at least a partial role in clearance.

In conclusion, this study has shown that MSG contains protective T cell epitopes which can be demonstrated by adoptive transfer experiments in the steroid-treated rat model of pneumocystosis. With the recent progress in cloning and expressing MSG genes (6–9), it should be possible to develop T cell clones, map their epitopes, and test their functional importance in vivo.

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References


