Pneumocystis carinii: Inhibition of Lung Cell Growth Mediated by Parasite Attachment

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Abstract

Pneumocystis carinii pneumonia is a significant cause of mortality in immunocompromised patients. Current concepts suggest that attachment of P. carinii to alveolar epithelium is required for development of pneumonia. We examined the mechanism of P. carinii adherence to cultured A549 cells, a permanent cell line derived from human alveolar epithelium. P. carinii adherence was quantified by measuring attachment of 51Cr-labeled P. carinii to cultured A549 cells. After 8 h of incubation, 37.4±4.2% of P. carinii were adherent to A549 cells. In the presence of agents known to impair cytoskeletal function, including 10-5 M cytochalasin B, 10-5 M colchicine, and 10-5 M trimethylolcholinic acid (TMCA), adherence was decreased from 57.4±4.2% to 9.3±3.4%, 12.5±3.6%, and 21.5±3.6%, respectively (P < 0.01, all comparisons). Secondly, we examined the effect of P. carinii on the function of A549 cells. P. carinii resulted in significant impairment of A549 cell growth, indicating P. carinii adversely affected the function of target lung cells. A P. carinii:A549 cell ratio of 50:1 resulted in 43.5±2.9% inhibition of A549 cell growth (P < 0.001). Additionally, TMCA, which significantly prevented attachment of P. carinii, reversed the impairment of A549 cell growth. These data demonstrate that P. carinii attachment to cultured lung cells can be quantified, is dependent on intact cytoskeletal function and is necessary for impairment of lung cell replication. (J. Clin. Invest. 1990. 85:391–396.) Pneumocystis carinii: pneumonia • parasite adherence • cytochalasin B • colchicine

Introduction

Pneumocystis carinii pneumonia is a frequent life-threatening complication of immune deficiency. Current concepts suggest growth of P. carinii organisms requires attachment of P. carinii trophozoites to the alveolar epithelial cells of the lung (1–7). P. carinii organisms appear to preferentially attach to type I alveolar epithelial cells, although small numbers may attach to the surface of type II alveolar epithelial cells (1, 4–7). There is no evidence to suggest that P. carinii enters the epithelial cell or has an intracellular phase in its life cycle; rather, the organisms remain on the surface of the cell. The mechanism of P. carinii attachment and how the lung cell is affected by this attachment is poorly understood.

Ultrastructural studies have provided important descriptive information regarding the attachment of P. carinii to alveolar epithelial cells (1–7). Although earlier reports suggested that P. carinii use specialized “filopodia” as organelles of attachment (8, 9), more recent studies suggest P. carinii attachment is not mediated by specialized structures (3–5); rather, attachment occurs as a result of close apposition (not fusion) of the cell membranes of P. carinii to the alveolar epithelial cell (3–6).

The study of P. carinii attachment to lung cells has been limited by the absence of an appropriate model that permits quantification of attachment and isolation of variables thought to be important in the attachment process. Our study includes the development of a novel method to quantify P. carinii attachment and suggests that P. carinii requires intact cytoskeletal function to adhere to target lung cells. Furthermore, P. carinii appears to adversely affect target cell function by impairing lung cell replication. This P. carinii-mediated impairment of lung cell growth requires attachment of the organism to the target cell surface since agents that prevent attachment also reverse this adverse effect on the target cell population.

Methods

P. carinii isolation and purification. P. carinii pneumonia was induced in Harlan Sprague-Dawley rats by immunosuppression with dexamethasone as previously described by Hughes and co-workers (10). Female Harlan Sprague Dawley rats (250 g; Harlan Sprague Dawley, Inc., Indianapolis, IN) were housed in open cages in a clean but not sterile environment. Drinking water containing a mixture of dexamethasone (2 mg/liter), tetracycline hydrochloride (500 mg/liter), and nystatin (200,000 U/liter) was provided ad lib. Rats were maintained on a diet containing 8% protein in order to intensify the severity of P. carinii disease (Teklad Corp., Madison, WI) (11).

After 8 wk of treatment, rats exhibited evidence of respiratory distress, wasting, and hair loss, and were killed by lethal injection with 26% pentobarbital intraperitoneally. After aseptically opening the thorax, rats were exsanguinated and the lungs perfused with HBSS, without calcium or magnesium, containing 0.6 mM EDTA, penicillin (100 μg/ml), gentamicin (4 μg/ml), and amphotericin B (0.5 μg/ml). Whole lung lavage was performed with sequential 10-ml aliquots of HBSS with immediate return of up to 50 ml of instilled fluid.

P. carinii organisms were isolated and purified by differential centrifugation adapted from the method described by Masur (12). Pooled lavage fluid (typically 45 ml) was centrifuged (800 g × 10 min) and P. carinii cysts were identified in the inflammatory cell pellets using a modified Grocott methenamine silver stain (13). The turbid supernatants containing predominantly P. carinii organisms were centrifuged (1,400 g × 30 min), the pellets resuspended in 1 ml of HBSS, and P. carinii organisms were quantified by the method of Bartlett et al. (14). Duplicate 10-μl aliquots of this suspension were directly spotted onto glass slides, air dried, and stained with Diff-Quik (Harleco, Dade Diagnostics, Aqua, Puerto Rico). Each drop occupied ~1 cm² on the slide, and the number of P. carinii trophozoites in 10 oil immersion fields was determined for each spot. The total number of P. carinii was calculated using the formula: P. carinii/ml = (P. carinii/field) 

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× (No. of fields/drop) × (10^7). Using the above method, average yield was 9.0 ± 0.4% × 10^6 purified P. carinii organisms from each rat lavaged. Further, P. carinii organisms represented 96 ± 5.5% of the cellular differential, with the remainder representing mostly nonivable inflammatory cells. Occasional rats had evidence of significant bacterial contamination noted on smears of the lavage, and these were discarded. Despite occasional evidence of other microorganisms on smears of the lavages, microbiologic evaluation of selected rat lavage isolates failed to culture any known pathogenic bacteria, fungi, or virus.

P. carinii adherence assays. Freshly isolated P. carinii (24 × 10^6 organisms) were incubated for 18 h in 1 ml of Dulbecco’s modified Eagle’s medium (Whittaker M. A. Bioproducts, Walkersville, MD) containing 20% heat-inactivated fetal calf serum, penicillin (10 μg/ml), gentamicin (4 μg/ml), and amphotericin B (0.5 μg/ml) and 50 μCi 51Cr/ml (679 mCi/mg, New England Nuclear, Boston, MA). This suspension was washed four times with medium to remove unincorporated 51Cr and adjusted to a concentration of 2.0 × 10^7 P. carinii per ml. Two million 51Cr-labeled P. carinii were incubated with confluent monolayers of A549 cells (American Type Culture Collection, Rockville, MD) grown in 24-well plates using DME containing 10% heat-inactivated fetal calf serum and penicillin, gentamicin, and amphotericin as noted above. After centrifugation onto the monolayers (1,400 × g × 5 min), 51Cr-labeled P. carinii organisms were incubated with the A549 cell monolayers for 0, 2, 4, and 8 h. After incubation, media were removed and A549 cells washed three times with HBSS to remove any unattached 51Cr-labeled P. carinii organisms. Adherence of P. carinii was defined as follows: % adherence = (A/A + B) × 100%, where A = dpm of 51Cr associated with the A549 cell monolayer and B = dpm of 51Cr-labeled P. carinii not adherent to the cellular monolayer.

To provide morphologic confirmation of P. carinii adherence to cultured A549 cells, 2 × 10^6 P. carinii were incubated with confluent monolayers of A549 cells grown on sterile plastic coverslips (Thermanox Tissue Culture Slips; Miles Laboratories, Inc., Naperville, IL) as described above. After incubating, the coverslips were removed, washed with HBSS and fixed in Trichol’s fixative containing 1% glutaraldehyde and 4% formaldehyde in 0.1 M NaPO4 buffer at pH 7.20 for 2 h. Scanning electron microscopy (SEM) was performed using an ETEC Autoscan microscope (Perkin-Elmer, Norwalk, CT).

To assess the importance of parasite cytoskeletal function on adherence, additional assays were conducted in the presence and absence of 10^−4 to 10^−5 M concentrations of cytochalasin B (Sigma Chemical Co., St. Louis, MO), colchicine (Eli Lilly Co., Indianapolis, IN), or trimethylcholinic acid (TMCA) (Sigma Chemical Co.) for 8 h, and the % adherence of 51Cr-labeled P. carinii was determined as noted above.

To verify that cytochalasin B, colchicine, and TMCA did not adversely affect P. carinii viability, 51Cr release (measure of cytoxicity) and [3H]leucine incorporation (measure of protein synthesis) were measured. The percent release of 51Cr (%R) from labeled P. carinii during 8 h of incubation in the presence and absence of 10^−3 M cytochalasin B, colchicine, or TMCA was determined as follows: %R = (A/A + B) × 100%, where A = dpm of 51Cr released into the medium and B = dpm of 51Cr associated with P. carinii. To determine the effect of these agents on protein synthesis, 2 × 10^6 P. carinii organisms were incubated with [3H]leucine (75 μCi/ml) (140 Ci/mmol, New England Nuclear) for 8 h in the presence and absence of 10^−3 M cytochalasin B, colchicine, or TMCA. After incubation, the P. carinii were washed three times with medium and sonicated. The sonicate was precipitated in ice-cold 10% trichloroacetic acid and the precipitable protein measured by scintillation counting.

A549 cell growth assay. The effect of P. carinii on the function of the A549 cells was measured by monitoring A549 cell replication. A549 lung cells were cultured in DME containing 10% heat-inactivated fetal calf serum, penicillin, gentamicin, and amphotericin B as noted previously. Confluent monolayers of A549 cells were grown on 25 cm² tissue culture flasks (Corning Glass Works, Parkridge, IL), removed by brief treatment with Trypsin Versene (1×) (Biofluids, Rockville, MD), and replated in fresh medium at a density of 30,000 cells per well in 6-well tissue culture plates (Nunc, Inc., Naperville, IL). After attachment of A549 cells to tissue culture plates (4 h), medium was replaced with fresh medium containing appropriate numbers of P. carinii organisms. Wells containing A549 cells without P. carinii organisms served as controls. After 72 h, the medium was removed and monolayers were washed thoroughly with three aliquots of HBSS. The A549 target cells were completely removed with Trypsin Versene (1×) and were enumerated by Coulter counter analysis (model ZF; Coulter Electronics, Inc., Hialeah, FL).

Impairment of target cell growth was expressed as a growth impairment (GI) index, as previously described (15), where 100% represents total inhibition of growth and 0% represents normal growth of A549 cells. The following formula was employed: GI Index = [(A - B)/A] × 100%, where A = A549 cell count in control wells, and B = A549 cell count in wells containing P. carinii.

Assessment of culture medium. To determine whether P. carinii consume excess media nutrients or release soluble growth inhibiting substances into the culture media, media incubated with A549 cells with or without P. carinii was assayed to determine its effect on further A549 cell growth. Media removed from A549 cell growth assays were filter sterilized using 0.45 μm disk filters (Acrodisc; Gelman Sciences, Ann Arbor, MI) to remove P. carinii organisms, added to 6-well culture plates containing newly seeded A549 cells (30,000 cell/well) and incubated for 72 h. A549 cells were then removed from the culture dishes and the GI indices determined as noted above.

Statistical methods. Data are expressed as means±SEM. Differences between experimental and control data groups were assessed using Student’s two-sample t test for two-sided alternatives in which P < 0.05 was defined as a statistically significant result.

Results

P. carinii adherence assays. P. carinii adherence to monolayer A549 cells was quantified using 51Cr-labeled P. carinii. Incubation of 51Cr-labeled P. carinii with A549 target cells for 2, 4, and 8 h (Fig. 1) resulted in 13.0±1.4%, 40.6±4.6%, and 37.4±4.2% adherence, respectively, (P < 0.001, latter two comparisons to control at 0 h). Ultrastructural examination by SEM revealed P. carinii organisms to be adherent to the A549 cell monolayer with P. carinii typically attaching in clusters to one in every four or five monolayer cells (Fig. 2). When 51Cr counts were detectable, adherent P. carinii were present; similarly, when no 51Cr counts were detectable, no P. carinii organisms were visible by SEM. Thus, adherence of P. carinii to cultured A549 cells could be quantified using 51Cr-labeled organisms and occurred in a time-dependent fashion and appeared to reach a maximum between 4 and 8 h.

Figure 1. Time course of P. carinii adherence to A549 cells. The 51Cr-labeled P. carinii were incubated with A549 cell monolayers for 2, 4, and 8 h. Data are expressed as percent adherence of P. carinii to A549 cell monolayers. *Denotes comparisons which were significantly different from controls (P < 0.05). Each data point represents six determinations and are expressed as mean±SEM.
Role of cytoskeletal components in adhesion. To determine the need for intact cytoskeleton function in the adherence of *P. carinii* to target cells, assays were conducted in the presence and absence of the antimotility agents cytochalasin B, colchicine, or the colchicine derivative, TMCA (Fig. 3). The adherence of *P. carinii* to the A549 cell layer was reduced from a control value of 37.4±4.2% to 9.3±3.4% (10⁻⁵ M cytochalasin B), 12.5±3.6% (10⁻³ M colchicine), and 21.5±3.6% (10⁻³ M TMCA) (*P* < 0.01, all comparisons). Further adherence assays were conducted using *P. carinii* or A549 cells with only a 1-h
preincubation of the cells with the antimotility agents at the same concentrations, followed by washing and incubating in the absence of the agents. Preincubation of the antimotility agents with either cell population failed to change adherence (data not shown), consistent with other studies which indicate the effects of these antimotility agents are "reversible" (16–18). Thus, agents that impair cytoskeletal function significantly reduced the adherence of the P. carinii organisms to A549 lung cells when present during the attachment process.

Additionally, the highest concentration of cytochalasin B, colchicine, and TMCA used (10–5 M), did not result in significantly greater release of 31Cr from treated P. carinii compared with control organisms (control = 29.1±4.3%; cytochalasin B = 32.5±5.0%; colchicine = 31.5±3.5%; TMCA = 30.0±3.5%; P > 0.10 all comparisons to control). Further, the incorporation of [3H]leucine into trichloroacetic acid precipitable protein by P. carinii organisms was not significantly altered in the presence of the drugs (control = 2,320±313 dpm; cytochalasin B = 2,091±381 dpm; colchicine = 2,627±642 dpm; TMCA 2,174±150; P > 0.10 all comparisons to control). Thus, agents that impair cytoskeletal function of P. carinii organisms and impair adherence of the organism to A549 cells do not interfere with the apparent viability or functional status of P. carinii organisms.

A549 target cell growth assay. Increasing concentrations of P. carinii organisms resulted in significant impairment of A549 cell growth as expressed by a dramatic increase in the GI index (Fig. 4). A P. carinii:A549 cell ratio of 1:10 resulted in a GI index of only 4.4±3.4% growth inhibition; whereas, at a ratio of 25:1, the GI index increased to 34.9±6.5% (P < 0.005); and at 50:1, increased to 43.5±2.9% (P < 0.001). No cytopathic effects were detectable in the target A549 cell population. Thus, the higher the number of P. carinii organisms the greater the impairment of A549 cell growth, indicating for the first time that P. carinii may seriously affect the normal function of target lung cell populations.

Additionally, heat inactivation of P. carinii substantially reduced the growth-inhibiting properties of P. carinii on A549 cells. After heat inactivation (56°C × 1 h), heat killed P. carinii resulted in a growth inhibition of only 8.2±3.5% compared to 33.2±15.5% growth inhibition from control P. carinii (P < 0.05), indicating that viable P. carinii are necessary for significant growth inhibition.

Effect of P. carinii treated culture media on A549 cell growth. Culture media incubated with A549 cells in the presence and absence of P. carinii was filter-sterilized to remove P. carinii organisms, and assayed to determine if P. carinii conditioned medium would impair A549 cell growth. Filter-sterilized media from incubations with A549 cells alone (controls) resulted in a GI index of 8.1±6.8%; similarly, filter sterilized media from incubations with A549 cells and P. carinii organisms resulted in a GI index of 6.4±4.9% (P = 0.8). Thus, there was no evidence of P. carinii-conditioned media adversely affecting A549 cell growth compared to control media. This suggests that P. carinii organisms do not impair A549 cell replication by consuming excess essential nutrients or releasing growth inhibiting toxins into the medium.

Role of adherence in growth impairment. To determine whether adherence of P. carinii is necessary for P. carinii to impair growth of A549 lung cells, the A549 cell growth assay was repeated with P. carinii in the presence and in the absence of TMCA (Fig. 5). TMCA was used because, unlike cytochalasin B and colchicine, it does not inhibit the growth of cultured cells (19). TMCA (10–5 M) reversed the effect of P. carinii on A549 cell growth with a reduction in the GI index from 56.2±2.9% (PC alone) to a GI index of 10.2±3.1% (PC with concentrations of TMCA during a 72-h incubation. Data are expressed as a growth impairment index (GI index). *Denotes comparisons that were significantly different from controls (no TMCA) (P < 0.05). Each data point represents six determinations and is expressed as mean±SEM.
that the parasites do not consume excess essential nutrients nor release stable growth impairing products into the culture medium. Further study of this P. carinii-mediated toxicity may provide important insights into the host-parasite interaction and improve our understanding of mechanisms operative in the disease process.

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References


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