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Evidence for a Protective Role of Pulmonary Surfactant Protein D (SP-D) against Influenza A Viruses

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Abstract
We tested the hypothesis that pulmonary surfactant-associated lectins—surfactant proteins A and D (SP-A, and -D)—contribute to initial protective mechanisms against influenza A viruses (IAVs). SP-D potently inhibited hemagglutination activity of several strains of IAV as well as causing viral aggregation. SP-D enhanced neutrophil binding of IAV and neutrophil respiratory burst responses to the virus. Neutrophil dysfunction resulting from IAV exposure was diminished when the virus was pre-incubated with SP-D. Each of these effects was mediated by the calcium-dependent carbohydrate-binding property of SP-D. Native SP-D preparations of both human and rat origin, as well as recombinant rat SP-D, had similar activity. SP-A also inhibited IAV hemagglutination activity. We have previously reported that related mammalian serum lectins (mannose-binding lectin [MBL] and conglutinin) have similar effects. SP-D was at least 10-fold more potent at causing hemagglutination inhibition than were SP-A or MBL. SP-D was shown to contribute to potent anti-IAV activity of human bronchoalveolar lavage fluid. These results suggest that SP-D—alone, and in conjunction with SP-A and phagocytic cells—constitutes an important component of the natural immune response to IAV infection within the respiratory tract. (J. Clin. Invest. 1994. 94:311–319.) Key words: influenza virus • neutrophil • surfactant protein D • surfactant protein A • hemagglutination

Introduction
Surfactant protein D (SP-D)1 is a member of the collectin subgroup of the mammalian C-type lectin superfamily, which includes another surfactant protein, surfactant protein A (SP-A), and at least two serum proteins, mannos-binding lectin (MBL), and conglutinin (1, 2). Each of these lectins has been shown to bind to a variety of microorganisms, and may have a role in host defense responses, both through inhibiting infectivity of various organisms directly, and by enhancing phagocyte activation (3–9). The evidence for this is strongest with MBL, in that low levels of this protein have been associated with a syndrome of increased susceptibility to infection during infancy (2). We have previously reported that MBL and conglutinin inhibit the infectivity and hemagglutination (HA) activity of influenza A viruses (IAV) by binding to carbohydrates present on the viral HA molecule (10, 11). These lectins also act as opsonins, promoting neutrophil activation in response to IAV. The strong structural relationship of these lectins with SP-D (e.g., 66% amino acid sequence identity of human SP-D with conglutinin) (12, 13), and the fact that SP-D accumulates in the lower respiratory tract at sites suitable to interact with IAV in vivo (14), prompted us to examine the activity of SP-D against IAV in vitro.

The collectins naturally occur as multimeric structures resembling C1q in having collagenous domains which tether multiple globular carbohydrate recognition domains (2, 15). SP-D is produced by alveolar type II pneumocytes and by Clara cells (nonciliated bronchiolar cells) (14). It has been shown to be a major E. coli binding protein in bronchoalveolar lavage fluid, and can bind to and aggregate a range of bacteria via its calcium-dependent carbohydrate binding activity (8). Rat and human SP-D have very similar sugar binding specificity and have similar interactions with bacteria (8, 12). In addition, SP-D has been shown to bind to alveolar macrophages (16) and to stimulate a chemiluminescence response in these cells (9).

We here report that SP-D of human and rat origin share with MBL and conglutinin the ability to potently inhibit HA activity of a variety of strains of IAV, and to enhance neutrophil activation by IAV.

Methods
Reagents. Formylmethionylleucylphenylalanine (FMLP), cytochalasin B, horseradish peroxidase-Type II, scoopoletin, superoxide dismutase (SOD), cytochrome-C, maltose, Ficoll, dextran, sodium citrate, maltosyl-agarose, and citric acid were purchased from Sigma Chemical Co. (St. Louis, MO) and Hypaque was obtained from Winthrop Pharmaceuticals (Des Plaines, IL). Dulbecco’s phosphate-buffered saline with or without Ca2+ or Mg2+ was purchased from Flow Labs (McLean, VA). Tris buffered saline contained either 2 mM Ca2+ or 10 mM EDTA, as indicated.

SP-D preparation. Native rat SP-D was isolated from the 10,000 g supernatant of bronchoalveolar lavage from silicotic rats (17). Human SP-D was isolated from the supernatant of lavage obtained from patients with human alveolar proteinosis (18). In each case, SP-D was purified by sequential saccharide affinity chromatography on maltosyl-agarose, and gel-filtration on 4% agarose (A15M; Bio Rad Laboratories, Richmond, CA). Rat SP-D showed a single disulfide-bonded component of 43 kD (reduced) after resolution by SDS-PAGE and silver staining. The human SP-D preparation showed a major band of 43 kD (reduced), and minor components corresponding to SP-D dimers, trimers and...
The ELISA assay to measure SP-D levels employed native human SP-D and the immunoglobulin fraction of a well-characterized rabbit anti–SP-D antiserum (8) to establish a standard curve. The assay detected concentrations between 5 and 250 ng/ml of human SP-D in a linear manner. Purified SP-D or BAL fluids were coated overnight on microtiter wells, then washed three times with PBS-Tween followed by blocking of nonspecific binding sites with 0.25% BSA in PBS-Tween. After further washing, the wells were incubated for 2 h with a 1:1,000 dilution of the anti–SP-D immunoglobulin preparation. After washing, a 1:1,000 dilution of goat anti–rabbit polyclonal IgG conjugated to horseradish peroxidase (HRP; Sigma Chemical Co.) was then added for 2 h, followed by washing and development using TMB peroxidase substrate (Bio Rad Laboratories). The reaction was stopped using \( \text{H}_2\text{SO}_4 \), and absorbance read at 450 nm. All samples were run in duplicate. Protein assays were performed using BCA protein assay according to manufacturer’s specifications (Sigma Chemical Co.).

Figure 1. SDS/PAGE of purified native human SP-D and SP-A. Human alveolar proteinosis SP-A and SP-D were reduced with 5% mercaptoethanol, followed by loading of 1 and 10 \( \mu \text{g} \), respectively, of the reduced SP-D and SP-A onto a 10% polyacrylamide gel. The gel was developed with Coomassie blue.

higher aggregates. The procedures for purification of these proteins have been previously published in greater detail (17, 18). As established in those publications these SP-D preparations were free of contamination with other proteins including SP-A based on SDS-PAGE and immunoblotting analysis.

Recombinant rat SP-D was derived from a full length clone of rat SP-D cDNA stably expressed in CHO K1 cells. The protein comigrated with native rat SP-D on SDS-PAGE under reducing and nonreducing conditions, showed the expected shift in apparent molecular mass after incubation with endoglycosidase F, bound efficiently to maltosyl-agarose, and eluted at the expected position on nondenaturing gel filtration chromatography. A more detailed description of the expression and purification of this protein is in preparation.

Human surfactant protein A (SP-A) was graciously provided by Drs. Virginia Shepherd and Zissis Chrones (Vanderbilt University, Nashville, TN). The SP-A was purified from alveolar proteinosis patients as described (19) and was shown to be at least 95% pure by SDS-PAGE analysis and by \( ^{35} \text{S} \) labeling followed by SDS-PAGE and autoradiography. Fig. 1 depicts the appearance of the SP-A used in these studies on SDS-PAGE. An additional lane showing the migration of native human SP-D is included for comparison. Note that the SP-A appears as a broad band of \( \sim 26-36 \text{ kD} \) as well as another band of \( \sim 62 \text{ kD} \). This is the typical appearance of purified SP-A derived from alveolar proteinosis patients when analyzed by SDS-PAGE under reducing conditions (19). Previous studies have demonstrated with amino acid sequencing and other methods that these bands represent monomeric and dimeric forms of SP-A (20).

Bronchoalveolar lavage (BAL) fluids were obtained from healthy volunteer donors with informed consent as approved by the Boston University School of Medicine Institutional Review Board for Human Research. Between 150 and 200 ml of normal saline was instilled for the lavage. Fluids retrieved from this procedure were subjected to an initial low speed centrifugation (150 g) to remove cells and other large particulates. Further processing of the fluid was carried out as described below.

Virus preparation. Virus stocks were grown in the chorioallantoic fluid of 10-d-old embryonated hen’s eggs, and purified on a discontinuous sucrose density gradient as previously described (21, 22). Virus stocks were suspended in Dulbecco’s modified phosphate-buffered saline (PBS), aliquoted and stored at \( \sim 70^\circ \text{C} \) until used. Potency of each virus stock was measured by hemagglutination assay, and titers of 1:8,000 through 1:32,000 (as indicated) hemagglutination units (HAU) were measured after samples were thawed from storage at \( \sim 70^\circ \text{C} \). Hemagglutinin (HA) titer were determined by titration of virus samples in PBS followed by addition of thoroughly washed human type O red blood cells. The A/Texas 77/H3N2 (Texas 77) and A/PR/8/34/H1N1 (PR8) strains of IAV were gracious gifts of Dr. Jon Abramson (Bowman-Gray School of Medicine, Winston-Salem, NC). The A/Brazil78/H1N1 (Brazil 78) and A/Mem71r-Bel11 strains were kindly provided by Dr. E. Margot Abramson (University of Melbourne, Melbourne, Australia) (23). The A/Bangkok 79/H3N2 (Bangkok 79) strain was a gift of Robert Webster, M.D. (St. Jude’s Hospital, Memphis, TN).

Incubation of SP-Ds with IAV stocks was carried out at 37°C in PBS with 2 mM Ca\(^{2+}\) except in indicated experiments in which maltose was added, or in which the buffer contained 10 mM EDTA and no added Ca\(^{2+}\). Where maltose was used to inhibit the carbohydrate binding activity of SP-D (or SP-A) the lectins were pre-incubated with 167 mM maltose for 15 min before adding the lectin to IAV. Preliminary experiments (data not shown) demonstrated that this concentration of maltose maximally reduced the HA inhibitory activity of SP-D without altering HA activity of the virus in the absence of SP-D or affecting neutrophil functions. In the former case, SP-D stocks were maintained in 10 mM EDTA and recalcified just prior to assays to minimize SP-D self-aggregation. Prior to incubation of the lectin with the virus, a low speed (150 g for 5 min) centrifugation of the virus preparation was carried out to remove any viral aggregates.
Assessment of virus aggregation. Virus aggregation was assessed by measuring changes in light transmission through suspensions of IAV after addition of various concentrations of SP-D. These measurements were performed on an SLM/Aminco 8000C spectrophotometer. The excitation and emission wavelengths were 350 nm and slits at 16, 0.25, 0.25, 2, and 2 nm. Previous experiments have established that a decrease in light transmission of vesicles in the size range of IAV particles indicates particle aggregation (24, 25).

Addition of SP-D to virus-free buffer containing 2 mM Ca⁺⁺ did not cause any decrease in light transmission.

Measurement of neutrophil activation. Neutrophils from healthy volunteers were isolated to >95% purity as previously described using dextran sedimentation, followed by a Ficoll-Hypaque gradient centrifugation for removal of mononuclear cells, and hypotonic lysis to eliminate contaminating erythrocytes (21). Cell viability was >98% as determined by trypan blue staining, and cells were used within five hours of isolation. H₂O₂ production was measured by the oxidation of scopoletin, and O₂⁻ assessed by the continuous monitoring of the SOD inhibitable reduction of cytochrome C as previously detailed (26). Neutrophil deactivation was assessed by first incubating cells with IAV for 10 min., followed by measurement of O₂⁻ production in response to FMLP (10, 21).

Measurement of influenza virus binding to neutrophils. Viral binding to neutrophils was measured by preparing FITC-labeled virus and incubating this preparation with neutrophils, followed by evaluation of cell-associated fluorescence using a flow cytometer. FITC stock was prepared at 1 mg/ml in 1 M sodium carbonate, pH 9.6. The FITC labeled virus was prepared by incubating concentrated virus stocks with FITC (10:1 mixture by volume of virus in PBS with FITC stock) for 1 h, followed by dialysis of the mixture for 18 hr against PBS. For fluorescence measurements, concentrated FITC-labeled virus was incubated for 30 min at 37°C with control buffer or various amounts of SP-D, followed by addition of 10-μl aliquots of these samples to neutrophils (10⁶ cells in 100 μl TBS containing either 2 mM Ca⁺⁺ or 10 mM EDTA, as indicated). In some samples SP-D was pre-incubated with maltose (167 mM) before mixing SP-D with virus. After allowing virus and neutrophils to interact for 15 min at 4°C, the neutrophils were washed, resuspended in virus-free PBS and fixed with 2% paraformaldehyde. Cell associated fluorescence was measured on a Becton-Dickinson FACS Scan 2 and analyzed using the Lysis II program.

Results

Effect of incubation with SP-Ds on hemagglutination activity of IAV. As presented in Table I, native rat and human SP-D inhibited HA activity of all H3 strains of IAV tested in a Ca⁺⁺-dependent manner. Recombinant rat SP-D had similar HA inhibitory effects. The Brazil 78 strain of IAV which is of the H1 subtype was also inhibited by the SP-Ds. If SP-Ds were pre-incubated with maltose they lost the ability to inhibit IAV HA activity. In contrast, pre-incubation with ≥160 mM of either galactose or N-acetyl glucosamine did not reduce HA inhibitory effects of SP-D (data not shown). These results indicate that HA inhibition by SP-D was mediated by its Ca⁺⁺-dependent, lectin activity, and are compatible with the known carbohydrate-binding specificities of SP-D. SP-D had much less activity against the PR-8 H1N1 strain of IAV (see Table I). The PR-8 strain of IAV differs from H3 strains in that it lacks high mannose oligosaccharide attachments on its HA and has no carbohydrate attachment near the sialic acid binding site (27, 28).

Another approach was taken to characterize the HA inhibitory effects of the SP-Ds that involved pre-incubation of much more concentrated preparations of IAV with escalating doses of the SP-Ds, followed by measurement of HA activity. As depicted in Fig. 2, both native rat and human SP-D caused dose-related inhibition of HA activity (ID₅₀, respectively, 1.5 and 3 μg/ml). For recombinant rat SP-D the ID₅₀ for HA activity of a similar concentration of IAV was 3.1 μg/ml (mean of four experiments; data not shown). The HA inhibitory effects were dependent on calcium, and inhibitable by pre-incubation of the lectins with maltose. Aliquots of these concentrated virus preparations treated with various doses of lectins were then used in assays of neutrophil binding to IAV (see below).

Rat and human SP-D cause aggregation of IAV. As depicted in Fig. 3, when either native rat or human SP-D were added to a stirred suspension of Texas 77 IAV in the presence of 2 mM Ca⁺⁺, a rapid decline in light transmission through the solution occurred. The effects of human and rat SP-D on light transmission were not seen in the presence of 10 mM EDTA. Pre-incubation of native rat SP-D with 167 mM maltose also abolished the effect. These findings indicate that SP-D aggregates IAV particles principally by binding to viral carbohydrates. Viral HA binding to carbohydrates on SP-D is unlikely, since HA binding to sialic acids is unaffected by Ca⁺⁺ chelation (29) (see Table I). In addition, the concentrations of maltose used in these experiments did not inhibit HA activity (see Table I). Recombinant rat SP-D caused similar aggregation of the Texas 77 strain of IAV in Ca⁺⁺-containing buffer. Maximal aggregation was achieved with either 613 ng/ml or 1.18 μg/ml of the recombinant protein in three experiments (data not shown).

SPD enhances neutrophil binding of IAV. Results of experimen...
addition of virus. (This amount of maltose did not alter the HA activity of IAV in the absence of SP-D). HA activity was significantly inhibited (P = 0.01) by 3.5 μg/ml rat SP-D or greater, but only in TBS with 2 mM Ca²⁺ without maltose. Human SP-D significantly reduced (P = 0.01) HA titers at all concentrations tested, including 1.7 μg/ml, but again only in TBS with 2 mM Ca²⁺ without maltose.

Fig. 2. SP-D inhibits the hemagglutination activity of IAV. HA titers of concentrated Bangkok 79 IAV alone or after incubation with various amounts of native rat or human SP-D (left and right panels, respectively) are shown. Results represent mean±SEM of four experiments in which Bangkok 79 IAV was pre-incubated for 30 min at 37°C in control buffers or buffers containing the indicated amounts of SP-D, followed by determination of HA titer using human type O red cells. Solid squares and open circles represent samples incubated in TBS with 2 mM Ca²⁺ and TBS with 10 mM EDTA, respectively. Open squares represent samples incubated in TBS with 2 mM Ca²⁺ also, but in this case SP-D was pre-incubated with 167 mM maltose before addition of virus. (This amount of maltose did not alter the HA activity of IAV in the absence of SP-D). HA activity was significantly inhibited (P = 0.01) by 3.5 μg/ml rat SP-D or greater, but only in TBS with 2 mM Ca²⁺ without maltose. Human SP-D significantly reduced (P = 0.01) HA titers at all concentrations tested, including 1.7 μg/ml, but again only in TBS with 2 mM Ca²⁺ without maltose.

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Fig. 3. SP-D promotes aggregation of IAV. Changes in percent light transmission occurring upon addition of SP-D to a stirred suspension of Texas 77 IAV (500 HAU/ml). Either native rat (left panel) or human SP-D (right panel) was added at 300 s. Assays were performed in TBS with 2 mM Ca²⁺, except where EDTA is indicated, in which case the assay was performed in TBS with 10 mM EDTA. Except for assays in EDTA, SP-D (which is stored in TBS containing 10 mM EDTA) was recalcified immediately before assay. Results represent mean±SEM of three similar experiments for the 366 ng/ml dose of rat SP-D, and the 125 and 250 ng/ml doses of human SP-D. The decline in light transmission 5 min. after addition of SP-D was significant (P = 0.01) for these samples. The remaining experiments are representative of two performed. In tracings designated “EDTA” 586 and 400 ng/ml of rat and human SP-D, respectively, were added at 300 s. In EDTA experiments no perceptible change in light transmission occurred upon addition of SP-Ds. In the tracing designated “Maltose,” 586 ng/ml of rat SP-D (in TBS with Ca²⁺) which had been pre-incubated with 167 mM maltose was added at 300 s. No perceptible response to the SP-D occurred in this setting.

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centrations of SP-D are those that were originally present upon incubation with virus. A 10-fold dilution was made upon addition of these IAV + SP-D samples to the neutrophil suspension. Dark squares represent experiments carried out in 2 mM Ca\(^{2+}\)-containing TBS. Open squares represent experiments carried out with SP-D which had been pre-incubated with 167 mM maltose before addition to virus. Open circles represent experiments carried out in TBS containing 10 mM EDTA. Results represent mean±SEM of 3–4 experiments. Mean neutrophil fluorescence was significantly increased \((P \leq 0.05)\) when virus samples which had been pre-incubated with 3.5 or 7 \(\mu\)g/ml of rat SP-D in Ca\(^{2+}\)-containing buffer. Concentrations of 1.7 and 3.5 \(\mu\)g/ml of human SP-D significantly \((P \leq 0.025)\) increased viral binding. These effects were not seen when either rat or human SP-D was pre-incubated with maltose, or when experiments were conducted in EDTA containing buffer. Note that SP-Ds were pre-incubated with IAV before addition to neutrophils in these experiments. When IAV was allowed to bind to neutrophils prior to addition of SP-D the lectin did not alter viral binding (data not shown).

of SP-Ds to cause viral aggregation. Neither native rat nor human SP-D was found to elicit \(\text{H}_2\text{O}_2\) production on their own at these concentrations (data not shown).

**SP-D protects neutrophils against deactivation caused by IAV.** Treatment of neutrophils with a wide variety of IAV strains results in impaired responsiveness of the cells to subsequent stimulation with various agonists (21, 30). The impairment, which we refer to as deactivation, involves chemotactic, respiratory burst, degranulation, and intracellular bacterial killing functions of the neutrophil. We have previously reported that pre-incubation of IAV with either conglutinin or MBL significantly reduces deactivation caused by the virus (10, 11). Using an assay for superoxide (\(\text{O}_2^{-}\)) production in response to FMLP as an index of deactivation, native rat SP-D was similarly protective (see Fig. 7). The neutrophils were first incubated for 10 min with Bangkok 79 IAV alone, or the same amount of IAV which had been pre-incubated with various concentrations of SP-D. FMLP (final concentration 10\(^{-7}\) M) was then added and the resulting \(\text{O}_2^{-}\) response measured. Neutrophils which had been treated with IAV showed a significantly reduced \(\text{O}_2^{-}\) response to FMLP compared to cells not exposed to virus. Neutrophils treated with IAV complexed with SP-D had a significantly greater \(\text{O}_2^{-}\) response to FMLP than did those treated with IAV.

**Figure 4.** SP-D enhances IAV binding to neutrophils. Mean neutrophil fluorescence after exposure to FITC-labeled IAV (Bangkok 79 strain) alone, or FITC-labeled IAV which had been pre-incubated with various concentrations of SP-D, is shown. Results of experiments using native rat and human SP-D are depicted, respectively, in the left and right panels. The same IAV + SP-D samples were used in these experiments as those described in Fig. 1. The concentrations of SP-D are those that were originally present upon incubation with virus. A 10-fold dilution was made upon addition of these IAV + SP-D samples to the neutrophil suspension. Dark squares represent experiments carried out in 2 mM Ca\(^{2+}\)-containing TBS. Open squares represent experiments carried out with SP-D which had been pre-incubated with 167 mM maltose before addition to virus. Open circles represent experiments carried out in TBS containing 10 mM EDTA. Results represent mean±SEM of 3–4 experiments. Mean neutrophil fluorescence was significantly increased \((P \leq 0.05)\) when virus samples which had been pre-incubated with 3.5 or 7 \(\mu\)g/ml of rat SP-D in Ca\(^{2+}\)-containing buffer. Concentrations of 1.7 and 3.5 \(\mu\)g/ml of human SP-D significantly \((P \leq 0.025)\) increased viral binding. These effects were not seen when either rat or human SP-D was pre-incubated with maltose, or when experiments were conducted in EDTA containing buffer. Note that SP-Ds were pre-incubated with IAV before addition to neutrophils in these experiments. When IAV was allowed to bind to neutrophils prior to addition of SP-D the lectin did not alter viral binding (data not shown).

**Figure 5.** Rat SP-D enhances neutrophil activation by IAV. Neutrophil \(\text{H}_2\text{O}_2\) responses elicited by Texas 77 IAV alone, or the same amount of virus which had been incubated with various concentrations of native rat SP-D, are shown. The same samples used in Fig. 2 to assess aggregation of Texas 77 IAV were used in these experiments. The original virus + SP-D samples were diluted from 1.5 to 2 ml by the addition of 0.5 ml of the same buffer used in Fig. 2 (i.e., either TBS with Ca\(^{2+}\) 2 mM [closed squares] or with EDTA 10 mM [open circles]) containing 4 \(\times\) 10\(^4\) neutrophils. Open squares again represent maltose treated SP-D samples. Results represent mean±SEM of three experiments for virus alone and virus treated with 367 ng/ml of SP-D (where \(\text{H}_2\text{O}_2\) production was significantly increased; \(P \leq 0.01\)), and two experiments for other data points.

**Figure 6.** Human SP-D augments neutrophil activation by IAV. Representative scopoletin fluorescence tracings are shown. A suspension of neutrophils was added (where abrupt downward deflection occurs) to solutions of TBS (with 2 mM Ca\(^{2+}\)) containing Texas 77 IAV alone (Control) or IAV which had been pre-incubated with either 125 or 250 ng/ml of human SP-D, as indicated. These tracings are representative of three or six (for the 250 ng/ml concentration) similar experiments. In these experiments pre-incubation of IAV with either 125 or 250 ng/ml of SP-D significantly increased the neutrophil \(\text{H}_2\text{O}_2\) response \((P \leq 0.025)\). No enhanced response was observed in similar experiments carried out in 10 mM EDTA (data not shown).
alone. No $O_2^-$ was generated during incubation of the neutrophils with IAV alone, or IAV complexed with SP-D (data not shown).

Addition of 440 ng/ml (final concentration) of native rat SP-D alone to neutrophils followed after 10 min by addition of FMLP led to a slight enhancement of the neutrophil $O_2^-$ response to FMLP (9±2% increase; $n = 3$; $P < 0.01$) as compared to control neutrophils. When FITC-labeled Bangkok 79 IAV was pre-incubated with this concentration of rat SP-D followed by addition of neutrophils to this mixture, the $O_2^-$ response to FMLP was increased by $31±4$% as compared with cells treated with IAV alone ($n = 4$; $P = 0.05$). The SP-D-opsonized IAV preparation also showed significantly enhanced binding to neutrophils (37±7%; $n = 4$; $P = 0.005$) compared to unopsonized IAV.

HA inhibitory activity of human bronchoalveolar lavage (BAL) fluids. An ELISA assay was developed to measure SP-D levels in BAL fluids obtained from healthy volunteer donors. We first measured the levels of SP-D present in BAL which had only been subjected to a 150 g centrifugation (hence still containing surfactant). The mean SP-D concentration in BAL fluid samples obtained from five donors was 139±29 ng/ml. The total protein concentration for these BAL fluid samples was 34±10 µg/ml. SP-D represented 0.5±0.1% of the total protein in these samples. The anti-SP-D immunoglobulin preparation at the dilution used in this assay had no significant reactivity with purified human SP-A. These SP-D values are similar to those that have previously been reported (18). We then subjected the BAL samples to a 10,000 g centrifugation. The vast majority of SP-A (99%) is insoluble and is concentrated in the surfactant pellet by this maneuver (31). The mean±SEM SP-D level of the 10,000 g supernatant was 104±31 ng/ml. Hence, as previously reported (18, 31) the majority of SP-D is soluble and not surfactant-associated. Based on our findings using purified SP-Ds, the concentrations of SP-D present in these BAL samples would be sufficient to show functional activity vs. IAV.

SP-D present in whole BAL bound to IAV as assessed using a dot blot assay (Fig. 8, Panel A). Sufficient SP-D was also present in the 10,000 g supernatant of BAL to show comparable binding to IAV (Fig. 8, Panel B). No binding of SP-D in BAL to control buffer alone or to BSA was detected. Nonspecific binding of immunoglobulins to the virus was also ruled out. Table II shows that the BAL samples also exhibited substantial HA inhibiting activity. Inclusion of maltose during the HA assay

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Figure 7. SP-D protects neutrophils against deactivation by IAV. Neutrophils were treated with control buffer, Bangkok 79 IAV alone (320 HAU/ml) or the same amount of virus which had been pre-incubated for 20 min with various concentrations of native rat SP-D as indicated. FMLP (10⁻⁴ M) was then added to all samples and the resulting superoxide response over 10 min measured using a continuous cytochrome C assay. The amount of superoxide produced in neutrophils treated with IAV preparations was divided by the amount produced in neutrophils not exposed to IAV to obtain the percent of Control $O_2^-$. Results represent mean±SEM of four experiments. Significantly more superoxide ($P < 0.05$) was produced in response to FMLP in all samples containing SP-D, as compared to that produced in samples treated with IAV alone.

Figure 8. Dot blot assay demonstrating binding of SP-D contained in whole human bronchoalveolar lavage fluids to IAV. Various dilutions of Mem71-lg Be1n IAV was blotted on to nitrocellulose and the blots were exposed to purified native human SP-D (Human SP-D), BAL fluids, or control buffer. After washing, blots were incubated with rabbit anti-human SP-D IgG followed by goat anti-rabbit IgG coupled to HRP as described in Methods. The figure depicts photographic film exposed to the blots after addition of substrate to produce a chemiluminescence reaction over areas where HRP was bound. A shows binding of human SP-D or SP-D present in the 150 g supernatant of BAL fluid to various dilutions of IAV (from undiluted [UD] to a 1:16 dilution). The undiluted IAV stock contained 32,000 HAU/ml and 1 mg/ml of protein. B assesses binding of SP-D present in BAL to IAV diluted either 16- or 32-fold, or to buffer or BSA (200 µg/ml) alone (i.e., no virus present in the latter two lanes). Note that SP-D in BAL did not bind to BSA (200 µg/ml) alone or buffer alone. Purified SP-D also did not bind in this setting (data not shown). In row 1 the supernatant of BAL which had been subjected to a 10,000 g centrifugation (to remove surfactant and SP-A) was used (denoted as BAL [pre-adsorption]). In row 2 the same BAL supernatant was used, but after it had been adsorbed with a maltosyl-agarose column (BAL [post-adsorption]) to reduce the amount of SP-D present (see text). The amount of total BAL protein used was the same in rows 1 and 2 (i.e., 100 µg/ml). In row 3, blots were incubated with buffer only (instead of BAL), followed by addition of antibodies (IgG alone). No chemiluminescence was evident in this row.
IAVS (2,400 HAU/ml of Mem71r-Belr strain) were preincubated in control buffer or buffer containing either bronchoalveolar lavage fluid (BAL) from healthy volunteer donors (n = 5 separate donors used), purified SP-A or purified human SP-D at the indicated concentrations. "BAL 150 g Supernatant" and "BAL 10,000 g supernatant" refer to the supernatant remaining after subjecting the BAL fluid to a either 150 or 10,000 g centrifugation for 1 h. Values represent mean±SEM percent inhibition of control HA titers (i.e., HA titer of BAL-, SP-A-, or SP-D-treated sample/BAL titer of control sample x 100) (n = 3–6 experiments). Where indicated BAL or SP-D was pre-incubated with a final concentration of 167 mM maltose before addition of IAV. When incubated in TBS + Ca²⁺ (without maltose), HA titers in all BAL-, SP-A-, or SP-D-treated samples were significantly reduced compared with control. * Significantly less inhibition (P ≤ 0.05) compared to BAL or SP-D without maltose. † Significantly less inhibition (P ≤ 0.05) compared to BAL before 10,000 g centrifugation. ‡ Significantly more inhibition (P ≤ 0.005) compared with either SP-A or SP-D alone at the same concentrations.

Table II. Inhibition of IAV Hemagglutination Activity by Bronchoalveolar Lavage Fluids, SP-D, or SP-A

<table>
<thead>
<tr>
<th>Condition</th>
<th>Percent reduction in HA titer in BAL- or SP-A-treated samples</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>TBS + Ca²⁺ + maltose</td>
</tr>
<tr>
<td>BAL 150 g supernatant</td>
<td>86±3%</td>
</tr>
<tr>
<td>BAL 10,000 g supernatant</td>
<td>69±6%</td>
</tr>
<tr>
<td>SP-A 3.8 µg/ml</td>
<td>59±6%</td>
</tr>
<tr>
<td>SP-A 7.6 µg/ml</td>
<td>61±8%</td>
</tr>
<tr>
<td>SP-D 113 ng/ml</td>
<td>50±0%</td>
</tr>
<tr>
<td>SP-D 225 ng/ml</td>
<td>56±6%</td>
</tr>
<tr>
<td>SP-D 450 ng/ml</td>
<td>84±3%</td>
</tr>
<tr>
<td>SP-A 3.8 µg/ml + SP-D 113 ng/ml</td>
<td>69±6%</td>
</tr>
<tr>
<td>SP-A 3.8 µg/ml + SP-D 225 ng/ml</td>
<td>78±3%†</td>
</tr>
</tbody>
</table>

Figure 9. Inhibition of IAV HA activity caused by BAL fluid before and after adsorption by maltosyl-agarose. The 10,000 g BAL supernatant pre- and post-adsorption by maltosyl-agarose (prepared as described in Fig. 8 B and D and text) was tested for its ability to inhibit HA activity of the Mem71r-Belr strain of IAV. BAL pre-adsorption is shown with open squares; post-adsorption BAL is shown with dark squares. The µg/ml of total BAL protein added is shown on the horizontal axis. (*) Statistically significant reduction (P = 0.025) in ability of post-adsorption BAL to inhibit HA activity.

Discussion

IAVs are major human and animal pathogens which undergo rapid evolution in their HA genes, enabling them to evade specific cell- and antibody-mediated immune responses (34). Given the substantial morbidity and mortality associated with influenza epidemics, it seems plausible that nonspecific, or "natural," host defense mechanisms may have evolved to protect against these viruses (35). The host response to IAV that occurs prior to the development of specific antibody and cell-mediated immunity is generally successful in that the virus remains confined to the upper respiratory tract, with viremia and viral pneumonia being uncommon (36). We have previously summarized data which support the contention that phagocytes (including neutrophils and macrophages) play a role in the early host defense against IAV infection (37). Also, bacterial superinfection, possibly resulting from IAV-induced phagocyte dysfunction, is the most important cause of morbidity and mortality during IAV epidemics (38). The findings presented herein suggest that the surfactant protein D may, by itself and in combination with phagocytes, also contribute to the initial host defense against IAV. This protein is particularly well situated to protect the lower respiratory tract from invasion by IAV.

SP-D of rat or human origin potently inhibits hemagglutina-

Protective Role of Surfactant Protein D against Influenza Viruses
tion activity of a variety of IAV strains, including representa-
tives of the two major subtypes currently circulating in the 
human population (i.e., H1 and H3). SP-D is also shown to 
cause aggregation of IAV particles. These effects are mediated 
through binding of the SP-D carbohydrate recognition domain 
to viral carbohydrates, as demonstrated by inhibition with malt-
ose or Ca\(^{2+}\)-chelation. The lack of effect of SP-D on the PR-
8 strain of IAV, which lacks glycosylation on the globular head 
of the HA (28) provides further evidence in support of this 
interpretation. Aggregation of viral particles by SP-D may re-
duce infectious particle numbers and enhance clearance by mu-
cociliary and phagocytic mechanisms. Neutrophils bind IAV to 
a greater extent when the virus is complexed with SP-D, perhaps 
in part due to viral aggregation. The concentrations of human 
or rat SP-D which cause enhanced respiratory burst responses 
parallel those which cause viral aggregation. This is consistent 
with the prior observation that larger IAV particles are a more 
effective stimulus for phagocyte respiratory burst activation 
(39). Pre-incubation of IAV with SP-D also protects the cells 
against the deactivating effects of the virus. These results imply 
that opsonization of the virus with SP-D enabled the neutrophil 
to process the virus in a manner less deleterious to the cell. This 
effect suggests that SP-D may be protective against bacterial 
superinfection in vivo.

The concentrations of SP-D required to mediate inhibition 
of IAV HA activity, or cause IAV aggregation are within the 
range of concentrations found in BAL fluids. Local concentra-
tions in the lung are certainly considerably higher than those 
found in unconcentrated BAL fluid. SP-A also inhibited IAV 
HA activity, although SP-A was more than 10-fold less potent 
than SP-D in this regard. Inhibition of IAV infectivity by similar 
concentrations of SP-A was recently reported by van Iwaarden 
et al. (32) as well. SP-D and SP-A exist in different compart-
ments in vivo, with SP-D being largely soluble, while SP-A is 
largely surfactant-associated. Since SP-A levels in whole BAL 
fluid exceed those of SP-D by ~10-fold, it may be that both 
lecitins contribute to IAV inhibitory activity in a cooperative 
manier (see Table II), with the former acting principally at the 
surfactant interface, and the latter in the fluid phase of the 
airways.

The serum proteins MBL and conglutinin, which bear sig-
ificant homology to SP-D, also have similar ability to inhibit 
HA activity and enhance neutrophil responses upon exposure 
to the virus (10, 11). Of note, SP-D exceeds these serum lectins 
in potency in these assays. Concentrations of MBL are 
required to achieve similar HA inhibition, or respiratory burst enhance-
ment exceed those of SP-D by 10-fold or more. While the HA 
inhibitory potency of conglutinin is similar to that of SP-D, to 
achieve optimal IAV aggregation or neutrophil respiratory burst 
enhancement 1.5 µg/ml (final concentration) of conglutinin are 
required as compared to between 250 and 500 ng/ml of SP-Ds. 
SP-D is also more potent than MBL or conglutinin at protecting 
neutrophils from deactivation by IAV. The lesser potency of 
MBL and SP-A as compared to SP-D and conglutinin may, in 
part, be attributable to the smaller size of the former molecules. 
It should be noted that SP-A and SP-D obtained from patients 
with alveolar proteinosis may differ in some respects from SP-
A or SP-D present in normal donors. In any case, our finding 
that siilicotic and recombinant rat SP-D and SP-D present in 
normal donor BAL had similar anti-IAV activities as the alveo-
lar proteinosis proteins suggests that our results are not attribu-
table to aberrances of the latter preparations.

A thorough understanding of lectin interactions with IAV 
may allow prediction of which viral strains are likely to be most 
virulent in vivo, or which hosts may be particularly vulnerable 
to adverse sequelae of influenza. It may be that strains of IAV 
with glycosylation patterns that do not allow lectin binding to 
the virus are more capable of causing viral pneumonia, viral 
infection of extrapulmonary tissues, or phagocyte deactivation 
and bacterial superinfection. Better understanding of the pre-
immune host response to IAV may lead to ways of predicting 
which patients are most at risk for complications of influenza 
(e.g., because of congenital or acquired differences in lectin 
production) and possibly to new modes of therapeutic inter-
vention.

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