Cigarette Smoke Can Activate the Alternative Pathway of Complement In Vitro by Modifying the Third Component of Complement

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Abstract

Cigarette smoking is associated with significant increases in the number of pulmonary mononuclear phagocytes and neutrophils. A potent chemoattractant for these cells is C5a, a peptide generated during complement (C) activation. We, therefore, investigated the possibility that cigarette smoke could activate the complement system in vitro. Our results show that factor(s) (mol wt < 1,000) present in an aqueous solution of whole, unfiltered cigarette smoke can deplete the hemolytic capacity of whole human serum in a dose-dependent manner. The particle-free, filtered gas phase of cigarette smoke is inactive. The smoke factor(s) do not activate serum C1, but do deplete serum C4 activity. Treatment of purified human C3 with whole smoke solution modifies the molecule such that its subsequent addition to serum (containing Mg/EGTA to block the classical pathway) results in consumption of hemolytic complement by activation of the alternative pathway. Smoke-modified C3 shows increased anodal migration in agarose electrophoresis, but this is not due to proteolytic cleavage of the molecule as evidenced by sodium dodecyl sulfate-polyacrylamide gel electrophoresis. In contrast to methylamine-treated C3, C3 treated with smoke is only partially susceptible to the action of the complement regulatory proteins Factors H and I. In addition, smoke-modified C3 has diminished binding to Factor H as compared with methylamine-treated C3. Finally, smoke-modified C3 incorporates [14C]methylamine which suggests that the thiolester bond may be intact.

These data indicate that aqueous whole cigarette smoke solution can modify C3 and activate the alternative pathway of complement, perhaps by a previously unrecognized mechanism. Should this occur in vivo, complement activation might partly account for the extensive pulmonary leukocyte recruitment observed in smokers.

Introduction

Cigarette smoking has been identified as the major causative factor in the development of pulmonary emphysema (1), but little is known of how cigarette smoke actually contributes to the pathogenesis of the disease. The characteristic histopathologic observation in the emphysematous lung is excessive tissue dissolution which is thought to result from an unrestrained proteolytic attack on lung interstitium, particularly the elastic fibers (reviewed in 2, 3). Neutrophils and mononuclear phagocytes (alveolar macrophages and monocytes) have been implicated as the source of this increased protease (primarily elastase) burden (4–6). The lungs of cigarette smokers are known to contain significantly increased numbers of these elastase-bearing phagocytes (7, 8), but the mechanism of this smoke-induced leukocyte recruitment is poorly understood.

Recently, efforts have focused on mechanisms whereby cigarette smoke may indirectly recruit phagocytic cells into the lung. Many of these studies have examined the elaboration of neutrophil chemoattractants by stimulated alveolar macrophages of smokers (9–12). In addition, proteolytic destruction of lung connective tissue may, in itself, contribute to the leukocyte recruitment. For example, proteolytically generated fragments of collagen (13) and fibronectin (14) have been demonstrated to be chemotactic for peripheral blood monocytes, and incubation of human elastin with neutrophil elastase results in the production of biologically active peptides which are also chemotactic for neutrophils and monocytes (15).

Less attention, however, has been paid to the possibility that neutrophil chemoattractants may be generated in the lung fluids of smokers as a direct result of complement (C) activation by inhaled cigarette smoke. It is well documented that when the complement cascade is activated, the fifth component of complement (C5) is proteolytically cleaved, resulting in generation of the chemotactically active peptide C5a (reviewed in 16). C5a is one of the most potent neutrophil and monocyte chemoattractants yet described and is biologically active at picogram quantities. Moreover, a recent report demonstrates that physiological concentrations of nicotine enhance neutrophil responsiveness to C5a (17).

We reported earlier that rats acutely exposed to cigarette smoke had significantly increased levels of chemotactic activity for neutrophils and monocytes in their lung fluids. However, prior depletion of circulating complement by treatment of the animals with cobra venom factor prevented this smoke-induced chemotactic activity (18). In addition, others have recently demonstrated consumption of complement hemolytic activity in serum following its incubation with cigarette smoke condensate (19, 20). Although suggestive, our preliminary studies

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1. Abbreviations used in this paper: C, complement; Ea, unsensitized rabbit erythrocytes; EA, sheep erythrocytes coated (sensitized) with goat anti-sheep erythrocyte antibody; EAC1, EA coated with human C1; EAC3b, sheep erythrocytes coated with human C3b; GVB2+, isoionic veronal-buffered saline (pH 7.4) containing 0.15 mM CaCl2, 0.5 mM MgCl2, and 0.1% gelatin; Mg-GVB, isoionic veronal-buffered saline (pH 7.4) containing 0.5 mM MgCl2 and 0.1% gelatin; NHS, normal human serum; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis.
and the others referred to above did not explore possible chemical mechanisms of complement activation by cigarette smoke.

Therefore, we undertook in vitro studies to investigate the effect of cigarette smoke on serum complement and purified complement components. In the present report, we show that incubation of whole human serum with an aqueous solution of unfiltered cigarette smoke depletes hemolytic activity. Serum C3 was not activated under these conditions; however, purified C3 preincubated with smoke solution and subsequently added to serum (containing 5 mM Mg-EGTA to block the classical pathway) did activate complement. These and other results to be discussed below suggest that aqueous cigarette smoke components are capable of triggering the alternative pathway of complement, probably by modifying C3.

Methods

Buffers. Buffers used were as follows: Dulbecco’s phosphate-buffered saline (PBS), pH 7.2; isotonic veronal-buffered saline (VB), pH 7.4; VB containing 0.15 mM CaCl$_2$ and 0.5 mM MgCl$_2$ (VB$^{++}$); VB$^{++}$ containing 0.1% gelatin (GVB$^{++.}$); VB containing 0.1% gelatin and 0.01 M EDTA (GVB$^{E}$); GVB$^{+}$ containing 2.5% (wt/vol) sucrose (SGVB$^{+}$); 0.1 M MgCl$_2$, 0.1 M EGTA adjusted to pH 7.0 (Mg-EGTA); and VB containing 0.5 mM MgCl$_2$ and 0.1% gelatin (Mg-GVB).

Purified complement proteins. Highly purified C3 was isolated using the method of Tack and Prahl (21). The C3 was then further purified by passage over a Sepharose 4B (Pharmacia Fine Chemicals, Inc., Piscataway, NJ) affinity column containing immobilized goat IgG antibodies (Dako Corp., Santa Barbara, CA) to human IgA, IgG, and IgM in order to remove traces of contaminating immunoglobulins. Small amounts of C3b and C3H$_2$O$_4$ were finally removed by chromatography on QAE-Sephadex (Pharmacia Fine Chemicals, Inc., Piscataway, NJ) according to the method of Janatova et al (22). C3 preparations were routinely rerechromatographed on QAE-Sephadex after prolonged storage at −85°C to remove the accumulated C3H$_2$O$_4$. Such purified C3 contained no immunochromatically detectable C5.

C3b was generated from purified C3 (2 mg/ml) by the addition of 1% (wt/wt) L-(cosalylamido 2-phenyl)ethyl chloromethyl ketone-treated trypsin ( Worthington Biochemical Co., Freehold, NJ) for 2.5 min at 22°C followed by 2% (wt/wt) soybean trypsin inhibitor (Sigma Chemical Co., St. Louis, MO) to stop the reaction.

Factors H and I were purified as described (23). The highly purified Factor H used in our binding experiment (see below) was a generous gift from Dr. Michael Pangburn (Scripps Clinic and Research Foundation, La Jolla, CA). Aggregated IgG. Human IgG was first isolated from serum by a single step procedure using DEAE Affi-Gel Blue, according to the manufacturer’s specifications (Bio-Rad Laboratories, Inc., Richmond, CA). IgG was then aggregated by incubation at 63°C for 20 min at a concentration of 10 mg/ml, as described (24). Particulate aggregates were removed by centrifugation at 200 g for 5 min and the supernate was used as soluble, aggregated IgG.

Sera and antisera. A pool of normal human serum (NHS) was obtained from young, apparently healthy, nonsmoking volunteers.

Monospecific, polyvalent antisera to C1q, C3, and C4 were raised in New Zealand White rabbits by multiple subcutaneous and intramuscular injections of 0.1 mg of purified antigen in Freund’s complete adjuvant. Antihuman C3 was also purchased from Cappel Laboratories (West Chester, PA). Antiserum to C1r and C1s were raised in goats, and also purchased from Atlantic Antibodies (Scarborough, ME).

Protein determinations. Protein concentrations were determined by the method of Lowry (25) using bovine serum albumin as a reference standard. The concentration of C3 was also determined by measuring absorbance at 280 nm using an $E_{1\text{cm}}^{\text{max}}$ of 9.7 as previously described (21).

Preparation of smoke solutions. Whole, aqueous, cigarette smoke solutions were prepared by a modification of the method of Carp and Janoff (26). The smoking apparatus consisted of a 50-ml plastic syringe fitted with a 3-way stopcock (Pharmaseal K-75, American Hospital Supply, Edison, NJ) to which the cigarette and a 0.1-ml glass capillary tube were attached. The capillary tube was placed into a 4-ml plastic tube containing 1.0 ml of PBS, and the tube was immersed in a 37°C water bath. All materials used in the preparation of smoke solutions (i.e., syringe, 3-way stopcock, capillary tube, plastic tube, and PBS) were sterile, pyrogen-free, and used only once. Smoke solutions were prepared using 2R1 Kentucky Reference cigarettes (a generous gift from Dr. D. Layton Davis, Tobacco and Health Research Institute, University of Kentucky, Lexington, KY) which were stored at −85°C. Before use, the cigarettes were thawed, and then rehumidified for 24 h at 60% relative humidity in a sealed chamber at room temperature. The smoking regimen, based on the standard smoking cycle (27), was as follows: 35 ml of tobacco smoke was drawn into the syringe in a 2-s puff and then slowly bubbled into the buffer. One puff of smoke per minute was drawn until a 5-cm butt length of cigarette was consumed (usually eight or nine puffs). In this manner, the smoke of six cigarettes was bubbled through 1.0 ml of PBS at 37°C, after which the solution was decanted to a new tube to remove the accumulated tar and large particulate matter. Since the final smoke solutions were slightly acidic, the pH was readjusted with 1.0 N NaOH to match that of the starting buffer. Smoke solutions prepared in this manner varied only ±10% in their ability to consume 50% of serum complement.

In some experiments, the whole aqueous smoke solutions were exhaustively dialyzed before use against PBS at 4°C using Spectra-Por 6 dialysis tubing (mol wt limit = 1000; Spectra Medical Industries, Inc., Los Angeles, CA) and these were designated as dialyzed smoke solutions. Aqueous solutions of the particle-free, gas phase of cigarette smoke were also prepared exactly as described above except that a Cambridge filter pad (44-mm diam, 1.5-mm thick, 0.22 μm pore size; Phipps and Bird, Inc., Richmond, VA) was inserted between the cigarette and the 3-way stopcock. The filter pad was replaced after each cigarette was consumed.

Aliquots of each smoke solution were tested for endotoxin contamination using the Limulus Amebocyte Lysate kit (M. A. Bioproducts, Walkersville, MD) and were found to be negative. After preparation, all smoke solutions were either stored at −85°C or used immediately.

Treatment of C3 and NHS with smoke solutions. Purified C3 and NHS were treated with smoke solutions, methylene, or PBS (the amounts are described in the appropriate section of Results) for 60 min at 37°C. The mixtures were then exhaustively dialyzed against PBS at 4°C, using a dialysis membrane with a mol wt limit of 50,000 (Spectra Medical Industries, Inc.), to remove unreacted, dialyzable smoke components or excess methylene. Preparations were stored at −85°C or used immediately.

Electrophoretic procedures. C3 and NHS, treated as above, were electrophoresed at 3-4 V/cm for 2.5 h at room temperature in 1% agarose (Seakem LE, FMC Corp., Rockland, ME) using 0.1 M barbital buffer (pH 8.6). After electrophoresis, C3 was visualized with a monospecific, polyvalent antihuman C3. Electrophoresis in polyacrylamide gel containing sodium dodecyl sulfate (SDS-PAGE) was performed according to the method of Laemmli (28). Staining of gels for proteins was performed using Coomassie Brilliant blue R-250 or Amido black 10B (Bio-Rad Laboratories, Inc.).

Hemolytic assays. Sheep erythrocytes sensitized with rabbit anti-sheep erythrocyte antibody (EA) and EA coated with human C1 (EAC1) were prepared according to established methods (29). Materials used in the preparation of EA and EAC1 were purchased from Cordis Laboratories, Inc. (Miami, FL). Unsensitized rabbit erythrocytes (E$_{R}$) were freshly isolated from a New Zealand White rabbit and then washed several times with cold Mg-EGTA. Hemolytic titration of serum C4 was performed as previously described (29). Depletion of serum hemolytic activity was measured by adding various amounts of smoke solutions, methylene, and/or PBS to 0.1 ml of NHS so that the final volume of the mixture was 0.2 ml. After a 60-min incubation at
37°C, 0.02 ml of the reaction mixture was removed and added to 0.1 ml of EA (5 x 10^6/m), the volume was adjusted to 0.4 ml with GVB2+, and the incubation was continued for an additional 60 min at 37°C. The hemolytic reaction was terminated by adding 1.6 ml of ice-cold GVB2+, the mixtures were centrifuged, and the percentage of hemolysis was measured as the absorbance of free hemoglobin in the supernate at 412 nm. The percent lysis of EA incubated with 0.01 ml of untreated NHS was used as a reference for 0% depletion of hemolytic activity. Since whole smoke solution has its own color, all hemolytic assays had smoke-color controls consisting of smoke solution, EA, and GVB2+. This absorbance value was subtracted from the smoke-treated experimental samples.

Activation of the alternative pathway was measured by pretreating 0.05 ml of purified C3 (1 mg/ml) with various amounts of smoke solutions, methylamine, and/or PBS so that the final volume was 0.1 ml. Following a 60-min incubation at 37°C, 0.1 ml of NHS containing 5 mM Mg-EGTA was added to each mixture and the reaction was allowed to proceed for an additional 60 min at 37°C. Next, 0.1 ml of each mixture was removed and added to 0.1 ml of Eq (5 x 10^6/m), the volume was adjusted to 0.4 ml with Mg-GVB, and the incubation continued at 37°C for 60 min. The reaction was stopped by addition of 1.0 ml of ice-cold Mg-GVB. After a low speed centrifugation, the absorbance of the supernate was measured as before at 412 nm.

**Immunochemical determination of C1 activation in serum.** NHS was incubated with an equal volume of whole smoke solution, PBS, or aggregated IgG (2 mg/ml) for 60 min at 37°C. These mixtures were then analyzed by single radial immunodiffusion against anti-C1r. The immunodiffusion slides contained 1% agarose in 0.03 M potassium phosphate (pH 8.0), 0.1 M NaCl, and 0.01 M EDTA, and the appropriate dilution of anti-C1r as previously described (30). Double (Ouchterlony) immunodiffusion analysis was also carried out on these samples using 0.8% agarose in veronal-buffered saline (pH 7.2) containing 2.5 mM CaCl2 and 0.09 M NaCl, and precipitin lines were developed with monospecific, polyvalent antihuman C1q, C1r, and C1s.

**Effect of Factors H and I on treated C3.** Samples of C3, 0.1 ml at 1.6 mg/ml, were treated with smoke solution, methylamine, or PBS. The treated C3 (160 µg) was then reacted for 45 min at 37°C with physiological concentrations of Factors H (50 µg) and I (3.4 µg) in a total volume of 0.2 ml. C3 cleavage products were subsequently visualized by SDS-PAGE under reducing conditions.

**Binding of Factor H to treated C3.** Factor H was radioiodinated (0.5 mCi/mg) using 125I-Bolton-Hunter reagent (New England Nuclear, Boston, MA) according to the manufacturer’s specifications. Sheep erythrocytes coated with C3b (EC3b) were prepared as described in (31). A competitive 125I-labeled Factor H binding assay was then carried out to test for activated C3 after treatment with smoke solution, methylamine, or PBS, according to the method of Pangburn and Müller-Eberhard (32). C3b, generated by treating C3 with trypsin (see above), was used as the positive control in this assay.

**Treatment of C3 with radiola beled methylamine or iodoacetamide.** 0.5 ml of C3 (2.5 mg/ml), pretreated with smoke solution, methylamine, or PBS as described above, was reacted with 4 mM [14C]methylamine (46 mCi/mmol; New England Nuclear) for 4 h at 37°C. Excess isotope was subsequently removed by dialysis against PBS at 4°C. Approximately 0.32 mg of each C3 was then added to 10 ml of Dimiscint (National Diagnostics, Somerville, NJ) and the uptake of labeled methylamine was determined by liquid scintillation counting.

Radioalkylation of C3 pretreated with smoke, methylamine, or PBS was carried out using [14C]iodoacetamide (23.6 mCi/mmol; New England Nuclear) according to the method of Janatova et al. (33).

**Results**

**Effect of aqueous cigarette smoke on serum hemolytic activity.** Fig. 1 demonstrates that aqueous, unfiltered cigarette smoke solution depletes the hemolytic capacity of NHS in a dose-dependent fashion and in a manner similar to that which occurs after treatment of serum with methylamine, a nucleophile known to activate the alternative pathway of complement by cleaving the thiolester bond in C3 (31). This smoke-induced depletion is not due to endotoxin contamination because smoke solutions were determined to be endotoxin negative using the Limulus assay. In addition, loss of serum hemolytic activity is not an artifact due to direct effects of smoke on the EA, because smoke-treated and untreated EA were equally susceptible to complement-mediated lysis by NHS. Dialyzed smoke solutions (1000-mol wt limit membrane) did not deplete serum complement activity. Similarly, the particle-free, aqueous gas phase of smoke had no effect on the hemolytic capacity of NHS. Finally, nicotine, ammonia, and cyanide, which are all present in relatively large quantities in cigarette smoke, failed to deplete serum hemolytic activity over a wide range of concentrations tested.

**Effect of smoke on purified C3.** Since smoke appeared to diminish serum hemolytic complement activity, we chose next to examine the effect of smoke on purified C3 because (a) C3 is the complement component found in the highest concentration in serum and is essential for the activation of both the classical and alternative pathways, and (b) smoke contains a number of substances (including methylamine) which can potentially break the thiolester bond in C3 and activate the alternative pathway. Fig. 2 demonstrates that incubation of purified C3 with smoke solution modified the activity of the protein such that its subsequent addition to serum caused consumption of complement. Before incubation with smoke, the C3 used in this experiment was rigorously purified, as described in Methods, to free it from contaminating C3b and C3(H2O), both of which can activate the alternative pathway.
Such purified C3, incubated with buffer, did not activate complement upon addition to serum. Since the C3 plus smoke mixtures were exhaustively dialyzed through large-pore membranes before their addition to serum, the resultant complement activation was due to modified C3 and not to residual active smoke components, because the latter are dialyzable through 1000-mol wt limit membranes (Fig. 1). Also, because the serum contained Mg-EGTA to prevent classical pathway activation, the smoke-modified C3 apparently activated the alternative pathway. Under the same conditions, methylamine-treated C3 (a known activator of the alternative pathway) gave similar results (Fig. 2).

When smoke-treated C3 was examined by immunoelectrophoresis, it showed increased anodal migration similar to that of C3b (Fig. 3). The smoke-induced change in electrophoretic mobility was not due to proteolytic cleavage of C3 because both the alpha and beta chains of smoke-treated C3 had an identical molecular weight as those of untreated, native C3 when analyzed by SDS-PAGE under reducing conditions (Fig. 4). When NHS was treated with smoke solution (then dialyzed) and examined by immunoelectrophoresis against anti-C3, increased anodal migration of C3 was again observed (data not shown). The smoke-induced electrophoretic modification of C3 could not be reproduced by treating C3 with methylamine (Fig. 3). In addition, the filtered gas phase of smoke and dialyzed whole smoke had no effect on the electrophoretic mobility of C3 (data not shown).

**Effect of smoke on serum C1 and C4.** To determine whether cigarette smoke solution could also activate the classical pathway, we examined the effect of smoke on C1 and C4 in whole serum. C1 activation was assessed by the radial immu-

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**Figure 2.** Effect of whole smoke and methylamine on purified C3. 50 µl of C3 (1 mg/ml) was pretreated with the indicated amounts of whole smoke solution (—), methylamine (CH₃NH₂) (-- - - - -- ), or PBS (— o —); the final volume of the mixtures was then adjusted to 0.1 ml with PBS. Following a 60-min incubation at 37°C, mixtures were exhaustively dialyzed against PBS at 4°C in a 0.5-ml conical microtube. 0.1 ml of NHS containing 5 mM Mg-EGTA was then added to each sample and incubated for 60 min at 37°C. 0.1 ml of each mixture was removed and added to 0.1 ml of FE (5 × 10⁷/ml), the volume was adjusted to 0.4 ml with Mg-GVB, and samples were incubated for an additional 60 min at 37°C. Hemolysis was measured as absorbance at 412 nm as before. Data are expressed as the percent consumption of hemolytic C in Mg-EGTA-treated serum, using untreated serum as the 0% reference (100% hemolysis).

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**Figure 3.** Electrophoretic mobility of C3 (3.2 mg/ml) treated with an equal volume of PBS (1), 1% trypsin (wt/wt) for 2.5 min at 22°C (2), an equal volume of whole smoke solution (3), or 50 mM methylamine (4). Following dialysis against PBS at 4°C, ~25 µg of C3 was applied to each well and electrophoresed in 1% agarose (in 0.1 M barbitral buffer, pH 8.6) for 2.5 h at room temperature. 0.1 ml of monospecific, polyvalent antihuman C3 was then added to each trough; C3 precipitates were visualized after a 12–24-h immunodiffusion period. The gel was subsequently stained with Coomassie blue after a 24-h soak in 0.9% NaCl plus 0.1% sodium azide (to remove nonprecipitated protein). Direction of C3 migration was toward the anode (+).

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**Figure 4.** SDS-PAGE of C3 (3.2 mg/ml) treated with an equal volume of PBS (lane 1), 1% (wt/wt) trypsin for 2.5 min at 22°C (C3b) (lane 2), an equal volume of whole smoke solution (lane 3), and 50 mM methylamine (lane 4). Following dialysis against PBS at 4°C, ~25 µg of C3 was applied to each lane and electrophoresis was carried out in 7% polyacrylamide under reducing conditions (1% 2-mercaptoethanol). The gel was subsequently stained with Coomassie blue. The 120- and 75-kD bands are the intact alpha and beta chains of C3, respectively. The alpha chain of C3b (alpha') is decreased by proteolytic cleavage to 110 kD.
The technique is based on the fact that C1 inactivator (Cl-INA) binds to Clr (but not the zymogen, Clr) when C1 is activated in serum. The resultant enzyme-inhibitor complex masks antigenic sites on the active enzyme (Clr). Therefore, a diminution of Clr antigenic reactivity in the test is an indication of Cl-INA-Clr complex formation, which in turn is an indication of prior Cl activation. Fig. 5 shows that incubation of serum with smoke solution did not result in a loss of anti-Clr reactivity, which was identical to that observed using buffered-treated serum. These results were obtained in both single radial immunodiffusion (Fig. 5 A) and Ouchterlony double diffusion tests (Fig. 5 B). In contrast, serum treated with aggregated IgG, a known activator of Clr and the classical pathway, showed a marked reduction in its reaction with anti-Clr in both tests (Figs. 5 A and B).

The effect of smoke on hemolytic levels of serum C4 is shown in Fig. 6. Aqueous solutions of unfiltered cigarette smoke totally depleted C4 activity of serum, as did treatment with aggregated IgG. Serum incubated with buffer retained full C4 activity. Although this result appears to be in conflict with the preceding data which indicated absence of classical pathway activation by cigarette smoke, the effect of smoke on serum C4 may be explained by the fact that an internal thiolester bond is present in this protein as well as in C3 (34). This point will be discussed in further detail below.

Effect of smoke on C3 cleavage by Factors H and I. Since it appeared that smoke treatment produced a modified form of C3 that is capable of activating the alternative pathway, we proceeded to test whether smoke-treated C3 was susceptible to inactivation by the complement regulatory proteins, Factors H and I. Factor H has the ability to bind to C3b and “C3b-like” C3 (a form of C3 activated by thiolester cleavage, e.g., methylamine), but not to native C3. After binding, Factor H acts as a cofactor for proteolytic cleavage of the alpha chain of C3 by Factor I (35). Fig. 7 demonstrates that smoke-treated C3 showed partial cleavage of its alpha chain (lane 4) qualitatively similar to (but not quantitatively comparable with) methylamine-treated C3, in which there was substantial cleavage of the alpha chain to the predicted 76- and 43-kD fragments (lane 2). Under the same conditions, C3 treated with buffer (lane 6) showed almost no cleavage of its alpha chain. In the case of the buffer control, the small amount of cleavage observed was probably due to C3(H2O), a “C3b-like” C3 that could have formed spontaneously during the incubation and dialysis period. Densitometric analysis of this gel revealed that alpha chain cleavage and appearance of the 76- and 43-kD products from C3 treated with smoke were intermediate in amount between those of methylamine and buffer-treated C3.

Binding of radioiodinated Factor H to smoke-treated C3. Since smoke-treated C3 was not quantitatively comparable with methylamine-treated C3 with respect to alpha chain cleavage by Factors H and I, we chose next to test whether C3 treated with whole smoke solution was capable of binding 125I-Factor H. In this assay, the binding of Factor H is competitive between EC3b and fluid-phase “C3b-like” C3 (produced by methylamine treatment) or true C3b. The greater the binding of 125I-Factor H to fluid-phase “C3b-like” C3 or C3b, the lower is its binding to EC3b. Table I shows that methylamine-treated C3, like C3b, competed for 125I-Factor H; however, smoke-treated C3 bound substantially less Factor H as did buffer-treated C3. This data supports the previous observation concerning cleavage of smoke-treated C3 by Factors H and I and is further evidence that smoke-treated C3, while capable of activating the alternative pathway of complement, is not entirely comparable in its functional behavior to methylamine-treated C3.

Effect of smoke on the incorporation of radiolabeled me-
Figure 7. SDS-PAGE analysis of the cleavage of treated C3 by Factors H and I. Samples of C3 (0.1 ml at 1.6 mg/ml) were pre-treated for 60 min at 37°C with an equal volume of whole smoke solution, PBS, or 50 mM methylamine (final concentration) and subsequently dialyzed against PBS at 4°C. Lane 1, C3 pretreated with methylamine; lane 3, C3 pretreated with whole smoke solution; lane 5, C3 pretreated with PBS. Aliquots of the above treated samples were also reacted with Factors H (50 μg) and I (3.4 μg) for 45 min at 37°C. Lane 2, C3-methylamine plus Factors H and I; lane 4, C3-smoke plus Factors H and I; lane 6, C3-buffer plus Factors H and I. ~10 μg of protein was applied to each lane and electrophoresed in 8.5% polyacrylamide under reducing conditions (1% 2-mercaptoethanol). The gel was subsequently stained with Amido black. The two chains of reduced native C3 are observed at 120 kD (alpha chain) and 75 kD (beta chain). Factors H- and I-generated cleavage products of the alpha chain of C3 are present at 76 kD (not readily observed in the figure) and 43 kD (observed in the figure). Additional Factor I-generated alpha chain cleavage products are seen below the 43-kD band. Factor H is observed as a single band at 150 kD; the concentration of Factor I was below the level of detection. Note that diminution of the alpha chain and subsequent appearance of the 43-kD cleavage product is minimal after reaction of buffer-treated C3 (negative control) with Factors H and I. By contrast, there is substantial diminution of the alpha chain and appearance of a 43-kD cleavage product after methylamine-treated C3 (positive control) is reacted with Factors H and I. Smoke-treated C3 undergoes partial alpha chain cleavage when incubated with Factors H and I, which is qualitatively but not quantitatively similar to methylamine-treated C3.

Table I. Binding of Radioiodinated Factor H to Smoke-Treated C3

<table>
<thead>
<tr>
<th>Fluid-phase competitor</th>
<th>Percent 125I-Factor H bound to EC3b</th>
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<tbody>
<tr>
<td>None</td>
<td>100</td>
</tr>
<tr>
<td>C3-PBS</td>
<td>71.4±5.7</td>
</tr>
<tr>
<td>C3-smoke</td>
<td>62.0±2.1</td>
</tr>
<tr>
<td>C3-methylamine</td>
<td>45.6±11.1</td>
</tr>
<tr>
<td>C3b</td>
<td>39.8±3.6</td>
</tr>
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</table>

The data are given as percent of 100 ng of 125I-Factor H (0.5 mCi/mg) bound to 5 x 10^-7 EC3b in the presence of 2.0 μg of fluid-phase C3 competitors in a total volume of 0.1 ml in SGVB. The C3, EC3b, and 125I-Factor H mixtures were incubated for 10 min at room temperature. The mixtures were then layered onto 0.3 ml of 20% sucrose in SGVB; cells were then rapidly pelleted using a microfuge. The bottoms of the microfuge tubes were cut off and the pellets counted for bound 125I. C3b was prepared by trypsinization of C3, as described in Methods, and used as the positive control. Values represent mean±SD of triplicate assays.

Discussion

In this paper, we present evidence showing that cigarette smoke is capable of activating the alternative pathway of complement in vitro. This finding confirms observations in two recent reports which showed that cigarette smoke condensate (tar fraction) activates complement in vitro (19, 20). Such results, along with our preliminary report of complement-dependent increases in chemotactic activity in the lung washings of smoke-exposed rats (18), raise the possibility that smoke-

Table II. Effect of Smoke on the Incorporation of [14C]Methylamine by C3

<table>
<thead>
<tr>
<th>Pretreatment of C3</th>
<th>Disintegrations per minute/0.32 mg C3</th>
<th>[14C]methylamine/C3 mol/mol</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>192,334±4726</td>
<td>1.10</td>
</tr>
<tr>
<td>Buffer (PBS)</td>
<td>189,014±3231</td>
<td>1.08</td>
</tr>
<tr>
<td>Whole smoke</td>
<td>132,425±4423*</td>
<td>0.76</td>
</tr>
<tr>
<td>Methylamine</td>
<td>13,322±1657</td>
<td>0.08</td>
</tr>
</tbody>
</table>

0.5 ml of C3 (2.5 mg/ml) was treated with 0.25 ml of whole smoke solution, 50 mM methylamine (final concentration), or 0.25 ml of PBS for 60 min at 37°C. Following an exhaustive dialysis at 4°C against PBS, the treated C3 was reacted with 4 mM [14C]methylamine (46 mCi/mol) for 4 h at 37°C as described in Methods. Values represent mean±SD of quadruplicate assays.

* Corrected for incorporation of label by nondialyzable smoke components (mol wt > 50,000). Values obtained after treating dialyzed smoke alone with 4 mM [14C]methylamine were subtracted from values obtained after reacting dialyzed mixtures of smoke and C3 with the isotope.)

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induced complement activation may also occur in humans. Should this prove true, chronic, low-grade activation of complement in the lung fluids of smokers could be an important initiating step in the pathogenesis of small airway inflammation in such individuals. Small airway inflammation, in turn, is thought to be a pathogenetic factor in the development of pulmonary emphysema (2, 3).

The factor(s) in cigarette smoke which deplete the hemolytic activity of serum and also produce a form of C3 that can activate the alternative pathway appear to have a mol wt < 1,000 and are found only in whole smoke. Numerous small nucleophiles, including methylamine and hydrazine (both of which can cleave thiolester bonds and produce “C3b-like” C3), are present in cigarette smoke (1, 36, 37). One would expect these substances to be found in the gas phase of smoke, which we report to be inactive, but at the pH of cigarette smoke (~6.0), volatile amines are protonated and thus are found in the particulate phase (1). However, the observed effect of smoke on complement cannot be due to the action of these individual smoke components since their concentration in cigarette smoke is far too low to explain the results. For example, using the published values of 4.6 μg methylamine and 30 ng hydrazine/cigarette (1, 37), our smoke solution, prepared as described in Methods, contains ~0.9 mM methylamine and 5.6 μM hydrazine (compare these concentrations with the effective concentration range for methylamine shown in Figs. 1 and 2). In addition, ammonia, at concentrations found in cigarette smoke, had no effect on serum complement, although it has previously been reported that higher concentrations (50 mM) of ammonia resulted in 50% production of “C3b-like” C3 after a 60-min incubation (31).

C3 treated with smoke, like methylamine-treated C3, has no change in molecular weight as determined by SDS-PAGE. In contrast to methylamine treatment, smoke-treated C3 has increased anodal mobility in agarose electrophoresis, indicating a more negatively charged molecule. Whether this is due to a secondary (artifactual) effect of cigarette smoke or is essential for the smoke-induced activating capacity of C3 is currently being investigated.

Smoke-treated C3 apparently does not bind Factor H as do “C3b-like” C3 or C3b and this may explain why C3 smoke is only partially susceptible to the action of Factors H and I in contrast to methylamine-treated C3. (Incidentally, poor binding of Factor H to smoke-modified C3 would theoretically facilitate complement activation.) Further, smoke-treated C3, like buffer-treated C3, incorporates radiolabeled methylamine but not iodoacetamide, while C3 treated with methylamine yields the opposite results. For all these reasons, it can be suggested that the modified form of C3 produced by cigarette smoke is probably different from that produced by methylamine, although both forms can activate the alternative pathway of complement. However, our data do not rule out the possibility that some C3 molecules undergo thiolester cleavage after exposure to cigarette smoke (see partial cleavage of smoke-treated C3 by Factors H and I, Fig. 7; see also the 25% decrease in incorporation of [14C]methylamine by C3 after smoke treatment, Table II).

Although smoke does not activate C1, indicating that it does not activate the classical complement pathway, smoke completely consumes serum C4. Since C4, like C3, contains an internal thiolester bond (34), cigarette smoke may directly consume C4 by cleaving its thiolester without activating the classical pathway. Alternatively, since C3 and C4 share a considerable degree of homology in the primary sequence of their alpha chains (38, 39), this could render both molecules susceptible to smoke-induced modifications by a thiolester-independent mechanism. Such possibilities remain to be tested.

We have shown that exposure of C3 to smoke solution produces a functionally modified form of the molecule which is capable of activating the alternative pathway of complement. It remains unclear whether this smoke-modified C3 is similar to methylamine-treated C3 in that its internal thiolester bond is broken or whether smoke treatment produces a heretofore unrecognized form of C3 which has the ability to activate the alternative pathway without cleavage of its thiolester. Clearly, however, smoke-treated C3 is not identical to methylamine-treated C3.

Cigarette smoke is a complex mixture of over 4,000 known chemicals (1, 36). We are currently attempting to fractionate smoke to separate possible complement activators from other “interfering” components. Future studies with isolated fractions of cigarette smoke may enable us to identify the chemical mechanism involved in smoke-induced complement activation.

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References


