The current epidemic of chronic obstructive pulmonary disease (COPD) has produced a worldwide health care burden, approaching that imposed by transmittable infectious diseases. COPD is a multidimensional disease, with varied intermediate and clinical phenotypes. This Review discusses the pathogenesis of COPD, with particular focus on emphysema, based on the concept that pulmonary injury involves stages of initiation (by exposure to cigarette smoke, pollutants, and infectious agents), progression, and consolidation. Tissue damage entails complex interactions among oxidative stress, inflammation, extracellular matrix proteolysis, and apoptotic and autophagic cell death. Lung damage by cigarette smoke ultimately leads to self-propagating processes, resulting in macromolecular and structural alterations — features similar to those seen in aging.

Tobacco-related diseases, including chronic obstructive pulmonary disease (COPD), account for 3.7% of the world burden of disability-adjusted life-years (DALYS), a measure of lost years of healthy life (1). Tobacco use, excessive alcohol consumption, and unhealthy diets and physical inactivity contribute to most preventable non-communicable diseases. These diseases are projected to impose a worldwide burden of $47 trillion health dollars by 2030. In contrast, it costs only $0.40 per individual per year to implement a program aimed at averting tobacco-related diseases that has the potential to save 25–30 million DALYS (1). Notwithstanding its preventable nature, the increasing prevalence, impact as the third leading cause death in the United States since 2008, and socioeconomic costs (1) call for vigorous research efforts to improve the understanding and, ultimately, management of COPD.

Under the umbrella definition of “decreased airflow that is not fully reversible” classically measured by the forced expiratory volume in one second (FEV1), COPD has a spectrum of clinical presentations, which affects accurate diagnostic phenotyping of patients as well as the design and validation of effective therapies (2). The typical clinical manifestations of the COPD syndrome include chronic bronchitis, a condition of large-airway inflammation and remodeling, and emphysema, a disease of the distal airways and lung parenchyma that manifests as loss of surface area for gas exchange. COPD decreases patients’ quality of life due to shortness of breath and chronic productive cough, which can progress over years to chronic hypoxemic and/or hypercarbic respiratory failure. Furthermore, systemic manifestations of COPD such as systemic inflammation, alterations of metabolism, cardiovascular events, and cancer contribute to the untimely death of these patients.

This Review emphasizes recent pathogenetic insights and emerging investigations into the complex and chronic nature of COPD (Table 1). These efforts have the added benefit of providing a window into lung biology, with a broader impact in the understanding of other non–tobacco-related pulmonary diseases.

Initiation
Tobacco smoke remains the key cause of COPD worldwide. Given that cigarette smoke contains thousands of injurious agents (3), its pathogenicity cannot be stringently studied one compound at a time. Aside from nicotine, heavy metals, and carcinogens, tobacco smoke leads to a significant exposure to oxidants. These include alkyl, alkoyl, and peroxy organic free radicals (causing lipid peroxidation), α,β-unsaturated aldehydes (such as acrolein and crotonaldehyde, which cause protein carbonylation and loss of sulfhydryls), and superoxide, N2O, and nitric oxide (which can generate peroxynitrite, leading to formation of dityrosine and/or 3-nitrotyrosine) (4).

While most studies have addressed medium- to long-term organonal responses to cigarette smoke, insights into immediate host responses to the inhalation of the toxic and oxidant components of smoke have been limited (Figure 1). Both in humans (5) and in rodents (6), tobacco smoke causes airway inflammatory responses within minutes or hours of exposure. One of the earliest manifestations is a breach in the vascular and airway barrier function (7), with brisk recruitment of circulating inflammatory cells to the lung (8). Indeed, oxidants present in the cigarette smoke trigger NF-κB–dependent inflammatory responses (9). The acute inflammatory response appears to be transient in nature and mediated by NF-κB, likely counteracted by regulatory networks that dampen NF-κB–dependent responses (10). Paradoxically, NF-κB may also participate in protection against cigarette smoke, as loss of function of the NF-κB p50 subunit augments cigarette smoke inflammatory responses (11).

Recent studies have implicated the host’s responses in the augmentation of lung injury by cigarette smoke. For example, cigarette smoke activates inducible nitric oxide synthase, leading to generation of oxidants, such as peroxynitrite (ONOO−), which has been linked to alveolar injury due to cigarette smoke (12). Another endogenous mediator of cell injury and inflammation co-opted in the early lung responses to cigarette smoke is the collagen degradation product proline-glycine-proline peptide (PGP), which engages CXCR2 receptors of neutrophils, therefore amplifying initial cigarette smoke-induced inflammation (13). In addition, the LPS in cigarettes may activate TLR4-expressing cells in the lung, leading to the activation of NF-κB responses (14). However, the...
Ddit4 (also known as DAPK3 or the Draper gene) is a stress-induced molecule (16) that might control how early lung responses control subsequent injury cells via activation of NADPH oxidase 3 (15). Increased oxidant generation in pulmonary capillary endothelial cells in the lung of apoptotic effector molecules induces transient apoptosis, and most importantly, against emphysema development (20). The master antioxidant transcription factor Nrf2 has been recently implicated in a broad range of lung responses involved in both the initiation and progression of lung injury due to cigarette smoke. Nrf2 controls more than 100 genes involved in antioxidant defenses, detoxification, and cellular physiology (19). Mice lacking Nrf2 show increased susceptibility to lung inflammation by acute cigarette smoke exposure, have upregulated RTP801 expression, and with chronic exposure also show increased susceptibility to alveolar cell apoptosis and development of emphysema (20).

**Progression**

The stage of progression of alveolar injury has attracted most of the research efforts in the COPD field (Figure 1). For more than 30 years, initiation and progression have been linked to extracellular matrix proteolysis, notably degradation of elastin by elastases, largely of inflammatory cell source. Key to this paradigm were the landmark discoveries of emphysema in α1-antitrypsin–deficient patients (21) and the induction of emphysema by intratracheal instillation of pancreatic elastase (22), as well as the finding that MMP-12–deficient mice are resistant to cigarette smoke–induced mouse emphysema (23). While extracellular matrix proteolysis is a central event in emphysema, it is apparent that it cannot explain the complexity of alveolar destruction in COPD.

A more complex picture of the mechanisms of alveolar destruction leading to emphysema has emerged in the past 12 years. Underlying the discovery of alternative molecular determinants and destructive processes was the important yet straightforward concept that the lung requires ongoing maintenance of its structures, notably during injury (24, 25). Variability in an individual’s ability to maintain lung structure and promote repair may explain the 20%–25% risk that smokers have of developing COPD as well as the reason that patients differ greatly in their clinical phenotypes and disease severity and progression. This concept has been verified in multiple models of emphysema (26) and, albeit at a descriptive level, supported by studies in lung samples from humans with COPD that demonstrate altered expression of multiple trophic/maintenance factors, including VEGF (27), Wnt signaling components (28), and adiponectin (29). This concept may also apply to the underlying mechanisms leading to the recently described disappearance of terminal airways in COPD (30).

We believe that mechanisms involved in the progression stage of COPD may be distinctly engaged in generating variable intermediate and clinically relevant disease phenotypes, such as emphysema, chronic airway disease (including chronic bronchitis and bronchiolitis), and systemic disease. Emphysema. In addition to enhanced lung elastolysis, the failure of the lung maintenance program in the parenchyma distal to the terminal bronchiole leads to a loss of alveolar cells by apoptosis in emphysema. The role of apoptosis was directly tested in mouse models of emphysema induced by a loss of VEGF function; in these models, caspase inhibitors preserved the integrity of alveolar septae (27, 31). Fueled by the identification of increased apoptotic cells in the parenchyma of human emphysema lungs (32), multiple mechanistic studies of the drivers and downstream consequences of this form of regulated cell death have emerged. Although it remains disputed which structural cell of the alveolus directs the process of alveolar destruction, it has been established that (a) apoptosis of both epithelial and endothelial cells occurs in models of emphysema (20, 27, 33), (b) direct instillation or overexpression in the lung of apoptotic effector molecules induces transient apoptosis of both epithelial and endothelial cells in the parenchyma of human emphysema lungs (32), multiple mechanistic studies of the drivers and downstream consequences of this form of regulated cell death have emerged. Although it remains disputed which structural cell of the alveolus directs the process of alveolar destruction, it has been established that (a) apoptosis of both epithelial and endothelial cells occurs in models of emphysema (20, 27, 33), (b) direct instillation or overexpression in the lung of apoptotic effector molecules induces transient

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**Table 1**

Levels of COPD complexity

<table>
<thead>
<tr>
<th>Level 1: Environment</th>
<th>Initiation</th>
<th>Progression</th>
<th>Consolidation</th>
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<tr>
<td>Infectious agents</td>
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<tr>
<td>Pollutants</td>
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<td>Tobacco</td>
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**Level 2: Genes**

- CHRNA3/S
- FAM13A
- HHIP
- SERPIN2
- XRCC5

**Level 3: Clinical phenotypes**

- Cancer
- Chronic bronchitis
- CVD
- Depression
- Emphysema
- Exacerbations
- Metabolic syndrome
- Osteoporosis
- Weight loss

**Level 4: Biomarkers**

- CRP
- Endothelial cell microparticles
- Exhaled condensate

**Level 5: Treatment**

- Bronchodilators
- Inhaled corticosteroids
- PDE4 inhibitors

Environmental etiologic factors, genetic basis, clinical phenotypes, biomarkers, and available treatments are listed. The pathogenetic stages of initiation, progression, and consolidation integrate these multiple levels, allowing for a comprehensive approach to understanding COPD. CRP, C-reactive protein. Data are from the American Journal of Respiratory and Critical Care Medicine (103).
airspace enlargement (33–36), and (c) specific induction of lung microvascular endothelial cell apoptosis is sufficient to cause a phenotype reminiscent of cigarette smoke–induced emphysema, including influx of inflammatory cells (37).

The induction of cell death in structural cells of the lung parenchyma (epithelial, endothelial, and possibly septal fibroblast cells) in response to cigarette smoke may be related to a loss of growth factors, oxidative stress injury, or intracellular response to stress imposed by noxious exposures (e.g., ER stress, ref. 38; or DNA damage, ref. 39). The ultimate outcome of these processes may differ depending on cell type; in contrast to small airway or alveolar epithelial or endothelial cells, alveolar macrophages are significantly more resistant to the apoptotic effect of direct cigarette smoke exposure or pro-apoptotic ceramides (40, 41), largely attributed to increased Akt signaling and increased ceramidase activity (42). Moreover, lung cells may respond to injury by activating pro-survival mechanisms such as autophagy, a response typically triggered by starvation. Lung epithelial cells (43), endothelial cells (44), fibroblasts (45), and alveolar macrophages (40, 45) all respond to cigarette smoke exposure by initiating autophagy signaling. Current evidence suggests that the abnormal persistence of such signaling or the inability to complete a physiological autophagic program may increase cellular stress such as ER stress, leading to caspase activation and apoptosis in diseased lungs (40, 44, 46).

Investigation into the interactions between upstream and downstream events related to alveolar cell apoptosis during emphysema onset and progression led to the key finding of self-amplifying injury loops involving apoptosis, oxidative stress, and inflammation (3–5). This concept may explain the progression of disease despite cessation of exposure to harmful initiators such as cigarette smoking (47). Paradigmatic of this interaction is the upregulation of pro-apoptotic sphingolipids in alveolar cells, including ceramides. Ceramides, which are induced directly by cigarette smoke or indirectly due to VEGF deprivation or oxidative stress, cause apoptosis of alveolar structural cells, self-amplify their own synthesis in a paracrine manner, increase oxidative stress, cause inflammation with activation of extracellular matrix proteases, and impair the clearance of apoptotic cells by lung alveolar macrophages (33, 35, 41, 48). Executioner caspases such as caspase-3 and elastases can in turn proteolytically activate endothelial monocyte-activating protein II (EMAPII), which has a dual action in the lung, causing endothelial cell caspase-dependent apoptosis as well as inflammation via CXCR3-dependent monocyte chemoattraction and activation (36). Additional positive interactions exist between extracellular matrix proteases, such as between cathepsin S and alveolar cell apoptosis (49), and between alveolar cell apoptosis and oxidative stress (50).

There is growing evidence that pulmonary and systemic inflammation, key events in COPD, may change in nature as the disease progresses. As mentioned above, inflammatory cell activation and influx may be operated by different mechanisms in the later phases of disease, when pathogen-associated molecular pattern-driven (PAMP-driven) processes (see below), abnormal apoptotic homeostasis (an apoptotic rate greater than the rate of clearance of dead cells by efferocytosis; ref. 51), and autoimmune responses become more prominently involved in inflammation (52, 53). Epigenetic dysregulation may contribute to excessive activation of proinflammatory cytokines and chemokines and steroid resistance (54, 55). As the lung maintenance program is eroded during chronic smoke exposure, inflammatory cells, including neutrophils, macrophages, and lymphocytes may change their phenotype. For instance, macrophages may switch toward an M2 phenotype, potentially limiting inflammation yet enhancing fibrosis and weakening their antibacterial functions (56). The concept of evolving inflammatory phenotype in the course of COPD is also supported by studies of lymphocyte involvement in the disease.

Figure 1
Pathogenetic factors organized based on their role in the initiation, progression, and consolidation of emphysema. (A) Initiation: Environmental agents trigger host cell responses, largely dominated by inflammation and oxidative stress. RTP801 is activated by cigarette smoke, largely due to oxidants, mediating inflammatory responses, oxidative stress, and alveolar cell death. DAMPs and PAMPs present in tobacco or generated endogenously may further enhance pathologic responses. Nrf2, by activating a host of antioxidant mediators, protects the lung and may promote lung repair processes. (B) Progression: Cigarette smoke disrupts alveolar maintenance, triggering apoptosis and autophagy; moreover, oxidants in tobacco and activated inflammatory and alveolar cells lead to extracellular matrix proteolysis, which further enhances inflammation and promote a feedback loop with apoptosis. Several of these interactions are facilitated by decreased expression of trophic/maintenance factors and endogenous mediators of alveolar destruction, including ceramide and EMAPII. (C) Consolidation: Over decades of exposure to cigarette smoke and endogenous amplifiers of destructive processes, there is progressive lung aging, with autoinflammatory stimuli generated through self-antigens or microbial/viral agents. TH17-positive cells, which are increased in COPD patients, may mediate the autoimmunity process. Macromolecular damage may lead to progressive telomere erosion and activation of p21CIP1/WAF1/SDI1 as part of the cell senescence response, which together may lead to a terminally injured lung.
Different populations of lymphocytes may aid in alveolar destruction; the presence of CD8 cells has been associated with disease severity (57), and oligoclonal CD4 cells have been detected in advanced COPD lungs (58), suggesting their participation in autoimmune responses in the consolidation stage of COPD. NK cells, which also accumulate in increased numbers in COPD lungs, may drive alveolar cell apoptosis via crosstalk with corresponding lung epithelial cell NKG2D ligands (59).

In addition, bacterial and viral infections, common causes of exacerbation in COPD patients and a major cause of pulmonary deterioration and mortality, may profoundly affect the nature of the immune response in the lung. Infectious agents, via PAMPs, may engage specific receptors, initiating host responses that lead to amplification of alveolar injury. Dysfunction of Nrf2 signaling in COPD patients (60) may compromise bacteria clearance (61). Influenza virus, a common cause of acute exacerbations, may synergize with cigarette smoke-induced responses via activation of the retinoic acid inducible gene–1 (RIG-1) helicase system. Indeed, double-stranded RNA viruses (e.g., influenza virus) activate innate signaling, culminating in the activation of interferon regulatory factor–3 and –7 (IRF-3 and -7) and the subsequent induction of antiviral genes, including NF-κB. This process was elegantly described by studies using the double-stranded RNA mimic, polyinosinic-polycytidylic acid (poly[I:C]) (62). Poly(I:C) cooperates with cigarette smoke to activate the RIG-1/MAWS/PRK/IL-18 pathway, causing excessive alveolar cell death, inflammation, and hastened and more extensive emphysema (62). These pathways may interact with endogenous danger-associated molecular patterns (DAMPs), such as high-mobility group box 1, to amplify early cigarette smoke–induced responses (63). Even elastin dissolution into chemotactic peptides during alveolar destruction further augments inflammatory responses independently of ongoing tobacco exposure (64).

Airway disease. Large and small (<2 mm diameter) airways are central sites of disease in COPD. While large airways undergo chronic inflammation with excessive mucus production, smaller airways are surrounded by inflammatory cells and wall fibrosis and exhibit intraluminal mucus accumulation (65). The discovery that cigarette smoke directly inhibits the CFTR function (66) provides a mechanistic link between smoking and abnormal mucous secretion by airway epithelial cells and suggests shared pathogenetic pathways between COPD and the chronic airway disease cystic fibrosis. Similar to another common chronic airway disease, asthma, COPD airways exhibit increased VEGF and IL-13 levels, which contribute to mucous gland hyperplasia and bronchial smooth muscle hyperreactivity (67). It is interesting to note that the chronic bronchitis phenotype of COPD in humans is characterized by increased VEGF in the large airways, while in emphysema there is a paucity of VEGF in the lung parenchyma (6–8). Moreover, the small airways in COPD appear to exhibit unique remodeling mechanisms. Indeed, inhibition of EMAPII attenuates small airway but not large airway remodeling in mice chronically exposed to cigarette smoke (36). Furthermore, both TGF-β and IL-1β were proposed as important mediators of airway remodeling in COPD explants (68). Recent studies revealed that small airways may disappear even earlier than surrounding alveoli (30), providing a common phenotype with that resulting from destructive processes occurring in emphysema. The mechanistic role of TGF-β initially suggested in ex vivo models (68) has been recently supported by the beneficial effects of anti-TGF-β antibodies or TGF-β inhibition with the angiotensin receptor inhibitor losartan on both airway remodeling and airspace enlargement due to cigarette smoke (69). Similarly, both molecular and pharmacological inhibition of IL-1β signaling attenuated airway remodeling and extracellular hyaluronan levels during cigarette smoke exposure (70).

Lung cancer. There is evidence of both active cell proliferation and apoptosis in COPD lungs, and the balance of cell proliferation to death may be a measure of (inadequate) repair in emphysema (71). However, a proliferative response in airway epithelial cells exposed to cigarette smoke (72) may be unwanted. When combined with the cancer-initiation properties of tobacco, these events may underlie the cancer-promoting effects of cigarette smoke–induced inflammation (73). Accumulation of macromolecular damage in the course of COPD might also explain the increasingly recognized association between lung cancer and emphysema (74).

Systemic manifestations. The lung-specific pathogenetic processes outlined above may apply to systemic organ dysfunction, as the lung allows passage of oxidants into the bloodstream (75). Right ventricular dysfunction due to pulmonary hypertension may occur due to oxidative and nitrosative stress, with reduced expression of vasodilators in COPD (12, 76–78). Prolonged exposures to cigarette smoke have an inhibitory effect on the bone marrow hematopoietic progenitor cell number and cycling function (79, 80), adversely impacting lung repair mechanisms in COPD. Finally, skeletal muscle wasting and decreased physical activity, which are major comorbidities in COPD, have been linked to both increased apoptosis and decreased vascular regeneration (81, 82).

Consolidation COPD may progress in patients despite smoking cessation, which challenges the concept of a direct link between ongoing exposure to cigarette smoke and the disease. This progression parallels persisting inflammatory responses (83), suggesting that additional mechanisms must account for the consolidation of the disease in genetically susceptible hosts, often after decades of active smoking (Figure 1). This change in the nature of inflammation in the course of disease is highlighted by the temporary nature of NF-κB activation in rodent lungs exposed to cigarette smoke (84). Two paradigms have emerged that might explain some of these observations: autoimmunity (52) and lung aging (85). These findings may derive from the profound lung alveolar damage (39) and airway remodeling imposed by chronic cigarette smoke exposure and bombardment by endogenous mediators of inflammation and cell injury. Understanding the mechanism of this persistence might have a far-reaching impact on the design and implementation of regenerative therapies.

It is conceivable that COPD becomes over time an autoinflammatory disease, possibly involving the inflammasome and related cytokines. Cigarette smoke activates the inflammasome in mouse lungs, involving ATP and its receptor P2X7 purinergic receptor (86). NLPR3 activation of caspase-1 (87) would lead to increased IL-1β (70) and IL-18 (62), both of which have been shown to participate in experimental models of cigarette smoke–induced inflammation and alveolar injury. Although DAMPs, including purines, can act early, in the initiation stage of COPD, the pathogenic roles of IL-18/IL-18 receptor signaling suggest that this inflammatory signaling occurs downstream of apoptosis, positioning it in the consolidation stage of alveolar injury (62).
There is growing evidence for a role of autoreactive T cells or auto-antibodies in the activation of specific immunity in COPD. Mice exposed to cigarette smoke exhibit lung infiltration by CD8 T cell oligoclonals (88), a finding that correlates with observations of oligoclonals of CD4 cells in lungs of patients with COPD (58). Auto-antibodies have been detected in lung tissues of patients with advanced disease (89), including antibodies targeted to immunogenic carbonylated proteins (90). The latter findings were duplicated in mice exposed to cigarette smoke (91). Furthermore, rats immunized with endothelial cells developed autoreactive antibodies and T cells, which mediated alveolar enlargement (92). Finally, elastin-reactive T cells have been detected in the peripheral blood of patients with established COPD (93). Although the precise nature of the auto-reactive antigens will require further study, there is evidence of profound alterations in the structure of the lung during the course of exposure to cigarette smoke, ultimately leading to replacement of the alveolar elastin framework by collagen (94). A shared theme with other autoimmune processes is the emergence of TH17 lymphocytes, which are found in increased numbers in COPD lungs (95). Their potential role in alveolar destruction was recently uncovered by the protection against inflammation and emphysema observed in IL-17RA–knockout mice (96).

The relentless lung injury due to oxidant exposure, along with the potential exhaustion of lung protective responses, ultimately leads to lung aging, with increased expression of markers of cellular senescence (97). Again, a common denominator for aging and cellular senescence is oxidative stress, resulting in macromolecular damage, including the increased expression of markers of DNA damage (39, 98) and adduct-modified proteins (99). The ultimate “biological clock” of cell turnover is controlled by telomerase, which preserves the shortening of ends of chromosomes during every round of mitosis. Lungs of patients with advanced emphysema have decreased telomere lengths in alveolar cells (97), which are paralleled by decreased telomere lengths in peripheral blood mononuclear cells (100, 101). The contribution of shortened telomeres was recently unraveled in investigations with the telomerase reverse transcriptase–knockout mice, which showed increased sensitivity to alveolar injury and airspace enlargement due to cigarette smoke, notably in late intercreses compared with early intercreses and wild-type mice (102).

The future
Key questions remain in our understanding of COPD, and these concern clinical phenomena, systemic manifestations of COPD, and the impact of exacerbations triggered by infections. These central clinical manifestations of COPD probably result from the interaction of disease-related genes with fundamental processes involving inflammation, thrombosis and hemorrhage, fibrosis, the immune response, proliferation, and apoptosis/necrosis, which underlie the so-called intermediate phenotypes (103). Development of model systems that can address mechanistically these interactions will remain vital for progress in COPD, validated by studies of the human disease. Some therapeutic strategies, such as restoring G1-antitrypsin activity or the use of Nrf2-dependent antioxidants, may directly antagonize destructive processes such as the activation of pro-apoptotic mediators and extracellular matrix proteolysis (36, 104); a significant challenge lies in restoring lung survival mechanisms without fueling oncogenesis. Recently described shortcomings in attempts to regenerate the lung in a murine model of COPD (105) remind us that approaches aimed at lung organ restoration (79) will require consideration of the extent of macromolecular damage imposed by decades of lung destruction (39, 98). The aggregate of these insights into the pathogenesis of COPD provide landmarks that should direct future investigations in COPD and targets for potential novel therapies.

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