Animal models of asthma

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Submitted 23 January 2009; accepted in final form 19 June 2009

Bates JH, Rincon M, Irvin CG. Animal models of asthma. Am J Physiol Lung Cell Mol Physiol 297: L401–L410, 2009. First published June 26, 2009; doi:10.1152/ajplung.00027.2009.—Studies in animal models form the basis for much of our current understanding of the pathophysiology of asthma, and are central to the preclinical development of drug therapies. No animal model completely recapitulates all features of the human disease, however. Research has focused primarily on ways to generate allergic inflammation by sensitizing and challenging animals with a variety of foreign proteins, leading to an increased understanding of the immunological factors that mediate the inflammatory response and its physiological expression in the form of airways hyperresponsiveness. Animal models of exaggerated airway narrowing are also lending support to the notion that asthma may represent an abnormality of the airway smooth muscle. The mouse is now the species of choice for asthma research involving animals. This presents practical challenges for physiological study because the mouse is so small, but modern imaging methodologies, coupled with the forced oscillation technique for measuring lung mechanics, have allowed the asthma phenotype in mice to be precisely characterized.

Allergic inflammation; mouse; airway smooth muscle; lung impedance

Asthma is a prime example of a “complex disease,” a term that has become popular for characterizing those pathologies that appear to have multifaceted etiologies and an entanglement of underlying mechanisms. The moniker of complex disease, however, often tends to belie a general lack of understanding about what is going on, and asthma is no exception. Indeed, asthma is more properly regarded as a syndrome than a disease because it is defined on the basis of clinical characteristics rather than underlying mechanisms, and there remains a great deal of controversy about which of the possible mechanisms are the most important (20). The principal characteristics of asthma are reversible airflow obstruction, hyperresponsiveness of the lungs to challenge with smooth muscle agonists, and airway inflammation. Not surprisingly, allergy heads the list of causative suspects, but it is by no means alone. Exercise, cold air, and emotional stress are also known triggers of asthma (1), leading to the notion that asthma itself represents merely the common clinical endpoint of a number of distinct disease processes (42, 53, 91).

As with most human diseases, studies in laboratory animals have produced much of what we currently think we know about the mechanisms responsible for asthma. Obviously, the relevance and validity of these studies are tied to how well we can produce accurate animal equivalents of human asthma. The development of such “animal models” is still very much a work in progress; although many of the various features of asthma have been convincingly recapitulated in animals, invariably every animal model misses some important aspect of the human syndrome (97). Also, very few animals spontaneously develop a condition with any similarity to asthma, the most reminiscent being an allergic syndrome in cats (72) and the condition known as heaves in horses (24). If asthma is a condition peculiar to humans for some reason, then having it manifest in its entirety in a laboratory animal may simply be impossible. On the other hand, given that we still do not fully understand what asthma in humans actually is, it remains difficult to know whether an animal really has it or not. Accordingly, much of the challenge in studying animal models of asthma lies in phenotyping them properly, particularly as asthma is defined in humans in terms of phenotype rather than underlying pathology. In this review, we therefore focus on the issue of how to assess the relevant function and structure in the lungs of laboratory animals, and what clues this has given us about the possible mechanistic underpinnings of asthma in humans.

Development of Animal Models

Early animal models of asthma were developed in a variety of species (90) and focused on the phenomenon of airways hyperresponsiveness (AHR), defined as excessive bronchoconstriction in response to a standardized challenge. The challenging agent was usually a smooth muscle agonist such as methacholine or histamine. These agonists were intended to mimic the actions of mediators released as part of an overexuberant immune response thought to be behind an attack of allergic asthma. In a similar vein, some animal models employed hyperpnea to mimic the asthma brought on by exercise (35). In any case, the motivation behind these models arose from the idea that asthma is primarily a matter of excessive shortening of the airway smooth muscle (98). This notion was also...
reflected in the predominant use of β-agonists to relax smooth muscle as frontline therapy for most asthma cases.

In the early 1980s, a greater awareness of the role of inflammation in asthma developed, driven by an increased understanding of allergic immunology together with observations that human asthmatics frequently exhibited marked symptoms when challenged with antigens of various kinds. The first inflammatory cell to be linked firmly to asthma pathogenesis was the eosinophil (27, 63), followed soon after by the T cell (66). This led to increasing use of corticosteroids in the routine treatment of asthma. Current asthma guidelines (1) call for combining corticosteroids with long-acting β-agonists in treating persistent asthma, which seems appropriate given its current definition comprising both obstructive and inflammatory components. In fact, it now seems that treating asthma with a β-agonist alone may be inadvisable. Indeed, monotherapy with some long-acting forms has recently been designated as unacceptable by the Food and Drug Administration.

As a result of studies with animal models, we now know that allergic asthma involves a complex interplay between the innate and adaptive immune systems. The involvement of adaptive immunity begins with naïve CD4 T cells that differentiate into T helper (Th) cells with the potential to regulate the fate, function, and location of a variety of other immune cell components. Different subsets of Th cells have been defined on the basis of the cytokines they secrete, with Th1 and Th2 cells being the best known examples. The production of Th1 cells is promoted primarily by IL-12, whereas differentiation into Th2 cells occurs in the presence of IL-4. In turn, Th1 cells produce IFNγ, whereas Th2 cells produce IL-4, IL-5, and IL-13. Allergic asthma is associated with a Th2 type of immune response, since the Th2 cytokines IL-4 and IL-13 are known to cause many of the features of the disease (36, 52). Th2 cytokines, particularly IL-5 and to certain extent IL-4, are also responsible for linking adaptive to innate immunity by promoting eosinophil proliferation in bone marrow and subsequent migration to the lung. Eosinophilia in the airways has long been linked to allergic asthma, although exactly why has been controversial. Nevertheless, their pivotal role in altering lung function in allergic mice has been demonstrated in experiments with eosinophil knockout animals (58).

The Th2 cytokine IL-4 is also implicated in the dominance of antigen-specific IgE over other antibody isotypes in allergic asthma. Although a number of cytokines such as IL-21 are associated with V8, V14 J7, and V2 (56). After the immune system has had a chance to mount a reaction against the antigenic protein, which takes several days, progressively low numbers, NKT cells have the capacity to rapidly produce high levels of various cytokines upon antigen recognition (16) and are found in the lung. Transfer of NKT cells from mice with allergic airway inflammation to naïve mice is sufficient to induce airway hyperreactivity (3).

Another CD4 subset that has emerged within the last two years is the Th17 cell. Both TGFβ and IL-6 are necessary, although not sufficient, for the differentiation of Th17 cells (17, 62, 86). These cells produce mainly IL-17, some IL-22 and IL-21, small amounts of IFNγ, and no IL-4 (46, 69). Although IL-17 is clearly detectable in allergically inflamed lungs, its role in the pathogenesis of the allergic response remains unclear. It seems that IL-17 may be needed for the initial development of airway inflammation in mice, but administration of IL-17 decreases eosinophilia. Also, adoptive transfer of Th17 cells induces neutrophilia and imparts resistance to steroid therapy, suggesting that IL-17 may function to promote neutrophil recruitment (55) and could therefore be particularly important in severe asthma (64). IL-17 has also been shown to interfere with epithelial cell production of eotaxin (77), which plays a major role in the recruitment and activation of eosinophils (70). IL-17 may thus regulate the eosinophil-neutrophil balance in the lung.

Despite the wealth of knowledge we now have about the role of immunity in allergic asthma, the complex pathophysiology of the disease caused researchers to have a difficult time agreeing over whether asthma is due primarily to altered immune status or to an abnormality of airway smooth muscle, or even both simultaneously (5). Work has thus proceeded along both fronts, driven by the respective convictions of the scientists involved. The development of animal models of asthma has been correspondingly schizophrenic. Nevertheless, the asthma world has come together on one thing regarding animal models: the species. As with most animal-based biomedical research, mice now dominate the scene because of the immunological and molecular tools available to study them, together with their obvious practical advantages related to cost and gestation period (23, 50). This is not to say, of course, that the mouse is the only species used to model asthma. Rat models have also been widely employed (21, 84), notably the Brown-Norway strain because of its propensity to exhibit a later allergic response following antigen challenge (84). Monkeys have also attracted attention as a useful model that bridges the gap to human relevance more than rodent species (28, 39, 61). Nevertheless, most asthma-related research continues to be pursued in the mouse. The development of transgenic mice that exhibit various lung pathologies is now a huge research enterprise, and mouse models of lung disease have been the subject of a number of recent reviews that cover their various pathophysiological features in detail (11, 23, 49, 50, 54, 67).

Mouse Models of Allergic Asthma

Allergic mouse models of asthma are generated by first sensitizing an animal to a foreign protein, most commonly ovalbumin. This is typically done by injecting the protein intraperitoneally along with an adjuvant, typically aluminum hydroxide, that serves to enhance the protein’s immunogenicity for reasons that are complex and not entirely understood (56). After the immune system has had a chance to mount a reaction against the antigenic protein, which takes several days,
the animal receives a further antigen exposure either directly to the lungs in the form of an aerosol or via postnasal drip following nasal instillation. This elicits an inflammatory reaction in the lungs characterized by an influx of eosinophils, epithelial thickening, and AHR. There is wide variation, however, in the precise recipe used by different investigators to perform these various steps. A typical scenario is to sensitize an animal with two intraperitoneal injections spaced a week or two apart, wait another week, and then challenge the animal by exposure to a 1% ovalbumin aerosol for 30 min each day for 3 days (87). The features of allergic inflammation typically reach their peak a day or two after the final exposure, although the time course of this process has not been fully characterized.

Many variations on all aspects of the above theme have been tried. For example, inhalational exposure to NO2 has been used as a means of bolstering the immune response in place of an intraperitoneal adjuvant (18), invoking the notion that airborne pollutants may play a role in the development of asthma in industrialized societies. Also, questions about the relevance of ovalbumin to human asthma have led to the use of antigens such as extracts of house dust mite (30) and aspergillus (40), which are more representative of those that occur naturally. Of particular recent interest has been the issue of chronicity of antigen exposure. The sensitization and challenge procedure described above is very acute, yet asthma in humans is typically a chronic disease that exhibits clear features of airway remodeling in its later stages (51). Researchers have thus been motivated to explore the effects of longer-term antigen exposures in mice. Somewhat disappointingly, extended exposure to ovalbumin aerosol in at least some mouse strains such as the BALB/c leads to tolerization, characterized by the waning of the inflammatory and hyperresponsive phenotypes (9, 94). There have been some reports that other antigens such as house dust mites produce a more extended asthma-like picture (30). Furthermore, simultaneous exposure to multiple antigens produces inflammation in the absence of adjuvant, and may even break through the tolerization barrier to produce a chronic phenotype (38). Nevertheless, the chronic model has yet to be optimized, particularly in view of the rather modest degree of AHR that has so far been demonstrated. Indeed, the generation of allergic mouse models of asthma remains something of a black art, no doubt due to the immensely complex and often poorly understood biological processes involved.

Fully elucidating the biological complexities of allergic mouse models of asthma is going to take some time, but it has moved ahead in recent years thanks to significant technical advances in phenotyping using both lung function assessment and imaging. The measurement of lung mechanical function in mice poses a particular challenge, of course, due to the small size of the animal (50). Measurement techniques developed in larger animals or humans frequently do not do well when scaled down to the mouse. The frustration this causes has led to a temptation to cut corners in the assessment of the mouse phenotype, the most problematic result of which has been the use of the parameter known as enhanced pause (Penh) to assess AHR (43). Due to its attractions of noninvasiveness and simplicity, as well as its availability in a commercial device (31, 32), Penh has become widely used in the study of AHR. Unfortunately, however, this use has invariably been inappropriate. Penh is merely a nonspecific feature of the breathing pattern and cannot be taken as a measure of lung mechanics, as has been discussed extensively elsewhere (7, 65). Of course, because the mechanical properties of the lung have an important bearing on the pattern of breathing (43), it is conceivable that a change in the breathing pattern, such as might be reflected in Penh, could signal the presence of some event worthy of more detailed investigation, such as activation of irritant receptors in the lung (2). On the other hand, one can just as easily screen for a change in breathing pattern using the obvious parameters of breathing frequency and/or tidal volume that seem to be just as efficacious as Penh (2) but do not suffer from the latter’s obscurity of meaning. Indeed, we would go so far as to suggest that the use of Penh be avoided completely (7). In any case, the pattern of breathing can never serve as a substitute for a measure of mechanical function.

Recently, two approaches have been used to extend unrestrained plethysmography by combining it with an independent noninvasive measure of changes in lung volume to estimate specific airway resistance (14, 76). These approaches are theoretically sound and may eventually prove useful for rapid screening, but the information they provide is noisy and subject to the vagaries of the animal’s spontaneous breathing pattern. The problem underlying these approaches has been ascribed to the “phenotyping uncertainty principle” (11), which states that there is a fundamental trade-off to be made between the maintenance of natural measurement conditions and precision. Penh sits at one extreme of this trade-off; a conscious animal that is free to choose its own pattern of activity and breathing exists in its most natural state, but the measurements of lung mechanics one can obtain under these conditions are noisy and very nonspecific.

At the other extreme of the phenotyping uncertainty principle lies the measurement of lung mechanics using the forced oscillation technique in anesthetized, tracheostomized, paralyzed mice (11). Here the animal’s breathing pattern is entirely under the control of the experimenter, and the upper airways are bypassed so that their mechanical properties do not interfere with those of interest (i.e., the lung). On the other hand, one is left with the conundrum that the extensive interventions employed to achieve this level of experimental control may have caused the mechanical properties of the lung to depart significantly from their natural state. Nevertheless, our understanding of AHR in mice has advanced considerably in recent years using this approach. Intermediate between the above two extremes of the phenotyping uncertainty principle lie methods, such as the measurement of transfer impedance in restrained animals (47), that are based soundly in physical theory, but which are subject to greater variation as a result of fewer constraints on the behavior of the animal.

Assessing Lung Mechanics in Mice

The assessment of lung mechanics in mice, or any experimental animal for that matter, boils down, in essence, to determining how difficult (or easy) it is to drive a given volume of gas into the lungs over a given period of time. At its most basic level, this process can amount simply to examining the peak airway pressures achieved during regular mechanical ventilation (11, 29, 50). Clearly, an increased peak pressure indicates an increased impedance to lung inflation, although in an entirely nonspecific manner. Much greater specificity is
achieved with the forced oscillation technique, in which an appropriate oscillatory flow signal is applied to the airway opening while the airway pressure is measured. In fact, flow signals containing a single dominant frequency determined by the number of breaths per minute have been used for decades to obtain estimates of lung resistance and elastance (6, 74). Numerical techniques based on the Fourier transform (13) have been used more recently to determine resistance and elastance at a number of different frequencies simultaneously when the oscillatory flow signal contains components at those frequencies. This multifrequency information about the respiratory system is encapsulated in a complex function of frequency known as the input impedance \( \text{Zin} \) (74). Being a complex function, \( \text{Zin} \) actually consists of two independent components known as the real part and the imaginary part, themselves both functions of frequency. The real part is known as the resistance because, at any particular frequency, it is equal in value to that of the conventional resistance that would be measured by oscillating the system at only that frequency. The imaginary part of \( \text{Zin} \) is known as the reactance, and is equal to conventional elastance scaled by the inverse of frequency. Obtaining \( \text{Zin} \) in a mouse is technically challenging because of the animal’s small size, but is now routinely achieved using either a commercially available device known as the Flexivent that is based around a computer-controlled piston oscillator (78) and a wavetube coupled to a loudspeaker (44).

Interpreting \( \text{Zin} \) is performed on the basis of a mathematical model of the lung with a structure that is interpretable in physiological terms. The appropriate model depends on the frequency range over which \( \text{Zin} \) is measured. When frequency is kept below \( \approx 20 \) Hz in mice, the model that has come to dominate the advanced investigation of lung mechanics is the so-called constant phase model introduced by Hantos and colleagues (45). This model has a particularly simple structure consisting of a single viscoelastic lung compartment served by a single airway. The airway has a resistance \( \text{Raw} \) that reflects its caliber and length. The mass of the air in the airway also provides the model with an inerterance, although this is so small in mice that it can be neglected below the 20-Hz maximum frequency typically used to determine \( \text{Zin} \) (44, 75, 87). The viscoelastic compartment is characterized by two parameters, \( G \) and \( H \), that account for the resistance and elastic properties, respectively, of the lung tissues. It has been found empirically that the impedance of the viscoelastic compartment has a frequency function, \( \text{Zin} \), that encapsulate the mechanical properties of the lung periphery. When the lung behaves in a mechanically homogeneous fashion, \( G \) and \( H \) reflect the intrinsic dissipative and elastic properties, respectively, of the tissues. Experimental studies have shown, however, that \( G \) invariably increases proportionately more than \( H \) when the lung becomes mechanically heterogeneous in such a way that ventilation is not apportioned to different regions of the lung according to the volume of each region, but rather becomes differentially affected due to inequalities in regional airway resistance and tissue stiffness. This finding is supported by theoretical analysis (8) and computational modeling (60, 82). An increase in the ratio \( GH \), known as hysteresivity (34), thus serves as a convenient marker for the presence of regional heterogeneities, which typically accompany most lung diseases and are also brought on transiently by induced bronchoconstriction (87). By contrast, if a fraction of the lung becomes isolated from the airway opening either by closure of the subtending airways or by atelectasis, collectively referred to as derecruitment, estimates of \( G \) and \( H \) will increase in the same proportion, leaving hysteresivity unchanged (4).

The parameters \( R_N \), \( G \), and \( H \) of the constant phase model together thus provide a means for distinguishing changes taking place in the conducting airways vs. changes in the lung periphery, and of inferring the presence of ventilation heterogeneity vs. derecruitment.

Figure 1 shows an example of the use of \( R_N \), \( G \), and \( H \) to infer details about the nature of AHR in mice. Allergic inflammation was induced in BALB/c mice using the standard 3-day challenge protocol described above. Groups of inflamed and control animals were then exposed for 40 s to an aerosol of methacholine and then switched immediately to a 5-min protocol during which \( \text{Zin} \) was measured approximately every 15 s. All three impedance parameters increased following cessation of the challenge, with the responses from the inflamed animals being substantially greater than those in the controls. The biggest difference between inflamed and control animals, however, occurred in \( H \), the parameter that measures lung elastance, indicating that the major effect of inflammation was manifest in the lung periphery. What is more, the ratio \( GH \) remained largely unchanged throughout, implying that the peripheral effect that occurred was due to derecruitment of lung units. Closure of small airways was subsequently confirmed directly using microcomputed tomography (59) on the basis of images such as those shown in Fig. 2, which also serve to illustrate that the airway tree of the mouse is structurally rather different to that of the human.

**Linking Structure to Function**

Despite the technical advance represented by the forced oscillation technique over earlier methods that yielded only single values for lung resistance and elastance, the parameters of the constant phase model still constitute an extremely simplistic view of a very complicated organ. Using only measurements of pressure and flow at the airway opening, it is not currently possible to reconstruct a more detailed picture of the lung than that contained in \( R_N \), \( G \), and \( H \). However, one can often use computer simulation to address questions about what might be causing experimentally observed changes in these parameters. Here one tries to construct as detailed a model of
the lung as anatomical and physiological knowledge will allow. The model is then used to simulate measurements of airway pressure and flow under conditions that match those used experimentally. The model thus serves as a virtual laboratory in which the feasibility of specific hypotheses can be addressed, providing in silico experimentation to complement conventional in vivo and in vitro experiments.

An example of this approach has been employed to investigate what might be responsible for the AHR observed in acutely inflamed BALB/c mice (87). A computational model of the mouse lung was constructed in which every airway in the entire airway tree was individually accounted for, with tissue units appended to each terminal bronchiole. The model was calibrated by adjusting its parameters until it exhibited changes in $R_N$, $G$, and $H$ similar to those seen in normal mice exposed to an aerosol of methacholine. To achieve this, however, it was not enough simply to simulate the effects of smooth muscle shortening by narrowing the model airways. Derecruitment of some fraction of the model lung was also required to mimic the increases in lung elastance observed experimentally. This was achieved by completely closing any airway in the model that narrowed to a specified critical radius, thereby mimicking the effects of formation of a liquid bridge across the airway lumen. In silico experimentation was then performed with the model in an attempt to make it hyperresponsive to methacholine in a way that matched experimental observations in allergically inflamed mice. Simply increasing the extent of smooth muscle shortening reproduced the observed changes in $R_N$ but failed to match the changes in $G$ and $H$. However, if the degree of shortening was kept the same as in control mice, but the epithelial layer in the model was thickened to match histological observations in the inflamed animals and the critical airway closure radius was also increased somewhat, the model was able to accurately mimic the behavior of all three impedance parameters (15). This suggests that the mechanism giving rise to AHR in the acutely inflamed BALB/c mouse does not involve an abnormality of the airway smooth muscle, but instead is linked to physical changes taking place in the periphery of the lung itself. Of course, results such as these are dependent on the validity of the model used to make the simulations, and no model is a perfect representation of reality. However, the model used in this case was based reasonably closely on the anatomy of the mouse lung, so the results of such simulations can lend strong support to one hypothesis over another.

The use of computational modeling can also raise important questions for future investigation. For example, geometric amplification of airway luminal narrowing can potentially occur either by epithelial thickening or by increased mucus secretion. Certainly, epithelial hypertrophy is a characteristic feature of allergic inflammation (87), but so is mucus hypersecretion. Mucus is produced by goblet cells in the airway epithelium, and the appearance of excessive mucus in the airways can lead to significantly impaired lung function. Furthermore, IL-13, one of the Th2 cytokines, is the major inducer of epithelial cell mucus (93), signaling through the IL-13/4 receptor-α complex (92) and epidermal growth factor receptor (96). We do not know which of these two factors, epithelial thickening or excessive mucus, is dominant in allergically inflamed mice. Experimenting with mice in which these two effects are individually manipulated would thus be an important area for future research.

Mouse Models of Excessive Airway Smooth Muscle Shortening

Although allergy and inflammation are widely recognized as central to the pathogenesis of asthma, there is a school of thought that places the principal blame for asthma squarely at the feet of the airway smooth muscle. Although the jury is still out on this question, considerable effort has been devoted to identifying animal models in which AHR results from excessive smooth muscle shortening. Related to this, some degree of airway smooth muscle hypertrophy has been identified in several chronic models of allergic inflammation (57, 79). The epithelium probably also plays an important role in ASM hyperplasia and subsequent AHR; selectively activating the nuclear transcription factor NF-κB in the airway epithelium results in airway smooth muscle hyperplasia and associated AHR (73). There is also evidence that the alterations in smooth muscle associated with inflammatory remodeling result in an increase in the speed of muscle contraction (80). It is not entirely clear why this should be associated with AHR and

Fig. 1. The parameters of respiratory input impedance ($R_N$, $G$, and $H$) expressed as fractional changes above baseline ($\Delta R_N$, $\Delta G$, and $\Delta H$, respectively) in control and allergically inflamed mice following a 40-s challenge with an aerosol of methacholine. MCh, the time of completion of delivery of methacholine. DI, the time of delivery of 2 deep lung inflations to 25 cmH2O. [From Wagers et al. (87).]
asthma, although there is speculation that it may allow the smooth muscle to more easily move into a latch state from which escape is difficult (33).

Enhanced shortening of airway smooth muscle has been induced by treating animals with cationic proteins (Fig. 3). Mice receiving an intratracheal instillation of poly-L-lysine were found to be hypersensitive to methacholine when it was delivered as an aerosol but not as an intravenous injection (48), implicating degradation of the normal barrier function of the epithelium as the causative factor. In other words, by making the underlying smooth muscle more accessible to agonists introduced into the airways, bronchoconstriction was enhanced both in terms of magnitude and speed of onset. Poly-L-lysine is an analog of the highly charged cationic proteins found in all inflammatory cells, particularly the eosinophil, suggesting a way in which the events accompanying allergic inflammation might lead to hypersensitivity and AHR (85).

Differences in the intrinsic contractility of airway smooth muscle also arise spontaneously between different strains of animal. The A/J mouse, for example, has been shown to be hyperresponsive, and, in particular, to exhibit bronchoconstriction that develops particularly rapidly following challenge (26, 88). There are also clear differences in natural airways responsiveness between different strains of rat, the Fischer and Brown-Norway strains being key examples (37, 89). Accord-ingly, we now recognize that genetic factors play a major role in determining AHR.

The Role of Lung Volume

The degree of airway smooth muscle shortening elicited by challenge with a standard dose of agonist is highly dependent on transpulmonary pressure. Most of the pulmonary airways are embedded in the lung parenchyma, which transmits transpulmonary pressure from the airway lumen to the pleural surface. The parenchymal attachments to the airway wall exert an outward tethering force that opposes shortening of the airway smooth muscle. The mitigating effect of these parenchymal forces is dramatically demonstrated by the way in which airways responsiveness is decreased by an increase in transpulmonary pressure of only a few cmH₂O (10, 22). Thus, a normoresponsive animal can be rendered very asthma-like simply by reducing its lung volume, just as has been demonstrated to be the case in human subjects (25). There are also some animal studies showing that chronic alterations in lung volume can lead to persistent changes in airways responsiveness, suggesting the possibility that airway smooth muscle somehow eventually adapts to a new lung volume (81). Of course, the forces of parenchymal tethering are not the only loads opposing airway smooth muscle shortening; the stiffness of the airway wall itself has also been shown to play a significant role in this regard (12, 68). In addition, the airway remodeling known to accompany some inflammatory conditions presumably has the potential to alter wall stiffness, but whether this tends to work for or against AHR remains controversial. It has recently been shown, for example, that there is no evidence of an effect on AHR due to mechanical changes in the airway wall of the acutely inflamed mouse (22). On the other hand, more chronic allergen exposures have shown evidence of an airway remodeling effect on AHR (79).

It must be mentioned that a great deal of animal work related to the hyperresponsiveness or otherwise of airway smooth muscle takes place in vitro, usually with tracheal smooth muscle, which is easily isolated and subjected to controlled force-length studies. The precision of the measurements that can be made with this preparation has led to the discovery of many intriguing phenomena such as the ability of smooth muscle to adapt its force generating capacity to baseline length (95). The biochemistry associated with the different phosphorylation states of the smooth muscle crossbridge is also rather complicated (41). It remains to be seen how significant some of these phenomena really are for asthma at the level of the entire animal or human subject. Nevertheless, computational modeling of the dynamic behavior of activated airway smooth muscle and its mechanical integration into the airway wall has shown promise in accounting for bronchoconstriction in vivo (10, 12, 22).
however, is often not considered. In fact, the degree of AHR is generally rather modest in most animal models considered to exhibit the phenomenon; a response in airway resistance of only two- to threefold above normal is not unusual (87, 88). In one sense, this may be apropos, given that much of human asthma also rumbles along chronically in a fashion that is perhaps more irritating than outright dangerous. On the other hand, in the subset of individuals who are afflicted with the severe form of the syndrome, the decrement in lung function accompanying an asthma attack can be extreme and even fatal. What distinguishes severe asthma from its more common milder form remains an area of active research that has so far yielded few answers. It is therefore perhaps not surprising that animal models of correspondingly severe AHR are also few in number. One reason for this may be the tendency for researchers to focus on single causes, in pursuit of the most important mechanism responsible for the AHR of asthma. It has recently been suggested, however, that the key to severe asthma may be the simultaneous presence of multiple mechanisms; allergically inflamed mice treated with poly-L-lysine were shown to exhibit an extreme level of AHR that was easily fatal and that could be attributed to the synergistic interaction between an enhanced ability of smooth muscle to narrow the conducting airways and an increased propensity for liquid bridges to occlude peripheral airways (9). The mechanism behind this synergy is illustrated in Fig. 4. A severe asthma-like phenotype can also be achieved by coexpressing IL-5 and eotaxin 2 in the same animal; IL-5 leads to systemic eosinophilia, whereas selective eotaxin 2 upregulation in the airway epithelium directs chemotaxis and markedly activates the diapedesing eosinophils (70). The physiological manifestations are nearly identical to the antigen/cationic protein model (9) supporting the multiple mechanism hypothesis for the genesis of extreme AHR.

Summary and Conclusions

Asthma still appears to be a uniquely human disease despite decades of attempts to recapitulate its features in animals. Nevertheless, animal models of AHR have provided many of the key insights we currently have about the possible causes of asthma and have served as invaluable test beds for pharmacotherapy. Mouse models of asthma now represent the bulk of the scientific industry in this field because they can be explored with the most complete range of biological reagents and genomic knowledge. Recent developments in physiological phenotyping and three-dimensional imaging have added greatly to the investigative armamentarium. Combined with computational models of lung function, we are now able to make some rather precise links between structure and function in the mouse lung that enable AHR to be tied to underlying mechanisms. Of course, at the end of the day we are still going to face the issue that a mouse is not a human, so there will always be a gap in our understanding that biological relevance and experimental ethics prevent us from bridging completely.

Fig. 3. The time course of bronchoconstriction in BALB/c mice following aerosolization of 3.125, 12.5, and 50 mg/ml methacholine. The open circles are saline-treated animals; the closed circles are animals treated with poly-L-lysine. The vertical dotted lines bracket parameter values obtained following deep lung inflations given to reestablish baseline conditions. Note the exaggerated response in R\textsubscript{L} compared with G and H in the animals treated with poly-L-lysine compared with controls. This contrasts with the effects of allergic inflammation shown in Fig. 1. *Significant difference in magnitude of response; + significant difference in timing of peak. P < 0.05. [From Bates et al. (15).]

Fig. 4. Mechanical and geometric mechanisms for airways hyperresponsive-ness. A: a modest degree of shortening of the airway smooth muscle (black ring) impinging on an airway wall that thickened is due to epithelial hypertrophy and/or mucus hypersecretion (gray annulus) and leads to a substantial degree in luminal area (white). B: the same degree of luminal narrowing can be caused by accentuated smooth muscle shortening in the presence of a normal airway lining. C: both mechanisms together lead to a dramatic reduction in luminal area and may even lead to complete airway closure.
Nevertheless, we expect the development and investigation of animal models of asthma to continue for some time into the future, where a great deal of new knowledge awaits.

GRANTS
This work was supported by National Heart, Lung, and Blood Institute Grants HL-67273, HL-75593, and HL-87788 and the Centers of Biomedical Research Excellence Grant P20-RR-15557 from the National Center for Research Resources.

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