relationship between bronchial hyperresponsiveness (BHR) and physical activity. Several hypotheses are invoked to explain this association, including a suggestion that physical activity reduces bronchial inflammation by altering airway physiology.

Their major hypothesis is that obesity reduces physical activity and that it is this reduction in physical activity which causes, in some mysterious way, the increase in BHR observed. Their proposed mechanism—that this lack of exercise is associated with a decrease in deep inspiration—is truly breathtaking.

We seek a much more obvious explanation, which is supported by the published evidence. In our recent survey reported in Thorax we demonstrated a highly significant association of body mass index with chronic cough. Other associations observed in this study infer that the cough of obesity is reflux in nature. If obesity leads to reflux-related respiratory symptoms, can this form of upper airway reflux cause BHR related to these symptoms? We cannot rule this out. Finally, obesity leads to reflux—in our recent survey reported in Thorax we demonstrated a highly significant association of body mass index with chronic cough. Other associations observed in this study infer that the cough of obesity is reflux in nature. If obesity leads to reflux-related respiratory symptoms, can this form of upper airway reflux cause BHR?

Unfortunately, Shaaban et al. do not provide us with any information concerning the incidence of classic reflux symptoms in their population. In a study of patients with dyspepsia and endoscopically proven gastroesophageal reflux by Bagnato et al., over one-third had significant BHR. These subjects had no personal/family history or symptoms suggestive of asthma.

However, about two-fifths of patients in the study by Shaaban et al. had asthma-like symptoms, defined as wheeze and sedentary breathlessness. We suggest that these patents could still have reflux-related symptoms as one-third of patients with chronic reflux cough, as demonstrated by pH monitoring, complain of exertional wheeze and dyspnoea.

With the rising levels of obesity in the population, the accurate recognition of the aetiology of the associated BHR is vital to avoid the spurious diagnosis of “late onset” asthma. Perhaps reflux asthma would be a better—but, as yet, unproven—term.

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Bronchial responsiveness and airway inflammation in trained subjects

We read with interest the paper by Shaaban and coworkers on the protective effect of physical activity against bronchial hyperreactivity (BHR) in the general population. The authors suggest that a beneficial effect of deep inspiration during exercise could account for the lower prevalence of BHR in physically active subjects compared with sedentary subjects, while the accompanying editorial favours an “anti-inflammatory” effect of exercise as the most plausible explanation.

We have studied lung function and airway cell biology in non-asthmatic amateur athletes and found that both modulation of airway responsiveness and downregulation of airway inflammation occur with training. At rest, the response to single-dose methacholine inhalation in the absence of deep breaths was significantly lower in amateur runners than in age-matched sedentary controls. Shortly after a marathon race the response to methacholine was further blunted, suggesting a causal relationship between endurance exercise and low bronchial responsiveness, possibly mediated by ventilation at increased lung volumes.

We have previously reported large numbers of neutrophils in induced sputum of runners. However, this finding was not associated with evidence of neutrophil activation after intense exercise, since expression of adhesion molecules by airway neutrophils decreased and the elastase concentration in sputum supernatants was unchanged after a marathon race compared with baseline. Similarly, inflammation in the airways was not associated with activation of the NFκB pathway in endurance-trained mice, while airway inflammation was found to decrease strikingly in ovalbumin-sensitised trained mice compared with sedentary mice. Exercise therefore appears as a model of tightly regulated airway inflammation, possibly secondary to exercise-induced mild bronchial epithelial damage. Along the same line, physically active smokers appear to be protected against lung function decline and the risk of developing chronic obstructive pulmonary disease compared with sedentary smokers, supporting a role for regular exercise in blunting airway inflammation.

We acknowledge that athletes, even at the amateur level, do not represent the general population. On the other hand, a publication bias may have favoured preferential reporting of exercise-associated BHR in athletes, especially those training under extreme environmental conditions (such as “ski asthma”) or exposed to irritants (such as swimmers). It is time to reconsider the beneficial effects of regular exercise as a strategy to preserve respiratory health.

Studies like that by Shaaban and coworkers will certainly help us to move in this direction.

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Proponibacterium acnes in granulomas of a patient with necrotising sarcoid granulomatosis

Necrotising sarcoid granulomatosis (NSG) was first described by Liebow in 1973. It is defined by three pathological features: the presence of a conglomerate mass of sarcoid-like granulomas; varying degrees of necrosis within the confluent granulomas; and vasculitis with granulomas and giant cells involving the walls of muscular arteries and veins. The relationship between NSG and classic sarcoidosis is controversial. In NSG hilar lymphadenopathy is not seen as frequently as in sarcoidosis, extrapulmonary involvement is rare and serum levels of angiotensin-converting enzyme (ACE) are not necessarily raised.

The cause of sarcoidosis is unknown, but it has been hypothesised that it results from exposure of a genetically susceptible individual to specific environmental agents. Abe et al. isolated Propionibacterium acnes (P acnes) in culture from sarcoidosis biopsy specimens, and recently the P acnes genome has been detected in sarcoid lymph nodes by
Numerous dots indicate the existence of *P. acnes* in situ hybridisation in the lung tissue of a patient with necrotising sarcoid granulomatosis. This is the first report of NSG with *P. acnes* DNA found in the granulomas of lung specimens. This may indicate an aetiological link between NSG and *P. acnes*, and it also suggests that NSG is an atypical sarcoidosis with a common aetiology. The clinical and pathological differences between these diseases could be explained by variability in the host response to *P. acnes* or the histological location of *P. acnes*, although further study would be necessary to arrive at more definite conclusions.

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**Figure 1** (A) Open lung biopsy specimen revealing necrotising granulomas (arrow) with giant cells aggregated in masses and distributed in a lymphangitic pattern. (B) Granulomatous vasculitis was also present: granulomas infiltrated the vascular walls and almost completely occluded the lumens aggregated in masses and distributed in a lymphangitic pattern. (C) Granulomatous vasculitis was also present: granulomas infiltrated the vascular walls and almost completely occluded the lumens aggregated in masses and distributed in a lymphangitic pattern. (D) Granulomatous vasculitis was also present: granulomas infiltrated the vascular walls and almost completely occluded the lumens aggregated in masses and distributed in a lymphangitic pattern.