THE IMPACT OF EXERCISE-INDUCED BRONCHOCONSTRICTION ON ATHLETIC PERFORMANCE – A SYSTEMATIC REVIEW

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ABSTRACT

Background: Exercise-induced bronchoconstriction (EIB) describes the phenomenon of transient airway narrowing in association with physical activity. Although it may seem likely that EIB would have a detrimental impact on athletic performance this has yet to be established.

Objectives: Therefore, the aim of this review is to provide a systematic appraisal of the current status of knowledge regarding EIB and exercise performance and to highlight potential mechanisms by which performance may be compromised by EIB. Data sources and study selection: PubMed/Medline and EBSCO databases were searched up to May 2014 using the search parameter: [(‘exercise’ OR ‘athlete’) AND (‘asthma’ OR ‘bronchoconstriction’ OR ‘hypersensitivity’) AND ‘performance’]. This search string returned 243 citations. After systematically reviewing all of the abstracts, 101 duplicate papers were removed, with 132 papers excluded for not including an exercise performance outcome measure. Results: The remaining ten studies that met the initial criteria were included in this review; six evaluated the performance of physically active individuals with asthma and/or EIB while four assessed the effects of medication on performance in a comparable population. Conclusion: The evidence concludes that whilst it is reasonable to suspect that EIB does impact athletic performance, there is currently insufficient evidence to provide a definitive answer.

Key points:

- Further work is required to establish the impact of exercise-induced bronchoconstriction on athletic performance.
- Future studies should address the impact of sport-specific protocols that are conducted in provocative environments.
- Disease severity and athletic standard need to be accounted for in the interpretation of future results.
1. BACKGROUND
At the London 2012 Summer Olympic Games, first and third place across all athletic track events were separated by 1% of the winner’s time [1]. Tiny margins decide success in elite sport and any small impingement on the ability of an athlete to perform maximally can affect competition outcome.

The capacity of the cardiovascular system is generally considered to be the limiting factor underpinning peak aerobic exercise performance [2], whilst in contrast the respiratory system is usually regarded as over-engineered for the demands during intense exercise [3].

Despite this, it is now established that a significant proportion of elite athletes exhibit an ‘abnormal’ airway response to intense exercise exposure. More specifically, up to half of certain cohorts of elite athletes such as swimmers and cross country skiers, appear to develop a post exercise transient narrowing of the airways [4]. This phenomenon, termed exercise-induced bronchoconstriction (EIB), can occur either in the presence or absence of other characteristic features of asthma [5]. Indeed, EIB in elite athletes has a distinct pathogenesis and athletes frequently exceed normal resting lung function [6]. Moreover it is common to encounter athletes with a significant reduction in lung function post exercise who perceive few respiratory symptoms.

It is now recognised that endurance athletes are susceptible to the development of airway dysfunction yet still win a disproportionately high percentage of Olympic medals [7, 8]. However, there is currently a dearth of scientific literature specifically addressing the impact of EIB on exercise performance. Although several studies have aimed to address this, many fail to account for key confounding factors. For instance, studies permitting a warm-up prior to exercise potentially initiates a refractory period resulting in significantly less severe or an absence of airway narrowing [9, 10].
Therefore, the aim of this review is to provide an overview of the current status of research in this field, characterize potential mechanisms by which EIB may impact exercise performance and provide recommendations for future research.

2. METHODS

A systematic evaluation of peer-reviewed literature from the PubMed/Medline and EBSCO databases was performed from January 1980 until May 2014 using the search parameter: [‘exercise’ OR ‘athlete’) AND (‘asthma’ OR ‘bronchoconstriction’ OR ‘hypersensitivity’) AND ‘performance’] (English and humans (Filter), loattrfull text (Filter), not review (Filter) AND has abstract (Filter). This initial search returned 243 papers. Two authors independently reviewed the titles and abstracts of potential studies and subsequently screened full-text study reports for inclusion. After systematically reviewing the literature, 101 duplicate papers were removed, with 132 papers excluded for not including an exercise performance outcome measure (Figure 1).

As research addressing the impact of EIB on exercise performance in elite athletes is relatively limited, individuals regularly participating in sport and/or physical activity were considered further. Included studies required subjects to have a previous physician diagnosis of asthma and/or objective evidence of EIB (i.e. ≥10% fall in FEV₁) following bronchoprovocation testing. Consequently, ten papers met the initial criteria; six purported to evaluate exercise performance in physically active individuals with asthma and/or EIB [11-16] while four assessed the effects of medication on performance in a comparable population [17-20]. These will be considered in more detail in the subsequent sections and form the basis of this review (Table 1). Furthermore, an appraisal of evidence using the Grading of Recommendations, Assessment, Development, and Evaluation (GRADE) [21] approach has been provided (Table 2).
3. RESULTS

3.1 DOES EIB IMPACT ON ATHLETIC PERFORMANCE?

The limitations to performance during sport are dependent on the nature of the exercise being undertaken. It is unlikely that EIB would affect athletic performance during an event lasting <1 min. However, during exercise of longer duration and with specific aerobic demands, an abnormal airway response could impede performance.

3.2 Maximal oxygen consumption

When evaluating an individual’s potential to perform successfully in endurance sport, maximal oxygen consumption (VO_{2max}) is often used as a surrogate measure for aerobic or cardiovascular capacity. As VO_{2max} is a good differentiator of aerobic performance in a heterogeneous group of individuals it is often used to characterize general athletic ability, i.e. VO_{2max} for a male individual of average athletic ability is ~45 mL.kg^{-1}.min^{-1} and it is unusual to encounter elite level endurance athletes with a VO_{2max} <65 mL.kg^{-1}.min^{-1}.

It has been argued that, in asthmatic athletes, VO_{2max} may be limited by factors other than cardiovascular capacity; i.e. by processes that slow the delivery of oxygen such as persistent airway narrowing, alveolar wall thickening and loss of elastic recoil. Certainly adult patients with severe asthma have reduced aerobic capacity [22] and asthmatic children appear to have a lower VO_{2max} and running performance than their non-asthmatic peers [23].

Despite this, Freeman et al. [14] reported that asthmatic athletes with moderate severity disease (i.e. FEV1<50% ) can still achieve a high VO_{2max} (i.e. >60 mL.kg^{-1}.min^{-1}). Moreover, cross-sectional studies show no significant differences in VO_{2max} between asthmatic and non-asthmatic runners [11, 13]. Indeed, some studies report higher VO_{2max} in elite-trained asthmatic cyclists, when compared with non-asthmatic cyclists [24].

However, the environment in which exercise is performed is important in the context of EIB. Stensrud et al. [15, 16] found that VO_{2max} and peak running speed (V_{peak}) was approximately
5% greater in adults with EIB (mean VO$_{2\text{max}}$: 49 mL kg$^{-1}$ min$^{-1}$) exercising in a humid environment (19.9°C and 95% relative humidity) when compared to results obtained when exercising in standard ambient conditions (20.2°C and 40% relative humidity) [16]. Moreover, VO$_{2\text{max}}$ and running performance was reduced by 6.5% in subjects with EIB exercising in a cold environment (-18°C and 39.2% relative humidity) [15]. Whilst these studies provide some evidence to suggest EIB has a deleterious impact on exercise performance, it is important to note that the study population was replicated, thus limiting the strength of evidence. Furthermore, a decrease in the endurance time of healthy physically active subjects has been observed during exercise in similar environmental conditions [25]. Moreover, the protocol employed was neither a traditional incremental exercise test nor a performance trial; running speed was set “by a combination of 95% of estimated maximum heart rate and the test leader’s evaluation of exhaustion after eight min” [15, 16]. Finally, athletes recruited for this study were well below the level expected for elite or competitive athletes (mean VO$_{2\text{max}}$: 48 mL kg$^{-1}$ min$^{-1}$). Indeed, only one study has examined VO$_{2\text{max}}$ in a cohort of subjects with EIB who could truly be considered of competitive athletic ability. Teixeira et al. [11] reported no difference in peak oxygen consumption determined by treadmill ramp protocol between untreated EIB positive (mean VO$_{2\text{max}}$: 63 mL kg$^{-1}$ min$^{-1}$) and EIB negative (mean VO$_{2\text{max}}$: 64 mL kg$^{-1}$ min$^{-1}$) marathon runners. This similarity in peak VO$_{2\text{max}}$ between subjects with and without EIB is consistent with an earlier retrospective cohort study conducted in ($n = 137$) army recruits [12] where no difference in baseline VO$_{2\text{max}}$ was observed between EIB positive and EIB negative groups. Moreover, performance in a two-mile time trial was not different between groups with both displaying similar features of athletic adaptation to the eight-week training programme.

In contrast to aerobic exercise, there have been no studies to date that address anaerobic capacity in athletes with EIB. Whilst Kippelen et al [26] reported no difference in VO$_2$ kinetics
between recreationally active EIB positive and trained negative controls during high-intensity exercise, no evidence is currently available in elite athletes.

3.3 Performance trials

Although VO₂max is a good predictor of aerobic exercise performance in the general population, the value of VO₂max in differentiating performance in a homogenous group of elite athletes is relatively limited [27]. This is likely due to contributing factors such as exercise, efficiency and motivation.

Several limitations arise from using an incremental or maximal exercise test as a performance outcome measure in trained athletes; specifically, the fact that the nature of the test does not reflect the demands of the sporting event.

Similar to Dickinson et al. [28], we have previously observed a significant improvement in lung function following the administration of inhaled salbutamol (400 µg) versus placebo in elite soccer players (n = 11) with asymptomatic EIB, however no difference was detected in multiple sprint times between trials [29]. In agreement, Kalsen et al. [30] observed similar findings in elite swimmers.

Furthermore, a recent study by Koch et al. [20] addressed the effects of administering 400 µg salbutamol on lung function, respiratory parameters and a 10 km time-trial performance in 49 trained cyclists (VO₂max range: 53-85 mL·kg⁻¹·min⁻¹) with (n = 14) and without (n = 25) EIB. They observed that although salbutamol improved resting lung function in both groups, no difference in performance or perception of effort was detected.

Whilst it is acknowledged that the evidence to date is extremely limited, there is no published data to suggest that EIB limits sport-specific exercise performance.
4. MECHANISMS BY WHICH EIB MIGHT IMPACT PERFORMANCE

There are a number of mechanisms by which bronchoconstriction may hypothetically impede athletic performance, however, if there is an impact then this likely arises due to the interplay between several factors (Figure 2).

4.1 Exercise airflow limitation

In both healthy and asthmatic individuals, a bronchodilatory response has been shown to occur with the onset of exercise [31]. However, it is unlikely that mild bronchodilation is of great significance in subjects susceptible to substantial bronchoconstriction (i.e. ≥25% fall in FEV₁).

In the context of EIB, airflow obstruction and symptoms classically develop in the period (e.g. 5-10 minutes) following exercise. Therefore, it has often been argued that EIB does not impact on in-exercise airflow and therefore is not relevant to exercise performance.

During exercise in healthy subjects, peak expiratory flow occurs close to the middle of the expiratory phase of respiration and there is a mild reduction in end-expiratory lung volume (EELV). However, in elite athletes who achieve high levels of ventilation there is evidence of encroachment of the tidal flow loop on the superimposed forced expiratory flow-limb. Moreover EELV may approach or surpass resting functional residual capacity (Figure 3). In recreationally active asthmatic individuals, Haverkamp et al. [32] reported an increase in average pulmonary resistance for the entire exercise duration and for the majority of subjects, expiratory flow limitation was evident. Furthermore, others have shown that EELV increases in exercising asthmatics [33].

Whilst it is accepted that limitations exist in the current methodology and techniques available to evaluate the impact of in-exercise airflow limitation [34], it seems possible that changes in maximum expiratory flow volume during exercise may influence airflow and the perception of dyspnoea experienced by athletes with EIB, as they adopt ventilatory strategies to overcome this constraint.
4.2 Increased work and oxygen cost of respiratory muscles

During exercise in healthy subjects, the oxygen consumption of respiratory muscles accounts for 5-10% of whole body oxygen consumption [35]. However, alterations in EELV, ventilation-perfusion ratio and airway calibre alter the work, and therefore oxygen consumption of the respiratory muscles [36].

In support of this concept, Aaron et al. [35] observed that the oxygen cost of breathing increases to 15% of total oxygen consumption in the presence of severe expiratory airflow limitation. This increased effort of breathing and associated oxygen cost has two potential consequences for exercise performance. Firstly, to drive a higher overall oxygen cost for a given external work load and secondly to potentially drive an earlier onset of respiratory muscle fatigue [37]. Increased respiratory muscle loading may also have a deleterious impact on exercise performance via the ‘respiratory muscle metaboreflex’ [38]. Strenuous exercise (>80% VO2max) in healthy individuals is associated with a reduction of diaphragmatic force following supra-maximal motor nerve stimulation—indicative of diaphragmatic muscle fatigue. This in turn may increase sympathetic vasoconstrictor outflow, reducing skeletal muscle blood flow [38]. It has been suggested that the restricted blood flow augments locomotor muscle fatigue and limits exercise capacity [37].

Furthermore, the recruitment of accessory respiratory muscles with diaphragmatic muscle fatigue has been shown to distort the chest wall and reduce the mechanical efficiency of breathing [37]. Therefore transient bronchoconstriction may increase respiratory muscle work during exercise resulting in respiratory muscle fatigue, compromised breathing efficiency and reduced oxygen availability to the working limb musculature.

4.3 Dyspnoea and perception of effort

Dyspnoea describes “a subjective experience of breathing discomfort that consists of qualitatively distinct sensations that vary in intensity”, however the precise genesis is complex
and remains incompletely understood [39]. It is now established that perceived exertion modulates exercise performance in healthy individuals and that the cues for sensation of perceived exertion are distinct from those of sense of effort [40]. Ventilation appears to be an important contributor to perceived exertion during exercise particularly at higher intensities [41].

It is postulated that dyspnoea begins with a physiological impairment (i.e. airway dysfunction), that leads to stimulation of pulmonary and extra-pulmonary afferent receptors; leading to afferent feedback where an uncomfortable or unpleasant sensation is perceived (i.e. chest tightness) [42]. More specifically, the sensation of dyspnoea may represent a conscious awareness of outgoing respiratory motor command. As the brainstem or motor cortex provides efferent commands to the ventilatory muscles a copy of this command is sent to the sensory cortex. The exchange between the motor and sensory cortex is termed ‘corollary discharge’ and is thought to be the mechanism by which conscious perception of effort arises [43]. This exchange has been proposed to determine the severity of dyspnoea and perception of effort during exercise. It is now possible to obtain surrogate indices of ventilatory drive and load imbalance [44] that can be used to characterize dyspnoea. Therefore it seems reasonable to posit that any condition that disproportionally increases the effort of breathing to the ventilatory load during exercise will amplify the perception of exertion causing premature termination of the exercise bout or a moderation of the intensity at which the exercise is performed.

To date, only one study has addressed the impact of inhaled short acting beta2-agonists versus placebo on central nervous system parameters [45]. Whilst no differences were observed on psychometric visual analogue scales and psychomotor performance tests between treatments, the cohort consisted of non-asthmatic trained triathletes (n = 23) (mean VO2max: 58 mL·kg⁻¹·min⁻¹) rather than asthmatic subjects.
4.4 Ventilation/perfusion mismatch

It is well recognised that a significant proportion (up to a third in some series) of healthy trained endurance athletes exhibit exercise-induced arterial hypoxemia (EIAH); i.e. a drop in oxygen saturation defined as mild 93–95% (or 3–4% <rest), moderate 88–93%, or severe <88% [46]. Haverkamp et al. [32] demonstrated EIAH during high-intensity exercise in physically active asthmatics (mean VO₂max: 48 mL·kg⁻¹·min⁻¹). The decreased effectiveness of gas exchange in this population may be a result of airway inflammation in both the airways and alveolar areas leading to a poorer alveolar ventilation-perfusion ratio, smooth muscle constriction, mucosal and interstitial oedema with infiltration and closing of the small airways [32].

Munoz et al. have previously demonstrated that in mild asthmatics, EIB provokes a greater ventilation-perfusion imbalance. They proposed that this is primarily due to post exercise increases in minute ventilation and cardiac output, benefiting partial pressure of oxygen in arterial blood. In addition, ventilation-perfusion imbalance likely reflects uneven airway narrowing and blood flow redistribution generating unique ventilation-perfusion patterns including the development of areas with low and high ventilation-perfusion ratios [47]. Consequently, depending on severity, EIB may cause ventilation-perfusion imbalance and may impair the respective uptake and elimination of O₂ and CO₂ into and from the blood.

5. EIB TREATMENT AND IMPACT ON PERFORMANCE

A significant amount of research has been conducted over the past two decades investigating how asthma treatment may impact on athletic performance. This is extremely pertinent given the fact approximately one fifth of swimmers and cyclists used beta2-agonists at the Summer Olympic Games in 2004 and 2008 [7]. It is now well established that pharmacological (e.g. inhaled corticosteroids in combination with a pre-exercise inhaled beta-2-agonists and if required leukotriene receptor antagonist / mast cell stabiliser) and possibly non-pharmacological interventions (e.g. fish oil
supplementation) are effective in protecting against airway constriction [48]. However, the mainstay of treatment for protection against EIB is the administration of an inhaled short acting beta2-agonist (e.g. salbutamol) 15 minutes prior to exercise [49].

5.1 Impact of medication in non-asthmatic athletes

Bedi et al. suggested that inhaled albuterol may provide a competitive advantage for non-asthmatic athletes resulting in speculation that beta-2 agonists are performance enhancing [50]. However, research to date provides little evidence that inhaled beta2-agonists in prescribed doses provide performance benefits in non-asthmatic subjects [51]. A recent systematic review and meta-analysis confirmed that no significant effects were detected for inhaled beta2-agonists on endurance, strength or sprint performance in healthy athletes. However, it was highlighted that systemic beta2-agonists may elicit positive effects on physical performance, although the quality of evidence is not currently sufficient to provide a definitive answer [52]. In contrast, clenbuterol is now prohibited by the World Anti-Doping Agency (WADA) given the anabolic effects associated with treatment [53].

In a sport-specific context, Sporer et al. [54] observed no effects of 800 µg salbutamol on 20-km time-trial performance in healthy elite cyclists (mean VO₂max: 67 mL·kg⁻¹·min⁻¹).

In contrast, some researchers have observed performance benefit; Van Baak et al [55] reported that 800 µg salbutamol significantly improved time-trial performance in well-trained amateur non-asthmatic male cyclists and triathletes (n = 16) by 1.9 ± 1.8% (P<0.05). However, lung function increased significantly following administration of salbutamol versus placebo suggesting some of the subjects may have had undiagnosed airway obstruction. More recently, Decorte et al. [56] observed an increase in muscular endurance following the inhalation of 800 µg salbutamol. Similarly, Kalsen et al. [30] showed an improvement in arm ergometer sprint performance and muscle strength following the inhalation of high doses of beta2-agonists in elite swimmers with and without airway hyperresponsiveness.
5.2 Impact of medication in asthmatic athletes

In asthmatic men (mean VO$_{2\text{max}}$: 43 mL·kg$^{-1}$·min$^{-1}$) Freeman et al. [19] observed no improvements in performance during progressive exercise following nebulization of 5mg of salbutamol despite a significant increase in resting and post-exercise FEV$_1$ [19]. Similarly, Ingemann-Hansen [57] showed no benefits of salbutamol on VO$_{2\text{max}}$ or endurance time to exhaustion within a comparable population.

In contrast, the administration of montelukast, a leukotriene antagonist, has been shown to improve physical performance and oxygen pulse while reducing the perception of strain at maximal workloads [17, 18]. However, the subjects recruited in these studies were only moderately trained individuals (mean VO$_{2\text{max}}$: 44 mL·kg$^{-1}$·min$^{-1}$) and even at fatigue did not attain a maximal perception of effort score. Thus it is difficult to extrapolate these findings to elite athletic performance.

5.3 Impact of supra-therapeutic medication doses

In non-asthmatic athletes, Dickinson et al. recently observed no effect on endurance, strength or power performance following 6-weeks 1600 µg inhaled salbutamol (WADA daily upper limit) [58]. Furthermore, no significant improvement in 5km time-trial performance was observed following one dose 1600 µg salbutamol in a comparable population [59]. Irrespective of the aforementioned findings, information is currently relatively limited regarding the effect of asthma medication administered at supra-therapeutic doses. Indeed long-term use of high doses of beta2-agonists have been shown to be highly anabolic in several animal studies and increase muscle strength in human studies [60] however this effect has yet to be investigated following chronic use of inhaled beta2-agonists. Potentially, very high doses of inhaled beta2-agonists provide the same performance-enhancing effects observed after oral ingestion [61].
6. DISCUSSION

6.1 Does EIB impact athletic performance?

This review reveals that it is not currently possible to draw a definitive conclusion as to whether EIB impacts athletic performance. Several methodological limitations contribute to this conclusion.

Firstly, the experimental designs of studies conducted to date have failed to adequately address the difficulties of studying athletic performance. VO\textsubscript{2max} is consistently reported as the primary outcome measure however the fact that VO\textsubscript{2max} is a poor predictor of athletic performance is often overlooked and sport-specific protocols are more appropriate. For instance, a high proportion of elite cyclists administer beta-2 agonists [7] as an ergogenic aid, yet any performance impact of this treatment will not be adequately detected by an indoor ramp exercise test.

In the very few studies that have employed a sport-specific challenge, EIB does not appear to impact performance [20]. However, it certainly appears that noxious environmental conditions (i.e. cold dry air) inhibit maximal aerobic capacity [15, 16]. However, this has yet to be determined in a sport-specific challenge.

Secondly, the athletic level of subjects varies significantly between studies. This point is highlighted throughout this review whereby athletic ability, as determined by VO\textsubscript{2max} ranges from 40-80 mL kg\textsuperscript{-1} min\textsuperscript{-1}; i.e. from recreationally active to elite level. The sustained ventilatory demand between elite and amateur events differs significantly. When considering exercise hyperpnoea is a key determinant of bronchoconstriction in susceptible individuals, extrapolating data derived from amateur or recreational athletes to the elite level is clearly not appropriate.

Thirdly, it is unclear whether the research to date has examined the best outcome measure. In this review we highlight the fact that central regulation (i.e. the central governor [40]) is a key
component dictating peak athletic performance. It is therefore important to note that studies to date have focused on physiological surrogates of performance rather than central outputs / measures of performance impediment such as rating of perceived exertion during exercise tasks.

6.2 Why do asthmatic athletes appear to outperform non-asthmatic elite athletes?

Elite level athletes with airway dysfunction have consistently been shown to outperform their healthy peers. However as this review highlights, there is no evidence that inhaled beta-2-agonists enhance athletic performance. Explanations for this disparity proposed include the fact that physiological changes / demands associated with EIB may represent an ‘extra’ training stimulus that non-asthmatic athletes do not experience [7]. Others have suggested that airway dysfunction develops in elite athletes throughout the course of their careers by virtue of greater training volume in conjunction with chronic exposure to noxious environmental conditions [4] (i.e. those with EIB may have trained harder and longer to gain a competitive advantage). Finally, the development of EIB may allow mechanical advantages in certain sports such as hyperinflation resulting in improved buoyancy and reduced drag co-efficient in swimmers.

6.3 Recommendations for future research

Further work is needed to explore the impact of EIB on athletic performance, with due consideration of the methodological issues mentioned. More specifically, it is vital that the design of future studies addresses the impact of sport-specific protocols that are conducted in provocative environments and in a homogenous population of athletes. Outcome measures should incorporate metrics of central respiratory drive and the perception of dyspnoea, which is likely to be relevant and pertinent to performance at the elite level. Moreover, further work is needed to explore the physiological consequences of EIB and asthma medication on in-
exercise pulmonary performance in sport-specific environments (i.e. using novel techniques of airflow limitation).

Finally, athletes included in future studies, should have a similar severity of disease. The degree of impairment to exercise performance is likely to be less in athletes with mild EIB (10 - 25% fall in FEV₁) when compared to athletes with more severe airflow limitation. This is relevant in the context of athletic groups ‘screened’ for EIB, many of whom have a mildly positive result to a bronchoprovocation test (e.g. eucapnic voluntary hyperpnoea) and in whom treatment recommendations may be made on the underlying premise that this will ‘improve’ their performance [62].

7. CONCLUSION

In conclusion, the development of EIB may have a deleterious impact on respiratory function during peak exercise however the underlying mechanisms and nature of this impediment remain to be determined. Whilst it is intuitive that EIB, when untreated, has a detrimental impact on elite athletic performance, this has not been established in the literature. Therefore, future work needs to establish the impact of bronchoconstriction on athletic performance and the development of treatment strategies to overcome this potential limitation.
TABLE HEADINGS

Table 1. Studies evaluating the consequences of exercise-induced bronchoconstriction on exercise performance parameters.

Table 2. Studies appraised using the Grading of Recommendations, Assessment, Development, and Evaluation (GRADE) criteria [21].
<table>
<thead>
<tr>
<th>First Author / Year</th>
<th>Population</th>
<th>Diagnosis</th>
<th>Methodology</th>
<th>Summary of findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Freeman et al. 1990 [13]</td>
<td>Asthmatic endurance trained runners (n = 4); mean ± SD VO_{2max}: 56 ± 5 mL·kg⁻¹·min⁻¹</td>
<td>Physician diagnosed asthma</td>
<td>Maximal treadmill exercise challenge; Incremental exercise challenge; 2-hr constant run at 70% VO_{2max}</td>
<td>No differences observed in VO_{2max} or running speed between asthmatic and non-asthmatic runners (P&gt;0.05)</td>
</tr>
<tr>
<td>Freeman et al. 1990 [14]</td>
<td>Asthmatic endurance trained runners (n = 16); mean ± SD VO_{2max}: 63 ± 6 mL·kg⁻¹·min⁻¹</td>
<td>Physician diagnosed asthma</td>
<td>Maximal treadmill exercise challenge; Sub-maximal treadmill exercise challenge</td>
<td>Mild to moderate asthmatics can achieve high VO_{2max} values comparable to healthy individuals; Severe asthma may inhibit VO_{2max}</td>
</tr>
<tr>
<td>Sonna et al. 2001 [12]</td>
<td>US Army recruits (n = 137); mean ± SD VO_{2max}: 46 ± 1 mL·kg⁻¹·min⁻¹</td>
<td>Maximal treadmill exercise challenge; EIB definition: ≥15% fall in FEV₁ post challenge</td>
<td>8-week basic Army training course</td>
<td>7% of the US Army recruits were diagnosed with EIB; EIB did not hinder physical performance gains during basic training</td>
</tr>
<tr>
<td>Stensrud et al. 2006 [16]</td>
<td>Patients with objective evidence of EIB (n = 20); mean VO_{2max}: 47 mL·kg⁻¹·min⁻¹</td>
<td>Maximal treadmill exercise challenge; EIB definition: ≥10% fall in FEV₁ post challenge</td>
<td>Maximal treadmill exercise challenge; Warm environment (19.9°C and 95% relative humidity)</td>
<td>Exercising in a humid environment improved VO_{2max} and V_{peak} in subjects with EIB (P&lt;0.05)</td>
</tr>
<tr>
<td>Stensrud et al. 2007 [15]</td>
<td>Patients with objective evidence of EIB (n = 20); mean VO_{2max}: 48 mL·kg⁻¹·min⁻¹</td>
<td>Maximal treadmill exercise challenge; EIB criteria: &gt;10% fall in FEV₁ post challenge</td>
<td>Maximal treadmill exercise challenge; Environment: Ambient conditions (20.2°C and 40% relative humidity); Cold environment (-18°C and 39.2% relative humidity)</td>
<td>Exercising in a cold environment decreased VO_{2max} and running speed in subjects with EIB (P&lt;0.05)</td>
</tr>
<tr>
<td>Study</td>
<td>Participants</td>
<td>VO2max (mean ± SD, mL.kg⁻¹.min⁻¹)</td>
<td>Challenge Method</td>
<td>Maximal exercise challenge</td>
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<tr>
<td>Teixeira et al. 2012 [11]</td>
<td>Elite runners (n = 20)</td>
<td>64 ± 6</td>
<td>Eucapnic voluntary hyperpnea challenge</td>
<td>Maximal treadmill exercise</td>
</tr>
<tr>
<td>Freeman et al. 1989 [19]</td>
<td>Asthmatic patients (n = 8); Non-asthmatic controls (n = 8)</td>
<td>43 ± 9; 41 ± 6</td>
<td>Physician diagnosed asthma</td>
<td>5mg nebulised salbutamol or saline placebo administered before a maximal exercise challenge performed on a bicycle ergometer.</td>
</tr>
<tr>
<td>Steinshamn et al. 2002 [17]</td>
<td>Asthmatic patients with EIB (n = 16); protocol replicated previous method [17].</td>
<td>45 ± 8</td>
<td>Treadmill exercise challenge</td>
<td>Maximal incremental exercise challenge was performed in ambient, sub-zero temperatures (-15ºC).</td>
</tr>
<tr>
<td></td>
<td>Asthmatic patients with EIB (n = 18); protocol replicated previous method [17].</td>
<td>48 ± 11</td>
<td>Treadmill exercise challenge</td>
<td>Protocol replicated previous method [17].</td>
</tr>
<tr>
<td>Koch et al. 2013 [20]</td>
<td>Cyclists with EIB (n = 14); Cyclists without EIB (n = 35)</td>
<td>65 ± 5; 66 ± 7</td>
<td>Eucapnic voluntary hyperpnea challenge</td>
<td>10 km time-trials 60 min after the administration of either 400 µg salbutamol or 400µg placebo.</td>
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</tbody>
</table>

**EIB**: Exercise-induced bronchoconstriction; **VO2max**: maximal oxygen consumption; **Vpeak**: peak running speed; **FEV1**: forced expiratory volume in one second.
<table>
<thead>
<tr>
<th>First Author / Year</th>
<th>Design</th>
<th>Limitations</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Overall quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Freeman et al. 1990</td>
<td>Observational</td>
<td>VO$_{2\text{max}}$ is a poor predictor of athletic performance</td>
<td>No important inconsistencies</td>
<td>Asthmatic subjects recruited rather than individuals with objective evidence of EIB</td>
<td>Low subject numbers</td>
<td>No further considerations</td>
<td>Very low</td>
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<td>Severity of airway obstruction not defined</td>
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<td>Freeman et al. 1990</td>
<td>Observational</td>
<td>No control group employed</td>
<td>No important inconsistencies</td>
<td>Asthmatic subjects recruited rather than individuals with objective evidence of EIB</td>
<td>Low subject numbers</td>
<td>No further considerations</td>
<td>Very low</td>
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<td>VO$_{2\text{max}}$ is a poor predictor of athletic performance</td>
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<td>Sonna et al. 2001</td>
<td>Observational</td>
<td>VO$_{2\text{max}}$ is a poor predictor of athletic performance</td>
<td>No important inconsistencies</td>
<td>Physically active army recruits rather than athletes</td>
<td>No serious imprecision</td>
<td>No further considerations</td>
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### Stensrud et al. 2006 [16]

**Randomised control trial**

Unable to blind study due to environmental conditions

No control group employed

VO$_{2\text{max}}$ is a poor predictor of athletic performance

Severity of EIB not defined

Standard of asthmatic subjects below standard expected for elite or competitive athletes (mean ± SD VO$_{2\text{max}}$: 47 mL kg$^{-1}$ min$^{-1}$)

No serious imprecision

No further considerations

Low

### Stensrud et al. 2007 [15]

**Randomised control trial**

Unable to blind study due to environmental conditions

No control group employed

VO$_{2\text{max}}$ is a poor predictor of athletic performance

Severity of EIB not defined

Standard of asthmatic subjects below standard expected for elite or competitive athletes (mean ± SD VO$_{2\text{max}}$: 47 mL kg$^{-1}$ min$^{-1}$)

No serious imprecision

No further considerations

Low


**Observational**

VO$_{2\text{max}}$ is a poor predictor of athletic performance

Severity of EIB not defined

Direct

Low subject numbers

No further considerations

Very low

### Medication

Freeman et al. 1989 [19]

**Randomised control trial**

VO$_{2\text{max}}$ is a poor predictor of athletic performance

No important inconsistencies

Asthmatic subjects recruited rather

No serious imprecision

No further considerations

Low
Severity of airway obstruction not defined than individuals with objective evidence of EIB

Standard of asthmatic subjects below standard expected for elite or competitive athletes (mean ± SD 43 ± 9 mL.kg⁻¹.min⁻¹)

No control group employed
VO₂max is a poor predictor of athletic performance
Severity of EIB not defined

No important inconsistencies

Standard of asthmatic subjects below standard expected for elite or competitive athletes (mean ± SD VO₂max: 45 ± 8 mL.kg⁻¹.min⁻¹)

No serious imprecision
No further considerations

Severity of EIB not defined

No important inconsistencies

Standard of asthmatic subjects below standard expected for elite or competitive athletes (mean ± SD VO₂max: 48 ± 11 mL.kg⁻¹.min⁻¹)

No serious imprecision
No further considerations

Severity of EIB not defined

No important inconsistencies

Direct

No serious imprecision
No further considerations

**EIB**, Exercise-induced bronchoconstriction; **VO₂max**, maximal oxygen consumption.
FIGURE HEADINGS

**Figure 1.** PRISMA flow chart representing search results.

**Figure 2.** Proposed schematic depicting how exercise-induced bronchoconstriction may impact exercise performance.

**Figure 3.** Maximal flow-volume loops during incremental maximal exercise: (a) endurance athlete without expiratory flow limitation; (b) endurance athlete with objective evidence of bronchoconstriction.
Figure 1.
Exercise-induced bronchoconstriction

- Increased respiratory muscle work
  - Respiratory muscle fatigue
  - Respiratory muscle metaboreflex
  - Fatiguing contractions of the diaphragm, expiratory and accessory muscles
  - Increased reflex activating metabolites
  - Increased group III/IV phrenic afferent discharge
  - Increased sympathetic efferent discharge

- Exercise airflow limitation
  - Alterations in ventilation mechanics
    - Decreased efficiency
    - Decreased lung capacity
    - Dynamic hyperinflation

- Increased perception of effort
  - Afferent feedback
  - Increased perception of effort

- Ventilation/perfusion mismatch
  - Reduced gas exchange
  - Arterial hypoxemia
  - Reduced oxygen kinetics?

- Symptoms
  - Dyspnoea
  - Shortness of breath
  - Chest tightness

- Reduced exercise capacity

Figure 2.
Figure 3.
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REFERENCES


