Cross refractoriness between sodium metabisulphite and exercise induced asthma

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Abstract

Background – Exercise and inhaled sodium metabisulphite are thought to cause bronchoconstriction in asthma through different mechanisms. The response to both stimuli becomes refractory with repeat challenge. The mechanism of refractoriness is unclear, although depletion of mast cell derived mediators or neurotransmitters has been suggested. Recent studies suggest a common mechanism involving release of inhibitory prostaglandins. If this is true, exercise and sodium metabisulphite induced bronchoconstriction should show cross refractoriness.

Methods – Thirteen subjects with mild asthma and previously established exercise and sodium metabisulphite induced bronchoconstriction performed two sodium metabisulphite challenges (giving a single dose previously shown to cause a 20% fall in FEV1) on one study day, and two exercise tests on another. The second challenge proceeded after recovery (FEV1 > 95% baseline) from the first. Subjects then attended on two further occasions when an exercise test was performed after sodium metabisulphite and a sodium metabisulphite challenge after exercise.

Results – When expressed as the percentage reduction in the area under the change in percentage FEV1 curve over 20 minutes (AUC) the response to exercise was reduced by a mean 62-3% (95% CI 46-5% to 78-1%) following a first exercise challenge, and by 50-7% (95% CI 27-8% to 73-6%) following a sodium metabisulphite challenge. The response to a sodium metabisulphite challenge was reduced by a mean of 80-2% (95% CI 68-9% to 91-5%) when it followed a sodium metabisulphite challenge, and by 37-3% (95% CI 15-1% to 59-5%) following an exercise challenge.

Conclusion – This study shows some cross refractoriness between exercise and sodium metabisulphite induced bronchoconstriction, in keeping with a partially shared mechanism of refractoriness.

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The response to a number of bronchoconstrictor challenges in asthma shows refractoriness – that is, a tendency to diminish with repeat challenge. This occurs most clearly after challenge(s) which cause bronchoconstriction through indirect mechanisms such as exercise,1 ultrasonically nebulised distilled water,2 adenosine 5’-monophosphate,3 and sodium metabisulphite,4 and is unusual with directly acting challenges such as histamine.5 6 The mechanism is unclear, although it has been suggested that refractoriness to exercise,1 ultrasonically nebulised distilled water,2 and adenosine 5’-monophosphate induced bronchoconstriction1 is due to depletion of mast cell derived mediators. Such a mechanism would not, however, explain refractoriness to sodium metabisulphite where bronchoconstriction is thought to involve neural reflexes and not mast cell mediator release.7 8

The cyclooxygenase inhibitor indomethacin inhibits the development of refractoriness to exercise,7 ultrasonically nebulised distilled water,10 and sodium metabisulphite,11 raising the possibility of a common mechanism of refractoriness involving the formation of inhibitory prostaglandins. We have tested the hypothesis that refractoriness to sodium metabisulphite and exercise induced asthma involves a common mechanism by seeking evidence of cross refractoriness. We measured the response to a single dose of sodium metabisulphite so that the response to exercise and sodium metabisulphite could be analysed in the same way.

Methods

SUBJECTS

Thirteen men aged 18 to 46 years with mild asthma requiring inhalation treatment only were recruited from the City Hospital asthma register. Four subjects were taking an inhaled corticosteroid regularly (beclomethasone 200–800 μg daily), one was taking inhaled sodium cromoglycate, and all took an inhaled β2 agonist as required. Beta2 agonists were withheld for six hours and sodium cromoglycate for 12 hours before each visit. Subjects were nonsmokers, had a forced expiratory volume in one second (FEV1) of greater than 60% predicted, and showed a fall in FEV1 after a preliminary exercise test of more than 15%. After verbal and written explanation of the study subjects gave signed consent to participate; the protocol was approved by the City Hospital ethics committee.

MEASUREMENTS

FEV1 was measured on a dry bellows spirometer (Vitalograph, Buckingham, UK) as the
higher of two successive readings within 100 ml. The provocative dose of sodium metabisulphite required to cause a 20% fall in FEV1 (PD20) was established using a MEFAR (Brescia, Italy) breath activated dosimeter to give doubling doses up to a maximum of 128 μmol as described. Exercise tests were performed at room temperature on an electric treadmill (Case 12, Marquette Electronics Inc, Milwaukee, USA). Subjects exercised for seven minutes whilst breathing dry air at room temperature through a mouthpiece connected to a Collins triple J valve, Douglas bag reservoir, and air cylinder. The speed and gradient of the treadmill were set at a level sufficient to maintain a heart rate of 80% of the predicted maximum for each subject. These settings were kept constant for all subsequent exercise tests.

PROTOCOL
Subjects attended on six occasions at the same time of day with visits separated by at least 48 hours. On the first two occasions they attended for a practice exercise test followed, on a separate day, by a sodium metabisulphite challenge to determine the sodium metabisulphite PD20. On the next occasion they attended for a single dose challenge with approximately the PD20 dose of sodium metabisulphite. FEV1 was measured before (baseline) and at intervals after the challenge. Subjects were rechallenged with the same dose of sodium metabisulphite after recovery (FEV1 > 95% baseline value) and FEV1 was then measured at intervals for 20 minutes.

Subjects reattended for two exercise challenges with the second challenge starting after recovery from the first (FEV1 > 95% baseline). FEV1 was measured before the exercise test (baseline) and at intervals after the first challenge and at the same times for 20 minutes after the second challenge. On the final two study days subjects attended for a sodium metabisulphite challenge followed by an exercise challenge, and an exercise challenge followed by a sodium metabisulphite challenge. The order of challenges was randomised.

ANALYSIS
Sodium metabisulphite PD20 was calculated by linear interpolation of the log dose-response curve. The airway response to exercise and the single dose challenge with sodium metabisulphite was expressed as the percentage change from the FEV1 recorded immediately before the challenge (baseline) and described as the maximum percentage fall in FEV1, and area under the change in FEV1, percentage curve over 20 minutes (AUC). Baseline FEV1, the maximum fall in FEV1, and AUC for the first and second challenges were compared within subjects by a paired t test and differences calculated with 95% confidence intervals (CI). A refractory index was derived by expressing the difference between the AUC for the first and second challenge as a percentage of the first AUC.

Results
Baseline FEV1 values before the first sodium metabisulphite and exercise challenges did not differ significantly. Baseline FEV1 values for the second sodium metabisulphite and exercise challenge were about 4% lower than the first, but did not differ significantly from each other. The median time to recovery was 50 minutes following a first exercise test, and 40 minutes following a first sodium metabisulphite challenge.

The maximum percentage fall in FEV1 after exercise was similar on the two occasions it was given as the first challenge (25.8% before exercise and 22.7% before sodium metabisulphite; table). When given as the second challenge the response to exercise was significantly reduced whether following an exercise test or sodium metabisulphite challenge.
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Discussion

We have shown that considerable refractoriness develops after both exercise and sodium metabisulphite induced bronchoconstriction in subjects with mild asthma, in keeping with previous studies. The new finding and main point of interest of the present study is the demonstration of cross refractoriness between exercise and sodium metabisulphite induced bronchoconstriction.

Cross refractoriness has been shown between exercise induced bronchoconstriction and bronchoconstriction induced by hypertonic saline, and adenosine 5'-monophosphate challenge has been shown to reduce the response to exercise and the early response to allergen in subjects with asthma, although it enhanced the response to allergen in atopic subjects. Cross refractoriness between different bronchoconstrictor challenges implies involvement of a common bronchoconstrictor pathway which becomes depleted, or activation of a common protective mechanism. The bronchoconstrictor response to exercise, adenosine 5'-monophosphate, and allergen is partially inhibited by antihistamine treatment, suggesting involvement of mast cell mediator release. Cross refractoriness could therefore be due to depletion of mast cell derived mediators. This would not, however, explain our finding of cross refractoriness between sodium metabisulphite and exercise induced asthma since, unlike exercise, sodium metabisulphite induced bronchoconstriction is not inhibited by antihistamines and is thought to be neurally mediated.

Cross refractoriness between sodium metabisulphite and exercise induced bronchoconstriction could be due to depletion of another pathway, common to the mechanism of bronchoconstriction of both challenges. Loss of airway smooth muscle responsiveness after the first challenge seems unlikely since the response to the directly acting spasmogen histamine is unchanged after exercise challenge in subjects refractory to exercise induced bronchoconstriction. Whether depletion of cholinergic pathways could explain cross refractoriness between sodium metabisulphite and exercise induced bronchoconstriction is also open to doubt since antimuscarinic agents have minimal effect on exercise induced asthma. If non-adrenergic non-cholinergic

metabisulphite (table, fig. 1). The mean maximum fall in FEV, was 13.6% after exercise (mean difference from first exercise challenge 12.2%; 95% CI 8% to 16.4%; p<0.001), and 14.3% after sodium metabisulphite (mean difference from response to exercise before sodium metabisulphite 8.4%; 95% CI 4.3% to 12.5%; p<0.001).

The response to sodium metabisulphite was also similar on the two occasions it was given as the first challenge (26.7% before sodium metabisulphite and 28.8% before exercise; table). When given as the second challenge sodium metabisulphite caused a significantly smaller fall in FEV, whether following sodium metabisulphite (10.1%; mean difference 16.6%; 95% CI 12.1% to 21.1%; p<0.001) or exercise (19.5%; mean difference 9.2%; 95% CI 1.8% to 16.7%; p<0.02; table, fig 2).

When expressed as a percentage reduction in the AUC over 20 minutes the response to exercise was reduced by a mean 62.3% (95% CI 46.5% to 78.1%; p<0.001) by a prior exercise test and by 50.7% (95% CI 27.8% to 73.6%; p<0.001) after a sodium metabisulphite challenge (figs 1 and 2). The response to a second sodium metabisulphite challenge was reduced by a mean 80.2% (95% CI 68.9% to 91.5%; p<0.001) after a first sodium metabisulphite challenge and 37.3% (95% CI 15.1% to 59.5%; p<0.01) after exercise (figs 1 and 2).

Figure 1 Percentage change in FEV, against time after exercise. Points represent mean with standard error of mean. Closed squares = before exercise; closed circles = before sodium metabisulphite; open squares = after exercise; open circles = after sodium metabisulphite.

Figure 2 Percentage change in FEV, against time after sodium metabisulphite challenge. Points represent mean with standard error of mean. Closed squares = before sodium metabisulphite; closed circles = before exercise; open squares = after sodium metabisulphite; open circles = after exercise.
excitatory neural pathways are involved in the bronchoconstrictor response to both exercise and sodium metabisulphite, cross refractoriness could result from depletion of neurotransmitters, as has been shown in animal studies in vitro following repeated neural stimulation.19

The alternative explanation for cross refractoriness between exercise and sodium metabisulphite induced bronchoconstriction is activation of a common protective mechanism such as the release of catecholamines or inhibitory prostaglandins. Catecholamine release during a first challenge is perhaps less likely since this would be expected to have a similar effect on a second challenge irrespective of the mechanism of bronchoconstriction, and previous studies have shown that the response to histamine is unaffected by prior exercise challenge.11 Furthermore, refractoriness to exercise develops without a rise in the concentration of circulating catecholamines.20 A role for inhibitory prostaglandins in the aetiology of refractoriness is supported by studies showing attenuation by indomethacin of refractoriness after exercise,2 sodium metabisulphite,4 and osmoregulated challenges.12,21 22 In man and allergen challenged sensitised guinea pigs,22 The most likely candidate is PGE2, since this is a major cycloxygenase metabolite of human airway epithelium23 and smooth muscle,24 and studies on canine and human airway smooth muscle in vitro show inhibition of cholinergic and non-cholinergic neurally induced contractions by low concentrations of PGE2.25,26 and inhibition of mediator release from immunologically activated human lung mast cells by higher concentrations.27 PGE2 has little or no direct relaxant or anticonstrictor effect on human airway smooth muscle in vitro24 so it would be expected to inhibit the response to bronchoconstrictor challenges acting through mast cell mediator release or neural pathways to a greater extent than those acting directly on airway smooth muscle. This view is supported by studies with inhaled PGE2 in asthma showing very effective protection against allergen,28 exercise,29 and sodium metabisulphite induced bronchoconstriction,30 but minimal effect on the response to the direct airway smooth muscle agonist methacholine.31

We did not show complete cross refractoriness between exercise and sodium metabisulphite induced bronchoconstriction, suggesting that other challenge specific mechanisms may contribute to refractoriness. There was, however, considerable cross refractoriness consistent with an important common component of refractoriness which may well involve inhibitory prostaglandins. The protective effects of frusenide in asthma (which preferentially inhibits the response to indirectly acting challenges32) can be attenuated by treatment with cyclooxygenase inhibitors,33-34 suggesting that there are similarities between the mechanism of action of frusenide and the mechanism of refractoriness. Inhibitory prostanooids, and PGE2 in particular, may therefore have a wider role as an endogenous protective factor against induced bronchoconstriction.

Adventitia

Breathless on Everest – I

“The problem with Everest is that it is just too damned high.” This is not an original or surprising statement, but is one I made to myself several times as I struggled up the Lhotope face, desperately breathless and wishing that the mountain was just 10 000 feet lower. Everest is, of course, very high. At 29 028 feet it is approaching the edge of the earth’s atmosphere and its summit often lies in the jet stream. Fiercose winds of 120 miles an hour or more, which can easily sweep men and all their belongings off the mountain, are commonplace. Because of its height the air is, of course, also very thin. The atmospheric pressure is 253 mm Hg, one third the level at sea level, and that is the level of hypoxia with which climbers have to contend.

The mountain was first climbed in 1953, largely as a result of the work of Dr Griffith Pugh, physiologist with the MRC. He had accompanied an expedition to the base of the mountain in 1951 and made calculations of likely water loss, heat loss, and need for oxygen supplementation. Pugh’s rules still hold. Most climbers use 21/2 min oxygen between Camp 3 at 23 500 feet and Camp 4 at 26 000 feet and then 3–4 litres of oxygen to get to the summit. No one had ever really tested Pugh’s rules, so when I was asked to join the British 40th Anniversary Everest Expedition and given the rather grandiose title of High Altitude Physiologist, I thought that I would try, using modern miniaturised equipment, to see how hypoxic people get at these altitudes and to determine the effect of added oxygen.

The British 40th Anniversary Expedition to Everest had three main aims. The first was to commemorate Hunt’s expedition and to follow the same route that Hillary and Tenzing had used in their triumphant ascent. The second was to raise money for the Himalayan Trust which was set up by Sir Edmund Hillary to provide schools and hospitals in the Khumbu region of Nepal but had recently become concerned with conservation in that increasingly desolate mountain region. The third was to put the first British woman on the summit of the mountain. As everyone now knows Rebecca Stephens made the summit in grand style, alone save for two Sherpas, and has got her name into the record books.

At the time I was invited to join the expedition I had very little climbing experience so I took myself on a crash training course which meant inveigling climbing friends into carting me up and down mountains. By the time I left for Everest I had terrified myself on rock crags and snow gullies all over Scotland and sheer limestone rock faces in Tuscany. I was not prepared, however, for the conditions I would experience on Everest.

We set off on 16 March 1993 from Kathmandu taking a bus to Jiri, a distance of 80 miles. The rest of the trip, approximately 150 miles, we had to walk. We arrived at Everest base camp, which is at 17 500 feet, on 7 April 1993 and sat there looking up at the majestic Khumbu icefall above us. The icefall has claimed more lives on Everest than any other part of the mountain. I set up my tent looking directly up the icefall, just to remind myself every day how frightening it can be. The icefall consists of huge blocks of ice moving at about three feet a day. They are very unstable and constantly falling. Each one may be as big as a cathedral. Between them there are crevasses 100 feet deep. We climbed the blocks of ice on jumars and crampons and crossed the crevasses on roped aluminium ladders. I crossed it six times and that was undoubtedly six times too many. To cross it we would leave at about 2.00 am and the great blocks of ice would be illuminated by the several head torches of the small party winding their way up this unstable environment.

ANDREW J PEACOCK

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