Syndrome X and hyperventilation

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Abstract
The cardiorespiratory responses to exercise and forced hyperventilation were measured in 17 unselected patients with syndrome X (angina, positive exercise test, normal coronary arteriogram, no other cardiovascular disease) and compared with those in 15 healthy subjects. Forced hyperventilation produced hypocapnia and metabolic alkalosis but no chest pain or electrocardiographic change. Patients with syndrome X showed reduced maximum oxygen consumption with an increased respiratory exchange ratio at peak exercise, confirming that exercise was limited by skeletal muscle perfusion—and thus that the increase in cardiac output with exercise is limited in syndrome X as in heart failure. Arterial carbon dioxide tension (Pco₂) homoeostasis during exercise was normal but the ventilatory cost of carbon dioxide excretion was increased in syndrome X (as in heart failure). End tidal Pco₂ measurements correlated only poorly with arterial Pco₂ in individual patients with syndrome X, providing a possible explanation for previous reports, based on end tidal Pco₂ of inappropriate hyperventilation.

Patients with syndrome X did not show inappropriate hyperventilation but they did show hyperventilation that was appropriate to maintain normal arterial Pco₂ in the face of reduced cardiac reserve.

Most patients with chest pain and normal coronary arteriography have non-cardiac pain that is musculoskeletal or oesophageal in origin. A few are thought to have true angina attributable by inference to an impaired coronary dilator response at the microvascular level (syndrome X) though in the absence of a directly documented vascular abnormality or known pathogenetic mechanism the existence of this syndrome is not universally accepted. Because of this uncertainty, some have ascribed the symptoms to hyperventilation, which would accord with the low end tidal partial pressure of expired carbon dioxide. Hyperventilation has indeed been shown to induce electrocardiographic ST segment depression, and alkalosis that is associated with hyperventilation can induce coronary vasoconstriction. To investigate the possible contribution of hyperventilation in these patients, we studied the cardiorespiratory response to exercise and hyperventilation in an unselected group of patients with syndrome X (defined as typical stable effort angina, a positive exercise test, and a normal coronary arteriogram in the absence of other cardiovascular disease).

Patients and methods

PATIENTS
We studied 17 patients (mean age 50 years, range 40–67 years; eight men), all of whom gave an independently confirmed history considered to be typical of effort angina (mean duration 3.9 years) with no other evidence of cardiovascular disease on clinical examination, chest radiograph, echocardiogram, or left ventricular angiogram; and in all of whom the coronary arteriograms were independently confirmed as entirely normal. No patient had hypertension or diabetes. Two patients smoked, and five were ex-smokers. Eleven patients were taking regular medication (four with β blockers, seven with calcium antagonists, four with long acting nitrates). The resting electrocardiogram was normal except in three patients with left bundle branch block. Exercise tolerance was limited in all patients, to the equivalent of New York Heart Association category II (15 patients) or III (two patients): in all patients treadmill exercise showed diagnostic ST segment depression (>1 mm horizontal or downsloping ST segment) or left bundle branch block (in five patients, exercise-related in two). All patients had normal respiratory function tests (spirometry, lung volumes, and transfer factor).

We studied 15 healthy individuals (mean age 47 years, range 22–75 years, eight men) as controls.

PROTOCOL
Patients and controls performed maximum treadmill exercise tests (Weber or standard Bruce protocol, selected as appropriate for exercise capacity) and 3 minutes of forced hyperventilation after familiarisation with the procedures (with 30 minutes rest periods preceding each procedure) ≥2 hours after a meal and >24 hours after stopping medication. The study was approved by the local ethics committee.

Respiratory gas exchange was measured as described elsewhere. During the first exercise test, expired gas from a face mask (Hans Rudolph 7900) was carried to a 7 litre mixing...
box (Airspec) to allow measurement of minute ventilation, oxygen consumption, and carbon dioxide elimination by argon dilution and mass spectrometry (Airspec 2200) corrected to standard temperatures and pressures dry. During the second exercise test and the forced hyperventilation test a capillary tube continuously sampled expired air from within the face mask to measure respiratory rate and end tidal oxygen and carbon dioxide, averaged over the sampling period of 30 seconds; end tidal carbon dioxide values were corrected as previously described. Heparinised arterial blood samples were taken during all these procedures (via an indwelling arterial cannula inserted 30 minutes before any measurement) for blood gas analysis in 13 of the 17 patients.

The relation of minute ventilation to minute carbon dioxide production during exercise was analysed by linear regression. Data are presented as mean (1 SD). Repeated measures in the same individual were analysed by analysis of variance. Other data were compared by paired or unpaired Student's *t* test as appropriate. Probability values of < 0.05 were regarded as significant.

### Results

During voluntary forced hyperventilation the respiratory rate increased (from 12.5 to 31.7 per minute), arterial PCO₂ and end tidal PCO₂ fell (from 36.4 to 23.5 mm Hg and from 33.3 to 20.6 mm Hg, respectively), and pH increased (from 7.4(0.04) to 7.6(0.10)), without chest pain or electrocardiographic change in any individual.

The table shows the exercise test data for patients with syndrome X. The patients had similarly limited exercise duration in both tests (10.2(2.6) and 9.9(2.5) minutes), and maximal oxygen consumption was less than in the controls (20.1(4.4) v 29.7(7.2) ml/min/kg, p < 0.001). Exercise was limited by chest pain in 14 (including two of the patients with fixed left bundle branch block and one with left bundle branch block induced by exercise), and by dyspnoea (a patient with fixed left bundle branch block), fatigue, and hypotension (40 mm Hg fall in systolic blood pressure in a patient with left bundle branch block induced by exercise) in one patient each. In all these patients there was diagnostic electrocardiographic evidence of myocardial ischaemia or left bundle branch block. At maximum exercise the respiratory exchange ratio (minute carbon dioxide/minute oxygen consumption) was increased to 1.1 (0.1) as in the healthy controls (compared with 0.7 (0.1) at rest). At peak exercise, there were very small reductions in arterial PCO₂ bicarbonate, and pH, reflecting a compensated metabolic acidosis; but arterial PCO₂ was normal at 50% maximum exercise. There was thus no evidence of inappropriate hyperventilation.

There was no relation between minute ventilation and minute carbon dioxide (VE/VECO₂) on exercise was linear in all syndrome X patients (all r ≥ 0.92) as it was in the healthy controls (figure); the slope (m) of this relation was, however, significantly steeper (32.3(4.6) v 26.2(5.0), p < 0.01), though the minute ventilation axis intercepts (c) were similar (2.6(1.9)/min v 2.6(2.2)/min). The ventilation required for carbon dioxide elimination was thus increased in patients—for example at a minute carbon dioxide production of 1 l/min, it was 34.9(4.2) compared with 27.9(4.3) l/min in the healthy controls (p < 0.001). In individual syndrome X patients end tidal PCO₂ correlated poorly with arterial PCO₂ both at

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**Table** Heart rate, systolic blood pressure, maximum oxygen consumption, respiratory exchange ratio (n = 17) and arterial and end tidal gas measurements (n = 15) from both exercise tests in patients with syndrome X. (The mean data for patients without left bundle branch block are given in italic and in parentheses, n = 12 and 9 respectively.)

<table>
<thead>
<tr>
<th></th>
<th>Rest</th>
<th>50% of maximum exercise</th>
<th>Maximum exercise</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate (per min)</td>
<td>81 (16) (74)</td>
<td>—</td>
<td>135 (26) (128)</td>
</tr>
<tr>
<td>SBP (mm Hg)</td>
<td>125 (13) (126)</td>
<td>—</td>
<td>160 (22) (160)</td>
</tr>
<tr>
<td>VO₂ max (ml/min/kg)</td>
<td>—</td>
<td>—</td>
<td>20 (4) (20)</td>
</tr>
<tr>
<td>RER</td>
<td>0.7 (0.7)</td>
<td>—</td>
<td>1.1 (0.1)</td>
</tr>
<tr>
<td>CO₂ ET (mm Hg)</td>
<td>35 (4) (35)</td>
<td>38 (4) (38)</td>
<td>37 (4) (37)*</td>
</tr>
<tr>
<td>BaseCO₂ (mm Hg)</td>
<td>37 (4) (37)</td>
<td>39 (4) (39)</td>
<td>36 (3) (36)*</td>
</tr>
<tr>
<td>pH</td>
<td>7.43 (0.04) (7.44)</td>
<td>7.42 (0.03) (7.41)</td>
<td>7.40 (0.06) (7.39)</td>
</tr>
<tr>
<td>Bicarbonate (mmol/l)</td>
<td>26 (1) (26)</td>
<td>25 (1) (25)</td>
<td>24 (3) (23)</td>
</tr>
<tr>
<td>Pao₂ (mm Hg)</td>
<td>92 (13) (92)</td>
<td>92 (7) (91)</td>
<td>100 (5) (106)</td>
</tr>
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</table>

SBP, systolic blood pressure; VO₂ max, maximum oxygen consumption; RER, respiratory exchange ratio (VO₂/VECO₂; CO₂ ET, end tidal PCO₂; BaseCO₂, arterial PCO₂; arterial PO₂.

*p < 0.05 compared with resting value (analysis of variance).
rest (r = 0.36, NS) and during exercise (r = 0.44, NS), though it correlated well during forced hyperventilation (r = 0.86, p < 0.001).

The findings did not differ in patients with left bundle branch block and without. For the 12 patients without left bundle branch block (mean age 49 years, range 40–67 years, five men) the exercise duration in the two tests were 10-1 (1-4) and 9-9 (1-8) minutes and exercise was limited by chest pain in 11 and fatigue in one. Haemodynamic changes, maximum oxygen consumption, and arterial gas changes during exercise were similar in the 12 patients without left bundle branch block (see table), as were the minute ventilation/minute carbon dioxide relations (slope, m = 33.2 (4.5), p < 0.01 compared with controls; ventilation axis intercept, c = 2.7 (1.9)/min).

Discussion
Since the advent of coronary angiography, reports on patients with "chest pain with normal coronary arteries" have accumulated, but the patients have not always been rigorously characterised nor have there been agreed criteria to define such patient groups. Among possible causal mechanisms for the symptoms, hyperventilation has been suggested with supportive evidence from measurements of end tidal PCO₂.

We therefore investigated a group of patients selected solely because they fulfilled pre-defined criteria for what has come to be known as syndrome X. We included patients with exercise induced or fixed left bundle branch block because in patients with syndrome X left bundle branch block can develop; analysis of the data showed similar findings whether or not patients with left bundle branch block were included.

Voluntary forced hyperventilation caused the expected reduction in arterial PCO₂ and respiratory alkalosis but it did not induce chest pain or electrocardiographic change. The patients were therefore studied during formal exercise testing. As in the healthy controls, arterial PCO₂ was maintained within a narrow range by matching the ventilatory clearance of carbon dioxide to its increased metabolic rate of production.13 At maximum exercise there was only a small reduction in arterial PCO₂, compensating for the onset of anaerobic metabolism.13,14 There was no evidence of altered arterial PCO₂ setpoint or inappropriate hyperventilation either at rest or during exercise. Inappropriate hyperventilation thus does not seem to be the cause of symptoms in patients with syndrome X as here defined.

Patients with syndrome X, however, showed reduced maximum oxygen consumption, with metabolic acidosis at peak exercise, implying that exercise was limited by perfusion to the exercising skeletal muscle as in heart failure; this is consistent with reported evidence of cardiac dysfunction in syndrome X.13,15 Moreover, the ventilatory cost of carbon dioxide clearance was increased in patients with syndrome X, a finding also characteristic of patients with congestive heart failure.17,18

The increase in carbon dioxide production resulting from the increased work of exercise, the increased respiratory exchange ratio as anaerobic metabolism developed at peak exercise, and the increased ventilation needed to excrete carbon dioxide all contributed to increased ventilatory cost of exercise that might be called appropriate hyperventilation because it maintained normal arterial PCO₂.

End tidal PCO₂ values correlated well with arterial PCO₂ values in the group as a whole, as was reported in healthy controls, but the correlation in individuals was poor (except during forced hyperventilation). This is consistent with the increase in physiological dead-space implied by the increased ventilatory cost of carbon dioxide excretion in these patients.10

Practically, it indicates that measurements of end tidal PCO₂ cannot be used as an indicator of arterial PCO₂ in individual patients with syndrome X, as was done in earlier studies that suggested that hyperventilation was a common cause of symptoms in patients with "chest pain" and normal coronary arteries.7

We found no evidence of inappropriate hyperventilation in patients with syndrome X, but we did find evidence of appropriate hyperventilation that seemed to result from an increase in physiological deadspace as is typical of other patients with reduced cardiac reserve.

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