Training and electrocardiographic abnormalities in the elderly

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Electrocardiograms (CM5 lead, where one lead is attached in the V5 position and one at the manubrium sterni, with the neutral lead on the back of the neck) have been recorded from 14 men and 25 women aged 60 to 75 years, at rest, during progressive bicycle ergometer exercise to 75 to 85 per cent maximum oxygen intake, and during the subsequent recovery phase. Earlier studies showing a high frequency of ischaemic electrocardiographic abnormalities in women are confirmed; it is suggested that this may reflect a high work load per unit mass of myocardium. Training induces an elevation of the ST segment at rest and during recovery, with a reduction in ST depression during work at a given heart rate, the exercise changes being related to the intensity and frequency of training selected by the subject. Possible explanations of the response to regular exercise include not only the development of the collateral circulation, but also a lessening of the hyperkalaemia of effort and a reduction in the work load per unit mass of myocardium secondary to hypertrophy or a change in the average dimensions of the heart.

The increased frequency of abnormal exercise electrocardiograms with ageing is well documented (Åstrand, 1965, 1969; Doan et al., 1965; Blackburn, 1969; Kasser and Bruce, 1969; Profant et al., 1972; Cumming et al., 1972, 1973). In men, it is generally accepted that there is a significant association between an altered waveform of the ST segment during exercise and myocardial ischaemia, patients with such an abnormality having an increased risk of a 'coronary' attack (Blomqvist, 1965; Blackburn, 1969; Bruce et al., 1969; Kasser and Bruce, 1969; Andersen et al., 1971). In elderly women (Table 1), there is the paradox of an equally high frequency of abnormal records despite a much lower risk of ischaemic heart disease than that seen in the men (Åstrand, 1965; Ostrander et al., 1965; Blackburn, 1969; Profant et al., 1972; Cumming et al., 1973).

Evidence regarding the influence of regular physical activity upon the electrocardiographic changes is conflicting. Saltin and Grimby (1968) noted that in athletes who continued sport into middle-aged and elderly life, ST changes were as common as in the general population, but lower frequencies of abnormalities were observed in ex-athletes. Roskamm et al. (1964) and Pyorala et al. (1967) had similar findings, but Holmgren and Strandell (1959) reported a relatively high frequency of electrocardiographic abnormalities in ex-athletes. Kavanagh and Shephard (1976) recently examined a substantial series of Masters’ class track athletes (age-specific track contests for competitors from the fifth decade (Masters class I), sixth decade (class II), seventh decade (class III), and eighth decade + (class IV); in this group, ST changes were somewhat less frequent than in the general population. Cross-sectional studies suggest benefit from moderate effort. Blackburn (1969) commented that the incidence of post-exercise ST depression was less in farmers than in relatively inactive rail clerks, while Salzman et al. (1969) found that 79 per cent of persons who showed an improvement in physical fitness after training also showed decreases in ST abnormalities during submaximal cycling, improvements being directly related to adherence to the exercise programme and resultant reductions in heart rate at a given load. Bruce et al. (1969) found a 10 per cent decrease of heart rate in submaximal work with a 40 per cent improvement of ST segmental voltages; on the other hand, ST depression was unchanged in symptom-limited maximal work. Several other authors (Mazzarella et al., 1966; Kilbom et al., 1969; Detry and Bruce, 1971; Costill et al., 1974) observed lesser abnormalities at a given work load, but no change in responses at a fixed heart rate after

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training. This implies that any effect of physical activity was the result of a reduction in the total cardiac work load rather than an improvement in myocardial oxygen supply. Kilbom (1971) found an increase in ST segment elevation during work and rest when middle-aged women were trained. However, the two subjects with initial ST depression did not improve.

Kavanagh et al. (1973) carried out experiments on a group of middle-aged ‘post-coronary’ patients. With training, some reversal of ST depression was found not only at a given work load, but also at a given heart rate; the favourable response was shown largely by younger patients who were able to push themselves to an intensive level of physical activity. Accordingly, it was thought of interest to examine the ST responses of a much older group of men and women, relating changes to the intensity of training undertaken.

**Subjects and methods**

The subjects were 14 men and 25 women aged 60 to 75 years. Though volunteers for a prereirement exercise class, the majority were quite sedentary when first enrolled (average bicycle ergometer prediction of aerobic power 22·6 ml/kg per min for the men and 21·0 ml/kg per min for the women, average of direct measurements on the treadmill 29·7 and 26·0 ml/kg per min for men and women, respectively). Apart from 3 men and 2 women, all were non-smokers. The group had also undergone a medical screening which had eliminated some 21 per cent before exercise testing on the usual clinical indications of unsuitability for participation in the exercise training programme. Specific contraindications to enrolment included uncontrolled metabolic disease, a diastolic pressure of more than 100 mmHg, heart failure, orthopaedic disabilities, regular medication other than laxatives or sedatives, history of angina or intermittent claudication, and major abnormalities of the resting electrocardiogram.

All electrocardiograms were recorded with the subjects in a resting but non-basal state. Strenuous physical exertion was avoided on the day of examination, and no meals or cigarettes were consumed within 2 hours of testing. Exercise was performed in an air-conditioned laboratory with a temperature of $22 \pm 2^\circ$C. The subjects wore shorts, and in the case of the women a light halter top. All individuals were familiar with the investigators, and had at least one familiarisation ride on the ergometer before the definitive tests.

The bicycle ergometer was a standard Von Döbeln/Monark design, pedalled at 50 r.p.m. A 12-minute progressive submaximum exercise procedure was followed, with 4 minutes at each stage. The initial loading was 40 to 50 per cent of the estimated maximum oxygen intake and subsequent loadings were adjusted to produce a final heart rate corresponding to 75 to 85 per cent of maximum oxygen intake. After exercise, the subjects remained seated at rest on the bicycle throughout the first 6 minutes of recovery.

During the exercise procedure, the electrocardiogram was recorded by CM5 leads, starting with an initial 6 minutes of rest on the bicycle, continuing through the final 10 seconds of each minute of exercise, and onto the sixth minute of recovery. A Simpliciscriptor electrocardiograph (model EK 100, Litton Medical Products) was used throughout. This apparatus meets American Heart Association specifications for high and low frequency response characteristics. Paper speed was set at 25 mm/s, and the gain was adjusted so that 1 mm of galvanometer deflection was equal to a signal of 0·1 mV. ST voltages were determined by simple inspection on a representative series of waveforms, values being averaged over a respiratory cycle. The isoelectric line was defined by joining consecutive PR intervals, and the ST displacement was recorded at the end of the ST segment, immediately before the T wave.\(^1\)

Exercise electrocardiograms were again recorded after 7 and 14 weeks of endurance training. Subjects met for an hour of supervised physical activity 4 times per week. The emphasis was on fast walking and jogging. An initial target pulse rate of 120 to 130 per minute was set, with progression to values of 140 to 150 per minute as the physical condition of the individual permitted. Class members were rated for the vigour and frequency of their participation, so that over the course of the study 4 self-

\(^1\)This measurement site avoids many of the artefacts that can produce ST depression in healthy young normal subjects (Lepeschkin, 1969).
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ST
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showing clinically
Subjects
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During
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averaged
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±10
97
±30
NS
<0.01
<0.05
<0.01
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<0.005).
At
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**Results**

**INITIAL STATUS**

Under resting conditions (Table 2), the men tended
to have a small elevation of the ST segment
(P < 0.01); in women the group average resting ST
voltage did not differ significantly from zero,
though 1 woman showed a substantial (> 0.1 mV)
depression.

During exercise, ST voltages became slightly
negative in both men and women, the trend for
the women being statistically significant at heart rates
averaging 118 and 135 beats per min (P < 0.01 and
< 0.05, respectively); however, the averaged data
showed no further increase of ST segmental
depression from the second to the final work load.

During recovery, there was a gradual return of the
ST voltage towards the isoelectric potential, but
this process was still incomplete 6 minutes after
stopping exercise.

Subjects showing clinically noteworthy ST
changes are listed in Table 3. Exercise showed 8
women and 3 men with ST depression greater than
0.1 mV. During recovery, 1 further man and 1
further woman developed abnormal records. On
the other hand, the 3 men and 4 of the 8 women
who showed ischaemia during exercise did not show
this phenomenon during recovery. Perhaps because
they were health-conscious volunteers, the majority
(79% men, 92% women) of our subjects were
non-smokers; nevertheless, the percentage of abnormal
electrocardiograms (29% in the men, 36% in the
women) is much as in previous surveys of sedentary
elderly people. Blackburn (1969) and Profant et al.
(1972) have also commented that the proportion
of abnormalities is influenced but little by smoking
habits.

**RESPONSE TO TRAINING**

The physiological response to training varied with
the self-selected intensity and frequency of effort;
gains of maximum oxygen intake for the high
frequency, high intensity group were as much as
30 per cent of the initial maximum oxygen intake,
whereas changes in the low frequency, low intensity
group were small and statistically insignificant
(Sidney and Shephard, 1976b).

**Resting electrocardiogram**

After 7 weeks of training, there was an insignificant
slowing of heart rate (2.9 beats/min). The resting
electrocardiogram now showed an elevation of the
ST segment; considering the results for both sexes
together, the average change was 0.03 mV
(P < 0.005). At 14 weeks, the slowing of the resting
heart rate (5-4 beats/min) was statistically significant

**Table 2: Initial data showing average deviations of ST segmental voltage (mean ± SD) during bicycle
ergometer exercise**

<table>
<thead>
<tr>
<th>Time of measurement (min)</th>
<th>Sex</th>
<th>ST segmental voltage (mV)</th>
<th>( f_h ) (beats/min)</th>
<th>Work load (Watts)</th>
<th>Significance of deviation from isoelectric potential</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rest, 6</td>
<td>M</td>
<td>±0.04 ±0.04</td>
<td>88 ±14</td>
<td>—</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>+0.01 ±0.08</td>
<td>80 ±13</td>
<td>—</td>
<td>NS</td>
</tr>
<tr>
<td>Exercise, 4</td>
<td>M</td>
<td>+0.03 ±0.08</td>
<td>103 ±14</td>
<td>36 ±15</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>-0.02 ±0.06</td>
<td>101 ±13</td>
<td>22 ± 9</td>
<td>NS</td>
</tr>
<tr>
<td>8</td>
<td>M</td>
<td>-0.02 ±0.06</td>
<td>118 ±10</td>
<td>66 ±21</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>-0.05 ±0.10</td>
<td>118 ±12</td>
<td>40 ±16</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>12</td>
<td>M</td>
<td>-0.02 ±0.09</td>
<td>140 ± 8</td>
<td>97 ±30</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>-0.04 ±0.09</td>
<td>135 ±10</td>
<td>59 ±16</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Recovery, 2</td>
<td>M</td>
<td>+0.04 ±0.13</td>
<td>110 ±12</td>
<td>—</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>-0.02 ±0.08</td>
<td>98 ±14</td>
<td>—</td>
<td>NS</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>+0.02 ±0.05</td>
<td>100±12</td>
<td>—</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>-0.01 ±0.08</td>
<td>89 ±12</td>
<td>—</td>
<td>NS</td>
</tr>
</tbody>
</table>

Note: Elderly subjects (M, 14; F, 25), 4 minutes per stage of progressive test carried to 75 to 85 per cent of maximum oxygen intake.

**Table 3: Occurrence of ischaemic changes in ST segment of electrocardiogram (≥ 0.1 mV depression) during bicycle
ergometer exercise (M=14, F=25). Intensity of effort was increased progressively to reach 75 to 85 per cent of
\( VO_2 \) max**

<table>
<thead>
<tr>
<th>Occurrence of ischaemic change</th>
<th>M</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exercise only</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Recovery only</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Exercise and recovery</td>
<td>0</td>
<td>5*</td>
</tr>
</tbody>
</table>

Total (exercise and/or recovery) 4 (29%) 9* (36%)

*One woman showed an electrocardiogram which indicated ischaemia at rest and during exercise and recovery.
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(P < 0.005). At this stage, there was a further small increase of ST segmental voltage (0.01 mV), but the change from 7 to 14 weeks was statistically insignificant. The extent of the positive deviation was apparently related to the extent of training undertaken; the high frequency, high intensity group showed an 0.06 mV elevation over 14 weeks (P < 0.05), and the high frequency, low intensity group also had an increase of 0.05 mV that was nearly significant, but there was no change in the low frequency, high intensity and low frequency, low intensity groups.

Exercise electrocardiogram

After 14 weeks of conditioning, the men showed decrements in exercise heart rates averaging 14 to 15 beats/min at each of the test work intensities, with increments in ST segment voltages averaging 0.04 to 0.06 mV (Table 4). In the women, the pre-training, post-training differences averaged 3 to 8 beats/min for heart rate and 0.01 to 0.07 mV for ST voltage; changes were significant for the first work levels only.

As a second approach to determining whether exercise ST segmental voltages were altered with physical training, we calculated linear regression equations relating ST segmental voltage to exercise heart rate for each individual. The ST segmental voltage corresponding to a fixed heart rate of 120/min was then predicted from these equations. A paired t test analysis of the data indicated that the potential of the ST segment at the fixed heart rate changed from -0.03 mV to +0.01 mV (N = 38, P < 0.01), with 7 weeks of training, and from -0.03 mV to +0.03 mV (N = 31, P < 0.025), with 14 weeks of training. The increase in potential of the ST segment was equal for both sexes (0.03 to 0.04 mV in men and 0.04 to 0.05 in women), but the pretraining, post-training differences were significant only for women. Classifying subjects according to the intensity and frequency of training, changes in the ST segmental voltage were significant only for the high frequency, low intensity group.

Eleven subjects initially showed ST depression greater than 0.1 mV during submaximal exercise. After 14 weeks of training, 5 of the 11 (4 women, 1 man) had improved to the point where the depression was less than 0.1 mV at the selected test heart rates. A further two men who failed to resolve their abnormality with 14 weeks of training did so over an additional 35 to 40 weeks of participation in the exercise class. One further woman had a substantial lessening of ST depression, although her final tracing was still clinically abnormal. The three remaining women showed no evidence of improvement. In addition, 1 woman and 1 man from the low frequency, low intensity group showed ischaemia in the final tests that had not been observed at the initial examination.

Recovery electrocardiogram

Considering all subjects, the 2-minute post-exercise tracing showed an insignificant elevation of the ST segment (0.01 mV) after 7 weeks of training. An additional 7 weeks of exercise class membership elicited no further changes. The largest elevations were seen in the high frequency, high intensity group (average 0.05 mV), but even this change was statistically insignificant. At the 6th minute of recovery, there was a 0.04 mV (P < 0.025) increase of ST voltage after 14 weeks of training. Changes were again largest for the high frequency, low intensity group (7 weeks, 0.06 mV, P < 0.025, 14 weeks, 0.05 mV, P < 0.025) and the high frequency, high intensity group (7 weeks, 0.05 mV non-significant; 14 weeks, 0.09 mV, P < 0.001).

Of the 6 subjects who initially showed clinically noteworthy ST depression after the ergometer test, 4 lost this sign as training proceeded.

Discussion

ABNORMALITIES IN WOMEN

The present data support the view of several earlier authors that electrocardiographic abnormalities are at least as common in elderly women as in men of

<table>
<thead>
<tr>
<th>Table 4 Changes in exercise heart rate and ST segmental voltage with 7 and 14 weeks of training: values are mean differences ± SD</th>
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<tbody>
<tr>
<td><strong>Work rate (Watts)</strong></td>
</tr>
<tr>
<td>-----------------------</td>
</tr>
<tr>
<td>Load 1</td>
</tr>
<tr>
<td></td>
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<td>Load 2</td>
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<td></td>
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<td>Load 3</td>
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<td></td>
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</tbody>
</table>

*P < 0.05, †P < 0.025, ‡P < 0.01, §P < 0.025, ‖P < 0.005, **P < 0.001.

Br Heart J: first published as 10.1136/hrt.39.10.1114 on 1 October 1977. Downloaded from http://heart.bmj.com/ on September 16, 2023 by guest. Protected by
of secondary differences (Anderson, 1969). Factors motivating subjects to volunteer for the present study were similar in men and women (Sidney and Shephard, 1976a), so it is unlikely that the proportion of abnormal electrocardiograms in the women was boosted by sample selection.

(b) Since women tend to have a more horizontal ST segment and a smaller T wave than men, it has been postulated that a given segmental depression is more obvious in female subjects (Lepeschkin, 1969). This could explain a high frequency of reported anomalies in ‘eyeball’ studies (Profant et al., 1972), but can hardly account for the series of carefully measured records showing frequent ST segmental depressions in elderly women. In the present experiments, data were referred to an isoelectric line drawn between consecutive PR segments; a small T wave and a more horizontal ST segment could conceivably have displaced our measurement site a little further into the diastolic phase in the women, but this would have reduced rather than increased the proportion of reported abnormalities.

(c) There have been suggestions that sympathetic activity is greater in women than in men at a given level of exercise. However, any difference must be small, since the heart rate at a given percentage of maximal effort is only a little higher in female than in men (Astrand and Ryhming, 1954). There seems no strong evidence that vasoregulatory asthenia, hyperventilation, postural effects, and related ST artefacts are more prevalent in women than in men.

(d) Women have a higher proportion of body fat than men. The frequency of abnormal ST segmental responses increases with obesity (Blackburn, 1969), though the impact of sex on this correlation does not seem to have been explored.

(e) It has been suggested that women may have as much atheroma as men, and may be as liable to ischaemia during exercise, yet rarely suffer fatal consequences of the disease (Dawber et al., 1957; Astrand, 1965; Kannel and Feinleib, 1972). This would accord with Anderson’s hypothesis that heart attacks reflect a ‘vulnerable myocardium’ (Anderson, 1973); hormonal factors and perhaps secondary differences in tissue mineral reserves serve to protect the female myocardium against ischaemia. At the age of 65, hormonal advantages of the female are fast waning. The present authors know of no good studies of atheroma and scarring of the myocardium in apparently healthy women; nor, unfortunately, are there data on the vulnerability of the tissues. However, angiograms often fail to support electrocardiograms showing apparent ischaemia in women (Likoff et al., 1966). A further possibility is that the female myocardium has a normal ‘vulnerability’, but constitutional and cultural factors keep women from the sudden increments in cardiac work load that could precipitate an ‘electrical’ death; an electrocardiographic test that requires elderly women to work to the same percentage of maximum effort as the men may be culturally unrealistic!

(f) The quadriceps is generally weaker in women than in men. Thus, it might be argued that attempts to perform the same relative work load with weaker muscles give rise to a greater tachycardia, a greater increase of systolic blood pressure, and a greater increase in cardiac work load. Kassar and Bruce (1969) suggest that if the heart of an elderly person is sufficiently loaded, ST segmental depression can arise in the absence of significant coronary disease. In the present experiments, the final blood pressure was only marginally higher in the women than in the men (194 versus 191 mmHg), while the final exercise heart rate was slightly lower for the women. Total cardiac work loads, therefore, cannot be blamed for the high proportion of ST abnormalities in the women.

(g) Perhaps the most intriguing possible explanation lies in weakness of the cardiac rather than the skeletal muscle. Women typically have smaller hearts than men, and the same absolute cardiac work load thus throws a greater relative strain on unit volume of cardiac tissue. It is most unlikely that the capillary supply per unit mass of muscle is greater in women; thus if a woman is taken to the same heart rate/pressure product as a man, the woman would inevitably be more liable to myocardial ischaemia. Such a deduction reinforces the conclusion of Cumming et al. (1973) that there is a need to revise criteria of abnormality for the female electrocardiogram.

**Identification of Myocardial Ischaemia**

The present data reinforce the point made by Cumming et al. (1972) in studies of male subjects, that it is desirable to study both exercise and recovery records if all cases with electrocardiographic manifestations of ischaemia are to be identified. The relative prognostic significance of changes during and after effort remains undecided. However, in view of the clinical success of simple submaximal recovery examinations such as the Master test, and
the reports of emergencies developing shortly after exercise (Bruce et al., 1968; Shephard and Kavanagh, 1975), recovery data should not be rejected.

**RESPONSES TO ENDURANCE TRAINING**

The training induced reduction in electrocardiographic abnormalities at a given heart rate and blood pressure is in accord with an earlier study of subjects who trained hard (Kavanagh et al., 1973). Many postcoronary programmes use relatively homoeopathic doses of exercise, and their failure to reduce ischaemia at a given heart rate or pulse-pressure product may reflect an insufficient intensity of physical activity. It is less certain that intense training can develop the collateral circulation, as might be expected from work with experimental animals. An alternative hypothesis is that a strengthening of the cardiac muscle and a reduction of mean ventricular radius decreases the work load sustained by unit volume of cardiac tissue.

The situation would then be the antithesis of that in the elderly untrained woman; electrocardiographic evidence of ischaemia would be reduced, but there would not necessarily be any correction of the underlying atheroma. A further possibility is that the well-trained subject releases less intramuscular potassium at a given intensity of effort (Blomqvist, 1969); this, also, would reduce ST changes.

The increase of voltage in the ST segment under resting conditions is not pathological—indeed Sjöstrand (1950) has related the phenomenon to a slow heart rate, while Saltin and Grimby (1968) have commented that endurance athletes show quite distinct ST elevation. Kilbom (1971) has previously seen the development of positive ST segmental voltages in middle-aged women in response to training. The significance of the phenomenon is unknown, though account should probably be taken of the altered resting status in deciding what is a clinically significant ST depression during exercise.

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**References**


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