Symptoms and quality of life in heart failure: the muscle hypothesis

Andrew J S Coats, Andrew L Clark, Massimo Piepoli, Maurizio Volterrani, Philip A Poole-Wilson

Heart failure is a common and debilitating condition with a high mortality. Reductions in mortality in this condition have been achieved by treatment with vasodilators, in particular the angiotensin converting enzyme (ACE) inhibitors, but there remains a considerable burden of disability and mortality. The other mainstay, diuretic treatment, has never been shown to reduce mortality in heart failure but its ability to relieve life threatening pulmonary oedema and to help relieve fluid retention and dyspnoea in chronic heart failure is not seriously questioned.

It is likely to be increasingly difficult to reduce mortality with newer treatments without specifically targeting the underlying processes leading to left ventricular dysfunction. There remain, however, many unresolved symptoms in optimally treated patients, with a correspondingly low quality of life. This is a valid area for therapeutic intervention, but one in which objective measurements of benefit have proved difficult to obtain. Therapeutic strategies have also been limited by our poor understanding of the physiological bases of the cardinal symptoms, dyspnoea and fatigue.

Measurement of quality of life in heart failure

Fletcher and colleagues have described the requirements of an ideal assessment of the quality of life (table). Questionnaires are the most common means of achieving such an assessment, and these can take the form of a general health survey—for example, the Nottingham health profile—or be specific to the condition under investigation—for example, the Yale scale, or the Minnesota living with heart failure questionnaire. Quality of life assessments have indicated improvement with some treatments, but on an individual patient basis there can be very poor correlation between the change in the quality of life scores and objective improvements in exercise tolerance. Also, different exercise protocols can show different changes with intervention, and there is little consensus as to whether incremental exercise tests or submaximal endurance tests better measure the symptoms limiting patients with chronic heart failure.

Symptoms limiting exercise in chronic heart failure

There is little correlation at rest or during exercise between measures of haemodynamics and either the symptoms limiting exercise in chronic heart failure or the degree of such limitation. Different symptoms may be produced by different forms of exercise tests or by different speeds of incremental exercise tests. Even in the same patient there is little correlation between the peak achieved pulmonary artery pressure and which of the two main symptoms limit different types of exercise. With ambulatory pulmonary arterial pressure monitoring it has been shown that limiting dyspnoea can occur at dramatically different peak pulmonary arterial pressures during different types of exercise. This argues against pulmonary arterial haemodynamic screens for potential agents of heart failure. If non-haemodynamic peripheral factors both limit exercise and are responsible for generation of symptoms, then other non-haemodynamic modes of action may be effective for relief of symptoms in chronic heart failure.

Muscle fatigue in chronic heart failure

Muscle fatigue can arise due to alterations in the supply of oxygen to exercising muscle or to a change in the muscle itself. Skeletal muscle can be abnormal in terms of total mass, structure, or metabolic or contractile function. Although restrictions in peak blood flow to exercising muscle have been shown in heart failure, it is not clear to what extent this is due to persistent vasoconstrictor drive, impaired endothelial dependent vasodilator function, or a reduction in either muscle capil-
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Figure 1  The muscle hypothesis of chronic heart failure. In this hypothesis an initial reduction in left ventricular function activates catabolic and reduces anabolic factors that cause skeletal myopathy. This in turn leads to exercise intolerance and sympathoexcitation, and through the combined effects of a persistent catabolic state and of profound inactivity, further worsen skeletal muscle structure and function, and may eventually lead to a progressive effect on remodelling of the left ventricle.

larity or muscle volume. Although there is agreement that in heart failure the total peak blood flow to an exercising limb is reduced, where the flow is expressed per unit muscle volume (such as by plethysmography) the results are contradictory. This suggests that an important part of the reduced flow may be due to a deficiency in the total bulk of muscle. Abnormalities of skeletal muscle histology,11 mitochondria,12 oxidative enzyme activities,13 and high energy phosphate handling14 have all been described in chronic heart failure, as has early muscle fatigue.15 Any one of these abnormalities could explain exertional fatigue in patients with heart failure, and undoubtedly a combination of factors is operative in some patients (fig 1). We have recently shown that one of the best correlates with exercise limitation in chronic heart failure is a measure of leg muscle volume,16 suggesting that some of the qualitative alterations of muscle may also be a reflection of skeletal muscle wasting. These structural and metabolic changes in skeletal muscle could also explain the delay in improvement in symptoms and exercise tolerance after effective haemodynamic improvement with ACE inhibitors17 or even cardiac transplantation.18

Dyspnoea in chronic heart failure
Dyspnoea is a difficult sensation to quantify and even more difficult to explain.19 There is a fairly close relation between the perception of dyspnoea and minute ventilation, perhaps through a perception of the respiratory effort required to generate and maintain a certain level of ventilation during exercise. There may also be a direct appreciation of the chemoreceptor drive to breathing or of the level of respiratory stimulating factors such as arterial lactate, pH, potassium, adenosine, carbon dioxide, and other unknown factors.

The normal mechanisms whereby ventilatory effort is matched to muscular work are imperfectly understood. Even less is known of the pathophysiology that produces both an excessive sensation of breathlessness and an exaggerated ventilatory response in heart failure.20 It has been argued that the increased ventilation seen during exercise in patients with heart failure must indicate the presence of wasted ventilation in the form of dead space ventilation because there is no substantial fall in arterial blood concentrations of CO₂. There is, however, often a modest fall in arterial CO₂ during exercise, and arterial O₂ rarely if ever falls in stable heart failure, so the factor that initiates and maintains the increased ventilatory response is not obvious. Something other than CO₂ retention or hypoxia must be driving both the sensation of breathlessness and the resultant exaggerated ventilatory response. We have studied this response by looking at the ventilatory equivalent for CO₂ (the slope between ventilation and the rate of CO₂ production) during exercise.21 This slope is increased in heart failure and the increase in slope correlates well with the severity of heart failure as assessed by peak O₂ consumption (fig 2).

It is not known which of several independent and complementary ventilatory control systems can account for the increased ventilation in heart failure, but early lactic acid release and acidosis, potassium release, respiratory muscle fatigue, non-asthmatic bronchial constriction, and a muscle ergoreflex22 have all been proposed. Circulating factors including lactate and acidosis do not seem necessary for this response and differences in potassium release do not seem to account for the altered response in heart failure (unpublished observations).

The frequent coexistence of dyspnoea and fatigue and the delay in resolution of dyspnoea after haemodynamic improvement raise the possibility that structural abnormalities of skeletal muscle may be involved in the generation of dyspnoea. Dyspnoea and increased ventilation could be manifestations of abnormalities in respiratory musculature similar to
those seen in skeletal muscle. Early respiratory muscle fatigue and histological changes in the diaphragm have been seen in chronic heart failure. An alternative explanation is that skeletal muscle signals contribute directly to the perception of both muscle fatigue and dyspnoea, not only through the excessive release of blood borne metabolic factors, but possibly also through an exaggerated neural signal from the muscle.

Mechanisms of skeletal myopathy in chronic heart failure

The cause of the changes in skeletal muscle in chronic heart failure is not clear, but research on the beneficial effects of exercise training on skeletal muscle function in heart failure has suggested that inactivity may play a part. Metabolic or hormonal derangements that favour catabolism over anabolism may also contribute to the myopathy. Possibilities include the release of tumour necrosis factor, abnormalities in the handling of intracellular thyroid hormone, and insulin resistance, as well as intestinal malabsorption. There is little agreement as to whether chronically impaired skeletal muscle flow is an important cause of the myopathy of heart failure, but the dissimilarity between the muscle changes of heart failure and the increased oxidative capacity seen in patients with peripheral vascular insufficiency make this unlikely. Similar changes are by contrast seen in chronic lung disease suggesting that many chronic disease processes involving prolonged inactivity may lead to a similar syndrome. The cachexia of neoplastic disease has some similarities with cardiac cachexia, and this syndrome can also be associated with unexplained dyspnoea.

Importance of skeletal myopathy in heart failure: the skeletal muscle ergoreflex

Small myelinated and unmyelinated fibres arise from poorly characterised work sensitive receptors within skeletal muscle. These fibres travel in the lateral spinohalamic tract and have been shown to mediate an ergoreflex effect constituting an increase in sympathetic outflow producing vasoconstriction in distant vascular beds and possibly a small increase in heart rate. They are sensitive to the metabolic state of the muscle but not to ischaemia alone; rather they are work sensors. Excess of intramuscular potassium may stimulate their activation. We have recently shown that these ergoreceptors are active in patients with heart failure and that they also substantially contribute to the ventilatory response to exercise in both normal people and patients with heart failure. Also they are ideal candidate receptors for the perception of muscular fatigue and that component of dyspnoea that cannot be explained by circulating metabolites.

Although one short report suggested that the sympathetic nervous response to ergoreceptor activation in heart failure was blunted, the vastly different baseline levels of sympathetic tone in the two patient groups in this paper made it difficult to interpret the incremental effect of ergoreflex activation. The muscle ergoreceptors have the properties necessary to link the abnormal skeletal muscle function of chronic heart failure to the fatigue, dyspnoea, hyperpnoea, and sympathoexcitation characteristic of this condition.

The muscle hypothesis of chronic heart failure

We propose that abnormal skeletal muscle in chronic heart failure is a central abnormality that can account for much of the pathophysiology and symptoms of the condition. This hypothesis proposes another cycle of deterioration similar to those of neuroendocrine activation. A reduction in left ventricular function sets in motion a series of metabolic events that lead to wasting of skeletal muscle and resultant abnormalities of muscular metabolism and function. In response to early metabolic distress in exercising muscle an exaggerated ergoreflex activation occurs that is perceived by the patient as both muscle fatigue and dyspnoea. This leads to excessive reflex sympathetic vasoconstrictor drive to non-exercising muscle beds and an excessive ventilatory response to exercise. Resolution of these symptoms and the exaggerated reflex responses would depend on resolution of the abnormal function of skeletal muscle, and would, therefore, be delayed after haemodynamic correction and would be most noticeable in treatments associated with a specific improvement in muscle function or exercise responses. Figure 1 shows the proposed sequence of events.

Consequences of a muscle hypothesis for the generation of symptoms in heart failure

Valid areas for investigation (and perhaps intervention) exist to prevent the cycle of events described in this hypothesis. One aspect of this theory is that the skeletal myopathy, by exaggerating the sympathetic nervous responses to exercise, may actually contribute to the progression of the disease process through an exaggeration of peripheral vasoconstriction and hence the afterload that develops and through the direct myotoxic effects of prolonged sympathoexcitation. Ventricular remodelling and the propensity to ventricular arrhythmias are both thought to be partly a result of such excessive and prolonged sympathetic activation. The precise mechanism of the persistent sympathoexcitation of chronic heart failure has never been established. The arterial baroreflex loop is not an ideal explanation because the gain of this reflex is known to be dramatically impaired in heart failure, and hence it would be unable to respond acutely by increasing sympathetic outflow. Impaired baroreflex activity would also not lead to such sympathetic activation. Other putative mechanisms such as activation of the renal renin-angiotensin system, low pressure receptors, cardiocardiac
reflexes, or intracranial pressure flow autoregulation cannot adequately explain the maintenance and progression of sympathetic activation. The ergoreflex given its sympathoexcitatory role when muscle physiology is perturbed may play an important part in this activation. Once activated, sympathetic tone may contribute to further catabolism of skeletal muscle and a progression of the deleterious cycle.

Improving symptoms in chronic heart failure

As we know so little of the genesis of the limiting symptoms in chronic heart failure, it is perhaps not surprising that we cannot accurately predict which treatments will improve symptoms in this condition. There can be dramatic discrepancies between haemodynamic effects, exercise indices, and quality of life scores. Only a more detailed understanding of the integrated pathophysiology underlying both the progression of the disease and the complex interaction between exercise responses and symptoms can allow more effective treatments to be designed. To do this we may have to look beyond the conventional haemodynamic remedies for heart failure. The benefits of exercise training on skeletal muscle structure and function and the parallel reduction in dyspnoea, ventilatory abnormalities, and sympathetic overactivity suggest that specific muscle treatments may help in the management of chronic heart failure.

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