Drug therapy in chronic heart failure

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Chronic heart failure is widely recognised as a common and escalating problem that causes major disability and often shortens life. Diuretics and digoxin have formed the mainstay of treatment for many years. Clinical trials have demonstrated that angiotensin converting enzymes and β-blockers, in selected patients, improve symptoms and reduce mortality. Angiotensin-II antagonists and spironolactone may also have a role in certain individuals. Newer pharmacological approaches to the management of this complex disease are being developed, but await full evaluation.

Heart failure is a complex clinical syndrome, most simply defined as cardiac dysfunction associated with symptoms. Chronic heart failure (CHF) is a common debilitating illness, associated with a high mortality. It affects between 0.5 and 3 million people in the UK and 0.4%–2% of the total European population. Prevalence increases sharply with age, affecting 6%–10% of those older than 65 years. CHF is thus a disease of the elderly, due to the increasing longevity of our population, and increased survival rates from acute myocardial infarction. The average age at presentation is 76 years, with men being at a 75% higher risk of developing heart failure than women.

AETIOLOGY

The most common cause of CHF is systolic dysfunction due to ischaemic heart disease. Other causes include hypertension and valvular disease. The aetiology of heart failure has changed over the years, reflecting successful treatment of valvular heart disease and improved survival of patients with hypertension and ischaemic heart disease. A frequent associated problem is atrial fibrillation, which may occur in up to 30% of new cases of heart failure. Altogether 20%–40% of CHF patients have normal systolic function and may have diastolic dysfunction, characterised by impaired ventricular relaxation, again more common in the elderly. Heart failure usually comprises a combination of both systolic and diastolic impairment.

IMPACT OF CHRONIC HEART FAILURE

Heart failure has a major impact on the quality of a patient’s life, worse than angina or chronic airways disease. Despite considerable advances in the management of CHF, it remains associated with a very poor prognosis, worse than many forms of cancer. Survival is variable and depends on, among other things, left ventricular ejection fraction, degree of neurohormonal activation, and functional class. In the Framingham cohort (1948–88), the median survival after the onset of heart failure was 1.7 years in men and 3.2 years in women. In severe heart failure, survival can be less than 50% at one year. Survival rates are far worse in clinical practice than those demonstrated in recent clinical trials, reflecting the highly selected nature of the patients randomised.

The morbidity and healthcare costs of heart failure are enormous. In the UK CHF is estimated to cost 1%–2% of the total spent on health care, more than £400 million per year; 75% of this expenditure is on hospitalisation, with CHF responsible for approximately 5% of all hospital admissions. Readmission for heart failure within three months of the initial hospital admission occurs in about one third of patients. The costs of drug treatment are comparatively small.

This review concerns the pharmacological treatments available for CHF patients, specifically those with left ventricular systolic dysfunction. Drug treatment should relieve symptoms, improve quality of life, slow disease progression, prevent hospital admission, and ideally prolong active life (box 1). Certain patient subgroups may also benefit from treatment with one of the

Abbreviations: ACE, angiotensin converting enzyme; ANP, atrial natriuretic peptide; BNP, brain natriuretic peptide; CARMEN, Carvedilol ACE Inhibitor Remodelling Mild CHF Evaluation; CHARM, Candesartan in Heart failure: Assessment of Reduction in Mortality and Morbidity; CHF, chronic heart failure; CIBIS, Cardiac Insufficiency Bisoprolol Study; COMET, Carvedilol or Metoprolol European Trial; CONSENSUS, Co-operative North Scandanavian Enalapril Survival Study; COPERNICUS, Carvedilol Prospective Randomised Cumulative Survival; ELITE, Evaluation of Losartan in The Elderly; ENABLE, Endothelin Antagonist Bosentan for Lowering Cardiac Events; IMPRESS, Inhibition of Metaloprotease by Omapatrilat in a Randomised Exercise and Symptoms Study of Heart Failure; MERIT-HF, Metoprolol CR/XL Randomised Intervention Trial in Heart Failure; NEP, neutral endopeptidase; NYHA, New York Heart Association; OPTIMAAL, Optimal Trial in Myocardial Infarction with the Angiotensin II Antagonist Losartan; OVERTURE, Omapatrilat Versus Enalapril Randomised Trial of Utility in Reducing Events; PRAISE, Prospective Randomised Amlodipine Survival Evaluation; RAAS, renin-angiotensin-aldosterone system; RALES, Randomised Aldactone Evaluation Study; RESOLVD, Randomised Evaluation of Strategies for Left Ventricular Dysfunction; SENIORS, Study of the Effects of Nebivolol Intervention on Outcome and Rehospitalisations in Seniors with Heart Failure; SOLVD, Studies of Left Ventricular Dysfunction; Val-HeFT, Valsartan Heart Failure Trial; V-HeFT, Vasodilator Heart Failure Trial
various surgical and device treatment modalities available. Specific areas of current interest include cardiac resynchronisation therapy, implantable cardioverter defibrillators, and left ventricular assist devices.

PATHOPHYSIOLOGY OF CHF
Heart failure is a complex syndrome characterised by symptomatic impairment, usually with exercise intolerance due to breathlessness or fatigue, with associated evidence of cardiac dysfunction. Fluid retention may lead to pulmonary and peripheral oedema. The primary abnormality must be impaired cardiac function, which then sets off a train of compensatory mechanisms, some of which are ultimately detrimental, worsening symptoms and reducing survival.

(1) Haemodynamic mechanisms
In response to reduced cardiac output a number of central and peripheral changes occur (fig 1). Centrally there is increased left ventricular filling pressure due to venous vasoconstriction, which attempts to maintain cardiac output by the Frank-Starling mechanism. In addition, the resistance vessels constrict to maintain systemic arterial blood pressure in the face of a reduced cardiac output. Excessive cardiac filling pressures lead to an increase in lung stiffness and ultimately pulmonary oedema, causing breathlessness (fig 2). Heart rate increases and mechanisms to increase inotropic support are activated. Cardiac dilatation and hypertrophy also occur. Remodelling occurs when the ventricle dilates becoming less spherical, resulting in a further fall in ejection fraction.

Changes also occur in the periphery. Peripheral vasoconstriction varies in different vascular beds; resistance in the coronary and cerebral vessels changes little, whereas that to the skin, gut, kidneys, and skeletal muscle increases.

Reduced blood flow to these tissues contributes to symptoms, particularly to skeletal muscle, which is a major cause of fatigue during exercise. Ultimately, the increased peripheral vascular resistance increases the load against which the heart ejects blood and this also has a detrimental effect on cardiac output.

(2) Activation of neurohormonal mechanisms
The mechanisms responsible for these haemodynamic changes are complex, but include the activation of neurohormonal systems. These systems are of considerable interest, as although they initially support cardiovascular homoeostasis, they ultimately appear to be harmful (fig 3).

The renin-angiotensin-aldosterone system (RAAS) is activated in heart failure, particularly in those patients taking diuretics, leading to increased concentrations of renin, angiotensin II, and aldosterone. Angiotensin II is a powerful vasoconstrictor of the systemic circulation that also inhibits vagal tone, promotes aldosterone production, and stimulates norepinephrine release from sympathetic nerve terminals. This leads to sodium and water retention, and potassium secretion via the kidneys and sweat glands. Angiotensin II also has important effects on cardiac myocytes and may contribute to the endothelial dysfunction seen in CHF patients. An important and often overlooked aspect of neurohormonal activation in heart failure is that there is a dialogue between the RAAS and the sympathetic nervous system, which is also activated. Increased activity of either system promotes the other.
Heart failure patients have an activated sympathetic nervous system and elevated concentrations of circulating catecholamines. Clinically, this manifests as tachycardia and signs of peripheral vasoconstriction. The adrenergic activation is initially useful, as it helps to maintain an acceptable cardiac performance, but long term adrenergic activation is detrimental—it is associated with exercise intolerance, haemodynamic abnormalities, and increased mortality. Increased sympathetic tone can potentiate renin and angiotensin activity, leading to salt and water retention, arterial and venous constriction, and increased ventricular preload and afterload. By increasing heart rate and promoting coronary vasoconstriction, catecholamine excess can also reduce myocardial blood flow. Norepinephrine is potently cardiotoxic and results in cardiac myocyte injury in concentrations found in the failing heart.

**MEDICAL TREATMENT OF HEART FAILURE**

Considerable advances have been made in the drug treatment of heart failure over the past few decades. The basic principles of CHF treatment are:

- To reduce the workload on the heart during episodes of acute decompensation through rest, salt and water restriction, and drug treatment.
- To suppress the harmful effects of the neuroendocrine compensatory mechanisms (RAAS, sympathetic nervous system and, in the severe stages of disease, vasopressin release).

Basic advice to patients should include information about their condition, as well as the importance of stopping smoking, reducing salt and alcohol consumption, monitoring weight and fluid intake, and exercising regularly. It must be remembered that heart failure rarely occurs as an isolated disease process and that management is often complicated by coexisting factors. The prevalence of diabetes, chronic lung disease, renal dysfunction, depression, and peripheral vascular disease is relatively high in the heart failure population. Often, elderly patients have comorbid arthritis or neurological disorders. There is evidence that specialist heart failure clinics in primary or secondary care, specialist nurses in the community, and multidisciplinary teams can improve the quality of life of patients with CHF and reduce the need for hospital readmission.

**PREVENTATIVE DRUG TREATMENTS**

Investigating the underlying cause of heart failure helps target effective treatment. Secondary prevention measures, including aspirin and intensive treatment of hypertension and dyslipidaemia, improve outcomes in patients with ischaemic heart disease, the commonest cause of CHF in the western world. Optimal blood glucose control in patients with diabetes should be encouraged using metformin, sulphonylureas and insulin as required. Warfarin may be beneficial in selected patients with heart failure—for example, those with atrial fibrillation or a left ventricular thrombus with no contraindication to chronic anticoagulation. Amiodarone and dofetilide appear safe and effective antiarrhythmic agents in CHF patients with atrial fibrillation. It has also been recommended (though not evidence based) that all patients with CHF receive immunisation against influenza (annually) and pneumococcus (once).

**DIURETICS**

Sodium and water retention is the hallmark of CHF and diuretics are mandatory treatment for patients with pulmonary or peripheral oedema. There are no alternative treatments that can replace diuretics once a patient has developed congestion. Loop diuretics revolutionised the treatment of heart failure, despite having detrimental effects on lipids, urate, and glucose metabolism. Loop (and thiazide) diuretics have not been evaluated in long term trials looking at mortality outcome, but improvements in cardiac function, exercise tolerance, and symptoms have been reported in a few short term studies. In severe heart failure, thiazide diuretics and metolazone have a synergistic effect with loop diuretics and may be used in combination. Diuretics activate the RAAS and should not be used in isolation, but should be given with an angiotensin converting enzyme (ACE) inhibitor.

Once fluid retention has resolved, treatment with diuretics should be maintained to prevent recurrent oedema, which may require frequent adjustments to the dose given. A clear explanation of their condition and the effect of diuretics, along with advice regarding weight monitoring and fluid...
intake, should enable informed patients to make appropriate changes to their diuretic dosage.

VASODILATORS
Improved survival in response to drug treatment was first shown in 1986 with the combination of isosorbide dinitrate and hydralazine.36 This combination has now been superseded by drugs that block the renin-angiotensin system. They may still have a role in patients who are intolerant of, or have contraindications to, ACE inhibitors. Other vasodilators, including the α-blocker prazosin37 and the calcium channel antagonist amlodipine,38 had neutral effects on mortality in patients with CHF.

DIGOXIN
Digoxin is a mildly positive inotrope (at higher concentrations) that also increases vagal tone and suppresses renin secretion from the kidneys (at lower concentrations). There is no prospective controlled trial of the effects of the addition of digoxin on symptoms. However, drug withdrawal studies have shown that digoxin improves exercise capacity, symptoms, and the need for hospital admission both on its own39 and when combined with an ACE inhibitor.40 The Digitalis Investigation Group reported no mortality benefit in CHF patients on digoxin treatment over three years.36 Digoxin may be used to control the ventricular response rate in those CHF patients with atrial fibrillation. It would also be reasonable to use digoxin in those patients who remain symptomatic despite adequate treatment with diuretics, ACE inhibitors, and β-blockers. Digoxin toxicity may occur with serum concentrations greater than 2 ng/ml, resulting in nausea and vomiting, cardiac arrhythmias, and neurological problems. Toxicity is more likely in the elderly and in those patients with renal impairment, hypothyroidism, and electrolyte disturbance or when used in combination with certain drugs, such as amiodarone, quinidine, or spironolactone. All other positive inotropes have had detrimental effects on outcome (see below).

ANGIOTENSIN CONVERTING ENZYME INHIBITORS
ACE inhibitors prevent the conversion of angiotensin I to angiotensin II, and also inhibit bradykinin degradation. ACE inhibitors shift the balance between the unwanted effects of angiotensin II and the beneficial vasodilatory and natriuretic effects of bradykinin (fig 3). The evidence that ACE inhibitors improve survival and symptomatic wellbeing in all grades of heart failure is now incontrovertible.34 35 36 It is well established that ACE inhibitors have beneficial effects in both the treatment and the prevention of heart failure, strongly supporting the neurohormonal theory.

In 1987, the double blind placebo controlled CONSENSUS study demonstrated a significant mortality benefit with enalapril (up to 40 mg/day) in patients with severe heart failure (26% enalapril v 44% placebo, p = 0.002).37 A significant improvement in the New York Heart Association (NYHA) classification, together with a reduction in heart size and a reduced requirement for other heart failure medication, was also observed in the enalapril group. The SOLVD trial provided further evidence of a mortality and morbidity reduction with enalapril (up to 20 mg/day) in patients with left ventricular dysfunction (≤ 35%) and mild to moderate CHF.38 Large, prospective, randomised double blind trials have demonstrated mortality and morbidity benefits in patients with symptomatic heart failure39 40, in those with asymptomatic left ventricular dysfunction,41 42 and in patients with left ventricular dysfunction complicating acute myocardial infarction.43 44 45 They have also been shown to reduce the need for hospital admission and have proved to be a highly cost effective treatment. Higher doses (32.5–35 mg/day) of lisinopril are more effective than lower doses (2.5–5 mg/day) in reducing the combined endpoint of death and the need for hospital admission.46

Concerns about ACE inhibitors, including the risk of hypotension and renal dysfunction, may lead to a reluctance of some general practitioners to start the drugs, despite a good understanding of their benefits.47 48 Contraindications to the use of ACE inhibitors include angio-oedema or anaphylaxis on previous exposure, pregnancy, and bilateral renal artery stenosis. Many of the common adverse effects, including hypotension, renal dysfunction, and hyperkalaemia can be prevented by careful initiation and titration of treatment, with subsequent monitoring of blood pressure, urea, creatinine, and electrolytes. A dry cough occurs in about 10% of patients, and is the most common reason for ACE inhibitor withdrawal. It may improve with time, or by reducing the dose used.

Angiotensin II blockers (AT1 antagonists)
Angiotensin II acts via a family of cell bound angiotensin receptors. The AT1 receptor has been shown to mediate the detrimental effects of angiotensin II in patients with heart failure and angiotensin II antagonists block the AT1 receptor. ACE is identical to kininase 2, a protease enzyme partly responsible for metabolism of the inflammatory mediator bradykinin. ACE inhibitors increase bradykinin levels, which may have beneficial vasodilator and fibrinolytic actions. Bradykinin is also thought to be responsible for the troublesome cough and angio-oedema seen in some patients.

There is also evidence of breakthrough angiotensin II activity as a consequence of alternative non-ACE dependent pathways (for example, chymase) despite therapeutic doses of ACE inhibitors.49 50 This is avoided with AT1 antagonists by virtue of their selective action on the angiotensin II type I receptor. It was hoped that these effects would be beneficial in patients with heart failure. Two randomised studies in patients with mild to severe heart failure suggested that losartan (an angiotensin II antagonist) and enalapril have similar effects on haemodynamics and exercise capacity.51 52

The ELITE study was designed to assess drug safety and tolerability and randomised 722 CHF patients over the age of 65 to losartan (50 mg once daily) or captopril (50 mg three times daily).53 After 48 weeks, losartan demonstrated a lower incidence of side effects and drug withdrawal than captopril and a significant reduction in the secondary endpoint of all cause mortality, mainly due to a reduction in the incidence of sudden death. However, the study was too small to reliably assess survival effects. The complex RESOLVD study comparing candesartan and enalapril did not support these findings.54

ELITE II compared the effects of captopril and losartan in 3152 patients with mild to severe heart failure and demonstrated no difference in mortality between the drugs.55 The Val-HeFT trial tested the hypothesis that the combination of valsartan in addition to an ACE inhibitor and β-blocker would improve outcome.56 A total of 5010 patients with symptomatic CHF were randomised to valsartan or placebo and followed up for an average of 1.9 years. Valsartan did not reduce mortality, but did significantly reduce the combined endpoint of mortality and morbidity, mainly by reducing heart failure admissions. The greatest effect seemed to occur in those patients who were not receiving an ACE inhibitor at baseline,57 and those patients who received triple neurohormonal blockade (with an ACE inhibitor, a β-blocker, and valsartan) appeared to do poorly on subgroup analysis. These findings need further investigation and should be interpreted with caution.

OPTIMAAL enrolled 5477 patients over 50 years old with confirmed myocardial infarction and acute heart failure...
during the acute phase, or a new Q-wave anterior infarction or reinfarction. Patients were randomised to losartan (titrated up to 50 mg once daily) or captopril (titrated up to 50 mg three times daily) and followed up for a mean of 2.7 years. There was a non-significant trend towards reduced all-cause mortality (the primary endpoint) in favour of captopril. Consequently ACE inhibitors should remain first choice treatment in patients following complicated myocardial infarction.

At present, no angiotensin II antagonist is licensed in the UK for use in heart failure, though valsartan has received its license in the United States. They may have a role in patients unable to tolerate unwanted effects with ACE inhibitors, particularly cough. The ongoing CHARM studies using candesartan may establish the role of the angiotensin II receptor blockers in CHF.

β-BLOCKERS

Prolonged neurohormonal activation has unwanted effects on the haemodynamics of the failing heart and also leads to apoptosis, arrhythmia, and reduced myocardial blood flow. For these reasons, drugs that interfere with the sympathetic nervous system have been extensively investigated in patients with CHF over the past 30 years. Over 15,000 patients have now been enrolled in placebo controlled trials looking at the effects of β-blockade in patients with congestive heart failure. These studies have confirmed, that like ACE inhibitors, long-term treatment with β-blockade can reduce symptoms, improve clinical status, and reduce the risk of death as well as the combined risk of death or hospitalisation.

Metoprolol is a second generation β1-selective antagonist with no intrinsic sympathomimetic activity. In CHF patients, it has been demonstrated to improve cardiac function, left ventricular remodelling and capacity for exercise, and lessen the symptoms of heart failure. The first placebo controlled, multicentre trial with a β-blocker in CHF was the Metoprolol in Dilated Cardiomyopathy trial. A total of 383 patients with symptomatic heart failure secondary to idiopathic dilated cardiomyopathy were randomised to metoprolol or placebo (up to 150 mg/day in divided doses) for 12 to 18 months. There was a non-significant, 7.2% absolute risk reduction in the combined endpoint of all-cause mortality and patient deterioration requiring listing for heart transplantation (20.1% placebo vs 12.9% metoprolol, p = 0.058) with a trend towards increased all-cause mortality in the metoprolol arm. A larger placebo controlled mortality study, the MERIT-HF study, followed randomising 3991 patients with CHF (NYHA II-IV) and an ejection fraction <40% to placebo or metoprolol CR/XL (controlled release/extended release up to 200 mg once daily—not licensed for use in patients with CHF in the UK) after a two week single blind run-in period. The study was stopped early after a mean follow up of one year on the recommendation of the Independent Safety Committee due to a significant 3.8% absolute risk reduction in all-cause mortality in the metoprolol group (7.2% vs 11%; adjusted p = 0.0062). Importantly, the average dose of metoprolol achieved in the MERIT-HF trial (159 mg) was larger than in the Metoprolol in Dilated Cardiomyopathy trial (108 mg).

Carvedilol is a non-selective β-adrenoceptor antagonist that also blocks α-receptors. Unlike other β-blockers it exerts novel antioxidant and antiproliferative effects, which may contribute to its beneficial effects in patients with cardiac disease. The United States Carvedilol Heart Failure Trials Program was an amalgamation of four separate multicentre trials involving 1094 patients with mild, moderate, and severe heart failure and ejection fraction <35%. Patients on stable background therapy were entered into one of the four placebo controlled studies on the basis of their ability to complete a six minute walk test. A predefined analysis of the combined data from all four trials demonstrated an overall mortality rate of 7.8% in the placebo group and 3.2% in the carvedilol group, during an average follow up of 6.5 months. The Australia/New Zealand Carvedilol Heart Failure Research Collaborative Group study enrolled patients on stable treatment including diuretics, an ACE inhibitor, and digoxin. The first phase was a six month submaximal exercise trial that enrolled 415 patients with stable heart failure of an ischaemic aetiology after a two to three week open-label run-in phase to carvedilol or placebo (up to 50 mg/day). The second phase, lasting an average of 19 months, demonstrated continued benefits on left ventricular dimensions and function and a 12.3% absolute risk reduction in the combined endpoint of mortality or hospitalisation (62.5% placebo vs 50.2% carvedilol, p = 0.02). While both the United States and Australia/New Zealand studies demonstrated a reduction in the combined endpoint of death and hospitalisation, only the United States study found a significant mortality reduction. Carvedilol is now licensed in the UK for reducing the clinical progression of heart failure and lowering the combined risk of morbidity and mortality.

The COPERNICUS Study Group recently published the results of an international multicentre trial looking at the effects of carvedilol on survival in severe CHF. A total of 2289 patients with heart failure symptoms at rest or on minimal exertion, on standard therapy, with an ejection fraction <25%, were randomised to carvedilol (n = 1136) or placebo (n = 1133) for a mean period of 10.4 months. Randomisation was stopped early in March 2000 on the recommendation of the Data and Safety Monitoring Board, because of a significant mortality benefit with carvedilol treatment. There was an estimated 7.1% absolute reduction in the risk of death in the carvedilol group at one year (18.5% placebo vs 11.4% carvedilol, p = 0.0014). The annual mortality rate of 18.5% in the placebo group, compared for instance with >60% in the CONSENSUS study, is lower than would be expected in non-selected patients.

CIBIS-II randomised 2647 patients with moderate to severe heart failure (mostly NYHA class III) and a left ventricular ejection fraction of 35% or less, on standard therapy with diuretics and ACE inhibitors, to placebo or bisoprolol a (β1-selective β-blocker) up to 10 mg a day. The Data and Safety Monitoring Board stopped the trial early, after a mean follow up of 16 months, because bisoprolol demonstrated a significant mortality benefit. There was an absolute risk reduction of 5.5% in all-cause mortality with bisoprolol compared with placebo (11.8% vs 17.3%, p=0.0001), as well as significant reductions in hospitalisations for heart failure, and in the combined endpoint of cardiovascular deaths and hospital admission for cardiovascular events.

In summary, patients with stable symptomatic heart failure due to left ventricular systolic dysfunction should be considered for β-blocker therapy once treatment with diuretics and ACE inhibitors has been optimised. Established contraindications include decompensated heart failure, reversible airways obstruction, advanced heart block, and symptomatic bradycardia or hypotension. Initiation and uptitration of β-blockers can be difficult in heart failure patients, often requiring persistence and specialist knowledge to enable target doses to be achieved. The patient’s heart rate, blood pressure, and clinical status should be monitored closely. It is important to reinforce that transient side effects are common and that the beneficial effects of β-blockers are sometimes only achieved after a number of months. In clinical practice withdrawal rates are greater than those seen among the highly selected patients enrolled in clinical trials.
Currently there are a number of studies addressing some unanswered questions regarding treatment with β-blockers. Patients with symptomless left ventricular dysfunction are being enrolled in the CARMEN study with carvedilol. The SENIORS study will compare the effects of nebivolol, a novel β₁-selective blocker with vasodilatory properties, against placebo in elderly patients with heart failure. The COMET trial, comparing carvedilol and metoprolol is the first large scale, event driven trial comparing non-selective and selective β-blockers. Over 3000 patients have been randomised and the results of the study should be available in 2003.29

**SPIRONOLACTONE**

Aldosterone levels commonly remain elevated in patients on an ACE inhibitor and may contribute to the worsening of heart failure. Spironolactone, a potassium sparing diuretic, is a competitive antagonist of aldosterone and has been shown to have additional beneficial effects in patients already treated with an ACE inhibitor.30 The RALES trial randomised 1663 patients with severe heart failure who were on standard therapy with diuretics, an ACE inhibitor with or without digoxin, to spironolactone or placebo. Spironolactone (25–50 mg a day) reduced absolute mortality by 11% (46% v 35% spironolactone) at a mean of two years. It also resulted in a significant decrease in the frequency of hospital admissions for worsening heart failure. Despite low rates of serious hyperkalaemia during the trial, the potential risk in clinical practice is greater, and renal function and potassium levels should therefore be monitored in patients on the combination of an ACE inhibitor and spironolactone.

**CALCIUM CHANNEL BLOCKERS**

The evidence that calcium channel blockers should be used for treating heart failure remains limited, but the non-rate limiting calcium antagonists appear safe and could be used to treat hypertension and angina in patients who also have CHF. The V-HeFT III study demonstrated a small symptomatic benefit with felodipine,31 while the PRAISE studies showed a neutral effect on mortality with amiodipine in patients with severe (NYHA III-IV) heart failure.32

**HARMFUL TREATMENTS**

It is important to be aware of medical treatments associated with adverse outcomes in CHF patients. Positive inotropes including the phosphodiesterase inhibitors milrinone,33 enoximone,34 and vesnarinone35 and xamoterol, an oral partial β₂-agonist36 all improved exercise tolerance and quality of life at the expense of increased early mortality. Flosequinan,37 an inodilator, and ibopamine,38 a dopamine agonist had similar outcomes. Class I antiarrhythmic drugs should also be avoided and patients should be advised to limit their use of non-steroidal anti-inflammatory agents that may aggravate heart failure.

**NEWER AGENTS**

There are a number of agents in the early stage of development, which may have a potential role in the management of CHF. The majority of these drugs attempt to restore the neurohormonal balance that is disrupted in patients with heart failure (fig 3). Atrial and brain natriuretic peptide (ANP and BNP) are hormones that have vasodilator, natriuretic, diuretic, and RAAS suppressing actions. ANP and BNP are degraded by the enzyme neutral endopeptidase (NEP or nephrilysin), and inhibitors of this enzyme have been investigated as possible treatments for CHF. In early studies NEP inhibitors demonstrated beneficial haemodynamic and neurohormonal effects, and improvements in exercise tolerance.70 71 However toxicity related to one agent, ecadotril, stopped the further development of these agents.72 Drugs that inhibit both ACE and NEP, including omapatrilat, have also demonstrated beneficial haemodynamic and neurohormonal effects in CHF patients.73 Omapatrilat caused fewer adverse events than lisinopril in the prospective double blind IMPRESS trial.74 A total of 573 patients with NYHA II-IV heart failure, who were already receiving an ACE inhibitor, were randomised to 20 mg lisinopril daily or 40 mg omapatrilat daily for 24 weeks. There was no significant difference in the primary endpoint, the improvement in treadmill exercise time, between the two groups. OVERTURE was a prospective, double blind trial that randomised over 5500 CHF patients to omapatrilat or enalapril.75 There was no significant difference in the primary endpoint, death, or hospitalisation for worsening heart failure, between the two groups.

Endothelin-1, a 21 amino acid peptide is a very powerful vasoconstrictor with anti-natriuretic, anti-diuretic, and positive inotropic effects. Plasma concentrations of endothelin-1 are increased in heart failure and higher concentrations are associated with worse symptoms and clinical outcome, more impaired haemodynamics and reduced left ventricular systolic function.76–79 Some endothelin receptor antagonists, including tezosentan, demonstrated favourable acute and chronic haemodynamic effects.80–83 The ENABLE program randomised a total of 1613 patients in Europe and Australia
(ENABLE 1) and North America (ENABLE 2), with NYHA class III-IV heart failure to the endothelin receptor antagonist bosentan 125 mg twice daily or placebo. Disappointingly, bosentan was no better than placebo in reducing the combined endpoint of death or hospitalisation for worsening heart failure, but was associated with an early increased risk of worsening heart failure.

There is also interest in other agents including erythropoietin analogues, cytokine antagonists, lower dose enoximone combined with a β-blocker, and arginine vasopressin antagonists. Further adequately powered trials enrolling patients typically of evidence in elderly patients over 70 years old, who make up a large majority of patients with heart failure, and women remain under-represented in most heart failure trials. Further adequately powered trials enrolling patients typically seen in daily clinical practice are needed to address these issues.

**UNANSWERED QUESTIONS**

There is not yet clear evidence of benefit from any of these treatments in patients with diastolic dysfunction—that is, overt heart failure with preserved left ventricular function. The optimal treatment of patients with asymptomatic left ventricular dysfunction also remains unclear. There is a lack of evidence in elderly patients over 70 years old, who make up a large majority of patients with heart failure, and women remain under-represented in most heart failure trials. Further adequately powered trials enrolling patients typically seen in daily clinical practice are needed to address these issues.

**CONCLUSIONS**

The pharmacological treatment of patients with CHF should include preventative measures addressing the underlying cause of the disease. Loop (and/or thiazide) diuretics are mandatory in those patients with evidence of pulmonary or peripheral congestion. If tolerated, ACE inhibitors and β-blockers reduce mortality rates and can prevent the progression of symptoms and the need for hospital in CHF patients. Low dose spironolactone increases survival in patients with severe heart failure (NYHA III-IV) with ongoing symptoms despite standard therapy. Further evidence is needed before the role of angiotensin-II antagonists can be confirmed. Effective implementation of these treatments on an individual basis should greatly improve the outcome of patients who develop heart failure.

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**Case report**

A 64 year old man suffered a non-Q-wave inferior wall myocardial infarction in 1994. He had a subsequent anterolateral ST elevation myocardial infarction in 1996 complicated by a ventricular fibrillation arrest and prolonged admission to the intensive treatment unit. After discharge he developed effort angina and subsequently left heart catheterisation and coronary angiography were undertaken. Severe three vessel coronary artery disease and global left ventricular systolic dysfunction were demonstrated. He went on to have quadruple bypass grafting and made a good recovery despite methyccillin resistant Staphylococcus aureus wound infection. He was discharged home taking furosemide 80 mg daily, enalapril 10 mg twice daily, atorvastatin 20 mg daily, and digoxin 125 mg daily. Allopurinol 200 mg daily was also prescribed for gout.

He did well until 2002 when he became increasingly breathless on exertion and was admitted in September with pulmonary oedema. His loop diuretic was changed to bumetanide 3 mg daily, as this is more predictably absorbed from a congested gut than furosemide, and metolazone 2.5 mg daily was added to increase his diuresis. After discharge he remained symptomatic and attempts were made to commence him on bisoprolol 1.25 mg daily. However he had symptomatic hypotension with a systolic blood pressure of 80 mm Hg and despite two attempts was unable to tolerate the drug. After a second admission with pulmonary oedema in October, spironolactone 25 mg daily was added to his drugs with careful monitoring of his renal function and potassium. He had mild renal impairment (creatinine 136 μmol/l), but his potassium remained within normal limits.

He remained limited on minimal exertion and was not considered suitable for heart transplantation. A 12-lead electrocardiogram demonstrated sinus rhythm, normal PR interval, left axis deviation, and left bundle branch block with a QRS duration of 180 ms. A transthoracic echocardiogram demonstrated a dilated left ventricle with an end diastolic dimension of 7.8 cm. There was severe global impairment of systolic function and septal motion was dyskinetic. He is currently awaiting implantation of a biventricular pacemaker system in an attempt to improve his symptoms.
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