Experimental approaches to the prophylaxis and treatment of ischaemic heart disease

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Summary
This paper describes some of the basic hypotheses on which our research in the cardiovascular field has been based. The difficulties and shortcomings of the experimental models used and our reliance on guidance from clinical science has been emphasized. The properties, effects and mode of action of the drugs resulting from our cardiovascular research have been described.

Introduction
The purpose of this paper is to describe our research aims and their implementation in the field of ischaemic heart disease. The research programme is divided into two areas. Firstly, the discovery of a means of controlling the development of coronary artery disease, and secondly the devising of agents that would control the complications of established coronary artery disease (Fig. 1). There are two requirements for effective research in the field of coronary artery disease. In the first instance, there is a need for reliable animal models of the disease and its complications. Whilst many forms of experimental atherosclerosis are available, their relevance to the human disease is debatable, particularly in relation to the development of therapeutic agents. The second requirement is an agreed understanding of the pathogenesis of coronary artery disease and its attendant complications. Unless clinicians can delineate realistic therapeutic goals, based on a knowledge of the causative mechanisms underlying the disease, then experimental research may easily proceed along misguided lines.

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Fig. 1. The basis of the separation of experimental research in ischaemic heart disease.
Prophylaxis of coronary artery disease

The experimentalist in this field is in the difficult position of wishing to devise an experimental model to test synthetic substances whilst there is disagreement as to what a particular animal model really means in terms of human disease, what therapeutic target is worthwhile, and what type of agent might be devised to achieve such a target. Whilst the mechanism of formation of the atherosclerotic plaque is debatable, clinical and epidemiological research suggests that certain factors predispose to plaque formation (Katz & Stamler, 1953). An obvious approach to the problem of the atherosclerotic plaque could be to delineate all the factors that favour its growth, and attempt to attenuate or even reverse the trend. In 1955, when our research in this field started, the factor that was most widely considered worthy of control was elevated serum lipids, particularly serum cholesterol. The association between hyperlipidaemia and ischaemic heart disease is based on four types of observation: (1) that arterial lesions contain lipid, (2) that induced hyperlipidaemia in animals leads to arterial lesions rich in lipids, (3) that patients with the symptom complex of ischaemic heart disease often have abnormal serum lipid levels, and (4) that hypercholesterolaemia at least is associated with increased risk of developing ischaemic heart disease (Oliver, 1967a). At that time, our research aim, therefore, was to discover an agent that would lower serum cholesterol and triglycerides effectively, and cause no serious side effects. Such an agent should meet most of the following requirements: (a) it should have a sustained and consistent effect in reducing elevated serum cholesterol and triglycerides or more correctly, it should consistently lower the SF0-20 and SF20-400 lipoprotein fractions, (b) this lowering of the serum lipids should be associated with a lowering of the accumulation of lipids in tissues, (c) there should be an increase in the excretion of neutral sterols in the stool, (d) the agent should not alter the patient's way of living, (e) there should be few toxic effects associated with the achievement of the hypolipidaemic action, (f) there should be few side effects, (g) precursors of cholesterol should not accumulate in the body, (h) an understanding of the mode of action of the drug is necessary. The reason for laying down such stringent requirements is that the final aim of treatment may be to give the agent to healthy hyperlipidaemic individuals who are assumed to have a greater than usual risk of developing vascular disease and who may therefore need to take the drug for the rest of their lives.

More recently, additional requirements have been laid down. In addition to demonstrating that an agent must lower elevated serum lipids effectively and safely, it is necessary to show that, if it is given in the pre-symptomatic or established phases of ischaemic heart disease, it alters the mortality from coronary heart disease. Some knowledge of its effect on the atherosclerotic plaque is also necessary.

Clofibrate*

Clofibrate meets many of the requirements laid down above, though naturally answers are still awaited from many of the studies with this compound. The activity of clofibrate was discovered in 1955 by Thorp, whilst examining a series of aryloxyisobutyricates with the following basic chemical structure (Fig. 2). Thorp showed that they all possess a novel type of hypolipaemic activity in experimental animals (Thorp, 1955). In normal rats and dogs the serum cholesterol was lowered by 60–70% compared to that of controls, following administration of these compounds, and the response was maintained for as long as the compound was administered. The cholesterol returned to normal levels on stopping treatment. Initially, therefore, the testing procedure was simply to determine the effect of these drugs on serum cholesterol in a variety of animals. One compound, the p-chlorophenol aryloxyisobutyric acid (CPIB*) was selected for clinical evaluation. Studies in monkeys showed that the addition of androsterone potentiated the hypolipidaemic action of CPIB and therefore the first studies in man were carried out with clofibrate in combination with androsterone. Subsequently, it was found that an equivalent hypolipidaemic effect could be achieved in man without the addition of androsterone (Oliver, 1963).

The effect of CPIB in man is to reduce elevated levels of serum lipoproteins, particularly the SF20-400 fraction and to a lesser extent the SF0-20 fraction. Studies in a mixed population of patients with coronary artery disease show that elevated serum

* Clofibrate (‘Atromid-S’—ICI) is the ethyl ester of chlorophenoxyisobutyric acid (CPIB).

Fig. 2. The structure of aryloxyisobutyrates and clofibrate.
cholesterol levels are reduced by 15–30% and elevated serum triglyceride levels by 30–40% (Oliver, 1967b). These effects persist for as long as treatment continues, and no escape is observed in patients who respond adequately. In addition, clofibrate reduces alimentary lipaemia, and increases lipoprotein lipase activity (Hood, Bedding & Corlander, 1963). Serum free fatty acids are lowered, but are still responsive to catecholamine stimulation. Clofibrate does not cause accumulation either in blood or tissues of unsaturated precursors of cholesterol such as desmosterol, and it brings about a reduction in the total cholesterol pool (Nestel, Hisch & Couzens, 1965; Hellman et al., 1963). This is accompanied by an increased biliary excretion of neutral sterols (Mitchell, Truswell & Brontestewart, 1967). Raised plasma fibrinogen is reduced by clofibrate, and decreased fibrinolytic activity is enhanced. Platelet stickiness, platelet turnover and platelet survival times are altered towards normal values (Oliver, 1967b). In clinical terms, visible tissue deposits such as lipaemic exudates in the retina of diabetics with retinopathy, or skin xanthomata may regress following treatment with clofibrate for 6–12 months (Duncan et al., 1968; Mishkel, 1967). The initial regression may be achieved without a marked reduction in the serum cholesterol, which suggests that it may take several months to reduce a large cholesterol pool. Oliver (1967) has suggested that the principal indications currently for the use of clofibrate are (1) patients with hypercholesterolaemia and hypertriglyceridaemia with or without xanthomata, (2) patients with fasting latescent plasma, (3) patients with hypercholesterolaemia with or without a familial predisposition to vascular disease, (4) diabetic patients with lipaemic diabetic retinopathy, in order to prevent further deterioration in visual acuity.

**Mode of action of clofibrate**

Any explanation of the mode of action of clofibrate must take into account the following observations. (1) Clofibrate does not decrease cholesterol synthesis in the liver when added to liver slices in vitro. This suggests, therefore, that it does not have a direct action on cholesterol synthesis at the hepatic level (Avoy, Swyrd & Gould, 1965; Thorp & Waring, 1962). (2) Clofibrate does not lower serum cholesterol in animals which have been thyroidec-tomized (Westerfeld, Richert & Ruegamer, 1968). (3) The volume distribution of CPIB shows that 96–98% is confined to the albumin space which is 8–12% of the total body weight. It is bound to the anionic binding sites of albumin. (4) In experimental animals, CPIB is not detected in hepatic bile, and most of an oral dose is excreted in the urine as the glucuronide (Thorpe, 1963). The problem, therefore, is to explain how an organic acid that is fully ionized in plasma, and is highly albumin-bound, that is excreted as the glucuronide via the kidney and is not metabolized in the body, can exert such a wide variety of effects (Fig. 3). The multi-system of effects of CPIB can be explained on the hypothesis that physiologically important albumin-bound endogenous acids are displaced from albumin by CPIB. Thus the continuous administration of CPIB results in a new equilibrium state in which these endogenous acids are either displaced to weaker ionic binding sites on albumin, or displaced altogether from albumin to cause a small increase in the concentration of ‘free form’ in the circulation. The free form

![Fig. 3. Metabolism of clofibrate.](http://pmj.bmj.com/)

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**Fig. 3. Metabolism of clofibrate.**
of these substances is responsible for their physiological effects. Physiologically important substances that are bound to albumin, which have been shown to be displaced by clofibrate, are thyroxine and its metabolites, free fatty acids, and 17-ketosteroid sulphates as well as pyridoxal phosphate and tryptophan.

There is much experimental work to support this hypothesis.

**Clofibrate and thyroxine**

Chang et al. (1967) have confirmed that clofibrate depresses the thyroxine binding to human pre-albumin at a concentration of $6 \times 10^{-8}$ m. Thyroxine-binding globulin is unaffected. Studies by Thorp (Thorp & Waring, 1962), and more recently by Westfield et al. (1968) have shown that clofibrate increases the concentration of radio-labelled thyroxine in the rat liver by 40%. There is no increase in the metabolic rate or alteration in thyroid function in these rats. In the thyroidectomized rat, the action of clofibrate on serum cholesterol is abolished. Furthermore, it has been shown that in the rat, the maximal effect of CPIB on the serum cholesterol corresponds with the period of maximal thyroid adrenocortical function. Clofibrate causes an increase in the weight of the liver in the rat and monkey, which is due to increased synthesis of hepatic protein. This is accompanied by a reduction in hepatic glycogen (Platt & Thorp, 1966). These changes are identical with those observed when thyroxine alone is administered. Studies by Harrison & Harden (1966) in hypercholesterolaemic patients with severe myxoedema showed that the serum cholesterol was unaffected by clofibrate until a small supplement of thyroxine is added. The hypolipidaemic effect of thyroxine treatment is significantly enhanced by the simultaneous administration of clofibrate. If thyroxine alone is subsequently withdrawn, a rebound of serum cholesterol to the pretreatment level, or higher, occurs.

Apart from its effects on thyroxine distribution, clofibrate also reduces the transport of free fatty acids from the fat organ to the liver. Barrett in our laboratories has shown that in vitro, CPIB will reduce the rate of free fatty acid release from epididymal fat pads incubated in plasma, as well as reduce the peak levels of free fatty acids following the injection of adrenaline and noradrenaline (Barrett, 1969a). It also abolishes the increase in cholesterol and phospholipids following ACTH stimulation in dogs. These observations suggest that part of the hypolipidaemic action of CPIB is due to depression of free fatty acid transport (Thorp, personal communication).

In summary, it seems that clofibrate acts primarily by displacing physiologically important substances from anionic binding sites on albumin. As a result, there is an increased uptake of thyroxine by the liver. The increased hepatic thyroxine activity causes an increase in protein synthesis. The accompanying reduction in hepatic glycogen increases the hepatic requirement for an alternative energy substrate, i.e. free fatty acids. However, in the presence of clofibrate, there is a depression of circulating free fatty acid levels. The combined effects of a diminished supply of free fatty acids to the liver and an increase in concentration of hepatic thyroxine, results in a marked drop in circulating levels of $\text{SF}_{20-400}$ lipoproteins which is the most marked and consistent effect of clofibrate in man. This summary of the mode of action of clofibrate is almost certainly incomplete. Space does not permit a consideration of the possible role of other important substances such as pyridoxal phosphate, androsterone sulphate, dehydroepiandrosterone sulphate and tryptophan.

**Subsequent developments**

Some scepticism exists in regard to this view of the mechanism of action of clofibrate. If the theory is true, then it should be possible to discover other agents which also lower serum lipids because they displace physiologically active substances from albumin. In the past 5 years, our experimental approach to the lowering of serum lipids has been to test drugs *in vitro*. A search was made for compounds displaying a greater selectivity and intensity of effect on the binding of thyroxine to albumin and pre-albumin than that of clofibrate. Furthermore, we wished to discover compounds with a greater degree of persistence *in vivo* than clofibrate, so that a lower daily dosage could be used. The first tests consisted of studies of the water solubility and half-life, in the rat, of candidate compounds of the aryloxyisobutyrate series. It was found that diminished water solubility was accompanied by an increase in the *in vivo* half-life.

Experiments were then carried out to determine the effects *in vitro* of selected compounds on the albumin binding of thyroxine. It was found that the compound ICI 54,856 was more effective than CPIB in increasing the amount of free thyroxine in an *in vitro* albumin-thyroxine system.

![ICI 54,856](attachment:image.png)

The methyl ester of ICI 54,856 has been taken forward for both animal and human studies. Its action is similar to that of clofibrate, but because of its increased persistence, it is effective at about
1/100th of the dose of clofibrate in lowering blood cholesterol in rat, dog and man. Its half-life in man is 30–40 days. An effective dose in man is 0·1 mg/kg, giving a circulating level of 60 μg/ml. 60–90% of ICI 54,856 is excreted by the kidney, partly as the free acid and partly as the glucuronide conjugate. Preliminary clinical trials with this novel compound show that it not only lowers the SF\textsubscript{20-400} lipoprotein fraction effectively, but also the SF\textsubscript{0-20} fraction, which in certain cases is resistant to CPIB treatment. (Barrett, 1969a). This drug is, however, in very early stages of clinical evaluation, and it is still in the experimental phase. It illustrates how an understanding of the mode of action of a drug can lead to further discoveries and that careful in vitro studies can be of predictive value for drug action in man.

The treatment of the complications of coronary artery disease

Angina pectoris

There is no animal model for the study of angina pectoris, because it is a symptom and therefore is peculiar to man. Any experimental approach must be related to clinical experience. It is widely accepted that anginal pain results from a disproportion between myocardial work and myocardial oxygen supply. Most experimental effort in the past has been devoted towards developing therapeutic agents which will increase myocardial blood flow. Whilst substances with potent dilating effects have been found, their clinical success appears to have been limited. Unless the obstruction to blood flow in diseased coronary arteries can be removed, it is probably more rational to attempt to improve the mechanical efficiency of the pump, rather than to improve blood flow through diseased arteries. All therapy for angina pectoris aims at reducing the work of the heart for a given external work-load, and no drug has been discovered that permits more total cardiac work to be done before pain. Thus, both weight-loss and trinitrin exert their beneficial effects in angina by diminishing the work of the heart. Initially, trinitrin was thought to act primarily by coronary vasodilatation, but this is not now considered to be the primary mode of action (Gorlin et al., 1959). Cervical sympathectomy also lowers the work of the heart as shown by Apthorp, Chamberlain & Hayward (1964). In their group of eight anginal patients who underwent bilateral cervical sympathectomy for severe angina pectoris, several patients increased their exercise tolerance two to three-fold following this operation. Similarly, Lindgren (1950) has reviewed the effects of cervical sympathectomy in eighty anginal subjects, and shown that 52% showed marked or moderate improvement through this operation.

These studies emphasize the role of endogenous catecholamines in increasing cardiac work.

Black in our laboratories initiated studies aimed at specifically interfering, by chemical means, with catecholamine activity on the heart. It was hoped that an agent could be discovered which would diminish the effects of catecholamines in increasing cardiac work in those patients with impaired myocardial blood supply (Black, 1967). A clue as to the type of chemical structure to be sought was provided by the discovery of Powell & Slater (1958), that the dichloro analogue of isoprenaline would block the effects of catecholamines on the heart. Though dichloroisoprenaline was introduced to clinical studies, there is no evidence that it was studied in angina pectoris. Thus it was Black’s hypothesis which really provided the impetus rather than the discovery of dichloroisoprenaline. The first effective agent discovered was pronethalol (‘Alderlin’, ICI). It was introduced into clinical trial specifically to determine whether an agent which prevented the action of catecholamines on the heart would be of benefit in angina pectoris (Black & Stephenson, 1962). It was shown to increase exercise tolerance in angina pectoris, and also to improve the anginal state clinically (Dornhorst & Robinson, 1962; Alleyne et al., 1963) Subsequently, pronethalol was withdrawn because it was found to cause lymphomata in the thymus of certain strains of mice. It was replaced by propranolol (‘Inderal’, ICI) which has a different chemical structure and is more potent (Black, Duncan & Shanks, 1965).

Mode of action in angina pectoris

Stimulation of cardiac beta-receptors results in an increased rate and force of myocardial contraction, as well as speed of conduction within the heart. As a result there is an increase in the oxygen consumption of the heart. In the healthy heart, blood flow to the myocardium is not a limiting factor, and therefore the heart is able to increase its work without incurring any damage. However, where the blood supply is limited, then there is reason to suppose that benefit may accrue from attenuating changes in the rate and force of myocardial contraction.

The mode of action of propranolol in reducing cardiac work is complex. The more important features are: (1) reduction in heart rate, (2) reduction in the speed of myocardial fibre shortening, (3) possibly a reduction in after-load. All these effects will reduce myocardial oxygen demand. They may be offset if the heart increases in size, as is associated with cardiac slowing and a reduction of myocardial contractility. Furthermore, there may be an increase in the systolic ejection time. An increase in either of these parameters will increase myocardial oxygen demand. It is thus not possible to predict which action will predominate in
an individual patient, so that some will improve and others may not (Fig. 4). Intensive study of propranolol in angina pectoris indicates that many patients with angina pectoris improve their exercise tolerance, reduce their daily incidence of anginal pain, and diminish their requirements for glycerol trinitrate (Amsterdam, Wolfson & Gorlin, 1968; Gianelly et al., 1967; Ginn & Orgain, 1966). It has been suggested that propranolol works in angina pectoris by a means other than beta-receptor blockade (Grandjean, 1967; British Medical Journal, 1969). Propranolol possesses the additional pharmacological property of membrane-stabilization for local anaesthesia (Vaughan Williams, 1966; Fitzgerald, 1960). Hence, it has been suggested that this property, by causing non-specific myocardial depression, will improve the anginal state. Clinical evidence that this is not so has been provided by the study of Robinson’s group in St George’s Hospital (Wilson et al., 1969). They have compared the dextro-isomer of propranolol which has equivalent membrane-stabilizing properties, but no beta-blocking properties, with racemic propranolol, which possesses both properties. They showed that dextrapropranolol, like saline, did not increase exercise tolerance in eight anginal patients, whilst racemic propranolol did do so. It is now very probable that the possession of membrane-stabilizing properties is irrelevant to the mode of action of racemic propranolol in the clinical dose range in which it is used (Barrett, 1969; Fitzgerald, unpublished observations).

Once the clinical effectiveness of propranolol had been established, the problem then arose as to what further discovery would be clinically desirable. Propranolol blocks other clinically important beta-receptors, particularly those in the lung and those subserving certain metabolic responses. A legitimate therapeutic aim might be an increase in selectivity of activity, so that only beta receptors in the heart were blocked. Furthermore, a compound might be discovered which possessed no membrane stabilizing or local anaesthetic properties. In order to discover such a compound, the following screening procedure was employed. Initially all candidate drugs were studied for their effects in antagonizing isoprenaline on the heart, peripheral blood vessels and lungs. The aim of these primary tests was to separate an effect on the heart from the effects on the lung and blood vessels. If a likely compound was found, it was then studied for its effects on the spike potential in the isolated frog nerve and also for its effects on intra-cardiac conduction. If the candidate compound still seemed interesting, its effects on lipolysis of the epididymal fat pad and on blood lactate levels were also studied. In addition, its possible effects on digitalis and halothane-catecholamine arrhythmias would be studied at this time. Such procedures are clearly far removed from the clinical situation, and their justification is based on a series of assumptions, some of which may be regarded as very tenuous.

**Practolol**

By such procedures, a new beta blocking compound, practolol (‘Eraldin’, ICI), was discovered. This compound is cardio-selective and has no membrane-stabilizing properties. It also differs from propranolol in that it has a degree of intrinsic stimulant activity. Practolol has a half-life in man of 10 hr compared with 2 hr for propranolol, and is minimally metabolized in man, whereas propranolol is extensively metabolized (Barrett et al., 1968; Dunlop & Shanks, 1968; Fitzgerald & Scales, 1968). Practolol has been undergoing clinical evaluation for some 2 years, and has already been shown to increase exercise tolerance in angina pectoris as well as to reduce the consumpnt of glycerin trinitrate and the incidence of anginal pain. These observations are of considerable academic interest, since they confirm Black’s original view that sympathetic activity may be deleterious to the heart of patients with angina pectoris. They also show that the membrane-stabilizing properties of propranolol are not necessary for its action in angina pectoris, since practolol is as effective as propranolol in increasing exercise tolerance, but does not have any local anaesthetic action (Wilson et al., 1969).

**Other clinical indications**

Clinical situations in which propranolol has been shown to be of value are given in the accompanying
table. Black has indicated that none of these was predicted from the initial experimental studies in animals (Black, 1967). This indicates yet again the tremendous value of accurate clinical observation and feed-back of these observations to the laboratory as a guide for further research. However, Black did predict that beta-blockade might be of value in the cardiac arrhythmias associated with myocardial infarction, since it was already known that this was a situation associated with high concentration of catecholamines in an area of local tissue necrosis or anoxia. Such a situation is known to be potentially arrhythmogenic, and studies over the last 10 years have indicated the high incidence of cardiac arrhythmias following myocardial infarction. Initially, studies with propranolol indicated that it would control or reverse cardiac arrhythmias in myocardial infarction, but subsequent extensive study both with propranolol and oxprenolol show that if the drugs are given routinely to patients with myocardial infarction they have no effect on the morbidity or mortality associated with this condition (Sowton, 1968). These observations proved to be very disappointing, since there had been considerable hopes that the prophylactic administration of a beta-blocking drug might diminish the initial high mortality due to cardiac arrhythmias in myocardial infarction, because animal investigations had shown that beta-blockade would protect against ventricular arrhythmias in experimental myocardial infarction (Fearon, 1967; Pentecost & Austen, 1966).

Myocardial infarction is associated with a state of high autonomic tone, and beta-blockade in this situation may affect the patient adversely for two reasons. Firstly, removal of sympathetic activity will permit vagal dominance of the heart, resulting in inappropriate cardiac slowing, a fall in cardiac output, hypotension and shock. The important role of the vagus in this situation has been demonstrated by Stannard, Sloman & Sangster (1968), who gave atropine at the same time as propranolol, and prevented the unwanted haemodynamic sequelae. In addition, severely affected hearts may require increased sympathetic activity to maintain a satisfactory degree of myocardial contractility. Blockade of enhanced sympathetic activity in this situation may reduce myocardial contractility below a critical level. It seems, therefore, that the original laboratory predictions for beta-blockade in cardiac arrhythmias associated with myocardial infarction have not been fulfilled. However, cardiac arrhythmias associated with anaesthesia, thyrotoxicosis, phaeochromocytoma, exercise and several other states, have been shown to respond favourably to beta-blockade (Gibson & Sowton, 1969). Recent studies with the cardio-selective beta-blocker, practolol, in post-infarction arrhythmias suggest that this compound may be of more value and safer to use in this particular situation, but much clinical study is yet to be done before such a conclusion can be proven (Jewitt, Mercer & Shillingford, 1969).

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