Dietary fish oil preserves cardiac function in the hypertrophied rat heart

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Abstract
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Keywords
rat, hypertrophied, function, heart, cardiac, dietary, preserves, oil, fish

Disciplines
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Abstract
Regular fish or fish oil intake is associated with a low incidence of heart failure clinically, and fish oil-induced reduction in cardiac remodelling seen in hypertrophy models may contribute. We investigated whether improved cardiac energy efficiency in non-hypertrophied hearts translates into attenuation of cardiac dysfunction in hypertrophied hearts. Male Wistar rats (n 33) at 8 weeks of age were sham-operated or subjected to abdominal aortic stenosis to produce pressure-overload cardiac hypertrophy. Starting 3 weeks post-operatively to follow initiation of hypertrophy, rats were fed a diet containing 10 % olive oil (control) or 5 % fish oil (ROPUFA w30 (17 % EPA, 10 % DHA)) + 5 % olive oil (FO diet). At 15 weeks post-operatively, ventricular haemodynamics and oxygen consumption were evaluated in the blood-perfused, isolated working heart. Resting and maximally stimulated cardiac output and external work were 60 % depressed in hypertrophied control hearts but this was prevented by FO feeding, without attenuating hypertrophy. Cardiac energy efficiency was lower in hypertrophy, but greater in FO hearts for any given cardiac mass. Coronary blood flow, restricted in hypertrophied control hearts, increased with increasing work in hypertrophied FO hearts, revealing a significant coronary vasodilator reserve. Pronounced cardiac dysfunction in hypertrophied hearts across low and high workloads, indicative of heart failure, was attenuated by FO feeding in association with membrane incorporation of n-3 PUFA, principally DHA. Dietary fish oil may offer a new approach to balancing the high oxygen demand and haemodynamic requirements of the failing hypertrophied heart independently of attenuating hypertrophy.

Key words: Fish oil; n-3 Fatty acids; Heart failure; Hypertrophy; Diet

Despite declining heart disease mortality rates, a continuing rise in heart failure incidence is of major clinical concern in many nations1. Regular fish (and hence n-3 PUFA) consumption is associated with low CVD mortality2,3. Antiarrhythmic effects first identified in animal models4 are evident in clinical studies showing reduced incidence of primary cardiac arrest5 and post-infarction arrhythmic deaths6, and represent the best-supported mechanism of action. However, in addition to arrhythmia prevention, alternative mechanisms of fish oil action, such as prevention of heart failure, are implicated directly from epidemiology7, and indirectly as reduced non-arrhythmic but otherwise unspecified cardiac deaths and post-infarction deaths8. These clinical outcomes are independent of dietary n-3 PUFA effects on classic risk factors such as plasma TAG and thrombogenic factors9 or blood pressure10. Animal studies have described fish oil-induced improvements in heart function and the prevention of fatal arrhythmia in non-disease states that are associated with the incorporation of the long-chain n-3 PUFA DHA into myocardial membranes4–10 and occur at lower intakes than for the vascular effects4.

Hearts isolated from rats fed fish oil exhibit significantly reduced oxygen consumption without compromise to cardiac output (CO) or external work13. This represents increased energy efficiency of oxygen use, increased coronary perfusion reserve, less damage in acute ischaemia and reduced myocardial stunning13,14,15. Cardiac hypertrophy is generally characterised by morphological and biochemical changes that predispose to myocardial ischaemia through disturbances of both oxygen supply and demand at the cellular level, which promote energetic failure and heart failure16,17. For example, high ventricular wall tension increases oxygen demand and restricts coronary blood flow, compromising oxygen delivery to myocardial mitochondria already jeopardised by high inter-capillary distances. The capacity of the hypertrophied cell to use oxygen has reportedly been further reduced by

Abbreviations: CO, cardiac output; FO, fish oil; FOH, rats with cardiac hypertrophy fed with a fish oil diet; MVO2, myocardial oxygen consumption; OO, olive oil; OOH, rats with cardiac hypertrophy fed with an olive oil diet; SV, stroke volume.

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a high ratio of contractile protein:mitochondria, and mitochondrial function itself may be compromised[18]. Consequently, an intervention that reduces myocardial oxygen consumption (MVO₂) without detriment to myocardial work output could improve cardiac function in hypertrophy and retard the progression to failure[19].

The present study tests the hypothesis that incorporation of n-3 PUFA into myocardial membranes through fish oil feeding will improve cardiac energy efficiency and attenuate cardiac dysfunction and heart failure in the hypertrophied heart.

Methods
A total of thirty-three male Wistar rats (CSIRO Human Nutrition, Adelaide, Australia) were used in the present study. The experiments were conducted according to the National Health and Medical Research Council of Australia, Guidelines for the Use of Experimental Animals, and were approved by the Animal Care and Ethics Committees of the University of Wollongong and CSIRO Human Nutrition.

Animal model of pressure-overload hypertrophy
Under ketamine (70 mg/kg) + xylazine (5 mg/kg) anaesthesia, twenty-eight rats were subjected to restrictive banding of the suprarenal abdominal aorta at 8 weeks of age using a blunt 23G needle as a template. The needle was tied tightly against the aorta above the renal arteries (causing visible kidney blanching), and then removed, leaving the suture in place to partially restore blood flow (visually confirmed). Another five rats were subjected to sham operation without aortic restriction. Heart function was assessed 15 weeks post-operatively.

Cardiac hypertrophy was determined from the heart weight:body-weight ratio and the heart weight:tibia length ratio. The latter index is not distorted by acute fluctuations in body weight or body-weight gain from fluid retention. Cardiac hypertrophy was defined as a ratio > 2 SD above the mean for rats of the same strain and age as those in the present study. The pooled mean of ventricle dry weight:tibia length from fifty-three rats previously recorded in this laboratory, not subjected to surgical intervention, was 0·376 (SD 0·045) g/cm. Cardiac hypertrophy was therefore defined for this strain and age of rat as a ventricle weight:tibia length ratio > 0·466 g/cm.

The pressure overload induced by aortic stenosis provokes a hypertrophic response within 4 d, and it has been well established by 3 weeks[20]. Because fish oil inhibits cardiac hypertrophy when presented before and during a hypertrophic stimulus[21,22], in the present study, the initiation of fish oil feeding was delayed until 3 weeks after surgery in order to focus interpretation on pathophysiology of the hypertrophied heart. The aortic stenosis model was preferred to alternative models of the following: aortic valve disruption (volume overload), which is unsuitable for evaluation in the isolated working heart preparation; and coronary artery occlusion (chronic myocardial infarction), which is prone to a high incidence of acute fatal arrhythmia and potentially confounding effects of tissue repair processes that may be influenced by dietary intervention independently of the effects on hypertrophic function.

Diets
Rats were fed an olive oil (OO) control diet for 4 weeks before surgery and 3 weeks post-operatively. They were then randomly allocated to either the fish oil (FO) diet or the OO control diet for 12 weeks. Sham-operated rats were fed the OO control diet throughout the experiment. Isoenergetic fully fabricated diets were prepared[16,23] containing 100 g/kg diet as fat. The OO (control) diet contained 10% (by weight) olive oil, and the FO diet contained 5% olive oil plus 5% fish oil. The olive oil was commercially available food-grade olive oil, refined to exclude antioxidant phytochemicals. The fish oil (ROPUFA® 30 n-3 EPA oil; DSM Nutritional Products, Basel, Switzerland) contained 16·8% EPA (20:5n-3) and 9·7% DHA (22:6n-3); 32·5% total n-3 fatty acids). Olive oil was selected for the isoenergetic control diet on the basis of the following: its main component (monounsaturated oleic acid (18:1)) being a physiologically neutral fat, and the oil providing sufficient PUFA to avoid essential fatty acid deficiency[10]. Although n-3 PUFA replace n-6 PUFA in myocardial membranes, the long-chain n-3 PUFA are incorporated in direct relationship to their absolute quantities in the diet and independently of dietary MUFA or n-6 PUFA dietary concentrations[23]. The main fatty acid components of the two diets were as follows: total SFA (OO 14%; FO 22%); total PUFA (OO 10%; FO 22%); oleic acid (18:1; OO 76%; FO 45%); linoleic acid (18:2n-6; OO 83%; FO 52%); EPA (FO 8·4%); DHA (FO 4·9%).

Isolated working heart
Rats were anaesthetised (pentobarbitone sodium 60 mg/kg intraperitoneally), and the right common carotid artery and the left femoral artery were cannulated to record arterial blood pressures proximal and distal to the aortic stenosis. Rats were then killed by rapid exsanguination and the hearts removed into ice-cold saline and prepared for working heart perfusion with porcine erythrocyte buffer (40% haematocrit in Krebs Henseleit solution) as described previously[13]. Briefly, the aorta, left atrium and pulmonary artery were cannulated and the heart perfused with Krebs Henseleit solution in Langendorff mode for 10 min for rat blood washout. Hearts were switched to working heart mode and perfused with erythrocyte buffer via the left atrium with 10 mmHg filling pressure (preload) and 75 mmHg diastolic aortic pressure (afterload, coronary perfusion pressure). A catheter introduced through silicone self-sealing tubing supplying the left atrium and fed into the ventricle via the mitral valve recorded left ventricular pressure. Working hearts were allowed to stabilise for 15 min.

Cardiac function
CO and coronary flow were measured at 5 min intervals by timed overflow, with coronary and aortic samples collected...
for the measurement of arterial and venous blood gases and pH. Stroke volume (SV) was calculated from the CO and heart rate. MVO$_2$ was derived from arterio-venous oxygen differences per unit coronary flow, and external work was derived from the CO and developed pressure (CO × mean aortic pressure) as described previously.$^{13,23}$ Cardiac energy efficiency (%) was calculated as the ratio of energy derived from MVO$_2$ (in kJ) to energy delivered as external work (in kJ) × 100. After equilibration, preload was changed over the range 5–20 mmHg by altering ventricular filling pressure, before returning pressure to 10 mmHg for recovery. Intraventricular pressure, aortic pressure and electrocardiogram were recorded continuously, while coronary and aortic flows were measured and blood samples were collected at 5 min intervals.

**Membrane fatty acids**

Hearts were rinsed in ice-cold saline, blotted dry and weighed. A small section of the ventricle free wall was weighed and retained for dry-weight estimation. The remaining ventricle was rapidly frozen using liquid N$_2$-cooled clamps and then frozen at −60°C until required for fatty acid measurement according to established methods.$^{23}$ Briefly, samples of the frozen tissue were pulverised over liquid N$_2$ and washed according to established methods.$^{23}$ In brief, samples of the frozen tissue were pulsed over liquid N$_2$ and washed according to established methods.$^{23}$

**Statistical analysis**

The CO, external work, coronary flow, oxygen extraction and MVO$_2$ were expressed 1/g ventricle dry weight. Results are expressed as means with their standard errors. For hemodynamic measures, two-way ANOVA was conducted with diet and hypertrophy main effects and diet × hypertrophy interaction. Individual comparisons between diet × hypertrophy were conducted using Scheffe’s post hoc $F$ test for multiple comparisons of individual means. Values in tables sharing a common superscript letter were not significantly different. The level of significance was considered at $P<0.05$.

**Results**

**Heart and body weights, blood pressure and cardiac hypertrophy**

Cardiac enlargement sufficient to classify as hypertrophy occurred in six of the fourteen rats in each of the dietary groups subjected to abdominal aortic stenosis. Rats with cardiac hypertrophy (designated OOH if fed the olive oil diet and FOH if fed the FO diet) had significantly greater ventricle weights and body weights than non-hypertrophied (designated OO or FO) or sham-operated rats (Table 1). Tibia length, body weight, body size or heart weight were not different between the dietary groups within either the hypertrophy or non-hypertrophy animals (Table 1). Less variation in tibia length than in body weight resulted in the ventricle length-tibia length being the clearer discriminator of hypertrophy.

### Table 1. Effect of diet and cardiac hypertrophy on animal characteristics and isolated working heart function at equilibrium

(Mean values with their standard errors)

<table>
<thead>
<tr>
<th>Diet group…</th>
<th>Sham control (n 5)</th>
<th>No hypertrophy (n 8)</th>
<th>Hypertrophy (n 6)</th>
<th>No hypertrophy (n 8)</th>
<th>Hypertrophy (n 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean (SEM)</td>
<td>Mean (SEM)</td>
<td>Mean (SEM)</td>
<td>Mean (SEM)</td>
<td>Mean (SEM)</td>
</tr>
<tr>
<td>Body weight (g)</td>
<td>477 (10)</td>
<td>442 (13)</td>
<td>504 (28)</td>
<td>468 (15)</td>
<td>486 (16)</td>
</tr>
<tr>
<td>Atria weight (g)</td>
<td>0.12 (0.02)</td>
<td>0.12 (0.01)</td>
<td>0.18 (0.03)</td>
<td>0.12 (0.01)</td>
<td>0.15 (0.02)</td>
</tr>
<tr>
<td>Ventricle weight (g)</td>
<td>1.66 (0.06)</td>
<td>1.49 (0.05)</td>
<td>2.23 (0.10)</td>
<td>1.69 (0.06)</td>
<td>2.10 (0.07)</td>
</tr>
<tr>
<td>Ventricle dry weight (g)</td>
<td>0.27 (0.01)</td>
<td>0.27 (0.01)</td>
<td>0.39 (0.01)</td>
<td>0.31 (0.01)</td>
<td>0.38 (0.02)</td>
</tr>
<tr>
<td>Dry:wet left ventricle weight</td>
<td>0.163 (0.005)</td>
<td>0.185 (0.004)</td>
<td>0.175 (0.007)</td>
<td>0.188 (0.002)</td>
<td>0.184 (0.002)</td>
</tr>
<tr>
<td>Tibia length (cm)</td>
<td>4.16 (0.01)</td>
<td>4.15 (0.04)</td>
<td>4.13 (0.07)</td>
<td>4.26 (0.07)</td>
<td>4.16 (0.06)</td>
</tr>
<tr>
<td>Ventricle weight:tibia length (g/cm)</td>
<td>0.40 (0.02)</td>
<td>0.359 (0.012)</td>
<td>0.545 (0.030)</td>
<td>0.397 (0.015)</td>
<td>0.503 (0.014)</td>
</tr>
<tr>
<td>Ventricle weight:body weight (g/100 g)</td>
<td>0.348 (0.010)</td>
<td>0.338 (0.011)</td>
<td>0.459 (0.031)</td>
<td>0.383 (0.017)</td>
<td>0.438 (0.032)</td>
</tr>
<tr>
<td>Heart weight:body weight (g/100 g)</td>
<td>0.373 (0.014)</td>
<td>0.356 (0.014)</td>
<td>0.488 (0.039)</td>
<td>0.386 (0.022)</td>
<td>0.443 (0.039)</td>
</tr>
<tr>
<td>Carotid systolic BP (mmHg)</td>
<td>146 (14)</td>
<td>161 (19)</td>
<td>220 (20)</td>
<td>166 (17)</td>
<td>218 (19)</td>
</tr>
<tr>
<td>Carotid diastolic BP (mmHg)</td>
<td>118 (13)</td>
<td>131 (17)</td>
<td>165 (13)</td>
<td>131 (12)</td>
<td>164 (15)</td>
</tr>
<tr>
<td>Femoral systolic BP (mmHg)</td>
<td>166 (13)</td>
<td>134 (16)</td>
<td>136 (13)</td>
<td>135 (19)</td>
<td>148 (15)</td>
</tr>
<tr>
<td>Femoral diastolic BP (mmHg)</td>
<td>125 (16)</td>
<td>100 (14)</td>
<td>111 (12)</td>
<td>114 (16)</td>
<td>118 (14)</td>
</tr>
<tr>
<td>Isolated working heart at equilibrium</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart rate (beats per min)</td>
<td>207 (26)</td>
<td>203 (23)</td>
<td>185 (24)</td>
<td>211 (14)</td>
<td>210 (19)</td>
</tr>
<tr>
<td>Cardiac output (ml/min per g dry weight)</td>
<td>82 (19)</td>
<td>108 (22)</td>
<td>56 (20)</td>
<td>128 (33)</td>
<td>127 (20)</td>
</tr>
<tr>
<td>Coronary flow (ml/min per g dry weight)</td>
<td>33 (4.7)</td>
<td>34.3 (6.2)</td>
<td>17.3 (3.9)</td>
<td>30.2 (6.1)</td>
<td>40.6 (4.8)</td>
</tr>
<tr>
<td>Ventricle dP/dt$_{max}$ (mmHg/s)</td>
<td>3070 (194)</td>
<td>3634 (503)</td>
<td>2389 (452)</td>
<td>3233 (388)</td>
<td>3236 (317)</td>
</tr>
<tr>
<td>Ventricle – dP/dt$_{max}$ (mmHg/s)</td>
<td>2460 (269)</td>
<td>2608 (355)</td>
<td>1632 (278)</td>
<td>2351 (292)</td>
<td>2560 (320)</td>
</tr>
</tbody>
</table>

OO, olive oil diet; OOH, rats with cardiac hypertrophy fed with an olive oil diet; FO, fish oil diet; FOH, rats with cardiac hypertrophy fed with a fish oil; BP, blood pressure.

$^{a,b}$Mean values within a row with unlike superscript letters were significantly different ($P<0.05$).
Femoral artery pressure was significantly lower (P<0.05) in rats subjected to aortic stenosis than in sham-operated controls and did not differ between the rats with or without cardiac hypertrophy or between the dietary groups (Table 1). Carotid artery pressure was significantly higher in animals with cardiac hypertrophy compared with those that failed to develop hypertrophy or sham-operated controls (P<0.05). There were no significant differences in carotid blood pressures between the dietary groups.

Equilibrium heart function

Under initial conditions of the 10 mmHg preload and the 75 mmHg afterload, there were no differences in spontaneous resting heart rate (Table 1). Uncorrected CO (ml/min) was significantly higher in hearts from fish oil-fed rats (FO 41.3 (SEM 10.9); FOH 48.1 (SEM 10.5)) than in hearts from rats fed olive oil (sham 20.1 (SEM 4.7); OO 29.0 (SEM 4.9); OOH 21.9 (SEM 7.6)) (P<0.01), with no significant interaction (P>0.05).

When corrected for ventricular mass, the CO of the hypertrophied OOH hearts was significantly lower per gram of tissue (P<0.05) than the OO, FO and hypertrophied FOH hearts (Table 1). The CO of the FOH (hypertrophied) hearts was not significantly different from that of the FO (non-hypertrophied) hearts. There was a significant (diet × hypertrophy) interaction for the maximum rate of ventricular relaxation (P=0.02) and the OOH hearts showed a significantly slower relaxation rate than all the other hearts (P<0.05; Table 1).

Coronary flow did not differ among the non-hypertrophied hearts but was significantly lower in the OOH hearts (P<0.01) than all the other hearts (Table 1). The MVO₂ at equilibrium in the FOH hearts was significantly lower per gram of tissue (P<0.05) and the OOH hearts showed a significantly slower relaxation rate than all the other hearts (P<0.05; Table 1).

The hypertrophied hearts had significantly lower energy efficiency than the non-hypertrophied hearts (two-way ANOVA (diet, hypertrophy), P<0.05) and the FO hearts had higher cardiac efficiency than the OO hearts (P<0.05). Cardiac energy efficiency was significantly lower in the OOH hearts than in the OO, FO and FOH hearts (Fig. 1).

Cardiac function under load

The CO, SV (Fig. 2) and external work (Fig. 3) increased with increasing filling pressure (preload) (P<0.0001). The hypertrophied hearts had a lower SV, CO or external work over the range of filling pressures (P<0.0001). A significant (diet × hypertrophy) interaction was evident for SV (P=0.01), CO (P=0.023) and external work (P=0.044). The OOH hearts had a significantly lower SV (P<0.0001), CO (P<0.0001) and external work (P<0.0001) than the OO hearts, while there were no significant differences between the FOH and FO hearts (SV P=0.10; CO P=0.082; external work P=0.51). Among the hypertrophied hearts, the FOH group had a significantly higher SV (P=0.008), CO (P=0.003) and external work (P=0.003) than the OOH hearts across the range of filling pressures.

Coronary flow and MVO₂ increased with increasing filling pressure (P<0.04; Fig. 2). A significant (diet × hypertrophy) interaction was evident for coronary flow (P=0.01) and MVO₂ (P=0.002). Coronary flow over the range of filling pressures was significantly lower in the OOH hearts than in the FOH (P=0.015) and OO hearts (P=0.005). The MVO₂ was not significantly different in the FOH hearts compared with the FO hearts (P=0.19), and there were no significant differences in coronary flow (P=0.12) or MVO₂ (P=0.059) between the OO and FO hearts. The cardiac energy efficiency of oxygen conversion into external work increased (P=0.0003) as external work was increased by increasing filling pressure (Fig. 3), and it was significantly lower in hypertrophy (P<0.0001) with both diets. No significant (diet × hypertrophy) interaction was evident for cardiac energy efficiency (P=0.87). Cardiac efficiency was higher in the FO hearts than in the OO and sham hearts (P=0.004) and higher in the FOH hearts than in the OOH hearts (P=0.004).

Membrane composition

The phospholipid fatty acid composition of the rat hearts was significantly influenced by the fat source in the diet. The FO
diet significantly increased the concentration of n-3 PUFA EPA and DHA and reduced the concentration of monounsaturated oleic acid and the n-6 PUFA arachidonic acid compared with the OO diet (Table 2). The total saturated fat concentration was unaltered but the total PUFA concentration and the unsaturation index were increased with FO feeding.

The hypertrophied OOH hearts incorporated significantly less linoleic acid ($P < 0.05$) with non-significant trends towards increased arachidonic acid ($P = 0.1$) and DHA ($P < 0.1$), producing, overall, a non-significant trend towards reduced n-6 PUFA ($P < 0.1$) and a small but significant increase in unsaturation index ($P < 0.05$) compared with the non-hypertrophied OO hearts (Table 2). There were no significant differences in phospholipid fatty acid composition between the hypertrophied (FOH) and non-hypertrophied (FO) fish oil hearts.

**Discussion**

The results of the present study demonstrate that incorporation of n-3 PUFA into myocardial cellular membranes to reduce MVO$_2$ in relation to the cardiac work output seen in normal hearts$^{(15)}$ translates into attenuation of cardiac dysfunction in the hypertrophied hearts. Fish oil feeding is known to limit the development of cardiac hypertrophy induced by a variety of stimuli$^{(21,22,24,25)}$, and until now, any attenuation of hypertrophied heart dysfunction by n-3 PUFA was always accompanied by, and could not be differentiated from, the prevention of ventricular remodelling$^{(21,24)}$. Our demonstration that hypertrophied heart function was improved independently of attenuating cardiac hypertrophy suggests that dietary fish oil may prevent cardiac dysfunction progressing to heart failure. The improved cardiac function, demonstrated not only at rest (as previously shown in association with hypertrophy attenuation$^{(21,24)}$), but over a wide range of loading conditions and functional measures, provides further support.

Fish oil improvement of cardiac function was largely limited to the hypertrophied hearts, even though myocardial energy efficiency was significantly greater with fish oil feeding in the non-hypertrophied hearts. Improved oxygen efficiency often does not translate into cardiac function improvement in healthy, well-oxygenated hearts$^{(15,24)}$. However,
energy-sparing effects consistently translate into improved heart function under cardio-depressant stress conditions such as myocardial ischaemia or cardiac hypertrophy. Attenuation of the decline in CO, the decline in cardiac relaxation and filling, the depressed external work and energy efficiency, over the entire Frank–Starling relationship, especially at high workload, illustrates the potential for translation across all stages of heart failure.

Increased ventricular stiffness impairs relaxation and filling, increases the work of contraction, impairs cardiac energetics, prolongs coronary artery compression and impairs coronary perfusion. It is therefore a major contributor to cardiac dysfunction. Retention of residual wall tension was less evident in the hypertrophied fish oil hearts by virtue of better cardiac relaxation. The hypertrophy-induced reduction in cardiac energy efficiency was attenuated by fish oil feeding, and the hypertrophied fish oil hearts did not display the depressed rate of relaxation, ventricle filling and CO usually associated with post-adaptive cardiac hypertrophy. Furthermore, the increase in coronary vasodilator reserve associated with fish oil in healthy hearts translated into an adaptive increase in oxygen delivery in the hypertrophied hearts during the demand of raised filling pressure, overcoming the normally depressed coronary flow and providing further advantage in pathophysiological conditions.

**Potential mechanisms of n-3 PUFA actions**

Impaired relaxation, contraction and energetic failure underpinning the pathophysiology of cardiac dysfunction and heart failure are all associated with altered myocardial Ca²⁺ homeostasis. In turn, energetic failure leads to altered Ca²⁺ handling, creating a vicious cycle. Conversely, numerous effects of the long-chain n-3 PUFA on myocardial Ca²⁺ regulation have biologically plausible correlates in the effects of dietary fish oil on hypertrophied heart function that might play a role in breaking that cycle. For example, pathological depolarisation and cardiac arrhythmias may be attenuated by the following: inhibition of phospholipase-C and to thereby attenuate inositol trisphosphate and diacylglycerol mobilisation from membrane phospholipids, which in turn attenuates Ca²⁺ sparks. Hypertrophied hearts could also benefit from increased coronary vasodilator reserve and reduced coronary artery compression, allowing for increased oxygen delivery.
heart dysfunction may be attenuated by the following: inhibiting mitochondrial Ca\(^{2+}\) overload to attenuate excessive oxygen consumption\(^1\),\(^15\),\(^20\); and limiting pathological increases in cytosolic free Ca\(^{2+}\) and cellular Ca\(^{2+}\) transients to reduce residual wall stiffness and improve relaxation and filling\(^29\),\(^30\).

In the present study, use of the isolated, blood-perfused, working heart model allows us to draw conclusions about the intrinsic role of membrane incorporation of the n-3 PUFA\(^1\),\(^4\),\(^14\),\(^15\),\(^27\),\(^32\),\(^33\) by excluding nervous or humoral influences or variable relationships between the availability of arachidonic acid with fish oil feeding implies a role for reduced eicosanoid production. However, earlier studies have shown that cyclo-oxygenase and lipoxygenase production of potent eicosanoids derived from arachidonic acid is not prevented by increases in membrane EPA when the DHA increase is inadequate\(^15\). While not denying a role for EPA in inflammatory and other signalling processes in blood, vasculature and other tissues\(^27\),\(^35\), membrane DHA status is consistent with clinical evidence of direct myocardial effects of dietary fish and fish oil. For whereas fish and fish oil reduce heart rate, heart failure and sudden cardiac death\(^5\),\(^7\),\(^8\),\(^10\),\(^34\) without preventing new (vascular) ischaemic events\(^8\),\(^10\), purified EPA used in a major clinical trial prevented new vascular ischaemic events without reducing sudden cardiac death\(^35\). The reduction in membrane arachidonic acid with fish oil feeding implies a role for reduced eicosanoid production. However, earlier studies have shown variable relationships between the availability of arachidonic acid in myocardial membranes and the ability of fish oils and other dietary fats to modulate myocardial eicosanoid production and pathophysiological responses such as cardiac arrhythmia\(^36\),\(^37\). Alternatively, DHA can inhibit the cyclo-oxygenase and lipoxygenase production of potent eicosanoids derived from arachidonic acid, and can also inhibit MAP kinase-stimulated Ca\(^{2+}\)-activated phospholipase-D to restrict arachidonic acid mobilisation from membrane phospholipid\(^27\). In addition to influencing eicosanoid mechanisms, n-3 PUFA can directly modulate cell membrane lipid and protein composition\(^14\),\(^15\). For example, ischaemic arrhythmias are not prevented by increases in membrane EPA when the DHA increase is inadequate\(^15\). While not denying a role for EPA in inflammatory and other signalling processes in blood, vasculature and other tissues\(^27\),\(^35\), membrane DHA status is consistent with clinical evidence of direct myocardial effects of dietary fish and fish oil.

<table>
<thead>
<tr>
<th>Table 2. Effect of cardiac hypertrophy and ROPUFA(^6) 30 dietary fish oil supplementation on myocardial total phospholipid fatty acid composition§</th>
<th>Percentage of occurrence¶</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diet</td>
<td>OO</td>
</tr>
<tr>
<td>-----------------------------------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Fatty acid</td>
<td>Mean</td>
</tr>
<tr>
<td>16:0</td>
<td>9.98</td>
</tr>
<tr>
<td>18:0</td>
<td>22.76</td>
</tr>
<tr>
<td>18:1:n-9 (OA)</td>
<td>4.71</td>
</tr>
<tr>
<td>18:2:n-6 (LA)</td>
<td>10.83</td>
</tr>
<tr>
<td>20:4:n-6 (AA)</td>
<td>12.72</td>
</tr>
<tr>
<td>20:5:n-3 (EPA)</td>
<td>20.56</td>
</tr>
<tr>
<td>22:4:n-6</td>
<td>0.05</td>
</tr>
<tr>
<td>22:5:n-3 or 24:0</td>
<td>0.41</td>
</tr>
<tr>
<td>22:5:n-6</td>
<td>1.72</td>
</tr>
<tr>
<td>22:6:n-3 (DHA)</td>
<td>9.32</td>
</tr>
<tr>
<td>Total fatty acids</td>
<td>93.88</td>
</tr>
<tr>
<td>Sum SFA</td>
<td>32.74</td>
</tr>
<tr>
<td>Sum MUFA</td>
<td>15.54</td>
</tr>
<tr>
<td>Sum PUFA</td>
<td>45.60</td>
</tr>
<tr>
<td>Total n-3 PUFA</td>
<td>19.10</td>
</tr>
<tr>
<td>Total n-6 PUFA</td>
<td>35.41</td>
</tr>
<tr>
<td>n-6:n-3</td>
<td>3.47</td>
</tr>
<tr>
<td>UI</td>
<td>193.7</td>
</tr>
</tbody>
</table>

OO, olive oil diet; OOH, rats with cardiac hypertrophy fed with an olive oil diet; FO, fish oil diet; FOH, rats with cardiac hypertrophy fed with a fish oil; OA, oleic acid; LA, linoleic acid; AA, arachidonic acid; ND, not detected; UI, unsaturation index.

* Mean values were significantly different for FO from those of the OO group \(P<0.05\).
† Mean values were significantly different for FOH from those of the OOH group \(P<0.05\).
‡ Mean values were significantly different for FOH from those of the OO group \(P<0.05\).
With more membrane manufactured as cells enlarge, hypertrophy-driven changes have the potential to alter myocardial phospholipid composition and modify responses to dietary intervention. Indeed, adaptive decreases in linoleic acid and increases in arachidonic acid, DHA and membrane total unsaturation that occur in response to a variety of stressors\(^{28}\), including pressure overload\(^{20,24}\) and catecholamine stress\(^{59}\), were observed in the present study. However, none of the adaptive changes were as great as changes induced by the FO diet. Any hypertrophy-associated increases in DHA and total unsaturation were overwhelmed by the dietary intervention and the arachidonic acid increase was reversed, such that no differences were evident between the hypertrophied and non-hypertrophied fish oil hearts.

**Comparison with therapeutic interventions to enhance cardiac function**

Dietary fish oil-induced improvement of hypertrophied heart contractile function has some similarities but many differences to therapeutic interventions used in the treatment of heart failure. For example, positive inotropic agents typically improve systemic haemodynamics and provide symptomatic relief in heart failure by increasing contractility at the expense of reduced energy efficiency\(^{40}\), increased risk of ischaemia\(^{41}\), cardiac arrhythmias and increased early mortality. Vasodilators relieve heart failure by reducing systemic blood pressure and unloading the heart, which the fish oil did not do in the present study. Moreover, the negative inotropic actions of the Ca\(^{2+}\) channel blocking class of vasodilators\(^{42}\) are inconsistent with the effects of fish oil. The angiotensin-converting enzyme (ACE) inhibitors and angiotensin receptor antagonists have no direct effect on myocardial contractile performance, acting principally by inhibiting hypertrophy and myocardial fibrosis\(^{43,44}\), whereas fish oil directly improved cardiac function without the effect on ventricular remodelling.

The greatest similarity of physiological mechanisms lies with adrenergic \(\beta\)-blockers, which reduce the heart rate and reduce the chronic hyper-adrenergic state that is typical of heart failure\(^{45}\), thereby reducing energy requirements. However, although fish oils can reduce intrinsic heart rate and prevent sudden death in rats associated with chronic isoprenaline treatment\(^{59}\), they have no \(\beta\)-adrenoceptor antagonist action\(^{46}\), and unlike \(\beta\)-blockers, heart rate reduction is achieved independently of sympathetic nervous activity\(^{12,16}\). In common with \(n-3\) PUFA, \(\beta\)-blockers facilitate chronic intrinsic improvements in cellular function in hypertrophied cardiomyocytes by improvements in cellular energetics\(^{49}\) or intracellular Ca\(^{2+}\) handling\(^{43}\).

Thus, dietary fish oil improved cardiac function in hypertrophied hearts in a manner that contrasts to the currently available suite of treatments used to stimulate the heart or counteract dysfunction. It appears to have an intrinsic effect on myocardial cellular function independent of cardiac unloading or slowed heart rate, to improve contractility without adverse energetic consequences and independently of modulating the hypertrophy.

**Therapeutic implications**

Epidemiological evidence showing a reduced risk of incident heart failure in people who regularly eat fish\(^{17,47}\) is now supported by a large secondary prevention clinical trial in which \(n-3\) PUFA supplementation reduced mortality and hospitalisation in heart failure in the context of usual care\(^{48}\), which included simultaneous treatment with many of the therapeutic agents discussed above. The findings of the present study begin to explain not only the original GISSI-HF outcomes but also a recent subset evaluation of the GISSI-HF trial\(^{48}\) which noted that the reduced number of hospital admissions due to cardiovascular reasons in heart failure was paralleled by improved cardiac function without the evidence of reverse remodelling\(^{49}\).

Animal and human studies have suggested that a ceiling exists for the cardiac effects of \(n-3\) PUFA\(^{50,51}\), which in animals reflects maximum incorporation of DHA into myocardial membranes\(^{23,52}\). Thus, the effects reported here for 5% dietary fish oil are probably achievable at lower intakes\(^{24,50,55}\), in line with even very low increases in fish oil intake, commensurate with human dietary intake, to increase myocardial membrane \(n-3\) PUFA incorporation\(^{23}\). An absence of \(n-3\) PUFA from the diet amplifies the cardiac depressant effects of ischaemic or hypertrophic stress. This can be overcome by incorporating fish oil in the diet to improve myocardial energetics, which is identified as a new candidate approach to the treatment of heart failure\(^{19}\). When combined with the amelioration of risk for sudden heart attack death and antihypertrophic actions, amelioration of heart failure adds a new dimension to the nutritional preconditioning properties of fish oil fatty acids\(^{16}\).

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**References**

Fish oil and heart failure


