Dissociation between exercise-induced reduction in liver fat and changes in hepatic and peripheral glucose homeostasis in obese patients with Non-Alcoholic Fatty Liver Disease

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Dissociation between exercise-induced reduction in liver fat and changes in hepatic and peripheral glucose homeostasis in obese patients with Non-Alcoholic Fatty Liver Disease

Running title: Exercise, liver fat and insulin sensitivity in obese patients with NAFLD

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Abstract

Non-Alcoholic Fatty Liver Disease (NAFLD) is associated with multi-organ (hepatic, skeletal muscle, adipose tissue) insulin resistance (IR). Exercise is an effective treatment for lowering liver fat but its effect on insulin resistance in NAFLD is unknown.

We aimed to determine whether supervised exercise in NAFLD would reduce liver fat and improve hepatic and peripheral (skeletal muscle and adipose tissue) insulin sensitivity. Sixty-nine NAFLD patients were randomised to 16 weeks exercise supervision (n=38) or counselling (n=31) without dietary modification. All participants underwent magnetic resonance imaging/spectroscopy to assess changes in body fat, and in liver and skeletal muscle triglyceride, before and following exercise/counselling. To quantify changes in hepatic and peripheral insulin sensitivity, a pre- determined subset (n=12 per group) underwent a two-stage hyperinsulinaemic euglycaemic clamp pre- and post-intervention. Results are shown as mean (95% CI).

Fifty participants (30 exercise, 20 counselling), 51 y (40, 56), BMI 31 kg/m² (29, 35) with baseline liver fat/water % of 18.8 % (10.7, 34.6) completed the study (12/12 exercise and 7/12 counselling completed the clamp studies). Supervised exercise mediated a greater reduction in liver fat/water % than counselling [\(\Delta\) mean change 4.7% (0.01, 9.4); \(P<0.05\)], which correlated with the change in cardiorespiratory fitness \(r = -0.34, P = 0.0173\).

With exercise, peripheral insulin sensitivity significantly increased (following high-dose insulin) despite no significant change in hepatic glucose production (following low-dose insulin); no changes were observed in the control group.

Although supervised exercise effectively reduced liver fat, improving peripheral IR in NAFLD, the reduction in liver fat was insufficient to improve hepatic IR.

Keywords: NAFLD, insulin resistance, exercise, liver fat and magnetic resonance spectroscopy.

Summary statement

In NAFLD, 16 weeks of supervised exercise effectively reduces liver fat and improves peripheral insulin resistance and cardiorespiratory fitness. Greater reductions in liver fat are needed to improve hepatic insulin resistance, requiring higher intensity or longer duration of exercise.
Introduction

Non-alcoholic fatty liver disease (NAFLD) is a spectrum of histopathological abnormalities which increase the risk of chronic liver disease, hepatocellular carcinoma and cardiovascular disease (1). NAFLD arises from accumulation of liver fat, frequently complicating obesity and other insulin-resistant states, co-existing with the metabolic syndrome (2, 3). NAFLD is associated with multi-organ (hepatic, skeletal muscle and adipose tissue) insulin resistance (IR) (4, 5).

Although certain anti-diabetes agents reduce liver fat (6, 7), the cornerstone of therapy is lifestyle modification through dietary intervention and/or physical activity (8, 9). Weight loss through dietary intervention has been shown to normalise moderate hepatic steatosis (12-13%) and hepatic IR (10, 7411). Considering that NAFLD patients tend to engage in less habitual leisure-time physical activity and be more sedentary, physical activity is also recommended (12, 13). Various modalities of exercise have been shown to be beneficial in reducing liver fat in NAFLD including aerobic (5, 14, 15) and resistance exercise (13), even without weight loss. A recent study addressing the dose-response relationship between aerobic exercise and reduction in liver fat suggests that even low volume, low intensity aerobic exercise can reduce liver fat without clinically significant weight loss (16). It is unclear to what extent reduction in liver fat following exercise is associated with improvements in hepatic and peripheral IR. This is of particular importance considering the high rates of incident type 2 diabetes mellitus (T2DM) in NAFLD patients.

We set out to determine the efficacy of supervised exercise training in reducing liver fat, and the relationship between reduction in liver fat and improvements in hepatic and peripheral IR using the gold standard method for measuring insulin resistance, a 2-step euglycaemic hyperinsulaemic clamp.

Experimental materials and Methods

Design

A 16-week randomised controlled trial of NAFLD patients, randomised to supervised moderate-intensity aerobic exercise or conventional counselling (control group) (Clinical Trials.gov 91NCT01834300).

Participants

Patients were recruited through hepatology clinics where they were undergoing routine clinical care from 4 teaching hospitals, and studied in 2 centres, in Guildford and Liverpool. NAFLD was diagnosed clinically by a hepatologist after exclusion of (steatogenic) drug causes, viral or autoimmune hepatitis (negative hepatitis B and C serology and auto-antibody screen), primary biliary cirrhosis and metabolic disorders (α1-antitrypin deficiency, Wilson’s disease).
Inclusion criteria were a diagnosis of NAFLD, being sedentary (<2 h/week low-intensity physical activity, no moderate- or high-intensity activity), non-smokers, with alcohol consumption <14100(females) and <21 (males) units/week. Exclusion criteria were T2DM, ischaemic heart disease or contraindications to exercise. Participants were excluded from follow-up assessment if they deviated 102from their habitual diet and lost excessive weight.

The study conformed to the Declaration of Helsinki and was approved by the local research ethics committees. All participants provided fully informed written consent.

Protocol

10669 patients were randomly assigned on a 1:1 basis using a computer-generated sequence to 16 weeks supervised exercise or conventional counselling (control group) using SAS v 9.1, PROC PLAN software (Statistical Analysis System Institute, NC, USA). Figure 1 shows the CONSORT diagram.

Supervised Exercise. After a familiarisation session, participants attended the university gymnasium weekly, wearing a heart rate monitor (Polar Electro Oy, Finland) and supervised by a trained exercise physiologist. Training intensity was based on individual heart rate reserve (HRR) ([Maximal HR 112during cardiorespiratory fitness testing] – [Resting HR]). Participants performed 3/week 30 min moderate (30% HRR) aerobic exercise (treadmill, cross-trainer, bike ergometer, rower) progressing 114weekly based on HR responses (5/week 45 min at 60% HRR by week 12). Throughout, participants were monitored via the Wellness System™ (Technogym U.K. Ltd., Bracknell, UK), which tracks exercise activity within designated fitness facilities or by repeated telephone or e-mail contact.

No dietary modifications were made, confirmed by standard 3-day food diaries collected immediately before and after the intervention and analysed for macronutrient intake.

Control Group. Participants were provided with advice about the health benefits of exercise in NAFLD but had no further contact with the research team. To minimise disturbance to behaviour, diet and physical activity were not monitored.

Measurements

Measurements were performed before and immediately after the intervention period. After overnight fast, venous blood was taken for measurement of glucose, liver function, lipid profile, adiponectin and leptin.

After full medical history and physical examination, a single person at each centre measured body weight, blood pressure, height, waist (umbilical) and hip (greater trochanter) circumference and performed bioimpedance analysis (Tanita BC-420MA, Tokyo, Japan).

Magnetic resonance methods were as previously described (17). Volumetric analysis of abdominal subcutaneous adipose tissue (SAT) and abdominal visceral adipose tissue (VAT) used whole-body axial T1-weighted fast spin echo scans (10 mm slice, 10 mm gap), the abdominal region being defined...
from the slices between the femoral heads, top of liver and lung bases. Proton magnetic resonance spectroscopy (1H MRS) quantified intrahepaticocellular lipid (IHCL) and intramyocellular lipid (IMCL). In liver 3 voxels of interest were identified at standardised sites avoiding ducts and vasculature. In skeletal muscle a single voxel was identified in each of the tibialis anterior and soleus muscles, avoiding bone, fascia and neurovascular bundle. Single voxel spectroscopy was conducted at each of these five sites: voxel size was 20×20×20 mm, TE (echo time) 135 msec, TR (repetition time) 1500 msec, with 64 acquisitions. 1H-MR spectra were quantified using the AMARES algorithm in the jMRUI-3.0 (18). Data were processed blind. Liver fat is expressed as the percentage of CH3 lipid signal amplitude relative to water signal amplitude after correcting for T1 and T2 (19), and intramyocellular lipid (IMCL) is expressed as CH2 lipid amplitude relative to total creatine amplitude after correcting for T1 and T2 (20). NAFLD was defined as mean IHCL > 5.3%, which corresponds in the present units (CH3/H2O) to the cut-off of 5.5% by weight advocated on the basis of a large healthy-population 1H MRS study (21) which took account of tissue density, water content and the relative proton densities of triglyceride and water to express IHCL as % by weight in terms more directly comparable with biochemical measurements. This cutoff is also in accordance with traditional definitions of fatty liver based on biochemical analysis (21). (Any IHCL value expressed here as x% CH3/H2O can be converted to y% by weight (i.e., 10 × y mg/g) by using y% = 97.1/[1 + (89.1/x%)]. Based on assumptions and data detailed in (21, 22)).

Clamp. Participants were instructed to avoid strenuous physical activity for 48 h. Upon arrival intravenous cannulae were inserted into both antebrachial fossae for blood sampling and infusion of stable isotopes, insulin and glucose. After unenriched blood samples, a primed infusion of [6,6-2H2] glucose (170 mg; 1.7 mg.min−1) was started. 5 baseline samples were taken from 100-120 min, when a 2-step hyperinsulinaemic–euglycaemic clamp commenced: insulin infusion at 0.3 mU.kg−1.min−1 (low-dose) for 120 min to measure insulin sensitivity of hepatic glucose production (HGP), then at 1.56 U.kg−1.min−1 (high-dose) for 180 min to measure insulin sensitivity of peripheral glucose uptake. Euglycaemia was maintained by adjusting a 20% glucose infusion, spiked with [6,6-2H2] glucose (758 mg·L−1 glucose for low-dose, 10 mg·L−1 high dose) according to 5 min plasma glucose measurements using a glucose oxidase method (Yellow Springs Analyser). Blood samples were taken every 30 min, except for every 5 min from 210-240 min (low-dose steady-state) and 390-420 min (high-dose steady-state).

Plasma glucose concentration and enrichment time-courses were smoothed using optimal segments analysis (23). HGP and glucose uptake (rate of disappearance, Rd) (μmol.kg−1.min−1) were calculated using non-steady-state equations (24), assuming a volume of distribution of 22% body weight. HGP was calculated at steady-state basally (90-120 min) and following low-dose insulin (210-240 min), corrected for fat-free mass and (since HGP is inversely related to [insulin]) multiplied by mean steady-state [insulin] (pmol.ml−1) at low-dose. Glucose Rd was calculated at steady-state following
168 high-dose insulin (390–420 min) and metabolic clearance rate (MCR) (ml.kg⁻¹.min⁻¹) was calculated at
169 basal and high-dose insulin steady-state (390–420 min) as (glucose Rd)/[glucose]. Glucose MCR and
170 Rd were corrected for fat-free mass and (since they are directly related to [insulin]) divided by mean
171 steady-state [insulin] (pmol.l⁻¹) at basal and high-dose.

172 Cardiorespiratory fitness assessment In Liverpool, cardiorespiratory fitness was assessed on a
173 treadmill ergometer following the Bruce protocol (25). Following 2 min warm up at 2.2 km/h on the
174 flat, initial workload was set at 2.7 km/h at 5° grade, then speed and grade increased step-wise every
175 minute. Heart rate and rate of perceived exertion were monitored throughout. VO₂peak was calculated
176 from expired gas fractions (Oxycon Pro, Jaeger, Hochberg, Germany) as the highest consecutive 15 s
177 rate in the last minute before volitional exhaustion, or when heart rate and/or VO₂ reached a plateau
178 (21). In Guildford, VO₂peak was performed on an electronically-braked bicycle ergometer (Lode;
179 Excaliber Sport, Groningen, the Netherlands) with breath analyser (Medical Graphics, St Paul, MN,
180 USA). Heart rate was measured throughout. After 2 min warm up at 50 W, resistance increased step-
181 wise at 20 W/min until volitional exhaustion (26). Cardiorespiratory fitness was defined as VO₂peak
182 identically at each facility (despite the different exercise modalities), expressed per kg body weight.

183 Biochemistry. Baseline plasma samples were analysed using an Olympus AU2700 (Beckman Coulter,
184 High Wycombe, UK) in Liverpool and an Advia 1800 Chemistry System (Siemens Healthcare
185 Diagnostics, Frimley UK) in Guildford, with standard proprietary reagents and methods: glucose with
186 hexokinase, total cholesterol and high-density lipoprotein (HDL) with cholesterol esterase/oxidase,
187 triglyceride with glycerol kinase and liver enzymes including alanine aminotransferase (ALT),
188 aspartate aminotransferase (AST) and gamma-glutamyltransferase (GGT) with International
189 Federation of Clinical Chemistry (IFCC) kinetic UV (without pyridoxal phosphate activation). Intra-
190 and inter-assay coefficients of variation were ≤10%. Low-density lipoprotein (LDL) was calculated
191 using the Friedwald formula. At a single centre, serum insulin, plasma adiponectin and leptin were
192 measured by RIA using commercial kits (Millipore Corporation, Billerica, MA; intra-assay CV 6%,
193 35%, 5% respectively), irisin by ELISA (Phoenix Pharmaceuticals, Inc. Burlingame, CA; intra-assay
194 CV 4.1%), fetuin-A by ELISA (Epitope Diagnostics, Inc. San Diego; intra-assay CV 4.8%) and serum
195 NEFA (Wako Chemicals, Neuss, Germany; inter-assay CV 3.0%). Glucose isotopic enrichment was
196 measured by GC-MS on a HP 5971A MSD (Agilent Technologies, Wokingham, Berks, UK) (27). IR
197 was quantified using HOMA2-IR (28). Indices of hepatic insulin resistance (Hepatic-IR) and adipose
198 tissue insulin resistance (Adipose-IR) were calculated (29, 30).

199 Diagnosis of metabolic syndrome was based on the National Cholesterol Education Program Adult
200 Treatment Panel III criteria (31). Ten-year cardiovascular risk was calculated using the 10 year
201 Framingham Risk Score (32).

202 Statistical Analysis
Power calculation. The primary outcome variable was IHCL (% fat/water). Based on mean IHCL of 20420%, we considered 30% relative difference between groups to be clinically significant, implying 205mean IHCL of 20% and 14% in the control and exercise groups respectively. Based on a 2-sample t- 206test, 5% 2-sided significance and standard deviation (SD) of 7.75% from previous studies, 56 patients 207(28 in each arm) were required to detect this 6% absolute IHCL difference with 80% power (27).

Statistical methods. For the primary comparison of supervised exercise vs. control, delta (Δ) change 209from pre-intervention was calculated and analysed using linear regression (ANCOVA), with pre data 210as a covariate (33). Linear regression assumptions were assessed using Q-Q plots and scatter plots of 211studentised residuals versus fitted values. Where linear regression assumptions were not met these 212were resolved using the natural logarithm transformation. For exploratory and comparison purposes 213any continuous demographic variable within each group was also estimated using a paired t-test. 214Correlations were quantified using Spearman’s Rank correlation coefficient (r_s). Data for continuous 215demographic variables are presented as median and inter-quartile range (IQR) and changes between 216supervised exercise compared to control are presented as mean (95% CI). Statistical analyses used 217Stata 13 (StataCorp. 2013. Stata Statistical Software: Release 13. College Station, TX: StataCorp LP). 218Unless otherwise stated, exact P-values are cited (values of “0.000” are reported as “<0.001”). Results 219are shown as mean (95% CI).

Results

Baseline characteristics Fifty patients completed the study [n=30 exercise (23 males, 7 female) and 222n=20 control (16 males, 4 female)] (Figure 1). The age of the participants was similar in the exercise 223[50y (46, 58), BMI 30.7 kg/m² (29.2,32.9)] vs. control groups [52y (46, 59), BMI 29.7 kg/m² 224(28.0,33.8)]. An equal number (n=15) completed the exercise in each centre (total exercise=30); 8 225controls completed in Liverpool and 12 controls completed in Guildford, Surrey (total controls n=20). 226Pre-intervention characteristics of the groups were similar with respect to age, VO_2peak, biochemical 227and metabolic characteristics, and body composition (Tables 1 and 2).

Changes in dietary intake: In the exercise group after 16 weeks, total energy intake and macronutrient 229composition remained unchanged compared with baseline: energy [0.4 MJ (-0.4, 1.2), P=0.40], 230protein [0.4 g (-11.6, 12.0), P=0.97], carbohydrate [6.4 g (-24.2, 37.0), P=0.34], sugars [-9.2 g (-27.2, 23130.0), P=0.41] and fat [9.8 g (8.5, 22.0), P=0.44].

Changes in body composition and biochemistry The primary outcome measure of IHCL in the 232exercise group was significantly reduced after 16 weeks: 19.4% (14.6, 36.1) vs. 10.1% (6.5, 27.1), but 233not in the control group: 16.0% (9.6, 32.5) % vs. 14.6 (8.8, 27.3). Supervised exercise mediated a 234greater IHCL reduction than in the controls [-4.7 % (-9.4, -0.01); P<0.05] (Table 2). Changes in ALT, 236AST and in GGT were not significant.
237SAT reduction with exercise was significantly greater than with control [-1.8L (= -3.0, -0.7); 238P=0.003], but changes in VAT were not [-0.7L (-1.6, 0.2); P<0.109], and nor were changes in IMCL in 239soleus and tibialis anterior (Table 1).

240The changes in fasting plasma insulin and HOMA2-IR [-0.5 (-1.0, 0.02; P=0.06] with exercise were 241not significantly different compared with control, nor were those in adiponectin, leptin, irisin or fetuin 242(Table 2).

243Changes in cardiorespiratory fitness Cardiorespiratory fitness (expressed as ml/kg/min) significantly 244improved in the exercise group after 16 weeks: 23.7 ml/kg/min (21.7, 27.8) vs. 32.3 ml/kg/min (27.6, 24538.0); there was no significant increase in the control group: 23.2 ml/kg/min (20.9, 25.6) vs. 23.1 246ml/kg/min (20.9, 26.9). Exercise mediated a greater improvement compared to control [7.3 ml/kg/min 247(5.0, 9.7); P<0.001].

248Cardiorespiratory fitness (expressed as absolute values in l/min) significantly improved in the exercise 249group after 16 weeks: 2.45 l/min (2.22, 2.69) vs. 3.05 l/min (2.77, 3.34); there was no significant 250increase in the control group: 2.31 l/min (2.05, 2.63) vs. 2.30 l/min (2.04, 2.57). Exercise mediated a 251greater improvement compared to control [0.72 l/min (0.42, 1.02); P<0.001].

252The greater fitness improvement was accompanied by greater reductions in total body weight [-2.5 kg 253(-3.9, -1.1); P<0.001], waist circumference [-3.0 cm (-5.5, -1); P<0.05] and percentage fat mass [-1.9% 254(-3.0, -0.7]; P<0.01] compared to control (Table 1). Changes in IHCL were significantly correlated 255with improvements in cardiorespiratory fitness (absolute and relative), total body weight and with 256 reductions in visceral and subcutaneous fat (Figure 2).

257Changes in peripheral and hepatic insulin sensitivity In the subset of 24 patients that underwent the 2- 258stage hyperinsulinaemic euglycaemic clamp, 12 patients in the exercise group and 7 patients in the 259controls completed the full clamp measurements. The changes in this exercise and control subset were 260similar to those seen in the whole group: [Liver fat, -9.3% (-18.1, 0.5) vs. 3.5% (-11.1, 3.9)] and 261VO2peak [7.7ml/kg/min (4.0, 11.1) vs. -1.4ml/kg/min (-4.4, 1.6)].

262Plasma glucose concentration at basal and during the clamp did not differ between interventions (data 263not shown). In the exercise group glucose infusion rate, corrected for [insulin], during the high-dose 264insulin infusion was higher post-exercise (P=0.009) (Figure 3a) but did not change in the control 265group. Following high-dose insulin infusion there was a significant increase in glucose Rd and MCR, 266corrected for [insulin] in the exercise group (P=0.02, P=0.004 respectively) with no significant 267change in the control group (Figure 3b and c). The change in glucose MCR was significantly different 268between groups (P=0.03).

269There was no significant difference with either intervention in HGP corrected for [insulin] at baseline 270or after low-dose insulin, (Figure 3d) or in the percentage decrease in HGP following low-dose insulin
in either the exercise group (pre-exercise 50.9±5.3%; post-exercise 55.3±6.4%) or the control group 272(pre 46.5±10.3%; post 56.0±8.5%).

Changes in glucose MCR, corrected for insulin, under basal conditions were significantly correlated 274with changes in fitness (r = 0.48, P = 0.04) but not in IHCL (r = 0.26, P = 0.28). After high-dose insulin, 275the correlation with IHCL did not reach statistical significance (r = 0.43; P = 0.18).

Discussion

We have demonstrated in a randomised controlled study that 16 weeks of supervised moderate- 278intensity aerobic exercise in NAFLD reduces liver fat and that this was correlated with an 279improvement in cardiorespiratory fitness. Using a 2-step euglycaemic hyperinsulinaemic clamp in 280conjunction with quantification of liver fat, we showed, for the first time in patients with NAFLD, that 281the exercise-induced reduction in liver fat was accompanied by enhanced skeletal muscle and adipose 282tissue insulin sensitivity, with no improvement in hepatic glucose production.

Various factors modulate liver fat, particularly regular physical activity (34, 35). Numerous studies 284have highlighted the therapeutic effects of endurance or resistance exercise in lowering liver fat in 285NAFLD, even without weight loss (15). However modest weight loss also has clinically significant 286effects on IHCL. In a study by Coker et al., measuring multi-organ insulin sensitivity in caloric 287restriction and exercise training (with and without weight loss), exercise with weight loss had the 288greatest effect both on visceral fat and hepatic glucose output suppression (36). However, liver fat was 289not measured, precluding direct comparison with the current study.

In the current study, exercising participants lost ~3% of body weight and this will have contributed to 291the reduction in IHCL. In a 2-week dietary intervention in NAFLD, ~4% weight reduction was 292associated with 42% reduction in liver fat (37) while in the LOOK-AHEAD study, lifestyle 293intervention in T2DM resulting in 1-5% weight change produced 33% reduction in hepatic steatosis 294(14). While there are clearly weight-dependent effects, the correlation between a reduction in liver fat 295and improvement in cardiorespiratory fitness in the supervised exercise group suggests that the latter 296also is a major driver of IHCL levels.

A significant improvement in peripheral (skeletal muscle and adipose) insulin sensitivity 298accompanied the reduction in liver fat following exercise. It is well documented that chronic exercise 299improves peripheral insulin sensitivity (38, 39). The improvement in peripheral insulin sensitivity 300following exercise training occurred without any change in intramyocellular lipid as has been shown 301in a previous study of overweight men (23). Petersen et al. (40), proposed that skeletal muscle IR 302promotes hepatic steatosis and metabolic syndrome, by altering post-prandial energy distribution, 303diverting glucose to the liver for de novo lipogenesis (DNL) and triglyceride synthesis. Furthermore, 304acute exercise through reversal of muscle IR, has been shown to reduce hepatic DNL by 30% and
hepatic triglyceride synthesis by 40% (41). In myostatin-null mice, increased muscle insulin sensitivity also protects against hepatic steatosis during high-fat feeding (42). Thus, skeletal muscle metabolism may influence hepatic triglyceride content and metabolism, with inter-organ ‘cross-talk’ between skeletal muscle, adipose tissue and liver (43). Although not measured here, myokines secreted by skeletal muscle after contraction appear to mediate this cross talk. Thus, a plausible mechanism in our study for the reduction in liver fat is enhanced peripheral insulin sensitivity and increased skeletal muscle glucose uptake reducing the flux of plasma glucose to the liver for triglyceride synthesis. The critical role of adipose IR in the metabolic and histological changes in NAFLD, as well as its reversal using thiazolidinediones, has also been demonstrated (29, 44). In this study, we showed that adipose-IR could also be improved with exercise training.

The lack of effect of the exercise programme on hepatic insulin resistance was surprising given the assumed links between liver fat accumulation and defective insulin suppression of glucose production (4, 45). Other studies have reported reduced hepatic steatosis and improved hepatic insulin resistance with weight loss following low calorie diets in NAFLD (10,11). However, in these studies liver fat was lower than in the current study and was reduced to normal by weight loss, from 12 to 2.5% (10) and from 12.8 to 2.9% (11). Although in our study there was a comparable loss of liver fat in the exercise group (9.3%) because the group had much higher liver fat levels at baseline (median 19.4%) many patients remained above the normal range after 16 weeks exercise. This suggests that greater reductions in liver fat are needed to improve hepatic insulin resistance, possibly to within the normal range. It is likely that this could be achieved by increasing the period of exercise supervision or the intensity of the exercise, or by caloric restriction (46). Sullivan et al. noted a similar dissociation between (reduced) liver fat and (unchanged) VLDL triglyceride synthesis rate, a metabolic pathway that also exhibits resistance to insulin, after exercise training in patients with NAFLD. Interestingly in the latter study, % liver fat was similar at baseline to the current study (5). Recent animal data may help provide a mechanistic explanation for the phenomenon of improved peripheral insulin sensitivity, reduced liver fat but impaired hepatic insulin sensitivity of glucose metabolism. This data suggests that within the liver glucose production and de novo lipogenesis have different insulin sensitivities: the gluconeogenic pathway is insulin-resistant (thus insulin cannot inhibit hepatic glucose production through gluconeogenesis) while the lipogenic pathway remains insulin-sensitive (thus insulin retains its ability to stimulate fatty acid synthesis) (47). This selective insulin resistance is explained by a bifurcation of the hepatic insulin signalling pathway: control of the repression of gluconeogenesis occurs through FoxO1, while a separate pathway controlling lipogenesis involves SREBP-1c (48). Although this cannot be tested in the current study, this mechanism would provide a plausible explanation for the dissociation of the effects of exercise on hepatic liver fat and hepatic glucose production.
We acknowledge limitations to the study. We used a per protocol analysis. The drop-out rate (19/69, 34128%) was higher than the anticipated 15-20%, 15 controls and 4 in the exercise group, apparently
mainly for practical reasons (e.g. time constraints, excessive research burden) but we believe the
disproportionately higher dropout rate in the control group reflects many participants’ underlying
desire to be randomised to the exercise program. The higher dropout rate in the control group is, we
cautiously argue, unlikely to bias our conclusion, and will of course not affect assessment of the effect
of the exercise intervention per se. A further imitation is that cardiorespiratory fitness was assessed at
study sites using two different modalities, treadmill vs. cycle ergometer. Whilst cardiorespiratory
fitness may be lower using cycle ergometry, the primary comparison was the change in fitness with
intervention, thus this is unlikely to bias our findings. This is likely due to the greater spread of
VO_{peak} results given the improvements post exercise training. While we believe our cohort is
representative of the general NAFLD population, there may be a selection bias with only the most
motivated patients consenting to participate in an exercise intervention study: this may underlie the
high dropout rate of controls. Accepting these limitations, the noteworthy strengths are the application
of whole body MRI and 1H-MRS, the most sensitive, non-invasive method to quantitate liver fat, and
measurement of corresponding changes in organ-specific insulin sensitivity. Using these gold standard
techniques we provide important insight into the mechanism by which exercise mediates reduction in
liver fat by enhanced peripheral (skeletal muscle) insulin sensitivity, without this reduction in liver fat
being paralleled by improved hepatic insulin sensitivity.

In summary, in patients with NAFLD exercise-induced reduction in liver fat is related to the
improvement in cardiorespiratory fitness and accompanied by an improvement of peripheral (muscle
and adipose) but not hepatic IR. The greatest benefit in normalising liver fat, improving both
peripheral and hepatic IR and potentially providing the greatest protection against incident T2DM,
may require increasing the duration and/or intensity of the exercise supervision, in conjunction with
caloric restriction.

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Declaration of interest
The authors have nothing to declare.
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Author contribution statement

DC, FSM, AMU and GJK conceived and designed the study protocol, obtained funding, were involved in collection and analysis of data and wrote the manuscript. VSS, CJP, HJ, MB, PR, MB, NCJ, ELT and JDB were involved in collection and analysis of data and contributed to the editing of the manuscript.

Clinical Perspectives

NAFLD represents a common obesity-related complication, increasing the risk of type 2 diabetes mellitus, cardiovascular disease and chronic liver disease. Exercise interventions are effective in reducing liver fat, even without significant weight loss.

We demonstrate exercise supervision is effective at reducing liver fat and this was related to an improvement in cardiorespiratory fitness. As expected exercise was associated with significant improvements in peripheral (skeletal muscle and adipose tissue) insulin resistance.

Surprisingly, despite significant reductions in liver fat with exercise, we did not observe an improvement in hepatic insulin resistance. We speculate that persisting elevated liver fat even after exercise training, means undiminished hepatic insulin resistance. Exercise training needs to be more prolonged or more intense to achieve a greater reduction in liver fat. These results have potential public health implications considering the associated long-term metabolic, hepatic and cardiovascular complications.
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444 Resistance exercise reduces liver fat and its mediators in non-alcoholic fatty liver disease


Figure legends

Figure 1. CONSORT diagram showing flow of participants through the study.

Figure 2. Black circles indicate individuals in the exercise group; open circles indicate individuals in the control group.

A) Relationship between reduction in liver fat (IHCL) and improvement in cardiopulmonary fitness (VO\textsubscript{peak} ml.kg\textsuperscript{-1}.min\textsuperscript{-1}) \((r= -0.34; P=0.02)\)

B) Relationship between reduction in IHCL and reduction in body weight \((r=0.48; P<0.001)\)

C) Relationship between reduction in IHCL and reduction in visceral adipose tissue volume \((r=0.37; P=0.008)\).

D) Relationship between reduction in IHCL and reduction in subcutaneous adipose tissue volume \((r=0.61; P<0.001)\).

Figure 3. Rates of a) glucose infusion (GINF) during high dose insulin, b) glucose uptake (Rd) during high dose insulin, c) glucose metabolic clearance (MCR) during high dose insulin and d) hepatic glucose production (HGP) during low dose insulin expressed relative to insulin, before (grey bars) and after (black bars) exercise or controls.

565
Assessed for eligibility ($n=100$)

Excluded ($n=31$) including
- Not meeting inclusion criteria ($n=24$)
- Declined to participate ($n=7$)

Randomised ($n=69$)

Supervised exercise training ($n=38$)
- Discontinued baseline assessment ($n=3$)
- Discontinued intervention ($n=4$)
- Did not maintain habitual diet ($n=1$)
- Declined post intervention assessments ($n=0$)
- Analysed ($n=30$)
  - Excluded from analysis ($n=0$)

Controls ($n=31$)
- Discontinued baseline assessment ($n=1$)
- Discontinued intervention ($n=0$)
- Did not maintain habitual diet ($n=1$)
- Declined post intervention assessments ($n=9$)
- Analysed ($n=20$)
  - Excluded from analysis ($n=0$)

Figure 2
<table>
<thead>
<tr>
<th>Within-group comparison</th>
<th>Between-group comparison</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ex Δ Change (Mean 95% CI)</td>
</tr>
<tr>
<td><strong>Pre Ex</strong></td>
<td><strong>Post Ex</strong></td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>95.6 (83.8-104)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>30.6 (29.0-32.9)</td>
</tr>
<tr>
<td>Waist (cm)</td>
<td>106 (101-112)</td>
</tr>
<tr>
<td>% fat mass</td>
<td>30.4 (25.9-32.1)</td>
</tr>
<tr>
<td>Systolic BP (mmHg)</td>
<td>135 (125-142)</td>
</tr>
<tr>
<td>Diastolic BP</td>
<td>83 (75-87)</td>
</tr>
<tr>
<td>VO₂peak(ml/kg/min)</td>
<td>23.7 (21.7-27.8)</td>
</tr>
<tr>
<td>ALT* (U/l)</td>
<td>45 (36-66)</td>
</tr>
<tr>
<td>AST* (U/l)</td>
<td>33 (25-47)</td>
</tr>
<tr>
<td>GGT* (U/l)</td>
<td>47 (35-62)</td>
</tr>
<tr>
<td>Cholesterol (mmol/l)</td>
<td>5.1 (4.7-5.7)</td>
</tr>
<tr>
<td>Triglycerides (mmol/l)</td>
<td>1.9 (1.4-2.63)</td>
</tr>
<tr>
<td>HDL (mmol/l)</td>
<td>1.2 (0.9-1.4)</td>
</tr>
<tr>
<td>LDL (mmol/l)</td>
<td>3.5 (3.0-3.9)</td>
</tr>
<tr>
<td>Chol:HD L ratio</td>
<td>4.6 (4.0-5.1)</td>
</tr>
<tr>
<td>Liver fat (%)</td>
<td>19.4 (14.6-36.1)</td>
</tr>
<tr>
<td>CH₃/water</td>
<td>9.8 (8.0-11.7)</td>
</tr>
<tr>
<td>SAT (l)</td>
<td>23.1 (19.4-32.0)</td>
</tr>
<tr>
<td>Abdominal fat (l)</td>
<td>33.2 (29.1-41.0)</td>
</tr>
<tr>
<td>VAT:SAT ratio</td>
<td>0.4 (0.3-0.6)</td>
</tr>
<tr>
<td>IMCL Soleus (CH₃/creatinine)</td>
<td>12.3 (9.0-16.8)</td>
</tr>
<tr>
<td>IMCL Tibialis Ante</td>
<td>9.0 (5.6-8.6)</td>
</tr>
<tr>
<td>Within-group comparison</td>
<td>Between-group comparison</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-------------------------</td>
</tr>
<tr>
<td>Pre Ex Median (IQR)</td>
<td>Post Ex Median (IQR)</td>
</tr>
<tr>
<td>11.2</td>
<td></td>
</tr>
</tbody>
</table>

Within-group comparisons use paired t-tests, p < 0.05 being taken as evidence of a significant change pre- to post-intervention: a negative change indicates reduction pre- to post. Between-group comparisons (final two columns) use linear regression (ANCOVA) comparing post-scores between groups correcting for pre-scores, Δ therefore indicates the difference between post-intervention means after correcting for pre-intervention scores: a negative difference indicates a lower mean for the exercise group compared with control. ^ indicates that a log transformation was necessary to meet the assumptions of linear regression; here, Δ is the ratio of geometric means post-intervention after correcting for pre-intervention scores, a ratio <1 indicating a lower mean in exercise group relative to control.
Table 2. Metabolic measurements in 50 patients before and after supervised exercise intervention (Ex; n=30) and control (Con; n=20) (reported as median and interquartile range as within group comparison). Mean delta changes with 95% confidence intervals (with significance values) are shown for each intervention and the delta changes are compared (between group comparison). *P<0.05.

<table>
<thead>
<tr>
<th>Test</th>
<th>Pre Ex Median (IQR)</th>
<th>Post Ex Median (IQR)</th>
<th>Pre Con Median (IQR)</th>
<th>Post Con Median (IQR)</th>
<th>Ex Δ Change Mean (95% CI)</th>
<th>Con Δ Change Mean (95% CI)</th>
<th>Δ Mean (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fasting glucose (mmol/l)</td>
<td>5.4 (4.8-6.1)</td>
<td>5.3 (4.9-5.7)*</td>
<td>5.6 (4.8-6.1)</td>
<td>5.5 (5.0-5.8)*</td>
<td>-0.15 (-0.30, 0.00)</td>
<td>-0.2 (-0.3, 0.0)</td>
<td>0.0 (-0.2, 0.2)</td>
</tr>
<tr>
<td>Fasting insulin (pmol/l)</td>
<td>131 (96-162)</td>
<td>115 (72-158)*</td>
<td>119 (96-193)</td>
<td>130 (95-195)</td>
<td>22 (-43, -1)</td>
<td>2 (-19, 23)</td>
<td>-26 (-55, 2)</td>
</tr>
<tr>
<td>HOMA2-IR</td>
<td>2.5 (1.8-3.0)</td>
<td>2.1 (1.3-2.9)*</td>
<td>2.2 (1.8-3.6)</td>
<td>2.5 (1.8-3.7)</td>
<td>-0.43 (-0.81, -0.05)</td>
<td>0.03 (-0.3, 0.4)</td>
<td>-0.5 (-0.4, 0.0)</td>
</tr>
<tr>
<td>Fasting FFA (mmol/l)</td>
<td>0.52 (0.45-0.60)</td>
<td>0.42 (0.35-0.59)</td>
<td>0.56 (0.39-0.71)</td>
<td>0.54 (0.32-0.65)</td>
<td>-0.04 (-0.11, 0.03)</td>
<td>-0.03 (-0.08, 0.03)</td>
<td>-0.03 (-0.1, 0.1)</td>
</tr>
<tr>
<td>Adipose–IR (mmol/l, pmol/l)</td>
<td>61 (48-88)</td>
<td>50 (30-86)*</td>
<td>55 (47-87)</td>
<td>60 (44-84)</td>
<td>-15 (-27, 2)</td>
<td>-0.5 (-17, 16)</td>
<td>-18 (-36, 0.5)*</td>
</tr>
<tr>
<td>Adiponectin (ng/ml)</td>
<td>5950 (3700-8100)</td>
<td>5450 (3550-7650)</td>
<td>6300 (5200-7950)</td>
<td>6650 (4950-9750)</td>
<td>-260 (-790, 269)</td>
<td>259 (-543, 1060)</td>
<td>-630 (-1497, 238)</td>
</tr>
<tr>
<td>Leptin (ng/ml)</td>
<td>9.2 (6.5-12.6)</td>
<td>7.1 (4.3-11.9)*</td>
<td>11.8 (7.0-18.5)</td>
<td>11.8 (6.9-19.0)</td>
<td>-1.7 (-3.0, -0.4)*</td>
<td>-0.3 (-1.5, 1.0)</td>
<td>-1.7 (-3.5, 0.1)</td>
</tr>
<tr>
<td>Irisin (ng/ml)</td>
<td>140 (128-171)</td>
<td>129 (121-173)*</td>
<td>140 (128-179)</td>
<td>145 (123-156)</td>
<td>-10.5 (-18.9, -2.1)</td>
<td>-5.4 (-16, 5.1)</td>
<td>-4.7 (-17, 8)</td>
</tr>
<tr>
<td>Fetuin-A *(μg/ml)</td>
<td>483 (412-518)</td>
<td>470 (397-506)</td>
<td>424 (393.8)</td>
<td>428 (394-477)</td>
<td>-1.9 (-15.5, 11.6)</td>
<td>-4.0 (27, 19)</td>
<td>-2.0 (-28, 24)</td>
</tr>
</tbody>
</table>

Within-group comparisons use paired t-tests, P<0.05 being taken as evidence of a change pre- to post-intervention: a negative change indicates reduction pre- to post.
Between-group comparisons use linear regression (ANCOVA) comparing post scores between groups whilst correcting for pre-scores, therefore indicates the difference between post intervention means after correcting for pre-intervention scores: a negative difference indicates a lower mean for the exercise group compared with control group.