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1	High-fat overfeeding impairs peripheral glucose metabolism and muscle microvascular
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51 Abstract

52 *Context:* The mechanisms responsible for dietary fat-induced insulin resistance of skeletal 53 muscle and its microvasculature are only partially understood.

54 *Objective:* To determine the impact of high-fat overfeeding on postprandial glucose fluxes,
55 muscle insulin signaling, and muscle microvascular eNOS content and activation.

56 Design: Fifteen non-obese volunteers consumed a high-fat (64%) high-energy (+47%) diet for 57 7 days. Experiments were performed before and after the diet. Stable isotope tracers were used 58 to determine glucose fluxes in response to carbohydrate plus protein ingestion. Muscle insulin 59 signaling was determined as well as the content and activation state of muscle microvascular 60 eNOS.

Results: High-fat overfeeding impaired postprandial glycemic control as demonstrated by 61 62 higher concentrations of glucose (+11%; P = 0.004) and insulin (+19%; P = 0.035). 63 Carbohydrate plus protein ingestion suppressed endogenous glucose production to a similar extent before and after the diet. Conversely, high-fat overfeeding reduced whole body glucose 64 65 clearance (-16%; P = 0.021) and peripheral insulin sensitivity (-26%; P = 0.006). This occurred despite only minor alterations in skeletal muscle insulin signaling. High-fat overfeeding 66 reduced eNOS content in terminal arterioles (P = 0.017) and abolished the increase in eNOS 67 68 Ser1177 phosphorylation that was seen after carbohydrate plus protein ingestion.

69 *Conclusion:* High-fat overfeeding impaired whole-body glycemic control due to reduced 70 glucose clearance, not elevated endogenous glucose production. The finding that high-fat 71 overfeeding abolished insulin-mediated eNOS Ser1177 phosphorylation in the terminal 72 arterioles suggests that impairments in the vasodilatory capacity of the skeletal muscle 73 microvasculature may contribute to early dietary fat-induced impairments in glycemic control.

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76	Précis								
77	Healthy vo	olunte	ers consum	ned a high-fat diet fo	or 7 day	s. The diet impaire	ed peripl	neral	glucose
78	clearance	and	abolished	insulin-stimulated	eNOS	phosphorylation	within	the	muscle
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101 Introduction

Excessive food intake and physical inactivity have driven the obesity epidemic, with obesity being a major risk factor for the development of insulin resistance and the metabolic syndrome 1. However, just a few days of excessive dietary fat intake can impair insulin action and glycemic control in healthy non-obese individuals 2-4. Understanding these early responses may provide insight into metabolic disease progression.

107

108 Impairments in glycemic control may be mediated by the inability of insulin to suppress 109 endogenous glucose production (EGP) in the liver and/or a reduction in insulin-stimulated 110 glucose uptake by peripheral tissues 5. The tissue-specific contributions to dietary fat-induced 111 impairments in glycemic control are only partially understood. One study reported a reduction 112 in hepatic insulin sensitivity and an increase in basal EGP in healthy men subjected to 5-days 113 of high-fat overfeeding 6. In that study, high-fat overfeeding had no effect on insulin-stimulated 114 glucose disposal, suggesting that changes in hepatic glucose metabolism precede that of 115 peripheral impairments. In contrast, another study reported a reduction in insulin-stimulated 116 leg glucose uptake after 3-days of high-fat overfeeding, but no change in hepatic insulin 117 sensitivity or EGP at baseline or following insulin administration 3. Each of these studies 118 utilized the hyperinsulinemic-euglycemic clamp, which, although useful, can be criticized due 119 to its failure to mimic postprandial conditions. Under clamp conditions, skeletal muscle is 120 responsible for 70-80% of glucose disposal, and EGP is completely suppressed. Following 121 glucose ingestion, splanchnic extraction and skeletal muscle glucose uptake make roughly 122 equal contributions to meal-derived glucose disposal (~30% each), and EGP is only partially 123 suppressed (50-60% decrease) 7-10. Thus, the processes governing postprandial glycemic 124 control are more dynamic/ complex than that of clamps. Moreover, as the normal route for 125 glucose entry into the body is via the gastrointestinal tract, it is necessary to determine the impact of high-fat overfeeding on postprandial glucose fluxes (i.e., meal-derived glucose entry
into the circulation, suppression of EGP, and stimulation of glucose disposal), which can be
achieved through the use of dual-glucose tracers 7,10.

129

As well as the classical actions of insulin on the myocyte, recent work has established a possible 130 131 role for the skeletal muscle microvasculature in insulin-mediated glucose disposal 11. 132 Physiological doses of insulin have been shown to increase skeletal muscle perfusion 12-14. 133 Inhibition of this hemodynamic action by L-NAME ($N(\omega)$ -nitro-L-arginine-methyl ester) or L-134 NMMA (NG-monomethyl-L-arginine acetate) has been shown to reduce skeletal muscle 135 glucose uptake 15-17. Alternatively, it has been shown that L-NAME administration can impair 136 glycemic control through inhibition of insulin secretion without changes in peripheral insulin 137 sensitivity 18, and another study reported that L-NMMA administration prevented vasodilation 138 during insulin/glucose infusion but did not alter whole body glucose uptake 19. Thus, the role of the skeletal muscle microvasculature in insulin-mediated glucose disposal requires 139 140 clarification. Animal studies suggest that impairments to insulin's microvascular action could 141 be a key early event in the development of insulin resistance in response to a high fat diet 20-22. 142 Importantly, the effect of insulin on the microvasculature is dependent on nitric oxide (NO) 143 synthesized in the endothelium of terminal arterioles. Terminal arterioles regulate the blood 144 flow in microvascular units (MVUs), which are the smallest functional elements to adjust 145 muscle blood flow in response to physiological signals and metabolic demands of the muscle 146 fibers 23. Each terminal arteriole delivers blood to \pm 20 capillaries 23. Increases in insulin 147 following meal ingestion activate eNOS by means of Ser1177 phosphorylation. This leads to the 148 production of NO, relaxation of the smooth muscle layer and vasodilation of the terminal 149 arterioles, thereby increasing blood flow in the MVU's 24. The content of NAD(P)H oxidase in 150 the endothelial layer of the terminal arterioles may reduce NO bioavailability, and therefore microvascular perfusion of the muscle, through NO scavenging by superoxide anions 25,26.
Despite the potential for impaired microvascular perfusion playing a role in lipid-induced
insulin resistance, no studies have investigated the protein content and activation state of eNOS
and NAD(P)H oxidase in response to a high fat diet.

The present study determined the role of EGP (primarily hepatic), oral glucose appearance, and whole-body glucose clearance in dysregulation of glycemic control after 7 days of high-fat overfeeding. To this end, we used stable isotope tracers to assess glucose fluxes in response to acute oral ingestion of an insulinotropic carbohydrate-protein mixture. We also determined the phosphorylation (activation) of key proteins involved in skeletal muscle insulin signaling as well as the protein content and activation state of eNOS and NADP(H) oxidase. We hypothesized that high-fat overfeeding would impair glucose clearance, not endogenous glucose production. We also hypothesized that high-fat overfeeding would impair insulin-stimulated eNOS Ser1177 phosphorylation, thereby identifying the muscle microvasculature as an early possible cause of dietary fat-induced insulin resistance.

176 Methods

177 **Participants**

Fifteen individuals (13 males/ 2 females) age 24 ± 1 y, height 176.1 ± 2.1 cm, body mass 77.15 ± 3.07 kg, and body mass index (BMI) 24.8 ± 0.6 kg/m² participated in this study. Participants were physically active, non-smokers, with no diagnosis of cardiovascular or metabolic disease, not taking any medication, and weight stable for ≥ 3 months. The study was approved by the local ethics committee and every participant provided written informed consent. Experimental trials were conducted from March 2015 to August 2016. Analysis was completed by May 2018. The study was registered at ClinicalTrials.gov (identifier: NCT03879187).

185

186 **Pre-testing**

Participants attended the laboratory for assessment of anthropometric characteristics (height,
body mass and BMI). This information was used to estimate resting energy expenditure (REE)
27. A standard correction for physical activity (1.6 and 1.7 times REE for females and males,
respectively) was applied to estimate total daily energy requirements. This information was
used to determine individual energy intakes for the diet intervention.

192

193 Experimental design

One-week after the pre-testing visit, participants returned to the laboratory to undergo metabolic testing. Participants then consumed a high-fat, high-energy diet for 7 days. The diet provided 4749 ± 181 kcal per day, with 188 ± 8 g [16% total energy (TE)] protein, 237 ± 8 g [20% TE] carbohydrate, and 333 ± 14 g [64% TE] fat intake. All foods were purchased and prepared by the research team. Participants were instructed to eat everything that was provided, not to eat any additional food, and to return any uneaten items so that diet values could be adjusted if necessary. All participants were informed about the importance of strict diet adherence. Adherence was checked by daily interviews that were conducted when participants
collected their food bundles. A second metabolic testing session was conducted the morning
after completing the diet.

204

205 Experimental protocol

206 Participants reported to the laboratory in the morning after an overnight fast (≥ 12 h), having 207 refrained from strenuous physical activity for \geq 48 h. After voiding and being weighed, a Teflon 208 catheter (Venflon; Becton, Dickinson, Plymouth, UK) was inserted into an antecubital vein of 209 each arm to allow blood sampling and isotope infusion. A baseline blood sample (10 mL) was 210 obtained before a primed constant infusion of [6,6-2H2]glucose was initiated and continued for 211 the duration of the experiment. Blood samples were divided between tubes containing EDTA 212 or a clotting catalyst (Sarstedt, Leicester, UK). Further blood samples were obtained 90, 105 213 and 120 min into the infusion (referred to as t = -30, -15 and 0 min in results/ figures). Biopsies 214 were obtained from the *vastus lateralis* under local anesthesia (Lidocaine 10 mg/mL) using a 215 5-mm Bergström needle, modified for use with manual suction. Two sections of muscle tissue 216 were blotted free of blood, snap-frozen in liquid nitrogen, and stored at -80°C. A third section 217 of muscle tissue was mounted in Tissue-Tek OCT (Sakura Finetek UK Ltd, Thatcham, UK) 218 and frozen in liquid nitrogen-cooled isopentane for cryo-sectioning and immunofluorescence 219 microscopy. Participants then consumed a carbohydrate plus protein solution. Further blood samples were obtained at 15, 30, 45, 60, 90- and 120-min post-ingestion, and additional muscle 220 221 biopsies were obtained at 30- and 120-min post-ingestion.

222

Two participants did not undergo the muscle biopsy procedure, so measurements of skeletal muscle insulin signaling are reported for n = 13. Data for immunofluorescence microscopy are reported for n = 12 and data for muscle glycogen are reported for n = 11 due to tissue quantity 226 limitations. Data for glucose kinetics are reported for n = 14 due to a technical issue with the 227 preparation of isotopes for one of the trials. All other data reported for n = 15.

228

229 Carbohydrate plus protein beverage

The test beverage was a 12.5% glucose solution (48.4 g glucose plus 1.6 g [U-13C]glucose in 400 mL of water) with 15 g whey protein (UltraWhey 90; Volac, Hertfordshire, UK). Whey protein was added to the solution as most meals will contain protein and to increase the insulinotropic effect of the beverage.

234

235 Blood analyses

Plasma samples were analyzed for triacylglycerol (TAG), total cholesterol, high-density
lipoprotein cholesterol (HDL), low-density lipoprotein cholesterol (LDL) (Horiba Medical,
Northampton, UK) and non-esterified fatty acids (NEFA; Randox, County Antrim, UK) using
a semi-automated analyzer (Pentra 400; Horiba Medical, Northampton, UK). Serum insulin
was determined using ELISA (EIA-2935; DRG instruments GmBH, Marburg, Germany).
Plasma glucose concentration and enrichment was determined using liquid chromatographytandem mass spectrometry (LC-MS/MS) as previously described 28.

243

244 Muscle analyses

245 Glycogen analysis

Glycogen content was determined as glycosyl units after acid hydrolysis and was measured
spectrophotometrically. The coefficient of variation for glycogen determination was 7-11%
(determined on multiple preparations from 3 separate biopsies).

249

250 Western blotting

To investigate skeletal muscle insulin signaling, commercially available antibodies were used
to determine the phosphorylation of key proteins (Akt Ser473 [Cell Signaling #4060], Akt Thr308
[Cell Signaling #13038], AS160 Ser588 [Cell Signaling #8730], and AS160 Thr642 [Cell
Signaling #8881] by SDS-PAGE and Western blotting as previously described 29

255

256 Quantitative immunofluorescence

Details of the specific quantification techniques can be found below, and all techniques have been described in detail previously, including antibody specificity experiments $_{30,31}$. All techniques used frozen muscle biopsy samples cryosectioned to a thickness of 5 µm, mounted onto uncoated glass microscope slides so that transverse orientated samples could be used for analysis. Two sections from each condition (pre and post high-fat overfeeding; 0 and 30 min) within a participant were placed on the same slide and analysis was performed in duplicate (two slides).

264

265 Sections were fixed in acetone and ethanol (3:1). For assessment of eNOS Ser1177/eNOS ratio, sections were triple stained with antibodies against eNOS (Transduction Laboratories, 266 267 Lexington, KY, USA), eNOS Ser1177 (Cell Signaling Technology, Beverly, MA, USA) and anti-a smooth muscle actin (aSMA; Abcam, Cambridge, UK). For assessment of NOX2 and 268 269 p47phox content, sections were double stained with either NOX2 or p47phox (kind gift from 270 Prof Mark Quinn, Montana State University) and anti-αSMA. All sections were then incubated 271 with appropriate secondary antibodies (Invitrogen, Paisley, UK) in combination with the 272 endothelial marker Ulex Europaeus-FITC conjugated (UEA-I-FITC; Sigma-Aldrich, UK).

273

Images were acquired using an inverted confocal microscope (Zeiss LSM-710, Carl Zeiss,
Germany) with a 40x NA oil immersion objective. Alexa Fluor 405 was excited using the 405

nm line of the diode laser and detected with 371-422 nm emission. FITC fluorescence was
excited with a 488 nm line of the argon laser and detected with 493–559 nm emission. Alexa
Fluor 546 and 633 fluorophores were excited with 543 nm and 633 nm lines of the helium–
neon laser and 548–623 nm and 638–747 nm emission, respectively. Identical settings were
used for all image capture within each participant.

281

282 All image analysis was performed using ImagePro Plus 5.1 (Media Cybernetics Inc, Bethesda, 283 MD, USA). Blood vessels were divided into either capillaries or arterioles using the α SMA 284 image. The endothelial (UEA-I-FITC) outline was then overlaid onto the corresponding 285 vascular enzyme image. Mean fluorescence intensity of the vascular enzyme signal was then quantified within the endothelial specific area. Diameter of the arterioles was determined on 286 287 calibrated images. Vessels larger than 20 µm in diameter were excluded to remove 3rd and 4th 288 order arterioles 32 from the analysis. As eNOS and eNOS Ser1177 phosphorylation had been 289 stained on the same sections it was possible to establish eNOS Ser1177/eNOS ratio on an 290 individual vessel basis, as the same endothelial outline could be placed over both eNOS and 291 eNOS Ser1177 images. The researcher was blinded to condition during imaging and analysis, 292 and all analysis was conducted by the same researcher. 6 ± 1 arterioles and 139 ± 11 capillaries were assessed per participant for eNOS content and eNOS Ser1177 phosphorylation. 8 ± 1 293 294 arterioles and 148 ± 11 capillaries were assessed per participant for NOX2 content. 8 ± 1 295 arterioles and 154 ± 14 capillaries were assessed per participant for p47phox content.

296

297 Calculations

Tracer calculations were performed using a non-steady state, single-pool model as previously described 33. Peripheral insulin sensitivity was calculated as the mean glucose clearance rate during the 2 h postprandial period divided by the mean serum insulin concentration over the 301 same period 34,35. Homeostatic model assessment of insulin resistance (HOMA-IR) was
302 calculated as described by 36. Area under the curve (AUC) was calculated using the trapezoidal
303 rule with zero as the baseline.

305 Statistics

All data are presented as means \pm standard error of the mean (SEM). A Shapiro-Wilk test was performed to test for normal distribution. Statistical analysis was performed using SPSS v23 for windows. Paired t-tests were used to make pre to post high-fat overfeeding comparisons where appropriate. All remaining data were compared using a two-way (trial x time) RM ANOVA, followed with Bonferroni-adjusted post-hoc t-tests where appropriate. Statistical significance was set at p < 0.05.

326 **Results**

327 Body mass and fasting blood parameters

The effect of high-fat overfeeding on body mass and fasting blood parameters is shown in Table 1. High-fat overfeeding increased body mass, plasma glucose, serum insulin, HOMA-IR, total cholesterol and HDL cholesterol. LDL cholesterol was unaffected by the diet, whereas TAG and NEFA decreased.

332

333 Postprandial plasma glucose and serum insulin

334 Plasma glucose and serum insulin increased in response to carbohydrate plus protein ingestion, 335 peaking at 30-45 min (Figure 1A and 1B, respectively). There was a main effect of trial (P =0.004) and a trial x time interaction (P = 0.012) for plasma glucose (Figure 1A), with plasma 336 337 glucose at 30- and 45-min time points being higher after high-fat overfeeding than before. 338 Postprandial plasma glucose AUC (0 - 120 min) increased by 11% after high-fat overfeeding 339 $(596 \pm 23 \text{ mmol/L per } 120 \text{ min before } vs. 663 \pm 19 \text{ mmol/L per } 120 \text{ min after; } P = 0.004)$. A 340 main effect of trial (P = 0.034) and a trial x time interaction (P = 0.009) were also evident for 341 serum insulin (Figure 1B), with serum insulin at the 45-min time point being higher after high-342 fat overfeeding than before. Postprandial serum insulin AUC increased by 19% after high-fat overfeeding $(34,164 \pm 4,525 \text{ pmol/L per } 120 \text{ min before } vs. 40,715 \pm 3,143 \text{ pmol/L per } 120$ 343 344 min after; P = 0.035).

345

346 EGP, oral glucose appearance, and whole-body glucose clearance

There was a trial x time interaction for EGP (P = 0.009; Figure 2A). High-fat overfeeding reduced EGP during fasting and early postprandial measurements, and carbohydrate plus protein ingestion suppressed EGP to a similar extent before and after high-fat overfeeding. Thus, alterations in EGP did not contribute to elevated glucose concentrations. Oral glucose 351 appearance was unaffected by high-fat overfeeding, although there was a tendency for a trial x 352 time interaction (P = 0.062; Figure 2B). Lastly, there was a main effect of trial (P = 0.025) for 353 whole body glucose clearance, with high-fat overfeeding resulting in reduced glucose clearance 354 rate (Figure 2C). When calculated for the entire 2 h oral glucose challenge, high-fat overfeeding 355 reduced time-averaged whole-body glucose clearance rate by 16% (3.2 ± 0.2 mL/kg/min post 356 vs. 3.8 ± 0.2 mL/kg/min pre; P = 0.021). Thus, elevated glucose concentrations observed after 357 high-fat overfeeding were due to reduced glucose disposal and not increased liver glucose 358 output.

359

360 Peripheral insulin sensitivity

361 Peripheral insulin sensitivity decreased by 26% after high-fat overfeeding (10.4 \pm 1.7 362 mL/kg/min/nmol/L post *vs.* 14.2 \pm 1.6 mL/kg/min/nmol/L pre; *P* = 0.006).

363

364 Skeletal muscle insulin signaling

365 The phosphorylation of key intermediates of the insulin signaling cascade is shown in Figure 3. Phosphorylation of Akt Ser473, Akt Thr308, AS160 Ser588 and AS160 Thr642 increased from 366 367 0 to 30 min after carbohydrate plus protein ingestion (P < 0.05). This response was not affected by high-fat overfeeding. Phosphorylation of Akt Ser473, Akt Thr308 and AS160 Thr642 decreased 368 369 from 30 to 120 min. However, phosphorylation of AS160 Thr642 was higher at 120 min than at 370 0 min. There was a trial x time interaction for AS160 Serss (P = 0.042). Before high-fat 371 overfeeding, phosphorylation of AS160 Ser588 remained elevated 120 min after carbohydrate 372 plus protein ingestion. After high-fat overfeeding, phosphorylation of AS160 Ser588 returned 373 to baseline at 120 min.

374

375 Muscle glycogen

High-fat overfeeding had no effect on muscle glycogen content. Fasting muscle glycogen content was 430 ± 37 mmol/kg dm before high-fat overfeeding and 398 ± 28 mmol/kg dm after. Carbohydrate plus protein ingestion did not affect muscle glycogen content either, such that values at 120 min were comparable to those seen at 0 min.

380

381 Total and phosphorylated eNOS

382 eNOS content of terminal arterioles and capillaries is shown in Figure 4. High-fat overfeeding reduced eNOS content within terminal arterioles by 6% (P = 0.017), whereas high-fat 383 384 overfeeding did not affect eNOS content within the capillaries (p = 0.197). High-fat 385 overfeeding also altered eNOS phosphorylation (Figure 5). Before high-fat overfeeding, carbohydrate plus protein ingestion increased eNOS Ser1177 phosphorylation within terminal 386 387 arterioles by 11%. This effect was no longer present after high-fat overfeeding (Figure 5C; trial 388 x time interaction, P = 0.007). A near identical response was observed when eNOS Ser1177 389 phosphorylation was normalized to eNOS content, with nutrient intake resulting in an 8% 390 increase in eNOS Ser1177/eNOS before high-fat overfeeding but no increase after (Figure 5D; 391 trial x time interaction, P = 0.039). A similar response was observed within the capillaries, with 392 a nutrient-stimulated 7% increase in eNOS Ser1177/eNOS before high-fat overfeeding but no increase after (Figure 5D; trial x time interaction, P = 0.013). 393

394

395 NAD(P)H oxidase

The protein content of the NAD(P)H oxidase subunits NOX2 (enzymatic subunit) and p47phox (main regulator subunit) was determined within terminal arterioles and capillaries. High-fat overfeeding had no effect on the content of either subunit of the NAD(P)H oxidase complex (Figure 6).

401 **Discussion**

402 The tissue-specific changes in glucose metabolism that underpin dietary fat-induced 403 impairments in glycemic control are not fully understood. The main observation of this study 404 was that 7 days of high-fat overfeeding led to an increase in postprandial glucose concentration 405 that was attributable to a reduction in whole body glucose clearance, not elevated EGP. This 406 suggests that peripheral tissue (such as skeletal muscle) is the primary site of early lipid-407 induced impairments in glucose metabolism. Despite this, we observed little to no change in 408 skeletal muscle insulin signaling, suggesting that mechanisms other than impaired insulin 409 signaling are responsible for the reduction in glucose clearance. Notably, high-fat overfeeding 410 abolished insulin-mediated eNOS Ser1177 phosphorylation in skeletal muscle terminal 411 arterioles, suggesting that reduced NO production leading to reduced perfusion of skeletal 412 muscle in response to insulin may be involved in mediating impaired glucose clearance in 413 response to 7 days of high-fat overfeeding.

414

415 High-fat overfeeding has frequently been applied in animal studies aiming to better understand 416 the mechanisms leading to obesity and insulin resistance. These studies are consistent in 417 suggesting that hepatic insulin resistance precedes that of skeletal muscle insulin resistance 37-418 40. Human research has produced equivocal findings, with both the liver 6 and skeletal muscle 419 3 suggested as the primary site of altered glucose metabolism. Brons et al. 6 reported a 26% 420 increase in basal EGP after high-fat overfeeding that resulted in a 0.46 mmol/L increase in 421 fasting glucose. In contrast, we observed a 7% decrease in basal EGP despite a significant 422 increase in fasting plasma glucose. The reason for this discrepancy is unclear, as both the 423 subject characteristics and the diet intervention were similar between the two studies. 424 Balancing EGP with exogenous glucose supply is an essential component of glycemic control, 425 and one that is impaired in type 2 diabetes 41. We found that the ability to suppress EGP in

426 response to carbohydrate plus protein ingestion was adequately maintained following high-fat 427 overfeeding. It is not possible to consider this observation in the context of the data from Brons 428 *et al.* 6 as their use of a hyperinsulinemic-euglycemic clamp led to the complete suppression of 429 EGP, regardless of diet. However, individuals with prediabetes also retain adequate 430 suppression of EGP despite postprandial hyperglycemia relative to individuals with normal 431 glucose tolerance 42. Thus, in the early stages of insulin resistance it seems that alterations in 432 hepatic glucose metabolism do not contribute to whole body impairments in glycemic control. 433 We also determined oral glucose appearance rate and found that it was not affected by 7 days 434 of high-fat overfeeding. The proportion of ingested glucose reaching the systemic circulation 435 after 2 h was 57% before high-fat overfeeding and 53% after, which is comparable to that 436 reported for healthy individuals 10.

437

438 High-fat overfeeding caused a 16% decrease in whole body glucose clearance, which is in close 439 agreement with the 20% decrease in leg glucose uptake recently reported after 3 days of 440 increased fat intake 3. This is also comparable to the physiology of prediabetes, where 441 postprandial hyperglycemia has been attributed to reduced glucose clearance, not increased 442 oral glucose appearance or increased EGP 42. Skeletal muscle is a major contributor to insulin-443 stimulated glucose disposal both under clamp conditions 43 and following glucose ingestion 7-444 10. We therefore determined the effect of high-fat overfeeding on components of the skeletal 445 muscle insulin signaling cascade. We focused our attention on Akt and the 160-kDa Akt 446 substrate (AS160; also known as TBC1D4); the latter has been identified as the most proximal 447 component of the insulin-signaling cascade linked to GLUT4 translocation 44-46 and an 448 important regulator of insulin-stimulated skeletal muscle glucose uptake 47. As insulin-449 stimulated AS160 activation is impaired in skeletal muscle of type 2 diabetics 48, this protein 450 could play a role in dietary lipid-induced impairments in muscle glucose uptake. In the present 451 study, carbohydrate plus protein intake led to a robust increase in the phosphorylation of Akt 452 Ser473 and Thr308 as well as AS160 Ser588 and Thr642. However, high-fat overfeeding had little 453 to no effect on basal or carbohydrate plus protein-stimulated phosphorylation of either protein. 454 Others have reported similar. For example, acute lipid-heparin infusion was found to decrease glucose disposal by 50%, without changes in Akt Ser473 phosphorylation 49, and high-fat 455 456 overfeeding reduced insulin-stimulated leg glucose uptake independent of changes in Akt 457 Thr308 or AS160 PAS phosphorylation 3. Collectively, these findings suggest that alterations in 458 Akt/AS160 activation do not play a role in early lipid-induced impairments in glucose disposal, 459 at least in response to high fat food intake for up to 7 days.

460

461 In the study by Lundsgaard et al. 3, reduced leg glucose uptake was attributed to increased 462 PDH-E1a Ser300 phosphorylation and a downregulation in oxidative glucose disposal. Insulin-463 induced GLUT4 translocation in combination with activation of PDH plays a key role in 464 determining insulin-stimulated glucose oxidation 50, which is an important route of glucose 465 disposal 51. It is well known that high-fat feeding inhibits PDH and reduces carbohydrate oxidation both at rest and during exercise 52,53. Thus, whilst we do not have data on PDH 466 activity or whole-body carbohydrate oxidation, it is likely that high-fat overfeeding inhibited 467 468 PDH and reduced oxidative glucose disposal, which could partially explain the reduction in 469 glucose clearance. Glucose taken up by skeletal muscle can also be diverted to glycogen 470 storage. In the present study, basal muscle glycogen was unaffected by high-fat overfeeding, 471 suggesting that the amount of carbohydrate provided was enough to maintain glycogen 472 synthesis rates in the high fat feeding period. Additionally, acute carbohydrate plus protein 473 ingestion did not stimulate glycogen synthesis either before or after high-fat overfeeding, 474 supporting the idea that ingested carbohydrate may have been partitioned towards oxidative 475 disposal.

477 High-fat overfeeding caused a small but significant 6% reduction in eNOS content in terminal 478 arterioles. Perhaps more importantly, high-fat overfeeding abolished the increase in eNOS 479 Ser1177 phosphorylation that was seen after carbohydrate plus protein ingestion. 480 Phosphorylation of eNOS at Ser1177 is essential to insulin-mediated-NO production by 481 endothelial cells 54. Given that increases in insulin-mediated skeletal muscle perfusion are NO 482 dependent 15, impaired eNOS Ser1177 phosphorylation may contribute to reduced glucose clearance following increased fat intake. Mechanistic support for the role of eNOS 483 484 phosphorylation in enhancing skeletal muscle perfusion and glucose uptake comes from 485 Kubota and colleagues 55. In a series of elegant experiments, these authors demonstrated that impairments in insulin-mediated eNOS phosphorylation led to reduced skeletal muscle 486 487 perfusion and impaired skeletal muscle glucose uptake. Moreover, restoration of insulin-488 mediated eNOS phosphorylation completely restored skeletal muscle perfusion and glucose 489 uptake in mice lacking endothelial IRS-2 and those fed a high fat diet 55. The possibility that 490 impairments in insulin's microvascular action may be an early event in the development of 491 lipid-induced insulin resistance is supported by recent animal work. Premilovac et al. 22 increased fat intake from 4.8% to 9.0% in Sprague Dawley rats and demonstrated that lipid-492 493 induced impairments in insulin-stimulated muscle glucose uptake originated solely from 494 impairments in insulin's microvascular actions. In that study 22, the insulin sensitivity of the 495 muscle fibers remained intact, which is in line with our observations in the present study. In 496 contrast, animals fed a 22% fat diet experienced insulin resistance in both the microvasculature 497 and the muscle fibers 22. Unlike the impairment in the insulin-induced activation of eNOS, 498 high-fat overfeeding did not change the protein content of endothelial specific NOX2 499 (enzymatic subunit) and of p47phox (main regulator subunit) of NAD(P)H oxidase in skeletal muscle terminal arterioles and capillaries, suggesting that 7 days of high-fat overfeeding does
not increase superoxide anion production by NAD(P)H oxidase.

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503 As discussed above, the finding that 7 days of high-fat overfeeding impaired insulin-mediated 504 eNOS phosphorylation in terminal arterioles could mean that a reduced ability to increase 505 muscle perfusion in response to carbohydrate plus protein ingestion was responsible for 506 reduced glucose clearance. However, we cannot be certain of this as we do not have a measure 507 of muscle perfusion. It is also difficult to demonstrate causality in human experiments, and thus 508 any observation of reduced muscle perfusion alongside impaired glucose clearance could be 509 an epiphenomenon. Thus, further work is required to confirm whether dietary lipid-induced 510 impairments in eNOS phosphorylation contribute to reduced glucose clearance. A further 511 limitation of the present study is the lack of a control group maintaining their habitual diet. 512 However, we had to consider the ethical correctness of including a separate control group that 513 would undergo numerous muscle biopsies. For this reason, we chose to compare our 514 experimental diet against our participant's habitual food intake, as we have done previously 4.

515

516 In conclusion, 7 days of high-fat overfeeding impaired whole-body glycemic control in healthy 517 non-obese individuals. This was due to reduced glucose clearance, not elevated EGP. The 518 reduction in glucose clearance occurred without an impairment in skeletal muscle insulin 519 signaling, suggesting that an alternative mechanism is responsible for this effect of the high-520 fat diet. This study is the first to show that the insulin-induced Ser1177 phosphorylation of eNOS, 521 which is known to lead to vasodilation of terminal arterioles after mixed meal ingestion, is 522 impaired after 7 days of high-fat overfeeding. An additional strength of this study is the use of 523 a carbohydrate-protein mixture to simultaneously create hyperglycemia and hyperinsulinemia,

524	which is more physiologically relevant than the clamp techniques that have been used in
525	previous studies.
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527	Data availability
528	The datasets generated during the current study are not publicly available but are available from
529	the corresponding author on reasonable request.
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741 **Table and figure legends**

Table 1. NEFA, non-esterified fatty acids; TAG, triacylglycerol; HDL, high-density lipoprotein cholesterol; LDL, low-density lipoprotein cholesterol; HOMA-IR, homeostatic model assessment of insulin resistance. Data presented are means \pm SEM (*n* = 15).

Figure 1. Plasma glucose (A) and serum insulin (B) before (pre) and after (post) 7 days of highfat overfeeding. Time points -30 - 0 min represent the final 30 min of the 2-h pre-infusion period. All subsequent time points are following the ingestion of carbohydrate plus protein (indicated by dotted line). Data presented are means \pm SEM (n = 15). *significantly different between trials at the annotated time point (P < 0.05).

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Figure 2. Endogenous glucose production (EGP) (A), oral glucose appearance (B), and wholebody glucose clearance rate (C) before (pre) and after (post) 7 days of high-fat overfeeding. Time points -30 - 0 min represent the final 30 min of the initial 2-h pre-infusion period. All subsequent time points are following ingestion of carbohydrate plus protein (indicated by dotted line). Data presented are means \pm SEM (n = 14).

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Figure 3. Phosphorylation of skeletal muscle Akt Ser473 (A), Akt Thr308 (B), AS160 Ser588 (C), and AS160 Thr642 (D) during fasting and following ingestion of carbohydrate plus protein, before (pre) and after (post) 7 days of high-fat overfeeding. Data presented are means \pm SEM (*n* = 13). AU, arbitrary units. †significantly higher than 0 min (*P* < 0.05). ‡significantly lower than 30 min (*P* < 0.05).

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Figure 4. eNOS content in terminal arterioles and capillaries before (pre) and after (post) 7 days
of high-fat overfeeding. A, representative confocal microscopy images of skeletal muscle

arterioles from pre- (a) and post-high-fat overfeeding (b). The skeletal muscle microvascular endothelium was revealed using Ulex europaeus-FITC conjugated lectin (UEA-I) (green). Skeletal muscle eNOS expression was revealed using Alexa Fluor 546 conjugated secondary antibody (red). Images not shown, arterioles and capillaries were differentiated using anti- α smooth muscle actin in combination with Alexa Fluor 405 conjugated secondary antibody. Bar represents 10 µm. B, mean fluorescence intensity of eNOS is summarized. Data presented as means ± SEM (n = 12). *Significantly lower than before high-fat overfeeding (P < 0.05).

773

774 Figure 5. eNOS phosphorylation in terminal arterioles and capillaries during fasting (0 min) 775 and 30 min after consuming carbohydrate plus protein, before (pre) and after (post) 7 days of high-fat overfeeding. A and B, representative confocal microscopy images of skeletal muscle 776 777 arterioles from pre- (A) and post-high-fat overfeeding (B), in the fasted (a) and stimulated (b) 778 state. The skeletal muscle microvascular endothelium was revealed using Ulex europaeus-779 FITC conjugated lectin (UEA-I) (green). Skeletal muscle eNOS Ser1177 phosphorylation was 780 revealed using Alexa Fluor 633 conjugated secondary antibody (red). Images not shown, 781 arterioles and capillaries were differentiated using anti- α smooth muscle actin in combination 782 with Alexa Fluor 405 conjugated secondary antibody. Bar represents 10 µm. C, mean 783 fluorescence intensity of eNOS Ser1177 is summarized. D, eNOS Ser1177 phosphorylation 784 normalized to eNOS content. Data presented as means \pm SEM (n = 12). \pm Significant increase 785 from 0 min (fasted) (P < 0.05).

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Figure 6. NOX2 and p47phox content in terminal arterioles and capillaries before (pre) and after (post) 7 days of high-fat overfeeding. A and B, representative confocal microscopy images of skeletal muscle from pre- (a) and post-high-fat overfeeding (b), illustrating NOX2 (A) and p47phox (B). The skeletal muscle microvascular endothelium was revealed using Ulex

- r91 europaeus-FITC conjugated lectin (UEA-I) (green). Skeletal muscle NOX2 and p47phox
- repression were revealed using an Alexa Fluor 546 conjugated secondary antibody (red).
- 793 Images not shown, arterioles and capillaries were differentiated using anti- α smooth muscle
- actin in combination with Alexa Fluor 405 conjugated secondary antibody. Bar represents 25
- μm. C, mean fluorescence intensity of NOX2 is summarized. D, mean fluorescence intensity
- of p47phox is summarized. Data presented as means \pm SEM (n = 12).