Acute dietary fat intake initiates alterations in energy metabolism and insulin resistance

Elisa Álvarez Hernández, … , Martin Hrabě de Angelis, Michael Roden


**BACKGROUND.** Dietary intake of saturated fat is a likely contributor to nonalcoholic fatty liver disease (NAFLD) and insulin resistance, but the mechanisms that initiate these abnormalities in humans remain unclear. We examined the effects of a single oral saturated fat load on insulin sensitivity, hepatic glucose metabolism, and lipid metabolism in humans. Similarly, initiating mechanisms were examined after an equivalent challenge in mice.

**METHODS.** Fourteen lean, healthy individuals randomly received either palm oil (PO) or vehicle (VCL). Hepatic metabolism was analyzed using in vivo $^{13}$C/$^{31}$P/$^1$H and ex vivo $^2$H magnetic resonance spectroscopy before and during hyperinsulinemic-euglycemic clamps with isotope dilution. Mice underwent identical clamp procedures and hepatic transcriptome analyses.

**RESULTS.** PO administration decreased whole-body, hepatic, and adipose tissue insulin sensitivity by 25%, 15%, and 34%, respectively. Hepatic triglyceride and ATP content rose by 35% and 16%, respectively. Hepatic gluconeogenesis increased by 70%, and net glycogenolysis declined by 20%. Mouse transcriptomics revealed that PO differentially regulates predicted upstream regulators and pathways, including LPS, members of the TLR and PPAR families, NF-$\kappa$B, and TNF-related weak inducer of apoptosis (TWEAK).

**CONCLUSION.** Saturated fat ingestion rapidly increases hepatic lipid storage, energy metabolism, and insulin […]

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Acute dietary fat intake initiates alterations in energy metabolism and insulin resistance

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CONCLUSION. Saturated fat ingestion rapidly increases hepatic lipid storage, energy metabolism, and insulin resistance. This is accompanied by regulation of hepatic gene expression and signaling that may contribute to development of NAFLD.

REGISTRATION. ClinicalTrials.gov NCT01736202.

paring subacute oral ingestion of fatty acids with different compositions found an increase in the glucose infusion rate only after polyunsaturated fatty acid ingestion (20).

To overcome possible limitations of the previous studies, such as the use of nonphysiological routes of lipid administration or mixed meals, which introduce protein and carbohydrates as confounders, we designed a translational study concept comprising a clinical trial involving healthy humans and a corresponding study involving nonobese nondiabetic mice. The randomized crossover clinical trial examined the effects of a single oral challenge with PO, which is mainly composed of saturated fatty acids (2), versus vehicle (VCL) ingestion on whole-body insulin sensitivity (WBIS) and hepatic insulin sensitivity. Moreover, the contributions of hepatic glucose fluxes, i.e., GNG, net glycogenolysis (GLYnet), and the futile exchange between glycogenogenic and glycogenolytic pathways (glycogen cycling) to EGP as well as the effects of these fluxes on hepatocellular lipids (HCL) and phosphorous-containing metabolites were analyzed using combined in vivo multinuclear $^{13}$C/$^{31}$P/$^{1}$H and stable isotope tracers to assess plasma glucose appearance rates and sources of EGP. In the mouse study, we examined the effects of a similar oral saturated fat challenge on insulin sensitivity and hepatic transcriptome changes.

Results

Studies in humans. A total of 14 young, lean, insulin-sensitive male volunteers (Figure 1 and Table 1) received either an oral dose of PO (~1.18 g/kg BW) or an identical volume of VCL on 2 occasions in random order, spaced by an 8-week interval.

PO results in increased circulating TG, glucagon, and incretins. After PO administration, TG in plasma rose by 59% (area under the time curve [AUC], \( P < 0.001 \)) and by 156% in chylomicrons (AUC, \( P = 0.009 \)) (Figure 2A). The AUC for plasma free fatty acids (FFA) (Figure 2B) and insulin concentrations (Figure 2C) was unchanged, while the AUC for plasma C-peptide was 28% higher after PO ingestion versus VCL (\( P < 0.005 \), Figure 2D). Of note, FFA were increased at 300, 420, and 480 minutes. Blood glucose levels were not different between PO- and VCL-treated groups (Figure 2E). Plasma glucagon rose by 41% (AUC, \( P < 0.0001 \)) only after PO ingestion (Figure 2F). Also, glucagon-like peptide 1 (GLP-1) and gastric inhibitory polypeptide (GIP) levels were markedly increased and remained elevated after PO ingestion (both \( P < 0.005 \)) (Supplemental Figure 2; supplemental material available online with this article; https://doi.org/10.1172/JCI89444DS1). Circulating levels of TNF-\( \alpha \), IL-6, fetuin-A, chemerin, omentin, and cortisol were not different between PO and VCL groups (\( P > 0.5 \) for all) (Supplemental Table 1).

PO induces insulin resistance at whole-body, liver, and adipose tissue levels. Insulin sensitivity was measured using hyperinsulinemic-euglycemic clamp tests in healthy humans. Steady state was reached (Supplemental Figure 1), and pertinent parameters were analyzed during this time. PO ingestion reduced WBIS by 25% compared with VCL treatment ($P = 0.0005$, Figure 3A). Furthermore, after PO, volunteers also showed a decrease of 22% ($P = 0.002$) in the rate of glucose disappearance (Rd), mostly due to a 33% ($P = 0.01$) reduction in glucose oxidation (GOX), while the rate of nonoxidative glucose disposal remained unchanged.
EGP increased by 10% \((P = 0.03)\) at 240 minutes and by 82% \((P = 0.008)\) at 480 minutes after PO compared with VCL treatment (Figure 3C). After PO, insulin-stimulated suppression of EGP was 15% lower than that detected after VCL \((P < 0.01, Figure\ 3D)\). Finally, insulin-stimulated suppression of FFA (Figure 3E) and TG (Figure 3F) was 30% \((P = 0.0001)\) and 80% \((P = 0.026)\) lower after PO than after VCL treatment.

**PO augments the contribution of GNG to EGP.** In order to further analyze the PO-induced increase in EGP, we measured the contributions of gluconeogenic and glycogenolytic fluxes. We found that GNG increased by approximately 70% \((P = 0.01)\), while GLYnet decreased by approximately 20% \((P < 0.05)\) after PO. Glycogen phosphorylase (GP) flux tended to be lower in the PO arm \((P = 0.085)\). The contribution of glycogen cycling to total GP flux was negligible in both study arms (Figure 4).

**PO increases lipid oxidation rates.** The respiratory quotient (RQ), defined as the rate of CO\(_2\) production/O\(_2\) consumption, was comparable in the period after PO or VCL ingestion but was reduced during the clamp only after PO (Figure 5A). Resting energy expenditure (REE) and lipid oxidation (LOX) rates increased markedly at 300 minutes after PO ingestion and remained elevated throughout the clamp (at 420 min) after PO (Figure 5, B and C). GOX decreased in the clamp following PO ingestion compared with GOX after VCL ingestion (Figure 5D).

**PO raises hepatic ATP and lipid content.** HCL and hepatic \(\gamma\)ATP increased by 33% \((P < 0.01)\) and 16% \((P = 0.009)\) from –120 minutes to 240 minutes after PO, respectively. At 240 minutes, \(\gamma\)ATP tended to be higher in the VCL arm \((P = 0.06)\), while HCL levels did not differ between PO and VCL arms. Hepatic inorganic phosphate (Pi) and \(\Delta\)Pi did not change after PO or VCL (Table 2).

**Studies in mice.** Two mouse cohorts received PO or VCL via gavage. One cohort then underwent hyperinsulinemic-euglycemic clamp tests in unrestrained mice. WBIS in mice trended toward a reduction \((P = 0.07)\) (Figure 7A). Whole-body glucose uptake, given by Rd, was not different between interventions (Figure 7B). Residual EGP during the clamp tended to be higher \((P = 0.06)\) (Figure 7C), and insulin-mediated suppression of EGP was 25% lower \((P = 0.04)\) after PO (Figure 7D), reflecting marked hepatic insulin resistance. PO ingestion did not affect insulin-induced FFA suppression \((P = 0.04)\) but reduced insulin-induced TG suppression by 83% \((P = 0.0039)\) (Figure 7F). Of note, glucose uptake in the gastrocnemius muscle (Supplemental Figure 3A) and white adipose tissue (Supplemental Figure 3B) as well as rates of glycolysis did not differ between the PO- and VCL-treated groups (Supplemental Figure 3C). Interestingly, PO administration did not change hepatic TG content in mice (Supplemental Figure 3D).

**PO causes changes in the expression of hepatic transcription factors.** Livers from both mouse cohorts were used for analysis on Affymetrix Gene 2.1 ST microarrays. A total of 273 and 327 differentially regulated probe sets were obtained from each cohort. Array data were deposited in the NCBI’s Gene Expression Omnibus (GEO) database (GEO GSE80259). Canonical pathways predicted to be differentially regulated by Ingenuity software after PO included TNF-like weak inducer of apoptosis (TWEAK) and aryl hydrocarbon receptor (AHR) under insulin-stimulated and noninsulin-stimulated conditions; phospholipase D and -myoinositol-5-phosphate signaling under insulin-stimulated conditions; and p38 MAPK, NF-kB, PPAR\(\alpha\), and OX40 under noninsulin-stimulated conditions (Figure 8, A and B, all \(P < 0.05)\). Several upstream regulators involved in hepatic fatty acid metabolism and inflammatory processes were predicted by Ingenuity software to be regulated by PO ingestion. These upstream regulators included LPS, which was activated with the most certainty, with a Z score of 1.9 and 2.3 under insulin- and noninsulin-stimulated conditions, respectively, TLR family members (TLR9 and TLR3 under insulin-stimulated conditions as well as TLR4 under noninsulin-stimulated conditions), PPAR\(\alpha\), and FOXO1 (all \(P < 0.01)\) (Figure 8, C and D).

### Table 1. Anthropometric and blood parameters of study participants

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Mean ± SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n) (men/women)</td>
<td>14 (14/0)</td>
</tr>
<tr>
<td>Age (yr)</td>
<td>25.8 ± 1.4</td>
</tr>
<tr>
<td>BMI (kg/m(^2))</td>
<td>22.5 ± 0.3</td>
</tr>
<tr>
<td>Waist circumference (cm)</td>
<td>79.5 ± 1.4</td>
</tr>
<tr>
<td>Lean BW (kg)</td>
<td>59.2 ± 2.1</td>
</tr>
<tr>
<td>Tg (mg/dl)</td>
<td>78.4 ± 9.8</td>
</tr>
<tr>
<td>FFA (µmol/l)</td>
<td>386.9 ± 32.9</td>
</tr>
<tr>
<td>ALT (U/l)</td>
<td>24.4 ± 3.2</td>
</tr>
<tr>
<td>AST (U/l)</td>
<td>26.0 ± 2.5</td>
</tr>
<tr>
<td>Fastiging blood glucose (mg/dl)</td>
<td>76.1 ± 2.0</td>
</tr>
<tr>
<td>2-hour postprandial blood glucose (mg/dl)</td>
<td>76.4 ± 4.7</td>
</tr>
<tr>
<td>HCL (% H(_2)O)</td>
<td>0.60 ± 0.09</td>
</tr>
</tbody>
</table>

### Table 2. Hepatocellular lipids and hepatic phosphorus-containing metabolites

- **Data represent the mean ± SEM.**
- \(n = 12.\) \(P < 0.01, \) \(P < 0.05, \) and \(P = 0.066,\) for –120 minutes compared with 240 minutes; \(P = 0.085,\) for the mean difference from –120 to 240 minutes between groups.

**Group**

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>–120</th>
<th>0</th>
<th>240</th>
</tr>
</thead>
<tbody>
<tr>
<td>VCL</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HCL (% H(_2)O)</td>
<td>0.97 ± 0.20</td>
<td>0.93 ± 0.23</td>
<td>1.04 ± 0.21</td>
</tr>
<tr>
<td>γATP (mmol/l)</td>
<td>2.86 ± 0.17</td>
<td>2.79 ± 0.14</td>
<td>3.17 ± 0.16 (^{1})</td>
</tr>
<tr>
<td>P (mmol/l)</td>
<td>2.35 ± 0.04</td>
<td>2.61 ± 0.07</td>
<td>2.61 ± 0.05</td>
</tr>
<tr>
<td>γATP/P</td>
<td>1.21 ± 0.07</td>
<td>1.07 ± 0.04</td>
<td>1.26 ± 0.10</td>
</tr>
</tbody>
</table>

**PO**

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>–120</th>
<th>0</th>
<th>240</th>
</tr>
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</tr>
</tbody>
</table>
PO induced a marked increase in plasma FFA concentrations in both humans and mice, but no alterations in circulating inflammatory markers or adipokines, such as TNF-α, IL-6, fetuin A, chemerin, and omentin. This finding indicates that the acute effects of PO are mediated by metabolic rather than endocrine changes and is partly in line with findings obtained after intravenous infusion of soybean oil, which showed no changes in circulating cytokines (19), or after ingestion of cream, which resulted in increased expression of TNF-α but not IL-6 (28). Interspecies differences could also be due to the administration of emulsified versus pure PO.

Notably, a single dose of PO markedly altered hepatic glucose fluxes and resulted in increased rates of GNG, reduced GLYnet, and a corresponding trend toward reduced GP flux. This finding extends our previous results on the effects of parenteral administration of polyunsaturated lipids (29). The present study also provides a comprehensive analysis of in vivo hepatic glucose and glycogen fluxes in humans that includes an assessment of glycogen cycling, which ensured correct estimation of GLY and GNG contributions to EGP (30).

In chronic insulin-resistant states, such as occurs in T2DM or type 1 diabetes mellitus (T1DM), elevation of GNG and EGP coexists with enhanced glycogen cycling (30). Surprisingly, oral lipid loading stimulated GNG and hepatic insulin resistance without affecting glycogen cycling, which remained negligible, as was reported for healthy humans in the fasted state (30, 31). This indicates that healthy humans can rapidly downregulate GLY under conditions of elevated GNG to avoid futile cycling. The absence of any effect on glycogen cycling might be due to the prevailing euglycemia and basal peripheral insulinemia compared with the hyperglycemic hypoinsulinemic conditions in the aforementioned study. Consequently, the augmented glycogen cycling observed in insulin-resistant states reflects chronically abnormal hepatic energy metabolism rather than an immediate physiological response to changes in dietary lipid supply. Of note, these changes occur in the presence of higher glucagon concentrations, which are likely due to FFA-induced glucagon stimulation (32). Specifically, the increase in GNG and the decrease in GLYnet upon PO administration could, at least in part, result from the increase in circulating glucagon. Even small changes in plasma glucagon can modify GLYnet independently of insulin in healthy humans (33).

Interestingly, ingestion of PO increased hepatic γATP and HCL concentrations, along with the induction of LOX and insulin resistance. Recently, we reported that a test involving ingestion of a mixed-meal with 35% fat content increases hepatic γATP exclusively in insulin-resistant obese humans but not in lean, nondiabetic or T2DM individuals (34). This suggests that the stimulatory effect of saturated fat on hepatic energy metabolism is dose dependent and may be linked to the onset of insulin resistance, but not the insulin sensitivity state per se. In line with this, recent studies report the

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**Discussion**

This study demonstrates that a single oral dose of saturated fat increases hepatic TG accumulation, insulin resistance, GNG, and ATP concentrations in the human liver. Ingestion of saturated fat also induces peripheral insulin resistance in skeletal muscle and adipose tissue. In mice, a single saturated fat load preferentially induces hepatic insulin resistance and also affects hepatic gene expression and signaling, which could contribute to the promotion of NAFLD.

Except for LPS, the analysis did not reveal whether the pathways and upstream regulators were activated or inhibited.

Under insulin- and noninsulin-stimulated conditions, PO resulted in upregulation of the following transcripts: miR1970, LPS-regulated genes (Ifit3, Clec4a3, Slc22a3, and C3ar1) (21–24), GO2, and Ar16, while Tweak gene expression was downregulated. Opposing regulatory transcripts, i.e., those that were upregulated under insulin-stimulated conditions and downregulated under noninsulin-stimulated conditions, were also found and included, for example, transcripts involved in cell growth (Map3k13 and Slc30a5) (25, 26) and FFA metabolism and development of NAFLD (fatty acid–binding protein 5 or Fabps) (27), as well as the predicted pseudogenes Gm3601I and H2-K2 (Figure 8E).

**Figure 2.** Time courses of circulating metabolites and hormones in humans. VCL (gray triangles) or PO (black circles) was administered at 0 minutes to lean, healthy men, and the hyperinsulinemic-euglycemic clamp was started at 360 minutes. TG circulating in plasma (solid line) and in chylomicrons (dashed line). The AUC was 59% and 156% higher after PO ingestion, respectively (A). Circulating FFA (B). Time courses for insulin (C), C-peptide (AUC 28% higher after PO) (D), blood glucose (E), and glucagon (AUC 41% higher after PO) (F). Values represent the mean ± SEM. n = 14; chylomicron TG n = 6. ***P < 0.001, **P < 0.005, and *P < 0.05, by paired, 2-tailed t test. Large asterisks refer to AUC differences; small asterisks refer to differences per time point.

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insulin resistance, while mice responded primarily with hepatic
impossible, given the current experimental design.
andance spectroscopy (MRS) monitoring, which would have been
may have been missed in the absence of continuous magnetic res-

gamma

fatty acid transport (42), which is in line with the
upregulation of fatty acid trapping by adipose tissue (37). In order to maintain
low levels of HCL, the liver can only make use of either
or of lipid export of apolipoproteins (39). Here, we show that a single dose of PO induces both mecha-
nisms, but nonetheless leads to increased HCL content.

upregulation of hepatic mitochondrial capacity in obese, insulin-
resistant, but not nondiabetic, humans in the absence of progressive
NAFLD (35) and that T2DM patients have decreased ATP turnover
associated with increased HCL (36). Collectively, these data suggest
that an acute rise in lipid availability and oxidation can upregulate
gamma ATP production and HCL deposition in young, healthy individu-
als with conserved mitochondrial plasticity, even after the develop-
ment of acute insulin resistance. The minor increase in HCL has to
be considered in the context of lean, insulin-sensitive individuals.
In this case, it is reasonable to assume that an increase in HCL after
just 1 dose of PO probably contributes to altered hepatic metabo-
lism. Furthermore, PO, like meal ingestion, probably induced indi-
nual time courses of increases, so that maximum gamma ATP increases
may have been missed in the absence of continuous magnetic reson-
ance spectroscopy (MRS) monitoring, which would have been
impossible, given the current experimental design.

Upon PO ingestion, human volunteers developed generalized
insulin resistance, while mice responded primarily with hepatic

upregulation of fatty acid transport of apolipoproteins (39). Here,
studies on insulin extraction used infusions of mainly polyunsat-
urated lipids, which revealed either unchanged or lower splanchnic
insulin extraction (17, 41). Likewise, oral intake of soybean oil
containing 61% polyunsaturated fatty acids resulted in increased
concentrations of both insulin and C-peptide (19).

The marked alterations in hepatic glucose fluxes in humans
and the predominant hepatic insulin resistance in mice raise inter-

interest in the effects of saturated fatty acids on hepatic gene expres-
sion, which has been previously examined mostly upon exposure
to polyunsaturated fatty acids (42). The present study showed that
a single dose of PO differentially regulated the canonical path-
ways TWEAK and AHR. The AHR pathway promotes NAFLD via
upregulation of fatty acid transport (42), which is in line with the
observed upregulation of Fadh5 and increase in HCL content in
humans. The TNF-related TWEAK is known to promote cell turn-
over homeostasis through the NF-kB and p38 MAPK pathways and
could serve as a biomarker of obesity and T2DM (43). TWEAK was
also found to be associated with reduced TG accumulation in pal-

Figure 3. Parameters of insulin resistance in human volunteers after VCL or PO during clamp experiments. VCL, white bars; PO, black bars. (A) WBIS, reflected by the M value. (B) Rd and its components GOX and nonoxidative glucose disposal (NOXGD). (C) EGP denoting hepatic insulin sensitivity at baseline (~180 min), after PO or VCL ingestion (240 min), and under insulin-stimulated conditions during the clamp (480 min). (D) Insulin-induced EGP suppression as an indicator of hepatic insulin sensitivity. (E) Insulin-induced FFA suppression reflecting adipose tissue insulin sensitivity and (F) the percentage of insulin-induced TG suppression. Data shown represent the mean ± SEM. n = 14. ****P < 0.0005, ***P < 0.01, and **P < 0.05; *P < 0.05, for GOX PO versus VCL. P values were determined by 2-tailed t test and ANOVA.
mitic acid–treated hepatocytes and may be involved in hepatic tissue repair (44). The present study cannot prove the activation status of TWEAK, thus making it difficult to interpret the observed downregulation of the Tweak gene and its receptor (Tnfrsf12a). However, recent findings of decreased circulating TWEAK concentrations correlating with obesity and concomitant NAFLD (45) could suggest that inhibition of the TWEAK pathway might increase susceptibility to hepatic injury.

The expression analyses further predicted PO-induced activation of LPS with or without insulin stimulation. This study also found upregulation of Ifi3, Clec4a3, Slc22a3, and C3ar1, genes regulated by LPS-stimulated macrophages. These data are in line with the reported increase in LPS concentrations upon high-fat ingestion (28, 46) resulting from lipid-induced disruption of the intestinal barrier (28). Interestingly, LPS is also known to decrease TWEAK signaling (47).

Downstream from LPS, NF-κB was found to be differentially regulated by PO. Generally known for its proinflammatory properties, NF-κB is also an important antiapoptotic factor (48, 49). LPS activation and TNF collectively lead to increased TLR4 expression, proinflammatory cytokine production, and inflammation, on the one hand, and NF-κB activation and cytoprotection on the other (48). NF-κB activation leads to modest and short-lived JNK activation, in turn inducing antiapoptotic genes, such as c-FLIP (a caspase 8 inhibitor) and X-linked inhibitor of apoptosis (48). As a result, the active NF-κB pathway is critical for LPS-induced resistance to hepatotoxicity (50). Additionally, a high-fat diet and obesity are associated with prolonged JNK activation and TNF-induced cell death (51, 52). This alludes to an adaptation, which is lost upon repeated and/or sustained exposure to hepatotoxic stimuli leading to NAFLD and steatohepatitis. Furthermore, the constitutive activation of NF-κB has been associated with severe hepatic and moderate peripheral insulin resistance (53). The present data suggest that a single PO challenge promotes pathways of LPS- and TLR4-mediated inflammation and cytotoxicity, which are buffered by the activation of NF-κB, which in turn contributes to insulin resistance. This study also found altered regulation of other putative cytoprotective mechanisms including the phospholipase C4 pathway, which is important for hepatic regeneration (54), and PPARα, which serves as both a canonical pathway and an upstream regulator protecting against NAFLD progression (29).

Our analyses of the differential regulation of single genes by saturated fat revealed several genes of interest. The observed upregulation of G0s2 may contribute to decreased TG clearance, thereby promoting NAFLD (55). The observed greater expression of Arl6 in the present study may also serve to protect against NAFLD,
because loss of function of this gene has been implicated in obesity, NAFLD, and diabetes in Bardet-Biedl syndrome (56). Of note, the descriptive nature of the transcriptome analyses and expression profiling carried out in our study does not allow final conclusions to be drawn as to causality. That is to say, given the results of this study, it is impossible to single out 1 gene, transcription factor, or pathway that is activated by acute exposure to PO and that can be causally linked to either insulin resistance or steatosis.

This study offers the advantages of the translational approach in mice and humans, the use of near-physiologic administration of saturated fat, and the comprehensive phenotyping of in vivo hepatic glucose and energy metabolism. This study’s limitations include the relatively small number of humans and mice, which might have prevented the detection of discrete effects of an acute lipid challenge. Nevertheless, the results demonstrate marked metabolic and transcripcional changes associated with PO treatment, even in these small groups. Another caveat of this study is that the hepatic expression data obtained in mice are not necessarily transferable to humans, as we could not obtain liver specimens from our human participants for ethical reasons. The lack of proteomic analysis of the targets identified by Ingenuity software constitutes another limitation of this study. Each pathway and regulator necessitates a thorough study of their role in the pathogenesis of insulin resistance and NAFLD, a task that must be tackled in subsequent studies.

The practical implication of this work is that the PO challenge used in this study most likely resembles the effects of ingestion of a meal rich in saturated fat, e.g., an 8-slice pepperoni pizza, containing approximately 16.72 g of saturated fat/110 g (57) or a meal consisting of a 110-g cheeseburger and a large portion of French fries, containing 18–25 g and 7–14 g per 1,000 kcal of saturated fat, respectively (58). One such meal would probably be sufficient to induce transient insulin resistance and impair hepatic metabolism, which necessitates the activation of hepatic defense mechanisms. Other simultaneously ingested biomacromolecules would exacerbate this metabolic challenge (59). The amount and types of fatty acids and carbohydrates in one such meal are in contrast to the diet recommendations of the American Diabetes Association (ADA). In this diet, daily intake of saturated fatty acids should not exceed 10% of total caloric intake. Furthermore, intake of mono- and polyunsaturated fatty acids and carbohydrates from vegetables, fruits, whole grains, and legumes is recommended (60). We presume that lean, healthy individuals are able to compensate adequately for excessive intake of saturated fatty acids, however, sustained and repeated exposure to such nutrients will ultimately lead to chronic insulin resistance and NAFLD. Recent studies have shown hepatic energy metabolism alterations and induction of insulin resistance in obese and lean patients after ingestion of a simple hypercaloric mixed meal containing 24 g of fat or a drink containing 1 g/kg BW of a mixture of palmitate and soybean oil (34, 61, 62). These results suggest that even lower doses of fatty acids are capable of inducing alterations similar to those observed with ingestion of pure PO.

In conclusion, the initial effects of ingestion of saturated fat include (a) augmented hepatic energy metabolism and lipid storage; (b) impaired hepatic insulin sensitivity, along with increased GNG flux; and (c) altered hepatic expression of genes regulating inflammatory and protective pathways, which predispose to and protect against the development of NAFLD.

Methods

Studies in humans

Volunteers. Fourteen lean, young male volunteers were enrolled in this randomized, controlled crossover trial (Figure 1 and Figure 9). Participants were recruited from March 2012 through December 2013. The sample size calculation was based on a 2-sided, paired t test, assuming a mean difference of EGP of 0.1 and a standard deviation of 0.11, resulting in a sample size of 12 to reach a power of 80%. The random allocation sequence was generated using SAS software (SAS Institute) by our in-house statistician. The possible order of treatments was randomly permuted in 2 blocks, with 1 extra block being generated to account for dropouts. Allocation was not concealed. Participants were enrolled and assigned to their treatment order by the study phy-
Volunteers drank 1.6 g/kg body water of 2H2O (99.9%; Sigma-Aldrich), approximately 684 kcal at 5:30 pm. At 8:00 pm, 10:00 pm, and 12:00 am, participants drank either VCL or PO within 10 minutes. Patients with more than 70 kg BW drank 92 g, and those with less than 70 kg BW drank 80 g of PO (~1.18g/kg BW; Biopalms; Landkrone) (65). To facilitate ingestion, PO was heated to 60°C, mixed with 1.84 g or 1.6 g emulsifier (Glice, Texturas; Albert y Ferran Adria), 9 or 8 g sugar-free vanilla syrup (Torani), and 81.2 or 70.4 ml bottled still water, for a PO mix of 92 g and 80 g, respectively. Oil test drinks were stirred constantly and served hot. For VCL administration, PO was substituted with 173.2 ml or 150.4 ml bottled still water, respectively. At 360 minutes, a hyperinsulinemic-euglycemic clamp began (10-min insulin bolus: 40 U/hour; continuous insulin infusion: 40 μU/m²/min; Insuman Rapid; Sanofi). Blood glucose concentration was adjusted to 90 mg/dl by adapting the glucose infusion rate (GIR) using 20% glucose (B. Braun AG) enriched with 2% D-[6,6-2H2]glucose, as described previously (19). Urine was sampled from –120 to 0 minutes, from 270 to 375 minutes, and from 390 to 510 minutes for the quantification of GNG and GLY. Blood was sampled at –60 and 360 minutes for assessment of GNG.

**Indirect calorimetry.** Indirect calorimetry (IC) was performed in the canopy mode using Vmax Encore 29n (CareFusion), as described previously (19), during baseline (at ~170 min), intervention (at 200 min), and steady-state clamp conditions (at 450 min) for 20 minutes after a 10-minute adaptation period. RQ, REE, and substrate oxidation rates were calculated as reported previously (19). Nonoxidative glucose disposal was calculated from the difference between rates of glucose disappearance and carbohydrate oxidation.

**Metabolites and hormones.** Blood samples were immediately chilled, centrifuged, and the supernatants stored at either −20°C or −80°C until analysis. Venous blood glucose concentration was measured immediately using the glucose oxidase method (EKF Biosen C-Line glucose analyzer; EKF Diagnostics) (19). TG concentration was analyzed enzymatically on a Roche Cobas c 311 Analyzer (Roche Diagnostics). Serum chylomicron content was determined from the TG concentration in the first fraction of density-gradient ultracentrifugation (66). FFA were assayed enzymatically (Wako) using orlistat to prevent in vitro lipolysis (19).Serum C-peptide, insulin, and plasma glucagon levels were measured by radioimmunoassay (EMD Millipore). Cortisol levels in serum samples were measured by immunoassay using a Siemens Immulite 2000XFI Analyzer (67). GLP-1 and GIP were measured by ELISA (TECOmedical; EMD Millipore) (64). ELISA was used to measure plasma concentrations of IL-6, TNF-a, fetuin-A (all using Quantikine HS ELISA kits from R&D Systems), omentin, and chemerin (19) using ELISA kits from BioVendor). Intrav..
Gas chromatography–mass spectrometry. Determination of atom percentage enrichment (APE) of $^2$H in blood glucose was done after deproteinization and derivatization to the aldonitrile-pentaacetate. The analyses were performed on a Hewlett-Packard 6890 gas chromatograph equipped with a 25 m/0.25 mm/0.12 μm CPSil5CB capillary column (Chrompack; Varian) and interfaced with a Hewlett-Packard 5975 mass selective detector. Selected ion monitoring was used to determine enrichments of fragments C3 to C6 (Supplemental Figure 1, A and B). Average mass units were 187 for endogenous glucose and 189 for D-[$^6,^6$-2H$_2$]glucose as described previously (19).

MRS. All measurements were conducted with the volunteers lying in a supine position within a whole-body 3.0 T Achieva MRI machine (Philips Healthcare). Twelve volunteers were studied, including all participants for whom flux measurements were obtained. The effects of PO or VCL on HCL and hepatic ATP concentrations were assessed at baseline and 360 minutes after intervention. For hepatic 1H MRS, a Q-body coil was used for shimming and HCL acquisition. Clinical T2-weighted turbo spin-echo (TSE) images were obtained in the transverse and coronal planes for localization and repositioning of the voxels used for HCL and ATP measurements. Respiratory-triggered $^1$H spectra were acquired with a single-voxel (30 × 30 × 20 mm$^3$) stimulated echo acquisition mode (STEAM) sequence. The variables were as follow: repetition time (TR) 3,000 ms, echo time 10 ms, and signal averages 16. To assure and interassay coefficients of variation (CV) for all cytokines were 2% to 7.2% and 4% to 14.4%, respectively.

Glucose and glucuronide $^2$H enrichment measurements by ex vivo $^2$H MRS. The positional enrichment of urinary acetaminophen glucuronide and plasma glucose, resulting from ingestion of $^2$H$_2$O and acetaminophen at the level of glucose 6-phosphate (G6P), was assessed as previously described (30) to estimate the contributions of GNG and GLY to EGP. Plasma glucose was derivatized to monoacetone glucose (MAG), while urinary acetaminophen glucuronide was converted into 5-O-acetyl monoacetone glucuronic lactone (MAGLA) (68). When plasma glucose enrichment was inadequate, urinary glucuronide enrichment was analyzed instead, since both methods yield identical estimates of EGP contributions (68, 69). In total, 9 participants yielded sufficient data for NMR analysis, 5 of them for glucuronide.

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Calculations. Total AUC were calculated using the trapezoidal method. FFA and TG suppression was calculated from 100 – (clamp FFA or TG concentrations × 100)/basal FFA or TG concentrations, respectively. Glycogen cycling, i.e., simultaneous fluxes through glycogen synthase (GS) and GP, was assessed. In order to measure glycogen cycling, isotopic tracer measurements of EGP, GNG, and GP fluxes must be supplemented by a measurement of GLYnet (in this case, 13C MRS) (71). GLYnet was calculated from the linear regression of hepatic glycogen content at –2, 0, 2, and 4 hours of PO or VCL ingestion using the least mean squares method. Rates of GLYnet were normalized to liver volume and BW and are expressed in μmol/kg/min (61).

Whole-body glucose disposal (M value) was calculated from glucose infusion rates during the clamp steady state. The rates of EGP (in μmol/kg/min) were calculated by dividing the tracer infusion rate of D-[6,6- 2H2]glucose times its enrichment to the hydrogens bound to carbon 6, by the mean percentage of enrichment of plasma D-[6,6-2H2]glucose and then subtracting the tracer infusion rate (19). To account for the incorporation of 2H from 2H2O by GNG during the overnight fast, background D-[6,6-2H2]glucose was determined before administration of 2H2O as well as at –185 minutes on the day of the study. Consequently, for determination of basal EGP and EGP at the end of the intervention, the background D-[6,6- 2H2] value before administration of 2H2O was used. GNG (in μmol/kg/min) was calculated from the difference between EGP and GLYnet. The fractional GP flux contribution to EGP, i.e., the fraction of EGP originating from GLY, was calculated as 1 – (H5/H2); where H5/H2 is the ratio of glucuronide position 5 to position 2 enrichment from 2H2O, keeping in mind that glucose derived from GLY is enriched with 2H in position 2, while glucose derived from GNG is enriched in positions 5 and 2. Absolute GP flux (in μmol/kg/min) was calculated from the equation: GP = EGP × (1 – H5/H2). Glycogen cycling was then calculated as: GP - GLYnet.

Figure 9. Human study design. Lean, healthy male adults randomly received either PO or VCL on 2 occasions. Hepatic metabolism was measured using in vivo 13C, 31P, 1H and ex vivo 1H MRS combined with 2H2O and acetaminophen ingestion before and during hyperinsulinemic-euglycemic clamps with D-[6,6-2H2]glucose–labeled 20% glucose infusion.
deoxyglucose-6-phosphate was separated from 2-[14C]deoxyglucose via ion-exchange columns (Poly-Prep AG1-X8; Bio-Rad) as previously described (72). Glucose uptake was calculated by multiplying the mean plasma glucose levels between 120 and 140 minutes of the clamp (mmol/ml) by 2-[14C]deoxyglucose tissue content (dpm/100 g tissue), divided by the 2-[14C]deoxyglucose plasma AUC in the same time frame. Radioisotopes were purchased from PerkinElmer and samples measured in an Ultima-Gold Scintillation Cocktail (Tri-Carb2910TR; PerkinElmer) (Figure 10). Whole-body glucose disposal (M value) was calculated from the tracer infusion rate, the specific activity of [3-3H]glucose, and BW.

**Biochemical analyses.** Blood glucose concentrations were assessed using a Contour hand-held glucometer (Bayer Vital). Plasma TG levels were determined by a colorimetric assay (Cayman Chemical), and plasma FFA levels were assessed with the FFA-HR(2)-Test (Wako). Hepatic TG levels were measured in whole-liver homogenates biochemically with the BioVision Assay.

**Experiments under insulin-stimulated conditions.** A permanent jugular vein catheter was placed under ketamine/xylazine anesthesia into a cohort of 19 mice with the aforementioned characteristics. Six to seven days later, the mice were fasted for 10 hours and then received 2 g/kg BM PO or VCL via gavage. Six hours later, unrestrained, conscious mice underwent hyperinsulinemic-euglycemic clamps. After 110 minutes of primed-continuous [3-3H]glucose infusion (1.85 kBq/min), a blood sample was collected to determine plasma insulin, glucose, and [3-3H]glucose concentrations for the calculation of basal EGP. A [3-3H]glucose infusion (3.7 kBq/min) containing insulin (15 pmol/kg/min; HumulinR; Lilly) was started. Blood glucose concentrations were measured every 10 minutes and target glycemia established by adjusting the GIR. At minute 120, 2-[14C]deoxyglucose (370 kBq) was injected intravenously to assess tissue-specific Rg rates. At the end of the experimental procedure, mice were euthanized by means of an intravenous ketamine/xylazine injection. Livers were collected, immediately snap-frozen in liquid nitrogen, and stored at −80°C. Blood was collected at culling, and plasma [3H] and [14C] radioactivity was determined in deproteinized plasma before and after [3H]2O evaporation to estimate glycolysis rates. In hepatic lysates, 2-[14C]deoxyglucose-6-phosphate was separated from 2-[14C]deoxyglucose via ion-exchange columns (Poly-Prep AG1-X8; Bio-Rad) as previously described (72). Glucose uptake was calculated by multiplying the mean plasma glucose levels between 120 and 140 minutes of the clamp (mmol/ml) by 2-[14C]deoxyglucose tissue content (dpm/100 g tissue), divided by the 2-[14C]deoxyglucose plasma AUC in the same time frame. Radioisotopes were purchased from PerkinElmer and samples measured in an Ultima-Gold Scintillation Cocktail (Tri-Carb2910TR; PerkinElmer) (Figure 10). Whole-body glucose disposal (M value) was calculated from the tracer infusion rate, the specific activity of [3-3H]glucose, and BW.

**Studies in mice**

**Animals.** Studies were conducted in lean, male, 14 week-old C57BL/6NTac mice (Taconic). Animals had ad libitum access to water and a standard chow diet. Mice were kept on a low-fat (LF) diet (13% of calories derived from fat, 17 kJ/g, Standard Diet 1310; Altromin), and were matched for body mass (BM) and littersmates. BM and composition (MiniSpec LF50; Bruker Optics) were measured 1 day prior to the start of the experiment. Animals were bred and housed in a temperature- and humidity-controlled environment including a 12-hour light/12-hour dark cycle, in compliance with the Federation of European Laboratory Animal Science Associations protocols.

**Experiments under noninsulin-stimulated conditions.** C57BL/6NTac mice (n = 20) with characteristics identical to those described above were fasted for 10 hours and given 2 g/kg BM PO (Landkrone) or water (VCL) per gavage. Lateral tail vein blood samples were obtained prior to treatment and 2 hours afterward. Six hours after treatment, mice were euthanized with isoflurane, and a vena cava blood sample was collected and centrifuged at 4°C, and plasma aliquots were immediately frozen in liquid nitrogen. Liver was dissected and immediately snap-frozen in liquid nitrogen (Figure 10).

**RNA isolation**

Snap-frozen liver samples from both cohorts were processed after administration of PO. Total RNA was isolated using the mRNAsely Mini Kit (QIAGEN). The Agilent 2100 Bioanalyzer was used to assess
RNA quality, and only high-quality RNA (RNA integrity number [RIN] >7) was used for microarray analysis.

Expression profiling
Total RNA (~30 ng) was amplified using the Ovation PicoSL WTA System V2 in combination with the Encore Biotin Module (both from NuGEN). Amplified cDNA was hybridized on Affymetrix Mouse Gene 2.1 ST array plates containing approximately 35,000 probe sets. Staining and scanning (GeneChip Scanner 3000 7G; Affymetrix) were performed according to the Affymetrix Gene Titan expression protocol and modified according to NuGEN’s Encore Biotin protocol.

Statistical transcriptome analysis
Expression Console software (version 1.3.0.187; Affymetrix) was used for quality control and to obtain annotated normalized robust multiarray average (RMA) gene-level data (standard settings included median polish and sketch-quantile normalization). Statistical analyses were performed using the statistical programming environment R implemented in CARMAweb (https://carmaweb.genome.tugraz.at/carma/). Genewise testing for differential expression was done using the limma t test, and a P value of less than 0.05 was set as the threshold to define sets of regulated genes. For a fold change greater than 1.3 times and a linear average expression greater than 4 were applied. Pathway analyses were generated using the Ingenuity Pathway Analysis (QIAGEN; www.qiagen.com/ingenuity), where the overlapping P value identifies transcriptional regulators that can explain observed gene expression changes. The activation Z score helps infer activation states of predicted transcriptional regulators, with values of 2 or more indicating activation and values of ~2 or less indicating inhibition.

General statistical analyses
Results are presented as the mean ± SEM and were compared using a 2-tailed Student t test or ANOVA adjusted for repeated measures, with Bonferroni’s testing as appropriate. Calculations were performed using GraphPad Prism, version 6.02 (GraphPad Software). A P value of less than 0.05 was considered statistically significant, unless otherwise indicated.

Study approval
All participants provided written informed consent before inclusion in the study, which was performed according to the Declaration of Helsinki of 2013 and approved by the ethics board of the Heinrich Heine University Düsseldorf. All animal experiments were approved by the Upper Bavarian district government (AZ 55.2.1.54-2532-4-11).

Author contributions
MR initiated the investigation, led the clinical experiments and wrote, reviewed, and edited the manuscript. EAH obtained and analyzed the data and wrote, edited, and reviewed the manuscript. SK obtained and analyzed the data, aided in designing the clinical study, and edited and reviewed the manuscript. AS obtained data and edited and reviewed the manuscript. PB designed the MRS study, obtained MRS data, and reviewed and edited the manuscript. EAH, SK, AS, and PB contributed equally to this project. YK interpreted MRS data. BN researched the clinical data and reviewed and edited the manuscript. CB and FC performed derivatization experiments and enrichment analysis for 1H-MRS. [GJ] performed the 1H-MRS analyses and reviewed and edited the manuscript. PN conducted laboratory analyses and reviewed and edited the manuscript. SN and JR supervised the mouse studies and edited and reviewed the manuscript. MI analyzed transcriptomics data and edited and reviewed the manuscript. All authors gave final approval of the version to be published.

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5. Erion DM, Shulman GI. Diacylglycerol-mediated


