Discovery of a Class of Endogenous Mammalian Lipids with Anti-Diabetic and Anti-inflammatory Effects

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SUMMARY

Increased adipose tissue lipogenesis is associated with enhanced insulin sensitivity. Mice overexpressing the Glut4 glucose transporter in adipocytes have elevated lipogenesis and increased glucose tolerance despite being obese with elevated circulating fatty acids. Lipidomic analysis of adipose tissue revealed the existence of branched fatty acid esters of hydroxy fatty acids (FAHFAs) that were elevated 16- to 18-fold in these mice. FAHFA isomers differ by the branched ester position on the hydroxy fatty acid (e.g., palmitic-acid-9-hydroxy-stearic-acid, 9-PAHSA). PAHSAs are synthesized in vivo and regulated by fasting and high-fat feeding. PAHSA levels correlate highly with insulin sensitivity and are reduced in adipose tissue and serum of insulin-resistant humans. PAHSA administration in mice lowers ambient glycemia and improves glucose tolerance while stimulating GLP-1 and insulin secretion. PAHSAs also reduce adipose tissue inflammation. In adipocytes, PAHSAs signal through GPR120 to enhance insulin-stimulated glucose uptake. Thus, FAHFAs are endogenous lipids with the potential to treat type 2 diabetes.

INTRODUCTION

Obesity and type 2 diabetes (T2D) are at epidemic proportions worldwide (Hu, 2011). The major pathogenic factors underlying T2D are resistance to insulin action in peripheral tissues and dysregulated insulin secretion. The Glut4 glucose transporter is the major insulin-regulated glucose transporter and mediates glucose uptake into skeletal muscle, heart, and adipocytes in response to rising insulin after a meal (Shepherd and Kahn, 1999). In humans and rodents with obesity or T2D, Glut4 is downregulated selectively in adipocyte tissue (AT) and not in muscle (Shepherd and Kahn, 1999). This alters AT biology leading to systemic insulin resistance (Abel et al., 2001). Glut4 knockout selectively in adipocytes in mice results in insulin resistance and increased T2D risk (Abel et al., 2001), whereas adipose-selective overexpression of Glut4 (AG4OX) lowers fasting glycemia and enhances glucose tolerance (Carvalho et al., 2005; Shepherd et al., 1993). These effects in AG4OX mice are mediated by glucose-dependent induction of lipogenesis in AT driven by ChREBP (Herman et al., 2012), a transcription factor that regulates both glycolysis and lipogenesis (Iizuka et al., 2004; Ma et al., 2005). ChREBP knockout in AG4OX mice completely reverses the enhanced glucose tolerance (Herman et al., 2012). Expression of ChREBP and lipogenic genes in AT is highly associated with insulin sensitivity in humans and rodents (Herman et al., 2012; Roberts et al., 2009) and increased de novo lipogenesis in AT has favorable metabolic effects including potentially increasing longevity (Bruss et al., 2010).

Elevated circulating fatty acids are generally associated with insulin resistance and glucose intolerance (Boden and Shulman, 2002). However, certain fatty acids such as dietary omega-3 fatty acids (Oh et al., 2010; Virtanen et al., 2014) and the endogenously produced palmitoleate (Cao et al., 2008) have favorable metabolic effects. Furthermore, large epidemiological studies show that an increased ratio of unsaturated to saturated fatty acids in serum triacylglycerols is associated with a reduced risk of T2D (Rhee et al., 2011; Risérus et al., 2009). Similarly, an increased ratio of monounsaturated to saturated fatty acids in the liver is associated with insulin sensitivity even with extensive hepatic steatosis (Benhamed et al., 2012). AG4OX mice have elevated circulating fatty acids and increased adiposity,
yet have lower fasting glycemia and profoundly enhanced glucose tolerance compared to controls (Carvalho et al., 2005; Herman et al., 2012; Shepherd et al., 1993). This raised the possibility that enhanced AT lipogenesis in response to Glut4 overexpression might drive the production of lipids which have favorable metabolic effects. Since Glut4 (Carvalho et al., 2005; Shepherd and Kahn, 1999) and ChREBP (Herman et al., 2012) expression are downregulated in AT in insulin-resistant humans and rodents, the production of these metabolically favorable lipids may be low in these states. To test these hypotheses, we performed lipidomic analysis of AT from wild-type (WT) and AG4OX mice. 

RESULTS

Identification of a Class of Glut4-Regulated Lipids

Using a quantitative mass spectrometry (MS) lipidomics platform (Saghatelian et al., 2004), we detected more than 1,400 ions in AT, 6% of which had a 2- to 4-fold difference between AG4OX and WT mice. A cluster of ions in AG4OX AT was elevated ≥16-fold (Figure 1A). The measured accurate mass of these ions enabled us to calculate their molecular formulas as C_{32}H_{61}O_{4} (509.4575), C_{34}H_{63}O_{4} (535.4732), C_{34}H_{65}O_{4} (537.4888), and C_{36}H_{67}O_{4} (563.5045). These formulas all contain a unique signature of four oxygen atoms indicating that these ions are members of a single lipid class. These formulas do not correspond to any known metabolite in the Metlin (Smith et al., 2005) and Lipid Maps (Sud et al., 2007) metabolite databases. We hypothesized these lipids might contribute to glucose-insulin homeostasis because of their abundance in AG4OX mice, in which improved glucose tolerance depends on enhanced AT lipogenesis (Herman et al., 2012). Therefore, we proceeded to determine the molecular structures and biologic effects of these lipids.

The mass differences among these ions suggested they contain fatty acids. Fragmentation of the 537 ion generated several product ions with masses of 255, 281, and 299 (Figure 1B), which correspond to palmitic acid (PA), octadecenoic acid, and hydroxy-stearic acid (HSA), respectively. The molecular formula of the 537 ion (C_{34}H_{65}O_{4}) does not contain any double bonds. This indicates that octadecenoic acid, which contains a double bond, results from fragmentation in the MS and is not part of the natural metabolite. Based on the chemical formula and the fact that this metabolite ionized only in the negative mode, the most reasonable structure for the 537 ion is an ester that combines PA and HSA to yield palmitic acid-hydroxy stearic acid (PAHSA) (Figures 1B and 1C). Based on this structural model and the masses detected for the other elevated ions, their structures are: palmitic acid-hydroxy palmitic acid (PAHPA, m/z 509), oleic acid-hydroxy stearic acid (OAHPA, m/z 563), and the 535 ion is a mixture of palmitoleic acid-hydroxy stearic acid (POHSA), and oleic acid-hydroxy palmitic acid (OAHPA) (Figure 1C). We refer to this class of natural-occurring lipids as fatty acid-hydroxy fatty acids (Figures 1C and 1D), abbreviated as FAHFAs. An additional ion, detected in positive ionization mode, was also upregulated in AG4OX AT (Figure 1A) but the molecular formula indicated it is not a FAHFA, and therefore we did not characterize it further.

Using a targeted MS approach, we identified 16 FAHFA family members in mouse serum that consisted of four fatty acids and four hydroxy-fatty acids in different combinations (Figure 1D). FAHFAs with PO, PA, or OA as the fatty acid moiety and HPA or HSA as the hydroxy-fatty acid moiety were most highly increased in AG4OX compared to WT mice (Figure 1D). Because PAHSAs were the most highly upregulated family member in AT of AG4OX (Figure 1A), we investigated their biologic effects.

Tissue Distribution of Total PAHSAs in WT and AG4OX Mice and Regulation by ChREBP

Targeted MS revealed PAHSAs in all tissues analyzed. In WT mice, total PAHSA levels are highest in brown adipose tissue (BAT) followed by subcutaneous (SQ) white adipose tissue (WAT), perigonadal (PG) WAT, and liver (Figure 1E). Total PAHSA levels are very low in heart and gastrocnemius muscle (data not shown). PAHSA levels vary >7-fold among tissues in WT mice (Figure 1E). In WT serum, total PAHSA levels are ~7 nM (Figure 1E). In AG4OX mice, total PAHSA levels are 16- to 18-fold elevated in SQ and PG WAT, 3-fold in BAT and ~2-fold in serum compared to WT mice (Figure 1E). In contrast, PAHSA levels in liver of AG4OX mice are ~30% lower than WT. Thus, Glut4 overexpression in AT results in broad systemic regulation of PAHSAs with tissue-specific alterations.

Because ChREBP regulates AT lipogenesis in AG4OX, we tested whether ChREBP regulates PAHSA levels in vivo. ChREBP knockout in normal mice reduces total PAHSA levels ~75% in PG- and SQ-WAT with no change in serum (Figure 1F). Knocking out ChREBP in AG4OX completely reverses the marked elevation in PAHSA levels in PG- and SQ-WAT and serum.

Tissue Distribution of Specific PAHSA Isomers and Regulation in WT and AG4OX Mice

We observed multiple peaks in the chromatograms that correspond to different PAHSA isomers with the ester connected to a different carbon of the hydroxy-fatty acid resulting in a branched lipid. Fragmentation of PAHSAs from AT using high collisional energy tandem MS (Moe et al., 2004) produced two ions at 127 and 155 (Figure 1B) indicating that the ester is at the 9th carbon of the HSA (Figure 1C). We refer to this isomer as 9-PAHSA, which was confirmed by chemical synthesis and coelution with 13C-9-PAHSA (Figure 2A). We also discovered PAHSAs with branched esters at carbons 5, 7, 8, 10, 11, 12, and 13 verified by comparison to synthetic standards (Figure 2A). Thus, there are at least eight PAHSA isomers. We achieved complete separation of all isomers except 13- and 12-PAHSA (Figure 2A), which we quantify together in all data sets.

We sought to determine which PAHSA isomers are upregulated in WAT and serum of AG4OX mice as an initial clue to which ones may have biologic activities that could affect glucose homeostasis. In WT serum, 13/12-, 11-, 10-, 9-, and 5-PAHSA are present at 0.4–2.5 nM, which is the range for signaling lipids such as prostacyclins, prostaglandins, steroids, and endocannabinoids. In WT WAT and BAT, 9-PAHSA is the most abundant isomer (Figure 2B). 13/12-, 11-, and 10-PAHSA are present at 20%–30% of 9-PAHSA levels and 8-, 7-, and 5-PAHSA are present at substantially lower concentrations (Figure 2B). Surprisingly, liver which is also a lipogenic tissue, has only...
Figure 1. Discovery and Characterization of a Class of Lipids (FAHFA)

(A) Comparative lipidomics of SQ white adipose tissue (WAT) from AG4OX and WT mice reveals the presence of a group of ions at m/z 509 (PAHPA), 535 (POHSA/OAHPA), 563 (OAHSA), and 537 (PAHSA) that are elevated 16- to 18-fold in AG4OX mice.

(B) Mass spectra of serum lipids from WT and AG4OX mice. The ions at m/z 537.4888, 509.4575, 563.5045, and 535.4732 are enriched in AG4OX mice.

(C) Structural formulas of PAHSA, PAHPA, OAHSA, and POHSA (not shown).

(D) Fatty Acid and Hydroxy Fatty Acid comparisons.

(E) Total PAHSA levels in SQ, PG, BAT, and serum from WT and AG4OX mice.

(F) Total PAHSA levels in SQ WAT, PG WAT, and serum from WT, AG4OX, ChREBP KO, and ChREBP KO x AG4OX mice.

(legend continued on next page)
13/12- and 9-PAHSA isomers in BAT or liver while fasting while 7- and 5-PAHSA are unchanged (Figure 2E). However, PAHSA are reduced in AG4OX liver indicating tissue-specific mechanisms for regulating uptake, synthesis, degradation or release.

This is further indicated by the tissue distribution of specific PAHSA isomers in WT mice. 13/12- and 9-PAHSA are present in all WT tissues examined (Figure 2C). 9-PAHSA is more abundant in AT than liver while 13/12-PAHSA is not. In contrast to 13/12- and 9-PAHSA, 5-PAHSA is restricted to AT, kidney, and serum (Figure 2C).

Physiologic Regulation of PAHSA with Fasting

We examined PAHSA regulation with fasting (Figure 2D). In the fed state, total PAHSA levels are highest in BAT; slightly lower in SQ and PG WAT; and substantially lower in liver, pancreas, and kidney (Figure 2D). Fasting increases PAHSA 2- to 3-fold in WAT and kidney and 65% in pancreas but does not alter the levels in BAT, liver, or serum (Figure 2D). Hence, PAHSA undergo tissue-specific regulation with fasting (Figure 2D). The fasting-induced increase in PAHSA in WAT is surprising since one would expect synthesis to be lower due to reduced lipogenesis and ChREBP with fasting. Indeed, in spite of elevated PAHSA levels, biosynthetic activity (described below) was not increased in WAT from fasted mice (data not shown). This may reflect inhibition of degradation or release. To better understand the mechanism, we determined fasting effects on PAHSA levels in AG4OX mice. Fasting further elevated PAHSA in WAT but not in BAT or serum (Figure S1 available online). Since PAHSA levels in AG4OX WAT are regulated by ChREBP-driven lipogenesis (Figure 1E) and lipogenesis is not increased with fasting, these data demonstrate an additional level of regulation and support the possibility that fasting inhibits PAHSA degradation or release.

We also investigated regulation of individual PAHSA isomers with fasting (Figure 2E). Although total PAHSA levels are unchanged in serum of fasted mice (Figure 2D), specific isomers (10-, 9- and 5-PAHSA) are modestly decreased (Figure 2E). In SQ and PG WAT, most of the isomers (13/12-, 11-, 10-, 9-, and 8-PAHSA) including the more abundant ones are increased with fasting while 7- and 5-PAHSA are unchanged (Figure 2E). Fasting had no effect on any PAHSA isomer in BAT or liver while all isomers were upregulated in kidney. In pancreas, 11- and 9-PAHSA are increased with fasting while 13/12- and 7-PAHSA are unchanged. Thus, PAHSA isomer levels undergo tissue-specific and isomer-specific regulation with fasting (Figure 2E). The abundance of different PAHSA isomers in the fasted state differs by 60-fold in a given tissue (compare 9- with 5-PAHSA in SQ WAT) (Figure 2E). These results suggest that fasting regulates pathways involved in synthesis, degradation, and/or release of specific PAHSA isomers in a tissue- and isomer-specific manner.

Regulation of PAHSA in Obesity and Insulin Resistance

We investigated PAHSA levels in insulin-resistant mice with high-fat-diet (HFD)-induced obesity (Figure 3A). After 9 weeks of HFD, mice were obese and diabetic (determined by GTT) (Figure S2A). HFD had differential effects on specific PAHSA isomers. 5- and 13/12-PAHSA were downregulated in HFD mice in serum, PG and SQ WAT and BAT (Figure 3A) although the difference did not reach significance for 13/12-PAHSA in PG WAT. Strikingly, 10-, 9-, 8-, and 7-PAHSA were increased in PG WAT of HFD-fed mice. Most of these isomers were decreased in SQ WAT and BAT and unchanged in serum (Figure 3A). Total lipid ion signal measured in SQ WAT was unchanged between chow- and HFD-fed mice (Figure S2B). 13/12- and 9-PAHSA were also decreased in liver (Figure 3A). These studies demonstrate: (1) 5-PAHSA and 13/12-PAHSA are consistently reduced in AT depots with HFD while other PAHSA isomers have opposite regulation among the depots (PG WAT versus SQ WAT and BAT) (Figure 3A); and (2) Only two of the five isomers in serum are reduced with HFD (Figure 3A). Thus, PAHSA undergo isomer-specific and tissue-specific regulation under insulin-resistant conditions in WT mice.

PAHSA Are Present in Food

To determine whether the changes in PAHSA levels in altered metabolic states could result from differences in dietary intake, we measured PAHSA levels in rodent and human foods. In chow and HFD, we found five of the seven isomers that are present in mouse AT, 13/12-, 11-, 10-, 9-, and 8-PAHSA, but not 7- and 5-PAHSA. However, the relative abundance among isomers was strikingly different from AT or serum with 10-PAHSA being most abundant in both diets (Figure 3B). Levels of all these isomers were substantially lower in HFD than chow (Figure 3B). Given that PAHSA increase in WAT during fasting (Figures 2D and 2E), regulation of tissue PAHSA levels does not simply reflect dietary intake. Similarly, the abundance of PAHSA isomers in serum and tissues (Figure 2C) does not correlate with predominant isomers.

(B) Structural analysis of the 537 ion from AG4OX WAT by tandem MS demonstrates that it is composed of palmitic acid (m/z 255) and hydroxy stearic acid (m/z 281). Octadecanoic acid (m/z 281) results from the dehydration of hydroxy stearic acid. Fragmentation at high collision energies produces two ions at m/z 127 and 155, identifying carbon 9 as the position of the hydroxyl group on hydroxy stearic acid, confirming the structure to be 9-PAHSA.

(C) Acyl chain carbon numbering scheme, molecular formula, mass and names of FAHFAs from the m/z 537 (PAHSA), m/z 509 (PAHPA), m/z 563 (OAHSA), and m/z 535 (POHS or OAHFA) ions.

(D) Constituent fatty acid and hydroxy fatty acid components of FAHFAs. Quantification of FAHFAs in rodent and human foods. In chow and HFD, we found five of the seven isomers that are present in mouse AT, 13/12-, 11-, 10-, 9-, and 8-PAHSA, but not 7- and 5-PAHSA. However, the relative abundance among isomers was strikingly different from AT or serum with 10-PAHSA being most abundant in both diets (Figure 3B). Levels of all these isomers were substantially lower in HFD than chow (Figure 3B). Given that PAHSA increase in WAT during fasting (Figures 2D and 2E), regulation of tissue PAHSA levels does not simply reflect dietary intake. Similarly, the abundance of PAHSA isomers in serum and tissues (Figure 2C) does not correlate with predominant isomers.
PAHSAs Are Synthesized in Mammalian Tissues

To determine whether PAHSAs are synthesized endogenously, we investigated PAHSA biosynthesis in liver and WAT lysates of normal mice. We detected PAHSA biosynthetic activity in both tissues, and it was markedly reduced by heat denaturation (Figure 3C). We also detected PAHSA biosynthesis in vivo. Gavage of mice with 9-hydroxy heptadecanoic acid (9-HHA), a hydroxy fatty acid not normally found in mammalian tissues resulted in synthesis of full-length FAHFAs containing a 9-HHA acyl chain (Figure 3C).

PAHSAs Are Present in Humans and Levels Are Reduced with Insulin Resistance

To determine if PAHSAs are present in humans and are regulated in disease states, we measured PAHSA isomers in serum and SQ

Figure 2. Identification and Quantification of PAHSA Isomers in Mouse Serum and Tissues

(A) Coelution of PAHSA isomers from serum and SQ WAT of WT and AG4OX mice with synthetic standards for individual PAHSA isomers. The peak for 5-PAHSA is shown in red in the inset. Note: different y axis scale for WT SQ WAT (Red numbers) versus AG4OX SQ WAT.

(B) Distribution and quantification of PAHSA isomers in serum and tissues of WT and AG4OX mice. “Ester position” refers to the location of the ester bond in PAHSA isomers. n = 3–5/group, *p < 0.05 versus WT (t test).

(C) Distribution and quantification of 13/12-, 9-, and 5-PAHSA isomers in serum and tissues of WT female FVB mice. n = 3–5/group. **Tissues with different letters are different from each other within the same isomer panel (p < 0.05, ANOVA).

(D and E) Total PAHSA levels (D) and PAHSA isomer levels (E) in serum and tissues of WT mice in fed or fasted (16 hr) states. “Ester position” refers to the location of the ester bond in PAHSA isomers. n = 3–5/group, *p < 0.05, **p < 0.07 versus fed (t test). #Tissues with different letters are different from each other for the fed state (p < 0.05, ANOVA).

Data are means ± SEM. See also Figure S1.

in chow (Figure 3B), suggesting that PAHSAs present in tissues are synthesized endogenously. The fact that 5-PAHSA is not present in mouse chow or HFD (Figure 3B) but is present in WAT, BAT, kidney, and serum (Figures 2A–2C, 2E and 3A) further supports this notion. We also found PAHSAs in all mouse tissues tested with different isomer distributions and abundance (Figure S2C).
WAT from insulin-sensitive and insulin-resistant nondiabetic humans. Subjects were middle-aged. BMI was increased in five out of six insulin-resistant participants (Table S2). Insulin resistance was demonstrated by a 61% reduction in glucose infusion rate during a euglycemic hyperinsulinemic clamp (Table S2). Serum triglycerides and free fatty acids in the fasting state were not different between groups (Table S2). Total PAHSA levels are reduced 40% in serum of insulin-resistant humans (Figure 4A). In serum of both insulin-sensitive and insulin-resistant humans, 9- and 10-PAHSA are most abundant and 13/12- and 5-PAHSAs are present at ~1/5 of these concentrations (Figure 4A). In insulin-resistant people, serum levels of all PAHSAs except 9-PAHSA are reduced by 40%–55% compared to insulin-sensitive people (Figure 4A). Serum concentrations of total PAHSAs and all isomers correlated remarkably strongly with insulin sensitivity measured by clamp (Figure 4B). Serum PAHSA levels did not correlate with levels of nonesterified fatty acids or triglycerides (data not shown) suggesting that PAHSA levels are regulated by different mechanisms.

In human SQ WAT, total PAHSA levels are reduced 70% (Figure 4C). We detected 13/12-, 11-, 10-, 9-, and 5-PAHSA isomers in these biopsies (Figure 4C). However, for technical reasons we were unable to quantify the levels of 11-PAHSA. 9-PAHSA levels were higher than all other isomers (Figure 4C) similar to mouse SQ WAT (Figures 2B, 2E and 3A). 13/12-, 10-, 9-, and 5-PAHSA concentrations in SQ WAT of insulin-resistant people were 60%–73% lower than in insulin-sensitive people (Figure 4C). Concentrations of total PAHSAs and of 9- and 5-PAHSA isomers

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**Figure 3. PAHSA Isomer Levels in Tissues of Mice on Chow or HFD, PAHSA Isomer Levels in Food, and PAHSA Biosynthesis In Vivo and in Tissues In Vitro**

(A) Quantification of PAHSA isomers in serum, SQ WAT, PG WAT, BAT, and liver of WT female FVB mice fed on chow or HFD for 9 weeks. “Ester position” refers to the location of the ester bond in PAHSA isomers. n = 3–6/group, *p < 0.05 versus chow (t test).

(B) Quantification of PAHSA isomers in mouse chow and HFD. n = 3/group.

(C) 9-PAHSA levels in liver and PG-WAT lysates incubated with palmitoyl-CoA and 9-hydroxy stearic acid and heat-denatured Controls. n = 3/group, *p < 0.05 versus heat-denatured Controls (t test).

(D) 9-palmitic-acid-hydroxy-heptadecanoic-acid (9-PAHHA) serum levels in mice 3 hr postgavage with 9-hydroxy-heptadecanoic-acid (9-HHA) or vehicle control. n = 3/group, *p < 0.05 versus vehicle (t test).

Data are means ± SEM. See also Figure S2.
in WAT correlate highly with insulin sensitivity (Figure 4D). Serum PAHSA levels correlated with WAT PAHSA levels only for 5-PAHSA (Figure 4E).

In summary, all PAHSA isomers detected are reduced in SQ WAT in insulin-resistant subjects and all but one are reduced in serum. Furthermore, PAHSA levels in serum and WAT correlate highly with whole-body insulin sensitivity. These effects parallel the effects in diet-induced obese mice in which all PAHSA isomers are reduced in SQ WAT, and 13/12- and 5-PAHSA are lower in serum compared to chow-fed mice (Figure 3A). Thus, the regulation of PAHSAs and their inverse correlation with insulin resistance is conserved between mice and humans.

PAHSAs Acutely Improve Glucose Tolerance and Ambient Glycemia

Since levels of PAHSA isomers correlate with insulin sensitivity (Figures 4B and 4D), we tested whether administration of PAHSAs could improve glucose homeostasis in obese, diabetic mice. We selected 9-PAHSA and 5-PAHSA because: (1) 9-PAHSA was the most abundant form in WAT and BAT in WT mice (Figure 2B) and in SQ WAT of humans (Figure 4C). (2) 9-PAHSA is the most strongly upregulated in serum and WAT of insulin-sensitive AG4OX mice (Figure 2B) and was downregulated (with other isomers) in WAT of insulin-resistant humans (Figure 4C). (3) 5-PAHSA was the most consistently downregulated in all adipose depots and in serum of insulin-resistant mice (Figure 3A) and in WAT and serum of insulin-resistant humans. Oral gavage of 5-PAHSA increased serum levels 3- to 5-fold in mice on chow and HFD (Figure S3). As shown in Figure 3A, baseline 5-PAHSA levels were 50%–80% lower in HFD-fed mice compared to chow-fed mice and 5-PAHSA gavage more than restored the levels (Figure S3). The elevation of serum 5-PAHSA levels after gavage in both chow and HFD-fed mice was similar to the elevation in serum of AG4OX mice (Figures 2B and S2). This indicated that PAHSAs are absorbed orally and this route of administration can be used to increase PAHSA concentrations for in vivo metabolic studies.

Acute oral administration of 5- or 9-PAHSA in insulin-resistant HFD-fed mice lowered basal glycemia at 30 min after PAHSA administration (Figure 5A, −30 to 0 min). Subsequently, glucose was administered and we observed improved glucose tolerance in PAHSA-treated mice with reduced area under the glucose excursion curve (Figure 5A). Because of the significant effect of PAHSAs on the baseline glucose levels in HFD-fed mice at 30 min after administration, we tested whether PAHSA action to lower baseline glycemia was sustained. 5-PAHSA or 9-PAHSA had a greater glucose-lowering effect than vehicle treatment in HFD-fed mice also at 2.5–3 hr after administration (Figure 5B). There was no difference in plasma insulin levels at this time in PAHSA-treated compared to vehicle-treated mice (data not shown). Since no food was available during this study, calorie absorption was not a variable and the results suggest that oral PAHSA administration improves insulin sensitivity. An insulin tolerance test showed lower glucose levels in PAHSA-treated mice compared to vehicle-treated mice for the study duration (120 min after insulin administration) which was largely because of the PAHSA effect to lower baseline glycemia (data not shown).

PAHSAs Stimulate Insulin and GLP-1 Secretion

To determine whether enhanced insulin secretion might contribute to the PAHSA-induced improvement in glucose tolerance (Figure 5A), we tested PAHSA effects on glucose-stimulated insulin secretion (GSIS) in vivo in aged chow-fed mice. 5-PAHSA improved glucose tolerance (Figure 5C) concurrent with acute enhancement of insulin secretion at 5 min after glucose administration (Figure 5D). This may result from direct effects on insulin secretion or from stimulation of incretin secretion since GLP-1 levels were also increased in PAHSA-treated mice at 5 min after glucose administration (Figure 5E). Thus, PAHSAs augment the acute stimulation of GLP-1 and insulin secretion in response to glucose and these effects most likely have an important role in the enhanced glucose tolerance following a single PAHSA dose.

To determine whether the stimulation of insulin secretion is a direct effect of PAHSAs on pancreatic beta cells, we incubated islets from human donors (Table S3, donors’ metabolic parameters) with 5-PAHSA and measured GSIS. 5-PAHSA had no effect on insulin secretion at 2.5 mM glucose but augmented the insulin secretion response at 20 mM glucose (Figure 5F). These data demonstrate that 5-PAHSA directly enhances GSIS in human islets. To determine whether PAHSAs directly stimulate GLP-1 secretion, we used the enteroendocrine cell line STC-1. Both 5- and 9-PAHSA rapidly stimulated GLP-1 secretion from STC-1 cells in a dose-dependent manner (Figure 5G). The effects are similar to those with the omega-3 fatty acid, α-linolenic acid, and a synthetic GPR120 ligand (Figure 5G). Thus, the rapid effects of PAHSAs to augment GSIS may be both direct effects on pancreatic beta cells and indirect effects through stimulation of GLP-1 secretion.

PAHSAs Enhance Insulin-Stimulated Glucose Transport and Glut4 Translocation by Activating GPR120

To further understand the mechanism(s) by which PAHSAs improve glucose homeostasis, we tested their effects on glucose transport in adipocytes. PAHSAs increased glucose transport at submaximal and maximal insulin concentrations (Figure 6A). Neither of the fatty acids that form the parent PAHSA structure, α-linolenic acid,
Figure 5. PAHSAs Improve Glucose Tolerance and Ambient Glycemia In Vivo and Augment Insulin and GLP-1 Secretion

(A) 4.5 hr after food removal, HFD-fed mice were gavaged with 5-PAHSA (top), 9-PAHSA (bottom) or vehicle control. 30 min later an oral glucose tolerance test (OGTT) was performed. n = 12–14/group, mice were on HFD for 42–52 weeks. *p < 0.05 versus vehicle at same time (t test). Area under the curve (AUC) was calculated from 0 to 120 min. *p < 0.05 versus vehicle (t test).

(B) 2.5 hr after food removal, HFD-fed mice were gavaged with 5-PAHSA (top), 9-PAHSA (bottom) or vehicle control. Glycemia was measured immediately before (time 0) and at 2.5 hr (5-PAHSA) or 3 hr (9-PAHSA) after PAHSA gavage. n = 12–14/group. *p < 0.05 versus vehicle (t test).

(C) 4.5 hr after food removal, aged, chow-fed mice (45-week-old) were gavaged with 5-PAHSA 30 min prior to an OGTT. n = 12–14/group. *p < 0.05 versus vehicle at same time (t test). Area under the curve (AUC) was calculated from 0 to 120 min. *p < 0.05 versus vehicle (t test).

(D and E) Serum insulin levels (D) and serum GLP-1 levels (E) 5 min postglucose challenge in chow-fed mice gavaged with 5-PAHSA or vehicle (glucose values shown in C). n = 12–14/group, *p < 0.05 versus vehicle (t test).

(F) Insulin secretion from primary human islets from two independent donors. Islets were incubated with low (2.5 mM) or high (20 mM) glucose ex vivo in the presence of 5-PAHSA (20 μM) or Control (KRB buffer). Diluent for 5-PAHSA was methanol (0.25%). n = 100 islets/condition, *p < 0.05 versus control 2.5 mM glucose (t test), #p < 0.05 versus all treatments at 2.5 mM glucose (t test), $p < 0.05 versus control and diluent at 20 mM glucose (t test).

(G) Active GLP-1 secretion from STC-1 cells in response to 5-PAHSA (5P), 9-PAHSA (9P), α-Linolenic Acid (ALA), GW9508 (GW), or vehicle control (CTL, DMSO). n = 4/group, *p < 0.05 versus vehicle (CTL) (t test). Data are means ± SEM. See also Table S3 and Figure S3.
palmitic acid or hydroxystearic acid, alone improved insulin-stimulated glucose transport (Figure 6B). The effects of PAHSAs on insulin-stimulated glucose transport occurred with both acute (30 min) and chronic (2–6 day) treatment and at concentrations as low as 500 nM (Figure S4A). PAHSAs did not alter total cellular Glut1 or Glut4 protein levels in adipocytes even after 6 days of incubation (data not shown).

Bioactive lipids can influence biology through binding to cell surface receptors such as G protein-coupled receptors (GPCRs) (Hara et al., 2013). The effects of PAHSAs on GLP-1 secretion and glucose transport are consistent with possible activation of GPCRs (Hirasawa et al., 2005). To determine whether PAHSAs activate GPCRs, we screened a panel of known lipid-activated GPCRs. Both 9-PAHSA (Figure 6C) and 5-PAHSA (data not shown) dose-dependently bind to and activate GPR120 which is also activated by ω-3 fatty acids and monounsaturated fatty acids (Hirasawa et al., 2005; Oh et al., 2010). Activation of GPR120 increases glucose transport and Glut4 translocation in adipocytes (Oh et al., 2010). To test whether GPR120 mediates the effects of PAHSAs, we knocked it down >95% in adipocytes with siRNA (Figure S4B). This completely reversed the enhanced effects of PAHSAs on insulin-stimulated glucose transport (Figure 6D). These data demonstrate that GPR120 mediates the effects of PAHSAs on insulin-stimulated glucose transport.

To determine the mechanism for enhancement of glucose transport with PAHSAs, we analyzed the effects on insulin-induced Glut4 translocation to the plasma membrane in adipocytes. In the absence of insulin, PAHSAs had no effect on Glut4 translocation (Figure 6F). However, PAHSAs enhanced Glut4 translocation at submaximal and maximal insulin stimulation.
A

CD80

CD86

CD40

MHCII

CD11c

% CD11c + MHCII+
% CD11c + CD86+
% CD11c + CD40+
% CD11c + CD80+

B

C

D

Control

9-PAHSA

Chow

HFD

IL-1β

TNF

E

Control

9-PAHSA

% TNF+ Macrophages
% IL-1β+ TNF+ Macrophages
% IL-1β+ Macrophages
% IL-1β+ Macrophages
concentrations (Figures 6E and 6F). These data indicate that PAHSAs augment insulin-stimulated glucose transport by enhancing Glut4 translocation. Knockdown of GPR120 in adipocytes completely blocked the effect of PAHSAs to augment insulin-stimulated Glut4 translocation (Figures 6E and 6F). We obtained this effect with two different GPR120 siRNAs indicating that it is a specific “on target” effect. These data together demonstrate that PAHSAs bind to and activate GPR120 (Figure 6C), which mediates the effects of PAHSAs on insulin-stimulated glucose transport and Glut4 translocation (Figures 6D–6F).

Administration of 9-PAHSA normalized the percentage of TNFα positive ATMs and reduced the percentage of IL-1β positive and double-positive ATMs (Figure 7D). Total ATM number was not reduced due to the short treatment duration (data not shown). This demonstrates that PAHSAs have anti-inflammatory effects in vivo. Therefore, reduced PAHSA levels in insulin-resistant states (Figures 3A and 4) could contribute to activation of the innate immune system, thus playing a role in AT inflammation and systemic insulin-resistance.

**DISCUSSION**

Adipose-Glut4 levels in humans are tightly associated with insulin sensitivity and lower Glut4 levels confer increased T2D risk (Carvalho et al., 2001; Shepherd and Kahn, 1999). Adipose-specific Glut4 overexpression in mice (AG4OX) causes beneficial metabolic effects which result from enhanced ChREBP-driven de novo lipogenesis in adipose tissue (AT) (Herman et al., 2012; Tozzo et al., 1995). Because this occurs even in the setting of obesity and elevated serum fatty acids (Herman et al., 2012; Tozzo et al., 1995), we sought to determine whether the augmented lipogenesis in AG4OX mice leads to production of lipid species that have favorable metabolic effects. Here we report the discovery of a class of mammalian lipids characterized by a branched ester linkage between a fatty-acid and a hydroxy-fatty acid or FAHFAs. The closest reported structures are (O-acyl)-omega hydroxy-fatty acids in the eye (Butovich et al., 2009), which are not branched.

Of the 16 FAHFA family members we report here, we extensively characterized the biology of PAHSA isomers that are present in many, if not all, tissues and in serum in normal mice and in human WAT and serum (Figures 1, 2, 3, and 4). Total PAHSA levels are highest in WAT and BAT that also have the greatest number of PAHSA isomers (Figure 2). Furthermore, PAHSA levels are highly elevated in serum, WAT and BAT of AG4OX mice that are obese but have markedly enhanced glucose tolerance (Herman et al., 2012; Shepherd et al., 1993). In parallel, nearly all PAHSA isomers are higher in serum and SQ WAT of insulin-sensitive humans compared to insulin-resistant humans.

In insulin-resistant, obese mice, PAHSA isomer levels show adipose-depot-specific regulation and all PAHSA isomers are lower in SQ WAT of obese compared to lean mice (Figure 3). These results in SQ WAT are similar to those in insulin-resistant

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**Figure 7. 9-PAHSA Inhibits LPS-Induced Dendritic Cell Maturation In Vitro and Proinflammatory Cytokine Production from Adipose Tissue Microenvironments In Vivo**

(A) LPS induces dendritic cell (DC) maturation (increased percentage of CD11c+ cells expressing costimulatory molecules, CD80, CD86, CD40, and MHCII). This LPS effect is reduced in the presence of 9-PAHSA (40 μM) compared to vehicle (DMSO) control. Quantification of CD11c+ cells which are positive for costimulatory molecules from the panel above. n = 3 mice/group.

(B) LPS-induced DC maturation is inhibited by increasing concentrations of 9-PAHSA. Red triangles represent vehicle for 9-PAHSA (DMSO, “-“) control. Red triangles represent vehicle for 9-PAHSA (DMSO, “-“) without LPS. n = 3 mice/group.

(C) LPS-induced cytokine secretion from DC’s is inhibited by increasing concentrations of 9-PAHSA compared to vehicle for 9-PAHSA (DMSO, “-“) control. Quantification D-percentage of AT macrophages expressing TNFα and IL-1β. Mouse fed on HFD or chow mice were gavaged for 3 days with 9-PAHSA (30 mg/kg for chow mice and 45 mg/kg for HFD mice) vehicle control, PG-WAT was harvested on day 4 and the stromal vascular cells were incubated in vitro with PMA, ionomycin and brefeldin for 5 hr. AT macrophages (CD45+CD11b+F4/80+) were stained intracellularly for TNFα and IL-1β.

(D) Flow cytometry representation of AT macrophages expressing TNFα and IL-1β. Mice fed on HFD or chow mice were gavaged for 3 days with 9-PAHSA (30 mg/kg for chow mice and 45 mg/kg for HFD mice) vehicle control, PG-WAT was harvested on day 4 and the stromal vascular cells were incubated in vitro with PMA, ionomycin and brefeldin for 5 hr. AT macrophages (CD45+CD11b+F4/80+) were stained intracellularly for TNFα and IL-1β.

(E) Quantification D-percentage of AT macrophages expressing TNFα, IL-1β or both. n = 5 mice/group. LPS concentration is 100 ng/ml for all panels. *p < 0.05 versus LPS-activated cells without PAHSA treatment (A–C) or control cells, same diet (E) by one-way (A–C) and two-way ANOVA (D). ^p < 0.05 versus all other groups by two-way ANOVA. Data are means ± SEM. See also Figure S5.
people (Figure 4). 5-PAHSA is consistently reduced in all adipose depots studied and in serum in insulin-resistant mice and humans. In insulin-resistant people, most PAHSA isomers are reduced in serum and AT and correlate highly with insulin sensitivity (Figure 4). ChREBP is required to maintain normal PAHSA levels in WT mice and elevated levels in AG4OX mice (Figure 1F).

In humans, ChREBP and lipogenic enzyme expression correlate strongly with insulin-sensitivity. Thus, the reduction in PAHSAs in insulin-resistant people may be mediated by suppressed ChREBP expression.

This class of lipids has multiple effects that improve glucose-insulin homeostasis which suggests that restoring PAHSA levels in insulin-resistant people could have beneficial metabolic effects. Oral PAHSA administration in insulin-resistant mice on a HFD rapidly lowers ambient glucose and also improves glucose tolerance (Figures 5A and 5B). This may result, at least in part, from enhanced glucose transport since PAHSAs augment insulin-stimulated glucose transport and Glut4 translocation directly in adipocytes in vitro (Figure 6). In addition, PAHSAs stimulate both insulin and GLP-1 secretion (Figures 5D–5G). Importantly, the effects of PAHSAs on insulin secretion are observed only under hyperglycemic conditions (Figure 5F). The enhanced GSIS in vivo most likely results from both direct effects on islet cells and indirect effects though GLP-1-stimulated insulin secretion.

Total PAHSA levels in tissues and serum are similar to concentrations of signaling lipids such as prostacyclins, prostaglandins, steroids, and endocannabinoids. PAHSAs are signaling lipids that directly bind to and activate GPR120 in a cell-based GPCR activity assay (Figure 6C). GPR120 activation appears to explain the effects on insulin-induced Glut4 translocation and glucose uptake in adipocytes (Figures 6B–6D) and may explain the effects on GLP-1 secretion and inhibition of inflammatory responses in immune cells. Thus, PAHSAs are endogenous GPR120 ligands and may also exert effects through other lipid-activated GPCRs.

PAHSAs have striking anti-inflammatory effects and largely block LPS-stimulated dendritic cell activation and cytokine production (Figure 7). Chronic, low-grade inflammation in AT plays an important role in obesity-related insulin resistance (Lumeng and Saltiel, 2011; Olefsky and Glass, 2010). Three days of PAHSA gavage in HFD-fed mice reduced the percentage of ATMs that express proinflammatory cytokines (Figure 7D). Therefore, anti-inflammatory effects of PAHSAs may promote insulin sensitivity and ameliorate other inflammatory diseases.

Insulin-sensitizing and anti-inflammatory effects and GLP-1 secretion are also observed with ω-3 fatty acids (Hirasawa et al., 2005; Oh et al., 2010). However, a major difference between ω-3 fatty acids and PAHSAs is that PAHSAs are synthesized endogenously as evidenced by PAHSA biosynthetic activity in adipose and liver lysates and incorporation of modified precursors into FAHFAs in vivo (Figures 3C and 3D). The potential importance of identifying an endogenous GPR120 ligand is demonstrated by the fact that loss of function mutations in GPR120 in humans promote obesity and insulin resistance (Ichimura et al., 2012). Thus, GPR120 is an important control point in the integration of anti-inflammatory and systemic insulin-sensitizing responses and is emerging as an important regulator of whole-body glucose-insulin homeostasis (Mo et al., 2013). It is the subject of ongoing preclinical investigation for the treatment of obesity-related insulin resistance, T2D, and inflammatory diseases (Mo et al., 2013; Oh et al., 2014).

Reduced PAHSA levels may contribute to diabetes risk since many PAHSA isomers are reduced in SQ WAT and serum of insulin-resistant rodents (Figure 3A) and humans (Figure 4). In humans, PAHSA levels in both serum and SQ WAT correlate highly with whole-body insulin-sensitivity (Figures 4B and 4D). Thus, reduced circulating PAHSA levels may serve as a biomarker for insulin resistance and T2D risk. PAHSAs lower glycemia, improve glucose tolerance and stimulate GLP-1 and insulin secretion in mice (Figure 5). This raises the possibility that restoring the reduced PAHSA levels in insulin-resistant humans could have therapeutic effects to prevent or ameliorate insulin resistance and T2D. The discovery of the FAHFAs is important also because their presence suggests uncharacterized biochemical pathways and enzymes that may be important in human physiology and disease. Changes in the levels of these metabolites and in their signaling pathways may provide important insights and new treatment avenues for metabolic and inflammatory diseases.

**EXPERIMENTAL PROCEDURES**

**Materials and Reagents**

All chemicals were purchased from Sigma-Aldrich (St. Louis, MO) unless otherwise stated.

**Culture and Differentiation of Cells**

ST3-L1 fibroblasts were cultured and differentiated as described (Norseen et al., 2012). STC-1 cells were maintained in DMEM supplemented with 10% FCS, pen/strep and maintained at 37 °C and 5% CO₂.

**Pancreatic Islets, GSIS, and GLP-1 Secretion Studies**

Human islets from nondiabetic donors were obtained from Prodo Laboratories (Irvine, CA). GSIS studies (Kowluru et al., 2010) and GLP-1 secretion from STC-1 cells (Hirasawa et al., 2005) were performed as described.

**Generation and Treatment of Bone-Marrow-Derived Dendritic Cells**

BMDCs were generated as described (Moraes-Vieira et al., 2014). Cells were incubated with 9-PAHSA 10 min prior to LPS (100 ng/mL) stimulation. CD11c, MHC II, CD40, CD80 and CD86 (all Biolegend) were detected by flow cytometry as described (Moraes-Vieira et al., 2014). Cytokine levels were measured by ELISA (Biolegend).

**9-PAHSA Biosynthetic Activity Assay**

Liver and PG-WAT tissue was Dounce homogenized in buffer A (10 mM Tris-HCL (pH 7.4), 250 mM Sucrose containing protease inhibitors (Roche)). Lysates were centrifuged at 1,200 g to remove incompletely lysed cells and debris. Lysates were then adjusted to 1 mg/ml protein and 100 μl was incubated with 100 μM palmityl-CoA and 100 μM 9-hydroxy stearic acid (PAHSA substrates) for 2 hr at 37°C. Control samples were heat denatured by boiling for 10 min prior to incubation with PAHSA substrates. After 2 hr the reaction was stopped by the addition of 300 μl cold buffer A followed by 400 μl of methanol (MeOH) and 800 μl of chloroform. Samples were vortexed and centrifuged at 1,200 g for 5 min. 9-PAHSA levels in the organic layer were measured by LC-MS.

**FAHFA Synthesis In Vivo**

Two hours post food removal C57Bl/J mice were gavaged with 25 mg/kg of 9-hydroxy heptadecanoic acid or vehicle. Three hours later mice were sacrificed and serum was collected. Serum lipids were extracted and 9-PAHHA levels measured by LC-MS.
Animal Studies and Measurement of Metabolic Parameters
Female AG4OX mice and WT FVB littermate controls (Shepherd et al., 1993) at 8–14 weeks old were used for FAHFA tissue distribution and regulation with fasting and HFD studies. ChREBP-KO, ChREBP-KO/AG4OX and control females (Herman et al., 2012) were used at 16–18 weeks. Mice were fed on chow (Lab Diet, 5008) or HFD (Harlan Teklad, TD.93075) for 9 weeks (female FVB) or 42–52 weeks (male C57BL6/J). OGTT’s were performed as described (Moraes-Vieira et al., 2014) after 5 hr food removal.

Anti-Inflammatory Effects of 9-PAHSA In Vivo
Male C57BL6/J mice on chow or HFD described above were gavaged once a day for 3 days with 30 mg/kg (chow) or 45 mg/kg (HFD) of 9-PAHSA or an equivalent volume of vehicle. On the 4th day PG stromal vascular fraction (SVF) cells were harvested and cultured for 5 hr with ionomycin, PMA and brefeldin at 37°C and the intracellular cytokine content was measured in gated CD45+, CD11b+, and F4/80+ cells as described (Moraes-Vieira et al., 2014).

Human Studies
Hyperinsulinemic-euglycemic clamp was performed in 13 nondiabetic subjects. SQ WAT biopsies were obtained from the peri-umbilical, abdominal region after an overnight fast.

Lipid Extraction
Lipid extraction was performed as described (Bligh and Dyer, 1959; Saghatelian et al., 2004). Tissues (60–150 mg) were Dounce homogenized on ice in a mixture of 1.5 ml MeOH, 1.5 ml chloroform and 3 ml citric acid buffer. PAHSA standards were added to chloroform prior to extraction. The resulting mixture was centrifuged and the organic phase containing extracted lipids was dried under N2 and stored at –80°C prior to solid phase extraction.

Lipidomic Analysis
Lipidomic analysis was performed using an Agilent 6220 ESI-TOF fitted with an electrospray ionization source with a capillary voltage of 3,500 V and fragmentor voltage of 100 V. A Gemini C18 reversed phase column (Phenomenex) and a C18 reversed phase guard column (Western Analytical) was used for LC-MS analysis in negative mode. In positive mode, a Luna C5 reversed phase column (Phenomenex) was used together with a C4 reversed phase guard column (Western Analytical). Drying gas temperature was 350°C, flow rate 10 l/min and nebulizer pressure 45 psi. Untargeted data were collected using an m/z of 100–1,500.

ADD
Structural characterization of the FAHFA’s of MS/MS was carried out on an Agilent 6510 quadrupole-time of flight MS.

Synthesis of PAHSA and PAHSA Standards
Detailed information on synthesis of PAHSA and PAHSA standards is outlined in the Supplemental Information. Multiple comparisons as specified in figure legends. All statistical analyses were performed with GraphPad Prism 5.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Extended Experimental Procedures, four figures, four tables, and one data file and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2014.09.035.

AUTHOR CONTRIBUTIONS
M.M.Y and I.S conceived of, designed, performed, and interpreted experiments and made figures. P.M.M.-V. designed, performed and interpreted the immunology experiments and made the figures. M.A.H. conceived of the untargeted lipidomics experiment and E.A.H. and M.A.H. designed and performed the untargeted lipidomics experiment. E.A.H. designed, performed and interpreted the data from the structure elucidation studies. E.A.H. and A.S. designed and performed chemical synthesis of the lipids. T.Z., I.S., E.A.H. and S.C. developed and applied new targeted lipidomics methods. J.L., A.S.D., O.D.P. assisted with animal studies. B.B.K., I.S., U.S., and A.H. designed, performed and interpreted the data from human studies. R.T.P, T.E.M. designed and performed Glut4 translocation assays. B.B.K. and A.S conceived of, designed and supervised the experimental plan and interpreted experiments. B.B.K., A.S., M.M.Y. and I.S. wrote the manuscript. M.A.H., P.M.M.V., O.D.P., T.E.M., and J.L. edited the manuscript.

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