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Author(s)	Takahashi, Haruya; Kamakari, Kosuke; Goto, Tsuyoshi; Hara, Hideyuki; Mohri, Shinsuke; Suzuki, Hideyuki; Shibata, Daisuke; Nakata, Rieko; Inoue, Hiroyasu; Takahashi, Nobuyuki; Kawada, Teruo	
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1	9-Oxo-10( <i>E</i> ),12( <i>Z</i> ),15( <i>Z</i> )-octadecatrienoic acid activates peroxisome	
2	proliferator-activated receptor $\alpha$ in hepatocytes	
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4	Haruya Takahashi <sup>1</sup> , Kosuke Kamakari <sup>1</sup> , Tsuyoshi Goto <sup>1,2</sup> , Hideyuki Hara <sup>1</sup> , Shinsuke	
5	Mohri <sup>1</sup> , Hideyuki Suzuki <sup>3</sup> , Daisuke Shibata <sup>3</sup> , Rieko Nakata <sup>4</sup> , Hiroyasu Inoue <sup>4</sup> ,	
6	Nobuyuki Takahashi <sup>1,2</sup> and Teruo Kawada <sup>1,2,#</sup>	
7		
8	<sup>1</sup> Laboratory of Molecular Function of Food, Division of Food Science and	
9	Biotechnology, Graduate School of Agriculture, Kyoto University, Kyoto Japan	
10	<sup>2</sup> Research Unit for Physiological Chemistry, Kyoto University, Kyoto, Japan	
11	<sup>3</sup> Kazusa DNA Research Institute, Chiba, Japan	
12	<sup>4</sup> Department of Food Science and Nutrition, Nara Women's University, Nara, Japan	
13		
14	<sup>#</sup> To whom correspondence should be addressed. Tel: +81-774-38-3751; Fax:	
15	+81-774-38-3752; E-mail: fat@kais.kyoto-u.ac.jp	
16		
17	Short title: 9-oxo-OTA activates PPAR $\alpha$ in hepatocytes	
18	Abbreviations: 9-oxo-OTA, 9-oxo-10( <i>E</i> ),12( <i>Z</i> ),15( <i>Z</i> )-octadecatrienoic acid;	
19	9-oxo-ODA, 9-oxo-10,12-octadecadienoic acid; BSA, bovine serum albumin; DMEM,	
20	Dulbecco's modified Eagle's medium; EtOH, ethanol; HAD, cis-10-heptadecenoic acid;	
21	LC-MS, liquid chromatography-mass spectrometry; LNA, linoleic acid; ALA,	
22	$\alpha$ -linolenic acid; PBS, phosphate-buffered saline; PPAR, peroxisome	
23	proliferator-activated receptor; QTOFMS, quadrupole-time-of-flight MS; UPLC, ultra	
24	performance LC	
25		

# 27 Abstract

Peroxisome proliferator-activated receptor (PPAR) $\alpha$  is mainly expressed in the liver 28and plays an important role in the regulation of lipid metabolism. It has been reported 29that PPARα activation enhances fatty acid oxidation and reduces fat storage. Therefore, 30 31PPARa agonists are used to treat dyslipidemia. In the present study, we found that 329-oxo-10(E),12(Z),15(Z)-octadecatrienoic acid (9-oxo-OTA), which is  $\alpha$ -linolenic acid 33 (ALA) derivative, is present in tomato (Solanum lycopersicum) extract. We showed that 9-oxo-OTA activated PPARa and induced the mRNA expression of PPARa target genes 3435in murine primary hepatocytes. These effects promoted fatty acid uptake and the secretion of  $\beta$ -hydroxybutyrate, which is one of the endogenous ketone bodies. We also 36 37 demonstrated that these effects of 9-oxo-OTA were not observed in PPARa-knockout 38 (KO) primary hepatocytes. To our knowledge, this is the first study to report that 9-oxo-OTA promotes fatty acid metabolism via PPARa activation and discuss its 39 40 potential as a valuable food-derived compound for use in the management of 41dyslipidemia.

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43 **Key words:** PPARα, oxylipin, hepatocyes, fatty acid metabolism, LC-MS.

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# 46 Introduction

Dyslipidemia, which results from obesity, is a recognized risk factor for serious disorders, including arteriosclerosis, hyperlipidemia, hepatic steatosis, insulin resistance, and cardiovascular disease [1-4]. Therefore, avoidance of dyslipidemia is important to help prevent these lifestyle diseases.

51Peroxisome proliferator-activated receptors (PPARs) are a family of ligand-activated 52transcription factors [5-9] comprised of three isoforms PPAR $\alpha$ , PPAR $\beta/\delta$ , and PPAR $\gamma$ . PPAR $\alpha$  is very important in the regulation of lipid metabolism [5-7] and is expressed at 53high levels in the liver where it promotes  $\beta$ -oxidation, ketogenesis, and lipid transport 5455[10, 11]. PPAR $\alpha$  activation induces the expression of PPAR $\alpha$ -regulated genes (e.g., carnitine-O-palmitoyltransferase 1 [CPT] and acyl-CoA oxidase [ACO]) and activates 56 $\beta$ -oxidation [12-14]. These effects result in increased energy expenditure and reduced 57fat storage [15]. It has been reported that PPARa activation enhances fatty acid 5859metabolism and decreases the levels of circulating and cellular lipids in obese diabetic 60 patients [9, 16]. Therefore, regulation of PPAR $\alpha$  activity is one of the important ways to 61 manage chronic diseases related to lipid metabolism dysfunction.

Tomato (Solanum lycopersicum) is one of the most extensively consumed crops 6263 worldwide. Numerous studies have shown that dietary tomato intake is associated with a reduced risk of chronic diseases [17–20]. Recently, we reported that tomatoes contain 64 65 the oxidized linoleic acid (LNA) derivatives 9-oxo-10,12-octadecadienoic acid (9-oxo-ODA) and 13-oxo-9,11-octadecadienoic acid (13-oxo-ODA), which can 66 function as PPARa agonists [21,22]. Oxo-ODAs are oxylipins, which include fatty acid 67 hydroperoxides, hydroxyl-, oxo-, epoxy, and keto fatty acids, divinyl ethers, aldehydes, 68 and jasmonic acid [23]. Tomato is believed to contain not only oxo-ODAs but also 69

many other beneficial oxylipins that enhance lipid metabolism and suppress diseases
such as arteriosclerosis, hyperlipidemia, and hepatic steatosis.

The aim of the present study was to identify the metabolite in tomato that enhances 72fatty acid metabolism and to determine its mechanism of action. In the present study, we 73 showed that 9-oxo-10(E),12(Z),15(Z)-octadecatrienoic acid (9-oxo-OTA), which is a 7475derivative of  $\alpha$ -linolenic acid (ALA), is present in tomato fruit extract. We showed that 9-oxo-OTA activates PPARa and induces the mRNA expression of PPARa target genes, 76which promotes fatty acid metabolism in murine hepatocytes. To the best of our 77knowledge, this is the first study to report that 9-oxo-OTA induces fatty acid 78metabolism through PPARa activation, and our findings suggest that 9-oxo-OTA could 79 be a valuable food-derived compound capable of improving dyslipidemia. 80

#### 82 Materials and methods

#### 83 Materials

Authentic 9-oxo-OTA was purchased from Cayman Chemicals (Ann Arbor, MI, USA). Authentic *cis*-10-heptadecenoic acid (HDA), which was used as an internal standard, was purchased from Sigma (St. Louis, MO, USA). All other chemicals were obtained from Sigma or Wako (Osaka, Japan). Buffers used were of high-performance liquid chromatography (HPLC) or liquid chromatography-mass spectrometry (LC-MS) grade. Tomatoes were obtained from a local market.

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# Analysis of 9-oxo-OTA by ultra-performance liquid chromatography-quadrupole time-of-flight mass spectrometry (UPLC-QTOF-MS)

The levels of 9-oxo-OTA were assessed as previously described, using a LC-MS system [24]. Briefly, each freeze-dried tomato (10 mg) was homogenized in 1 mL of extraction solvent (99.5% EtOH containing 1  $\mu$ g/mL HDA) with a mixer. After centrifugation (15,000 rpm, 10 min, 4°C), the supernatant was collected for use as an extract. The extract was filtered through a 0.2- $\mu$ m-pore polyvinylidene difluoride (PVDF) membrane (Whatman, Brentford, UK), and the filtrate was used for LC-MS.

99 LC-MS was performed using a Waters Acquity UPLC system (Milford, MA) 100 coupled to a Xevo QTOF-MS equipped with an electrospray source operating in 101 negative ion mode (ESI negative mode), with a lock-spray interface for accurate mass 102 measurements. Leucine enkephalin was employed as the lock-mass compound. It was 103 infused directly into the MS at a flow rate of 30  $\mu$ L/min and a concentration of 200 104  $\mu$ g/mL (in 50% acetonitrile, 50% water, 0.1% formic acid). The capillary, sampling cone, 105 and extraction cone voltages were set at 2700 V, 20 V, and 1 V, respectively. The source and desolvation temperatures were 120°C and 450°C, respectively. The cone and
desolvation gas flow rates were set at 50 and 800 L/h, respectively.

An aliquot of the extracted sample (3 µL) was injected into an Acquity UPLC 108 BEH-C18 reversed-phase column (column size,  $2.1 \times 100$  mm; particle size, 1.7 µm). 109 110 Mobile phases A (water and 0.1% formic acid) and B (acetonitrile and 0.1% formic 111 acid) were used. The column temperature was set to 40°C. The buffer gradient consisted 112of 30% to 50% B for 0-4 min, 50% to 85% B for 4-14 min, 99% B for 14-17 min, and 113 30% B for 3 min, at a flow rate of 300 µL/min. Data were acquired with MassLynx software (Waters). External mass calibration was performed following 114the 115manufacturer's protocol.

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# 117 Analysis of β-hydroxybutyrate by UPLC-QTOF-MS

The medium from a culture of primary hepatocytes (50  $\mu$ L) was mixed with 50  $\mu$ L of acetonitrile. After centrifugation (15,000 rpm, 10 min, 4°C), the supernatant was collected for use as an as an extract. The extract was filtered through a 0.2- $\mu$ m-pore polyvinylidene difluoride (PVDF) membrane (Whatman), and the filtrates were used for LC-MS.

In the MS system, the capillary, sampling cone, and extraction cone voltages were set at 2600 V, 15 V, and 3 V, respectively (ESI negative mode). The source and desolvation temperatures were 120°C and 450°C, respectively. The cone and desolvation gas flow rates were set at 50 and 800 L/h, respectively.

We used a BEH Amide column (column size,  $2.1 \times 100$  mm; particle size,  $1.7 \mu$ m). The column temperature was set to 40°C. The buffer gradient consisted of 95% B for 0–4 min, 95% to 50% B for 4–10 min, 50% B for 10–16 min, and 95% B for 4 min, at a 130 flow rate of 300 µL/min. Data were acquired with MassLynx software (Waters).

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#### 132 Luciferase assay

Luciferase assays were performed as previously described, using a GAL4/PPAR 133 chimera system [25]. We transfected p4xUASg-tk-luc (a reporter plasmid), 134135pM-hPPARa (an expression plasmid for a chimeric protein containing the GAL4 136DNA-binding domain and each human PPAR-ligand-binding domain), and pRL-CMV 137(an internal control for transfection efficiency) into monkey CV1 kidney cells by using 138Lipofectamine (Life Technologies Japan Ltd.) according to the manufacturer's protocol. 139Luciferase activity was assayed using the dual luciferase system (Promega, Madison, 140 WI, USA) according to the manufacturer's protocol.

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### 142 **Preparation of mouse primary hepatocytes**

143All the animal experiments were approved by the Kyoto University Animal Care Committee. C57BL/6J male mice (wild type and PPAR $\alpha^{-/-}$ , free-feed) were anesthetized 144with intraperitoneal administration of pentobarbital, and the liver was perfused with 145146liver perfusion medium (Life Technologies Japan Ltd.). Then, the liver was treated with liver digestion medium (Life Technologies Japan Ltd.). After filtration through a 147148100- $\mu$ m nylon mesh filter, hepatocytes were isolated by repeated centrifugation at 50  $\times$ 149g for 3 min (3 times). The isolated hepatocytes were cultured in type-1 collagen-coated 12-well plates at a cell density of  $2.0 \times 10^5$  cells/well. After a 5-h incubation at 37°C in 1505% CO<sub>2</sub> in 20% serum DMEM (1.0g/L glucose), the hepatocytes were cultured in 151152serum-free DMEM (1.0g/L glucose) with or without 9-oxo-OTA for 5 h (mRNA assay) or 8 h (LC-MS and RI assay), and then used for mRNA quantification and LC-MS 153

assays.

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#### 156 Quantification of mRNA expression levels

157Total RNA was prepared from primary hepatocytes using Sepasol (Nacalai Tesque), 158according to the manufacturer's protocol. Using M-MLV reverse transcriptase (Life 159Technologies Japan Ltd.), total RNA was reverse transcribed. To determine mRNA 160 expression levels, real-time quantitative RT-PCR analysis was performed with a Light 161 Cycler System (Roche Diagnostics) using SYBR green fluorescence signals as 162described previously [25, 26]. The oligonucleotide primer sets for mouse 36B4 (internal 163 control) and the PPAR $\alpha$  target genes were designed using a PCR primer selection 164program on the website of the Virtual Genomic Center at GenBank and the sequences 165are shown in Table 1. All mRNA expression data are presented as ratios relative to the 166 control in each experiment.

167

# Measurement of <sup>14</sup>C-Palmitic acid uptake and <sup>14</sup>CO<sub>2</sub> production in murine primary hepatocytes

170These experiments were conducted as previously described with some modifications [27-29]. In the measurement of <sup>14</sup>C-palmitic acid uptake, the cells were washed with 171PBS (500 µL/well) and then incubated in assay buffer (<sup>14</sup>C-palmitic acid, 2 µCi/well; 172fatty acid free BSA, 2.5%; and normal palmitic acid, 200 µM) for 3 min at room 173174temperature (1 mL/well). The experiment was stopped by washing the cells 3 times with PBS (500 µL/well). The cells were lysed in 0.1% Triton-X solution (200 µL/well). The 175176 radioactivity in the cell lysate was counted for 1 min in 2 mL of scintillation solution to measure fatty acid uptake. In the measurement of  ${}^{14}CO_2$  production, mice primary 177

hepatocytes were incubated in assay buffer (DMEM containing <sup>14</sup>C-palmitic acid, 2 178179µCi/well; glucose, 1.0g/L; L-carnitine, 200 µM; fatty acid free BSA, 2.5%; and normal palmitic acid, 200 µM) for 8h. The medium was transferred to a 50-mL tube. An 180 181 uncapped-2mL sample tube containing a piece of filter paper soaked in 0.1N NaOH was placed inside a 50-mL tube. After the tube was sealed, 200µL of 70% perchloric acid 182was added to the medium sample to release <sup>14</sup>CO<sub>2</sub>. The saturated filter paper containing 183trapped <sup>14</sup>CO<sub>2</sub> was assessed for radioactivity in a liquid scintillation counter (LS6500, 184185Beckman Coulter, CA, USA).

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#### 187 Statistical analyses

Data are presented as mean  $\pm$  standard error of the mean (SEM). Differences between groups were compared with the Student's *t*-test (for two groups) and one-way analysis of variance (ANOVA) followed by Tukey's test. *P* values less than 0.05 were considered statistically significant.

#### 193 **Results**

#### 194 Identification of 9-oxo-OTA in tomato fruit extract

195First, we explored the oxylipins, which are metabolized from ALA in plants (Fig. 1), 196 in the tomato extract using LC-MS. The results showed that the extract of tomato fruit 197 homogenate obtained under heating (37 °C, 30 min) contained 9-oxo-OTA (Fig. 2 A; Rt 198= 7.91 min, m/z = 291.20, Fig. 2 B, C). The 9-oxo-OTA peak was detected as an 199unfragmented deprotonation ion  $(C_{18}H_{27}O_3^-, [M-H]^-, Fig. 2 C)$  by electrospray negative 200 ionization mass spectrometry. In the tandem mass spectrometry (MS/MS) data, we 201confirmed that the fragment ion detected in the tomato extract (m/z = 185, 125, and202 121) matched the pure 9-oxo-OTA sample (Fig. 2D).

Quantitative analysis of the 9-oxo-OTA content in each part of the tomato fruit homogenate under liquid nitrogen (gelatinous tissue: low limit of quantification (LLOQ), sarcocarp: LLOQ, peel: LLOQ; Table 2) or heating (gelatinous tissue: approximately 0.46  $\mu$ g/g of tissue weight, sarcocarp: 0.23  $\mu$ g/g of tissue weight, peel: 0.62  $\mu$ g/g of tissue weight; Table 2) revealed that the amount of 9-oxo-OTA in all tissues was increased by heat treatment.

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# 9-oxo-OTA activated PPARα and induced the mRNA expression of PPARα target genes

9-oxo-OTA is analogous to 9-oxo-10,12-octadecadienoic acid (9-oxo-ODA, Fig. 1) and 13-oxo-9,11,15-octadecatrienoic acid (13-oxo-OTA, Fig. 1). In our previous studies showed that 9-oxo-ODA is involved in PPAR $\alpha$  activation [22] and 13-oxo-OTA activates PPAR $\gamma$  [24]. Therefore, we hypothesized that 9-oxo-OTA activates PPAR $\alpha$ and/or PPAR $\gamma$ . To determine the effect of 9-oxo-OTA on PPAR $\alpha$  and PPAR $\gamma$  activity, we 217first performed a luciferase ligand assay. In the PPARa activation level, 9-oxo-OTA was stronger than 13-oxo-OTA (Fig. 3 A). In the PPARy activation level, 13-oxo-OTA was 218stronger than 9-oxo-OTA (Fig. 3 B). We also showed that 9-oxo-OTA increased 219luciferase activity in a time dependent manner (Fig. 3 C), and this luciferase activity 220was inhibited by PPARa antagonist (GW6471) treatment (Fig. 3 D). PPARa is 221222expressed in liver where it promotes  $\beta$ -oxidation, ketogenesis, and lipid transport [10, 22311] and important in the regulation of lipid metabolism [5-7]. To elucidate the 224contribution of 9-oxo-OTA to lipid metabolism in hepatocytes, murine hepatocytes were 225cultured in medium containing 9-oxo-OTA. In murine hepatocytes, we showed that the expression of PPAR $\alpha$  mRNA was increased by 9-oxo-OTA treatment (approximately 226227 1.8-fold, Fig. 4 A). Furthermore, the mRNA expression levels of PPARa target genes, such as Cpt1a, Acs, Hmgcs2, and Cd36, were increased by 9-oxo-OTA treatment 228229(approximately 1.5-fold, 1.4-fold, 1.5-fold, and 1.4-fold, respectively, Fig. 4 B-E). We 230also demonstrated that the 9-oxo-OTA-mediated effect on PPARa target gene 231expression was not observed in PPARα-KO primary hepatocytes (Fig. 4 B-E). These findings suggest that 9-oxo-OTA promotes the mRNA expression of the genes involved 232233in fatty acid oxidation via PPARa activation.

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# **9-oxo-OTA promoted fatty acid oxidation in primary hepatocytes**

236 PPAR $\alpha$  is the major regulator of fatty acid metabolism [5-7], suggesting that 237 9-oxo-OTA induces fatty acid oxidation via PPAR $\alpha$  activation. We investigated whether 238 9-oxo-OTA influences not only mRNA expression levels but also lipid metabolism. The 239 results showed that <sup>14</sup>C-palmitic acid uptake was increased by 9-oxo-OTA treatment 240 (approximately 1.4-fold, Fig. 5 A), suggesting enhanced fatty acid metabolism. We also

showed that <sup>14</sup>CO<sub>2</sub> production was increased by 9-oxo-OTA treatment (approximately 2412421.5-fold, Fig. 5 B). β-Oxidation is one of the primary modes of fatty acid metabolism, 243which converts fatty acids to ketone bodies.  $\beta$ -hydroxybutyrate, which is one of the ketone bodies, is an important marker of fatty acid oxidation. We investigated whether 2449-oxo-OTA effects the production of  $\beta$ -hydroxybutyrate in murine primary hepatocytes 245and demonstrated that  $\beta$ -hydroxybutyrate secretion was increased by 9-oxo-OTA 246treatment (approximately 1.5-fold, Fig. 5 C). The enhancement of fatty acid uptake and 247secretion of  ${}^{14}CO_2$  and  $\beta$ -hydroxybutyrate induced by 9-oxo-OTA were not observed in 248PPARa-KO primary hepatocytes (Fig. 5). These findings suggest that 9-oxo-OTA 249promotes fatty acid metabolism in primary hepatocytes via a PPARα-dependent 250251pathway.

#### 253 **Discussion**

254In the present study, we analyzed the metabolites in tomato extract by using LC-MS 255and identified 9-oxo-OTA. Our previous study showed that ALA, which is a 9-oxo-OTA precursor, was present in tomato extracts [30]. These data suggested that 2562579-oxo-OTA can be derived from ALA via a non-enzymatic and/or enzymatic reaction. It 258is possible that ALA is susceptible to oxidation at C9 via auto-oxidation and/or 259lipoxygenase is involved in the production of 9-oxo-OTA in tomato fruits. Our previous 260study also revealed that 9-oxo-ODA is localized primarily in the fruit peel [30]. 261However, here we showed that 9-oxo-OTA exists in not only peel but also in the 262gelatinous tissue and sarcocarp. Further examinations are necessary to elucidate the 263differences in the localization of 9-oxo-OTA and 9-oxo-ODA. We also revealed that the 264amount of 9-oxo-OTA in tomato extract is increased by heating. This finding suggested 265that heat processing increases the amount of 9-oxo-OTA in processed tomato foods. 266Tomato fruit contains not only 9-oxo-OTA but also many other compounds. Therefore, 267it is difficult that the effect of tomato on fatty acid oxidation is explained by only 9-oxo-OTA via PPARa dependent pathway. To evaluate 9-oxo-OTA in vivo, it is 268269necessary to process selective breeding (tomato fruit having high 9-oxo-OTA content) and/or to improve processing condition (e.g. tomato juice having high 9-oxo-OTA 270271content).

272 PPAR $\alpha$  is expressed principally in the liver, and it is a key regulator of various 273 aspects of lipid metabolism, including  $\beta$ -oxidation, ketogenesis, and lipid transport [10, 274 11]. PPAR $\alpha$  agonists are used in the treatment of dyslipidemia [9, 16]. Although we 275 previously reported that 9-oxo-ODA contributes to the activation of PPAR $\alpha$  [21], little 276 is known about the effect of 9-oxo-OTA on PPAR $\alpha$ . The structure of 9-oxo-OTA is similar to that of 9-oxo-OTA. Therefore, we hypothesized that 9-oxo-OTA also has the
ability to activate PPARα.

279Here, we demonstrated for the first time that 9-oxo-OTA activates PPAR $\alpha$  in luciferase ligand assay. There is the possibility that 9-oxo-OTA activate PPAR $\alpha$  both 280281directly and/or indirectly. In case of the direct action, 9-oxo-OTA acts as PPARa ligand. 282In case of the indirect action, the metabolites of 9-oxo-OTA act as PPARa ligands. We 283also showed that 9-oxo-OTA increased the mRNA expression of Cpt1a, Acs, Hmgcs2, 284and Cd36, which are PPAR $\alpha$  target genes that are involved in fatty acid metabolism 285[12-14, 31-33, 35-36]. These effects of 9-oxo-OTA on the expression of PPARa target genes were not observed in PPARa-KO primary hepatocytes. These findings suggested 286287that 9-oxo-OTA promotes fatty acid metabolism in murine hepatocytes via a 288PPAR $\alpha$ -dependent pathway. However, the expression of Cd36 was trend toward significance in PPARa-KO primary hepatocytes treated with 9-oxo-OTA. We confirmed 289290that 9-oxo-OTA also activates PPAR $\gamma$  in luciferase ligand assay. It is well known that 291the expression of Cd36 is induced by not only PPAR $\alpha$  but also PPAR $\gamma$  activation [34-36]. Therefore, there is the possibility that the effect of 9-oxo-OTA on PPAR $\gamma$ 292293 activation contributes to trend toward increment of Cd36 expression.

Our present study also showed that the effect of 9-oxo-OTA on fatty acid oxidation occurs through alteration of not only mRNA expression levels but also metabolism. Fatty acids are taken up by hepatocytes primarily via CD36 [32]. We showed that fatty acid uptake was increased by 9-oxo-OTA treatment. This effect was not observed in PPAR $\alpha$ -KO primary hepatocytes, suggesting that 9-oxo-OTA promotes fatty acid uptake via PPAR $\alpha$  activation. We also showed that the mRNA expression level of *Hmgcs2* (3-hydroxy-3-methylglutaryl-CoA synthase 2) was increased by 9-oxo-OTA treatment.

301 *Hmgcs2* is the key enzyme involved in ketone body formation [33].  $\beta$ -Hydroxybutyrate 302 is a major component of ketone bodies and a marker of fatty acid oxidation. Therefore, we hypothesized that 9-oxo-OTA also promotes the production of  $\beta$ -hydroxybutyrate in 303 murine hepatocytes. The LC-MS analysis revealed that β-hydroxybutyrate secretion 304 305 from murine primary hepatocytes was increased by 9-oxo-OTA treatment. In the RI experiment using <sup>14</sup>C-palmitic acid, we showed that <sup>14</sup>CO<sub>2</sub> production was increased by 306 9-oxo-OTA treatment. The increment of  ${}^{14}CO_2$  production and  $\beta$ -hydroxybutyrate 307 308 secretion suggest that 9-oxo-OTA promoted lipid metabolism in wild-type murine hepatocytes. In contrast, 9-oxo-OTA treatment decreased <sup>14</sup>CO<sub>2</sub> production and 309 β-hydroxybutyrate secretion in PPARα-KO primary hepatocytes. Our present study is 310 311 focused on the effect of 9-oxo-OTA on PPARa activation and its downstream targets. 312But, there is the possibility of the effect of 9-oxo-OTA on other pathways including 313 PPARγ activation. The previous study showed that acetoacetyl-CoA synthetase (AACS) 314is a PPARy target gene [37]. AACS is a ketone body-specific ligase and might play an 315important role in the provision of acetyl-CoA for lipogenesis [38]. 9-oxo-OTA activates both PPARa and PPARy. In the PPARa KO hepatocytes treated with 9-oxo-OTA, there 316317 is the possibility that fatty acid oxidation via PPAR $\alpha$  dependent pathway disappeared and lipogenesis proceed via PPAR $\gamma$  dependent pathway. In this study, we showed that 318 319 9-oxo-OTA promotes fatty acid metabolism via PPARa activation and increases the expression of PPARa target genes in murine hepatocytes. Further examination is 320 321necessary to determine the effect of 9-oxo-OTA on human metabolism.

In conclusion, 9-oxo-OTA contained in tomato fruit activates PPARα and induces the mRNA expression of PPARα target genes. This promotes fatty acid oxidation in murine hepatocytes. These findings provide the first evidence that 9-oxo-OTA induces fatty acid metabolism through a PPARα-dependent pathway and may contribute to the
development of improved treatments for dyslipidemia patients. Our findings suggest
that 9-oxo-OTA might be a valuable, food-derived compound for maintaining health.

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# 448 **Figure legends**

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449 Fig. 1. Oxo-OTAs, Oxo-ODAs and their precursors (fatty acids).
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#### 451 Fig. 2. Identification of 9-oxo-10(E), 12(Z), 15(Z)-octadecatrienoic acid (9-oxo-OTA)

452 by LC-MS. (A) The structure of 9-oxo-OTA. (B) Extracted ion chromatogram (m/z =

291.20) data and (C) full mass data for 9-oxo-OTA in an authentic sample and a tomato
extract sample. (D) MS/MS data for 9-oxo-OTA in an authentic sample and a tomato
extract sample.

456

Fig. 3. 9-oxo-OTA activated PPARa. (A) Effect of 9-oxo-OTA and 13-oxo-OTA 457 $(30\mu M respectively)$  on PPAR $\alpha$  activity with a luciferase reporter assay. (B) Effect of 4589-oxo-OTA and 13-oxo-OTA (10µM respectively) on PPARy activity with a luciferase 459reporter assay. (C) Effect of 9-oxo-OTA on PPARa activity in a dose-dependent manner. 460 461 (D) Evaluation of PPAR $\alpha$  activity with a luciferase reporter assay in cells treated with 462 9-oxo-OTA (30 µM) and/or GW6471 (PPARa antagonist, 100 nM). Data are presented as mean  $\pm$  SEM (n = 5). Symbol for p < 0.05 between the means of the 3 groups. OTA; 463 464 9-oxo-OTA, GW; GW6471.

465

466 Fig. 4. 9-oxo-OTA activated the expression of PPARα target genes in murine 467 hepatocytes. Effect of 9-oxo-OTA (30 μM) on (A) *PPARα* expression levels in murine 468 primary hepatocytes. Effect of 9-oxo-OTA (30 μM) on (B) *Cpt1a*, (C) *Acs*, (D) *Hmgcs2*, 469 and (E) *Cd36* expression in wild-type (WT) or PPARα<sup>-/-</sup> (KO) primary hepatocytes. 470 Data are presented as mean  $\pm$  SEM (n = 5–6). \*\**p* < 0.01, \*\*\**p* < 0.001 vs. control.

- 472 Fig. 5. 9-oxo-OTA promoted fatty acid metabolism in murine hepatocytes. (A) The
- 473 uptake of <sup>14</sup>C-palmitic acid and (**B**) <sup>14</sup>CO<sub>2</sub> production in murine WT or KO primary
- 474 hepatocytes treated with 9-oxo-OTA (30  $\mu$ M). (C) The secretion of β-hydroxybutyrate
- 475 from murine WT or KO primary hepatocytes treated with 9-oxo-OTA (30 μM). Data are
- 476 presented as mean  $\pm$  SEM (n = 3–5). \*p < 0.05, \*\*p < 0.01 vs. control.



Figure 1. Takahashi H, et al.



Figure 2. Takahashi H, et al.



Figure 3. Takahashi H, et al.











Figure 4. Takahashi H, et al.







Figure 5. Takahashi H, et al.

gene	forward primer	reverse primer
Ppara	TCGCGTACGGCAATGGCTTT	TCTTCATCCCCAAGCGTAGGAGG
Cpt1a	CTCAGTGGGAGCGACTCTTCA	GGCCTCTGTGGTACACGACAA
Acs	ACATCCACGTGTATGAGTTCTACGC	AGTAGACGAAGTTCTCACGGTCGAT
Hmgcs2	AATCAGTGGAAGCAAGCTGGA	GTCCAGGGAGGCCTTCAAAA
Cd36	GATGTGGAACCCATAACTGGATTCAC	GGTCCCAGTCTATTTAGCCACAGT
36B4	TCCTTCTTCCAGGCTTTGGG	GACACCCTCCAGAAAGCGAG

Table 1. Oligonucleotide primers used for mRNA analysis.

tissue	liquid nitrogen	heating
gelatinous tissue	LLOQ	0.46±0.10
sarcocarp tissue	LLOQ	$0.23 \pm 0.05$
peel tissue	LLOQ	$0.62 \pm 0.12$

Table 2. Quantitative analysis of 9-oxo-OTA in tomato fruit homogenate under liquid nitrogen or heating (at  $37^{\circ}$ C for 30 min;  $\mu$ g/g fresh weight).

LLOQ: low limit of quantification. Data are presented as mean  $\pm$  SEM (n = 3).