Saturated phosphatidic acids mediate saturated fatty acid–induced vascular calcification and lipotoxicity

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Recent evidence indicates that saturated fatty acid–induced (SFA-induced) lipotoxicity contributes to the pathogenesis of cardiovascular and metabolic diseases; however, the molecular mechanisms that underlie SFA-induced lipotoxicity remain unclear. Here, we have shown that repression of stearoyl-CoA desaturase (SCD) enzymes, which regulate the intracellular balance of SFAs and unsaturated FAs, and the subsequent accumulation of SFAs in vascular smooth muscle cells (VSMCs), are characteristic events in the development of vascular calcification. We evaluated whether SMC-specific inhibition of SCD and the resulting SFA accumulation plays a causative role in the pathogenesis of vascular calcification and generated mice with SMC-specific deletion of both Scd1 and Scd2. Mice lacking both Scd1 and Scd2 in SMCs displayed severe vascular calcification with increased ER stress. Moreover, we employed shRNA library screening and radiolabeling approaches, as well as in vitro and in vivo lipodisomic analysis, and determined that fully saturated phosphatidic acids such as 1,2-diesteroyl-PA (18:0/18:0-PA) mediate SFA-induced lipotoxicity and vascular calcification. Together, these results identify a key lipogenic pathway in SMCs that mediates vascular calcification.

Introduction
Vascular calcification is a major complication in patients who are aging, have diabetes, or have chronic kidney disease (CKD) and is an active process that differentiates vascular smooth muscle cells (VSMCs) into osteoblast-like cells (1, 2). This process is highly regulated by transcription factors involved in osteogenic differentiation, such as RUNX2, MSX2, and ATF4 (3–5). Several in vitro and in vivo studies have shown lipids to play a causative role in the pathogenesis of vascular calcification in addition to inorganic phosphate, inflammatory cytokines, and oxidative stress (6–13). Treatment with unsaturated fatty acids (UFAs) inhibits vascular mineralization and osteogenic differentiation (14, 15), whereas oxidized lipids, such as oxysterols and oxidized phospholipids, elicit procalcific effects in vascular cells (9, 16, 17). In addition to this evidence, we also previously reported that saturated fatty acids (SFAs) and calcium simultaneously accumulate during osteogenic differentiation of vascular cells (18, 19).

Ectopic accumulation of excess lipids, called lipotoxicity, plays a central role in the pathogenesis of cardio-metabolic diseases, including diabetes, obesity, atherosclerosis, and vascular calcification (20–24). However, evidence from a number of experimental systems is emerging, stating that SFAs and UFAs have distinct contributions to lipotoxicity (25, 26). SFAs such as palmitic acid (16:0) and stearic acid (18:0) induce apoptosis, oxidative stress, and ER stress in a variety of mammalian cell lines (including hepatocytes, macrophages, and VSMCs), whereas UFAs such as oleic acid have no or minimal lipotoxic properties (5, 25–30). In addition, cotreatment with UFAs blocks SFA-mediated lipotoxic effects (25, 29). However, the specific metabolite of SFAs that induces lipotoxicity, the mechanism underlying SFA-mediated lipotoxicity, and how UFAs block SFA-mediated lipotoxicity are largely unknown.

Proper intracellular SFA and UFA balance is controlled by a lipogenic enzyme called stearoyl-CoA desaturase (SCD) (31, 32). SCD catalyzes the conversion of SFAs to monounsaturated FAs, mainly 16:0 into palmitoleate (16:1n-7), and 18:0 into oleate (18:1n-9) (33, 34). This introduction of a double bond markedly impacts several chemical properties, including a decrease in melting point and an increase in solubility. The activation of SCD therefore neutralizes SFA-mediated lipotoxicity (5, 25). The expression of SCD is highly regulated by a number of hormonal and dietary factors (33, 35–37). Recently, we found that the accumulation of SFAs by either supplementation with exogenous SFAs or inhibition of SCD induces mineralization of VSMCs (5, 19). In addition, SFA-mediated lipotoxicity and vascular calcification were completely blocked by an acyl-CoA synthetase inhibitor and were attenuated by the shRNA-mediated inhibition of fatty acid elongase-6 (Elov6) (5), suggesting that stearoyl-CoA (18:0-CoA) or its metabolite contributes to mineralization of VSMCs. The majority of 18:0-CoA is utilized as substrate for acyltransferases, such as glycerol-3-phosphate acyltransferase (GPAT) and 1-acylglycerol acyltransferase (AGPAT), and incorporated into phospholipid and neutral lipid fractions. However, which metabolite of 18:0-CoA mediates the lipotoxic and procalcific effects has not been identified.
In this study, we examined whether (i) accumulation of SFAs in VSMCs through SCD reduction is a common feature in vascular calcification and (ii) whether the inhibition of SCD specifically in SMCs causes vascular calcification in vivo. Finally, we identified SFA-derived metabolites that are responsible for lipotoxicity and vascular calcification.

**Results**

**Reduced SCD activity is common in VSMCs of murine models of vascular calcification.** The reduction of klotho, an aging-related protein, is considered to be a major reason that patients with CKD have a markedly higher prevalence of vascular calcification (38, 39). We previously reported that accumulation of SFAs, particularly 18:0, through SCD inhibition induced vascular calcification in vitro (5). To determine whether the inhibition of SCD induced by klotho deficiency is a major event in vascular calcification, we used immunomagnetic cell sorting to isolate VSMCs from klotho-deficient (Klotho−/−) mice with the SMHC-GFP gene (SMMHCGFP; Klotho−/−) and WT mice (SMMHCGFP; Klotho+/+). As expected, Klotho−/− mice developed severe hyperphosphatemia, high fibroblast growth factor 23 (FGF23) circulating levels, and aortic medial calcification (Supplemental Figure 1, A–C; supplemental material available online with this article; doi:10.1172/JCI82871DS1). Quantitative PCR (qPCR) analysis showed that VSMCs highly expressed Scd1 and Scd2 isoforms, but both of these isoforms were significantly reduced in Klotho−/− mice compared with WT controls (Figure 1A). Scd1 and Scd2 protein levels were also diminished in Klotho−/− mice (Figure 1B). Consistently, A9 desaturase activity was significantly reduced in Klotho−/− mice (Figure 1C). In addition, desaturation index (16:1/16:0 and 18:1/18:0) levels decreased in the aortic medial layers of Klotho−/− mice (Figure 1D). Since excess calcitriol and inorganic phosphate are major contributors to vascular calcification induced by klotho deficiency (40, 41), we examined whether calcitriol, phosphate, and their combination affects SCD expression in human VSMCs. Two SCD isoforms (SCD1 and SCD2) are present in the human genome (42). Human VSMCs express SCD but not SCD5 protein (data not shown). Treatment of VSMCs with high phosphate (2.0 mM) and calcitriol reduced levels of SCD mRNA by 59% and 33%, respectively (Figure 1E). Moreover, a combination of high phosphate plus calcitriol reduced SCD mRNA, activity, and protein by over 80% (Figure 1, E–H).

We also examined whether aortic SCD activity and expression are reduced in another murine model of vascular calcification, DBA/2J mice with CKD (12). DBA/2J mice with 5/6 nephrectomies developed increased levels of serum creatinine, phosphorus, FGF23, and severe medial calcification consistent with a CKD phenotype. CKD DBA/2J mice had significantly lower aortic Scd1 and Scd2 expression compared with DBA2J with sham operations (normal kidney function, Supplemental Figure 2, A–G). CKD reduced the activity of SCD by 88% (Supplemental Figure 2H). We previously reported that a 50% reduction of SCD activity is sufficient to induce mineralization of VSMCs (5). Thus, we concluded that aortic 18:0 accumulation by a reduction of SCD1 and SCD2 isoforms is a significant event in the development of vascular calcification. In support of the results from the animal studies, human subjects with stage 3 and 4 CKD had higher levels of serum 18:0 but not oleic acid, as well as a consequent lower desaturation index that is reflective of SCD activity compared with subjects with normal kidney function, suggesting that CKD reduces SCD activity in humans (Supplemental Figure 2, I–K).

**Dual inhibition of SCD1 and SCD2 induces mineralization, osteogenic differentiation, and ER stress of VSMCs in vitro and in vivo.** We previously reported that treatment of VSMCs with an SCD pan inhibitor (SCDi), CAY10566, induced mineralization and osteogenic differentiation. However, which SCD isoform contributes to vascular calcification has not been determined. To determine which SCD isoform is more important in mediating vascular calcification, SCD1 and SCD2 isoforms were inhibited in VSMCs using shRNAs. Simultaneous inhibition of Scd1 and Scd2, but not Scd1 or Scd2 single inhibition, induced mineralization and alkaline phosphatase (ALP) activity (a marker for osteogenic differentiation) of VSMCs. In this study, we used the mouse VSMC, MOVAS-1, line because other VSMC lines — such as human primary VSMCs, human VSMC SM2, mouse and rat primary VSMCs, and bovine calcifying vascular cells — were either not resistant to or morphologically altered in the presence of lentiviral shRNA and puromycin selection.

**Scd1 and Scd2 shRNA efficiently reduced levels of Scd1 and Scd2 mRNA (Supplemental Figure 3A).** Interestingly, simultaneous inhibition of Scd1 and Scd2 was required for inducing mineralization and osteogenic differentiation of VSMCs (Supplemental Figure 3, B–D). We previously reported that 18:0-induced ER stress contributed to vascular calcification (5). Scd1/2 dual inhibition, but not Scd1 or Scd2 single inhibition, drastically induced levels of phospho-protein kinase R-like endoplasmic reticulum kinase (p-PERK), CCAAT-enhancer-binding protein homologous protein (CHOP), spliced X-box binding protein (sXBP-1), and activating transcription factor-4 (ATF4) protein and mRNA, indicating that Scd1/2 dual inhibition is required for causing ER stress (Supplemental Figure 3, E and F). We also found that Scd1/2 dual inhibition induced the production of inflammatory cytokines in VSMCs. These data suggest that dual inhibition of Scd1 and Scd2, rather than the inhibition of a single Scd isoform, contributes to mineralization of VSMCs, as well as the activation of ER stress.

To examine whether ER stress directly contributes to vascular calcification, human VSMCs were treated with 2 chemicals that specifically induce ER stress, tunicamycin (a specific inhibitor of N-linked glycosylation) and thapsigargin (a noncompetitive inhibitor of the sarco/ER Ca2+ ATPase). Both ER stress inducers significantly induced mineralization of VSMCs (Supplemental Figure 3G). We also examined which of the 3 ER stress pathways contributes to SCD inhibition–induced mineralization. As shown in Supplemental Figure 3, H–J, Atf6, Ire1, and Perk shRNAs reduced the expression of their target genes by 52%, 81%, and 68%, respectively. In addition to Atf4 and Chop knockout (5), Atf6, Ire1, and Perk knockdown attenuated SCD inhibition–induced mineralization (Supplemental Figure 3, K–M). In addition, Atf6 and Perk knockdowns reduced mineralization even in the absence of SCDi (Supplemental Figure 3, K–M). These results suggest that all 3 ER stress response pathways synergistically induce vascular calcification in response to SCD inhibition.

Since our in vitro studies indicated that dual inhibition of Scd1 and Scd2 was required for inducing vascular calcification,
we next determined the role of SCD1 and SCD2 dual inhibition in the pathogenesis of vascular calcification in vivo. Since global Scd1 and Scd2 double KO mice are embryonic lethal, we generated SMC-specific Scd1 and Scd2 dual inducible KO (SMC-Scd1/2 KO) mice by crossing Scd1/2 double floxed mice with SMMHC-CreER(T2) mice. Only males were used for the studies because the SMMHC-CreER(T2) transgene was inserted onto the Y chromosome. The SMC-specific Scd1/2 dual conditional mice were i.p. injected with tamoxifen or vehicle to generate SMC-specific Scd1/2 dual inducible KO (SMC-Scd1/2 KO) and control mice. There were no differences in body weight, serum cholesterol, serum triacylglycerol (TAG), serum calcium, serum creatinine, and serum phosphorus between SMC-Scd1/2 KO mice and control mice (data not shown).

SMC-specific Scd1/2 KO mice displayed severe vascular calcification in both aortic sinus and arch lesions, whereas vascular calcification was not observed in the control mice (Figure 3, A–D). Atherosclerotic lesions were not found in either the control or SMC-Scd1/2 KO mice (data not shown). Consistent with histological analysis, aortic calcium content was significantly (>4.8-fold) higher in SMC-Scd1/2 KO mice (Figure 3D). In support of these data, we found that SMC-Scd1/2 deficiency increased mRNA levels of osteogenic makers such as Alp, osteocalcin (Ocn), osteopontin (Opn), and type III phosphate transporter-1 (Pit1), and reduced SMC markers such as Sm22 and Smmhc (Figure 3E). Along with the induction of vascular calcification, SMC-specific Scd deficiency also induced ER stress in the VSMCs of SMC-Scd1/2 KO mice (Figure 3, E–G). qPCR and immunoblot analysis revealed that mRNA and protein levels of ER stress markers such as Atf4, Chop, sXbp-1, Gadd34, and Atf3 were significantly increased in the VSMCs of SMC-Scd1/2 KO mice (Figure 3, E and F). We previously demonstrated that CHOP-induced cell apoptosis contributes to the pathogenesis of vascular calcification (9). Immunofluorescence analysis showed that apoptotic cell death and CHOP expression is significantly higher in the aortic sinuses of SMC-Scd1/2 KO mice compared with control mice (Figure 3, G–I).
of SCD, results in ER stress and vascular calcification (5). In addition, our previous studies indicated that SCD inhibition and 18:0-mediated ER stress and vascular calcification were completely abolished in the presence of Triascin C, a potent pan inhibitor of acyl-CoA synthetase (5). 18:0 must be converted to 18:0-CoA to elicit lipotoxic effects. Since over 70% of generated 18:0-CoA from VSMCs treated with SCDi and Scd1/2 shRNA is incorporated to acyl-lipid fractions as a substrate for acyltransferases, we hypothesized that an acyl-lipid containing 18:0 mediates SFA-induced ER stress and vascular calcification. To identify a 18:0 metabolite responsible for lipotoxicity and vascular calcification, we used several inhibitors or activators such as ACAT (Sandoz 58-035), ceramide synthase (Fumonisin B1), fatty acid oxidation (etomoxir), and a DGAT1 inhibitor (A-92250) to block 18:0 metabolism. Although Triascin C completely inhibited 18:0-induced calcification, none of these inhibitors blocked 18:0-mediated vascular calcification (Figure 4A). In addition, Gpat4, Agpat3, and Agpat5 knockdowns significantly induced mineralization in control VSMCs containing empty shRNA, whereas Gpat4, Agpat3, and Agpat5 knockdowns were the only acyltransferase knockdowns that prevented SFA-induced vascular calcification (Figure 4A).

Transient knockdowns of acyltransferases in human cells using siRNA and shRNA were also performed. However, the efficiency of knockdown was more effective and consistent using stable cell lines than transient knockdowns. Since VSMCs are difficult to transfect with plasmids and oligos, the knockdown efficiency was only 17%–55% and there were considerable variations in the knockdown efficiencies between genes. To induce vascular calcification and ER stress, we treated all acyltransferase knockdown VSMCs with 200 μM 18:0. Treatment with 18:0 significantly reduced mineralization in control VSMCs expressing empty shRNA, whereas Gpat4, Agpat3, and Agpat5 knockdowns induced mineralization in control VSMCs containing empty shRNA. We further examined the contribution of Gpat4, Agpat3, and Agpat5 in SCD inhibition–mediated vascular calcification and lipotoxicity. While Gpat4 and Gpat1 knockdowns comparably reduced Gpat activity in VSMCs (Supplemental Figure 5, A and B), Gpat4 but not Gpat1 knockdown attenuated mineralization of VSMC in both the presence and absence of 18:0 and SCDi (Supplemental Figure 5, C–E). Since Gpat4 knockdown reduced Ocn and Opn expression, 18:0-mediated induction of osteogenic differentiation was reduced in Gpat4 but not Gpat1 knockdown cells (Supplemental Figure 5, F and G). In addition, Gpat4 knockdown reduced the expression of Pit1, which contributes to vascular calcification, resulting in the reduction of phosphate uptake (Supplemental Figure 5, H and I). SCD inhibition– and 18:0-mediated ER stress was attenuated in Gpat4 but not Gpat1 knockdown cells (Supplemental Figure 5, J–N). Treatment with 18:0 and SCDi increased levels of ATF4 and CHOP protein and mRNA in control cells, whereas Gpat4 knockdown significantly attenuated the induction of ATF4,
and Ocn, as well as Pit1, were reduced in Agpat3 and Agpat5 knockdown cells treated with SCDi (Supplemental Figure 7, C–H). In addition, Agpat3 knockdown completely reduced SCDi inhibition–mediated induction of ATF4 and CHOP proteins (Supplemental Figure 7I). Levels of ATF4 and CHOP proteins were also substantially lower in Agpat5 knockdown cells treated with SCDi, although Agpat3 knockdown was more effective than Agpat5 knockdown (Supplemental Figure 7J).

GPAT4, AGPAT3, and AGPAT5 catalyze the initial 2 steps of de novo glycerolipid biosynthesis and generate phosphatidic acid (PA) in the ER (Figure 5A and ref. 43). In addition, the inhibition of acyl-CoA:cholesterol acyltransferase-1 (ACAT1) and ACAT2 responsible for cholesteryl ester (CE) synthesis, diacylglycerol acyltransferase-1 (DGAT1) and DGAT2 responsible for triacylglycerol synthesis, monoacylglycerol acyltransferase-1 (MOGAT1) responsible for diacylglycerol (DAG) synthesis, and serine palmitoyltransferase (SPTLC1) responsible for sphingolipid and ceramide synthesis...
did not affect 18:0-induced vascular calcification and lipotoxicity. These results suggest that PA or its downstream metabolite, glycerophospholipid containing 18:0, contributes to 18:0-induced lipotoxicity. These results suggest that PA or its downstream metabolite, glycerophospholipid containing 18:0, contributes to 18:0-induced lipotoxicity. These results suggest that PA or its downstream metabolite, glycerophospholipid containing 18:0, contributes to 18:0-induced lipotoxicity. These results suggest that PA or its downstream metabolite, glycerophospholipid containing 18:0, contributes to 18:0-induced lipotoxicity. These results suggest that PA or its downstream metabolite, glycerophospholipid containing 18:0, contributes to 18:0-induced lipotoxicity.

To determine whether PA or glycerophospholipid contributes to lipotoxicity, we next modified Lipin, a PA phosphatase (PAP) enzyme that participates in the third step of de novo glycerolipid synthesis and catalyzes the conversion of PA to DAG in the ER. Lipin2 is a major isoform expressed in VSMCs (Supplemental Figure 8A). Treatment with Lipin2 shRNA reduced mRNA levels of Lipin2 and PAP activity by 78% and 46%, respectively (Supplemental Figure 8, B and C). The remaining activity is likely derived from Lipin1 and PAP2 enzymes. In Lipin2 knockdown cells, the effects of 18:0 and SCD inhibition on mineralization and osteogenic differentiation were significantly enhanced compared with the control cells (Supplemental Figure 8, D–G). In addition, Lipin2 knockdown augmented ER stress (Supplemental Figure 8, H–K). These results suggest that PA contributes to SFA-induced mineralization and lipotoxicity.

SFAs such as 18:0 and 16:0 preferentially accumulate in PA. Based on the acyltransferase and Lipin shRNA screening, PA is a potential candidate for mediating SFA-induced lipotoxicity. We therefore hypothesized that 18:0 is preferentially incorporated into the PA fraction. Since control human VSMCs converted over 50% of 18:0 and 16:0 into MUFAs, and because VSMCs treated with SCDi had a complete inhibition on the conversion of SFAs to MUFAs (Figure 5, B and C), SFA-specific lipid partitioning is determined only in VSMCs treated with SCDi. Under SCD inhibition, 18:0 and 16:0 are preferentially incorporated into the PA fraction in VSMCs (Figure 5, D–F). The incorporation of 18:0 and 16:0 into the PA fraction was increased by >15-fold and >9-fold, respectively. Upon SCD inhibition, SFA distribution into lysophosphatidylcholine (LPC), phosphatidylserine (PS), and lysoPA (LPA) also increased, whereas distribution into phosphatidylcholine (PC) and sphingomyelin (SM) decreased (Figure 5, E and F). In addition, SCD inhibition did not affect the lipid distribution of 18:1n-9 (Supplemental Figure 9, A and B). Consistently, 18:0 distribution in Scd1/2 knockout MOVAS-1 cells was nearly identical to the distribution in VSMCs treated with SCDi (data not shown).

Fully saturated PAs such as 1,2-distearoyl PA (18:0/18:0-PA) and 1,2-dipalmitoyl PA (16:0/16:0-PA) accumulate in SCD knockdown VSMCs and the aortic medial layers of SMC-Scd1/2 KO mice. We next performed an LC-MS–based lipidomic quantitative analysis to determine which lipids are increased in human VSMCs in response to SCD inhibition in vitro and in vivo. According to the analysis, over 270 lipid species were detected in VSMCs (Supplemental Table 3). Similar to the results of the “4C:18:0” distribution study, in vitro SCD inhibition most drastically increased absolute levels of PA by 23-fold compared with the control, followed by LPC and DAG levels (Figure 5G). SCD inhibition increased all fully saturated PAs (PA containing 2 SFAs in both the sn-1 and sn-2 position) such as 16:0/18:0-PA and 18:0/18:0-PA, myristostearoyl-PA (14:0/18:0-PA), and 16:0/16:0-PA. Both 16:0/18:0-PA and 18:0/18:0-PA were most drastically increased by over 200-fold in VSMCs treated with SCDi. Levels of fully unsaturated PAs such as dioleoyl (18:1/18:1) and partially unsaturated PAs such as palmitoyloleoyl (16:0/18:1) were not altered in VSMCs treated with SCDi (Figure 5H).

Similar to the in vitro results, in vivo quantitative lipidomic analysis was able to detect over 250 lipid species (Supplemental Table 4). Since cell-sorted–VSMCs were insufficient for the lipidomic analysis, the lipid contents in the medial layer of aortas were compared between SMC-Scd1/2 KO and control mice. The in vivo lipidomic analysis revealed that levels of PA significantly increased in the medial layer of aortas of SMC-Scd1/2 KO mice compared with control mice (Figure 5I). However, levels of phosphatidylcholine, CE, and TAG were reduced in SMC-Scd1/2 KO mice (Figure 5I). SMC-Scd1/2 KO deficiency increased levels of fully saturated PAs such as 16:0/16:0-PA and 18:0/18:0-PA by 11-fold and 63-fold, respectively, whereas levels of partially...
and fully unsaturated PAs were not changed (Figure 5J). Similar to SMC-Scd1/2 KO mice, CKD DBA/2J mice had significantly higher levels of fully saturated PAs, such as 16:0/16:0-PA, 16:0/18:0-PA, and 18:0/18:0-PA (Supplemental Figure 9C), compared with sham-operated DBA/2J mice.

We previously found that 18:0 preferentially accumulates in the ER, which is highly associated with ER stress (5). We therefore hypothesized that SFAs, such as 18:0, accumulate in the form of PAs in the ER. To determine our hypothesis, we isolated the ER fraction from VSMCs treated with SCDi and analyzed levels of PA. LC-MS/MS analysis showed that fully saturated PAs such as 16:0/16:0-PA and 18:0/18:0-PA accumulate in the ER. Levels of fully saturated PAs in the ER fraction were >10-fold higher than in the whole cell lysates (Supplemental Figure 9D).

UFAs such as 18:1n-9 reduce fully saturated PAs, resulting in a drastic attenuation of ER stress and vascular calcification. Cosupplementation with UFAs such as 18:1n-9 has been shown to rescue SFA-induced lipotoxicity through an unknown mechanism (25). Supplementation with exogenous UFAs such as 18:1n-9, linoleic acid (18:2), and docosahexaenoic acid (22:6) completely blocked SCD inhibition–induced mineralization (Figure 6, A and B) and osteogenic differentiation (Figure 6C). In addition to SFA-induced mineralization, calcitriol-induced mineralization was inhibited by 18:1n-9 supplementation (Figure 6, D and E). UFA treatment completely blocked ER stress induced by SCD inhibition (Figure 6, F–H). We next examined whether UFAs modulate the lipid partitioning of SFAs. Borate-coated TLC analysis showed that UFA cotreatment completely rescued the accumulation of 14C-18:0 into the PA fraction by SCD inhibition (Figure 6, I and J). In addition, quantitative LC-MS analysis showed that supplementation with 18:1n-9 completely normalized levels of fully saturated PAs such as 18:0/18:0-PA and 16:0/16:0-PA increased by SCD inhibition (Figure 6K).

Fully saturated PAs are disfavor substrates for LIPIN2, which contributes to its accumulation in SCD1/2-inhibited VSMCs. We
as a recombinant LIPIIN2 protein, whose activity was completely inhibited by N-methylmaleimide. The conversions of PAs to DAGs by the recombinant LIPIN2 were detected by LC-MS/MS. We used 16:0/16:0-PA and 18:0/18:0-PA as fully saturated PAs, 16:0/18:1-PA and 18:0/18:1-PA as partially unsaturated PAs, and 18:1/18:1-PA and 18:2/18:2-PA as fully unsaturated PAs. As shown in Figure 7A, LIPIN2 showed preferential specificity for fully unsaturated PAs (18:1/18:1-PA and 18:2/18:2-PA), followed next by partially unsaturated PAs (18:0/18:1-PA and 16:0/18:1-PA). Fully saturated PAs, 16:0/16:0-PA and 18:0/18:0-PA, were disfavor substrates for LIPIN2 and displayed only 1% of Lipin2 activity when compared with 18:1/18:1-PA (Figure 7A).

Fully saturated PAs such as 18:0/18:0-PA, but not unsaturated PAs or partially unsaturated PAs, induced vascular calcification and lipotoxicity. To obtain direct evidence that fully saturated PAs determined a mechanism by which SCD inhibition causes massive accumulation of fully saturated PAs in VSMCs, and UFA supplementation rescues the accumulation of fully saturated PAs. SCD inhibition and 18:0 treatment did not affect the expression and the activity of GPAT, AGPAT, and Lipin in VSMCs (data not shown). It has already been reported that GPAT4, AGPAT3, and AGPAT5 comparably utilize both saturated fatty acyl-CoAs such as 18:0-CoA and unsaturated acyl-CoAs such as 18:1-CoA. AGPAT3 and AGPAT5 also use both saturated and unsaturated LPA. We therefore hypothesized that LIPIIN2 uses unsaturated PAs more efficiently than fully saturated PAs, which results in the specific accumulation of fully saturated PAs such as 18:0/18:0-PA and 16:0/16:0-PA in Scd1/2 knockdown VSMCs. To determine the substrate preference of LIPIIN2, the lysate of HEK-293 cells infected with adenoviruses expressing human LIPIIN2 was used as a recombinant LIPIIN2 protein, whose activity was completely inhibited by N-methylmaleimide. The conversions of PAs to DAGs by the recombinant LIPIIN2 were detected by LC-MS/MS. We used 16:0/16:0-PA and 18:0/18:0-PA as fully saturated PAs, 16:0/18:1-PA and 18:0/18:1-PA as partially unsaturated PAs, and 18:1/18:1-PA and 18:2/18:2-PA as fully unsaturated PAs. As shown in Figure 7A, LIPIIN2 showed preferential specificity for fully unsaturated PAs (18:1/18:1-PA and 18:2/18:2-PA), followed next by partially unsaturated PAs (18:0/18:1-PA and 16:0/18:1-PA). Fully saturated PAs, 16:0/16:0-PA and 18:0/18:0-PA, were disfavor substrates for LIPIIN2 and displayed only 1% of Lipin2 activity when compared with 18:1/18:1-PA (Figure 7A).
induce vascular calcification and lipotoxicity, human VSMCs were treated with 18:0/18:0-PA and 16:0/16:0-PA as fully saturated PAs, and 18:1/18:1-PA and dilinoleyl-PA (18:2/18:2-PA) as a fully unsaturated PA. These PAs showed distinct effects on mineralization, osteogenic differentiation, ER stress, and lipotoxicity in VSMCs. As shown in Figure 7, B and C, the fully saturated PAs 18:0/18:0-PA and 16:0/16:0-PA significantly increased calcium content. In particular, 18:0/18:0-PA was very potent at inducing mineralization (Figure 7B). It increased calcium content as low as 2.5 μM in a dose-dependent manner (Figure 7C). At a 10 μM concentration, 18:0/18:0-PA increased calcium content by over 5-fold compared with the vehicle. The fully unsaturated PAs 18:1/18:1-PA and 18:2/18:2-PA and the partially unsaturated PAs 18:0/18:1-PA and 16:0/18:1-PA had no effect on mineralization (Figure 7D). Consistently, 18:0/18:0-PA but not the partially saturated PAs 18:0/18:1-PA and 16:0/18:1-PA induced mRNA and protein expressions of osteogenic markers such as \textit{Ocn}, \textit{Opn}, and \textit{Pit1} (Figure 7, E–G), and ER stress makers such as \textit{Atf4} and \textit{Chop} (Figure 7, H–J). Cosupplementation with 18:1/18:1-PA did not affect 18:0/18:0-PA-induced mineralization (data not shown). The data suggest that the inhibitory effect of UFAs on SFA-induced mineralization requires the competition of free UFAs with free SFAs on the acylation to prevent the formation of fully saturated PAs. To
concluded that the accumulation of 18:0 by a significant reduction of SCD by phosphate and calcitriol is a major event in the development of vascular calcification.

In addition, we demonstrated that dual inhibition of Scd1 and Scd2 directly causes vascular calcification, along with massive accumulation of SFAs both in vitro and in vivo. This is direct evidence that SFA-induced lipotoxicity through Scd1 and Scd2 repression in VSMCs plays a causative role in the pathogenesis of vascular calcification. There are several mechanisms by which SFA accumulation induces vascular calcification. Recent studies suggest that ER stress-mediated cell death contributes to the pathogenesis of vascular calcification (9). Atf4 and Chop knockdown attenuates mineralization and osteogenic differentiation of VSMCs in response to SFA treatment and other inductive factors (5, 6, 9). In addition to saturated fat, oxidized lipids and TNFα— which induce ER stress, tunicamycin, and thapsigargin— also induce mineralization of VSMCs. We previously reported that in vivo CHOP deficiency and treatment with chemical chaperones such as 4-phenylbutyric acid and tauroursodeoxycholic acid attenuate CKD-dependent vascular calcification and inhibit cell apoptosis (6, 9). This study provides direct evidence that accumulation of SFAs by SCD inhibition induces vascular CHOP expression upon ER stress response activation, which in turn causes vascular apoptosis, thereby inducing vascular calcification. Our current study provides in vivo evidence that the accumulation of SFAs in VSMCs through SMC-specific SCD deficiency induces ATF4-mediated osteogenesis (5, 48) and CHOP-mediated cell death (9, 49), which synergistically lead to ectopic mineralization and osteogenic differentiation of VSMCs. We also found that, in addition to the PERK arms, the ATF6 and IRE1 arms of the ER stress response contribute to SFA-induced mineralization via unknown mechanisms. Identification of specific mechanisms by which the ATF6 and IRE1 pathways induce vascular calcification will be required in future studies.

In addition to vascular calcification, the ectopic accumulation of SFAs and their lipotoxic effects, including ER stress, oxidative stress, and apoptotic cell death, are implicated in the pathogenesis of other metabolic diseases such as diabetes, obesity, and atherosclerosis (20, 21, 29, 50, 51). However, the molecular mechanism by which an SFA metabolite elicits these lipotoxic effects has not been identified. Our current findings are the first to demonstrate that fully saturated PAs generated through de novo glycerolipid synthesis are the major metabolites responsible for SFA-induced lipotoxicity in VSMCs. The first step to identifying the toxic metabolite was screening inhibitors and the shRNA library to identify 18:0-metabolizing enzymes responsible for lipotoxicity to consolidate potential SFA metabolites. The metabolism of 18:0 mainly includes fatty acid oxidation, desaturation, and acylation. We previously reported that (i) 18:0 most potently induced ER stress in VSMCs, (ii) acyl-CoA formation was required for 18:0-induced ER stress, and (iii) the conversion of 18:0 to 18:1n-9 by SCD was protective for 18:0-induced ER stress (5). Etomoxir, a fatty acid oxidation inhibitor, did not affect SFA-induced lipotoxicity, suggesting that fatty acid oxidation is a minor contributor in SFA-induced lipotoxicity. In addition, over 70% of 18:0-CoA is incorporated into lipid fractions by acyltransferases. We therefore hypothesized that a metabolite derived from SFAs through acylation mediates
the lipotoxicity. Twenty acyltransferases were present in VSMCs. Based on the shRNA screening, we found that, out of these 21 acyltransferases expressed in VSMCs, GPAT4, AGPAT3, and AGPAT5 contribute to SFA-induced mineralization and ER stress. GPAT and AGPAT constitute the first 2 reactions of de novo biosynthesis of glycerolipids and generate PAs by utilizing saturated acyl-CoA, including 18:0-CoA, as a substrate. We next examined whether PA or its downstream metabolite is responsible for SFA-induced lipotoxicity in VSMCs. LIPIN, a PA phosphatase, catalyzes the third step of de novo glycerolipid synthesis by converting PA to DAG. The shRNA-mediated knockdown of Lipin2, which is the major isoform in VSMCs, augmented SFA-induced lipotoxicities, such as mineralization and ER stress, in contrast to Gpat4, Agpat3, and Agpat5 knockdowns. These results suggest that PA is the major metabolite responsible for SFA-induced lipotoxicity. In addition, the evidence from the shRNA library screening, radioactive SFA metabolite responsible for SFA-induced lipotoxicity. In addition to knockdowns. These results suggest that the ATF4-CHOP axis of the ER stress response contributes to mineralization induced by fully saturated PAs.

We also demonstrated the mechanism for the accumulation of SFAs into the PA fraction and the mechanism by which UFAs block SFA-induced lipotoxicity. Lipin2 is a major enzyme that metabolizes PA into DAG in VSMCs. Lipin2 shows a preferential response to fully saturated PAs such as 18:1/18:1-PA and 18:2/18:2-PA compared with fully saturated PAs such as 18:0/18:0-PA and 16:0/16:0-PA. This preference causes accumulation of PA fractions in SCD-inhibited VSMCs. Even when excess SFAs such as 18:1n-9 and 18:2 are treated with VSMCs, they never accumulate as PAs. Fully saturated PAs are therefore metabolized less than unsaturated PAs. UFAs elicit distinct roles in the context of lipotoxicity (22, 25). Cosupplementation with UFAs such as 18:1n-9 and 22:6 acid has been known to block SFA-induced lipotoxicity (25, 29). However, the mechanism by which UFAs attenuate SFA-induced lipotoxicity is poorly understood. Cotreatment with UFAs such as 18:1n-9 and 22:6 acid completely blocks mineralization, osteogenic differentiation, and ER stress of VSMCs induced by SCD inhibition and calcitriol. We also find that UFA cotreatment completely normalizes the preferential distribution of SFAs to the PA fraction and absolute levels of fully saturated PAs in SCD-inhibited VSMCs. The Lipin2 substrate specificity accounts for the mechanism by which UFA cotreatment reduces levels of PA, resulting in the attenuation of SFA-induced lipotoxicity. Partially saturated PAs are better substrates for Lipin2 than fully saturated PAs. In addition, partially saturated PAs such as 18:1/16:0-PA and 18:0/18:1-PA do not show lipotoxic effects in contrast to fully saturated PAs. These results demonstrate that the insertion of at least one molecule of UFA into PA increases Lipin2 activity, thereby preventing the accumulation of fully saturated PAs and their lipotoxicity. SCD inhibition also unexpectedly increases fully saturated DAGs, suggesting that other LIPIN isoforms and PAP2 can efficiently utilize fully saturated PAs. In addition, we find that DGAT2 cannot utilize fully saturated DAGs as a substrate when saturated acyl-CoA is another substrate. The occurrence of lower substrate specificities of LIPIN2 and DGAT2 toward fully saturated PAs and DAGs, respectively, is consistent with Goldman and Vagelos’s hypothesis that at least one UFA is required for triglyceride synthesis (52). This explains why SCD inhibition and excess SFA increase levels of PAs and DAGs, but also reduces levels of TAGs.

The current study suggests that the intracellular location of fully saturated PA synthesis and accumulation is important for SFA-induced lipotoxicity, since GPAT4 but not GPAT1 knockdown inhibits SFA-induced lipotoxicities such as ER stress and mineralization. Although both GPAT1 and GPAT4 catalyze the same reaction that converts glycerol-3-phosphate to 1-lysophosphatidic acid, GPAT1 and GPAT4 are localized in the mitochondria outer membrane and ER, respectively (53, 54). In addition, both isoforms are able to utilize saturated fatty acyl-CoAs, such as 18:0-CoA, as a substrate (55). The accumulation of fully saturated PAs in the ER fraction was more than 10-fold higher than in the whole cell fraction. AGPAT3, AGPAT5, and Lipin2 are also located in the ER (56–58). These data suggest that the ER is a major organelle for synthesizing and accumulating fully saturated PAs, which directly lead to the lipotoxicity, particularly ER stress. However, it is still possible that GPAT1, rather than GPAT4, contributes to other SFA-induced lipotoxicities such as oxidative stress and mitochondrial dysfunction.

In addition to VSMCs, lipotoxic effects induced by SFAs have been observed in a variety of cell types, including cardiomyocytes, fibroblasts, pancreatic β-cells, hepatocytes, and macrophages (25, 28–30, 51, 59). The accumulation of fully saturated PAs by exogenous SFA addition or SCD inhibition is not limited to VSMCs. We detected that hepatocytes and 3T3 fibroblasts accumulate fully saturated PAs in response to SCD inhibition. Interestingly, recent studies have demonstrated that hepatic fully saturated PAs such as 16:0/16:0-PA, but not unsaturated PAs derived from de novo glycerolipid synthesis, inhibit the mTOR complex 2, leading to insulin resistance (60, 61). Therefore, fully saturated PAs may not only play a critical role in vascular calcification, but also in many other cardiometabolic diseases.

In conclusion, our findings reveal that increased fully saturated PAs through de novo glycerolipid synthesis promotes mineralization of VSMCs under procalcific conditions, including CKD. As indicated in Figure 8, we propose that CKD and klotho deficiency reduce SCD expression by increased inorganic phosphate and calcitriol. Reduced SCD activity leads to an increase in SFA levels in VSMCs. SFAs such as 18:0 and 16:0 preferentially accumulate in PA fractions via the reaction of GPAT4 and AGAPT3/5 in the ER. UFAs such as 18:1n-9 compete with SFAs on the reaction of GPAT4 and AGAPT3/5 to inhibit the formation of fully saturated PAs. The accumulated fully saturated PAs cause ER stress, resulting in the induction of the pro-osteogenic factor ATF4 and the proapoptotic factor CHOP. The pro-osteogenic and proapoptotic effects of the 2 ER stress effectors synergistically lead to vascular calcification. In addition to the PERK arm, the IRE1 and ATF6 arms of the ER stress response also contribute to SFA-induced mineralization via a currently unknown mechanism.
Methods

Animals. C57Bl6 Scd1 conditional KO (lox/lox) mice (32), C57Bl6 klotho-deficient mice (39), and C57Bl6 SMMHC-CreER(T2) mice (62) were generated as previously described. C57Bl6 Rosa-tomatoTd reporter (63), C57Bl6 SMHC-Cre/GFP mice (64), and DBA/2J mice were obtained from the Jackson Laboratory. Similar to Scd1lox/lox mice, Scd2lox/lox mice were generated by floxing the exon 3 encoding the catalytic site of this enzyme (Supplemental Figure 10). In brief, DNA pools from a mouse strain 129/SV BAC library (ResGen Inc.) were screened by PCR using a primer pair that amplifies the exon 4 portion of the Scd2 gene. A genomic fragment was isolated from the Scd2-BAC clone and introduced into a plasmid that carries the MCI1-TK selection cassette. A mini-targeting vector was constructed by cloning 4 contiguous PCR fragments: (i) a homology to the region immediately upstream of exon 3, (ii) a region containing a loxP-flanked exon 3, (iii) an FRT-flanked pGK promoter/EM7 promoter-NEO-pGhApA cassette, and (iv) a homology to the region immediately downstream of exon 3. The targeting vector was linearized and introduced by electroporation into murine SV129 RI embryonic stem cells. Karyotypically normal ES clones were micro-injected into C57BL/6 blastocysts to produce chimeric founders.

The targeting vector contains a loxP-flanked exon 3, (iii) an homology to the region immediately downstream of exon 3, (ii) a region that contains a loxP-flanked exon 3, (iii) an FRT-flanked pGK promoter/EM7 promoter-NEO-pGhApA cassette, and (iv) a homology to the region immediately downstream of exon 3. The targeting vector was linearized and introduced by electroporation into murine SV129 RI embryonic stem cells. Karyotypically normal ES clones were micro-injected into C57BL/6 blastocysts to produce chimeric founders at the University of Wisconsin-Madison Biotechnology Center’s Transgenic Animal Facility. The Scd2lox/lox mice were crossed with Rosa-Flp mice and backcrossed 10 times with C57Bl6 mice. Scd1lox/lox and Scd2lox/lox mice were bred to generate Scd1/2lox/lox mice. The resulting Scd1/2lox/lox mice were intercrossed with C57Bl6 SMMHC-CreER(T2) mice to obtain SMMHC-CreER(T2), Scd1lox/lox; Scd2lox/lox mice. Since the SMMHC-CreER(T2) transgene was inserted on the Y chromosome, only males were used in this study. Five-week-old males were i.p. injected with either 1 mg tamoxifen in vegetable oil or the vehicle for 5 consecutive days. The mice treated with tamoxifen and vehicle were used as SMC-specific Scd1/2 KO mice and control mice, respectively. After the injections, the mice were maintained on a special diet (TD10364) for 10 weeks. For cell sorting, SMC-Scd1/2 KO mice were crossed with Rosa26-tomatoTd reporter mice.

Cell cultures. Human aortic VSMCs were obtained from Lonza. MOVAS-1 was provided by Mansoor Husain at the University of Toronto (Toronto, Canada) (5, 6). These VSMCs were cultured in DMEM containing 10% FBS and 2 mM inorganic phosphate in the absence/presence of 200 μM 18:0-BSA complex, 300 nM CAY10566 (Cayman Chemical), 100 nM calcium (Cayman Chemical), 0.1 μM thapsigargin (Fisher Scientific), and various doses of PA (methyl-β-cyclodextrin complex, Avanti Polar Lipids Inc.). Seven days after the treatments, cells were stained with Alizarin red to identify calcified cells. Calcium deposition in the plates and the aortas was quantified as previously described (6, 9, 18).

RNA analysis. qPCR assays were performed using an Applied Biosystems StepOne qPCR instrument. Quantitative expression values were calculated from an absolute standard curve method using the plasmid template (Open Biosystems; GE Healthcare) containing each of the target gene cDNA. Primer sequences that are fully validated were obtained from the Primer Bank (http://pga.mgh.harvard.edu/primerbark; Harvard University, Cambridge, Massachusetts, USA) except for sXBP-1 and Ocn, as shown in Supplemental Table 5.

Determination of 14C-18:0, 14C-16:0, and 14C-18:1n-9 distribution. Isolated aortas were digested to single cells by digestion at 37°C in collagenase buffer (3.2 mg/ml collagenase II, 0.7 mg/ml elastase, 0.2 mg/ml soybean trypsin inhibitor) in Hank’s buffered saline solution. Aortas were harvested under sterile conditions following flushing of the vasculature system with sterile heparinized PBS, and minced prior to the digestion. Single cell suspension was sorted based on GFP and tomatoTd expression for klotho mice and SMC-Scd1/2 KO mice, respectively. Sorting was performed on a MoFlo high-speed cell sorter (Beckman Coulter) at the University of Colorado flow cytometry and sorting core facility.

Immunoblot analysis. Cell and tissue lysates were prepared using RIPA buffer (Cell Signaling Technology). The samples were separated by SDS-PAGE, transferred to a nitrocellulose membrane, and immuno-blotted with the following antibodies: p-PERK (clone 16F8), p-eIF2α (clone 119A11), CHOP (clone D46F1), and ATF4 (clone D4B8) from Cell Signaling Technology; CREB2 (clone C-20), GADD153 (clone B-3), SCDD1 (clone S-18), SCDD2 (clone C-16), and GAPDH (clone V-18) from Santa Cruz Biotechnology Inc.; GPAT4 (clone NB100-2389) and SCDD5 (clone 45800002) from Novus Biologicals; and GPD1 (clone 4613) from ProSci. Samples were visualized using horseradish peroxidase-coupled to appropriate secondary antibodies, with enhancement by an ECL detection kit.

Calcium content in cultured cells and aortas. Calcium deposition in the plates and the aortas was quantified as previously described (6, 9, 18).

Immunomagnetic cell sorting. Isolated aortas were digested to single cells by digestion at 37°C in collagenase buffer (3.2 mg/ml collagenase II, 0.7 mg/ml elastase, 0.2 mg/ml soybean trypsin inhibitor) in Hank’s buffered saline solution. Aortas were harvested under sterile conditions following flushing of the vasculature system with sterile heparinized PBS, and minced prior to the digestion. Single cell suspension was sorted based on GFP and tomatoTd expression for klotho mice and SMC-Scd1/2 KO mice, respectively. Sorting was performed on a MoFlo high-speed cell sorter (Beckman Coulter) at the University of Colorado flow cytometry and sorting core facility.

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 Generation of GPAT4- and LIPIN2-overexpression VSMCs using the Tet-on gene expression system. Human GPAT4 and human LIPIN2 cDNA were purchased from Open Biosystems and cloned into a pLenti-CMV-Zeo vector (Addgene). VSMCs expressing TetR (TetR-VSMCs) were generated by infecting lentiviruses containing TetR and a single colony isolation with blasticidin (5 μg/ml). TetR-VSMCs were infected with lentiviruses containing either human GPAT4 or LIPIN2. Colonies were selected by treatment with 100 μg/ml zeocin for 7 days. To overexpress GPAT4 and LIPIN2, TetR-VSMCs expressing GPAT4 and LIPIN2 were treated with 1 mg/ml doxycycline (Dox) or vehicle for 24 hours and then treated with either 300 nM CAY10566 or 200 μM 18:0.

 Determination of 14C-18:0, 14C-16:0, and 14C-18:1n-9 distribution. Control VSMCs were pretreated with either vehicle or 300 nM SCDDi. After preincubation, VSMCs were treated with 1 or 5 μg C14-stearic acid or 14C-palmitic acid-BSA complex for 6 hours. Cells were harvested with PBS with buthylated hydroxytoluene (50 μg/ml). Total lipids were extracted by Bligh & Dyer method under acidic conditions. The extracted total lipids were subjected to 2.3% boric acid or 14C-palmitic acid–BSA complex for 6 hours. Cells were harvested with PBS with buthylated hydroxytoluene (50 μg/ml). Total lipids were extracted by Bligh & Dyer method under acidic conditions. The extracted total lipids were subjected to 2.3% boric acid–coated TLC. The TLC was developed twice with chloroform/ethanol/water/triethylamine (30:35:7:35), as previously described (66). The developed TLC was autoradiographed overnight. The resolving lipid fractions were scraped off the plate, and radioactivity was measured using a liquid scintillation counter.
**LC-MS lipidomic analysis.** Lipid analysis was performed by LC electron spray ionization-MS/MS using a Shimadzu LC-20AD LC system and an Applied Biosystems 3200 qTrap mass spectrometer with a turbo-ionspray source (350°C). For PA and phosphatidylglycerol (PG), LC was performed on a LC-NH2 50 × 2.1 mm column (Agilent Technologies) as previously described. Solvents C and D consisted of tetrahydrofuran/methanol/water (30:20:50 and 75:20:5, respectively), both containing 10 mM ammonium formate. Columns were heated to 50°C. LPA, lysophosphatidylserine (LPS), lysophosphatidylinositol (LPI), PE, phosphatidylserine (PS), PG, and PI were analyzed with negative ion mode, whereas positive ion mode was maintained for 5 minutes, restored to 100% Solvent A by a 1 minute linear gradient, maintained at 100% Solvent B for 5 minutes, returned to 0% solvent B at 11 minutes, and maintained for 5 minutes. Negative ion mode was used for PA and PG analysis. For other lipid analysis, LC was performed on a Zorbax C18, 1.8 μm, 50 × 2.1 mm column (Agilent Technologies) as previously described. Solvents C and D consisted of tetrahydrofuran/methanol/water (30:20:50 and 75:20:5, respectively), both containing 10 mM ammonium formate. Columns were heated to 50°C. LpA, lyso-phosphatidylserine (LPS), lyso-phosphatidylinositol (LPI), PE, phosphatidylserine (PS), PG, and PI were analyzed with negative ion mode, whereas positive ion mode was used for phosphatidylcholine (PC), LPC ceramide (CER), DAG, TAG, CE, and SM. The detailed mass spectrometry conditions such as collision energy, declustering potential, and mass transition are obtained prior to participation.

**Generation of adenoviruses.** Adenoviruses containing human LIPIN2 cDNA, DGAT2 cDNA, Scal shRNA, and GFP shRNA were generated using a ViraPower Adenoviral Expression System (Invitrogen).

**SCD activity.** SCD activity was measured as previously described (31, 32, 67, 68). For the microsomal SCD assay, microsomes were incubated with 14C-18:0-Coa (5 μCi) and NAAD (20 mM) for 5 minutes. For the cell assay, cells were incubated with 14C-18:0 or 14C-16:0-BSA complex for 16 hours. After the reaction, the extracted and saponified fatty acids were separated by 10% silver nitrate–impregnated TLC using chloroform/methanol/acetic acid/water (90:8:1:0.8) as the developing solution. The plates were autoradiographed. Fractions were scraped off of the plate, and radioactivity was measured using a liquid scintillation counter.

**PAP and DGAT activity.** PAP activity was measured on cell extracts, as described previously with a modification (69, 70). Cell extracts from HEK293AD cells treated with adenovirus containing human LIPIN2 were used as a crude LIPIN2 enzyme. Cells were harvested with a buffer containing 0.25 M sucrose, 2 mM dithiothreitol (to stabilize PAP1 activity), protease inhibitor mixture, phosphatase inhibitor cocktail, and 0.15% Tween 20. We designed an optimum assay for PAP using 100 mM Tris/maleate buffer (pH 6.5), 5 mM MgCl2, 2 mg/ml fatty acid–poor bovine serum albumin, and 0.6 mM PA, which was dispersed in 0.4 mM egg PC and 1 mM EDTA that was used to prepare the lipid substrate. The conversion of PA to DAG by PAP was analyzed by LC-MS. PAP activity in the LIPIN2 enzyme was completely inhibited in the presence of 5 mM N-ethylmaleimide. The PAs 18:0/18:0, 16:0/16:0, 18:1/18:1, 16:0/18:1-PA, and 18:2/18:2 were purchased from Avanti Polar Lipids. DGAT activity was measured on microsomal fraction, as described previously, with a modification (71). Microsomal fraction from HEK293AD cells treated with adenovirus containing human DGAT2 was used as a crude DGAT2 enzyme. Crude DGAT2 (1 μg) was incubated for 5 minutes at 37°C with 25 μM 18:0-Coa and 200 μM DAGs (18:0/18:0-DAG and 18:1/18:1-DAG) from Nu Check. The conversion of DAG to TAG by DAGT2 was analyzed by LC-MS. The conditions of LC-MS were described in the above section of the LC-MS–based lipidomic analysis.

**Biochemical analysis.** Serum cholesterol, triglyceride, phosphorus, and calcium were analyzed using colorimetric enzyme assay kits (Pointe Scientific). Serum FGF23 was measured using an FGF23 ELISA kit (Immuneassays International).

**Human subjects.** Serum samples were collected from 10 patients with stage 3 and 4 CKD (mean eGFR 34 ± 7 ml/min/1.73 m2) and mean age (57 ± 7 years), and 10 age-matched patients with normal kidney function (mean eGFR 80 ± 12 ml/min/1.73 m2) and mean age (59 ± 14 years) after a minimum 4-hour fast. Trained personnel at the University of Colorado Clinical and Translational Research Center collected the serum samples. All patients gave informed consent for their blood specimen to be used for research purposes. Glomerular filtration rate (GFR) was estimated using the 4-variable modification of diet in renal disease prediction equation. All other parameters, such as serum cholesterol, phosphorus, and calcium, were previously reported (9). The nature, benefits and risks of the study were explained to the volunteers, and their written informed consent was obtained prior to participation.

**Statistics.** Data were collected from more than 2 independent experiments and were reported as the mean ± SEM. Statistical analysis for 2-group comparison was performed using a 2-tailed Student’s t test, or one-way ANOVA with a Student-Newman-Keuls post-hoc test for multigroup comparison. Significance was accepted at P < 0.05.

**Study approval.** Animal experiments were approved by the Institutional Animal Care and Research Advisory Committee of the University of Colorado Boulder. All human participants were enrolled in a separate clinical research study, and the procedures were approved by the Colorado Multiple Institutional Review Board or the Institutional Review Board of the University of Colorado Boulder.

**Author contributions**

M. Masuda, SMA, and M. Miyazaki designed and performed the main experiments, and ALK and KO provided technical assistance for the main experiments. MK, JMN, and SO provided essential mouse lines and ALK and KO provided technical assistance for the main experiments. MK, JMN, and SO provided essential mouse lines and ALK and KO provided technical assistance for the main experiments. MK, JMN, and SO provided essential mouse lines and ALK and KO provided technical assistance for the main experiments. MK, JMN, and SO provided essential mouse lines and ALK and KO provided technical assistance for the main experiments. MK, JMN, and SO provided essential mouse lines and ALK and KO provided technical assistance for the main experiments. MK, JMN, and SO provided essential mouse lines and ALK and KO provided technical assistance for the main experiments.

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