Thyroid hormone responsive protein Spot14 enhances catalysis of fatty acid synthase in lactating mammary epithelium

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Abstract    Thyroid hormone responsive protein Spot14 has been consistently associated with de novo fatty acid synthesis activity in multiple tissues, including the lactating mammary gland, which synthesizes large quantities of medium chain fatty acids (MCFAs) exclusively via FASN. However, the molecular function of Spot14 remains undefined during lactation. Spot14-null mice produce milk deficient in total triglyceride and de novo MCFAs that does not sustain optimal neonatal growth. The lactation defect was rescued by provision of a high fat diet to the lactating dam. Transgenic mice overexpressing Spot14 in mammary epithelium produced total milk fat equivalent to controls, but with significantly greater MCFAs. Spot14-null dams have no diminution of metabolic gene expression, enzyme protein levels, or intermediate metabolites that accounts for impaired de novo MCFAs. When [13C] fatty acid products were used in vitro using crude cytosolic lysates, native FASN activity was 1.6-fold greater in control relative to Spot14-null lysates, and add back of Spot14 partially restored activity. Recombinant FASN catalysis increased 1.4-fold and C = 14:0 yield was enhanced 4-fold in vitro following addition of Spot14. These findings implicate Spot14 as a direct protein enhancer of FASN catalysis in the mammary gland during lactation when maximal MCFA production is needed.—Rudolph, M. C., E. A. Wellberg, A. S. Lewis, K. L. Terrell, A. L. Merz, N. K. Maluf, N. J. Serkova, and S. M. Anderson. Thyroid hormone responsive protein Spot14 enhances catalysis of fatty acid synthase in lactating mammary epithelium. J. Lipid Res. 2014. 55: 1052–1065.

Supplementary key words fatty acid synthase catalysis • [13C] fatty acid quantification • medium chain fatty acids • lactation

Mammalian cells manufacture fatty acids de novo using a distinct pathway to synthesize fatty acids from acetyl and malonyl esters of CoA catalyzed by the dimer of FASN (EC 2.3.1.85) (1). FASN is absolutely essential for production of de novo fatty acids in mammals (2). The biological requirement for FASN is highlighted by the fact that FASN is detected in most normal human tissues and that homozygous deletion of FASN in the mouse results in embryonic lethality (3, 4). The importance of the de novo pathway is underscored by the complex biological functions of its fatty acid products, including biosynthesis of phospholipids, energy storage via esterification into triglycerides (TAGs), use as energy substrates for β-oxidation, providing both endocrine and nuclear hormone signaling ligands, and for critical posttranslational modifications of proteins (5–12). Perhaps the most demanding setting for FASN catalysis is during lactation, when extremely high quantities of de novo fatty acids are transmitted to the offspring as substrate for the growth of the young (13–15).

Regulation of the principle enzymes of de novo fatty acid synthesis, ATP citrate lyase (ACLY) (EC 2.3.3.8), acetyl-CoA carboxylase (ACC) (EC 6.4.1.2), and FASN, has been shown to occur at the level of enzyme gene expression (16), translation/degradation (17–19), phosphorylation (20–22), assembly into multimeric complexes (23–25), and allosteric activation/inhibition (18, 23, 26, 27). Recently, evidence for the control of de novo fatty acid synthesis by small effector proteins has emerged. Thyroid hormone responsive protein “Spot 14” (THRSP) (Spot14, S14) and only family

Abbreviations: ACC, acetyl-CoA carboxylase; ACLY, ATP citrate lyase; DGAT, diacylglycerol O-acyltransferase; HCD, high carbohydrate diet; HFD, high-fat diet; MCFAs, medium chain fatty acid; MEC, mammary epithelial cell; MMTV LTR, Mouse Mammary Tumor Virus Long Terminal Repeat; Spot14-R (MIG12), Spot14-related protein; TE2, thioesterase 2; THRSP, thyroid hormone responsive protein Spot 14; TAG, triglyceride (triacylglyceride).

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member Spot14-related protein (Spot14-R) (MIG12, Mid1IP1) are postulated to function as effector proteins for the de novo fatty acid synthesis pathway (28, 29). Spot14 was discovered in 1981 and was established to be a small acidic protein of 17 kDa with an isoelectric point of 4.85 (30). It has no known enzymatic function, the primary sequence predicts no established functional domains, and the crystal structure has identified three antiparallel α-helices in an asymmetrical dimer (28, 31). Spot14 mRNA is sharply induced in the liver of streptozotocin-induced diabetic rats by insulin treatment, as well as in livers of fasted rats refed carbohydrates and acutely treated with thyroid hormone (32, 33). Its gene expression in the liver was decreased in rats fed a diet containing long chain PUFAs, and in the lactating mammary gland following administration of trans-10 cis-12 conjugated linoleic acid (34–36). In 2003, we observed that Spot14 was induced in unison with 75 metabolic genes in the mouse mammary gland at secretory activation (37). More recently, we showed regulation of Spot14 mRNA by sterol response element binding protein 1 in mammary epithelial cells (MECs) depleted of the adipose compartment, and that Spot14 mRNA in MECs was not suppressed by provision of a high fat diet (HFD) to the dam (13). Others have shown evidence, independent of rodent models, for Spot14 involvement during milk fat production in dairy cows (36, 38). Subsequent analysis of the Spot14-null mouse firmly established Spot14 as necessary for the de novo synthesis of medium chain fatty acids (MCFAs) characteristic of milk TAG (39). Hence, in lactation, Spot14 appears to enhance de novo synthesis of the MCFAs that contribute to the total milk energy content needed for neonatal growth.

Both activation and inhibition of de novo fatty acid synthesis by Spot14 have been reported, depending upon the lipogenic tissue type and the cellular context (28, 29, 31, 39–42). For example, the rate of de novo fatty acid synthesis in the Spot14-null mammary gland was decreased over 2-fold, implicating Spot14 as an activator of de novo fatty acid synthesis in this tissue (39). Conversely, Spot14-null hepatic de novo fatty acid synthesis was increased 2-fold under lipogenic stimuli, suggesting Spot14 is an inhibitor of this process in the liver (40). Coexpression of Spot14 with Spot14-R was determined to be inhibitory, as Spot14 heterodimerizes with Spot14-R to prevent Spot14-R association with ACC and decrease activation of the hepatic de novo fatty acid synthesis pathway (45). In adult neural stem cells, Knobloch et al. (29) demonstrated cytosolic localization and coimmunoprecipitation of Spot14 and Spot14-R, and that Spot14 overexpression decreased de novo synthesis of fatty acids in low proliferating cells. Colbert et al. (28) used liver-specific antisense Spot14 knock down in chow-fed mice resulting in approximately 30% decreased $^3$H$_2$O incorporation into de novo lipid. This work was supported by efforts from Aipoalani et al. (41), who knocked down both Spot14 and Spot14-R in primary hepatocyte culture and observed a 65% reduction of $[^14]$C-acetate incorporation into de novo fatty acids. Thus, overlapping and conditional roles for Spot14 regulation of fatty acid synthesis have been demonstrated (29, 41). In mammary cancer cells, antisense Spot14 knock down decreased $^{14}$C-acetate incorporation into lipid nearly 60%, supporting a role for Spot14 in de novo fatty acid synthesis activation (44). Cumulatively, these published findings indicate that Spot14 may both positively or negatively regulate the de novo fatty acid synthesis pathway depending on the tissue-specific or cellular context, as well as the presence of Spot14-R. The lactating mammary gland lacks expression of Spot14-R, meaning that the MEC-specific function of Spot14 is independent from Spot14-R. In the lactating mammary gland, Spot14 is considered an activator of de novo fatty acid synthesis because the milk produced by Spot14-null dams contains 60% less MCFAs (39), which are unique products of mammary alveolar de novo fatty acid synthesis during lactation (15). Zhu et al. (39) determined that the activities of enzymes upstream of FASN in the biosynthetic pathway were present and functioning in the Spot14-null mammary gland during lactation, including malic enzyme, 6-phosphogluconate dehydrogenase, glucose-6-phosphate dehydrogenase, ACC, and FASN. Importantly, in vivo tracer studies using $^2$H$_2$O incorporation into milk MCFAs revealed that the rate of fatty acid synthesis in Spot14-null mice was decreased approximately 2.5-fold (39). Taken together, these observations indicate that the catalytic rate of FASN in the lactating mammary gland is diminished in the Spot14-null mouse.

Here we report that tissue-specific overexpression of Spot14 in MECs increased the amount of MCFAs known to originate from de novo synthesis, without altering the percent milk fat by volume or influencing offspring growth. Spot14-null milk production was not diminished, but milk from Spot14-null dams had significantly reduced fat percentage leading to restricted pup growth. We found that this loss in total milk fat and subsequent inability to support litters was rescued by provision of a diet high in fat to the Spot14-null dams. Further, we tested the hypothesis that Spot14 increases the in vitro catalytic rate of both recombinant FASN and of native FASN in MEC cytosolic lysates using a recently developed method that specifically quantifies synthesis of $[^13]$C fatty acid products (45). Together, these results suggest the molecular function of Spot14 in MECs is to enhance FASN catalysis during lactation when de novo fatty acid synthesis is needed to achieve optimal MCA content in the milk.

**MATERIALS AND METHODS**

**Animals**

Mice were maintained in the Center for Comparative Medicine. The Institutional Animal Care and Use Committee of the University of Colorado Denver approved all animal procedures. C57Bl/6 and FVB control mice were purchased from Jackson Laboratories (Bar Harbor, Maine). Spot14-null mice maintained on the C57B6/J background were a kind gift of Dr. Cary Mariash, and details regarding generation of these mice are described elsewhere (40). The Transgenic and Knockout Core Facility at the University of Colorado Denver generated the transgenic mice overexpressing Spot14 driven by the mouse mammary tumor virus long terminal repeat (MMTV-Spot14). The pCMV-SPORT6 vector containing mouse Spot14 was purchased from Open Biosoys
Acid pellets were resuspended in 30 μl of 1% pentafluorobenzyl alcohol from Integrated DNA Technologies (Coralville, IA) or from Thermo Fisher Scientific. Tail DNA with the following primers: forward, 5′-TAG-TCT-GGG-ACG-TCG-TAT-GGG-TA3′ to generate a product of 200 base pairs.

**Milk fat and fatty acid composition analysis**

Milk was collected at lactation days 4 and 10 using vacuum suction, and milk fat analysis was conducted as previously described (46). Weigh-suckle-weigh experiments were conducted according to Dempsey et al. (47). Milk protein concentrations were determined using the Pierce 660 protein assay (Thermo Fisher Scientific). Milk lactose concentrations were quantified using the lactose assay kit according to the manufacturer’s instructions (Abcam, ab83384). HPLC grade reagents were purchased from Sigma-Aldrich, St. Louis, MO. Neutral lipid extraction was adapted from Hutschins et al. (48, 49), and pentafluorobenzyl derivatization was according to Zarini et al. (50). Mouse milk was thawed at 37°C for 10 min, vortexed, and 5 μl of whole milk was transferred to 2 ml Eppendorf tubes containing 0.5 ml of 2:1 v/v methanol/water; samples were acidified with 40 μl of 1 M HCl, vortexed into solution, cleared by centrifugation at 10,000 g, and supernatant was transferred into borosilicate glass tubes. Milk fat was extracted twice using 1 ml of 2:1 v/v isooctane/ethyl acetate, and samples were vortexed vigorously, allowed to settle for 5 min at room temperature, and then centrifuged at 500 g for 1 min at room temperature. Samples were taken to dryness using a Savant SpeedVac concentrator and lipid pellets were resuspended in 0.5 ml of 100% methanol.

Blended internal standards (1 μg) containing [D]-decanoic acid, [D]-lauric acid, [D]-myristic acid, 1,2,3,4-[13C]palmitic acid, [D]-stearic acid, [D]-oleic acid, [D]-arachidonic acid, and [D]-docosahexanoic acid (purchased from Sigma-Aldrich; Cambridge Isotopes, Andover, MA; or Cayman Chemical Co., Ann Arbor, MI with 99 atom % 13C and 99 atom % D, respectively) were added to each sample. Samples were saponified by addition of 0.5 ml of 1 M NaOH at 37°C for 1 h mixing at 20 min intervals; and samples were acidified with 0.6 ml of 1 M HCl to a pH between 2.0 and 4.0 and mixed by vortex. Saponified fatty acids were extracted twice with 1.0 μl of isooctane; 300 μl of the saponified fatty acid fraction was transferred to new glass tubes, and samples were taken to dryness under N2. Saponified fatty acid pellets were resuspended in 30 μl of 1% pentafluorobenzyl bromide in acetonitrile and 30 μl of 1% N,N-diisopropylethylamine in acetonitrile to initiate derivatization and incubated at room temperature for 30 min (50). Samples were dried under N2, resulting pellets were resuspended in 1 ml of isoctane, vortexed, and 50 μl was transferred into vials for GC-MS. Sample analysis was performed by negative ion chemical ionization GC-MS using the Finnigan DSQ GC-MS system (Thermo Finnigan, Thousand Oaks, CA), with a ZB-1 column (15 m, 0.25 mm inner diameter, 0.10 mm film thickness; Phenomenex) using methane as reagent gas. The GC temperature gradient was 125 to 230°C at 20°C/min, 230 to 245°C at 5°C/min, and finally 245 to 300°C at 30°C/min, before being held at 310°C for 1 min. Data were acquired in full scan mode to identify fatty acids of acyl chain length from 10 to 22 carbons. Peak areas of the analyte or of the standard were measured, and the ratio of the area from the analyte-derived ion to that from the internal standard was calculated.

**NMR metabolomic analysis**

Lipid and aqueous metabolomic analysis of both whole gland and of MEC preparations were carried out according to Rudolph et al. (51). The MEC and tissue extracts were analyzed using a Bruker 500 MHz DRX NMR spectrometer. All metabolite concentrations were assessed as micromoles per gram of tissue.

**Adipose depleted MEcs**

MECs were prepared as previously described (13) with the following modifications: mammary digestion buffer included 5 ml of Ham’s F12 medium (Life Technologies, Grand Island, NY), 5 mg trypsin (Gibco #840-7250), 10 mg collagenase (Boehringer #1088793), 50 mM NaF, and 5.0 mM sodium orthovanadate (heat activated). Tissues were minced and digested in 5 ml of buffer as previously described. After incubation for 25 min in a 37°C shaker at 200 rpm, 1 ml of FCS was added to the digested cell suspension, brought up to 15 ml with ice-cold PBS, and spun at <300 g for 10 min. Washing was repeated two additional times with PBS. Approximately 200 μl of the resulting 500 μl of packed organoids was lysed in 1 ml of TRIZol (Life Technologies) for RNA analysis and the other portion was lysed in 1 ml of lysis buffer for protein, consisting of 50 mM Tris (pH 7.4), 150 mM NaCl, 2.0 mM EDTA, 50 mM NaF, 5.0 mM sodium orthovanadate, 1 Triton X-100, 1% deoxycholate, and 0.1% SDS, to which 0.57 mM PMSF, 20 μl/ml EDTA-free Inhibitor cocktail (Roche), and 1.0 mM DTT was added. Samples were homogenized using a Brinkman Polytron, and lysate was centrifuged at 13,000 g for 10 min at 4°C. Protein concentrations were determined using Pierce 660 protein assay (Thermo Fisher Scientific).

**Quantitative real-time PCR analysis**

Supplementary Table II lists the primers, probes, and amplicon formula weights for all genes used in this study. The primers and probes were purchased for use with TaqMan reaction chemistry from Integrated DNA Technologies (Coralville, IA) or from Applied Biosystems (Life Technologies) with 5′ FAM fluorophore and either 3′ Black Hole or minor groove binding quenchers, respectively. The VERSO cDNA synthesis kit (Thermo Fisher Scientific) was used, and each reaction consisted of 0.5 μg total RNA in 11 μl of nuclease-free water, to which 1.0 μl of blended 3:1 v/v random hexamers to oligo dT was added and samples were incubated at 70°C for 5 min. Samples were collected with a quick spin, and 9 μl of a mix containing 4 μl 5× buffer, 2 μl of dNTP blend, 1 μl of RT enhancer, and 1 μl of VERSO enzyme was added per reaction. The cDNA was diluted to 50 μl, and 2.5 μl was used as template into the quantitative (q)PCR reaction (representing 25 ng total RNA per reaction). A qPCR master mix was made containing 12.5 μl of Absolute Fast qPCR Mix-Lox Rox (Thermo Fisher Scientific), 1.25 μl of 20× primer/probe mix (10 μM
forward, 10 µM reverse, and 5 µM probe), and 6.25 µl of nuclease-free water per reaction well, to which 2.5 µl of cDNA per reaction was added. qPCR data were collected on the Applied Biosystems 7500 Fast or the 7900 HT thermocycler (Life Technologies). PCR products representing the unique product (the ampiclon) for each gene were amplified from cDNA as above, and amplicons were purified using the Qiagen MinElute PCR purification kit (Qiagen). Concentrations of purified double stranded amplicons for each gene were measured using the NanoDrop (Thermo Fisher Scientific). The molarity of each amplicon was calculated using the equation:

\[
\text{mol/liter} = \frac{\text{amplicon \, ug}}{2 \times 308.95 \times R \times N}
\]

where 308.95 g/mol is the average molecular weight of a nucleotide base and N represents the nucleotide length of the amplicon sequence (multiplied by 2 for double stranded). The resulting molar amount was converted to molecules/l by multiplying by \(6.022 \times 10^{23}\), and subsequently to molecules/µl by dividing by \(1 \times 10^5\). Serial dilutions were performed to generate a standard curve to quantify the amount of DNA (2 µl) by heat shock, single colonies were selected and screened for inclusion of the transgene, and a high expressing bacterial clone containing the pET11b-Spot14 plasmid was chosen. The plasmid was submitted for sequencing, and indicated that Spot14 had 100% identity to the original sequence.

Preparation of recombinant FASN

The DH10Bac bacterial stock (DH10-FASN-His) containing the poly-His-tagged FASN baculovirus construct was generously provided by Dr. Stuart Smith and Andreziet Witkowski (Children’s Hospital Oakland Research Institute). BD BaculoGold Bright DNA (Life Technologies) and 2 µg of purified FASN-His BacMid DNA were cotransfected into SF9 cells to generate FASN-His baculovirus. SF9 cells (500 x 10^6) were transduced with FASN-His baculovirus at an MOI of 2-3. SF9 pellet was resuspended in 5 ml of buffer A containing 250 mM potassium phosphate (pH 7.0), 0.2 mM DTT, 10% glycerol, and 2× EDTA-free protease inhibitors (Roche Applied Science); passed through a glass dounce homogenizer using 25 strokes; and the cellular debris was cleared by ultracentrifugation at 100,000 g for 30 min. Nickel-NTA beads (5 ml) (Qiagen) were packed into a column, and the beads were equilibrated in 40 ml of buffer A. The cleared SF9 lysate was added to the column by gravity drip, and the column was washed with 20 ml buffer A + 50 mM imidazole (repeated twice). FASN-His was eluted from nickel-NTA using 10 ml buffer A + 200 mM imidazole, and the eluate was transferred to dialysis tubing for equilibration in 1 l of buffer A + 2 mM DTT overnight at 4°C. Equilibrated FASN-His was concentrated in successive rounds using an Amicon Ultra-4 centrifugal filter unit (Millipore, Billerica, MA) at 3,000 g for 20 min at 4°C. FASN-His enzyme concentration was measured using a 280 nm extinction coefficient of 482,200 M⁻¹cm⁻¹ (dimer, without disulfides).

Preparation of recombinant Spot14

The pCMV-SPORT6 vector containing mouse Spot14 was purchased from Open Biosciences (Thermo Fisher Scientific), and the plasmid was purified using the Qiagen HiSpeed plasmid maxiprep kit (Qiagen) according to the manufacturer’s protocol. The purified plasmid containing mouse Spot14 was modified by PCR to introduce engineered restriction digestion cut sites into the sequence (forward: GGA-ATT-CCA-TAT-GAT-GCA-AGT-GCT-ACC-CTG-CCC-TGT-CAT-TTC; reverse: CGG-GGA-TCC-GGC-TCA-CTA-CAG-GAC-GCT-CCC-TGT-CAT-TTC). The PCR product was cleaned up using the Qiagen MinElute PCR purification kit according to manufacturer’s protocol (Qiagen) and was run on a 1% agarose gel to isolate the desired product. Gel-extracted PCR product (2 µg) and 2 µg of pET11b plasmid DNA were cut using Ndel and BamHI restriction enzymes in React buffer 3 in a total volume of 100 µl at 37°C overnight according to manufacturer’s instructions (Life Technologies). Digested DNA was cleaned up with the Qiagen DNA Mini purification kit (Qiagen), and DNA was ligated in a 15 µl volume using 8 µl of Spot14 insert, 1 µl vector, 3 µl 5× ligation buffer, 1 µl 10 mM ATP, and 1.5 µl T3 ligase at 10°C for 3 h. DH5α competent bacteria (25 µl) were transformed with ligated DNA (2 µl) by heat shock, single colonies were selected and screened for inclusion of the transgene, and a high expressing bacterial clone containing the pET11b-Spot14 plasmid was chosen. The plasmid was submitted for sequencing, and indicated that Spot14 had 100% identity to the original sequence. 

Bl21 competent bacteria were transformed and plated, and single colonies were selected as above. Small preparations of single colonies were induced with IPTG, and production of the Spot14-His protein was evaluated by immunoblot using the anti-Spot14 antibody. The colony with the greatest amount of Spot14-His production was maintained for additional studies. Bl21-Spot14-His bacteria were grown in 1 l of LB media under standard conditions containing 50 µg/ml ampicillin, were induced at an optical density of 0.6 with 0.5 mM IPTG, and the growth was maintained at 27°C overnight. Following protein induction, bacteria were collected by centrifugation at 6,000 g for 15 min and LB was decanted. Bacteria were resuspended in 40 ml of buffer containing 50 mM potassium phosphate (pH 7.0), 50 mM NaCl, 10 mM imidazole, and 1 mM DTT; and cells were lysed by sonication using 40 pulses at 90% power. Lysates were cleared by centrifugation at 15,000 g for 20 min, and the supernatant was transferred to a 50 ml conical tube. Purification of Spot14-His was performed essentially as the FASN-His protein, and the molecular weight was 19,388 g/mol and the extinction coefficient used was 12,090 M⁻¹cm⁻¹.

Standard and quantitative immunoblotting

MEC lysates were prepared as above and proteins were resolved using standard SDS PAGE techniques, transferred to normal or low fluorescence PVDF membrane (Millipore), and incubated overnight at 4°C with primary antibodies. Antibodies were purchased from Cell Signaling Technology: ACC (#3676), pACC ser79 (#3661), ALCY (#4332), pALCY ser45 (#4331), AMPK (#2603), pAMPK thr172 (#2535), and CyPA (#2175); from Santa Cruz Biotechnology: FASN (SC-20140), goa anti-rabbit-HRP (SC-2004), KRT18 (SC-28264), and SLC25A1 (sc-86392); from Fitzgerald Industries International: PLIN1 (70R-PR004), PLIN2 (20R-AP002); from Abcam: Spot14 (ab56193); CSN2 was a kind gift of Margaret Neville (University of Colorado); and thioesterase 2 (TE2) was a kind gift of Stuart Smith (CHORI, San Francisco, CA). Recombinant FASN and Spot14 proteins were used to generate a standard curve to quantify the amount of FASN or Spot14 present in MEC lysates. Triplicate dilutions ranging from 0.15 to 1.0 µg of FASN or Spot14 protein were run using standard SDS PAGE techniques, transferred to low fluorescence PVDF membrane, and incubated overnight at 4°C with primary antibodies as above. Quantitative immunoblots were incubated in 1:15,000 goat anti-rabbit IRDye 800CW secondary (LI-COR Biosciences) for 2 h at room temperature. Infrared immunoblots were scanned using the LI-COR Odyssey system and data were analyzed using LI-COR Image Studio 2.0 software (LI-COR Biosciences) for 2 h at room temperature. Infrared immunoblots were scanned using the LI-COR Odyssey system and data were analyzed using LI-COR Image Studio 2.0 software (LI-COR Biosciences).
GC-MS as described above. Following initiation, 200 μg FASN were also provided a high carbohydrate/low fat diet to stimulate reduced growth of the offspring, and we have confirmed that the lactation phenotype in Spot14-null dams results in reduced growth of the offspring. Extending those findings, Spot14-null mothers reared by Spot14-null dams (Fig. 1A). These results indicate that Spot14-null dams consuming normal chow or a HCD cannot generate enough milk energy density to sustain the optimal growth of their offspring, but that incorporation of maternal dietary fat into the milk TAG restores this energy deficit.

RESULTS

HFD restores and high carbohydrate diet impairs growth of pups reared by Spot14-null dams

The TAG macronutrient content in milk provides the primary source of calories needed for the growth of the offspring, particularly in mice that produce about 30% milk fat by volume. Maternal HFD has been shown to decrease de novo fatty acid synthesis in mammary alveolar cells during lactation, while a high carbohydrate/low fat diet stimulates this pathway (13, 52). Zhu et al. (39) showed that the lactation phenotype in Spot14-null dams results in reduced growth of the offspring, and we have confirmed these published data by observing reduced growth of offspring reared by Spot14-null dams (Fig. 1A). To eliminate any effect due to the genetic background of the pups, both control and Spot14-null dams were cross-fostered with wild-type pups, and litters were standardized to seven or eight pups. Extending those findings, Spot14-null mothers were also provided a high carbohydrate/low fat diet to stimulate de novo fatty acid synthesis 1 day prior to parturition, and the lactation defect worsened (Fig. 1A; high carbohydrate diet (HCD), diamonds; 3.9% kcal fat). The Spot14-null lactation phenotype was rescued by maternal provision of a diet rich in fat 1 day prior to parturition (HFD, triangles; 46% kcal fat) (Fig. 1A). These results indicate that Spot14-null dams consuming normal chow or a HCD cannot generate enough milk energy density to sustain the optimal growth of their offspring, but that incorporation of maternal dietary fat into the milk TAG restores this energy deficit.

Mammary epithelial Spot14 overexpression does not impair mammary gland development or lactation

To evaluate the effect of transgenic Spot14 on mammary gland development and lactation, mice were generated that overexpress Spot14 driven by the mammary epithelial specific MMTV LTR. Spot14 transgene expression was confirmed by immunohistochemistry directed against the HA tag engineered into the transgene (supplementary Fig. 1A). Transgene expression was also quantified by qPCR, and revealed that the highest MMTV-Spot14 expression was detected in the mammary gland, followed by occasional detection of the transgene in the lung of some mice; however, no expression of the transgene was detected in the ovary, uterus, heart, hypothalamus/pituitary, or liver (supplementary Fig. IC, D). Qualitative evaluation of whole mounts from virgin day 5 and nulliparous mammary glands indicated subtle differences in branching may exist in MMTV-Spot14 females, but terminal end buds were hollow and appeared similar to controls (supplementary Fig. IC, D). Examination of whole-mounted mammary glands from mid-pregnant and mid-lactating control and MMTV-Spot14 females, as well as analysis of histological sections at parturition day 14 and lactation day 4, revealed no obvious differences in gland morphology and organization (supplementary Fig. 1D, E). Consistent with competent development and function of the mammary gland was the normal growth of offspring reared by MMTV-Spot14 transgenic dams (Fig. 1B). Although the weights diverged slightly upward for pups reared by the MMTV-Spot14 dams, no significant difference in pup growth was observed, suggesting that MMTV-Spot14 mothers do not supply excessive nutrients to the offspring.

Spot14-null dams, but not MMTV-Spot14 dams, produce lipid-poor milk

To determine the influence of Spot14 on milk fat content, control, Spot14-null, and MMTV-Spot14 transgenic milk was collected at lactation day 4 and the fat content was analyzed as previously described (46). Control dams synthesized milk containing approximately 35% milk fat by wet volume, however the fat content in milk from Spot14-null mice was significantly decreased, approximately 1.6-fold \((P = 0.001)\) (Fig. 2A). In contrast to the Spot14-null dams, milk fat percentage from the MMTV-Spot14 dams was equivalent to controls, indicating that MEC overexpression of Spot14 did not influence total TAG amount. The protein and lactose contents of the Spot14-null milk and the MMTV-Spot14 milk were not altered relative to control milk (data not shown). A weigh-suckle-weigh approach was used to determine the amount of milk produced in a 2 h period (47). Milk production in the Spot14-null dams was equivalent to controls, suggesting milk synthesis and secretion is not obstructed (Fig. 2B).

The NMR metabolomic lipid analysis of the Spot14-null MECs, dephospho sphingomyelins (53), was consistent with reduced milk lipid by volume, indicated by nearly 30% decreased intracellular TAG content.
Spot14 increases fatty acid synthase activity in MECs to production of lipid-poor milk, rather than defective lactation that could result from impaired gland development, impaired secretory activation, or obstructed secretion as is seen in other models. This observation is further supported by the ability of the HFD to compensate for the lipid-poor milk produced by Spot14-null mothers provided the control diet (Fig. 1A; open circles vs. triangles). (Table 1; \(P = 0.01\)). Similar results were obtained with analysis of the whole mammary gland (Table 1). The PUFA components in the Spot14-null MECs and whole gland were also significantly decreased, but the MUFA:PUFA ratio was unchanged (Table 1). No differences were observed for the phospholipids quantified. These results indicate that the Spot14-null lactation phenotype is due to production of lipid-poor milk, rather than defective lactation that could result from impaired gland development, impaired secretory activation, or obstructed secretion as is seen in other models. This observation is further supported by the ability of the HFD to compensate for the lipid-poor milk produced by Spot14-null mothers provided the control diet (Fig. 1A; open circles vs. triangles).

![Fig. 1. Pup weights of offspring from control, Spot14-null, and MMTV-Spot14 mothers. A: Litters were standardized to seven or eight wild-type pups per dam, and \(n = 3–5\) dams per genotype and diet. Dams were provided standard chow, HCD (3.9% kcal fat), or HFD (46% kcal fat) 1 day prior to parturition. Growth of pups reared by Spot14-null dams on the chow diet (open circles) diverged from that of controls (solid circles) at lactation day 11 (\(P \leq 0.05\)). Maternal HFD (solid triangles) provided 1 day prior to parturition restored the pup growth deficit observed in offspring suckling Spot14-null dams provided chow (open circles). Pups reared by dams provided HCD grew at the lowest rate. B: Pup growth of offspring reared on control or dams overexpressing Spot14 driven by a mammary epithelium specific promoter (MMTV-Spot14); the difference is not significant.](image)

![Fig. 2. Percentage of milk fat by volume at lactation day 4. A: Milk was collected and analyzed by creamocrit as described. Spot14-null (Spot14\(^{-/-}\)) milk fat was reduced approximately 35% relative to controls, while milk fat percentage from dams overexpressing Spot14 in the mammary epithelium (MMTV-Spot14) was not different from controls. All animals were maintained on standard rodent chow. B: Weigh-suckle-weigh analysis of total milk production was not different in control and Spot14-null mice.](image)
Overexpression of Spot14 increases MCFA composition of milk TAG

In contrast to the adipose tissue and the liver, de novo synthesis of fatty acids in the mammary gland generates great amounts of MCFA that are defined by the saturated acyl chain length ranging between 10 and 14 carbons \((C = 10 \text{ to } 14)\) \((54)\). Conversely, the long chain fatty acids in milk TAG, those with an acyl chain length of \(C \geq 16\) carbons, have been shown to originate from either dietary sources or from mobilization of TAG reserves of the adipose tissue \((15)\). Accordingly, “preformed” fatty acids originate from sources other than the de novo fatty acid pathway of the MEC and possess an acyl chain length of \(C \geq 16\) carbons. Fatty acids with \(C = 16\) can be synthesized de novo in the MEC, or they might be synthesized by the liver and adipose tissue, or they could be present in the diet. Therefore, \(C = 16\) represents fatty acids derived from both de novo and preformed sources.

Milk was collected from control, Spot14-null, and MMVT-Spot14 overexpressing dams as previously described \((46)\) and the neutral lipids were extracted, saponified with NaOH to release individual fatty acids, and the fatty acid composition was quantified by GC-MS according to previous methods \((45, 48, 50)\). Loss of Spot14 resulted in significantly decreased MCFA from the milk TAG at lactation day 10 \((\text{Fig. 3A)}\), and although the PUFA appeared to be decreased, the difference did not reach significance. These data were consistent with the NMR lipid analysis of extracts prepared from whole mammary glands or MECs. Conversely, overexpression of Spot14 in MECs from transgenic mice resulted in significantly increased MCFA in milk TAG at lactation day 4 \((\text{Fig. 3B)}\). This finding indicates that overexpression of Spot14 in vivo during lactation is sufficient to elevate the de novo synthesized MCFA products incorporated into milk TAG, especially in early lactation when MCFA are not yet largely present \((13)\). Interestingly, the levels of linoleic acid \((18:2)\) and arachidonic acid \((20:4)\) were also elevated in milk from MMVT-Spot14 transgenic mice. Although this result is consistent with loss of PUFA in Spot14-null tissues and milk, we do not understand the functional relevance (if any) of this observation at present. Collectively, when Spot14 was lost during lactation, the de novo MCFA yield was diminished, and when Spot14 was artificially overexpressed in vivo, the de novo MCFA yield was increased.

Decreased MFCA in Spot14-null milk does not result from reduced lipogenic mRNA, proteins, and enzyme phosphorylation status

The de novo fatty acid pathway is regulated at multiple levels including gene expression, enzyme abundance, and posttranslational control, such as phosphorylation. Control and Spot14-null MECs depleted of mammary adipose were prepared in order to avoid confounding analysis due to adipocyte contamination \((53)\). Gene expression was quantified for multiple members of the glycolytic and fatty acid synthesis pathways at lactation days 4 and 10 in these MECs. mRNA levels were equivalent for the 18 metabolic pathway genes evaluated, and some mRNAs were significantly increased \((\text{supplementary Fig. II)}\). This finding indicated that loss of Spot14 in MECs did not influence expression of these genes. Overexpression of Spot14 in the MMVTSpot14 MECs also did not alter mRNA levels of ACLY, ACC, or FASN \((\text{data not shown)}\). Analysis of total protein levels in adipose-depleted MECs from control and Spot14-null mice showed no noteworthy differences for the mitochondrial citrate transporter \((\text{SLC25A1)}\), ACLY, ACC, FASN, or diacylglycerol O-acyltransferase \((\text{DGAT1)}\)

\[(\text{Fig. 4)}\]. Loss of any of these proteins could account for a reduction in intermediate metabolites that supply substrate for FASN catalysis; and, if FASN levels were decreased in Spot14-null MECs, then reduced MCFA might result. Importantly, levels of TE2 \((\text{EC = 3.1.2.14})\), the MEC-specific
Spot14 increases fatty acid synthase activity in MECs

Phosphorylation of ACC at Ser79 is associated with catalytic inhibition in vitro (21). The phosphorylation status of ACLY and ACC was evaluated using quantitative immunoblotting (Fig. 5C). Interestingly, phosphorylated ACLY (activated) and phosphorylated ACC (inhibited) were both significantly increased in Spot14-null MECs (1.8- and 2.4-fold, respectively). Regulation of FASN activity by phosphorylation is beginning to emerge. It has been reported that mTORC mediates threonine phosphorylation at positions 1029 and 1033, which negatively regulates FASN in the mouse liver (12). Currently, no commercial antibodies exist; its analysis requires immunoprecipitation of FASN and use of phosphorylated threonine-specific antibodies (12). No difference was observed in the phosphorylated enzyme responsible for releasing MCFA products from FASN during lactation (55), was maintained at the mRNA level (supplementary Fig. I, Lac 10) and appeared slightly greater at the protein level (Fig. 4). While a reduction of TE2 could account for the diminished MCFA observed in the Spot14-null milk, no difference in TE2 mRNA or protein was detected.

Quantitative immunoblot measurements were used to evaluate the total abundance of ACLY and ACC; both were significantly increased in Spot14-null MECs (1.4- and 1.7-fold, respectively), and total FASN trended upward but did not reach significance (Fig. 5A, B). In addition to total enzyme levels, phosphorylation of the de novo fatty acid synthesis enzymes potently regulates their activity. Phosphorylation of ACLY at Ser454 has been shown to activate the enzyme (20); while, phosphorylation of ACC at Ser79 is associated with catalytic inhibition in vitro (21). The phosphorylation status of ACLY and ACC was evaluated using quantitative immunoblotting (Fig. 5C). Interestingly, phosphorylated ACLY (activated) and phosphorylated ACC (inhibited) were both significantly increased in Spot14-null MECs (1.8- and 2.4-fold, respectively). Regulation of FASN activity by phosphorylation is beginning to emerge. It has been reported that mTORC mediates threonine phosphorylation at positions 1029 and 1033, which negatively regulates FASN in the mouse liver (12). Currently, no commercial antibodies exist; its analysis requires immunoprecipitation of FASN and use of phosphorylated threonine-specific antibodies (12). No difference was observed in the phosphorylated enzyme responsible for releasing MCFA products from FASN during lactation (55), was maintained at the mRNA level (supplementary Fig. I, Lac 10) and appeared slightly greater at the protein level (Fig. 4). While a reduction of TE2 could account for the diminished MCFA observed in the Spot14-null milk, no difference in TE2 mRNA or protein was detected.

**Fig. 3.** Milk fatty acid composition. A: Spot14-null (Spot14−/−) and control dams were milked at lactation day 10 when contribution of de novo synthesized fatty acid contribution in milk is maximal. Total milk lipid was extracted and saponified into individual fatty acids that were quantified by GC-MS. MCFAs (C = 10:0–14:0) and C = 16:0, known to originate from de novo synthesis, were all significantly reduced in Spot14−/− milk. B: MMTV-Spot14 and control dams were milked at lactation day 4 when de novo synthesized fatty acid contribution in milk is moderate. Fatty acid composition was quantified as in (A). Milk from MMTV-Spot14 overexpressing dams contained significantly greater de novo fatty acids (C = 10:0–16:0) than controls. Linoleic and arachidonic acids (18:2 and 20:4) were also significantly increased in MMTV-Spot14 milk at lactation day 4. (*) denotes significantly different fatty acids by Welch’s t test.
threonine form of immunoprecipitated FASN using an anti-phospho-threonine antibody that is not site specific (supplementary Fig. IVB).

Intermediate metabolites are not decreased in Spot14-null MECs

The phosphorylation status of ACLY and ACC suggested that the MECs from the Spot14-null dams should have increased acetyl-CoA (the product of ACLY) and decreased malonyl-CoA (the product of ACC). NMR metabolomics was used to quantify 35 aqueous metabolites from both the MECs and whole mammary gland lysates (WTL; contains luminal milk and adipose contents) of control and Spot14-null dams at lactation day 10, and revealed that acetyl-CoA was significantly increased (supplementary Table I). Malonyl-CoA levels were not significantly different in Spot14-null glands, despite the increased inhibitory phosphorylation of ACC (Fig. 5C). Hence, adequate levels of both acetyl-CoA and malonyl-CoA substrates required for FASN catalysis were present in Spot14-null milk. Perilipin 2 (PLIN2, adipophilin), β casein (CSN2), and cytokeratin 18 (KRT18) are MEC-specific proteins; and, perilipin 1 (PLIN1) is an adipose-specific protein whose absence indicates sufficient depletion of adipocyte compartment during the MEC preparation.

Spot14 enhances recombinant FASN catalysis in vitro

In order to test the hypothesis that Spot14 may directly increase the activity of FASN, recombinant Spot14 and FASN were prepared and FASN kinetic assays were conducted, as previously described with minor modifications

Fig. 4. Immunoblot for de novo fatty acid synthesis pathway proteins. MECs depleted of the adipose compartment (MECs) were isolated from Spot14-null (Spot14−/−) and control dams at lactation day 10. The mitochondrial citrate transporter (SLC25A1), ACLY, ACC, FASN, and DGAT1 were equivalent in Spot14−/− relative to controls. TE2 was slightly increased in Spot14−/− MECs. The proteins of the de novo fatty acid synthesis pathway are intact, which does not account for the decreased MCFAs composition in Spot14-null milk. Perilipin 2 (PLIN2), adipophilin, β casein (CSN2), and cytokeratin 18 (KRT18) are MEC-specific proteins; and, perilipin 1 (PLIN1) is an adipose-specific protein whose absence indicates sufficient depletion of adipocyte compartment during the MEC preparation.

Fig. 5. Quantitative immunoblots from day 10 control and Spot14-null MECs. A: Scanned fluorescent images of total and phosphorylated forms of ACLY, ACC, and AMPK, with cyclophilin A as a loading control. The presence of Spot14 denotes genotype. B: Background adjusted intensities plotted as the ratio of total ACLY, ACC, and FASN to cyclophilin A loading control. Total ACLY and ACC are significantly increased in the Spot14-null MECs. * P < 0.01. C: The ratio of phosphorylated to total ACLY, ACC, and AMPK was calculated from the background adjusted intensities. Phosphorylated ACLY and ACC are significantly increased and phosphorylated AMPK trends upward in the Spot14-null MECs. *P ≤ 0.01.
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The reaction velocity was determined using varied amounts of $[13\text{C}]_2$-malonyl-CoA with constant 20 $\mu\text{M}$ $[13\text{C}]_2$-acetyl-CoA and 200 mM NADPH (Fig. 6C). In the presence of Spot14, the reaction velocity was greater than FASN alone, reaching an apparent $V_{\text{max}}$ of 220 nmol $[13\text{C}]$ fatty acids synthesized min$^{-1}$ mg$^{-1}$ compared with 191 nmol $[13\text{C}]$ fatty acids synthesized min$^{-1}$ mg$^{-1}$. Interestingly, the apparent $K_m$ was reduced nearly 45% from 12.5 $\mu\text{M}$ to 6.9 $\mu\text{M}$ $[13\text{C}]_2$-malonyl-CoA in the presence of Spot14. Although kinetics under steady-state catalysis have been shown to follow a typical overall Michaelis-Menten rate law for each of the three substrates, acetyl-CoA, malonyl-CoA, and NADPH (57), the apparent $K_m$ data must be interpreted for the overall reaction kinetics and not the individual reactions leading to palmitate, such as substrate loading/unloading or acyl hydrolysis. The reduction in apparent $K_m$ would be consistent with release of fatty acid products shorter than C = 16:0 (C = 14:0) from the multi-component enzyme. Accordingly, the distribution of $[13\text{C}]$ MCFA (C = 10–14) was increased nearly 4-fold relative to

![Graph A](image1.png)

**Fig. 6.** Recombinant Spot14 increases the activity of recombinant FASN in vitro. A: FASN activity increased 1.4-fold with addition of Spot14 in 40× molar excess in the presence of 20 $\mu\text{M}$ $[13\text{C}]_2$-acetyl-CoA and $[13\text{C}]_2$-malonyl-CoA. Uniformly incorporated $[13\text{C}]$ fatty acid products of enzyme catalysis were quantified using GC-MS. B: FASN activity was increased in a Spot14 concentration-dependent manner. C: Spot14 increased the apparent $V_{\text{max}}$ and apparent $K_m$ for the overall reaction using constant 20 $\mu\text{M}$ $[13\text{C}]_2$-acetyl-CoA and 200 mM NADPH, while varying the amount of $[13\text{C}]_2$-malonyl-CoA. D: At the 20:10 $\mu\text{M}$ ratio of $[13\text{C}]_2$-acetyl-CoA to $[13\text{C}]_2$-malonyl-CoA, the total $[13\text{C}]$ fatty acid yield was increased 1.8-fold in the presence of Spot14, and the proportion of C = 10–14 $[13\text{C}]$ fatty acids was increased nearly 4-fold relative to FASN alone.

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FASN alone (Fig. 6D), and the total [13C] fatty acid yield increased 1.8-fold over the entire reaction. Overall, addition of Spot14 to recombinant FASN in vitro enhanced the rate of catalysis by increasing the product yield of \( C = 10-14 \) [13C] fatty acids.

**Native FASN activity is reduced in crude lysates from Spot14-null MECs**

The MECs from lactation day 10 control and Spot14-null dams were prepared, and crude cytosolic extracts were generated for FASN activity assays as previously described (45). MECs were prepared from the whole mammary gland to avoid any FASN and Spot14 that could be contributed by mammary adipocytes present in the whole tissue (53). For the purpose of the kinetic calculations, the amount of cytosolic FASN per microgram of total cytosolic protein from control and Spot14-null MECs was quantified relative to the regression curve using quantitative immunoblotting for recombinant FASN (supplementary Fig. III). Based on the calculated amount of FASN per microgram of MEC total cytosolic protein, equivalent amounts of FASN, as a function of cytosolic lysate volume, were loaded into each activity assay. FASN activity assays were performed as above using 20 \( \mu \)M [13C] \( \sim \) acetyl-CoA and 20 \( \mu \)M [13C] \( \sim \) malonyl-CoA substrate concentrations. The activity of FASN from Spot14-null MEGs relative to control lysates containing Spot14 (1.5 ± 0.16 nmol vs. 2.4 ± 0.31 nmol [13C] fatty acids synthesized sec\(^{-1}\) mg\(^{-1}\); Fig. 7A) and, the reaction velocity in control lysates containing Spot14 was significantly increased 1.6-fold relative to Spot14-null lysates (Fig. 7B). When recombinant Spot14 was added to the Spot14-null lysates in 40 molar excess, a 20% increase in the catalytic rate was observed (1.9 ± 0.08 nmol vs. 1.5 ± 0.16 nmol [13C] fatty acids synthesized sec\(^{-1}\) mg\(^{-1}\); Fig. 7A); however, the reaction velocity increase did not reach significance \( (P = 0.08; \text{Fig. 7B}) \). These observations suggest that recombinant Spot14 added back to Spot14-null MEC lysates could enhance the rate of native FASN catalysis relative to Spot14-null lysates; however, alternative forms of regulation may influence FASN catalysis because Spot14 add back could not fully restore activity.

**DISCUSSION**

Analysis of Spot14-null and transgenic mice overexpressing Spot14 during lactation was conducted to investigate the molecular function of Spot14 in a tissue type that is devoid of confounding influences of Spot14-R. In contrast to other mouse models with severe lactation failure, such as the DGAT 1-null, prolactin receptor +/−, STAT5a-null, ELF5 +/-/H1002, alpha-lactalbumin-null, and galanin-null mice (58–63), the lactation phenotype in Spot14-null dams is mild and no lactation defect was observed for MMTV-Spot14 overexpressing dams (Fig. 1A, B). Consistent with this mild phenotype is the observation that Spot14-null dams produce milk and lactate sufficiently (Fig. 2B), but that the total lipid content of the milk is reduced (Fig. 2A). Pup growth deficit was rescued simply by providing a HFD to the lactating dams (Fig. 1A, HFD). While reduced milk TAG would be consistent with lower levels of DGAT1, no differences in DGAT1 mRNA (supplementary Fig. II) or protein levels (Fig. 4) were observed. The MCFA (C = 10:0 to 14:0) unique to mammary epithelium de novo fatty acid synthesis were significantly reduced by 60% in Spot14-null milk (Fig. 3A). Importantly, none of the mRNA and total protein levels for de novo fatty acid synthesis pathway enzymes were lower in Spot14-null MECs, that could account for the reduction of MCFA, revealing that Spot14 does not regulate mRNA or protein abundance in lactating MECs.
We have found that Spot14 stimulates the in vitro activity of recombinant FASN (Fig. 6A) and native FASN present in MEC cytosolic lysates (Fig. 7A). The in vitro results are consistent with the in vivo observations from the Spot14-null and the MMTV-Spot14 overexpressing mice. Loss of Spot14 function resulted in significantly decreased MCFAs (Fig. 3A), while MMTV-Spot14 overexpressing dams produced milk with significantly increased MCFAs (Fig. 3B). MCFAs were increased in the MMTV-Spot14 dams at lactation day 4, a time when the de novo pathway is not yet maximal (13), indicating that Spot14 overexpression stimulated de novo fatty acid synthesis without influencing levels of preformed fatty acids. Moreover, overexpression of Spot14 increased MCFA and altered the composition of fatty acids present in the milk TAG (Fig. 3B) without affecting the total amount of TAG in the milk (Fig. 2A). Together, these data are in agreement with work that showed the in vivo rate of $^3$H$_2$O incorporation into de novo synthesized lipids was decreased 2.5-fold in mammary glands of Spot14-null dams (39). Collectively, these experiments suggest the lactating mammary epithelium has evolved a tissue-specific molecular function for Spot14 as an enhancer of FASN catalysis.

Kim et al. (43) demonstrated that Spot14-R facilitated multimerization of ACC in vitro; however, they observed no Spot14 effect on the assembly or activity of ACC. We could not rule out the possibility that Spot14 may function to facilitate dimer formation or stabilization of FASN to stimulate catalysis in vivo. Nondenaturing native gels indicated that FASN existed as a dimer in both control and Spot14-null samples (supplementary Fig. IV). With the exception of TE2, the independent enzyme that converts product specificity of FASN from $\text{C} = 16:0$ to MCFA (15, 64), little is understood regarding proteins that might alter catalytic rate and fatty acid product distribution. Consistent with increased product release, the presence of Spot14 in vitro resulted in an approximately 4-fold increase in $\text{C} = 14:0$ (Fig. 6D) and a nearly 2-fold decrease of apparent $K_m$ (Fig. 6C). Spot14 itself does not have hydrolase activity using the model TE2 substrate myristoyl-CoA (supplementary Fig. IIID), and Spot14 did not bind acetyl- or malonyl-CoA substrates (not shown). These results suggest that Spot14 might augment release of fatty acids via the resident thioesterase domain of FASN. Nagget et al. (65) showed that the resident thioesterase domain from FASN relies entirely on its covalent linkage to FASN for function, as the recombinant “free” thioesterase domain had no activity toward S-acyl-FASN. We speculate that Spot14 could foster a more productive interaction of the resident thioesterase domain with the acyl carrier domain of FASN to stimulate release of its fatty acid products.

This observation could also indicate that expression of Spot14 in the lactating MEC may function to support the intervention of TE2 to liberate high quantities of MCFAs, rather than acting predominantly with the resident TE domain of FASN. At the present time, we cannot distinguish between Spot14 interaction with endogenous resident TE domain of FASN or if Spot14 enhances TE2 interaction to release MCFA from FASN in vivo. FASN was immunoprecipitated from control and Spot14-null mammary gland lysates, but neither Spot14 nor TE2 were detected in the FASN pull-down, indicating that any potential FASN/Spot14/TE2 complexes associate transiently. That TE2 was not detected in FASN pull-down was consistent with previous work that failed to observe a stable FASN/TE2 complex from rat mammary gland lysates (66). In that study, Mikkelsen et al. (66), used a clever experimental approach to demonstrate that a stable complex formed only when FASN was acylated on the 4′-phosphopantetheine thiol group of the ACP domain that was actively engaged in chain elongation, after which, the complex rapidly dissociated following acyl hydrolysis. It is possible that active acyl chain elongation on FASN is also required for association of Spot14 with FASN, but this hypothesis was not addressed in the present studies.

When recombinant Spot14 was added back to null cytosolic lysates, the catalytic rate was not fully restored to that from control lysates containing native Spot14 (Fig. 7A). Given the increased phosphorylation of ACY and ACC in Spot14-null MECs (Fig. 5C), we hypothesized that FASN could be more highly phosphorylated in Spot14-null MECs during lactation. Phosphorylation of FASN at Thr1029 and Thr1033 (mouse) by mTORC1 has been shown to inhibit FASN activity (12); however, no significant difference in total phosphorylated threonine was observed using an anti-phosphothreonine antibody (supplementary Fig. IVB). This observation indicates that the regulation of FASN by mTORC1 may not be a feature in lactating MECs as was observed in liver (12). Site-specific antibodies are needed to properly analyze the phosphorylation status of FASN as it pertains to enzyme activity. The fact that addition of recombinant Spot14 to null lysates did not restore FASN catalysis to levels observed in control lysates suggests that other proteins and/or posttranslational modifications may regulate FASN in vivo to preserve the dynamic balance between preformed fatty acids and those synthesized by FASN.

It is interesting to note that none of the aqueous intermediate metabolites of glycolysis and the TCA cycle were decreased in Spot14-null MECs (supplementary Table I), consistent with the previous observation that enzyme activity upstream of FASN using in vitro assays was intact in the Spot14-null mammary gland (39). Furthermore, a small number of these metabolites were significantly increased in the Spot14-null MECs, including acetyl-CoA, aromatic amino acids, glutamate, pyruvate, and succinate; while malonyl-CoA trended higher in Spot14-null mammary glands (supplementary Table I). This observation indicates that ample substrate is present in vivo for FASN catalysis, and supports the conclusion that Spot14 functions to enhance the rate of FASN catalysis during lactation in vivo. Lipogenesis in the lactating mammary gland represents a highly specialized function that maintains the dynamic balance of preformed fatty acids with activity of the de novo fatty acid synthesis pathway to preserve a relatively constant milk fat content unique to each mammalian species. Our findings implicate Spot14 as a direct protein enhancer of FASN, enhancing MCFA synthesis, a function
that appears to be unique and critical to the mammary gland during lactation.\textsuperscript{54}

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