Fetuin-A modulates lipid mobilization in bovine adipose tissue by enhancing lipogenic activity of adipocytes

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ABSTRACT

Fetuin-A (FetA) is an adipokine and free fatty acid (FFA) carrier linked to adipose tissue (AT) function in monogastrics and ruminants. In dairy cows, plasma and AT FetA decrease after parturition, coinciding with reduced lipogenesis and increased lipolysis. In monogastrics, FetA enhances lipogenesis, but its role on lipid mobilization of ruminants is unclear. We hypothesized that FetA modulates lipid mobilization in bovine AT by enhancing the lipogenic activity of adipocytes. Our objective was to determine the effects of FetA on lipogenesis and lipolysis in cultured primary adipocytes from dairy cows. Preadipocytes from the tailhead subcutaneous AT depot were induced to differentiate in a 7-d coculture in vitro model. The effects of FetA on lipolytic responses of adipocytes were evaluated after a 2-h β-adrenergic stimulation with 1 µM isoproterenol (ISO) alone or combined with 0.1 mg/mL of FetA (FetA+ISO), and in cells treated with medium alone (CON) or with 0.1 mg/mL of FetA (FetA). Lipogenic responses of adipocytes treated with CON or FetA from d 5 to 7 of differentiation were assessed by fatty acid (FA) uptake quantification and triacylglycerol (TAG) accumulation, and the gene and protein expression of lipogenic markers. Bovine adipocytes abundantly expressed FetA gene and protein and secreted 48 ± 3.5 ng/DNA relative fluorescence units (RFU). Adrenergic stimulation with ISO increased lipolysis compared with CON, as reflected in the release of glycerol (0.12 ± 0.04 vs. 0.04 ± 0.02 nM/DNA RFU) and FFA (15 ± 13 vs. 6.2 ± 2.4 nM/DNA RFU). Lipolysis induced by ISO was attenuated by the addition of FetA (FetA+ISO) as reflected by lower glycerol (0.06 ± 0.04 nM/DNA RFU) and FFA (5.7 ± 2.7 nM/DNA RFU) release compared with ISO alone. Compared with CON, FetA enhanced lipogenic responses as demonstrated by higher FA uptake and increased accumulation of TAG. Exposure to FetA upregulated 1-acylglycerol-3-phosphate acyltransferase-2 (AGPAT2) gene expression and protein content, as well as its activity. Adipocytes exposed to FetA increased the secretion of the metabolite of AGPAT2, phosphatidic acid. In conclusion, FetA attenuates lipolytic responses and enhances lipogenesis in bovine adipocytes. The upregulation of the rate-limiting lipogenic enzyme AGPAT2 by FetA suggests a potential pathway by which this adipokine promotes TAG synthesis in adipocytes. These findings suggest that FetA is a potential target for lipid mobilization modulation in AT of dairy cows.

Key words: adipocyte, dairy cow, lipogenesis, lipolysis

INTRODUCTION

Increased lipid mobilization in adipose tissues (AT) is the major metabolic adaptation to negative energy balance in periparturient dairy cows. During lipolysis, one molecule of triacylglycerol (TAG) is broken down into 3 free fatty acid (FFA) molecules and 1 molecule of glycerol through the activity of adipose triglyceride lipase, hormone-sensitive lipase, and monoglyceride lipase. Released FFA are either re-esterified into TAG within the adipocytes or used as an energy source elsewhere in the body (Vernon and Pond, 1997). Lipogenesis involves the biosynthesis of TAG from fatty acids (FA) either synthesized from FFA released from blood TAG or de novo within the adipocytes. Glycerol phosphate pathway is the major pathway for de novo synthesis of TAG by adipocytes, which consists in a stepwise addition of fatty acyl groups catalyzed by distinct enzymes: glycerol-3-phosphate acyltransferase (GPAT), 1-acylglycerol-3-phosphate acyltransferase (AGPAT), lipins (phosphatidate phosphatases), and diacylglycerol acyltransferase (DGAT) to finally form TAG (Takeuchi and Reue, 2009). Adipocyte TAG reserves serve as the main source of energy to offset negative energy balance in periparturient cows (McNamara, 1995).

Around parturition, lipolysis is enhanced and lipid synthesis is reduced in adipocytes by physiological factors such as decreased plasma insulin and glucose
concentrations, impaired insulin sensitivity in AT and other peripheral tissues, and increased concentrations of catecholamines, growth hormone, and glucocorticoids (Bauman and Currie, 1980). Also, a decrease in the transcription and activity of key lipogenic enzymes limits the ability of adipocytes to synthesize FA and TAG thus enhancing FFA release from AT (McNamara, 1995). The relative balance between lipolysis and lipogenesis in AT controls the release of FFA from adipocytes and is critical to ensure a successful transition from gestation to lactation in dairy cows. Dysregulated lipid mobilization leads to a massive release of FFA into circulation that, coupled with an excessive accumulation of lipids in the liver and exacerbated pro-inflammatory responses, predisposes periparturient dairy cows to health events (Vernon and Houseknecht, 2000; Contreras et al., 2018). Currently, a major challenge in the field of periparturient dairy cow management is the adequate regulation of AT lipolysis and lipogenesis rates to ensure that released FFA are fully metabolized for energy needs, thus avoiding the detrimental lipid accumulation in blood and tissues.

We recently determined the dynamics of a novel adipokine, fetuin-A (FetA; α-2-Heremans-Schmid glycoprotein; encoded by AHSG gene), in serum and AT of dairy cows during the periparturient period (Strieder-Barboza et al., 2018). Our results indicated that FetA is a negative acute-phase adipokine in the subcutaneous AT and that its plasma concentrations were negatively associated with markers of AT lipid mobilization including BHB and BCS loss. In the same study, we reported FetA gene and protein expression in bovine adipocytes, suggesting a potential autocrine role for this adipokine (Strieder-Barboza et al., 2018). Even though FetA was shown to be expressed by bovine AT, its secretion by primary adipocytes has not yet been reported. In humans and mice, FetA secretion by visceral and subcutaneous AT was augmented during obesity and in animals fed a high-fat diet (Jialal et al., 2015; Pérez-Sotelo et al., 2017). In monogastric adipogenic models, this effect is probably related to the lipogenic properties of FetA as this protein is a carrier of lipids in plasma and facilitates the incorporation of exogenous FA into intracellular TAG (Kumbla et al., 1989; Cayatte et al., 1990). Because low FetA in plasma and AT coincided with low lipogenesis and high lipolysis rates in periparturient cows, and given the fact that FetA was associated with pro-lipogenic states in nonruminant species, we hypothesized that FetA modulates lipid mobilization by enhancing lipogenic activity in bovine adipocytes. Our objectives were to determine the effect of FetA on lipogenesis and lipolysis, and to identify potential mechanisms by which this adipokine modulates lipid mobilization of bovine adipocytes.

MATERIALS AND METHODS

All animal procedures were approved by the Michigan State University Animal Care and Use Committee.

Tissue Collection and Processing

Subcutaneous AT from the tailhead depot from 7 nonlactating, nongestating, multiparous Holstein dairy cows was collected in Krebs-Ringer modified buffer supplemented with HEPES 10 mM (pH = 7.4) at a local slaughterhouse as previously described (Strieder-Barboza et al., 2018). In brief, AT (1 g) was digested with 5 mL of collagenase type II solution (2 mg/mL; Worthington Biochemical, Lakewood, NJ) and then centrifuged (800 × g, 10 min, 4°C) to separate the primary adipocytes from the stromal vascular fraction (SVF). Primary mature adipocytes were washed in 5 mL of Krebs-Ringer modified buffer with 4% BSA (Millipore-Sigma, Sigma-Aldrich, St. Louis, MO), centrifuged, and then retained for use in transwell inserts for inductions using a coculture protocol (Thelen et al., 2018). The SVF was then sequentially filtered through 100- and 40-μm cell striainers (Falcon, Corning, Corning, NY) and centrifuged. The resulting cell pellet was resuspended and incubated in erythrocyte lysis buffer. After another centrifugation, resultant cells were resuspended in basal preadipocyte medium containing Dulbecco’s modified Eagle’s medium: F12 (Corning), 10% fetal bovine serum (Corning), 2 mmol/L of L-glutamine (Corning), 1% (vol/vol) antibiotic-antimycotic (Corning), 44.05 mmol/L of sodium bicarbonate (Sigma-Aldrich, St. Louis, MO), 100 μmol/L of ascorbic acid (Sigma-Aldrich), 33 μmol/L of biotin (Sigma-Aldrich), 17 μmol/L of pantothenate (Sigma-Aldrich), and 20 mmol/L of HEPES (Corning) and incubated at 37°C in a humidified atmosphere with 5% CO2 with medium replacement every 48 h (Strieder-Barboza et al., 2018). Preadipocytes were obtained by outgrowth of plastic-adherent cells from the SVF cells after 2 serial passages in culture flasks (Corning).

Cell Induction and Differentiation

Expanded preadipocytes populations were seeded in 6-, 12-, or 24-well plates (Corning) and allowed to proliferate to confluency. Preadipocytes were induced to differentiate after 48 h at 100% confluency (d 0) using a coculture model (Thelen et al., 2018). Briefly, 900 mature adipocytes/cm² were placed in 0.4-μm transwell inserts (Greiner Bio-One, Kremsmünster, Austria) over the attached preadipocytes for the first 5 d of differentiation. For differentiation induction, cells were incubated with basal medium supplemented
with 10% fetal bovine serum, 5 µmol/L of troglitazone (Cayman Chemical, Ann Arbor, MI), 0.5 mmol/L of 2-isobutyl-1-methylxanthine (AdipoGen Life Sciences, San Diego, CA) and the following reagents from Sigma Aldrich: 5 µg/mL of insulin, 10 mM acetate, and 1 µmol/L of dexamethasone for 7 d. The 2-isobutyl-1-methylxanthine and dexamethasone were used only during the first 48 h of induction, and medium changes were performed every 48 h. Adipocyte lipid accumulation was assessed quantitatively in triplicates per experimental unit using a 96-well plate for the AdipoRed assay (Lonza, Allendale, NJ) and a Synergy H1 Microplate Reader (Biotek, Winooski, VT). Fetuin-A dose of 0.1 mg/mL was used for all metabolic assays and was established based on previous studies with adipocytes (Heinrichsdorff and Olefsky, 2012; Pal et al., 2012), as well as on a dose response assay (0, 50, 100, 250, 500, and 1,000 µg/mL of FetA) having the gene expression of lipogenic and inflammatory genes as the outcome.

**Lipolysis Assay**

Induction of lipolysis was performed using the non-selective β-adrenergic receptor agonist isoproterenol (ISO; Sigma-Aldrich). Briefly, cultured adipocytes (n = 6) were removed from 6-well plates (Corning) using trypsin (Thermo Fisher Scientific, Waltham, MA), seeded in triplicate at 1 × 10⁵ cells/well in black well-clear bottom 96-well plates (Nunc, Roskilde, Denmark), and allowed to attach for 4 h at 37°C in a humidified atmosphere with 5% CO₂. Next, the reagent of the QBT Fatty Acid Uptake assay containing a fluorescent-labeled FA analog was added to the cells (100 µL/well). Kinetic uptake was measured every 20 s for 1 h using a Synergy H1 Microplate Reader (Biotek). Triplicate values per sample were averaged, subtracted from blank, and divided by basal values (CON) to calculate the area under the curve for each treatment. Values are expressed as fold change over CON.

**Fatty Acid Uptake Assay**

Fatty acid uptake analysis was performed using the kinetic QBT Fatty Acid Uptake Assay (Molecular Devices, Sunnyvale, CA). Cultured adipocytes (n = 6) were detached by trypsinization and seeded as described for the lipolysis assay. Viability after trypsin detachment was assessed by Trypan blue exclusion. Adipocytes were then serum-starved overnight and pre-incubated with 0.1 mg/mL of FetA, 10 nM insulin (insulin, positive control; Sigma-Aldrich) or serum-free basal preadipocyte medium (CON; basal FFA uptake) during 30 min at 37°C in a humidified atmosphere with 5% CO₂. Next, the reagent of the QBT Fatty Acid Uptake assay containing a fluorescent-labeled FA analog was added to the cells (100 µL/well). Kinetic uptake was measured every 20 s for 1 h using a Synergy H1 Microplate Reader (Biotek). Triplicate values per sample were averaged, subtracted from blank, and divided by basal values (CON) to calculate the area under the curve for each treatment. Values are expressed as fold change over CON.

**Triacylglycerol Accumulation**

Intracellular TAG accumulation in adipocytes was assessed by Adipogenesis Assay Reagent (AdipoRed, Lonza) and the lipid droplet staining HCS LipidTox (Life Technologies) analyzed through confocal microscopy imaging. Cultured adipocytes were incubated with basal adipocyte differentiation medium supplemented with 0.1 mg/mL of FetA for 48 h (d 5 to 7 of the differentiation protocol). Basal TAG accumulation was determined in adipocytes incubated with basal differentiation medium without addition of FetA (CON).

**Lipid Droplet Staining and Confocal Microscopy Imaging.** After 48 h-incubation with or without FetA, adipocytes (n = 8) were seeded in duplicates/treatment per cow in a glass bottom 24-well plate (Corning) at a concentration of 20,000 cells/cm². Then DAPI (NucBlue, Life Technologies) and HCS LipidTox with Alexa Fluor 594 (Life Technologies) were used to visualize adipocytes’ nuclei and intracellular TAG, respectively. These dyes were used according to
the manufacturer’s protocols. Briefly, adipocytes were fixed with 4% paraformaldehyde (Electron Microscopy Sciences, Hatfield, PA) in PBS (1× PBS) for 20 min at room temperature. Two hundred fifty microliters of HCS LipidTOX neutral lipid stain 1:200 in 1× PBS was added to each well (Corning) and incubated at room temperature for 30 min. The buffer was removed and 200 µL of DAPI (300 nM in 1× PBS) was added to each well and incubated at room temperature for 5 min. Images were acquired using an Olympus Fluoview 1000 Confocal Laser Scanning Microscope (Olympus America Inc., Center Valley, PA) configured on an IX81 inverted microscope and FV10-ASW software (version 4.2.3.6) using a PLAPON 60×/1.42 oil objective. Alex Fluor 594 fluorescence (577 nm excitation/609 nm emission) was excited with the 543 nm HeNe gas laser, detected using a BP 560IF emission filter, and displayed in red color. The DAPI fluorescence (358 nm excitation/461 nm emission) was excited using the 405 nm diode laser, detected using a 430 to 470 nm band pass emission filter, and displayed in blue color. Images were obtained using sequential single confocal XY scan mode. Control images for TAG accumulation included preadipocytes and nonstained adipocytes. Relative fluorescence intensity of intracellular TAG per cell was determined by ImageJ (https://imagej.net/Welcome) software and calibrated by cell number.

**Adipogenesis Assay Reagent.** AdipoRed is a solution of Nile red, a hydrophobic stain, that fluoresces when partitioned into lipid droplets of differentiated adipocytes. Briefly, adipocytes (n = 5) in triplicates/treatment per cow in 24-well plates (Corning) were washed once with 1× PBS, incubated with AdipoRed assay reagent for 20 min at room temperature, and then analyzed for fluorescence intensity (excitation/emission = 485/572 nm) using a Synergy H1 Microplate Reader (Biotek). Preadipocytes from each experimental unit on d 0 served as negative controls. Relative fluorescence units of adipocytes were calibrated by their corresponding preadipocyte RFU and are presented as fold change over preadipocyte lipid accumulation.

**Gene Expression Analysis of Adipogenesis and Lipogenesis Markers**

After treating adipocytes (n = 5) for 48 h with or without 0.1 mg/mL of FetA, culture medium was removed, and cells were rinsed twice with ice-cold 1× PBS. Adipocytes’ RNA was extracted using Promega simplyRNA Cells Kits (catalog no. AS1390, Promega, Madison, WI) in the Maxwell RSC Instrument (Promega) as described previously (Strieder-Barboza et al., 2018). This protocol started with the addition of 100 µL of 1-thioglycerol/homogenization solution to each of the 2-wells/treatment per cow of a 12-well plate with adipocytes and then transferred to a microfuge tube. Next, 200 µL of lysis buffer was added and homogenate was vortexed and then placed in Maxwell RSC Cartridges, which were previously loaded with 10 µL of DNase I. Purity, concentration, and integrity of AT and cells’ RNA were evaluated using an Agilent Bioanalyzer 2100 (Agilent Technologies, Santa Clara, CA). All samples had an RNA integrity number ≥6. Conversion to cDNA was performed using the Applied Biosystems High Capacity cDNA Archive Kit (Applied Biosystems, Foster City, CA). Transcriptional studies were performed on the cDNA samples using quantitative PCR reactions on QuantStudio 7 Flex (Applied Biosystems). All qPCR reactions were performed in duplicates, and no-template controls were included on each plate for each TaqMan gene expression assay.

Gene expression data of 5 endogenous control genes (B2M, EIF3K, GAPDH, RPLP0, and RPS9) were analyzed using qBase+ analysis software, which calculates the stability of endogenous control genes (M-value). Following qBase+ analysis of gene expression data, endogenous control genes EIF3K and RPS9 were ranked best. The Cq values of the target genes (ADIPOQ, AGPAT2, AHSG, CD36, CEBPβ, DGAT1, DGAT2, FABP4, FATP1, GAPT1, GAPT2, LPIN1, and PPARG) were converted to normalized relative gene expression as described previously (Hellemans et al., 2007; Contreras et al., 2017a). The quantitative PCR assays were conducted in duplicates with TaqMan gene expression assays from Applied Biosystems, except FetA (AHSG), which was provided by Integrated DNA Technologies (IDT, Coralville, IA) as described previously (Strieder-Barboza et al., 2018; Supplemental Table S1; https://doi.org/10.3168/jds.2018-15808).

**Western Blotting**

Western blots were performed as previously described (Strieder-Barboza et al., 2018). Protein from cultured adipocytes (n = 8) was extracted from 3 wells/treatment per cow of a 12-well plate using ice-cold RIPA buffer (Teknova, Hollister, CA) containing protease (Roche, San Francisco, CA) and phosphatase (Thermo Fisher Scientific) inhibitors. Estimation of protein content was carried out using the Pierce BCA Protein Assay Kit (Thermo Fisher Scientific). Fetuin-A content was analyzed using reducing conditions. Briefly, samples were added to a reducing buffer containing 10 mmol/L of dithiothreitol and 5% β-mercaptoethanol and denatured at 95°C for 4 min. Twenty micrograms of protein was separated by electrophoresis on a 4 to
20% SDS-PAGE gel and transferred to a polyvinylidene difluoride membrane. The membrane was blocked in Tris-buffered saline solution with 0.01% Tween-20 (TBST) and 5% fat-free milk. Membranes were incubated with a conjugated polyclonal rabbit anti-bovine AGPAT2 antibody (C-terminal region catalog no. ARP4G363_P050, Aviva Systems Biology Corporation, San Diego, CA) in 1 µg/mL concentration in TBST-5% milk for 16 h at 8°C. αβ-Tubulin (1:1,000; catalog no. 2148; Cell Signaling Technology, Danvers, MA) served as loading control for adipocyte protein. Membranes were then exposed to horseradish peroxidase substrate (catalog no. WBLUR0100, Millipore, Darmstadt, Germany) and visualized by chemiluminescence using the ChemiDoc Touch Imaging System (Bio-Rad, Hercules, CA). Band densitometry was determined using the Image Lab software (Bio-Rad). Values of AGPAT2 protein content are expressed as means of relative band density using αβ-tubulin band intensity values as calibrator.

**Phosphatidic Acid Analysis**

Phosphatidic acid (PA) concentrations in adipocytes were measured using the enzymatic fluorometric PicoProbe Phosphatidic Acid Assay Kit (catalog no. K748, BioVision Inc., Milpitas, CA) following the manufacturer’s instructions. Briefly, after treating adipocytes (n = 5) with or without 0.1 mg/mL of FetA for 48 h, culture medium was removed, and cells from 3 wells/treatment/cow of a 12-well plate were rinsed twice with ice-cold 1× PBS, scraped from cell culture plates, and collected in 1× PBS. For lipid extraction, 1 × 10⁶ adipocytes were mixed with 1 mL of assay buffer (BioVision) and sonicated using an ultrasonic liquid processor (Misonix, Farmingdale, NY) for 2 min at 4°C. Protein content was estimated at this point using a BCA Protein Assay Kit (Thermo Fisher Scientific). Following this, 3.75 mL of chloroform/methanol/12 N HCl (2: 4: 0.1 vol/vol; Sigma-Aldrich) was added to the cell homogenate and mixed thoroughly. Then, 1.25 mL of chloroform was added to the solution, vortexed for 30 s, and added to 1.25 mL of 1 M NaCl (Sigma-Aldrich). Followed by centrifugation, the lower organic layer containing solubilized lipids was collected and transferred to a glass tube. Chloroform was evaporated in a vacuum concentrator (Savant SPD121P, Thermo Fisher) for 2 h at 40°C. Before performing the analysis, samples were solubilized in 50 µL of 5% Triton X-100 solution. For PA measurement, 10 µL of sample was added to a 96-well plate and incubated for 1 h at 45°C with a compound that hydrolyzes PA. Samples were then incubated for 30 min at 37°C in the presence of a developer and enzyme mix that converts a nonfluorescent probe to a fluorescent product (Ex/Em = 535/587 nm) that can be quantified (Synergy H1 Microplate Reader, Biotek). Concentrations of PA were calculated using the manufacturer’s PA standard (BioVision) and L-A-phosphatidic acid sodium (P9511, Sigma-Aldrich). Phosphatidic acid values were obtained by subtracting the background RFU from each sample and then calibrated by the total protein concentration (nmol of PA per ng of protein).

**Statistical Analysis**

Data were analyzed using JMP Statistical Software (SAS Institute Inc., Cary, NC). Normality of the variables was checked using the Kolmogorov-Smirnov test (P < 0.05). Nonnormally distributed variables (FFA concentrations in lipolysis assay, FA uptake assay, adipocytes AGPAT2 protein content) were natural logarithm transformed. One-way ANOVA pairwise comparisons were performed using the Tukey’s post hoc test. Mean differences were considered significant when P ≤ 0.05 and tendencies when P < 0.10. Average values throughout the text and figures are showed as mean ± SEM.

**RESULTS**

**Bovine Adipocytes Abundantly Secrete Fetuin-A**

The capacity of primary bovine adipocytes to express and secrete FetA into the medium was assessed. We determined that these cells secrete 48 ± 3.5 ng of FetA/DNA RFU per 2-h incubation in serum-free basal medium and express AHSG and FetA protein at a similar level as those observed in AT of periparturient cows (P > 0.05; Supplemental Figure S2; https://doi.org/10.3168/jds.2018-15808).

**Fetuin-A Attenuates Lipolytic Responses in Bovine Adipocytes**

After assessing the ability of bovine adipocytes to secrete FetA, we evaluated the effect of FetA on lipolytic responses of bovine adipocytes. β-Adrenergic stimulation with ISO for 2 h increased the release of glycerol (0.12 ± 0.04 vs. 0.04 ± 0.02 nM/DNA RFU, P = 0.003) and FFA (15 ± 13 vs. 6.2 ± 2.4 nM/DNA RFU, P = 0.04) compared with CON (Figure 1). Remarkably, ISO-induced lipolysis was attenuated by FetA (FetA+ISO) as reflected in lower glycerol (0.06 ± 0.04 nM/DNA RFU, P = 0.02) and FFA (5.7 ± 2.7 nM/DNA RFU, P = 0.01) release compared with ISO alone (Figure 1).
FETuin-A and Lipid Mobilization in Bovine Adipocytes

Fetuin-A Enhances Lipogenesis in Bovine Adipocytes

Lipolysis and lipogenesis are continuous processes occurring simultaneously within adipocytes. Thus, we evaluated the effects of FetA on both lipolytic and lipogenic activity. We assessed lipogenesis by quantifying adipocyte FA uptake and accumulation of intracellular TAG. The FetA exposure during 1.5 h (0.5 h pre-incubation and 1-h kinetic assay) increased FA uptake by 1.5 times compared with CON (P = 0.02; Figure 2). The effect of FetA on FA uptake was similar to that of insulin (P = 0.65; Figure 2B), increasing FA uptake by 1.7 times compared with CON (P = 0.004; Figure 2B). In agreement with the higher incorporation of FA, we observed that adipocytes treated with FetA for 48 h increased lipogenic capacity compared with CON (P < 0.05). Adipocytes in the FetA group had a higher accumulation of TAG as quantified with the neutral lipid fluorescent stain HCS LipidTOX (P = 0.003; Figure 3A–3C) and with the quantitative AdipoRed assay (P = 0.04; Figure 3D–3E).

Fetuin-A Upregulates AGPAT2 Expression and Activity in Bovine Adipocytes

To explore potential mechanisms by which FetA enhances lipogenic function in bovine adipocytes, we evaluated the effect of the adipokine on key regulators of adipogenesis (CEBPB, ADIPOQ, and PPARG), FA uptake (FABP4, FATP1, and FAT/CD36), and lipogenic enzymes controlling de novo TAG synthesis in adipocytes, such as AGPAT2, DGAT1, DGAT2, GPAT1, GPAT2, and LPIN1. We observed that FetA increased the transcription of AGPAT2 (P = 0.02; Figure 4A) and AHSG (P = 0.05; Figure 4B) and tended to increase the expression of CEBPB (P = 0.07; Figure 4C) and FATP1 (P = 0.07; Figure 4D). No differ-

Figure 1. Effect of fetuin-A (FetA) on β-adrenergic induced lipolysis in bovine adipocytes (n = 6). Concentrations of (A) glycerol and (B) free fatty acids (FFA) released in culture medium after a 2-h lipolysis induction with 1 µM isoproterenol (ISO). Concentrations of glycerol (nM) and FFA (nM) were standardized by the cellular DNA content [DNA relative fluorescence units (RFU)] measured by CyQUANT (Life Technologies, Carlsbad, CA). Values are showed as mean ± SEM. *Statistically different (P < 0.05). CON = control, no reagents added.

Figure 2. Effect of fetuin-A (FetA) on fatty acid (FA) uptake by bovine adipocytes (n = 6). (A) Example of the kinetic QBT FA uptake (Molecular Devices, Sunnyvale, CA) output from bovine adipocytes over 1-h assay. RFU = relative fluorescence units. (B) Area under the curve (AUC) from FA uptake assays of bovine adipocytes treated with 0 (CON) or 0.1 mg/mL of FetA (FetA) or 10 nM insulin (insulin). Bovine adipocytes were treated and pre-incubated with treatments for 30 min at 37°C before the assay. Values are expressed as AUC using CON FA uptake as the calibrator (means ± SEM). Bars with different letters (a,b) are significantly different (P ≤ 0.05).
ence was observed in the gene expression of PPARG, ADIPOQ, DGAT1, DGAT2, GPAT1, GPAT2, LPIN1, FAT/CD36, and FABP4 between FetA and CON (P > 0.05). In agreement with the increase in AGPAT2 gene expression, AGPAT2 protein content was also increased by FetA compared with CON (P = 0.02; Figure 4B).

The enzymatic activity of AGPAT2 during TAG synthesis transfers a FA to the lysophosphatidic acid molecule to produce PA. We assessed the effect of FetA on AGPAT2 activity by quantifying the concentrations of PA in bovine adipocytes treated with 0.1 mg/mL of FetA for 48 h. Adipocytes treated with FetA syn-

![Figure 3. Effect of fetuin-A (FetA) on triacylglycerol accumulation in bovine adipocytes. (A) Laser scanning confocal microscopy imaging (60×) of lipid droplets in cultured bovine adipocytes (n = 8) treated with 0 (CON; bottom panel) or 0.1 mg/mL of FetA (FetA; upper panel). Scale bars: 20 µm. (B) Preadipocytes and nonstained cells served as controls. Lipid droplets of bovine adipocytes were stained with Alexa Fluor 594 (HCS LipidTox, Life Technologies, Carlsbad, CA; red fluorescence) and nuclei were stained with DAPI (NucBlue, Life Technologies; blue fluorescence). Scale bars: 20 µm. (C) Plotted relative fluorescence units (RFU) measured through ImageJ (https://imagej.net/Welcome) software (means ± SEM) and calibrated by number of cells per image. *P ≤ 0.05. (D) Relative fluorescent units of AdipoRed (Lonza, Allendale, NJ) assay analysis in cultured bovine adipocytes (n = 5) treated with 0 (CON) or 0.1 mg/mL of FetA (FetA), and respective preadipocytes (PREADIPO) on d 0. Bars with different letters (a–c) are significantly different (P ≤ 0.05). (E) Representative microscopy images (20×) of lipid droplets (dark red dots) in bovine adipocytes treated with 0 (CON) or 0.1 mg/mL of FetA (FetA), and preadipocytes (PREADIPO) assayed with AdipoRed. Scale bars: 100 µm.
thetized more PA (0.27 ± 0.002 nmol/ng of protein) compared with CON adipocytes (0.20 ± 0.01 nmol/ng of protein; \( P = 0.060 \); Figure 4F).

**DISCUSSION**

The balance between lipogenesis and lipolysis in AT of periparturient dairy cows is critical to ensure a healthy and successful transition to lactation. Thus, reducing lipolysis or promoting lipogenesis are important therapeutic alternatives to improve periparturient cows’ health. In our previous study, we observed that the expression of FetA and other lipogenic markers was downregulated in subcutaneous AT after parturition (Strieder-Barboza et al., 2018). Also, previous studies in nonruminants have consistently demonstrated the
strong lipogenic properties of FetA (Cayatte et al., 1990; Pérez-Sotelo et al., 2017). To date, however, the mechanisms by which FetA increases TAG synthesis in adipocytes have not been characterized. In this study, we report for the first time that FetA modulates lipid mobilization in isolated and cultured bovine adipocytes and identify AGPAT2 activity as the potential mechanism for the lipogenic responses induced by FetA.

Previous studies demonstrate abundant mRNA and protein expression of FetA in AT of periparturient, mid-, and late-lactation dairy cows, as well as in cultured bovine adipocytes (Zachut et al., 2017; Strieder-Barboza et al., 2018). However, none of these studies had identified the source of FetA within AT. Here we demonstrate that primary adipocytes from subcutaneous AT of dairy cows not only express but also secrete FetA. Even though secretome studies of rodents’ visceral and gonadal AT also detected significant amounts of FetA (Roca-Rivada et al., 2011), this is the first report of FetA secretion by bovine adipocytes. The synthesis of FetA is dynamic and depends on the metabolic status. In humans and rodents, FetA secretion decreased during fasting, weight loss, and anorexia and increased during weight gain and obesity (Jialal et al., 2015; Pérez-Sotelo et al., 2017). In dairy cows, serum concentrations and AT gene and protein expression of FetA decreased during early lactation and coincided with negative energy balance state (Strieder-Barboza et al., 2018). The changes in the FetA expression and secretion patterns in the AT of rodents, bovines, and humans during lipid mobilization suggest a role for FetA in the modulation of lipogenesis and lipolysis in adipocytes.

In the present study, FetA attenuated lipolytic responses induced by β-adrenergic stimulation (ISO). Our results suggest that the effect of FetA during induced-lipolysis in bovine adipocytes was insulin-independent because no insulin was added to the medium before and during β-adrenergic stimulation. Previous studies indicate that FetA may control insulin signaling in AT and is an attractive candidate gene for disturbed adipocyte lipolytic function in obesity and insulin resistance in humans (Dahlman et al., 2004). Adipose tissue lipolytic responsiveness and sensitivity to adrenergic agents, including catecholamines, are increased in dairy cows around parturition (Bell, 1995). Lipolysis increases the concentrations of FFA in plasma. Because high lipolytic rates are considered a risk factor for the development of diseases during the periparturient period (Grummer, 1993), attenuating lipolysis or regulating insulin function (or both) in early lactation would be particularly beneficial to dairy cows that develop hypoinsulinemia concurrently with a state of insulin resistance (Contreras et al., 2017b). Even though FetA seems to be involved in the pathways that control lipolytic responses in bovine adipocytes, the mechanisms by which this adipokine may play anti-lipolytic roles in dairy cows remain to be established. Additionally, since lipolysis and lipogenesis occur simultaneously within adipocytes, these results may also suggest that the responses observed could be directly related to enhanced lipogenesis.

In the present study, we observed that incubation of bovine adipocytes with FetA increased FA uptake compared with nontreated cells and induced similar responses to those observed in cells treated with the lipogenic hormone insulin. Although speculative at this time, FetA could have stimulated the translocation of plasma membrane FA transporters (i.e., FATP1 and CD36) similar to the known effects of insulin on FA trafficking. Another possibility is that FetA could have bound FA and translocated them into the intracellular compartment. Fatty acid transporters, such as albumin and FetA, are known for carrying lipids in plasma and facilitating FA uptake by cells (Glatz et al., 2010). Previously, FetA was reported to translocate into the cytoplasm of 3T3-L1 and human adipocytes (Dasgupta et al., 2010) and to accumulate in vesicles in the cytosol (Reynolds et al., 2005); nevertheless, its relationship with FA transport was not reported in these studies. Fetuin-A carries high amounts of cholesterol, cholesterol-esters, TAG, and FA in plasma, which correspond to almost 33% of its molecule (Kumbla et al., 1989, 1991). In rabbit and human cells, FetA increased the incorporation of exogenous FA into intracellular TAG by almost 50-fold compared with albumin (Cayatte et al., 1990). Similarly, we observed that FetA increases TAG accumulation in cultured bovine adipocytes compared with nontreated cells. This suggests that increased FetA-stimulated FA uptake by adipocytes might have enhanced the intracellular synthesis of TAG. Due to its lipogenic properties, increased serum concentrations and AT expression and secretion of FetA have been associated with obesity and obesity-related disorders in humans and animals (Chen et al., 2009; Jialal et al., 2015; Pérez-Sotelo et al., 2017). In dairy cows, we previously reported a decrease in serum FetA concentrations and its expression in AT from 2 wk prepartum to 10 DIM (Strieder-Barboza et al., 2018). This period of decreased FetA coincided with the downregulation in the transcription of lipogenic enzymes in the subcutaneous AT and with a marked increase of serum FFA concentrations in periparturient cows. Evidence from global gene expression profiling studies revealed that A-HSG might be involved in regulating energy metabolism in dairy cattle (Chen et al., 2011), and in lipid accumulation in several AT depots in beef cattle (Robinson and Oddy, 2004). Together, these results provide evidence...
that FetA may promote lipogenesis in vitro and in vivo in the AT of cows.

Although it is clear that FetA promotes the synthesis of TAG, the mechanisms involved are yet to be studied. Whether the stimulation of TAG accumulation in bovine adipocytes is due to the entry of FA into cells or whether FetA might directly stimulate lipogenic enzymes is not known. To address these questions, we evaluated the gene expression of key enzymes involved in the TAG synthesis through the glycerol-3-phosphate pathway in adipocytes: (1) GPAT, which is involved in glycerol-P synthesis; (2) AGPAT, a catalyst for the first step in the formation of PA; (3) phosphatidate phosphohydrolases (lipin), which form diacylglycerol, and DGAT (Coleman and Lee, 2004; Takeuchi and Reue, 2009). In our study, FetA upregulated the expression of AGPAT2 and increased AGPAT2 protein content in adipocytes. The AGPAT2 is the predominant AGPAT isoform in AT and catalyzes acylation of its strict substrate lysophosphatidic acid to PA (Takeuchi and Reue, 2009). In cows, AGAP play an important role in the TAG synthesis in the mammary gland (Mistry and Medrano, 2002), but have not yet been reported in AT. Peroxisome proliferator-activated receptor γ agonists induce AGPAT2 activity (Blanchard et al., 2016) that it is required for TAG accumulation in mature adipocytes (Gale et al., 2006). Knocking down AGPAT2 decreased gene expression of adipogenesis regulators such as PPARγ and C/EBPB. Also, the absence of AGPAT2 delayed the expression of mature adipocyte markers such as FABP4 and GLUT4, and reduced TAG accumulation in adipocytes (Blanchard et al., 2016). The crucial function of AGPAT2 for adipogenesis is highlighted by the near complete absence of AT and a range of metabolic changes, such as extreme insulin resistance, in humans with congenital generalized lipodystrophy as a consequence of AGPAT2 deficiency (Agarwal and Garg, 2006). These results demonstrate that other AGAPT family members (AGAPT1 and AGAPT3–10) cannot compensate for AGPAT2 activity, thus being a specific and rate-limiting enzyme for TAG synthesis (Agarwal et al., 2002; Takeuchi and Reue, 2009). Although we demonstrate that FetA enhances AGPAT2 activity, future studies are needed to evaluate if this activation is a consequence of increased FA uptake mediated by FetA or a direct signaling effect.

The AGPAT2 controls adipogenesis through modulation of the synthesis of phospholipids and TAG precursors, especially PA (Gale et al., 2006). Impaired AGPAT2 activity affects the availability of PA for TAG synthesis but not overall PA synthesis nor utilization of PA for phospholipid synthesis (Gale et al., 2006). In our study, we evaluated the effect of FetA on AGPAT2 activity by measuring adipocyte concentrations of PA. We observed that FetA increased not only gene expression and protein content of AGPAT2 in bovine adipocytes, but also PA concentrations. Phosphatidic acid was highlighted as one of the main lipid regulators of lipid droplet size, an important lipid-storage organelle (Fei et al., 2011). Our results suggest that FetA increases AGPAT2 activity, and therefore PA availability for TAG synthesis in bovine adipocytes. Knowing that FetA may modulate TAG synthesis precursors in adipocytes provides valuable insights into potential targets for modulating the flux of lipids in that AT of dairy cows.

It is important to mention that we analyzed the effect of FetA on a limited pool of genes involved in adipogenesis, FA uptake, and lipogenesis. Although we identified an effect of FetA on a few of these selected genes, it is possible that other lipogenic markers could be driving the lipogenic effect of FetA. Future studies using genetic manipulation of AGPAT2 and AHSG expression in AT are needed to conclusively affirm that AGPAT2 signaling is required for the lipogenic effect of FetA in bovine AT.

**CONCLUSIONS**

Results from this study demonstrate that FetA modulates lipid mobilization by attenuating lipolytic responses and enhancing lipogenesis in bovine adipocytes. Fetuin-A upregulates the expression and activity of AGPAT2, a rate-limiting lipogenic enzyme, and suggests a potential mechanism by which this adipokine promotes TAG synthesis in adipocytes. Our study provides novel knowledge on how FetA promotes TAG synthesis in bovine adipocytes and opens the possibility of using FetA as a potential therapeutic target for the modulation of lipid mobilization in AT of periparturient dairy cows and humans with similar AT disorders.

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