S-Nitrosylation of Fatty Acid Synthase Regulates Its Activity Through Dimerization

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One Sentence Summary: Fatty acid synthase is S-nitrosylated during adipogenesis.

Keywords: S-nitrosylation, Fatty acid synthase, Dimerization, Adipocyte, Adipogenesis.

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ABSTRACT

Nitric oxide (NO) regulates a variety of physiological processes, including cell proliferation, differentiation, and inflammation. S-nitrosylation, an NO-mediated reversible protein modification, leads to changes in the activity and function of proteins. In particular, the role of S-nitrosylation during adipogenesis is largely unknown. We hypothesized that the normal physiological levels of NO, but not the excess levels generated under severe conditions such as inflammation, may be critically involved in the proper regulation of adipogenesis.

We found that endogenous S-nitrosylation of proteins was required for adipocyte differentiation. By performing a biotin-switch assay, we identified fatty acid synthase (FAS), a key lipogenic enzyme in adipocytes, as a target of S-nitrosylation during adipogenesis. Interestingly, we also observed that the dimerization of FAS increased in parallel with the amount of S-nitrosylated FAS during adipogenesis. In addition, we found that exogenous NO enhanced the dimerization and the enzymatic activity of FAS. Moreover, site-directed mutagenesis of three predicted S-nitrosylation sites indicated that S-nitrosylation of FAS at Cys\textsuperscript{1471} and Cys\textsuperscript{2091}, but not at Cys\textsuperscript{1127}, increased its enzymatic activity.

Taken together, these results suggest that the S-nitrosylation of FAS at normal physiological levels of NO increases its activity through dimerization and may contribute to the proper regulation of adipogenesis.
INTRODUCTION

Fatty acid synthase (FAS), which is highly expressed in adipose tissue, liver, and lactating mammary glands, catalyzes the synthesis of palmitate from acetyl-CoA and malonyl-CoA in the presence of NADPH (1–3). FAS is active as a homodimer, and each monomer has seven separate functional domains, including malonyl/acetyltransferase (MAT), β-ketoacyl synthase (KS), β-ketoacyl reductase (KR), dehydrase (DH), enoyl reductase (ER), thioesterase (TE), and acyl carrier protein (ACP)(4). Although the monomer contains all the activities required for palmitate synthesis, dimer formation is essential for FAS function (5–9). Several studies have reported that FAS is involved in the regulation of adipose tissue mass (1, 10–18) and a key enzyme regulating energy metabolism and a metabolic oncogene (19, 20). Therefore, FAS has been considered as a potential target for the development of anti-obesity and anti-cancer drugs (21–25). Expression of FAS is transcriptionally regulated by the sterol regulatory element binding protein-1c (SREBP-1c) and by upstream stimulatory factors 1 and 2 (USF1 and USF2) in response to feeding or to insulin (26, 27). So far, however, nothing is known about the post-translational modification of FAS except that it is ubiquitinated (28, 29).

Nitric oxide (NO) is a critical signaling molecule that is involved in a number of physiological and pathological processes. When overproduced, NO is known to have harmful effects on cell function and survival (30–32). In contrast, within the physiological concentration range, NO plays a critical role as a regulator of cellular signaling pathways (33–35). A major mechanism mediating the biological function of NO is protein S-nitrosylation, a posttranslational modification of proteins involving the addition of an NO$^+$ to a cysteine thiol group of the proteins. Therefore, under physiological conditions, S-nitrosylation can affect a number of cellular signaling pathways by inducing conformational changes of the protein and affecting protein-protein interactions and protein functions (36-39).
NO is reported to improve the β-oxidation of fatty acids through reversible protein S-nitrosylation (40). NO is also known to impair the anti-lipolytic action of insulin in obesity through S-nitrosylation (41) and to enhance adipogenesis in primary human preadipocytes (42). However, little is known about the target proteins of S-nitrosylation during adipogenesis.

In the present study, we showed that FAS is S-nitrosylated during adipogenesis and that the S-nitrosylation of FAS increases its activity by enhancing dimerization, indicating that S-nitrosylation of FAS contributes to adipogenesis.
RESULTS

Protein S-nitrosylation is required for adipocyte differentiation

To investigate whether NO is involved in the process of adipocyte differentiation, we first evaluated NO production during adipocyte differentiation (Fig. 1A). The production of NO during adipocyte differentiation was estimated by measuring the total concentration of its oxidation metabolites, nitrite and nitrate, using an NO assay kit. Compared with the early differentiation period of adipocytes (days 0-2), there was a significant increase in total NO level in the supernatants of cells taken at the later differentiation period (days 6-8) (Fig. 1A). We next investigated whether protein S-nitrosylation, a major post-translational modification by NO, might have a unique role in adipogenesis. Protein S-nitrosylation can occur even at low levels of NO and can affect the activity and stability of specific proteins (36-39).

Immunofluorescence (IF) assay using an antibody against nitroso-cysteine showed that protein S-nitrosylation was higher in differentiated adipocytes (day 9) than in undifferentiated preadipocytes (Fig. 1B). S-nitrosocysteine (SNOC), an NO donor, demonstrated a biphasic effect: increased adipogenesis at a low concentration (50 µM) and decreased adipogenesis at a high concentration (200 µM) (Fig. 1C). 2-(4-carboxyphenyl)-4,4,5,5-tetramethylimidazoline-1-oxyl-3-oxide (cPTIO) and 2-phenyl-4,4,5,5-tetramethylimidazoline-1-oxyl-3-oxide (PTIO), well-known NO scavengers, inhibited adipogenesis in a concentration-dependent manner (Fig. 1D, 1E). The better NO scavenger, cPTIO showed slightly more efficient inhibition of adipogenesis than PTIO at the same concentrations (Figs. 1D and 1E). To test the effect of NOS inhibitor on adipogenesis, we performed same experiments with NNA, a broad NOS inhibitor. We found that there was no effect of NNA on adipogenesis (Supplementary Fig. 1A). In agreement with a previous report (43), we could not see the expression of iNOS and nNOS during adipogenesis, whereas eNOS protein band was detectable only at the later differentiation stage (Day 14) (Supplementary Fig. 1B).
Increased S-nitrosylation and dimerization of FAS in differentiated adipocytes

Following a previous report on fatty acid metabolism and S-nitrosylation (40), we analyzed the levels of FAS, one of the essential proteins for lipid droplet formation during adipogenesis, in differentiated adipocytes. We confirmed that the levels of nitrosocysteine and FAS in more differentiated adipocytes (day 12) were higher than those in less differentiated adipocytes (day 2) (Fig. 2A). The yellow color in the merged image of nitrosocysteine (green) and FAS (red) images implied that FAS was likely to be S-nitrosylated (SNO-FAS) (Fig. 2A). We also performed the immunostaining for nitrosocysteine with negative controls to reaffirm antibody and detection specificity (Supplementary Fig. 2). To confirm that the biotin-switch assay allows detection of SNO-FAS, the lysate of undifferentiated adipocytes was subjected to the assay after incubation in SNOC with or without the assay components, including methyl-methanethiosulfonate (MMTS), ascorbate, and N-[6-(biotinamido)hexyl]-3′-(2′-pyridyldithio)-propionamide (biotin-HPDP) (Fig. 2B). In addition, we tested whether FAS in adipose tissue homogenates could be S-nitrosylated. We eliminated S-nitrosocysteine by treatment with UV-light and used this sample for negative control. We found that FAS was nitrosylated in human adipose tissue and the SNO-FAS in lysates was eliminated by UV-light (Fig. 2C).

To test if SNOC-induced FAS nitrosylation is cPTIO sensitive, we incubated the cells with cPTIO before SNOC treatment and performed biotin switch assay. We found that SNOC-induced FAS nitrosylation was cPTIO sensitive (Supplementary Fig. 3). It has been reported that DEA-NONOate, another NO donor, can nitrosylate proteins (44). Thus we tested the effect of DEA-NONOate on FAS nitrosylation. Similar to SNOC, DEA-NONOate nitrosylated FAS (Supplementary Fig. 3). We then compared the relative amount of SNO-FAS at different stages of adipocyte differentiation (days 0-14). Interestingly, we found that
the SNO-FAS level was increased during adipocyte differentiation (Fig. 2D). We also found that FAS dimer formation was increased during the later stage of adipocyte differentiation (day 9) (Fig. 2E).

**Dimerization and activation of FAS are mediated by NO**

To identify whether FAS dimerization can be directly induced by NO, we examined the effect of exogenous NO on the dimerization level of FAS by non-denaturing PAGE and western blot analysis. We found that SNOC significantly increased FAS dimerization (Fig. 3A). FAS dimer disappeared when cell lysates were boiled under reducing conditions, which suggested that FAS dimerization might be mediated through disulfide bonds. We then tested whether NO could influence FAS activity. Similar to the result with dimerization, the enzymatic activity of FAS was significantly higher in the lysate of ADSCs treated with fresh SNOC than in that treated with old SNOC, from which NO had dissipated (Fig. 3B).

**FAS is S-nitrosylated at Cys\(^{1471}\) and Cys\(^{2091}\)**

FAS contains several cysteines in its primary sequence. Recently, Cys\(^{1127}\), Cys\(^{1471}\), and Cys\(^{2091}\) have been predicted as candidate sites for FAS S-nitrosylation by a mass spectrometry-based proteomics study (40). Thus, we examined whether these three cysteines may be involved in FAS S-nitrosylation. We generated mutants in which each of the cysteines was mutated to alanine (C1127A, C1471A, and C2091A). We also included Cys\(^{161}\), in the active site of FAS, as a negative control for FAS activity. HEK293 cells were transiently transfected with FLAG-tagged wild-type FAS (FAS-WT) or its cysteine to alanine mutants and were exposed to SNOC. Cell lysates were subsequently subjected to the biotin-switch assay. The treatment of cells with SNOC resulted in S-nitrosylation of FAS-WT and
the C161A and C1127A mutant proteins. In contrast, neither the C1471A nor the C2091A mutant was nitrosylated by SNOC (Fig. 4A). The dimerization of FAS-WT and the C1127A mutant was enhanced by SNOC, but the dimerization of C161A, C1471A, or C2091A did not increase (Fig. 4B), demonstrating that S-nitrosylation of FAS at Cys\textsuperscript{1471} or Cys\textsuperscript{2091} is important for FAS dimerization. Consistent with these results, the SNOC-induced increase in activity of the C161A, C1471A, or C2091A mutants was significantly lower than that of FAS-WT (Fig. 4C). In addition, we examined the effect of cysteine to alanine mutation itself upon the basal enzymatic activity, and there was no significant difference versus FAS-WT (Supplementary Fig. 4).
DISCUSSION

There are conflicting views regarding the role of NO in adipogenesis. Several papers have reported that NO suppresses adipocyte differentiation, but others have shown that NO can enhance adipogenesis (42, 43, 45-47). This discrepancy seems reasonable considering our results demonstrating that the production of NO was relatively low even though it increased significantly during adipogenesis (Fig. 1A) and that SNOC has a biphasic effect on adipogenesis (Fig. 1C). Taken together, these data lead us to assume that a small amount of free NO is required for adipogenesis, whereas a larger amount of NO may suppress adipogenesis.

Protein S-nitrosylation is a major reversible posttranslational modification by NO that regulates protein function and stability (36–39, 48–53). Interestingly, it has been reported that protein S-nitrosylation is increased in adipose tissue in a model of obesity (41). To explore the biological role of S-nitrosylation in adipogenesis, we first focused on FAS, the key enzyme of fatty acid synthesis. We found that FAS was S-nitrosylated under physiological conditions and the level of SNO-FAS was gradually increased during adipogenesis (Fig. 2). FAS became more active following S-nitrosylation, indicating that S-nitrosylation plays an important role in adipogenesis. It has been reported that NO regulates mitochondrial fatty acid metabolism through protein S-nitrosylation regulation (40). Following a mass spectrometry (MS)-based proteomics study, VLCAD [very long chain acyl-coenzyme A (CoA) dehydrogenase] was proposed as a candidate S-nitrosylated protein involved in fatty acid metabolism. Similar to SNO-FAS, the activity of SNO-VLCAD was found to be increased by S-nitrosylation, implying a stimulating role of S-nitrosylation for fatty acid oxidation. These studies suggest that S-nitrosylation might stimulate both adipogenesis and
fatty acid oxidation. Therefore, the balance of the S-nitrosylation level might be a key signaling pathway regulating obesity.

Previous reports have shown that protein S-nitrosylation affects protein activity by enhancing protein dimerization of Drp1 (52) or interrupting protein dimerization of eNOS (54). FAS formed dimers following S-nitrosylation. So far, there is no cysteine residue which is known to be involved in FAS dimerization. Although we showed that the Cys\textsuperscript{1471} and Cys\textsuperscript{2091} residues of FAS are important for dimerization and activation by NO as targets for S-nitrosylation, neither cysteine residue is in the active site. However, interestingly, Cys\textsuperscript{1471} is in the center of the interdomain of FAS, which is known to be important for catalytically active dimer formation (55).

NO scavengers significantly inhibited adipogenesis. We suggest that the inhibition of SNO-FAS formation leading to FAS inactivation could be a possible signaling pathway by which NO scavengers inhibit adipogenesis. During adipocyte differentiation, the decrease in fatty acid synthesis is known to interrupt the cross-stimulation of C/EBP\textgreek{a} and PPAR\textgreek{y}, and leads to insufficient adipogenesis (56–58). We assume that these events may have occurred in our experimental system, at least in part, when exposed to NO scavengers. Based on the data showing no effect of NOS inhibitor on adipogenesis (Supplementary Fig. 1A), we have a hypothesis that NO during adipogenesis in the present experiments could be coming from intracellular ‘SNO pool’ such as S-nitrosoglutathione (GSNO), SNOC and other SNO-proteins, as suggested before (34). Though there are no previous reports about the higher thiol oxidation states of FAS, it is still possible that the further thiol oxidation of FAS can alter its activity in our experimental system. It is because reversible S-nitrosylation may facilitate further oxidation of the same cysteine thiol (59, 60). In addition to affecting adipogenesis, fatty acids produced by FAS are known to stimulate other signaling pathways, including autophagy and miRNA signaling (61, 62). FAS has also received much attention regarding
the regulation of its activity during cancer development (19-23). As a key enzyme regulating energy metabolism and as a metabolic oncogene, FAS has been considered as a potential target for the development of anti-obesity and anti-cancer drugs. Taken together, our results suggest that the regulation of FAS activation by S-nitrosylation may contribute to the regulation of a variety of physiological or pathological events, including adipogenesis and cancer.
Conflict of Interest

The authors state no conflicts of interest.
MATERIALS AND METHODS

Reagents and antibodies

The NO scavengers 2-Phenyl-4,4,5,5-tetramethylimidazoline-1-oxyl-3-oxide (PTIO) (P5084) and 2-(4-carboxyphenyl)-4,4,5,5-tetramethylimidazoline-1-oxyl-3-oxide (cPTIO) (C221) were obtained from Sigma–Aldrich (St. Louis, MO, USA). The NO donor, diethylamine diazeniumdiolate diethylammonium salt (DEA-NONOate) (D5431) and a NOS inhibitor, Nω-Nitro-L-arginine (L-NNA) (N5501) were also obtained from Sigma–Aldrich (St. Louis, MO, USA). S-nitrosocysteine (SNOC) was prepared as described previously (63). Lipofectamine 2000 (11668019) and Lipofectamine LTX with Plus Reagent (15338030) were obtained from Life Technologies (Carlsbad, CA, USA). Anti-nitrosocysteine (ab94930), and anti-FAS (ab22759) antibodies were purchased from Abcam (Cambridge, UK), mouse monoclonal anti-FLAG (F3166) antibody from Sigma–Aldrich (St. Louis, MO, USA), and anti-GAPDH (SC-25778) from Santa Cruz Biotechnology (Dallas, TX, USA).

Cell culture and adipose tissue

HEK293 cells were grown in DMEM (Life Technologies, Carlsbad, CA, USA) containing 10% fetal bovine serum at 37°C under 5% CO₂/95% air. Human adipose-derived stem cells (ADSCs) were purchased from Lonza (Walkersville, MD, USA). Confluent ADSCs were incubated in ADSC differentiation medium containing 1 µg/ml insulin, 1 µM dexamethasone, 0.5 mM IBMX, 1 µM troglitazone (called as IDXT), and 10 % FBS. The culture medium was changed every 2 days. Insulin (I5500), IBMX (I5879), dexamethasone (D8893), and troglitazone (T2573) were purchased from Sigma–Aldrich (St. Louis, MO, USA).
USA). Flash frozen human subcutaneous adipose tissue (T-SQFX-FF) was purchased from Zen-Bio (Research triangle park, NC, USA).

NO assay

For quantitative determination of the nitric oxide level, a Nitric Oxide Assay Kit (Colorimetric, ab65328) from Abcam (Cambridge, UK) was used. Total nitrate and nitrite levels were measured in a two-step process. The first step converts nitrate to nitrite with nitrate reductase. The second step uses Griess Reagents to convert nitrite to a deep purple azo compound. The amount of azochromophore reflects the amount of nitric oxide in the samples. The absorbance at 550 nm was immediately recorded and compared to the absorbance of a freshly prepared standard curve of sodium nitrite.

Immunocytochemistry

Adipocytes were fixed for 1 h in 10% formalin in PBS followed by permeabilization in 0.1% Triton X-100 for 10 min. Blocking was performed in PBS containing 10% normal goat serum for 1 hour. The fixed cells were incubated using anti-FAS or anti-nitrosocysteine overnight in the blocking solution. The cells were washed and incubated for 1 h with either Alexa-Fluor-488-conjugated goat anti-mouse antibody (A10680) or Alexa-Fluor-594-conjugated goat anti-rabbit antibody (A11012) (Molecular Probes, Carlsbad, CA, USA). After washing, the cells were mounted in a mounting medium containing DAPI. Cells were observed using an EVOS FL cell imaging system (Life Technologies, Carlsbad, CA, USA). Formalin solution (HT501128), goat serum (G9023), and fluoroshield mounting medium with DAPI (F6057) were purchased from Sigma–Aldrich (St. Louis, MO, USA).

FAS constructs
A plasmid encoding human FAS (SC127829, NM_004104) was obtained from Origene (Rockville, MD, USA) and subcloned into the pcDNA3.1 vector (Life Technologies, Carlsbad, CA, USA). Point mutations were introduced to change each cysteine in the FAS gene (Cys\(^{161}\), Cys\(^{1127}\), Cys\(^{1471}\), or Cys\(^{2091}\)) to an alanine. The primer sequences for PCR-directed mutagenesis of FAS Cys\(^{161}\) to Ala were: forward, 5' - CGCAGATGGAGACACATTCCCTCATCCAGGCT - 3'; reverse, 5' - GCCATTCGGAGCTGAGGGCTGTGCTCTCCAGGCT - 3'. For mutagenesis of FAS Cys\(^{1127}\) to Ala, the primer sequences were: forward, 5' - CCGGACCAAGCCCTCCCTCTCCCTTCTCC - 3'; reverse, 5' - CCGGTCGCTCCAGAAGGCTTCGCTCTCTCGTCC - 3'. For mutagenesis of FAS Cys\(^{1471}\) to Ala, the primer sequences were: forward, 5' - CCAGCCGTGGAGCGTGCCCTCCGGTCGGTCTCC - 3'; reverse, 5' - CTGGAGAGTGCTGCTGCTCCGTCCCTCC - 3'. For mutagenesis of FAS Cys\(^{2091}\) to Ala, the primer sequences were: forward, 5' - CGCACCACTCAGCAGCCCTCCCTCCCTCC - 3'; reverse, 5' - CGCAGAGGCTGGAGGCTGCTGCTGCTCCGTCCCTCC - 3'.

**Oil Red O staining**

Adipocyte differentiation was assessed using an Oil Red O stain (O0625, Sigma–Aldrich, St. Louis, MO, USA) as an indicator of intracellular lipid accumulation. After ADSC differentiation to adipocytes, cells were washed twice with PBS, fixed with 10% formalin in PBS for 1 h, and then washed with 60% isopropanol, before being allowed to dry completely. Adipocytes were stained with 0.2% Oil Red O reagent for 10 min at room temperature and washed with \(\text{H}_2\text{O}\) four times. Each sample was eluted with 100% isopropanol for 10 min and absorbance was measured at 500 nm using a spectrophotometer. To visualize the nucleus,
adipocytes were counterstained with a hematoxylin reagent (H3136, Sigma–Aldrich, St. Louis, MO, USA) for 2 min and washed twice with H2O. The level of adipocyte differentiation was observed using an inverted phase microscope.

**Biotin-switch assay**

Cell lysates or adipose tissue lysates were prepared in HENTS buffer (100 mM HEPES, 1 mM EDTA, 0.1 mM neocuproine, 1% Triton X-100, and 0.1% SDS, pH 7.4). The biotin-switch assay was carried out as described previously (64) with a slight modification. Briefly, free thiols in the sample were blocked by incubation with 10 mM methymethanethiosulfonate (MMTS) at 50°C for 15 min. The MMTS was then removed by acetone precipitation, and the pellet was resuspended in HENS buffer (HEN plus 1% SDS). S-nitrosothiols were selectively reduced with 20 mM ascorbate and the reformed free thiols were labeled with 1 mM N-[6-(biotinamido)hexyl]-3′-(2′-pyridyl)dithio)-propionamide (HPDP-biotin) for 1 h at RT. The biotinylated proteins were then collected on avidin agarose beads, which were then washed three times with neutralization buffer (20 mM HEPES-NaOH pH 7.4, 100 mM NaCl, 1 mM EDTA, 0.5% Triton X-100). Proteins were eluted from the beads by SDS-PAGE loading buffer and subjected to immunoblot analysis. Neocuproine (N1501), MMTS (64306), and ascorbate (A7506) were purchased from Sigma–Aldrich (St. Louis, MO, USA). EZ-Link HPDP-biotin and NeutrAvidin agarose beads were from Thermo Scientific (Waltham, MA, USA).

**FAS activity assay**

The FAS activity assay was performed as previously described (65) with minor modifications. In brief, cell lysates were mixed with acetyl-CoA and NADPH in 0.2 M potassium phosphate buffer, 0.4 mM EDTA (pH 7.0) and incubated at 30°C for 10 min. The
enzymatic reaction was initiated by adding 20 μl of malonyl-CoA solution (0.2 mM), and the decrease in OD was measured every 1 min for 30 min via kinetic measurements obtained on a microplate reader set at 340 nm. Based on the results, the overall FAS enzyme activity was estimated by calculating the NADPH oxidation, using $\varepsilon = 6220 \text{ M}^{-1}\text{cm}^{-1}$. Acetyl-CoA (A2056), NADPH (N1630), and malonyl-CoA (M4263) were purchased from Sigma–Aldrich (St. Louis, MO, USA).

**Statistical analysis**

The statistical analyses of the data were analyzed with the Student’s t test or by one-way analysis of variance (ANOVA). Data are expressed as means ± S.D. of at least three independent experiments. P-value of less than 0.05 was considered statistically significant.
REFERENCES


Figure Legends

Figure 1. Adipocyte differentiation requires protein S-nitrosylation.

A. Total NO produced by differentiated adipocytes at each period (0-2 days, 3-5 days, or 6-8 days) was measured using an NO assay kit. C-E. ADSCs were grown in the IDXT condition and treated with the indicated concentration of SNOC (C), PTIO (D) or cPTIO (E). Lipid droplets in adipocytes were stained with Oil Red O dye and quantified. Data were normalized by setting the insulin control as 1 (mean ± S.D.). **p<0.01 by t test. *p<0.05, ##p<0.01 compared to IDXT by t test. Data are representative of three individual experiments.

B. On day 9 following IDXT incubation, immunofluorescence staining was performed on the control and the 9-day differentiated adipocyte samples using an anti-nitrosocysteine antibody.

Figure 2. Increased S-nitrosylation and dimerization of FAS in differentiated adipocytes.

A. Immunofluorescence to detect expression of FAS (red) and S-nitrosylated proteins (green) in differentiated adipocytes (on day 2 and 12). Merged images (yellow) demonstrate the overlap between the localization of FAS and S-nitrosylated proteins. DAPI staining (blue) was used to identify cell nuclei. Scale bar = 100 µm. B. To confirm the S-nitrosylation of FAS and that the biotin-switch assay detects SNO-FAS properly, the lysate of undifferentiated ADSCs was subjected to a biotin-switch assay with or without the components of the assay as described in Materials and Methods and immunoblotting following incubation with the NO donor S-nitrosocysteine (SNOC) or Old SNOC. C. The lysates of human adipose tissue (200 mg) were irradiated with UV-light or incubated in the dark for 10 min. Then, the lysates were subjected to a biotin-switch assay with or without ascorbate. UV-irradiated sample was used as a negative control by eliminating S-nitrosocysteine. Data are representative of three individual experiments. D. The lysates of cells at different stages of adipocyte differentiation (days 0-14) were subjected to a biotin-
switch assay followed by immunoblotting for SNO-FAS (top), total FAS (middle), and GAPDH (bottom). And the ratio of SNO-FAS/Total FAS expression was quantified. Data were normalized by setting ‘Day 0’ as 1 (mean ± S.D.). *p<0.05 compared to ‘Day 0’ by t test. E. Immunoblotting of cell lysates carried out under non-denaturing conditions shows the dimerization of FAS during adipogenesis (days 3 and 9). The ratio of FAS dimer/FAS monomer was quantified. Data were normalized by setting ‘Day 3’ as 1 (mean ± S.D.). *p<0.05 compared to ‘Day 3’ by t test. Data are representative of three individual experiments.

**Figure 3.** Exogenous NO induces dimerization and activation of FAS.

A. SNOC-induced FAS dimer formation in ADSCs was observed on a non-denaturing PAGE and by western blot analysis. SNOC-induced dimer of FAS was well detected when cell lysates were not boiled under non-reducing conditions (left) and still detected when cell lysates were boiled under non-reducing conditions (middle), but not detected when cell lysates were boiled under reducing conditions (right). B. The lysates of ADSCs were exposed to SNOC (25 µM) or old SNOC and subjected to non-denaturing PAGE (upper) or the FAS activity assay (lower). *p<0.05 compared to old SNOC by t test. Error bars indicate S.D.. Data are representative of three individual experiments.

**Figure 4.** FAS is S-nitrosylated on Cys\(^{1471}\) and Cys\(^{2091}\).

A, B. HEK293 cells were transfected with FLAG-tagged wild-type FAS (FAS-WT) or its cysteine mutants (C161A, C1127A, C1471A, and C2091A) and exposed to SNOC or Old SNOC. The cell lysates were subsequently subjected to a biotin-switch assay (A) or to non-denaturing PAGE (B). After the blots were developed with a primary antibody against
FLAG, they were stripped and re-probed with a primary antibody against FAS. And the ratio of SNO-FAS/Total FAS expression (A) or FAS dimer/FAS monomer (B) was quantified. Data were normalized by setting Old SNOC for each group as 1 (mean ± S.D.). *p<0.05 by t test. C. The lysates of cells transfected with FAS-WT and its cysteine mutants were exposed to SNOC (25 µM) or Old SNOC and subjected to the FAS activity assay. Data were normalized by setting the old SNOC samples for each group as 1 (mean ± S.D.). **p<0.01 compared to FAS-WT SNOC by t test. Data are representative of three individual experiments.
Fig. 1

A

Total NO production (nM)

Day 0-2 Day 3-5 Day 6-8

B

Nitro-Cys
DAPI
Day 0 Day 9

C

Relative Oil Red O

Insulin SNOC 50µM SNOC 100µM SNOC 200µM

D

Relative Oil Red O

Insulin PTIO 25µM PTIO 50µM PTIO 100µM

E

IDXT + cPTIO 50µM

IDXT

IDXT
Fig. 2

A

Day 2

Day 12

NitroCys
FAS
Merge
DAPI

B

SNO
- FAS
Total FAS

C

SNO-FAS
Total FAS
GAPDH
Input

MW kDa
315
250
315
250
180
140
95
42
55
72

Ascorbate
+ + - + +
+ - + +
+ + + +

Biotin
- + + + +
+ + - + +
+ + + - +
+ + + +

SNO-Cys
MMTS

+ + - + +
+ + + +

UV
- + + +

IB: α-NitroCys

Ascorbate
- - + +

GAPDH
- - + +

Input

Total FAS

SNO-protein (IB: α-NitroCys)
Fig. 2 (Continue)

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<td>Reduced &amp; boiled</td>
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</tr>
</tbody>
</table>

SNOC

- FAS dimer
- FAS monomer

GAPDH

B

<table>
<thead>
<tr>
<th>Condition</th>
<th>MW (kDa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Untreated</td>
<td>-</td>
</tr>
<tr>
<td>Old SNOC</td>
<td>-</td>
</tr>
<tr>
<td>SNOC</td>
<td>-</td>
</tr>
</tbody>
</table>

Specific activity of FAS (nmole NADPH/min/mg protein)

- Untreated
- Old SNOC
- SNOC

* Significant difference
Fig. 4

A

<table>
<thead>
<tr>
<th></th>
<th>PrEDNA</th>
<th>FAS WT</th>
<th>C161A</th>
<th>C127A</th>
<th>C1471A</th>
<th>C2091A</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNOC</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Input (5%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**IB : α-FLAG**

- SNO-FAS
- Input (5%)

**IB : α-FAS**

- SNO-FAS
- Input (5%)

GAPDH

**SNO-FAS / Total FAS**

(IB : α-FLAG, ratio of Old SNOC)

<table>
<thead>
<tr>
<th></th>
<th>FAS WT</th>
<th>C161A</th>
<th>C127A</th>
<th>C1471A</th>
<th>C2091A</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>1.5</td>
<td>2</td>
<td>2.5</td>
<td>1</td>
</tr>
</tbody>
</table>

* indicates significant difference from control.
Fig. 4 (Continue)

**FAS dimer / FAS monomer**  
(IB : α-FLAG, ratio of Old SNOC)

<table>
<thead>
<tr>
<th></th>
<th>FAS WT</th>
<th>C1127A</th>
<th>C1471A</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>0.5</td>
<td>1.0</td>
<td>15</td>
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<tr>
<td>20</td>
<td>2.5</td>
<td>3.0</td>
<td>4.0</td>
</tr>
</tbody>
</table>

IB : GAPDH
IB : α-FLAG
IB : α-FAS

**SNOC**

- FAS WT
- C161A
- C1127A
- C1471A
- C2091A

* by guest, on November 10, 2017 www.jlr.org Downloaded from
Fig. 4 (Continue)

C

Relative activity of FAS (Ratio of Old SNOC)

<table>
<thead>
<tr>
<th></th>
<th>Old SNOC</th>
<th>SNOC</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT FAS</td>
<td>0.97</td>
<td>1.00</td>
</tr>
<tr>
<td>C161A</td>
<td>1.65</td>
<td>1.68</td>
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<tr>
<td>C1127A</td>
<td>1.39</td>
<td>1.42</td>
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<tr>
<td>C1471A</td>
<td>1.28</td>
<td>1.30</td>
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<tr>
<td>C2091A</td>
<td>1.18</td>
<td>1.20</td>
</tr>
</tbody>
</table>

** Statistical significance calculated by two-tailed Student’s t-test, p < 0.01.