A classic metabolic concept posits that insulin promotes energy storage and adipose expansion, while catecholamines stimulate release of adipose energy stores by hydrolysis of triglycerides through β-adrenergic receptor (βARs) and protein kinase A (PKA) signaling. Here, we have shown that a key hub in the insulin signaling pathway, activation of p70 ribosomal S6 kinase (S6K1) through mTORC1, is also triggered by PKA activation in both mouse and human adipocytes. Mice with mTORC1 impairment, either through adipocyte-specific deletion of Raptor or pharmacologic rapamycin treatment, were refractory to the well-known βAR-dependent increase of uncoupling protein UCP1 expression and expansion of beige/brite adipocytes (so-called browning) in white adipose tissue (WAT). Mechanistically, PKA directly phosphorylated mTOR and RAPTOR on unique serine residues, an effect that was independent of insulin/AKT signaling. Abrogation of the PKA site within RAPTOR disrupted βAR/mTORC1 activation of S6K1 without affecting mTORC1 activation by insulin. Conversely, a phosphomimetic RAPTOR augmented S6K1 activity. Together, these studies reveal a signaling pathway from βARs and PKA through mTORC1 that is required for adipose browning by catecholamines and provides potential therapeutic strategies to enhance energy expenditure and combat metabolic disease.

Introduction

Adipose tissue depots in both humans and rodents perform a remarkable spectrum of contrasting functions, including sequestration of excess caloric energy in the form of triglyceride in white adipose tissue (WAT) and rapid conversion of oxidative energy to heat for survival in the cold in brown adipose tissue (BAT). Various fat depots also secrete specific signature profiles of proteins and other factors that play key roles in overall systemic energy metabolism and glucose homeostasis (1). A general paradigm in the field is that 2 major opposing hormone systems, insulin (Ins) and catecholamines, are prominent regulators of these adipocyte functions. The catecholamines norepinephrine and epinephrine activate β-adrenergic receptors (βARs) to increase cAMP levels and cAMP-dependent protein kinase A (PKA) activity. PKA phosphorylates and regulates several important targets in adipocytes, including hormone-sensitive lipase and the lipid droplet-associate perilipins, which collectively promote triglyceride hydrolysis and liberation of free fatty acids (2–4). The opposing metabolic regulator is Ins, which antagonizes the action of the catecholamines to stimulate lipolysis by activating phosphodiesterases that degrade cAMP (5) and by activating lipid synthesis pathways through actions of the protein kinase AKT (6).

Signaling by catecholamines to stimulate lipolysis in WAT provides fatty acid substrates to fuel peripheral tissues, while, in BAT, lipolysis provides fatty acid substrates for generating heat. Brown adipocytes are highly enriched in mitochondria and express the unique protein uncoupling protein-1 (UCP1), which “uncouples” the mitochondrial proton gradient from adenosine triphosphate (ATP) production during fatty acid oxidation to produce thermal energy (7). The result is net energy expenditure. UCP1-containing “brown-like” adipocytes can also be recruited within WAT depots through prolonged βAR stimulation of lipolysis (8, 9) and are called “brite” or “beige” adipocytes (10, 11). Studies using mouse models show that the increase in these beige adipocytes closely correlates with resistance to obesity (12, 13), and in humans, the amount of detectable brown/beige adipocytes is significantly correlated with reduced percent body fat and circulating triglycerides, as well as greater Ins sensitivity (14–21). As with lipolysis, where Ins opposes the actions of βARs, Ins also appears to antagonize catecholamine stimulation of adipose browning (22) secondary to its suppression of lipolysis.

A major signaling node for the anabolic actions of Ins that strongly promote lipogenesis and protein synthesis downstream of AKT are the mTOR complexes. mTOR is a 250 kDa conserved Ser/Thr kinase that regulates cell growth and metabolism in response to environmental cues such as growth factors and nutrients, in addition to Ins (23). There are 2 structurally and functionally distinct mTOR-containing protein complexes, mTORC1 and mTORC2, required for activation of AKT (24–26). A defining characteristic of mTORC1 is its inhibition by the macrolide antibiotic rapamycin (26, 27), and it contains the partner protein RAPTOR (regulatory-associated protein of mTOR). A well-characterized downstream target of mTORC1, but not mTORC2, is p70 ribosomal S6 kinase 1 (S6K1), which is thus highly activated by Ins.
Ins activation of mTOR would suggest that catecholamines may oppose this action and inhibit this complex. Indeed, data showing that the cAMP pathway can inhibit mTOR exists not only in adipocytes (28, 29), but also in other cells such as lymphoblasts (30) and smooth muscle cells (31, 32). However, catecholamine action to increase the abundance of beige adipocytes within WAT entails increased biosynthetic capacity, including enhanced mitochondrial and cell protein mass, for which mTOR1 is a central player. We therefore further investigated this connection and show here that β-adrenergic stimulation activates the mTORC1/S6K1 pathway. Remarkably, we found that βAR-mediated browning of white adipose depots in mice requires the activation of mTORC1 and S6K1 through a mechanism that is distinct from that of Ins and growth factors. Specifically, both mTOR and the mTORC1 component RAPTOR can be phosphorylated by PKA, and interference with phosphorylation of RAPTOR by PKA prevents activation of S6K1 by β-agonists, without affecting its stimulation by Ins. These findings illustrate the existence of a new signaling pathway from PKA to mTORC1 and its requirement for the sympathetic nervous system–driven (SNS-driven) recruitment and expansion of the UCP1-containing beige cells in WAT.

Results

βAR agonists activate mTORC1 in adipocytes

During preliminary studies comparing the effects of Ins and βAR agonists on signal transduction and metabolism in adipocytes, we observed that when 3T3-L1 adipocytes were treated with the nonselective βAR agonist isoproterenol (Iso), S6K1 became phosphorylated, which is best known to be activated by Ins. We then compared the abil-
AKT-dependent pathway (33). We compared signaling events mediating S6K1 activation in response to Iso versus Ins in 3T3-L1 adipocytes. As shown in Figure 2A, Ins (10 nM) increased phosphorylation of AKT (Thr308, Ser473) and S6K1 (Thr389), as established by others (6, 34). Phosphorylation of S6K1 by Ins was fully suppressed by rapamycin. In contrast to Ins, Iso activated S6K1 but did not trigger phosphorylation of AKT\textsuperscript{T308} or AKT\textsuperscript{S473} (Figure 2B). In addition, proline-rich AKT substrate 40 (PRAS40) was phosphorylated by Ins treatment but not by Iso. We also observed that GSK3\textalpha{} and -\textbeta{} were phosphorylated by either Ins or Iso, which was unaffected by rapamycin. ULK1, another established mTORC1 target (35), was also activated by both Ins and Iso. To further confirm the activation of mTORC1 by Iso, the pan-mTOR inhibitor torin1 (36) blocked both Iso- and Ins-stimulated S6K1 activation (Supplemental Figure 4). We also found that rapamycin treatment in L1 adipocytes did not affect Iso-mediated lipolysis (Supplemental Figure 5). Considering that G protein–coupled receptors including \beta{}ARs can also activate MAPKs such as ERK and p38 MAPK (37–41), we examined these pathways. As shown in Figure 2, C and D, the MEK inhibitor PD98059 (PD) and the p38 MAPK inhibitor SB202190 (SB) fully blocked activation of their respective kinases following Iso stimulation, but these inhibitors had no effect on Iso-mediated S6K1 phosphorylation.
Cold-temperature challenge. A cold-temperature challenge is a classic maneuver that triggers an increase in SNS outflow to tissues including white and brown fat and generates heat from stored energy (42–45). When placed in a cold environment, mice exhibit a moderate decrease in core body temperature within the first few hours, followed by a temperature plateau. In this early phase, the animal generates heat largely through muscle shivering, which is followed by nonshivering thermogenesis (NST) in BAT (7). Mice were treated with rapamycin or vehicle for 48 hours prior to the cold (4°C) exposure. There was a dramatic increase in S6 phosphorylation in interscapular BAT (iBAT) (Figure 3A) and in s.c. inguinal WAT (iWAT) (Figure 3B). In mice that were pretreated with rapamycin, this phosphorylation was fully blocked. Similar results were observed in gonadal WAT (gWAT; not shown). Figure 3C shows that core body temperature dropped slightly in both groups as expected, but after 7 hours, the rapamycin-treated group exhibited a further decline in temperature, which became significant by 10 hours. At this time point, UCP1 protein level in iBAT tended to be lower with rapamycin treatment, with no effect on Ins signaling as measured by AKT phosphorylation (Supplemental Figure 6). However, given the whole-body exposure to rapamycin, this temperature change cannot be ascribed specifically to adipose tissue. In general, during this

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**P < 0.01

** Figure 3. Elevating SNS activity via cold exposure activates mTORC1 signaling in BATs and WATs. Male mice (n = 5/group) received either vehicle or rapamycin (Rapa; 4 mg/kg BW) by i.p. injection for 2 days followed by exposure to 4°C or 23°C for 6 hrs. (A and B) iBAT (A) and iWAT (B) were collected, homogenized, and assayed by immunoblot for p-S6 and total S6. GAPDH served as an internal control. Bar graphs show image quantification of p-S6 normalized to total S6. One-way ANOVA followed by post-hoc group comparisons; **P < 0.01. (C) Core body temperature was monitored and recorded for vehicle- or Rapa-treated mice during cold exposure. One-way ANOVA followed by post-hoc Bonferroni analysis. **P < 0.01. (D) iBAT and iWAT were dissected from cold-acclimated mice receiving either vehicle or Rapa for immunoblotting detection of UCP1 and internal-control GAPDH. (E) Core body temperature was recorded for vehicle- or Rapa-treated mice during cold adaptation (n = 4). C, control. One-way ANOVA followed by post-hoc Bonferroni analysis. *P < 0.05.
The presence of these cells becomes quite evident. Following longer exposure times, the robust levels of UCP1 in iWAT were greatly diminished by rapamycin. Although there is no apparent difference in iBAT UCP1 by Western blot between the vehicle and rapamycin groups, there nevertheless could be a net increase in total tissue mass and protein, including more overall UCP1. However, the weights of the iBAT were not different between the groups (4°C saline: 0.085 ± 0.021 mg vs. 4°C rapamycin: 0.077 ± 0.014 mg) (n = 4; t test; P = 0.605). Additionally, the rapamycin-treated cold-adapted mice displayed lower core body temperature, shown in Figure 3E.

Selective stimulation of adipose tissue by β3AR agonist. To more specifically activate βAR signaling only in adipose tissue, we used the well-established model of β3AR activation with the highly selective agonist CL316,243 (CL), in which chronic β3AR agonist treatment leads to the robust browning of WAT that is often accompanied by increased energy expenditure and resistance to obesity and its metabolic complications (48–51). Mice received CL either with or without rapamycin for 1 week as detailed in Methods.
ods. Upon dissection, gross examination of the adipose tissues revealed noticeable differences among the treatment groups. Adipose tissues from the CL-treated mice appeared to have a slightly darker complexion than those of the vehicle or rapamycin groups (Figure 4A). Histologically, CL treatment decreased lipid droplet size in iBAT and caused a substantial increase in multilocular adipocytes within the iWAT, along with a robust increase in UCP1+ staining (Figure 4B). Rapamycin dramatically suppressed the numbers of UCP1-staining cells in iWAT, and rapamycin alone was observed to have no effect. In gWAT, a similar suppression of UCP1 by rapamycin was observed as in iWAT but to a lesser extent. Figure 4C presents Western blots showing that, similar to cold exposure, CL treatment led to rapamycin-sensitive S6 phosphorylation in all 3 adipose depots. iWAT exhibited the most robust increase in UCP1 expression, which was prevented by rapamycin. Levels of NDUFS4, a component of respiratory complex I, measured here as a surrogate indicator of mitochondrial mass, was also increased in a rapamycin-sensitive manner. This is consistent with the need for significant mitochondrial biogenesis to support the increased respiration and uncoupling that occurs in brown adipocytes. In addition to Ucp1, peroxisome proliferation-activated nuclear receptor-α (PPARα), which is abundant in brown adipocytes, was increased in iWAT and gWAT by CL treatment, as were a number of established PPARα gene targets involved in fatty acid

Figure 5. Adipose-specific deletion of Raptor impairs brown/beige adipocytes during cold challenge. (A) Representative images of the iBAT, iWAT, and gWAT tissues from female floxed control (fl/fl) and adipose-specific Raptor KO (adKO). (B) Raptor, p-S6 and total S6, p-AKT (S473), total AKT, mitochondrial proteins UCP1, NDUFS4, COX4, endothelial marker VE-cadherin, and internal loading control GAPDH were detected by Western blotting in fat tissues of female Raptor<sup>fl/fl</sup> control and the adipose tissue-specific KO (adKO) at room temperature (RT) or cold exposure.
oxidation. The same pattern was observed in iWAT and gWAT: increased by CL and essentially blocked by rapamycin (Supplementary Figure 7). Taken together, these data from cold exposure and CL treatment suggest that mTORC1 signaling is necessary for the browning of WAT.

Adipose-specific deletion of Raptor blocks acquisition of brown/beige adipocytes. We generated mice with adipose-specific deletion of Raptor by crossing Raptor<sup>fl/fl</sup> mice with adiponectin-Cre-expressing mice. The Raptor<sup>fl/fl</sup> and Raptor<sup>Cre</sup> adiponectin-Cre (herein referred to as adKO) mice were subsequently subjected to a cold challenge. As reported earlier by Hall and colleagues (52), the adipose depots lacking RAPTOR were smaller in size (Figure 5A). They also contained fully mature adipocytes and expressed adipogenic markers such as PPAR<sub>γ</sub> and fatty acid binding protein 4 (aP2) (Supplemental Figure 8, A and B). Further analysis by Western blotting confirmed that Raptor was deleted in adipose tissues (Figure 5B), while it remained intact in other tissues (not shown). As expected with a lack of RAPTOR, levels of p-S6 were almost undetectable. For reasons that are unclear, total S6 was also reduced in the Raptor<sup>-null</sup> adipose tissues, while total amounts of AKT and VE-cadherin were unchanged between control and adKO adipose tissues. Importantly, the response to the cold challenge was lost in the adKO adipose tissues. The levels of UCP1 and indicators of mitochondrial content such as NDUSF4 and COX4 were much lower in all adipose depots of the Raptor<sup>-null</sup> mice after chronic cold exposure, while another mitochondrial marker, ATP5A, was unchanged in these adipose tissues between the adKO and Raptor<sup>fl/fl</sup> control mice. We also noted that, unlike treating adult mice with rapamycin, there was a blunted response observed in iBAT of adKO mice.
pressed by H89. Since cPKA can phosphorylate RAPTOR (Figure 6C), which is 150 kDa in size (54), we expressed myc-tagged RAPTOR in HEK293 cells and treated the cells with Fsk or Iso, with or without H89. Following IP of myc-RAPTOR and Western blotting with the PKA substrate antibody, Fsk and Iso treatment induced robust phosphorylation of myc-RAPTOR (Figure 6E), which was blocked by H89 treatment. We compared Iso with Ins treatment and examined this phosphorylation. While Iso clearly increased the PKA-motif phosphorylation of mTOR (Figure 6F) and Raptor (Figure 6G), there was no such effect of Ins treatment. These results indicate that the mTORC1 components mTOR and RAPTOR can be directly phosphorylated by PKA.

We next mutated the 3 Ser in mTOR (S1276, S1288, and S2112) and the Ser in RAPTOR (S791) to Ala. First, shown in Figure 6H, the triple mutant of mTOR, designated as S3A, could no longer be phosphorylated in vitro by cPKA. Next, WT mTOR and the S3A mutant were introduced into HEK293 cells. Following treatment of the cells with Iso for 30 minutes, mTOR was IP and analyzed for phosphorylation. Figure 6I shows that the S3A mutant of mTOR is refractory to the cAMP-elevating agents in cells. Second, in sharp contrast with WT RAPTOR, the Ser791A mutant could no longer be phosphorylated in vitro by cPKA (Figure 6J). We then compared the effects of WT RAPTOR and S791A RAPTOR in cells. In addition, although Ser792 is known to be an AMPK phosphorylation site likely because the deletion of Raptor occurs prenatally at the time the cells are becoming adipocytes during late gestation (53) and adiponectin-Cre begins to be expressed. This genetic evidence confirms a role for mTORC1 in the process of adipose browning and also suggests that there is a role for mTORC1 in the original development of the interscapular depot, which may or may not be dependent on PKA signaling.

**mTOR and its component RAPTOR are phosphorylated by PKA**

We next tested the possibility that mTORC1 might itself be a target of PKA. By inspecting the amino acid sequences of mTOR and RAPTOR across species, we indeed found that all mammalian mTOR proteins contain 3 strong and completely conserved RRXS motifs (Figure 6A) and that there is 1 motif in RAPTOR, as well (Figure 6B). Figure 6C shows that purified catalytic subunit of PKA (cPKA) can phosphorylate mTOR and RAPTOR, detected with an antibody that recognizes phosphorylated Ser or Thr in the context of the RRXS/T motif. Next, we expressed a myc-tagged mTOR in HEK293 cells and treated these cells either with Fsk or Iso for 30 minutes in the presence or absence of H89. As shown in Figure 6D, both Fsk and Iso increased PKA-substrate motif phosphorylation of myc-mTOR, which was blocked by H89. Figure 6D also shows a heavily phosphorylated band of approximately 150 kDa, which was also phosphorylated following PKA activation and suppressed by H89. Since cPKA can phosphorylate RAPTOR (Figure 6C), which is 150 kDa in size (54), we expressed myc-tagged RAPTOR in HEK293 cells and treated the cells with Fsk or Iso, with or without H89. Following IP of myc-RAPTOR and Western blotting with the PKA substrate antibody, Fsk and Iso treatment induced robust phosphorylation of myc-RAPTOR (Figure 6E), which was blocked by H89 treatment. We compared Iso with Ins treatment and examined this phosphorylation. While Iso clearly increased the PKA-motif phosphorylation of mTOR (Figure 6F) and Raptor (Figure 6G), there was no such effect of Ins treatment. These results indicate that the mTORC1 components mTOR and RAPTOR can be directly phosphorylated by PKA.

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Mutations of RAPTOR PKA site Ser791 have functional consequences for βAR signaling

To determine if phosphorylation of RAPTOR S791 is necessary for S6K1 activation mediated by βAR signaling, a lentivirus harboring an RAPTOR shRNA was used to infect NIH3T3 cells in order to knock down endogenous RAPTOR, as shown in Figure 7A. Using this maneuver, WT myc-RAPTOR or the S791A mutant myc-RAPTOR was introduced into HEK293 cells and were subsequently treated with Iso or Fsk. Compared with the robustly phosphorylated WT myc-RAPTOR, phosphorylation was eliminated in the mutants (Figure 6K) to an equal degree. The residual phosphorylation that is seen is not RAPTOR, but rather is a protein that is slightly smaller than RAPTOR and is captured nonspecifically by myc antibody-conjugated beads alone, since cells lacking a myc-RAPTOR construct still contain this faint phosphorylated band. Since mutation of Ser791 to Ala already disrupted the PKA-dependent phosphorylation of RAPTOR, AMPK and Ser792 are unlikely to be involved in β-agonist-dependent phosphorylation of RAPTOR. From this series of experiments, we conclude that mTOR and RAPTOR are direct substrates of PKA.

Discussion

The SNS is well known to activate PKA through βARs not only for lipolysis and uncoupled respiration in brown adipocytes through UCP1, but also for increasing the net amount of browned adipocytes present within white fat depots (8, 56–58). It has been repeatedly noted in laboratory animal models that the ability to generate these beige adipocytes is closely correlated with resistance to weight gain and preserved Ins sensitivity when challenged with a high-fat diet (48–50, 59–61). With the current intense interest in brown/beige adipose tissue in adult humans as a therapeutic target for increasing energy expenditure in metabolic disease, understanding the molecular events that drive their appearance and maintenance are critical. While the underlying cell lineage distinctions or molecular cues need further clarification (reviewed by ref. 62), increasing this UCP1-expressing adipocyte population is clearly a process that requires considerable synthetic and remodeling capacity within the tissue. This remodeling includes producing additional mitochondria and respiratory chain components, fatty acid oxidizing capacity, and, of course, UCP1. Here, we have shown that a direct signaling pathway from PKA to mTORC1 is involved in the β-adrenergic recruitment of these beige adipocytes in response to cold adaptation or to treatment with the β-AR agonist CL. In addition, our studies in mice lacking RAPTOR in adipocytes show impaired expression of UCP1 and other mitochondrial components, suggesting that there may be a role for mTORC1 even in the early development of the iBAT depot.

We previously showed that, downstream of βARs and PKA, other kinases are activated in adipocytes, which together coordinate to maximize fuel metabolism. These kinases include ERK (39) to contribute to overall lipolysis (41), as well as p38 MAPK (40, 63). While these MAPKs, particularly p38, have an important functional role for driving the transcription of the Ucp1 and Pgc1a genes (60, 63, 64), blockade of these MAPKs did not interfere with βAR activation of mTORC1. Instead, key components of mTORC1, specifically mTOR and RAPTOR, are directly phosphorylated by PKA. In mTOR, mutation of the 3 candidate PKA sites prevented its phosphorylation. Similarly, mutation of Ser791 in the RAPTOR PKA site eliminated its phosphorylation and functionally prevented activation of S6K1 by Iso but not by Ins.

In addition to UCP1 and mitochondrial genes, key components of the machinery necessary for the metabolic needs of brown and beige cells are also under the control of mTORC1. For example, PPARα is a master nuclear receptor for fatty acid β oxidation, and PPARα has been shown to also participate in UCP1 expression either directly or indirectly through ERα (65–70). In addition, several downstream target genes of PPARα and ERα are similarly under the control of mTORC1. These include Pdk4, Cpt1b, Fabp3, and Elov3. Clearly, a number of molecular events are set in motion to achieve the complex cellular process of adipose browning; some are dependent upon mTORC1, while others are not. Given that translational control is one of the hallmarks of biological regulation by mTOR (71), it would not be surprising if, along with direct transcriptional events, a layer of translational transcripts were significantly lower. Together, these results argue for an important role of PKA phosphorylation of RAPTOR in the acquisition of the Ucp1-expressing cell phenotype.

(55), some protein kinase substrate sequence prediction algorithms (e.g., Kinexus PhosphoNet) include Ser792 as a candidate PKA site. Therefore, we also generated the double-mutant S791A/S792A. All 3 of these myc-RAPTOR constructs (WT, S791A, and S791A/S792A) were introduced into HEK293 cells and were subsequently treated with Iso or Fsk. Compared with the robustly phosphorylated WT myc-RAPTOR, phosphorylation was eliminated in the mutants (Figure 6K) to an equal degree. The residual phosphorylation that is seen is not RAPTOR, but rather is a protein that is slightly smaller than RAPTOR and is captured nonspecifically by myc antibody-conjugated beads alone, since cells lacking a myc-RAPTOR construct still contain this faint phosphorylated band. Since mutation of Ser791 to Ala already disrupted the PKA-dependent phosphorylation of RAPTOR, AMPK and Ser792 are unlikely to be involved in β-agonist-dependent phosphorylation of RAPTOR. From this series of experiments, we conclude that mTOR and RAPTOR are direct substrates of PKA.
regulation turns out to be part of this mTORC1-driven effect on adipose tissue. This will be an important direction to explore in expanding these studies in the future.

The finding that PKA directly activates mTORC1 to drive adipose browning is both unexpected and rather extraordinary for several reasons. First, Ins and catecholamines are best known for their ability to antagonize one another’s metabolic effects, yet Ins is a well-known activator of mTORC1 for promoting growth and energy storage. From much work conducted in this field, inhibiting mTORC1 with rapamycin seems to be beneficial in terms of metabolism under some circumstances, but there are also complications. For example, while mTORC1 inhibition with rapamycin can increase longevity in mice (72, 73), chronic rapamycin treatment causes glucose intolerance and hyperlipidemia in both mice (74) and humans (75, 76); this appears to be caused by disturbances in mTORC2 function (77). Mice with aP2-driven deletion of Raptor in adipose tissue were reported to be resistant to diet-induced obesity, with smaller adipose tissue, increases in UCPI, and apparent improved Ins sensitivity (52, 78, 79). However, unlike our results described here, these earlier studies did not assess the effects of RAPTOR on the recruitment of brown adipocytes in response to cold or β-agonist stimulation. Importantly, the use of the aP2-Cre driver has subsequently been largely abandoned because it has been discovered that, in this animal, Cre recombinase can be expressed in other nonadipose tissues, including the brain, and other tissues during early development (64, 80, 81). Therefore, one cannot rule out the possibility that the loss of RAPTOR in some other tissue or cell type might contribute to the reported lean phenotype (52).

While it remains to be tested, our results here may contribute to understanding some other observations in the literature that suggest cAMP can activate mTORC1. For instance, thyroid-stimulating hormone (TSH) has been reported to activate S6K1 in thyroïd epithelia in a cAMP-dependent manner (82), and in pancreatic islets, GLP-1-dependent increases in cAMP — in the context of elevated glucose — were observed to activate S6K1 (83), which the authors concluded was due to cAMP-evoked intracellular Ca2+ release. More recently, in a disease model of primary pigmented nodular adrenocortical disease (PPNAD), which inactivates the regulatory subunit of PKA, it was reported that mTORC1 activity was increased, resulting in prevention of apoptosis (84). In 2 of these studies, the authors pointed to PRAS40 phosphorylation as a mediator, but whether this is the sole factor involved was not tested. In our studies here in adipocytes, phosphorylation of PRAS40 does not occur in response to β-agonist stimulation, thus eliminating it as a regulatory factor.

We believe that our study makes 2 important biological observations. First, that mTORC1 is a necessary component in the process of the browning of white adipose depots. The second point is that there exists a signaling pathway from PKA to mTORC1 activation via direct phosphorylation of mTOR and RAPTOR. Given these findings and the ubiquitous expression of mTOR and PKA in many cell types, we speculate that this PKA-mTORC1 pathway may have broader significance by regulating a variety of cell signaling events in other tissues. Therefore, our studies illustrate that an inherently catabolic process of energy expenditure in adipose tissue utilizes what is most commonly considered to be the molecular driver of growth and anabolic metabolism. When considered together, our findings suggest that the control of mTORC1 activity, and its contributions to metabolic regulation, deserves fresh appraisal.

**Methods**

**Reagents and antibodies.** Iso, Fsk, Ins, dexamethasone (Dex), isobutylmethylxanthine (IBMX), and the small molecule kinase inhibitors H89, SB, and PD were obtained from Sigma-Aldrich. The β,AR-selective agonist CL was a gift from Elliott Danforth Jr. (American Cyanamid Co., Pearl River, New York, USA). Rapamycin and torrin1 were from LC Laboratory. KT5720 was from Enzo Life Sciences. The protease inhibitor cocktail (complete Mini) and a phosphatase inhibitor cocktail (PhosSTOP) were from Roche Diagnostics. The following antisera were obtained from Cell Signaling Technology: p-S6 (Ser^240/244), catalog 2215, total S6 (catalog 2217), p-S6K1 (Thr^389, catalog 9205), total S6K1 (catalog 9202), p-AKT (T^308, catalog 9275), p-AKT (Ser^473, catalog 4060), total AKT (catalog 9272), p-PRAS40 (Thr^446, catalog 2640), total PRAS40 (catalog 2610), p-ULK1 (S^757, catalog 6888), total ULK1 (catalog 8054), p-ERK (Thr^202/Tyr^204, catalog 4370), total ERK (catalog 9102), p38 MAPK (catalog 9202), p-GSK3 (S^9 and S^21, catalog 8566), total GSK3 (catalog 5676), phospho-RRXS/T motif antibody (PKA substrate, catalog 9624), RAPTOR (catalog 2280), RICTOR (catalog 9476), “myc tag” (catalog 2272), COX4 (catalog 4844), and β-actin (catalog 4967). Other antisera used included UCP1 (catalog ab23841) and ATP5A (catalog ab14748) (both from Abcam). p-p38 MAPK (catalog ref-V121-A) and purified PKA catalytic subunit were obtained from Promega, plus NDUFS4 (Pierce Biotechnology, catalog PA5-21677), VE-cadherin (catalog sc-28644) and GAPDH (catalog sc-25788) (both from Santa Cruz Biotechnology). The EZview Red Anti-c-myc Affinity Gel (catalog E6654) and the secondary antibodies conjugated with alkaline phosphatase (anti-rabbit, catalog A3677; anti-mouse, catalog A3562) were from Sigma-Aldrich.

**Animal experiments.** Male C57BL/6j mice were obtained from the Jackson Laboratory. Experiments were conducted in mice that were 12–14 weeks of age. For the cold-challenge experiments, mice were administered rapamycin (5 mg/kg body weight [BW]/day) delivered in 100 μl/20 g BW or vehicle (75% saline, 10% ethanol, 10% PEG300, 5% Tween 80) once a day for 2 consecutive days. On the third day, the mice were housed individually and placed at 4°C for up to 10 hours, with fresh bedding and free access to water. Control mice were maintained at room temperature (23°C). For chronic cold challenge, mice were placed at 4°C for 7 days with daily rapamycin (2.5 mg/kg BW/day) administration. For selective activation of β,AR, mice were injected i.p. with CL (1 mg/kg BW/day) (86, 87) with or without rapamycin (2.5 mg/kg BW) for 1 week, after which they were euthanized, and iBAT, iWAT, and gWAT were dissected and immediately frozen in liquid nitrogen. Mice with a Raptordon/° allele (JAX-031888), as well as adiponectin-Cre (JAX-010830), were obtained from the Jackson Laboratory and bred to generate mice with a Raptor adipose-specific deletion. Tail DNA genotyping and Western blotting of lysate from adipose tissue, liver, and heart were used to confirm the specific deletion of Raptor in adipose.

**Adipocyte cell culture.** 3T3-L1 preadipocytes were obtained from ATCC. They were maintained and differentiated into adipocytes as described (88), with modifications (89). HEK293 cells were purchased from ATCC and cultured in DMEM with 10% FBS. Human primary s.c. adipocytes were provided by Zen-Bio. NIH3T3 cells were purchased from ATCC and cultured in DMEM supplemented with 10%
calf bovine serum. Prior to treatment, cells were washed twice with serum-free DMEM and maintained overnight in DMEM with 0.1% fatty-acid-free BSA cells. HIB-1B preadipocytes were cultured and differentiated as described (87, 90). HIB-1B cells stably expressing RAPTOR WT, S791A (alanine) or S791D (aspartic acid) were generated as previously described (91, 92).

Plasmids and mutagenesis. The WT myc-mTOR and myc-RAPTOR mammalian expression vectors were from Addgene. To mutate candidate P6A sites in mTOR, the following primers were used to change the Ser residues to Ala: S2112A forward: 5′-TTT AGA CGG ATC GCC AAG CAG CTA-3′ and S2112A reverse: 5′-TAG CTG CTT GGC GAC TGG CTC GCT GAA-3′; for S1276A/S1288A/S2112A triple mutation, the pair of primers forward: 5′-CTG CTA GGC TGG GCT CTT CTG GAG GTC TTC AGA CGG ATC GCC AAG CAG GCG TCC TAC TCC TCC CTC TCC AAC TCC TCC CTC AAC TCC CTC TCC-3′ and reverse: 5′-CTT CAG CAG CTC CAG ACG CAA GCG TCG CAG CAG-3′ was used and combined with 5′-GCC AGA AGG GTC GGC AAG GAC GAC TGG-3′. To mutate RAPTOR Ser791 to Ala, the primers were RAPTOR S791A forward: 5′-GAC GAC ATG CGC GCC GCC GCC TCC TAC TCC TCC CTC AAC TCC CTC-3′ and RAPTOR S791A reverse: 5′-GGG GGT GAG GGA GGT GAG GCC GCC GCC CAT CTT GTC. For the RAPTOR S791A/S792A mutation, forward: 5′-GAC AAG ATG CGC GCC GCC GCC TCC TAC TCC TCC CTC AAC TCC CTC-3′ and reverse: 5′-GGG GGT GAG GGA GGT GAG GCC GCC GCC CAT CTT GTC-3′. To mutate RAPTOR S791 to aspartic acid (S791D), we changed S791A forward primer from 5′-GGG GGT GAG GGA GGT GAG GCC GCC GCC CAT CTT GTC-3′ to 5′-GGG GGT GAG GGA GGT GAG GCC GCC GCC CAT CTT GTC-3′. To mutate RAPTOR S791 to aspartic acid (S791D), we changed S791A forward primer from 5′-GGG GGT GAG GGA GGT GAG GCC GCC GCC CAT CTT GTC-3′ to 5′-GGG GGT GAG GGA GGT GAG GCC GCC GCC CAT CTT GTC-3′. The WT myc-mTOR and the S791A mutant RAPTOR, or myc-tagged mTOR cells or tissues were lysed Transfection, IP, and Western blotting. Cells or tissues were lysed and sonicated in buffer containing 25 mM HEPES (pH 7.4), 150 mM NaCl, 5 mM EDTA, 5 mM EGTA, 5 mM glycero phosphate, 0.9% Triton X-100 (Sigma-Aldrich), 0.1% Igepal CA630, 5 mM sodium pyrophosphate, and 10% glycerol, plus 1 tablet each of complete protease inhibitor cocktail and PhosSTOP phosphatase inhibitors per 10 ml. For IP, HEK293 cells were transfected with myc-tagged mTOR or RAPTOR expression plasmid using lipofectamine 2000 (Invitrogen) according to the instructions provided. Cells were collected and lysed in the above buffer, and 2 mg total protein was incubated with anti–c-myc affinity gel overnight (Sigma-Aldrich). Beads were washed thoroughly in lysis buffer. For the in vitro PKA phosphorylation assay, 2 mg total protein was incubated with anti–c-myc affinity gel for 4 hours. Beads were washed twice with the lysis buffer and once with the PKA reaction buffer (40 mM Tris-HCL [pH 7.4], 20 mM magnesium acetate, 0.2 mM ATP), and PKA catalytic subunit was added to the beads. Protein phosphorylation by PKA was measured with PKA substrate antibody (phospho-RXXS/T motif antibody; Cell Signaling Technology). FLAG-tagged WT RAPTOR and the S791A mutant RAPTOR, or myc-tagged WT RAPTOR and S791D mutant, were introduced into HIB-1B cells, and single clones were selected with 5 μg/ml blastcidin for FLAG-RAPTOR cells or 100 μg/ml hygromycin for myc-RAPTOR cells to generate stable cell lines. Unless otherwise indicated, 50 μg total protein was resolved in 4%–20% Tris-glycine gels (Invitrogen) and transferred to nitrocellulose membranes (Bio-Rad) and incubated overnight at 4°C with specific primary antibodies. Secondary antisera conjugated with alkaline phosphatase were used for specific protein detection. To repro- blots, membranes were stripped in buffer (62.5 mM Tris, 2% SDS, 100 mM β-mercaptoethanol) at 37°C for 60 minutes. Image acquisition was performed on a Typhoon FLA 9000 imager (GE Healthcare), and data were analyzed using ImageQuant TL software.

RNA isolation and quantitative PCR. RNA from adipose tissues were first extracted by Trizol and purified by RNA mini-columns from Qiagen. Reverse transcription and SYBR green quantitative PCR (qPCR; Invitrogen) were performed according to the manufacturer protocols. Target primer sequences are shown in Supplemental Table 1.

Tissue histology and microscopy. Adipose tissues were fixed with 4% paraformaldehyde in PBS, dehydrated, embedded in paraffin, and sectioned (5-μm thickness). Sections were stained with H&E and examined under bright-field microscopy with a Nikon 80i. UCPI staining was performed as described previously (60, 93).

Statistics. All data are presented as mean ±SEM. Comparisons between groups were assessed by 1-way ANOVA with Tukey’s post-hoc test or unpaired Student’s t test (2-tailed) as indicated in figure legends.

Study approval. All animal experiments were approved by the Sanford Burnham Prebys Medical Discovery Institute’s Institutional Animal Care and Use Committee in accordance with the 8th edition of the NIH Guide for the Care and Use of Laboratory Animals.

Note added in proof. While this manuscript was in press, an independent report confirmed our finding of a positive role for mTORC1 signaling in the browning of white fat (94).


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