The Emerging Importance of Mitochondria in White Adipocytes: Neither Last nor Least

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The growing recognition of mitochondria’s crucial role in the regulation of white adipose tissue remodeling and energy balance underscores its significance. The marked metabolic diversity of mitochondria provides the molecular and cellular foundation for enabling adipose tissue plasticity in response to various metabolic cues. Effective control of mitochondrial function at the cellular level, not only in thermogenic brown and beige adipocytes but also in energy-storing white adipocytes, exerts a profound influence on adipose homeostasis. Furthermore, mitochondria play a pivotal role in intercellular communication within adipose tissue via production of metabolites with signaling properties. A more comprehensive understanding of mitochondrial regulation within white adipocytes will empower the development of targeted and efficacious strategies to enhance adipose function, leading to advancements in overall metabolic health.

Keywords: White adipocyte; Mitochondria; Adipose tissue remodeling; Metabolic health; Signaling metabolite

INTRODUCTION

Mitochondria, often referred to as the “powerhouses of the cell,” are remarkable organelles essential for cellular energy production and a host of other vital functions [1-4]. Present in the cytoplasm of eukaryotic cells, these double-membrane-bound structures are derived from ancient bacteria that were engulfed by early eukaryotic cells in a symbiotic relationship. Their primary and most renowned role lies in generating adenosine triphosphate (ATP) through oxidative phosphorylation, a process that efficiently converts nutrients, such as glucose and fatty acids, into usable energy. This metabolic pathway occurs within the inner mitochondrial membrane and is fueled by the electron transport chain and the citric acid cycle. However, mi-
Mitochondria’s significance extends beyond energy production; they are also involved in regulating cellular metabolism, controlling calcium homeostasis, participating in apoptosis (programmed cell death), and contributing to various signaling pathways [5-9]. Understanding the intricacies of mitochondria and their functions is crucial for comprehending cellular physiology, energy metabolism, and their broader implications for human health and disease.

White adipocytes are a crucial component of human body’s metabolic machinery. As specialized cells primarily found in adipose tissue, these adipocytes play a central role in energy regulation and storage [10-12]. Their primary function revolves around the storage of triglycerides, which are synthesized from excess dietary fats and carbohydrates. In times of energy surplus, white adipocytes store triglycerides in the form of lipid droplets, effectively acting as energy reservoirs [13,14]. Conversely, during periods of energy demand, such as fasting or physical activity, these adipocytes release stored triglycerides through lipolysis, providing a vital source of energy for the body’s various physiological processes [15]. Beyond their role in energy storage, white adipocytes also secrete a variety of hormones and adipokines that impact systemic metabolism, inflammation, and appetite regulation [16,17].

Thermogenic brown and beige adipocytes possess a high number of mitochondria, guaranteeing their capacity to uphold distinct bioenergetic processes and effectively transform stored energy into heat [18-20]. Meanwhile, a growing body of evidence is substantiating the critical role of mitochondria in white adipocytes, despite the presence of fewer mitochondria in these cells [21,22].

Here, we review the current knowledge about the role of mitochondria in white adipocytes, focusing on the mitochondrial regulation of adipocyte formation and function. We also discuss the emerging evidence linking mitochondrial signaling metabolites and white adipocyte identity.

**ROLE OF MITOCHONDRIA IN THERMOGENIC ADIPOCYTES**

Mitochondria play a pivotal role in the metabolic dynamics of both brown and beige adipocytes, specialized fat cells that hold significant implications for energy homeostasis and metabolic health [23,24]. Brown adipocytes, prominent in rodents and human infants but also present in human adults, are characterized by a high number of mitochondria and an abundance of iron-rich proteins, giving them their distinctive brown color [25-27]. These mitochondria contain a unique mitochondrial protein called uncoupling protein 1 (UCP1), which enables the dissipation of proton gradients generated during oxidative phosphorylation (OXPHOS) [28,29]. This uncoupling mechanism facilitates the generation of heat rather than ATP, contributing to the thermogenic capacity of brown adipocytes. As a result, brown adipose tissue acts as a natural thermal regulator, aiding in maintaining body temperature and increasing energy expenditure.

In recent years, beige adipocytes have garnered attention for their capacity to transform into a “brown-like” phenotype in response to various stimuli, such as cold exposure or certain hormonal cues [18,20]. These cells emerge within white adipose tissue (WAT) depots and share similarities with brown adipocytes, including the presence of multilocular lipid droplets and increased mitochondrial content. The mitochondria in beige adipocytes also express UCP1, albeit to a lesser extent than classical brown adipocytes, contributing to their thermogenic capability [20,29]. Notably, creatine, a naturally occurring compound primarily known for its role in cellular energy metabolism, has also garnered attention for its potential involvement in thermogenesis [30]. While traditionally associated with enhancing muscular performance by replenishing ATP, recent research has indicated that creatine may impact thermogenesis through its interactions with mitochondria [31,32]. Studies have suggested that creatine supplementation could enhance mitochondrial bioenergetics and heat production [31,32]. In both brown and beige adipocytes, the central role of mitochondria in thermogenesis and metabolic regulation underscores their importance in maintaining overall energy balance and metabolic health.

**ROLE OF MITOCHONDRIA IN WHITE ADIPOCYTE FUNCTION**

White adipocytes, despite having a lower abundance of mitochondria compared to other types of adipocytes, rely on functional mitochondria as crucial guardians to maintain health and regulate various aspects of adipocyte biology. At a homeostatic level, mitochondria play essential roles in ensuring cellular survival, controlling adipocyte differentiation, maintaining glucose and lipid homeostasis, and orchestrating metabolism of branched-chain amino acids (BCAAs). Additionally, functional mitochondria in white adipocytes significantly contribute to glucose homeostasis and adipokine secretion, and are involved in the browning process, which transforms white adipocytes into thermogenic brown-like adipocytes.
Like other mitochondria-containing cells, white adipocytes heavily depend on ATP produced by mitochondria to sustain essential metabolic processes such as triglyceride synthesis, gluconeogenesis, and fatty acid re-esterification for survival [33]. This tight relationship is attributed to the mitochondrial energy center, serving as both the hub for cellular energy metabolism and the exclusive site for essential enzymes involved in these anabolic processes [34-36]. For instance, mitochondria play crucial roles in lipogenesis and lipolysis in white adipocytes. This organelle produces critical metabolic intermediates, like citrate, which is essential for the lipogenic process to function optimally [21]. Additionally, mitochondria provide the required environment for lipogenesis and lipolysis, particularly accommodating medium-chain fatty acids [37]. ATP production by mitochondria governs the efficiency of these metabolic pathways. Furthermore, mitochondria actively participate in phospholipid synthesis, further highlighting their multifaceted role in lipid metabolism [38].

Leucine, isoleucine, and valine are BCAAs and are catabolized only in mitochondria. Given the important influence of adipocytes on BCAA metabolism, mice with deletion of mitochondrial branched-chain amino acid aminotransferase 2 (Bcat2) exhibit an intriguing metabolic profile despite consuming more food. Specifically, these mice show elevated plasma levels of BCAAs and a remarkable reduction in adiposity and body weight. This reduction is accompanied by an increase in energy expenditure, significant improvement in glucose tolerance and insulin sensitivity, and notable protection against diet-induced obesity [39].

Emerging research emphasizes the pivotal role of mitochondria in governing glucose homeostasis within white adipocytes, particularly through their influence on insulin actions. Notably, when mitochondrial respiratory inhibitors or uncoupling reagents are administered, there is a significant reduction in insulin-stimulated glucose uptake in adipocytes [40].

Additionally, in rats fed a high-fat diet (HFD), mitochondrial biogenesis and mitochondrial DNA (mtDNA) copy numbers are downregulated in WAT, leading to elevated glucose levels and diminished glucose uptake by white adipocytes [41].

Mitochondria also play a significant role in adipokine synthesis and secretion, with mitochondrial biogenesis being closely linked to this process. Stimulation of mitochondrial biogenesis through treatments like endothelial nitric oxide synthase activation or overexpression of nuclear respiratory factor-1 (Nrf1) can enhance the synthesis of adiponectin, a critical adipokine [42,43]. Interestingly, adiponectin overexpression reciprocally leads to an increase in mitochondrial density in white adipocytes, creating a regulatory loop between mitochondria and adipokine synthesis [44].

With the advancement of sequencing technologies, a few studies have investigated the heterogeneity of white adipocytes in mice and humans using single-nucleus RNA sequencing (snRNA-seq) [45-47]. Furthermore, despite limited evidence, it is suggested that the heterogeneity of white adipocytes may be determined by the activity of mitochondria, such as a subpopulation of white adipocytes in mice displaying higher expression of genes associated with mitochondrial activity [45]. Moreover, a snRNA-seq study on human white adipocytes revealed two distinct clusters: one with higher gene expression related to mitochondrial capacity and enhanced mitochondrial function, and the other with lower gene expression related to mitochondrial function and reduced mitochondrial activity, respectively [47].

The mitochondrial unfolded protein response (UPRmt) has a protective effect on white adipocytes, as it serves as a mechanism for mitochondria to manage stress and maintain homeostasis [48,49]. Several studies have demonstrated that UPRmt also plays a crucial role in fat metabolism and protects against obesity. In a cohort study, UPRmt-related genes, including C/EBP homologous protein (Chop), heat shock protein 60 (Hsp60), caspase-9, and Htra2 serine peptidase (ClpP), are significantly reduced in higher-weight identical twins [50]. Moreover, several genes of the UPRmt-related sirtuin family, namely Sirt1/2/3, are involved in the regulation of adipose tissue metabolism [51-56]. It is worth noting that not all UPRmt-related genes are beneficial to the organism, such as ClpP and Park2 (the PARKIN-encoding gene) [57,58]. Deletion of these two factors has been found to result in a beneficial metabolic phenotype.

**WHITE ADIPOCYTE MITOCHONDRIAL HOMEOSTASIS AND METABOLIC HEALTH**

While the above evidence implies an important role of mitochondrial homeostasis in maintenance of white adipocyte function and health, mitochondrial homeostasis is often disturbed by stress conditions, such as aging, obesity, and other metabolic challenges. When mitochondria turn dysfunctional, they can have adverse effects on white adipocytes and the metabolic homeostasis of the organism. A number of studies have demonstrated that dysfunctional mitochondria in white adipocytes can lead to abnormal lipid metabolism, inflammation, impaired glucose metabolism, dysregulated adipokine secretion, and apo-
For example, adipocyte-specific overexpression of amyloid-beta precursor protein (App) has been linked to white adipocyte mitochondrial dysfunction. This results in the inhibition of catecholamine-stimulated lipolysis, leading to significant white adipocyte hypertrophy, increased inflammation, and fibrosis in WAT. These mice also show insulin resistance, glucose intolerance, and sensitivity to obesity [59]. Another critical player in white adipocyte mitochondrial function is the mitochondrial dicarboxylate carrier (mDIC), mainly expressed in WAT and encoded by the Slc25a10 gene. mDIC transfers succinate from the mitochondrial matrix to the cell membrane, where it interacts with the succinate receptor (SUCNR1), blocking the production of cyclic adenosine monophosphate (cAMP) and subsequently suppressing the hormone-sensitive lipase pathway, leading to decreased lipolysis. Dysfunctional mDIC-mediated succinate transport in white adipocytes during obesity results in the release of fatty acids from WAT to the liver, causing adipocyte dysfunction and liver lipotoxicity, thereby impacting the development of nonalcoholic fatty liver disease and systemic insulin resistance [60].

Mitochondrial dysfunction may have complex effects on the system’s metabolism, such as transcription factor A, mitochondrial (TFAM). Knocking out Tfam specifically in adipocytes leads to mitochondrial dysfunction, decreases weight gain induced by age or diet, impairs glucose tolerance, lowers energy expenditure, promotes inflammation and apoptosis in WAT, and reduces circulating adiponectin and leptin levels. These mice are protected from obesity but are sensitive to hepatosteatosis, hypertension, and cardiac dysfunction [61]. When ferritin mitochondrial (Ftmn), which is a mitochondrial matrix protein that chelates iron, is overexpressed specifically in adipocytes, mice are subjected to dietary challenges, resulting in leaner body weights. However, these mice exhibit dysfunction of adipose metabolism evidenced by glucose intolerance, reduced adiponectin levels, increased damage from reactive oxygen species (ROS), elevated levels of growth/differentiation factor 15 and fibroblast growth factor 21, and decreased circulating and intracellular adiponectin levels. Interestingly, despite the severe adiposity dysfunction in the transgenic mice, a noteworthy outcome is observed—they display significant β-cell proliferation, suggesting a beneficial mitochondria-induced adipose-pancreatic inter-organ signaling axis [62].

Moreover, mitochondrial dysfunction induced by the deficiency of genes such as CR6-interacting factor 1 (Crf1), FUN14 domain-containing protein 1 (Fundc1), and NADH dehydrogenase (ubiquinone) iron-sulfur protein 4 (Ndufs4) has been associated with increased inflammatory cell infiltration, enhanced inflammatory gene programs and insulin resistance/glucose intolerance [63-65]. Additionally, thymidine kinase 2 (Tk2)-deficiency dramatically reduces circulating levels of leptin and resistin [66]. White adipocytes exposed to mitochondrial dysfunction induced by the administration of oligomycin A and antimycin A and by knockdown of mitochondrial transcription factor A (mtTEA) exhibit a dose-dependent downregulation of adiponectin expression and impaired glucose homeostasis [40]. Iron metabolism also plays an essential role in mitochondrial function, and the manipulation of iron-related genes can lead to mitochondrial dysfunction and abnormal changes in white adipocytes, affecting systemic energy homeostasis. For example, overexpression of Asn-Glu-Glu-Thr (mitoNEET), an outer mitochondrial membrane protein and iron-sulfur cluster transfer protein, causes mitochondrial dysfunction, resulting in increased adipose tissue mass due to enhanced lipid uptake and storage. These mice also display enhanced insulin sensitivity and higher expression of adiponectin [67]. Furthermore, the response of these mice to an HFD challenge is dynamic, with short-term exposure leading to enhanced browning signature in subcutaneous WAT and limited expansion, but with prolonged exposure causing diminished browning and rapid WAT expansion and weight gain [68]. Studies on other iron-related genes, ferritin heavy chain (Fth), and Ftmn, have also provided evidence for the crucial role of mitochondrial iron in WAT functionality [62,69].

The evidence obtained from rodent models is exciting; however, data from human studies are even more valuable regarding translational implications. Numerous population studies have provided compelling evidence for the significant association between adipocyte mitochondrial function and obesity. For instance, lower body weight and higher mitochondrial respiration rate and mitochondrial number in WAT show strong correlations [70]. Comparative studies involving identical twins have revealed that the obese twins had significantly lower mitochondrial mtDNA than their non-obese counterparts, despite both having the same mtDNA sequence [71]. A research study conducted in Japanese and Italian populations has demonstrated a significant association between mtDNA variation at position 15497 guanine/adenine (Mt15497G→A) and lipid metabolism as well as obesity-related factors. These factors include body weight, body mass index, waist circumference, hip circumference, and intra-abdominal fat [72,73]. Another study has investigated the status of subcutaneous WAT mitochondrial OXPHOS in four groups: non-obese individuals, non-obese individuals with type 2 diabetes mellitus (T2DM), nonobese obese individuals, and
obese individuals with T2DM. In both non-obese and obese T2DM subjects, mitochondrial transmembrane potential, inorganic phosphate utilization, and electron transport chain activity are significantly reduced compared to the corresponding non-obese individuals without T2DM. Notably, the mitochondrial index, a measure of mitochondrial function, is significantly higher in lean individuals with T2DM than in lean individuals without diabetes. Furthermore, in people with obesity and T2DM, respiratory chain activities (specifically, complex I, I to III, II to III) and phosphorylation capacity of white adipocyte mitochondria are markedly lower than in obese individuals without T2DM [74]. Collectively, these findings indicate that obesity itself can have a detrimental impact on white adipocyte mitochondrial function, leading to various metabolic disturbances associated with obesity.

FUNCTIONAL ROLE OF MITOCHONDRIA IN WHITE ADIPOCYTE BROWNING/BEIGING

The white-to-beige adipocyte transition, known as browning or beiging, is a process driven by the expression of specific genes that confer a more thermogenic and metabolically active phenotype, characterized by the induction of thermogenin (UCP1) expression and the formation of brown adipocytes. This transition is induced by various environmental cues, such as cold exposure, β-adrenergic stimulation, or metabolic stress, and is associated with increased mitochondrial biogenesis and function, leading to increased energy expenditure and thermogenesis [75-77].

In addition to the crucial role of mitochondria in white adipocyte function, a growing number of studies suggest that mitochondrial regulation is pivotal for the cell lineage fate of adipocyte progenitors and the formation of mature white adipocytes. Given the above-mentioned mitochondrial role in BCAA metabolism, a substantial increase in leucine catabolism and expression of enzymes involved in the BCAA catabolic pathway observed with increasing mitochondrial numbers during adipogenesis [75,76]. Adipogenesis is also concomitantly accompanied by heightened mitochondrial biogenesis, activity, and the generation of ROS [21,33]. Consequently, the differentiation process reprograms the energy status from glycolytic to oxidative metabolism, thus acquiring and sustaining a mature adipocyte phenotype [77]. Consequently, any malfunction in mitochondrial metabolism or disruption in ROS production, induced either through genetic manipulation of mitochondrial constituents or employment of pharmacological inhibitors, is sufficient to perturb adipogenesis, evidenced in studies using cultured preadipocyte cell lines and the stromal vascular fraction of adipose tissues [78-80].

With recent discoveries of adipose tissue-resident adipose progenitor cells (APCs), more precise and targeted evidence pertaining to mitochondrial regulation of their lineage potential has come to light [81]. Of note, unbiased quantitative proteomic analysis of murine APC populations has revealed significant depot- and sex-dependent differences in the expression of mitochondrial proteins and regulators of lipid metabolism [82,83].

Functionally, impaired mitochondrial activity (achieved by a physiologically relevant reduction via ectopic expression of the mitoNEET protein, the above-mentioned outer mitochondrial membrane protein governing oxidative capacity) exerts a profound influence on the behavior of platelet-derived growth factor receptor (PDGFR) β+ adipose progenitors, driving them to adopt a pro-inflammatory phenotype at the cost of their adipogenic capacity in adult mice [84].

MITOCHONDRIAL ROLE IN DETERMINING WHITE ADIPOCYTE IDENTITY

White adipocytes possess remarkable phenotypical plasticity and dynamism, allowing them to undergo a transformative process known as browning or beiging when exposed to certain conditions such as cold temperatures, activation of β-adrenergic receptors, exercise or diet [85-89]. During this process, white adipocytes adopt characteristics similar to brown adipocytes, resulting in the induction of beige adipocytes. Evidently, a crucial aspect of this browning/beiging process involves increasing the number of mitochondria and enhancing mitochondrial biogenesis. This mitochondrial enhancement is a key factor in facilitating the transition of white adipocytes towards a more thermogenic and metabolically active phenotype, thus promoting energy expenditure and metabolic health [90].

Mitochondrial double-membraned structures encompass a large variety of chemical reactions that occur within an organism to sustain life, including the breakdown of nutrients and the synthesis of essential molecules [2,91]. Metabolites, the small molecules involved in these metabolic processes, serve as building blocks, energy sources, and signaling molecules that regulate various cellular activities [92,93]. Emerging evidence is showing that mitochondrial signaling metabolites act as intricate messengers, transmitting information about cellular status, environmental cues, and metabolic processes, ensuring the efficient cell fate determination of different cells, including adipocyte lineage cells. By influencing enzymatic activities, gene expression, and signaling pathways, signal metabolites orchestrate a wide array of physiological responses, spanning from adaptation to stress to the coordination of growth and differentiation [5]. Succinate, a key metabolite in the tricarboxylic acid cycle, has recently garnered significant attention for its intriguing role in the modulation of WAT physiology [60,94]. Emerging research has unveiled a novel connection between succinate and the conversion of white adipocytes to beige adipocytes, a process known as white-to-beige adipocyte conversion. It is pro-
posed that succinate is involved in the activation of pathways related to beige adipogenesis and thermogenesis [95].

Our recent work highlighted that mitoprotease Lon protease 1 (LONP1)-dependent metabolic enzyme turnover controls white-to-beige adipocyte conversion via fine-tuning intracellular succinate concentration [96]. We observed an increase in mitochondrial protein turnover, coupled with dynamic adjustments in the levels of the pivotal mitoprotease LONP1, in response to thermogenic stimulation. We uncovered the selective degradation of succinate dehydrogenase B (SDHB), a crucial element of mitochondrial complex II, by LONP1. Following thermogenic stimulation, this LONP1-mediated SDHB degradation led to an elevation in intracellular succinate levels. The resultant rise in the succinate to α-ketoglutarate ratio fostered enhanced histone methylation on thermogenic genes, thereby promoting the shift from white-to-beige adipocytes. Notably, elevating LONP1 expression induced heightened intracellular succinate levels, effectively rejuvenating beige cell conversion and adaptive thermogenesis in aged adipocytes and mice.

**CONCLUSIONS**

In recent years, extensive research has been directed towards unraveling the intricacies of mitochondrial regulation in maintaining metabolic homeostasis within adipocytes, which includes the emerging body of evidence that highlights the notable importance of mitochondria within white adipocytes (Fig. 1). These advancements have greatly enhanced our comprehension of the pivotal roles played by mitochondria in ensuring the overall health of adipose tissue. The orchestration of mitochondrial activity is indispensable in shaping the functionality of mature adipocytes, while also exerting a profound influence on the microenvironment within WAT.

The exploration of therapeutic applicability of these findings remains at the forefront in the field of WAT mitochondrial research. A comprehensive analysis of the temporal progression...
Each WAT mitochondrial dysfunction model documented in prior research becomes instrumental in ascertaining the optimal window for initiating therapeutic interventions aimed at remedying the observed mitochondrial aberrations. Understanding the complex interplay between mitochondrial function and adipocyte physiology as well as its implication in the development of pathological WAT microenvironment is another forefront of adipocyte mitochondrial biology.

In addition, mitochondrial metabolism is emerging as a key hub of signal metabolite production, linking mitochondrial functional normality to active regulation of various biological processes, such as cell fate determination, beyond traditional views of mitochondrial metabolism that is passively adapted to meet energy demands of the cell. Further understanding of the intricate interplay between signaling metabolites (e.g., succinate), adipocyte biology, and energy metabolism holds promise for the development of innovative therapeutic strategies aimed at enhancing energy expenditure and combating obesity.

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

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