Review

The role for adipose tissue in weight regain after weight loss

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Summary

Weight regain after weight loss is a substantial challenge in obesity therapeutics. Dieting leads to significant adaptations in the homeostatic system that controls body weight, which promotes overeating and the relapse to obesity. In this review, we focus specifically on the adaptations in white adipose tissues that contribute to the biological drive to regain weight after weight loss. Weight loss leads to a reduction in size of adipocytes and this decline in size alters their metabolic and inflammatory characteristics in a manner that facilitates the clearance and storage of ingested energy. We present the hypothesis whereby the long-term signals reflecting stored energy and short-term signals reflecting nutrient availability are derived from the cellularity characteristics of adipose tissues. These signals are received and integrated in the hypothalamus and hindbrain and an energy gap between appetite and metabolic requirements emerges and promotes a positive energy imbalance and weight regain. In this paradigm, the cellularity and metabolic characteristics of adipose tissues after energy-restricted weight loss could explain the persistence of a biological drive to regain weight during both weight maintenance and the dynamic period of weight regain.

Keywords: Adipogenesis, dieting, obesity, weight regulation.

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Introduction

Over 60% of adults and close to 20% of children in the United States are overweight or obese (1,2). Weight loss strategies are only transiently effective for most people, as the vast majority of individuals who attempt to lose weight are not able to achieve and maintain a 10% reduction over a year (3). Over a third of lost weight tends to return within the first year and the majority is regained back within 3 to 5 years (4,5). A number of reasons have been proposed for the high recidivism rates (5,6), but there is substantial evidence for a biological drive to regain weight after weight loss (7,8). The objective of this review is to summarize the contribution of white adipose tissue to this biological drive and discuss how changes in its cellularity and metabolic characteristics may facilitate weight regain.

The biological drive to regain weight

The biological control of body weight involves a complex feedback loop between the brain and periphery. The brain receives signals from the periphery regarding long-term energy stores (i.e. adipose tissue triglyceride) and short-term nutrient availability (i.e. immediate availability of circulating nutrients) and based upon these integrated signals, adjusts energy balance to meet both the long-term and short-term objectives of energy homeostasis. This feedback system adapts when energy intake is cognitively (in humans) or forcefully (in animal models) restricted.

In a previous review (7), we summarized the adaptations to energy-restricted weight loss that are thought to promote weight regain (Fig. 1). This adaptive response involves coordinated changes in the brain, gut, muscle, liver, adipose
tissue and neuroendocrine system, which culminate in a concerted effect on energy balance. Peripheral signals create an ‘anabolic’ neural profile in the hypothalamus and hindbrain, increasing appetite and sending neuroendocrine efferent signals to enhance metabolic efficiency in peripheral tissues. Metabolic requirements decline as a function of (i) lost mass, (ii) reduced consumption of food and (iii) increased metabolic efficiency of peripheral tissues. Peripheral tissues clear circulating nutrients more effectively and utilize fuels more efficiently to produce the energy they need. Signals from the periphery convey to the brain that energy stores are depleted and nutrient availability is low and these signals integrate in key circuits of the hypothalamus and hindbrain that serve as the primary control centres for energy balance regulation. The response to these integrated signals is that appetite increases and the expenditure of energy declines. We have referred to this quantitative difference between the caloric value reflecting appetite and expenditure requirements as the energy gap (9–11). To maintain the reduced weight, food intake must be cognitively (in humans) or forcefully (in animals) restricted to the level that expended energy is suppressed. During weight maintenance after weight loss, this energy gap reflects the magnitude of the daily burden that thwarts cognitive efforts to maintain the reduced weight. When efforts to restrict intake fail, overfeeding occurs, and the relapse to obesity begins. This pressure to continue to overfeed generally persists until the lost weight returns. In some cases, the biological pressures may lead to weight gain that surpasses the original weight.

A fundamental understanding of this energy gap, dictated solely by biological pressures, has emerged from preclinical studies of weight regain in diet-induced obesity (DIO) models. The energy gap at the maintenance-relapse transition is influenced in predictable ways by diet composition (12), by the length of time in weight maintenance after weight loss (10) and by physical activity levels (13). Weight regain driven solely by this biological pressure reflects a first-order growth curve (4,11,13) such that the energy gap at the maintenance-relapse transition is influenced in predictable ways by diet composition (12), by the length of time in weight maintenance after weight loss (10) and by physical activity levels (13).
energy gap diminishes as the relapse to obesity progresses. As such, the magnitude of the energy gap is greatest at the nadir weight after weight loss (9,11,13). Furthermore, this energy gap does not dissipate with time in weight maintenance. Rather, studies indicate that the magnitude of the energy gap gradually increases the longer an animal maintains their reduced weight with an energy-restricted diet (10). The implications from these observations are that the biological pressures may strengthen with time during weight maintenance and with the amount of weight lost.

White adipose tissue is a critical node in the homeostatic system that controls body weight and it plays a particularly important role in the biological drive to regain lost weight. Over the past several decades, adipose tissue has been recognized as a dynamic, multifunctional organ with a number of different types of cells (14,15). It houses the majority of stored energy as triglyceride, which is thought to be the primary targeted parameter for regulation in long-term energy homeostasis. The adipocyte serves its primary purpose of long-term storage of energy and as weight is gained, lost and regained, adipocytes and their support cells must undergo a substantial amount of remodelling to accommodate the gain or loss of stored energy (16). As an integrated node in the feedback system, adipose tissues must send and receive important signals to and from the brain and other peripheral tissues to appropriately adjust the level of stored energy.

Changes in adipocyte cellularity

Adipocyte size: highly modified
Weight loss is accompanied by a dramatic reduction in the size of adipocytes (Fig. 2), which is reversed when weight is regained (11,13,17–19). An individual adipose depot contains adipocytes that vary with respect to their size, and a size frequency distribution provides a clear picture of this variability within a depot. Because adipose depots exhibit differing cellularity profiles, a frequency distribution is often more informative than an average diameter. Studies in both humans and rodents suggest that adipocyte size is the most changeable aspect of cellularity characteristics in studies of weight loss and regain. During weight loss, energy stores are mobilized from adipocytes and adipocytes become smaller. During weight gain and weight regain, energy is accumulated and adipocytes become larger. The broad range for adipocyte size provides enormous flexibility for the amount of energy that can be stored at any one time. However, as adipocytes change size with the mobilization or accumulation of energy, the extracellular matrix must be remodelled to accommodate the change or a considerable mechanical strain will be imposed upon the adipocytes (16). Mariman has hypothesized that weight loss causes cellular stress in adipocytes, resulting in an altered metabolic profile that would relieve the stress via increased storage of lipid (8). From this perspective, one portion of the biological drive to regain weight could be based in the mechanical and molecular changes that are working to relieve the cellular stress and mechanical strain of the adipocyte.

Adipocyte number: modified unidirectionally
Weight loss does not lead to any discernible change in the number of adipocytes in adipose tissue (11,13,17–19) (Fig. 2). The number of adipocytes in a normal, healthy individual remains relatively constant throughout adulthood (20), but there are conditions in which the number of

![Figure 2](image_url)
adipocytes in particular adipose depots may increase. Our studies in a rodent paradigm of weight loss and regain suggest that the metabolic conditions during the relapse to obesity may provide the conditions that promote hyperplasia. Early in the relapse process, we observed the emergence of a population of very small (<20 μm) adipocytes, which was accompanied by an increase in total number of adipocytes in the depot (9). This increase in cell number persisted throughout the relapse process as all of the adipocytes became larger. We have speculated that this increased cell number partially explains animals in this model surpassing their pre-weight loss weight following relapse (11,13). While substantiating the temporal changes in cell size frequency distribution and total cell number in humans presents a logistical challenge, a hypercellularity phenomenon with similar characteristics has been reported in post-obese humans (21). Even so, this relapse-induced hyperplasia of adipose tissue, if it does occur, is likely limited to individuals who have a genetic predisposition for obesity. We have yet to observe an increase in cell number in diet-resistant rats or in DIO mice, which tend to relapse to their previous weight. Regardless, increasing the number of adipocytes in a depot in effect increases the overall capacity of that depot for triglyceride storage, and what flexibility exists for changing cell number appears to be unidirectional. There is very little evidence that the number of adipocytes is ever reduced under normal metabolic conditions associated with changes in weight. The implication is that increasing the number of adipocytes in a depot represents a permanent increase in the overall capacity of that depot to store triglyceride.

**Adipocyte turnover: a tightly controlled balance**

Because the number of adipocytes was observed to be relatively stable in normal, healthy adults, it was long thought that the adipocytes produced by puberty represented the population of cells that persisted throughout life. Tracer studies have discounted this notion by revealing that new adipocytes are being produced and mature adipocytes are being cleared with some regularity (22,23). A wide demographic study of Swedish adults observed that the turnover rate for adipocytes is approximately 8–10% per year. The generation of new adipocytes involves two distinct steps: (i) the proliferation of preadipocytes and (ii) the differentiation of preadipocytes into functioning adipocytes, capable of storing and releasing energy. The clearance of mature adipocytes is less understood, but is known to involve the recruitment of macrophages. The crown-like structures that are observed in adipose tissues represent adipocytes targeted for clearance, surrounded by the recruited macrophages (24). While the regulatory mechanisms for the generation and clearance of adipocytes are very different, they must be tightly linked to some global regulatory system that keeps them balanced, otherwise adipocyte number would be much less stable. The development of obesity is accompanied by a higher absolute amount of turnover, which is reflected in their greater fat mass and higher number of total adipocytes in their depots (16). The generation of new cells and clearance of mature cells remains, in general, balanced at a higher level in the obese. When adjusted for the difference in fat mass, the actual rate of cell turnover per unit fat mass is similar. At present, we do not know how adipocyte turnover is affected with weight loss or during the process of weight regain. However, if hyperplasia does occur, there must be some transient imbalance between new cell generation and mature cell clearance to account for the difference in cell number. Our ongoing studies will likely clarify how and when this balance is altered to elicit the hyperplasia we observed in our rodent paradigm of weight regain.

**Metabolic capacity of the adipocyte**

**Changes in global gene expression**

Adipose tissues experience a global down-regulation of gene expression in obese subjects in response to energy-restricted weight loss (25), which includes all of the key metabolic pathways. However, this effect is partly reversed at the transition to weight maintenance. With weight maintenance and during weight regain, an expression profile that would enhance energy conservation and the repletion of energy stores emerges (25–31). Markers of oxidative stress and inflammatory cytokines, which are also known to suppress appetite and increase expenditure, decline (29,32,33). The impaired induction of lipogenesis by insulin, glucose and feeding associated with obesity (34–37) resolves after energy-restricted weight loss (9,29,38–40). Finally, the enhanced metabolic response to ingested energy enhances nutrient clearance during weight maintenance and during sustained periods of overfeeding. These adaptive responses in the adipocyte prime the tissue to replete energy stores when nutrients once again become readily available.

**Metabolic changes linked to adipocyte size**

Insulin sensitivity is inversely related to size of the adipocyte (41). Compared with large adipocytes, small adipocytes exhibit higher rates of insulin-stimulated glucose uptake, higher levels of glucose oxidation and a lower sensitivity to antilipolytic action of insulin (42–44). In addition, smaller adipocytes exhibit a lower basal and catecholamine-induced lipolysis, have a lower rate of turnover of stored lipid and express genes favouring energy storage (28,45,46). The higher lipolytic capacity and triglyceride turnover in larger adipocytes is associated higher levels of Adipocyte triglyceride lipase (ATGL), Hormone sensitive lipase (HSL) and Lipoprotein lipase (LPL) (47–50). De novo lipogenesis is also down-regulated as
adipocytes increase in size (51–54). Varlamov et al. (55) suggested that this relationship between cell size and metabolic function serves to protect against lipid overload and continual expansion, which could eventually have deleterious consequences for the health of the cell. It was suggested that when the adipocyte size approaches a critical threshold in an individual (~100 μm), the capacity to take up and store circulating nutrients becomes diminished. If such a threshold exists, the implication is that an adipose tissue depot has a limited capacity to store energy, based upon the number of adipocytes it contains. Once that capacity is reached, the generation of new adipocytes (increasing the total capacity for storage) is the only avenue for storing more energy in the depot.

**Functional changes of the adipocyte?**

Beyond the cellularity characteristics, there is growing evidence to suggest that adipocytes have the capacity to alter their metabolic profiles and engage in wholesale changes in function, given the right metabolic context (14). White adipocytes have been observed *in vivo* to undergo transdifferentiation into brown adipocytes, which serve to dissipate, rather than store energy (56–58). Likewise, white adipocytes in the mammary gland have even been reported to transdifferentiate into glandular milk-producing epithelial cells during lactation, an effect that reverses after involution (57). These observations provide a novel perspective of the versatility of adipocytes that was once unappreciated. At present, few studies have considered such dramatic functional transformations in the context of weight loss, weight maintenance and weight regain studies. Given the metabolic extremes that can occur with weight loss and weight regain, it would be prudent for future studies to consider the extent to which adipocytes might be altered with energy restriction and gross overfeeding.

The versatility of metabolic profiles of adipocytes in changing environments may partly depend on the origins of the adipocytes. New adipocytes may primarily arise from resident preadipocytes and progenitors of the mesenchymal lineage, but recent findings demonstrate that bone marrow-derived progenitors (BMP) of the hematopoietic lineage can also migrate out of the skeleton and differentiate into adipocytes (59–62). Although this phenomenon needs to be demonstrated in humans, they may have an important role during weight regain if hyperplasia occurs. For instance, the observations of preferential homing and differentiation in visceral depots and lower leptin expression than white adipocytes suggest than BMP adipocytes could be a detriment to energy balance and metabolic health (60). The behaviour of these adipocytes during and after weight loss has not been determined, but would be essential for hypothesizing their relative role in energy balance and weight regain.

**Neuroendocrine signals affecting adipose tissue**

Energy-restricted weight loss from obesity is accompanied by a reduced sympathetic (SNS) tone (63–68) and reduced thyroid hormone levels (63,69–71). In contrast to the effects on SNS, the effect on thyroid hormones is observed less consistently and/or is more transiently tied to the early stages of weight loss (69,72,73). Collectively, these neuroendocrine changes can act upon adipose tissues to affect the size and number of resident adipocytes (Fig. 2). The SNS has established effects on the metabolic state and cellularity of adipose tissues (74,75) and a decline in SNS tone in this tissue could explain the shift in metabolic state favouring the uptake and deposition ingested energy, as well as the hyperplasia. Other studies indicate that both preadipocytes and adipocytes are responsive to Thyroid Stimulating Hormone (TSH) and thyroid hormones in a similar fashion (76–80). Both the SNS and thyroid hormones have inhibitory effects on preadipocyte proliferation and stimulatory effects on preadipocyte differentiation. As such, a decline in SNS tone and thyroid axis activity during weight maintenance may provide permissive conditions for preadipocyte proliferation, while the reversal of these neuroendocrine inputs during weight regain could underlie the hyperplasia. While these neuroendocrine inputs provide a plausible explanation for both metabolic and cellularity adaptations with weight loss and regain, their actual contribution to the adaptive response in adipose tissues requires further study.

**Adipose signals for long-term energy stores**

Leptin and insulin are often referred to as ‘adiposity signals’ because their levels generally reflect fat mass. Fasting levels of both hormones decrease with the decline in adiposity that occurs with weight loss (Fig. 2). The decline in leptin is more intuitive because it is secreted directly from adipocytes. The impact on insulin is indirect, reflecting the improvement in insulin sensitivity that occurs with weight loss (81–84). Interestingly, a number of studies have observed that leptin and insulin are actually reduced to a greater extent than would be expected for the amount of fat mass (11,21,65,85,86). We speculate that this may occur because leptin, and perhaps insulin, levels reflect both the amount of stored lipid and the size of the constituent adipocytes (87). Smaller adipocytes secrete less leptin and result in lower circulating levels for a given fat mass. Smaller adipocytes are also more insulin sensitive (45,46), which presumably means they require lower circulating levels of insulin to impart the same metabolic control. The reduction in cell size and the loss of total fat mass, therefore, may contribute independently to the decline in leptin and insulin. If new, very small adipocytes are generated early in the relapse process, the impact of cell size could be
compounded. Regardless, the integrated adiposity signal conveyed to the brain is that the total energy reserves are low and that the adipocytes are far below their maximal capacity to store energy. The changes in these hormones directly contribute to enhanced hypothalamic expression of arcuate nucleus (ARC) neuropeptide Y (88–92) and agouti-related peptide (91,92), as well as decreased expression of proopiomelanocortin (89) (Fig. 1). These changes are the hypothalamic hallmark of an ‘anabolic’ state, leading to a positive energy imbalance and weight gain. Although the concept that the adiposity signals reflect both total stores and the fraction of maximal capacity filled is consistent with observations in weight loss studies, it needs to be tested more rigorously.

What complicates the role of leptin and insulin as ‘adipose signals’ is that their relationship to adiposity is maintained only during energy balance and the correlations only apply to fasted levels of the hormones. When an energy imbalance occurs, leptin and insulin reflect the metabolic state (anabolic or catabolic) of adipose tissue, as it deposes or mobilizes energy. Overfeeding increases circulating levels of leptin and insulin (93) and, with persistent overfeeding during weight regain, both leptin and insulin resolve long before the weight is fully regained (7). For this reason, leptin and insulin, by themselves, do not appear to sustain the signal of energy depletion as weight is being regained.

**Nutrient availability as a reflection of the capacity to store excess energy**

To complement the signal of energy depletion from these hormones, we have proposed that signals reflecting nutrient availability play a more critical role during the dynamic phases of weight regain (7). Signals could be either the nutrients or their surrogate neuroendocrine signals. The improvement in systemic metabolic regulation is often accompanied by lower fasting levels of glucose, free fatty acids (FFAs) and triglycerides (TGs), and more consistently yields reduced postprandial excursions of glucose and TGs with potentiated postprandial reductions in FFAs (7). This wholesale, consistent change in circulating nutrients undoubtedly imparts some homeostatic influence on the signals of nutrient status (Fig. 1). Levels of glucose are detected by nutrient-sensing systems in both the periphery (94–98) and brain (96,99), with consequences to energy balance and fuel utilization in the periphery. Triglycerides may even be sensed via their putative effects on leptin and insulin transport across the blood–brain barrier (9,100,101). FFAs are sensed, such as glucose, in the central and peripheral nutrient-sensing systems and can reduce subsequent food intake when infused into the gut (102,103), into the circulation (104) or directly into the brain (105,106). The cellular and metabolic adaptations in adipose tissues certainly contribute to the attenuated postprandial excursions of circulating nutrients following weight loss. The consequence to systemic metabolism is that postprandial glucose excursions would be attenuated and the postprandial suppression of circulating FFAs would be potentiated.

**The ‘nutrient clearance’ hypothesis for the dynamic phase of weight regain**

This hypothesis suggests that the energy gap between appetite and expended energy persists during weight regain as a function of the capacity of adipose tissue to clear and store excess energy (7) (Fig. 2). Early in relapse, the adipose tissue’s capacity to clear excess energy is pitted against the rate at which nutrients are ingested and absorbed. As weight regain progresses, the adipocytes gradually increase in size and their capacity to clear excess energy diminishes. Excursions of glucose and TGs become larger and the suppression of FFA under dynamic (postprandial) states of metabolism would gradually become attenuated. Once the adipocytes near a critical threshold of size and the maximal capacity for stored energy is approached, the rate of weight regain would diminish. As the pre-weight loss weight is once again achieved, or surpassed if adipocyte hyperplasia has occurred, the fasting and postprandial levels of circulating nutrients would once again reflect the high levels observed with the insulin resistant state.

This simplistic hypothesis integrates the long-term adipose signals, reflecting the level of ‘stored energy’, with short-term signals of nutrient availability, which essentially reflect the ‘capacity to store energy’. Both signals are fundamentally rooted in the cellular and metabolic profiles of adipose tissues. Conceptually, the long-term signals provided by leptin and insulin would establish the global ‘anabolic’ tone in the hypothalamus, hindbrain and peripheral tissues. In this anabolic context, circulating nutrients and their surrogate neuroendocrine signals would become more important under postprandial conditions and during extended bouts of overfeeding while the weight is being regained. The convergence of these long-term and short-term signals in the energy homeostatic circuits of the brain would then dictate the magnitude and persistence of the energy gap.

The fundamental ideas behind this hypothesis are not entirely novel and they certainly present a simplified picture of the feedback system. Decades of research and numerous publications have provided a basic understanding of the key nodes of the homeostatic system controlling body weight and of adipocyte biology. Practically, the picture becomes much more complex as the integrated feedback signal from adipose tissues includes feedback from multiple adipose depots that have different metabolic and cellularity characteristics (15). Dieting and weight regain tend to alter
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visceral adipose depots more than subcutaneous depots (107–109), but less is understood about the interplay between depots, how they collectively establish a capacity-related ‘threshold’ for adipocyte size and about their relative contribution to the signals of energy depletion and nutrient availability during weight maintenance and weight regain. Furthermore, there is a large variability between individuals with respect to the metabolic and cellular characteristics of their adipose depots (22). This variation may translate into different ‘thresholds’ for adipocyte size and, consequently, different maximal capacities for a given adipocyte number. Even so, the value of this hypothesis is that it provides a basic explanation for the persistence of the energy gap driving weight regain in both static (during weight maintenance) and dynamic (during weight regain) phases of the relapse to obesity. In addition, it frames the integration of long-term signals for stored energy and short-term signals of nutrient availability in a manner that links both to the cellular and metabolic characteristic of adipose tissues.

Conclusions

Adipose tissues represent a key node in the homeostatic system that regulates body weight. Weight loss from caloric restriction results in substantial changes that prime adipose tissues to take up and store ingested energy. In combination with the other adaptations in this homeostatic system, these changes in adipose tissues present a significant challenge for successful weight loss maintenance. Weight loss awakens the body’s defence system in a manner that is persistent, saturated with redundancies and well-focused on the objective of restoring the body’s depleted energy reserves. Successful, long-term weight loss requires recognition of the strength and persistence of these biological pressures and a better understanding of how they may be countered with environmental, behavioural and pharmaceutical interventions. Adipose tissues and, more specifically, the adipocytes may provide an important target for developing interventions, given their critical role in the adaptive response. To be effective, interventions aimed at preventing weight regain will likely need to be as comprehensive, persistent and redundant as the biological adaptations they are attempting to counter.

Conflict of interest statement

The authors declare no conflict of interest.

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