Progress in the Molecular Understanding of Central Regulation of Body Weight by Estrogens

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Objective: Estrogens can act in the brain to prevent body weight gain. Tremendous research efforts have been focused on estrogen physiology in the brain in the context of body weight control; estrogen receptors and the related signals have been attractive targets for development of new obesity therapies. The objective is to review recent findings on these aspects.

Methods: Recent studies that used conventional and conditional knockout mouse strains to delineate the cellular and molecular mechanisms for the beneficial effects of estrogens on body weight balance are reviewed. Emerging genetic tools that could further benefit the field of estrogen research and a newly developed estrogen-based regimen that produces body weight-lowering benefits also are discussed.

Results: The body weight-lowering effects of estrogens are mediated by multiple forms of estrogen receptors in different brain regions through distinct but coordinated mechanisms. Both rapid signals and “classic” nuclear receptor actions of estrogen receptors appear to contribute to estrogenic regulation of body weight.

Conclusions: Estrogen receptors and associated signal networks are potential targets for obesity treatment, and further investigations are warranted.

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Introduction

A dramatic decline in circulating 17β-estradiol (E2) in postmenopausal women has been associated with development of obesity, type II diabetes, and the metabolic syndrome (1). While supplementation of E2 may ameliorate these risks, the application of estrogen replacement therapy in postmenopausal women has been very controversial. Since E2 can act upon several forms of estrogen receptors (ERs), and these ERs are coupled with complex intracellular signals, the body weight-lowering benefits produced by E2 are often associated with increased risks of reproductive endocrine toxicity and breast cancer (2). Obviously, one solution to this dilemma would be to target selective ER populations and/or ER-coupled signals that produce body weight benefits without unwanted side effects. Therefore, tremendous efforts have been focused on identifying the critical ER isoforms, the specific action sites of ERs, and the ER-coupled intracellular signals that are required for estrogenic actions on body weight control. It needs to be noted that actions of E2 both in the peripheral tissues and in the brain are important for the regulation of energy homeostasis, as demonstrated by studies using systemically or centrally administrated E2 (3). Since the peripheral actions of E2 have been extensively reviewed elsewhere (4), this review focuses on the central actions of E2 in the control of body weight.

ERα in the Brain Regulates Multiple Aspects of Energy Homeostasis

It has been well established that estrogens play an essential role in preventing body weight gain. For example, the withdrawal of endogenous estrogens by ovariectomy (OVX) in female animals leads to body weight gain and hyperadiposity, and this obese phenotype can be prevented by E2 replacement (5-10). The estrogenic effects on body weight homeostasis are believed to be primarily mediated by estrogen receptor-α (ERα), one of the “classical” estrogen receptors. Humans or mice with mutations in the ERα (Esr1) gene are obese (11,12). Further, deletion of ERα in mice blocks the antiobesity effects of E2 replacement (7). Early studies showed that microinjections of E2 into various brain regions change animal’s feeding behavior and body weight (13,14), suggesting that ERα expressed in the brain is important for the regulation of body weight balance. This notion was further supported by recent observations from various genetic mouse models. For example, Xu and Clegg crossed mice carrying loxP-flanked ERα alleles (ERαlox/lox) (15) to the Nestin-Cre transgenic mice (16) to produce mice lacking ERα only in the brain (17). Female mutant mice develop obesity, characteristic of increased body weight and body fat. Obesity in these mice is associated with hyperphagia, decreased energy expenditure, and decreased physical activity, which may all contribute to the
ERx is abundantly expressed in multiple brain regions that are implicated in the regulation of body weight balance. These include the ventrolateral portion of the ventromedial hypothalamus (VMH), the arcuate nucleus (ARC), the medial preoptic area (MPOA), and the nucleus of solitary tract (NTS) (18). Thus, an important question is which ERx population(s) in the brain are critical for the regulation of energy homeostasis. To this end, several groups have used genetic approaches to dissect out the physiological roles of ERx in various brain regions in the context of body weight control.

ERx in the VMH

As previously mentioned, abundant ERx is concentrated in the ventrolateral subdivision of the VMH (18). The VMH (also known as the VMN) is an important component of the neural circuits responsible for the homeostatic regulation of body weight (19). Accumulating evidence indicates a significant role of ERx in the VMH in mediating estrogenic actions on body weight balance. For example, Musatov et al. used shRNA-mediated gene silencing approach to knock down ERx in the VMH, while ERx expression in the adjacent ARC and other hypothalamic regions is shown to be unaffected (20). Animals with VMH-specific ERx knock-down are less sensitive to E2-induced weight loss and develop obesity characteristic of increased visceral fat (20). The obese syndrome is likely caused by decreased physical activity and impaired thermogenesis, whereas food intake of these animals are not directly affected (20).

In parallel, Xu and Clegg crossed ERxlox/lox mice and SF1-Cre mice, a VMH-specific Cre mouse line (21), to generate mice lacking ERx only in SF1 neurons. Notably, since SF1 only colocalizes with 50% of ERx-positive neurons in the VMH, these crosses achieved deletion of 50% ERx in the VMH (17). Nevertheless, female mutant mice show modest body weight gain and significant increases in body fat with a preferential increase in the visceral fat (20). Interestingly, these obese phenotypes are associated with normal food intake but profound decreases in brown adipose tissue (BAT)-mediated thermogenesis.

These observations in different models highlighted a significant role of VMH ERx signaling in regulating thermogenesis. This notion is further supported by Martinez de Morentin’s recent findings that injections of E2 into the VMH promote BAT-mediated thermogenesis in a feeding-independent manner (22). These authors further demonstrated that effects of E2 in the VMH on thermogenesis are mediated through inhibition of the AMP-activated protein kinase (AMPK) pathway (22).

Notably, Correa et al. recently developed a mouse model with NKX2-1, a transcription factor, deleted in VMH SF1 neurons (23). Deletion of NKX2-1 in SF1 neurons results in loss of 26% ERx-positive neurons in the VMH (23). Interestingly, female mice carrying this mutation develop profound obesity, associated with decreases in physical activity but normal BAT-mediated thermogenesis and food intake (23). Further, the authors used the Designer Receptors Exclusively Activated by Designer Drugs (DREADD) approach to show that stimulation of VMH neurons promotes physical activity in mice (23). Together, these data support a possibility that at least a subset of VMH ERx neurons function to stimulate physical activity, and therefore to prevent body weight gain.

Thus, multiple studies have demonstrated a critical role of VMH ERx in preventing body weight gain in females. It is clear that actions of VMH ERx do not regulate food intake, but stimulate energy expenditure. It appears that different subsets of VMH ERx neurons regulate different components of energy expenditure. Thus, some VMH ERx neurons primarily stimulate BAT-mediated thermogenesis to burn excess energy, whereas at least a subset of VMH ERx neurons promotes physical activity to dissipate energy.

ERx in the ARC

Abundant ERx is also expressed by neurons in the ARC (18). The ARC contains 2 distinct neural populations. These are neurons expressing pro-opiomelanocortin (POMC) and those expressing neuropeptide Y (NPY) and agouti-related peptide (AgRP). While POMC neurons synthesize and secrete an anorexigenic peptide, α-melanocyte-stimulating hormone (α-MSH), to activate melanocortin receptors, NPY/AgRP neurons release orexigenic peptides, NPY, and AgRP (24-26). Notably, AgRP is the endogenous antagonist of the melanocortin receptors (24-26). POMC and NPY/AgRP populations are believed to be the primary central regulators of energy homeostasis (27,28).

Olofsson et al. demonstrated that estrous-dependent fluctuations in circulating E2 in female mice are negatively correlated with expression of NPY and AgRP in the hypothalamus and the amount of daily food intake (29). These authors further showed that central administration of E2 inhibits NPY/AgRP neurons and suppresses food intake (29). Importantly, the E2-induced anorexia in female mice is blunted when NPY/AgRP neurons are selectively ablated (29). This study indicates that NPY/AgRP neurons are functionally required for the inhibitory effects of E2 on food intake. However, these authors also found that NPY/AgRP neurons express none to minimal levels of ERx (29). Thus, E2 may regulate these NPY/AgRP neurons indirectly via presynaptic neurons that express ERx; alternatively, E2 may directly regulate NPY/AgRP neurons through other ERs.

Notably, about 20–30% POMC neurons in the ARC co-express ERx (17,30,31). Using electron microscopy, Gao et al. reported that E2 can increase excitatory synaptic inputs onto ARC POMC neurons, which is associated with increased miniature excitatory postsynaptic current (32). Similarly, Malaya et al. reported that E2 stimulates POMC neurons by rapidly uncoupling GABAergic receptors from the G-protein-gated inwardly rectifying K+ channels (33). To further determine the physiological significance of ERx in POMC neurons, Xu and Clegg crossed ERxlox/lox mice and POMC-Cre mice (34) to generate mice lacking ERx only in POMC neurons (17). Female mice lacking ERx only in POMC neurons develop hyperphagia and modest body weight gain (17). Together, these observations indicate that ERx in POMC neurons is physiologically relevant in the regulation of food intake (17).

ERx in the DRN

ERx is abundantly expressed in the dorsal raphe nuclei (DRN) (18). Cao et al. further demonstrated that the majority of these ERx-
positive neurons in the DRN are serotonin (5-HT) neurons (35). Consistent with earlier results that E2 increases neural activities (demonstrated by c-fos immunoreactivity) in the DRN (36,37), Cao et al. showed that propyprazol triol (PPT, a selective ERz agonist) activates identified DRN 5-HT neurons via an ERz-dependent mechanism (35). Interestingly, Santollo et al. reported that microinjections of E2 into the DRN decreases food intake in rats (38). To further examine the roles of ERz in DRN 5-HT neurons, Cao et al. crossed ERzlox/lox mice and TPH2-CreER to generate mice lacking ERz only in 5-HT neurons (35). Interestingly, while these mutant mice show comparable basal food intake and body weight, they are resistant to estrogenic effects to suppress binge-like eating (35). These results suggest that ERz expressed by DRN 5-HT neurons primarily functions to suppress binge-like eating, while its roles in the basal feeding behavior may be minor.

**ERz in the NTS**

ERz is also present in the brain stem, including the NTS (18). Geary et al. showed that E2 replacement in wild-type mice suppresses food intake and potentiates CCK-induced satiation, which are accompanied by increased activity in NTS neurons (7,39). Interestingly, these responses are all abolished in mice lacking ERz (7,39). Further, it is shown that direct administration of E2 in the NTS potentiates CCK-induced satiety signals (40). Collectively, these findings support the notion that ERz in the brain stem, such as in the NTS, may be another physiologically important site to mediate the E2-induced anorexia.

Certainly, the physiological functions of ERz in other brain regions have not been fully revealed. For example, ERz is abundantly expressed in the amygdala (41). Earlier studies showed that injections of E2 into the amygdala decrease body weight in rats, effects that remain in rats with large hypothalamic lesions (42), suggesting a potential role of amygdala ERz in body weight control. In addition, Santollo et al. reported that microinjections of E2 into the MPOA decreases food intake in rats (38). Further, accumulating evidence indicates that E2 regulates food-associated reward (43), suggesting a role of ERz (or other ERs) expressed by brain reward centers (e.g., the nucleus accumbens and the lateral hypothalamus) (18). The functions of these ERz populations (among others) warrant further validation with genetic models. Despite the incomplete genetic mapping for ERz functions in brain regions, an interesting segregation model already started to emerge (Figure 1). Thus, ERz in the VMH enhances energy expenditure by stimulating BAT-mediated thermogenesis and/or physical activity; ERz in the ARC, DRN, NTS, and perhaps other regions, prevents body weight gain primarily by suppressing energy intake. These segregated ERz populations may function complementarily to mediate the full spectrum of estrogenic effects on female energy homeostasis. Supporting this possibility, Xu and Clegg have shown that female mice lacking ERz in both VMH SF1 neurons and in ARC POMC neurons develop hyperphagia and decreased thermogenesis which result in more robust obesity compared to modest obesity seen in mice with ERz deletion only in POMC neurons or in SF1 neurons (17).

**ERz in male brains**

It is clear that actions of ERz also prevent obesity in males. For example, ERz gene deficiency results in obesity in male mice (12,44) and in men (45,46). In addition, administration of E2 or its analogs reduces body weight in male mice (32,47). The major male sex hormone, testosterone, can be converted into E2 by aromatase, and both male and female aromatase knockout mice develop obesity (48). Notably, abundant aromatase is expressed by the brain (49), which makes it possible that ERz in male brains could be exposed to high levels of E2 despite the lack of circulating estrogens. Consistent with this notion, Xu and Clegg showed that male mice lacking ERz in the brain develop obesity (17), arguing that brain ERz also regulates male energy balance as it does in females. However, deletion of ERz in VMH neurons, POMC neurons, or DRN neurons, although produces feeding and/or body weight phenotypes in females, fails to affect male energy homeostasis. Thus, it is speculated that different brain ERz population(s) may be responsible for estrogenic actions on body weight balance in males, which remain to be identified.

**ERz-Coupled Intracellular Signals**

In addition to the sites of ERz actions, another major question in the field is what intracellular signals mediate ERz effects on body weight balance. ERz-coupled intracellular events can be divided into several modes. First, sub-sets of intracellular ERz are concentrated on the cytomembrane and in the cytosol, where it regulates rapid signaling pathways, including the PI3K/Akt pathway and the AMPK pathway (Figure 2A,B). Park et al. found that E2 stimulates the PI3K/Akt cascade in VMH neurons (50). Similarly, Malyala et al. reported that E2 activates ARC POMC neurons in an PI3K-dependent manner (33). As mentioned above, Martinez de Morentin et al. found that E2 inhibits the AMPK pathway in VMH neurons via an ERz-dependent mechanism and this inhibition mediates estrogenic actions to stimulate thermogenesis (22). Together, these observations support a model that ERz-initiated rapid signaling pathways, including PI3K and AMPK, mediate estrogenic actions to prevent body weight gain. However, it is worth noting that the roles of these...
As a nuclear receptor, ERα can also form a complex with other nuclear receptors or transcription factors to regulate gene transcription via directly binding to the EREs (51). Nevertheless, since the rapid signals initiated by cytosolic ERα are presumably also eliminated in these MOER mice, the possible contribution of the cytosolic ERα to energy homeostasis still remains unknown (Figure 2B).

As a classic nuclear receptor, ERα can also translocate to the nucleus to directly bind to the estrogen response elements (EREs) on the target genes and regulate gene transcription (Figure 2C). These ERE-dependent actions of ERα, however, do not appear to mediate the anti-obesity effects of E2. Park et al. generated a NERKI mouse model in which E207A/G208A mutations were introduced in the DNA binding domain of ERα, which abolish ERα-ERE binding (52). In these mice, metabolic phenotypes affected in ERα knockout mice including body weight, glucose homeostasis, energy expenditure, and physical activity, are restored to nearly normal levels (50), suggesting that the ERE-dependent ERα functions are not required to maintain body weight.

As a nuclear receptor, ERα can also form a complex with other nuclear receptors or transcription factors, which regulates gene transcription in an ERE-independent manner (Figure 2D). Little is known, however, about whether the ERE-independent ERα functions are involved in estrogenic effects on body weight control. Zhu et al. showed that hypothalamic ERα interacts with a nuclear receptor co-activator, namely steroid receptor co-activator-1 (SRC-1), and that deletion of SRC-1 blunts effects of E2 to reduce body weight and food intake, and to stimulate energy expenditure (53). These suggest that ERα’s nuclear receptor properties may still be required for estrogenic actions on energy homeostasis. Based on these observations and those from NERKI mice (50), it is speculated that the ERE-independent ERα functions may contribute to estrogenic effects on body weight control. Future studies, therefore, are warranted to identify the nuclear receptors and transcription factors that form a complex with ERα to mediate estrogenic actions on body weight balance.

### ERβ and Body Weight Balance

Compared to ERα, estrogen receptor-β (ERβ), another classic ER has received less attention at least in the context of body weight balance. An earlier study by Ohlsson et al. reported that chow-fed mice with global deficiency in ERβ show normal body weight and fat mass compared to wild-type mice (54). In addition, the authors reported that mice with compound knockout of both ERα and ERβ develop obesity with the same severity as mice only lacking ERα (54). Consistent with this, both Santollo et al. (55) and Roesch (9) found that an ERβ agonist, diaprylpropionitrile (DPN), has no effect on food intake and body weight in chow-fed O VX rats, while PPT (the ERα agonist) at similar doses can significantly reduce food intake and body weight. While these earlier studies suggest a minor role of ERβ in body weight control in chow-fed animals, Foryst-Ludwig et al. demonstrated that ERβ knockout mice, when fed a high-fat diet (HFD), developed obesity compared to HFD-fed wild-type mice (56). This increased sensitivity to diet-induced obesity is associated with normal food intake, but increased energy expenditure and decreased fat oxidation (56). Consistently, Yepuru et al. developed new selective ERβ agonists (β-LGNDSs), and found these agonists attenuate HFD-induced body weight gain associated with increased energy expenditure (57). Thus, the current data suggest that ERβ may play an important role in preventing obesity when animals are challenged by obesogenic diets, while ERα’s functions in animals fed on regular chow diets appear to be minimal. Certainly, the ERβ-mediated control of energy homeostasis warrants further investigation. For example, the action sites of ERβ on energy balance remain to be confirmed, although both Foryst-Ludwig et al. and Yepuru et al. suggested a contribution from ERβ in the peripheral tissues (56,57).

### GPR30 and Body Weight Balance

GPR30 (also known as GPER) is a G protein-coupled estrogen receptor, bound to the cell membrane. In vitro studies confirmed that E2 binds to GPR30. Body weight phenotypes among several independent GPR30 knockout mouse lines are controversial. For example, both Haas et al. (58) and Sharma et al. (59) observed obese phenotypes in male and female GPR30 knockout mice, which were generated by Wang et al. (60); however, Liu et al. reported no difference in body weight in the same GPR30 knockout strain (61). Otto et al. constructed an independent GPR30 knockout line, and
found no obese phenotypes in female mutants (62). Interestingly, another GPR30 knockout line generated by Martensson et al. showed reduced body weight only in males, but not in females (63). More recently, Davis et al. carefully characterized Wang’s GPR30 knockout mice and reported that both male and female mutants are significantly heavier than wild-type littermates, which appears to depend on reduced energy expenditure independent of physical activity, but not on food intake (64). Importantly, body weight-lowering effects of E2 are attenuated in OVX GPR30 knockout mice compared to OVX wild-type mice (64). The discrepancy from these studies may be attributed to different strategies to construct the GPR30 knockout alleles, different genetic background mice were maintained on, and/or different facility environment, etc. Nevertheless, observations from Wang’s GPR30 knockout line are expected that this ERα knockout mice, when combined with various Cre drivers that target distinct brain regions, has ultimately allowed the field to definitively examine the physiological roles of ERα in these various brain regions in body weight control (17,35). Of course, the ERα<sup>lox/lox</sup> mouse line has also been widely used to examine metabolic functions of ERα in peripheral tissues, including the adipose tissue (65) and the liver (66). Notably, an ERα<sup>lox/lox</sup> mouse line has been developed and validated by Antal et al. (67). With emerging evidence for a significant role of ERβ in preventing diet-induced obesity, it is expected that this ERα<sup>lox/lox</sup> mouse line will be heavily used in the field to dissect out critical ERβ populations for its beneficial effects against obesity.

Lee et al. recently developed and validated an Esr1<sup>Cre</sup> mouse line that expresses Cre recombinase in ERα-positive cells (68). The authors have combined this Cre mouse line with viral vectors that express optogenetic channels, e.g., channelrhodopsin and halorhodopsin, to achieve photostimulation or photoinhibition of ERα-positive neurons selectively in the VMH. In addition to the site specificity (only targeting VMH ERα neurons), more importantly, this strategy allows manipulations of neural activity with various scales of strength and with a high temporal resolution. Taking these advantages, the authors elegantly demonstrated that while weak activation of VMH ERα neurons instantly triggers sexual behavior, strong activation of the same neurons initiates attack (68). Certainly, this Esr1<sup>Cre</sup> mouse line notes a long-awaited tool for the field of estrogen research. With this tool, combined with the optogenetic and pharmacogenetic (DREADD) viruses, investigators can examine the roles of ERα neural activity in any given brain region in the regulation of food intake, energy expenditure, thermogenesis, and physical activity, etc. In addition, this Esr1<sup>Cre</sup> mouse can be crossed to various loxed mouse alleles to delete genes of interest only in ERα-positive cells. These would allow further dissection of intracellular

**Figure 3** Electrophysiological recordings in identified ERβ-positive 5-HT neurons in the DRN. (A) Brightfield, (B) fluorescence for GFP, (C) for TOMATO, and (D) merge from a recorded neuron in the DRN of the brain slice prepared from a TPH2-CreER/Rosa26-tdTOMATO/ERβ-eGFP mouse. (E) A representative trace showing that DPN treatment (300 nM, bath perfusion) decreased firing frequency of ERβ-positive 5-HT neurons in the DRN. (F) The trace of the only ERβ-positive 5-HT neuron that showed increased firing frequency upon DPN treatment (300 nM, bath perfusion). Summary data of the effects of DPN (300 nM) on (G) firing frequency and (H) resting membrane potential in 13 ERβ-positive 5-HT neurons in the DRN. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]
signals in ERα-positive cells that may mediate estrogenic functions, including body weight control.

Other genetic tools that may benefit the estrogen field include a transgenic ERα-eGFP mouse line developed by Matsuda et al. (69). The authors have confirmed that in these ERα-eGFP mice, an enhanced GFP protein is expressed largely in ERα-positive neurons (as indicated by endogenous ERα immunoreactivity), although a few GFP-labeled neurons are found to be ERα negative (69). Similarly, Milner et al. developed and validated a transgenic ERβ-eGFP mouse line, which expresses the enhanced GFP protein in ERβ-positive cells (70). Obviously, these eGFP models can replace the conventional staining procedures (e.g., immunohistochemistry or in situ hybridization) required to visualize ER-positive neurons; GFP can also be easily combined with another staining protocol to determine if ER-positive cells express other proteins or mRNAs. In addition to these histological applications in usually fixed tissues, investigators can also use GFP as a marker to directly identify ER-positive cells in un-fixed tissues. For example, flow cytometry can be used to sort out highly purified cells that express ERα or ERβ; the similar approach has been applied to purify NPY/AgRP neurons which facilitated the discovery of novel factors that regulate these neurons and food intake (71). In addition, investigators can prepare fresh brain slices from ERα-eGFP or ERβ-eGFP mice and perform electrophysiological recordings in identified ER-positive neurons. One step further is to cross these eGFP transgenics onto mice that express another fluoroscent reporter in a selective neural population. For example, we have crossed the ERβ-eGFP mice with TPH2-CreER/Rosa26-tdTOMATO mice (35) to generate mice carrying all 3 transgenic alleles. In the tri-genetic offspring, TPH2-CreER/Rosa26-tdTOMATO/ERβ-eGFP mice, tamoxifen injections (3 mg/injection, intraperitoneal, twice 24 hours apart) induced the strong red fluorescence (TOMAT0) exclusively in TPH2-positive neurons (5-HT neurons; enhanced GFP is expressed only in ERβ-positive neurons); thus, double-labeled neurons (yellow) are identified as ERβ-positive 5-HT neurons (Figure 3A–D). With this tool, we performed electrophysiological recordings in ERβ-positive 5-HT neurons in the DRN and examined effects of DPN (the ERβ agonist) on firing properties of these neurons. We found that DPN decreased the firing frequency in 12 out of 13 neurons we recorded (see typical trace in Figure 3E and summarized data in Figure 3G), while the other neuron showed increased firing frequency upon the DPN treatment (see the trace in Figure 3F and summarized data in Figure 3G); resting membrane potential of all neurons were not altered (Figure 3H). These findings are interesting, as the ERβ agonist inhibits most of ERβ-positive 5-HT neurons in the DRN, while the ERα agonist (PPT) has been shown to activate DRN 5-HT neurons (35). More importantly, the fact that all double-labeled neurons responded to the ERβ agonist provided the proof of the principle that the ERα-eGFP and ERβ-eGFP mouse models could be used in combination with other genetically labeled reporter lines to allow experiments in highly selective neural populations. These will no doubt advance our understanding about molecular and cellular actions of these ERs.

Therapeutic Potential

While the body weight-lowering effects of estrogens have been well established in animal models, the application of estrogen replacement therapy to treat/prevent obesity in humans (including post-menopausal women) has been hampered due to increased risks of reproductive endocrine toxicity and breast cancer associated with the conventional estrogen replacement therapy (2). One idea to overcome these issues is to develop estrogen analogs that may only target specific ER isoforms or specific sites of ER actions that produce body weight benefits. Indeed, Finan et al. recently developed a GLP-1-estrogen conjugate, which uses glucagon-like peptide-1 (GLP-1) as a “carrier” to deliver estrogens preferentially to GLP-1 receptor-enriched regions, including the hypothalamus (47). The authors demonstrated that systemic administration of this GLP-1-estrogen conjugate, in both male and female mice with diet-induced obesity, substantially reduces body weight and food intake, and improves the glucose tolerance and insulin sensitivity (47). Most importantly, the common side effects associated with estrogen replacement therapy (e.g., reproductive endocrine toxicity and breast cancer risks) are avoided in GLP-1-estrogen treated mice, presumably because estrogens are not delivered to the reproductive organs and the breast (47). Further mechanistic studies indicate that the body weight-lowering effects of the conjugate stem from both GLP-1 and estrogens; interestingly, both ERα and ERβ are required for estrogen-mediated benefits since genetic deletion of either of these ERs blunts effects of the conjugate (47). Subsequently, Cao et al. found that GLP-1-estrogen also delivers bioactive estrogens to the DRN and substantially suppresses binge-like eating in female mice partially through acting upon ERα expressed by DRN 5-HT neurons (35). The development of this GLP-1-estrogen conjugate is certainly an exciting step forward in the field, as this conjugate itself or its modified forms could potentially become a therapy for obesity and/or binge eating. Further, these studies proved the concept that estrogens could be conjugated with other peptides or molecules to achieve target-specific delivery and therefore avoid unwanted side effects while maintaining their beneficial effects.

Another strategy is to conjugate E2 with selective estrogen receptor modulators (SERMs) which function to antagonize E2 actions selectively in the breast and reproductive organs. Kim et al. recently tested such a conjugate, equine estrogen (CE) and bazedoxifene (BZA) (72). Oral administration of CE and BZA produces profound body weight loss through stimulating lipid oxidation and energy expenditure in HFD-fed OVX mice (72). Importantly, this regimen does not cause increases in uterine weight in mice (72), because BZA antagonizes estrogenic actions in the uterus (73,74). The newly developed ERβ agonists (β-LGNDs) may also carry therapeutic potentials. As mentioned above, Yepuru et al. demonstrated that administration of β-LGNDs in HFD-fed mice significantly prevents body weight gain and improves glucose tolerance (57). Unlike E2, these agonists do not stimulate proliferation of a human endometrial adenocarcinoma cell lines in vitro, and neither do these agonists increase uterine weight in female rats (57), suggesting minimal side effects at least in the reproductive organs. While further investigations are needed to test the potential toxicities in other tissues (e.g., the breast), these newly developed ERβ agonists represent an alternative strategy: to selectively target ERβ as potential therapies for obesity.

Conclusion

The body weight-lowering effects of E2 are likely mediated by multiple forms of ERs. In particular, ERα in different brain regions regulates distinct aspects of female energy balance, and these different
ERα populations may provide well-coordinated responses in food intake, thermogenesis, and physical activity to ultimately prevent body weight gain. While ERα in male brains is also essential for the maintenance of normal body weight, the exact sites of actions remain to be identified. With regard to ERα-coupled intracellular signals, both rapid signals (e.g., PI3K and AMPK) and “classic” nuclear receptor actions on gene expression appear to contribute to estrogenic regulation of body weight, although the picture of this complex signal network is still not clear. Effects of other ERs (ERβ and GRP30) on body weight balance may have been under-appreciated in the past; fortunately, revisits of various knockout models started to reveal previously unrecognized roles of these receptors, and these warrant further investigations. Another area that deserves more careful investigations in the future is the interactions of E2 and other sex hormones (e.g., progesterone) in the context of feeding and body weight control, as emerging evidence from human studies indicates that the interactions of E2 and progesterone, rather than either alone, influence feeding behaviors in cycling women (75).

It needs to be recognized that the applications of various genetic mouse models, e.g., conventional knockout strains, ERαlox/lox MOER, and NERKI, have substantially advanced our current understanding about where and how estrogens regulate body weight balance. Further, several additional relevant genetic mouse tools, including ERαlox/lox, Est1Cre, and GFP reporters, can be added into our toolbox. These tools are expected to facilitate our future research efforts to complete mapping critical ERα/ERβ sites for body weight controls, to identify the intracellular signals or target genes that mediate estrogenic actions on body weight, to reveal the physiological relevance of ERα neural activities in controlling feeding behavior and/or energy expenditure, and to gain molecular insights regarding how ERα/ERβ neurons may be regulated. Of course, this too will be extremely useful to validate the newly developed estrogen-based compounds as potential obesity therapies and to facilitate the identification of new targets and development of new compounds.

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