20 YEARS OF LEPTIN

Leptin and reproduction: past milestones, present undertakings, and future endeavors

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Abstract

The association between leptin and reproduction originated with the leptin-mediated correction of sterility in ob/ob mice and initiation of reproductive function in normal female mice. The uncovering of a central leptin pathway regulating food intake prompted the dissection of neuroendocrine mechanisms involving leptin in the metabolic control of reproduction. The absence of leptin receptors on GnRH neurons incited a search for intermediary neurons situated between leptin-responsive and GnRH neurons. This review addresses the most significant findings that have furthered our understanding of recent progress in this new field. The role of leptin in puberty was impacted by the discovery of neurons that co-express kisspeptin, neurokinin B, and dynorphin and these could act as leptin intermediates. Furthermore, the identification of first-order leptin-responsive neurons in the premammillary ventral nucleus and other brain regions opens new avenues to explore their relationship to GnRH neurons. Central to these advances is the unveiling that agouti-related protein/neuropeptide Y neurons project onto GnRH and kisspeptin neurons, allowing for a crosstalk between food intake and reproduction. Finally, while puberty is a state of leptin sensitivity, mid-gestation represents a state of leptin resistance aimed at building energy stores to sustain pregnancy and lactation. The mechanisms underlying leptin resistance in pregnancy have lagged; however, the establishment of this natural state is significant. Reproduction and energy balance are tightly controlled and backed up by redundant mechanisms that are critical for the survival of our species. It will be the goal of the following decade to shed new light on these complex and essential pathways.

Key Words

leptin

gonadotropin-releasing hormone

reproduction

hypothalamus

neurotransmitters

Introduction

The survival of any species depends on its ability to reproduce, otherwise it will become extinct. Reproduction is an energy-demanding process and its physiological costs vary widely in the animal kingdom, but there is no doubt that the burden of reproduction lies with females. The ability to reproduce starts at different ages in humans but invariably involves a period of sexual maturation that culminates with the first menstrual cycle. In most inbred strains of mice, the first ovulation occurs between 6 and 8 weeks of age; however, in humans, puberty is attained in females at ~12–13 years of age. While the onset of puberty hallmarks the reproductive lifespan of a female, pregnancy, parturition, and lactation are the crux of
reproductive biology and remain the most physiologically complex and energy-demanding life processes.

The adipose mass of an organism represents an energy storage reservoir. In organisms that undergo periods of hibernation or torpor, buildup of the adipose mass sustains the low metabolic rates and is essential for reproductive function, as best exemplified in ground squirrels (Forger et al. 1986). Except in marsupials, torpor in mammals is not a requirement for survival, but the ability to fatten up at critical periods is an essential component of normal physiology. In this review, we will address the leptin-mediated effects and associated mechanisms that pertain to the accumulation of adipose mass at critical times during the reproductive events of a female, namely at puberty and pregnancy.

The seminal discovery that leptin is secreted from adipocytes (Zhang et al. 1994) raised an interest in the mechanisms of food intake and adipose tissue accumulation and their impact on the obesity epidemic. However, the link between adipose tissue and reproduction predates obesity, which often confounds and aggravates studies of leptin and the reproductive axis. The inspirational papers of Gordon Kennedy pioneered the link between energy intake and reproduction by placing these two pathways within the hypothalamic network (Kennedy & Mitra 1963a,b,c). Later, Frisch & McArthur (1974) unveiled their critical weight hypothesis and demonstrated that perturbations in the adipose mass affect female fertility negatively (Vigersky et al. 1977, Frisch et al. 1980, 1981). While much of important physiology was unveiled in these early days, it is unfortunate that the groundbreaking parabiosis experiments of the late Doug Coleman (1973) between db/db and ob/ob mice did not involve investigation of the reproductive axis during the months’ long response of ob/ob mice to the db/db-originated leptin. Rose Frisch studies were critical in providing a basis for the role of leptin in reproduction and were analogous to Coleman’s groundbreaking parabiosis experiments that provided a framework for the roles of leptin and leptin receptor. It was not until leptin rescued the sterility of ob/ob mice (Chehab et al. 1996, Mounzih et al. 1997) and advanced the onset of puberty in normal mice (Ahima et al. 1997, Chehab et al. 1997, Yura et al. 2000) that Kennedy and Frisch’s works on the link between the adipose tissue and reproduction were revived and a tying metabolic factor inducing a state of leptin resistance during pregnancy, between these two pathways.

Overall, initial studies have emphasized two essential physiological functions of leptin pertaining to its role in reproduction, namely in puberty and pregnancy. Although leptin affects both processes, their mechanisms are distinct in that the former is a state of leptin sensitivity highlighted by a leptin surge in rodents and primates (Ahima et al. 1998, Suter et al. 2000), while the latter imparts on increasing food intake via a leptin-resistant state (Mounzih et al. 1998). Thus, this review will focus on the progress aimed mostly at increasing our understanding of a leptin-sensitive state in puberty and to a lesser extent a leptin-resistant state during pregnancy.

Leptin in puberty

Our knowledge of the mechanisms and cues that control the onset of puberty have considerably increased in the 20th century with much of the work originating from rodent and primate animal models. There is no doubt that the onset of puberty is gated by multiple metabolic factors, which altogether inform the reproductive system about the extent of energy stores in an organism in order to activate the GnRH pulse generator. In fact, increases in adipose tissue mass advance the onset of menarche (Wattigney et al. 1999, Kimm et al. 2001), whereas fat depletion delays it (Frisch et al. 1980). Paradoxically, leptin-deficient subjects are obese but, due to the lack of leptin, fail to advance to puberty unless treated with exogenous leptin (Montague et al. 1997, Farooqi et al. 1999). These studies highlight the critical role of leptin in the onset of human reproductive function. There is also little doubt that any serious puberty-triggering factor must be able to influence directly or indirectly the secretion or regulation of GnRH. A requirement for such a factor is that mutations in its gene, whether in human and/or mouse models, must result in infertility or significant
Leptin and reproduction

In a milieu of elevated leptin levels at puberty, it is conceivable that leptin and insulin together influence GnRH secretion. The critical role of IGF1 is also shown by the postnatal lethality of Igf1 null mice (Liu et al. 1993, Powell-Braxton et al. 1993). However, a mixed genetic background of Igf1 null mice rescues their lethal phenotype and these mice exhibit growth retardation and a failure to attain puberty (Liu & LeRoith 1999). In human, homozygous deletion of the IGF1 gene resulted, as in genetically heterogeneous mice, in postnatal survival and retarded growth but normal, if not, early puberty (Woods et al. 1996, 2000). Thus, unlike in NIRKO mice, IGF1 does not appear to be critical and necessary for the onset of puberty but remains an important factor as it correlates with growth. An elegant experiment (Divall et al. 2010) targeted the deletion of Ir or Igf1r on GnRH neurons and determined puberty and fertility outcomes in knockout mice. Mice with a targeted deletion of Ir on GnRH neurons displayed normal puberty and fertility. However, male and female mice with a deletion of the Igf1r on GnRH neurons showed delayed puberty but normal fertility. Interestingly, IGF1 administration advanced puberty in normal but not Igf1r knockout mice. Although Igf1r knockout mice showed delayed but not a block in puberty, IGF1R signaling on GnRH neurons appears to be an important contributor to the timing of puberty.

Orexigenic and anorexigenic factors

Orexigenic factors such as ghrelin, NPY, MCH, agouti-related protein (AgRP), and orexin constitute a food intake-regulatory circuit that could influence the hypothalamic–pituitary–gonadal (HPG) axis. Ghrelin, an endogenous ligand for the GH secretagogue receptor (Howard et al. 1996), is similar to leptin in that they are both secreted from the periphery, the former from stomach and the latter from adipose tissue, and exert their major effects on the arcuate nucleus (ARC; Cowley et al. 2001, Hewson et al. 2002), including the HPG axis (Steiner et al. 2003, Kluge et al. 2007). The fact that leptin and ghrelin are antagonistic in their effects on food intake, their respective anorexigenic and orexigenic roles make them ideal candidates to regulate the reproductive system in times of nutritional variation. Ghrelin exerts a negative effect on GnRH (Fernandez-Fernandez et al. 2005), whereas leptin stimulates it or facilitates its secretion (Nagatani et al. 1998, Reynoso et al. 2003). However, unlike leptin-deficient mice, ghrelin knockout mice are neither obese nor reproductively impaired (De Smet et al. 2006). Similarly, orexin knockout mice suffer no energy or

reproductive dysfunction. Leptin fulfills both of these requirements as shown by the infertility of leptin-deficient mice and humans and their leptin-mediated rescue (Chehab et al. 1996, Farooqi et al. 1999, Gibson et al. 2004) as well as its indirect stimulation of GnRH (Nagatani et al. 1998, Reynoso et al. 2003) and the potentiation of luteinizing hormone (LH) release from pituitary gonadotropes in the presence of GnRH (Avelino-Cruz et al. 2009).

Leptin also appears to exert GnRH-independent effects on the reproductive system as shown by its ability to induce ovulation in Gnrh (Gnrh1)-deficient mice (Barkan et al. 2003), an intriguing but interesting finding. Overall, leptin is a metabolic factor that bridges the regulation of the fat mass with reproduction.

The central question is the nature of the critical metabolic factor(s) that signals to GnRH neurons the appropriate physiological readiness to initiate GnRH release and activate the puberty cascade. Nutritional and metabolic factors have long been suspected to participate in the process and could elicit a direct or indirect release of GnRH. While mouse and human studies convincingly demonstrate an essential role of leptin in puberty (Chehab et al. 1996, 1997, Ahima et al. 1997, Cheung et al. 1997, Farooqi et al. 1999), the absence of signaling-competent leptin receptor expression on GnRH neurons strongly suggests that an intermediary factor or pathway mediates the essential effects of leptin on the activation of GnRH neurons. Potential intermediate candidates are insulin, insulin-like growth factor 1 (IGF1), growth hormone (GH), ghrelin, neuropeptide Y (NPY), orexin, melanin-concentrating hormone (MCH), adiponectin, kisspeptin, and possibly gut peptides. Which of these factors is the single, if any, stimulus and what are its backup or co-factors? From an evolutionary perspective, it makes sense that leptin is a metabolic factor that bridges the regulation of GnRH neurons. From an evolutionary perspective, it makes sense that metabolic factor(s) that signals to GnRH neurons the appropriate physiological readiness to initiate GnRH release and activate the puberty cascade. Nutritional and metabolic factors have long been suspected to participate in the process and could elicit a direct or indirect release of GnRH. While mouse and human studies convincingly demonstrate an essential role of leptin in puberty (Chehab et al. 1996, 1997, Ahima et al. 1997, Cheung et al. 1997, Farooqi et al. 1999), the absence of signaling-competent leptin receptor expression on GnRH neurons strongly suggests that an intermediary factor or pathway mediates the essential effects of leptin on the activation of GnRH neurons. Potential intermediate candidates are insulin, insulin-like growth factor 1 (IGF1), growth hormone (GH), ghrelin, neuropeptide Y (NPY), orexin, melanin-concentrating hormone (MCH), adiponectin, kisspeptin, and possibly gut peptides. Which of these factors is the single, if any, stimulus and what are its backup or co-factors? From an evolutionary perspective, it makes sense that leptin is a metabolic factor that bridges the regulation of the fat mass with reproduction.

Insulin/IGF1

The insulin/IGF1/GH axis is the hallmark of anabolic hormones that affect GnRH neurons at puberty. This is best demonstrated by the infertility of neuron-specific insulin receptor knockout female mice (NIRKO), which exhibit a 90% reduction in LH but remain responsive to a GnRH agonist (Bruning et al. 2000). Thus, neuronal expression of the insulin receptor appears to be critical for reproduction. In a milieu of elevated leptin levels at puberty, it is conceivable that leptin and insulin together influence GnRH secretion. The critical role of IGF1 is also shown by the postnatal lethality of Igf1 null mice (Liu et al. 1993, Powell-Braxton et al. 1993). However, a mixed genetic background of Igf1 null mice rescues their lethal phenotype and these mice exhibit growth retardation and a failure to attain puberty (Liu & LeRoith 1999). In human, homozygous deletion of the IGF1 gene resulted, as in genetically heterogeneous mice, in postnatal survival and retarded growth but normal, if not, early puberty (Woods et al. 1996, 2000). Thus, unlike in NIRKO mice, IGF1 does not appear to be critical and necessary for the onset of puberty but remains an important factor as it correlates with growth. An elegant experiment (Divall et al. 2010) targeted the deletion of Ir or Igf1r on GnRH neurons and determined puberty and fertility outcomes in knockout mice. Mice with a targeted deletion of Ir on GnRH neurons displayed normal puberty and fertility. However, male and female mice with a deletion of the Igf1r on GnRH neurons showed delayed puberty but normal fertility. Interestingly, IGF1 administration advanced puberty in normal but not Igf1r knockout mice. Although Igf1r knockout mice showed delayed but not a block in puberty, IGF1R signaling on GnRH neurons appears to be an important contributor to the timing of puberty.

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reproductive peculiarities (Chemelli et al. 1999). Interestingly, knockout mice for NPY are fertile and lean (Erickson et al. 1996a). However, ob/ob mice deficient in NPY or in the NPY-Y4 receptor show an attenuation of obesity and improved fertility (Erickson et al. 1996b, Sainsbury et al. 2002), suggesting that a lack of NPY or its Y4 receptor alleviates the restraint placed by a deficiency of leptin on the reproductive axis and, most importantly, demonstrating a central role for NPY in reproduction. Although NPY has long been known to stimulate GnRH secretion (Khorram et al. 1988, Sutton et al. 1988), it was also found to have both inhibitory and stimulatory effects on GnRH, via the NPY-Y1 and NPY-Y4 receptors respectively (Roa & Herbison 2012). Thus, it would be revealing to determine the physiological states and mechanisms under which NPY exerts this variable effect on GnRH neurons. Another observation that links NPY to GnRH is their migration origin. Both GnRH and NPY neurons originate from the olfactory placode and migrate into the CNS during embryogenesis (Hilal et al. 1996), implying that their mutual relocation to the hypothalamus could be somewhat evolutionary and functionally linked.

AGRP is expressed in leptin-responsive neurons of the ARC and exerts significant effects on energy intake when overexpressed in transgenic mice (Ollmann et al. 1997), but has no obvious effects on adiposity or fertility in Agrp knockout and double Agrp/Npy knockout mice (Qian et al. 2002, Xu et al. 2005). However, ablation of AGRP-expressing neurons in leptin-deficient ob/ob and leptin receptor-deficient db/db mice, remarkably, restores fertility (Israel et al. 2012, Wu et al. 2012, Sheffer-Babila et al. 2013). Thus, the findings that NPY and AGRP play critical roles in reducing sterility in leptin- or leptin-signaling-deficient states (Erickson et al. 1996b, Israel et al. 2012, Wu et al. 2012) uncover an essential link between energy balance and reproductive pathways. While it is reasonable to assume that Agrp/Npy projections into GnRH neurons might serve as the long-sought leptin intermediate, a more plausible hypothesis is that they represent a secondary alternate reproductive pathway, namely because single or double Agrp/Npy knockout mice without leptin or leptin receptor mutations fail to exhibit any reproductive defect. However, Agrp stimulates GnRH release (Stanley et al. 1999) and thus this secondary pathway is fundamental and may underlie yet unexplained mechanisms of fertility in ob/ob mice such as when food restricted or bred on mixed genetic backgrounds (Lane & Dickie 1954, Ewart-Toland et al. 1999, Qiu et al. 2001). The facilitating and rescuing effects of this secondary pathway may also explain fertility in a female with a leptin receptor mutation (Nizard et al. 2012).

MCH is another orexigenic peptide expressed in the lateral hypothalamus, where leptin-responsive neurons are located. However, Mch knockout mice are lean, hypophagic, and remain fertile (Shimada et al. 1998), owing most probably to the presence of enough adipose tissue that secretes leptin. Furthermore, ob/ob mice lacking Mch remain infertile despite attenuation of their obesity caused by an increase in energy expenditure and not decreased hyperphagia (Segal-Lieberman et al. 2003).

Anorexigenic factors also affect the reproductive system. It is well known that perturbations in energy distribution and balance such as in lipodystrophy and anorexia nervosa cause interruptions of the menstrual cycles. Notable anorexigenic factors secreted by the gut in response to nutrient ingestion include protein tyrosine (PYY), pancreatic polypeptide, cholecystokinin (CCK), glucagon-like peptide 1 (GLP1), oxyntomodulin (OXM), and apolipoprotein A-IV (apoA-IV). Pyy knockout mice are obese and fertile (Batterham et al. 2006). Cck knockout mice are normal and fertile (Lacourse et al. 1999). Glp1r knockout mice show impairments in behavioral and stress responses and minor reproductive disturbances, such as reduced gonadal weights in males and slight puberty delay in females, but overall normal fertility (MacLusky et al. 2000). Similar to GLP1, OXM results from the proglucagon (Gcg) gene, acts as an agonist at GLP1R and stimulates weight loss (Kosinski et al. 2012), but has not yet been shown to exhibit any reproductive effect. Similarly, apoA-IV, which is secreted by the small intestine in response to fat absorption, was suggested as a satiety factor but apoA-IV (Apoa4) knockout mice showed only mild dyslipidemia and no reproductive defects (Weinstock et al. 1997).

POMC neurons that activate the anorexigenic arm of the central leptin pathway are direct leptin targets in the ARC. Mutations of the POMC gene in human and its ablation in mice result in obesity but no apparent effects on the reproductive axis (Krude et al. 1998, Yaswen et al. 1999). In addition, targeted deletion of the leptin receptor on POMC neurons resulted in no reproductive defects (Balthasar et al. 2004, van de Wall et al. 2008). However, deletions of both leptin and insulin receptors from POMC neurons caused insulin resistance and reduced fertility, a condition that resembled characteristics of the polycystic ovary syndrome (Hill et al. 2010). Thus, POMC neurons also mediate normal function of the reproductive axis in females. α-MSH is the POMC-derived ligand that binds to MC4R located on second-order leptin-responsive hypothalamic neurons, which stimulate the anorexigenic arm of the pathway. Mc4r knockout mice are obese and subfertile (Huszar et al. 1997) and, interestingly, their...
reproductive dysfunction can be rescued with increasing exercise (Irani et al. 2005). This observation is consistent with the ability of a melanocortin receptor antagonist to reverse reduced food intake in leptin-treated ob/ob mice, without, however, affecting the activation of their reproductive axis, suggesting a dissociation of the melanocortin pathway from the reproductive system (Hohmann et al. 2000). These observations are also consistent with normal fertility of adult obese subjects carrying heterozygous and homozygous deleterious mutations in the MC4R gene (Faroogi et al. 2003). Thus, while melanocortin neurons could play a secondary role in reproduction, they are unlikely to serve as the critical leptin intermediates.

Overall, it appears that while the orexigenic and anorexigenic factors listed above could play single handedly a critical role in fertility, it is more likely that altogether they would be the gatekeepers of reproduction. To gain further insight into their individual role, a more informative approach would be required to generate double and triple knockout alleles in a single mouse and then determine their combined effects on obesity and reproduction. While this strategy is quite laborious for traditional knockout strategies, the recent advent of genomic editing technologies such as the CRISPR/Cas9 system (Wang et al. 2013) makes it likely to generate such informative experimental model systems.

Kisspeptins

The exciting discovery of the kisspeptin system offered a great candidate for a leptin intermediate acting between leptin-responsive neurons and GnRH neurons. Kisspeptin, a product of the KISS1 gene, is expressed in various mammalian species, predominantly in a large population of neurons in the ARC and to a lesser extent in the preoptic area (POA), the rostral periventricular region of the third ventricle, as well as in scattered regions of the brain outside of the hypothalamus (Lehman et al. 2010). Kisspeptin neurons connect to NPY and POMC neurons (Backholer et al. 2010), which are first-order leptin-responsive neurons in the hypothalamus, enhancing the coordination of the nutritional leptin axis to its reproductive counterpart.

Kisspeptin emerged as a primary puberty candidate factor following the findings that subjects with hypogonadotropic hypogonadism carry mutations in the kisspeptin receptor GPR54 (KISS1R; de Roux et al. 2003, Seminara et al. 2003, Semple et al. 2005). The first indication that kisspeptin could indeed be the long-sought leptin intermediate was demonstrated in leptin-deficient ob/ob mice that express low levels of Kiss1 mRNA, that are then stimulated to be increased with leptin treatment (Smith et al. 2006). Targeted deletions in mice of Kiss1 or the Kiss1 receptor Gpr54 gene revealed infertility and abnormal sexual maturations (d’Anglemont de Tassigny et al. 2007, Lapatto et al. 2007), affirming a critical role for the kisspeptin system in the activation of the reproductive axis. Consistently, the finding that i.c.v. administration of kisspeptin stimulated maturation of the reproductive axis in hypoleptinemic states of prepubertal rats and hyperleptinemic fai/fia Zucker rats with a mutation in the leptin receptor (Navarro et al. 2004), suggesting that Kiss1 advances puberty when administered centrally and that its GnRH-triggering function is upstream of GnRH neurons, which is what would be expected from a leptin intermediate. Most importantly, kisspeptins are potent stimulators of GnRH release (Irwig et al. 2004, Thompson et al. 2004, Messager et al. 2005, Pielecka-Fortuna et al. 2008). While these above findings were close to sealing the nature of the leptin intermediate, it was surprisingly found that deletion of the leptin receptor from Kiss1 neurons (Donato et al. 2011) or ablation of >95% of kisspeptin neurons before puberty resulted in normal puberty and fertility (Mayer & Boehm 2011). Furthermore, transgenic expression of the kisspeptin receptor in neurons of leptin receptor-deficient mice failed to rescue their sterility (Cravo et al. 2013). Thus, these studies dampened the excitement of a kisspeptin–leptin intermediate hypothesis, despite the fact that a small amount of kisspeptin might actually be sufficient to trigger puberty (Popa et al. 2013). While kisspeptin expressing neurons are unlikely to be the leptin intermediate neurons, they nevertheless play a critical role in reproduction, perhaps in the maintenance of fertility after puberty. Thus, it would be interesting to tease out the effects of kisspeptin at puberty and subsequently for the maintenance of fertility. In addition to leptin, insulin receptors are located on kisspeptin neurons and could contribute to their effect on reproduction. Interestingly, deletion of the insulin receptor from kisspeptin neurons resulted in delayed puberty, but thereafter, normal fertility (Qiu et al. 2013), adding insulin as a potential mediator of reproduction from kisspeptin neurons.

The involvement of kisspeptin in sexual maturation and reproduction extends to an exciting set of neurons in the ARC, which co-express kisspeptin, neurokinin B, and dynorphin. These neurons, termed KNDy, project on GnRH neurons, stimulate GnRH release (Hrabovszky et al. 2010, Ramaswamy et al. 2010), and get activated during the LH surge (Merkley et al. 2012). Furthermore, central administration of a NK3 receptor (NK3R) agonist in
prepubertal females elicits LH secretion, whereas infusion of an NK3R antagonist delays vaginal opening and LH pulse amplitude (Navarro et al. 2012, Li et al. 2014). The proportion of kisspeptin neurons that express the long form of the leptin receptor (LeprB) is controversial and ranges from 6% (Louis et al. 2011) to 40% (Smith et al. 2006), implying that leptin action on KNDy neurons could be minimal or substantial. Analogous to the fact that a small amount of kisspeptin is required to trigger puberty (Popa et al. 2013), it is equally conceivable, but remains to be determined, that a small number of KNDy neurons expressing LeprB could elicit GnRH release. The small proportion of neurons required to trigger reproductive function is reminiscent of the fact that implantation of preoptic slices containing only 3–140 GnRH neurons is enough to correct hypogonadism in Gnrh-deficient mice (Silverman et al. 1985). While STAT3 signaling is not required for reproduction (Bates et al. 2003), only 15% of KNDy neurons in the ARC phosphorylate STAT3 upon leptin exposure (Cravo et al. 2011). Thus, KNDy neurons that bind leptin, express LeprB, but suppress STAT3 phosphorylation will be critical for unraveling a leptin-kisspeptin STAT3-independent pathway for initiating reproductive function.

Premammillary ventral nucleus

An illuminating finding that throws new light on the search for a leptin intermediate was the identification that the premammillary ventral nucleus (PMV), a site in the hypothalamus that exhibits broad expression of receptors involved in energy metabolism, encompasses leptin-responsive neurons (Donato et al. 2010). Lesions in the PMV resulted in decreased activation of GnRH and Kiss1 neurons, causing low estrogen secretion and deficient feedback of GnRH on proestrus (Donato et al. 2013), demonstrating that the PMV is a critical area that mediates the effects of leptin on reproduction. Another tantalizing finding is the uncovering that neurons directly responsive to leptin, located in various brain sites including the PMV, POA, ARC, and dorsomedial hypothalamus, release the neurotransmitter nitric oxide (Donato et al. 2010). While it has been previously known that targeted inactivation of the mouse nitrous oxide synthase (Nos1) gene caused infertility in the presence of normal leptin levels (Gyurko et al. 2002), recent studies have demonstrated that deletion of the neuronal Nos gene, or its pharmacological inhibition, blunted the ability of exogenous leptin to restore fertility to ob/ob mice (Bellefontaine et al. 2014). In the same study, leptin has been found to coordinate fertility by acting on neurons in the POA to induce NO synthesis via activation of neuronal neuronal nitric oxide synthase (nNOS). Thus, leptin-responsive nitric oxide-releasing NOSs define a new class of PMV neurons that provide input to the neuroendocrine regulation of reproduction.

Overall, it appears that the identification of a single critical leptin intermediate, which indirectly affects GnRH secretion, remains to be fully elucidated, although the primary candidates are KNDy neurons and NO leptin-responsive neurons. Consistent with the notion of redundancies in reproductive pathways, it is quite likely that a single leptin intermediate may not exist, but rather that a web of neuronal connections coordinate the firing of a complex system that is fundamental to species survival. The experimental dissection of these pathways will prove to be difficult but not impossible considering the increasing use of genome-editing tools to knock in and knock out multiple genes in the same cell and generate corresponding mouse models (Wang et al. 2013). Another possibility, which remains to be fully explored and distinct from the stimulatory hypothesis of a leptin intermediate, is the removal of a restraint imposed on GnRH neurons, such as from epigenetic mechanisms. All these approaches should help us in the following decade to expand on novel findings and derive conclusive mechanisms for the reproductive side of leptin.

Leptin in maternal nutrition during pregnancy

The adipose tissue mass plays another role in reproduction, essentially for building up adequate energy reserves to sustain a pregnancy and for the subsequent energy demands of lactation. The first evidence of leptin playing a role in pregnancy originated from the report that pregnant women secrete elevated levels of leptin from the placenta into the maternal circulation (Masuzaki et al. 1997). The effects of leptin during pregnancy were subsequently assessed in leptin-treated and mated male and female ob/ob mice, thus resulting in ob/ob pregnancies that were controlled with exogenous leptin (Mounzih et al. 1998). In this study, withdrawal of leptin treatment at 0.5, 6.5, 10.5, and 19.5 days post coitum (p.c.) of pregnant ob/ob mice did not affect implantation, gestation, or parturition. However, the food intake of ob/ob females continuously treated with leptin during pregnancy resulted, as of day 10.5 p.c., in increased food intake compared with previous days, demonstrating an attenuation effect of the leptin treatment and the establishment of a leptin-resistant state. Consistently, during the late pregnancy of rats, surges in food intake are associated...
with elevated plasma leptin levels and central administration of leptin during this period is less effective at reducing food intake compared with cyclic rats, again demonstrating a period of reduced leptin sensitivity (Johnstone & Higuchi 2001). The onset of leptin resistance in pregnancy was appropriately associated with decreased STAT3 phosphorylation in the VMH (Ladyman & Grattan 2004, 2005) and by the dysregulation of AgRP/NPY and melanocortin neurons (Ladyman et al. 2009). Thus, mid-gestation in the mouse represents the beginning of a leptin-resistant state that could conceivably be derived from synaptic plasticity and reprogramming of neuronal projections into the hypothalamus. These changes would be aimed at establishing a body weight set point that results in increased food intake and adipose mass accumulation. Maternal food intake, whether increased or decreased, was found to program postnatal leptin expression, as demonstrated by elevated leptin expression in adipose tissue and plasma secretion in female pig offspring, whose mothers were allowed for higher food consumption during the second quarter of pregnancy (Eckert et al. 2000). While decreased maternal nutrition during pregnancy can have devastating effects on fetal reprogramming, one effect is that the postnatal leptin surge (Ahima et al. 1998)
is severely attenuated by maternal under nutrition (Delahaye et al. 2008), presumably resulting in delayed growth and puberty.

The mechanisms underlying leptin resistance in pregnancy may be underlined by the same mechanisms as in obesity; however, the triggering factors in either case are likely to be distinct owing to the differences of both physiological states. The onset of leptin resistance in pregnancy is a natural process, which could be carried post partum in subsets of women with obesity (Gunderson & Abrams 2000). The timing for the onset of leptin resistance in pregnancy is predictable and therefore could ease the uncovering of the triggering mechanisms and the associated factors.

Conclusions and future perspectives

Puberty and mid-gestation are physiological states of leptin sensitivity and leptin resistance respectively. Studies centering on understanding the mechanisms that underlie the onset of puberty, and to a lesser extent those involved in leptin resistance during pregnancy, have been exhaustive in the past 20 years. Our knowledge of the triggering neurons has been substantial and potential leptin intermediates have emerged and enlightened the role of leptin in reproduction. First, the location of a previously unappreciated site for leptin action in the PMV opens new avenues to investigate neuronal projections, synaptic plasticity, and neurotransmitters that signal the timing and firing of GnRH neurons to trigger the reproductive cascade. Secondly, the characterization of the kisspeptin system, specifically the KNDy neurons, which are upstream of GnRH, and the potential role leptin plays in these neurons continue to be an exciting pathway to decipher and dissect. Thirdly, the revealing role of AgRP/NPY neurons that influence GnRH neurons is a critical step that bridges the central nutritional pathway elicited by the binding of leptin to first-order neurons to the leptin-responsive neurons in the reproductive axis. The essential criteria for neurons to qualify, as leptin intermediary neurons, is that they would have to respond to leptin, initiate leptin signaling, not necessarily via STAT3 phosphorylation, and stimulate GnRH release most probably through KNDy neurons. In addition, site-specific deletion of the leptin receptor from these neurons would have to result in sterility, irrespective of the presence or absence of obesity. Furthermore, ob/ob and db/db mice should display dysregulation of these intermediary neurons, stemming from their hypoactivation in leptin or leptin signaling deficiency.

While reproductive disturbances are largely dissociated from common obesity, nutritional factors and reproduction are closely connected, as exemplified in states of negative energy balance, when food intake corrects amenorrhea. For example, exogenous leptin induces menstruation in hypothalamic amenorrhea and is thus an appropriate fertility treatment for this disorder (Welt et al. 2004, Chou et al. 2011). Furthermore, in lipodystrophy, leptin treatment, withdrawal, and reinstatement have effects on the progression, interruption, and regain of puberty (Kamran et al. 2012).

A proposed pathway that recapitulates the salient findings summarized in this review is shown in Fig. 1 and outlines the essential neuronal connections that involve leptin, insulin, and IGF1 or their surrogate neurons to converge onto GnRH neurons. As central pathways are critical for any leptin-mediated effect, the focus of future years will be on new first-order sets of leptin-responsive neurons in the PMV and other brain regions that project on GnRH neurons. These neurons will undoubtedly uncover mechanisms that will begin to unravel the complexities of a redundant system essential for species survival.

Finally, a provocative question is whether the primary function of leptin pertains to its metabolic regulation of the reproductive axis. Then conceivably, dysregulation in segments of this pathway, perhaps those that involve the leptin intermediate(s), could cause overweight disorders. While innumerable and challenging arguments would certainly be raised on either side of this hypothesis, it remains worthy of consideration. We still have intriguing questions to ask for many more years and additional lessons to be learnt from the role of leptin in the metabolic control of neuroendocrine reproductive biology.
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