Leap of Faith: Does Serum Luteinizing Hormone Always Accurately Reflect Central Reproductive Neuroendocrine Activity?

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Abstract

The function of the central aspects of the hypothalamic-pituitary-gonadal axis has been assessed in a number of ways including direct measurements of the hypothalamic output and indirect measures using gonadotropin release from the pituitary as a bioassay for reproductive neuroendocrine activity. Here, methods for monitoring these various parameters are briefly reviewed and then examples presented of both concordance and discrepancy between central and peripheral measurements, with a focus on situations in which elevated gonadotropin-releasing hormone neurosecretion is not reflected accurately by pituitary luteinizing hormone release. Implications for the interpretation of gonadotropin data are discussed.

Introduction

A central neuronal component to the control of anterior pituitary function for the regulation of reproduction was appreciated long before hormones from either location could reliably be measured or were even completely identified. Hypophysectomy stopped female reproductive cycles and decreased libido. Replacement via the injection of pituitary extracts restored female cycles and libido [1]. A role for the hypothalamus was confirmed when electrical stimulation of this area was found to induce ovulation in female rabbits [2]. Rosalyn Sussman Yalow shared the Nobel Prize in Physiology and Medicine in 1977 for her work with Solomon Bernson to develop radioimmunoassays. Immunoassays opened a new world for endocrinology in which the dynamics of hormone release took a leading role because the sensitivity and relative simplicity of these methods made it possible to reliably analyze more frequent samples. In 1970, Dierschke et al. [3] published serum concentrations of luteinizing hormone (LH) released from the anterior pituitary of primates as a function of sample frequency, demonstrating the now classic sawtooth pattern of LH pulses. That paper concluded that the rhythmic pulsatile release of LH might be due to ‘intermittent signals from the central nervous system’. This observation was made around the time the first hypothalamic releasing factors were being sequenced by brute force biochemistry. In 1971, Matsuo et al. [4] published the sequence of what is now called gonadotropin-releasing hormone (GnRH), which is acknowledged to be the primary central factor regulating pituitary gonadotropin synthesis and release. Andrew V. Schally and Roger Guillemin, the heads of the laboratories that isolated the initial hypothalamic releasing factors for
pituitary hormones, rounded out the Nobel Committee’s selections for Physiology and Medicine in 1977.

Ernst Knobil and colleagues [5] conducted classic ablation/replacement studies that demonstrated the effects of pulsatile versus continuous GnRH on gonadotropin levels (fig. 1). In these studies, the medial basal hypothalamus was lesioned to eliminate endogenous GnRH release, and then GnRH was replaced intravenously in different patterns. The observations that the administration of GnRH in a continuous manner, rather than mimicking the presumed pulsatile release pattern, decreased circulating LH and follicle-stimulating hormone (FSH) levels within a few days, and that this decrease was reversible upon return to pulsatile administration, demonstrated the importance of the episodic nature of the central signal. This finding remains the basis for the current use of long-acting GnRH agonists to treat both precocious puberty and sex steroid-dependent cancers [6–8] versus episodic administration for fertility restoration [9]. To this day, the generation of episodic hormone release in reproductive neuroendocrinology remains a topic of investigation and lively debate.

**Direct Monitoring of Central Neuroendocrine Activity**

The two methods most commonly used to directly monitor the activity of the central reproductive neuroendocrine system are multiunit activity (MUA) and sampling pituitary portal blood for GnRH (fig. 2). By the first method, MUA is measured using an array of electrodes deployed in a brain area of interest. The electrodes record action potential firing from several neurons in their vicinity, and peaks in this signal are generated when multiple neurons fire in a coordinated manner. MUA measurements, originally in the monkey [10] and subsequently in other species [11–14], demonstrated that peaks in neuronal activity within the medial basal hypothalamus were typically highly correlated with LH pulses. The phrase ‘GnRH pulse generator’ was coined to describe this neuronal activity. There are two drawbacks of MUA records, i.e. the inability to identify the cell type or types being recorded, and at least one degree of separation (release of GnRH itself) between the peaks in neuronal activity and the release of LH.

The second method for directly monitoring reproductive neuroendocrine activity is measuring GnRH released into pituitary portal blood. Portal sampling bridges some of the gap between medial basal hypothalamic MUA and LH release by directly monitoring GnRH release. This technically difficult approach of accessing pituitary portal vessels is required because GnRH has a very short half-life due to enzymatic destruction in the blood; further, the small volume of portal blood is considerably diluted in the cavernous sinus and jugular blood. As such, GnRH cannot be measured in the peripheral circulation. This method was first applied in the rat [15, 16], but the small size of this species mandated the removal of the pituitary, precluding the simultaneous measurement of LH and limiting interpretable sampling to the periovulatory period.

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**Fig. 1.** Administration of hourly pulses of GnRH maintains typical levels of both LH (closed symbols) and FSH (open symbols) in ovariectomized rhesus monkeys. In contrast, continuous GnRH suppressed both hormones within about a week. Adapted from Belchetz et al. [5] with permission from the AAAS.
when a large release of GnRH occurs. Sheep proved to be an excellent model for this approach; the large size allowed multiple simultaneous samples from both pituitary portal and jugular blood in fully conscious animals, permitting direct correlation of the two measurements within the same animal [17]. High-frequency sampling of portal blood suggested that GnRH release in sheep and MUA peaks in primates have a similar shape with a sharp onset to a peak, a high plateau below peak values, then a precipitous decline to baseline [18] (fig. 2). Portal sampling provides an excellent measure of the integrated output of GnRH at the median eminence; it is this GnRH release that regulates the pituitary gland. Of note, recent studies in in vitro preparations using methods that allow more localized monitoring of GnRH release have also identified secretion in the perisomatic region [19] and at the intersection of two GnRH neuronal processes [20]. GnRH released in these areas likely serves neuromodulatory functions that have been suggested in several studies [21–23], and may be under different regulatory control than neuroendocrine release of GnRH for pituitary regulation [24]. With regard to the neuroendocrine release of GnRH, microdialysis and push-pull perfusion have also provided more localized measurements of release near the median eminence [25–27]. The triple crown of simultaneous recording of MUA peaks or other electrical activity, GnRH release and LH release remains unclaimed; however, the high correlation of the neuronal measurement, whether MUA or GnRH release, with LH pulses leads to the reasonable postulate that LH pulses can be used as a bioassay for central reproductive neuroendocrine activity.

Fig. 2. Two methods for the direct monitoring of reproductive neuroendocrine activity. a MUA measurements in the infundibular region of an ovariectomized rhesus monkey; peaks in MUA directly precede LH pulses. Adapted with permission from O’Byrne et al. [28]. b Simultaneous sampling of pituitary portal blood from an ovariectomized sheep at 30-second intervals and jugular blood at 10-min intervals demonstrated a similar correlation between GnRH pulses and LH pulses. Adapted with permission from Moenter et al. [18]. These figures have been scaled so that the x-axis is the same to facilitate comparison.

**Matches and Mismatches between GnRH Release and LH Release**

In many biological states, the assumption that LH pulses reflect GnRH release patterns is likely a valid one. It remains, however, an assumption. Discrepancies between these measures can affect the interpretation of results based on observation of LH release alone.

**The Preovulatory GnRH Surge**

One such mismatch comes during the preovulatory period, when the action of sustained elevated levels of circulating estradiol from the mature follicle(s) triggers one of the rare positive feedback events in physiology, the preovulatory GnRH surge [16, 29–32]. Direct measurements of the GnRH surge in sheep indicate that it begins coincidently with the LH surge, but that it persists for a con-
siderably longer duration (fig. 3a). Blockade of GnRH receptors before the onset of the surge or at different points during the LH surge eliminates or shortens the LH but not the GnRH surge, indicating that GnRH is a prerequisite for the duration of the LH surge [33]. Although the existence of a surge mode of GnRH release in humans remains a topic of debate [34–36], an estradiol-induced GnRH surge has been observed in rhesus monkeys [29], suggesting that at least some Old World primates exhibit this phenomenon; the presence of at least an episodic pattern of GnRH is acknowledged to be necessary for the generation of LH surges even in primates [37]. Of note, with regard to mismatches between GnRH and LH, if a GnRH surge does not exist in humans, this would be an example of a mismatch in which a sustained LH increase is a false positive for increased central reproductive neuroendocrine activity.

Potential central roles for GnRH beyond its established action to induce the LH surge may provide insight into the extended nature of the GnRH versus the LH surge in sheep. Specifically, the duration of the GnRH surge is quite similar to that of proceptive sex behavior in ewes. This suggests a possible role for extended-duration GnRH release in mating. Indeed, the administration of GnRH antagonists after the LH surge is complete can reduce receptivity in ewes [21]. This additional biological action provides a physiological rationale for GnRH release that is unaccompanied by LH release. The time of ovulation appears more related to the onset of the LH surge [38], and ovulation does not require the full spontaneous LH surge [39], suggesting that it is initiated by a relatively short duration of exposure to LH. In contrast, the behavior needed to fertilize an oocyte requires continued exposure to GnRH. These observations indicate that not all release from GnRH neurons is detected by monitoring pituitary hormone in the peripheral circulation, and that GnRH has additional roles beyond the regulation of gonadotropes.

**High-Frequency Episodic GnRH Release**

The clear mismatch in the duration of GnRH and LH release during the preovulatory surge is a once-a-cycle
phenomenon in females, and is a quantitative rather than a qualitative change. Examples of mismatch when GnRH release is episodic and the examination of LH alone produces a qualitatively different interpretation about the central function may more substantially affect how we think about the hypothalamic-pituitary-gonadal axis. One example of this is illustrated in figure 3 [32]. During the midfollicular phase (fig. 3b), 5 GnRH/LH pulse pairs are evident. In contrast, during the late follicular phase just before surge onset, this relationship is less clear and LH release alone does not provide an accurate portrait of GnRH release (fig. 3c). There are several possible explanations for this. First, the amplitude of GnRH release is lower and may less effectively produce clear increases in LH that can be detected. Second, the GnRH release frequency is slightly higher, providing less time for the clearance of LH. The longer half-life of LH would preclude complete clearance from the circulation, thus obscuring pulses. Third, estradiol modifies the GnRH pulse shape [40], potentially altering pituitary responsiveness [41, 42]. Fourth, it is possible that the readily releasable pool of LH is depleted by high-frequency input. Finally, there is a possible technical reason, i.e. that the removal of a portion of portal blood for GnRH measurement may diminish the signal at the pituitary so that well-coordinated LH pulses are not produced. There are other examples in which a high-frequency, clearly episodic GnRH release is associated with circulating LH levels that do not reflect this activity, including thyroidectomized ewes (fig. 4a) [43], long-term castrate rams (fig. 4b) [44], and rams following treatment with the broad-spectrum opiate antagonist naloxone (fig. 4c) [45]. This last example is particularly compelling, because lower-frequency GnRH pulses are accompanied by clear LH pulses (time -3 to 0 h; fig. 4c) in the same animal in which high-frequency GnRH is accompanied by an apparently apulsatile LH release. Unlike the example in figure 3b and c above, the shift in GnRH frequency in response to naloxone in figure 4c is rapid, and the degradation of the LH pulse signal is not likely attributable to any of the above limitations of portal blood sampling. Taken together, these examples indicate that LH is not always a reliable biomarker when GnRH release frequency is high.

The most recent example of a possible mismatch between GnRH and LH release comes from a study of the development of GnRH release during the prenatal through adult period. This study used fast-scan cyclic voltammetry (FSCV) to monitor GnRH release in the median eminence in brain slices. This method takes advantage of the oxidation of GnRH on a carbon fiber microelectrode that can be positioned within a relatively discrete brain region [20]. A GnRH-specific signal was
detected with FSCV in the median eminence, and also in the preoptic area when electrodes were placed near the apposition of two identified GnRH fibers or that of a GnRH fiber and soma. No signal was detected in the median eminence of hypogonadal mice lacking GnRH, demonstrating a specificity of this method for the GnRH decapeptide. Another recent study of cultured GnRH neurons also demonstrated release in the perisomatic region, using uptake of the fluorescent dye FM1-43 [19]. An advantage of these newer approaches is their ability to detect a release within very localized regions; a disadvantage is that both are currently accomplished in an in vitro preparation of either cultured cells (FM1-43) or acutely prepared brain slices (FSCV). This precludes simultaneous monitoring of LH, introducing another potential caveat that must be considered. Of note in this regard, for adult male mice, the frequency of median-eminence GnRH release detected with FSCV in vitro [47] is similar to that of LH pulse frequency in vivo [48, 49]. When FSCV was used to monitor GnRH release frequency locally in the median eminence in brain slices made from mice aged from embryonic day 18 (E18) through postnatal week 1, somewhat unexpected results were obtained (fig. 5) [47]. A very-high-frequency release was observed from E18 through 1 week of age (∼5 release events/h on E18, ∼15 events/h within 24 h of birth, ∼5 events/h at 1 week). The frequency then dropped to almost 0 by 2 weeks of age. The high-frequency release during the first week was further demonstrated to be vesicle mediated and independent of the neuromodulator kisspeptin. These observations were initially surprising, as LH was undetectable at these ages, similar to previous observations [50–52], despite the pituitary expression of both the gonadotropin subunit genes and GnRH receptor [47]. Evaluation of LH release in vivo in response to a single injection of GnRH revealed no secretion at 1 week of age, when the GnRH release frequency was high in the median eminence, but a marked increase in response to GnRH at 2 weeks of age, when GnRH release was essentially absent. Although the pairing together of the in vitro GnRH and in vivo LH measures in different mice of the same ages must be done with caution, one possible explanation for these observations is that high-frequency endogenous GnRH release reduced the responsiveness of the pituitary gland with regard to LH release, which then recovered when the GnRH release frequency dropped about 2 weeks postnatally. Of interest, pituitary follistatin mRNA levels were elevated in control mice at 1 week of age, but declining thereafter [47]; follistatin mRNA is upregulated by high-frequency GnRH [53, 54], adding further evidence that the high GnRH frequencies observed in brain slices at this age may also be received in vivo by the pituitary gland and affect its function even though this is not reflected in LH release. The LH levels in mice through these stages of development suggest a physiological situation in which the GnRH frequency is sufficiently high to lead to pituitary shutdown, similar to both continuous GnRH (fig. 1) and high-frequency GnRH (3–5 pulses/h) in ablation/replacement studies [55].
GnRH, rather than low GnRH driving a disproportionately high LH release. While the relatively long half-life of LH clearly contributes to the obscuring of clear pulses in the circulation, there are other aspects to consider. These include changes in pituitary responsiveness to GnRH as well as other neuroendocrine and peripheral factors (e.g. steroids) that co-vary under these experimental conditions and may play important roles in sculpting the LH output. This could be achieved in part through changes in the GnRH receptor number or myriad postreceptor processes that are reviewed elsewhere [7, 56–59]. Increases in LH sample frequency may help clarify pulses in instances in which this variable is limiting, but in the examples shown this is typically adequate. Alternatively, the use of assays for the free α-subunit, which has a much shorter half-life than the full LH dimer, may be another way to determine if it is the LH half-life alone that leads to a ‘blurring’ of the LH pulse pattern [60]. FSH, which has a longer half-life than LH, has been ignored for this discussion, as it would be even more problematic in terms of reflecting central neuroendocrine activity.

If LH Does Not Always Reflect GnRH Release, Does the Observation of Low LH Levels Always Indicate a Lack of GnRH Release?

This is a question that is critical to sculpting our interpretation of many studies in which LH is the primary variable measured. Direct measurement of central neuroendocrine activity remains difficult. The increasing use of genetic manipulations to probe further up the hypothalamic-pituitary-gonadal axis and study cells that are afferent to the GnRH neuron means that more steps occur between the manipulation and the release of LH. Further, many of these studies are done in small laboratory species because of the power of genetics, a trend that is likely to continue, since CRISPR/CAS9 methodologies make other species more accessible to genetic manipulation [61]. The small blood volume of many of these species means that investigators must attempt to interpret an episodic pattern of LH, and thus GnRH release, from a limited number of samples. Together the biological ‘distance’ of the manipulation from LH measurement and the relatively low resolution of this measurement can contribute to potential inaccuracies in the interpretation.

Studies of the kisspeptin neuronal population in the arcuate nucleus serve to illustrate different ways to interpret the same data set. Arcuate kisspeptin neurons have been postulated to be the source of GnRH pulse generation [62, 63]. These cells are also called KNDy neurons because of their coexpression of both neurokinin B (NKB) and dynorphin, two other neuromodulators thought to be important in fertility control. The anatomical connections of these cells with each other and with GnRH neurons – in combination with their expression of receptors for NKB and dynorphin but not kisspeptin receptors, which are expressed by GnRH neurons – have led to the following model. KNDy neurons act as an interconnected network that is activated by release of NKB and activation of neurokinin 3 receptors. This activation initiates kisspeptin release, which can induce GnRH release both at the soma and at the terminals [24, 64, 65], with the latter release leading to increased LH secretion. The activation of KNDy neurons is also thought to initiate subsequent dynorphin release to terminate their increased activity.
kisspeptin release and eventually GnRH release. Repeated cycles through this series of events would lead to episodic activation of GnRH neurons, hence a ‘pulse generator’.

Many studies have tested various aspects of this model for pulse generation with a wide range of results depending on how the studies were done. For example, NKB and the neurokinin 3 receptor agonist senktide have been reported to increase, decrease or have no effect on LH release [66–70]. While differences in species and endocrine milieux can no doubt explain some of the range of results, it is also possible that a mismatch exists between central reproductive neuroendocrine actions and what the pituitary reports with LH release. Few studies have combined the administration of KNDy peptides with direct measures of central reproductive neuroendocrine output. One study in goats serves to illustrate different directions of movement of a central response (MUA) and LH release.

Wakabayashi et al. [71] monitored LH in response to NKB administered into the lateral ventricle of ovariectomized goats. NKB caused a decrease in LH levels and an apparent loss of the pulsatile pattern (fig. 6). If this had been the only measurement made, a logical conclusion would have been that NKB reduced GnRH release. GnRH itself was not measured in this study, but MUA peaks within the arcuate nucleus that were coincident with LH release were monitored. NKB caused a marked increase in the frequency of peaks in MUA spikes coincident with the drop in circulating LH. Thus, at least one central reproductive measure was increased despite the drop in LH. One possibility is that NKB activated KNDy neurons and this caused a large release of dynorphin, shutting down KNDy neurons and subsequent GnRH release. Another possibility is that NKB activated KNDy neurons, causing a large release of kisspeptin that induced GnRH release, but at

Fig. 7. Selective deletion of estrogen receptor-α from kisspeptin neurons causes early vaginal opening and elevated LH levels that subsequently return to control values but are not associated with estrous cyclicity. ** p < 0.001; *** p < 0.0001. From Mayer et al. [73] with permission. VO = Vaginal opening; WT = wild type.
such a high frequency that the pituitary output was blurred. Finally, it is possible that NKB itself directly activated GnRH release, which has been demonstrated to occur in male mice [72]. While these possibilities remain to be investigated, the simultaneous measurements of both the hypothalamic and pituitary levels of the axis opened the door to additional interpretations.

Another interesting example is from studies in which estrogen receptor-α was selectively deleted from kisspeptin-expressing neurons using targeted expression of cre recombinase, so-called KERKO mice [73]. Female KERKO mice exhibited early vaginal opening and elevated LH levels at 15 and 25 days of age, but no difference at 35 days of age (fig. 7). Despite similar single-point LH values, KERKO animals did not exhibit estrous cyclicity. Kisspeptin neurons are postulated to mediate steroid feedback to the GnRH system; thus the early vaginal opening and elevated LH levels appeared consistent with a loss of negative feedback leading to a precocious activation of GnRH release. But is the subsequent drop in LH due to eventual failure of the KNDy drive to GnRH neurons, or is it due to suppression of the pituitary in response to persistent high-frequency GnRH release? This question cannot be answered from the present data.

**Conclusion**

While LH clearly does not always accurately reflect central reproductive neuroendocrine activity, particularly when the central system is driven at a high frequency subsequent to experimental manipulation, for most physiologic measures, it remains a good bioassay. Exciting new genetic approaches make it possible to activate the central system to extents greater than typical physiology. These approaches are of value in trying to understand the neuronal networks underlying control of fertility, but high levels of central activation can add complexities that need to be considered. As the field implements these exciting new methodologies, one challenge is to make use of output measures that reflect, as much as possible, the nuanced patterning of the reproductive neuroendocrine system. Increased use of new low-volume assays for LH to monitor pulse patterns in mice [49], and optogenetic methods for activating central neuroendocrine pathways in vivo [74], may help in this regard. Moving genetic studies into larger species as CRISPR/CAS9 becomes more standard is another possible approach. While we need to be cautious in drawing absolute conclusions about the central function from examining only the pituitary output, interpretation of results should continue to be pushed to creative limits, so long as speculation is identified as such. Acknowledging multiple alternative mechanisms and interpretations does not diminish the importance of scientific findings. Rather, it strengthens its scholarly aspects and broadens thinking in the field.

**Acknowledgements**

This paper was supported by the National Institutes of Health/Eunice Kennedy Shriver National Institute of Child Health and Human Development (R01 HD34860, R01 HD41469, P50 HD28934). The author thanks Laura Burger, Eden A. Dulka, Carol F. Elias, Fred J. Karsch, Pei-San Tsai, Elizabeth Wagenmaker, Luhong Wang and Nancy L. Wayne for comments during manuscript preparation.

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