Mathematical modelling of endocrine systems

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DOI:
10.1016/j.tem.2019.01.008

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Citation for published version (Harvard):
Zavala, E, Wedgwood, KCA, Voliotis, M, Tabak, J, Spiga, F, Lightman, SL & Tsaneva-Atanasova, K 2019, 'Mathematical modelling of endocrine systems', Trends in Endocrinology and Metabolism, vol. 30, no. 4, pp. 244-257. https://doi.org/10.1016/j.tem.2019.01.008

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Hormone rhythms are ubiquitous and essential to sustain normal physiological functions. Combined mathematical modelling and experimental approaches have shown that these rhythms result from regulatory processes occurring at multiple levels of organisation and require continuous dynamic equilibration, particularly in response to stimuli. We review how such an interdisciplinary approach has been successfully applied to unravel complex regulatory mechanisms in the metabolic, stress, and reproductive axes. We discuss how this strategy is likely to be instrumental for making progress in emerging areas such as chronobiology and network physiology. Ultimately, we envisage that the insight provided by mathematical models could lead to novel experimental tools able to continuously adapt parameters to gradual physiological changes and the design of clinical interventions to restore normal endocrine function.

Understanding the Complexity of Endocrine Regulation Demands an Interdisciplinary Approach

Endocrine axes are the perfect example of complex physiological regulatory systems involving multiple levels of organisation (e.g., central nervous system, secretory glands, tissues, cells, hormones) and timescales [e.g., monthly rhythms, circadian (see Glossary) oscillations, ultradian fluctuations, fast responses]. These systems typically exhibit nonlinear responses, possess multiple components with several feedback loops, and are involved in crosstalk interactions with each other and other body systems (e.g., the immune and nervous systems, the digestive and reproductive apparatus). Endocrine axes are also highly dynamic, with hormone levels exhibiting complex temporal behaviour over short and long timescales that combines sensitivity with robustness, which allows adaptability to physiological challenges. More importantly, dysregulation of these dynamic processes (particularly when it is irreversible) can lead to disease.

Since the seminal work by Norbert Wiener in the mid-20th century, mathematical modelling has helped physiologists to understand how concepts such as negative feedback are key to homeostasis. In endocrinology, new mechanisms of dynamic active regulation have been uncovered to explain the ability to anticipate events and to quickly react to stimuli. Instead of stabilising set points within a certain range, endocrine axes generally control dynamic phenomena (e.g., hormone rhythms, neuron firing, body temperature). Notably, the efforts to uncover the regulatory mechanisms that sustain this ‘homeodynamics’, their robustness in the face of disturbances, their plasticity to adapt to new dynamic regimes (allostasis), and their disruption during disease have largely benefited from mathematics. Some of these benefits have been already described in several reviews. The review in [1] covers general principles of modelling in neuroendocrinology using the growth hormone system as an example, while a recent review by the same authors addresses the contributions of modelling to hypothalamic–pituitary–gonadal regulation.

Highlights

Combining appropriate mathematical models with carefully designed experiments offers great potential to understand complex endocrine regulation at multiple levels of organisation.

Understanding the mechanisms underlying coordinated, rhythmic insulin secretion requires novel mathematical and computational methods that consider the pancreatic islet as a network of beta cells.

Mathematical models, in combination with experimental physiology, have uncovered the mechanisms by which glucocorticoid hormones exhibit normal ultradian pulsatility and respond rapidly to stressors, including during inflammation.

Supported by optogenetic experiments, a hypothalamic neural network comprising kisspeptin secretory neurons has been postulated as driving pulsatile GnRH dynamics involved in the regulation of the reproductive cycle.

The dynamic clamp, a hybrid system integrating electrophysiological measurements with mathematical modelling, enables the interactive manipulation of key parameters in real time.

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pituitary neurosecretory systems [2]. The review in [3] describes in detail several mathematical modelling tools such as types of equations, analysis of their dynamic behaviour (e.g., bistability, oscillations), and approaches to deal with biological noise and systems with multiple timescales in view of applications in endocrinology. This demonstrates a growing interest in the use of quantitative tools and methods to investigate complex hormone dynamics, particularly in relation to stress, reproduction, and metabolism [4–9]. However, an increased appreciation of the insight that mathematical modelling can bring to experimental research could better inform the design of novel interdisciplinary approaches aimed at untangling the complexity of endocrine regulation.

In this review, we show how combining mathematical models with the appropriate experimental set up amounts to the best tool available to understand this complexity. Through examples from the metabolic, stress, and reproductive axes, we illustrate how models can provide insight on dynamic hormone regulation spanning several spatiotemporal scales and the key role that these quantitative models could play in the advancement of chronomedicine. Rather than present the vast and diverse array of mathematical models used in endocrinology, which we feel may be overwhelming, we choose instead to demonstrate how models have been used to answer specific questions. We also discuss an example of a new class of hybrid approaches: the dynamic clamp in electrophysiology, where real-time integration of mathematical modelling with experimental techniques can be used to understand the behaviour of secretory cells. Lastly, through a discussion of open research questions at the intersections between the metabolic, stress, and reproductive axes, we give a perspective of the field and how experimental and clinical research can benefit from mathematical modelling approaches.

The Metabolic Axis: From Mechanisms of Secretion to Beta Cell Coordination and Beyond

Given its strong association with diabetes, insulin secretion by pancreatic beta cells (Figure 1) has been the subject of intense study for almost a century [10]. The primary secretory pathway of glucose-stimulated insulin secretion is associated with complex patterns of electrical activity across the plasma membrane, which allow Ca\(^{2+}\) ions to enter the cell and trigger the secretory machinery. This electrical activity is coupled to cell metabolism, which acts as a glucose sensor by raising the intracellular ATP:ADP ratio, causing K\(_{\text{ATP}}\) channels to close, depolarising the membrane and driving it towards its threshold for action potential initiation [11]. Mathematical models provide an ideal framework to investigate the complex interaction between metabolic and electrical pathways in beta cells over the diverse timescales at which these processes occur.

The majority of mathematical models of beta cell behaviour are based on the Chay–Keizer model [12]. This model, which describes electrical activity and Ca\(^{2+}\) dynamics, has subsequently undergone a plethora of modifications, including those to incorporate glycolytic and mitochondrial components. The primary goal of these models is to elucidate the mechanisms giving rise to pulsatile insulin secretion with a mean period of ~5 min observed in rodents, dogs, and humans [13,14]. To this end, many models consider oscillations in Ca\(^{2+}\) and metabolic activity under the assumption that one of these essentially sets the overall period of the pulses ([15] and references therein). However, the development and subsequent analysis of the dual-oscillator model [16] highlighted that these two mechanisms may actually work cooperatively to generate rhythmic insulin secretion (i.e., that Ca\(^{2+}\) and glycolytic activity can oscillate independently of one another but together give rise to oscillations on the timescale typically observed in experiments). This model has thus been an invaluable tool for studying the interactions of these processes and highlights the importance of understanding the timescales over which they
**Glossary**

**Allostasis:** the adaptive processes by which a physiological regulatory system re-establishes homeostasis (typically with increased fragility) to compensate for physiological disruptions.

**Bistability:** the coexistence of two stable equilibrium states that, under certain conditions, may be observed in a dynamical system.

**Chronomedicine:** a novel approach to medicine that focuses on understanding the natural rhythms of the body for the prevention, diagnosis, and treatment of diseases. Associated terms are chronodisruption (when the timing of perturbations is key to the disruption of a physiological rhythm), chronotherapy (when the timing of treatment is key to restoring health), and chronobiology (when referring to rhythms not exclusive to humans).

**Circadian:** a biological rhythm displaying an oscillation period of about 24 hours.

**Insulin resistance:** a decrease in the sensitivity of target tissue to insulin.

**Nonlinear response:** the response of a regulatory system in which the change of the output is not proportional to the change in the input.

**Oestrous:** refers to the onset of a reproductive cycle in most mammals.

**Relaxation oscillator:** an oscillator that achieves its rhythmicity from repetitive cycles of accumulation and discharge (e.g., electrical activity, neurotransmitter concentration).

**Robustness:** the ability of a system to preserve its dynamic behaviour while coping with perturbations.

**Sensitivity:** the ability of a system to respond rapidly (and/or with large excursions) to stimuli.

**Syncytium:** the conceptualization of a cohort of multiple cells exhibiting such a degree of interconnectedness and synchronised behaviour that they may be understood as if they were a single cell (e.g., multinucleate cell).

**Ultradian:** a biological rhythm displaying an oscillation period of less than 24 hours.
The dual-oscillator model has since been modified to incorporate Ca$^{2+}$ feedback to glycolytic activity. This improved integrated oscillator model [15,18] further highlights that neither oscillations in Ca$^{2+}$ nor metabolism establish the overall rhythmicity in beta cells by themselves [19,20] and exemplifies how models can be developed in light of new experimental evidence.

One of the striking features of beta cells is that within islets they exhibit tight synchronisation of regular oscillations in electrical activity, while isolated cells oscillate irregularly ([21] and references therein). This phenomenon has been mathematically modelled by considering the islet as a network of beta cells. Under the heterogeneity hypothesis [22], variability in individual cells is ‘smoothed’ by intercellular interactions so that the network may be thought of as the average of the cells in it. This has led to the idea that islets are essentially a syncytium, with no single cell dictating the overall network response. However, this notion has been challenged by novel optogenetic experiments that show that silencing the activity of a single (specific) cell can disrupt electrical rhythms across the entire islet [23]. The presence of these so-called hub cells can be understood through the application of computational graph theory to the islet. Graph theoretic models place importance on the presence and nature of interactions within islets rather than the dynamics of individual beta cells [24]. Such models emphasise the dependency of these interactions on the extracellular concentrations of glucose [25] and that heterogeneous coupling could give rise to networks supporting hub cells [26], features that would be difficult to understand without an underlying model. Despite the success of using graph theory in this system, there is currently no experimental nor mathematical model that explains the results from the hub cell silencing experiment, but it is likely that combining the two approaches will be necessary to do so.

Alongside secretory deficiencies, insulin resistance is one of the primary mechanisms associated with the development of type 2 diabetes [27]. To investigate this, a recent phenomenological model [28] describes whole-body responses to insulin resistance including upregulation of beta cell function on short and medium timescales and changes to beta cell mass over longer timescales. Importantly, the model predicts the effect of temporary weight gain and loss as well as medical procedures such as gastric bypass surgery. The study introduces the notion of a threshold for decreases in insulin sensitivity: small decreases can be compensated for effectively whereas larger decreases cannot. In particular, the model highlights how feedback mechanisms to counter insulin resistance can contribute to the development of diabetes once the threshold has been crossed. The related concept of personal fat thresholds [29] is already being used to develop diet plans for diabetic patients; mathematical modelling has the potential to further support such interventions. Critically, analysis of the mechanisms in the model that establish the threshold explain why preventing diabetes is significantly easier than reversing it, exemplifying how models can be used not only to design therapeutic interventions (see Box 1 for an example), but also to direct public policy.

Figure 1. The Metabolic Axis. Regulation of blood plasma glucose levels is achieved primarily through the complementary actions of the hormones insulin, glucagon, and somatostatin. Insulin promotes the absorption of glucose from the blood by the liver and peripheral tissues, thus lowering the blood glucose concentration. In these tissues, glucose is then converted to glycogen or fat and subsequently stored. Glucagon plays the opposite role to insulin, encouraging tissues to transform these substrates back into glucose for secretion into the bloodstream. Somatostatin inhibits the secretion of insulin and glucagon by, respectively, beta and alpha cells, both of which reside in multicellular structures known as the islets of Langerhans, which are located in the pancreas. Mathematical models of beta cell behaviour typically account for the electrical activity originating from ion channels involved in insulin secretion. Recent models have also accounted for beta cell metabolism, including, for example, the glycolytic activity and mitochondrial components shown in the ‘dual-oscillator model’ (see text).
The Hypothalamic–Pituitary–Adrenal (HPA) Axis: A Choreography between Hormone Rhythms and the Stress Response

The body’s response to stress is mediated by several hormones, a crucial one being cortisol. Cortisol belongs to a group of glucocorticoid steroid hormones with a broad spectrum of context-dependent effects. Because they are rapidly secreted in response to physical and psychological stressors, they are commonly known as stress hormones. In the clinic, synthetic glucocorticoid hormones are widely prescribed for their anti-inflammatory effects as well as in hormone replacement therapy [42]. The circulating levels of glucocorticoids – cortisol in humans, corticosterone in rodents (CORT) – are dynamically controlled by the activity of the hypothalamic–pituitary–adrenal (HPA) axis (Figure 2), which is characterised by the rhythmic secretion of corticotropin-releasing hormone (CRH) and arginine vasopressin (AVP) from the paraventricular nucleus of the hypothalamus (PVN), adrenocorticotropic hormone (ACTH) from the pituitary, and CORT from the adrenal glands. Despite cumulative evidence showing the importance of CORT rhythms for immunological, cognitive, reproductive, and metabolic functions [42,43], little attention has been paid to developing the dynamic aspects of glucocorticoid drug therapies. From a theoretical point of view, understanding how the HPA axis sustains rhythmic activity while simultaneously eliciting fast, transient, and proportionate responses to stressors constitutes a major challenge.

One of the key steps in understanding the dynamic activity of the HPA axis relates to the causal relationship between ACTH and CORT secretion. A pioneering mathematical model addressed this challenge by accounting for several steps of the signalling pathway: the activation of a putative ACTH receptor in the membrane of adrenocortical steroidogenic cells, its relay via cAMP in the cytosol, the mitochondrial import of cholesterol (the substrate for CORT biosynthesis), and the synthesis and secretion of CORT [44]. The model was fitted to adrenal secretory rates of cortisol and blood ACTH concentrations measured in dogs subjected to intravenous infusions of ACTH. Importantly, this model predicted changes in adrenal sensitivity between small versus large pulses of ACTH, a phenomenon that has been further identified and investigated in other mammals. Subsequent models considered the feedback loops that glucocorticoids exert at the level of the pituitary and hypothalamus [45,46]. These models offered qualitative predictions of feedback-generated ultradian oscillations in CORT levels and suggested possible ways to include circadian modulation. In this sense, [45] showed that coupling this feedback mechanism with a central nervous system-driven pulse generator enables both ultradian and circadian variability in hormone secretion. These early models also
Endogenous glucocorticoids (CORT) are vital hormones involved in many physiological processes that are key to homeostasis and survival (e.g., mediating the stress response, anti-inflammatory and immunosuppressive effects, regulation of glucose expenditure). The circulating levels of CORT are controlled by the HPA axis. Corticotropin-releasing hormone (CRH) and arginine vasopressin (AVP) stimulate the release of CORT.

(Figure legend continued on the bottom of the next page.)
aimed at explaining specific physiopathological changes such as stress, infusion of synthetic glucocorticoids, and adrenalectomy. Interestingly, the model in [46] also proposed a bistability mechanism that would explain the allostatic transition of the HPA axis subjected to chronic stress.

Although the models in [45,46] demonstrate the possibility of ultradian oscillations generated through negative feedback, the predicted frequency of these oscillations significantly differs from the near-hourly oscillations observed in humans. It was not until the work by Walker et al. [47] that the mechanisms underlying ultradian oscillations were correctly predicted as originating from the negative feedback loops between the pituitary and adrenal glands, while the hypothalamic drive provides the source of circadian modulation. This model predicted near-hourly oscillations of ACTH and CORT secretion supported by in vivo data, even in the presence of a constant hypothalamic CRH signal. Subsequent experiments confirmed this model prediction [48,49], which demonstrated that a previously hypothesised hypothalamic ‘pulse generator’ [50] is not essential to generate ultradian glucocorticoid oscillations.

While recent mathematical models of the HPA axis have focused on the role of glucocorticoid dynamics in mental health [51,52], others have investigated the stress response, the role of nuclear receptors, and inflammation [53–55]. Understanding how healthy adrenal glands achieve rapid CORT secretion while simultaneously preventing their uncontrolled release in response to stressors is key to explaining the dysregulation observed in endocrine disorders such as Addison’s disease and Cushing’s syndrome (Box 2). In this direction, the work in [55] combined experimental physiology and mathematical modelling to predict how surges of ACTH may be decoded by the adrenal gland, hypothesising that the control mechanism may comprise an intra-adrenal negative feedback loop mediated by the glucocorticoid receptor. The organisation of the molecular mechanisms involved in such intra-adrenal regulation was postulated in [54], distinguishing between slow genomic and fast non-genomic signalling pathways. These mechanisms were mathematically modelled as a regulatory network that not only predicted the transient dynamic responses observed during the stress response but explained how the adrenal glands can decode ACTH pulses of different magnitudes, including those observed during inflammation.

The Reproductive Axis: Uncovering the Mechanisms of GnRH Pulsatility
Hormone signals within the hypothalamic–pituitary–gonadal (HPG) axis (Figure 3) are critical for reproduction, with a key regulatory process being the pulsatile release of gonadotropin-releasing hormone (GnRH) from the hypothalamus onto the pituitary gland. Mathematical models have provided insight into how GnRH pulsatility controls the synthesis and secretion of gonadotropic hormones [luteinizing hormone (LH) and follicle stimulating hormone (FSH)] from the pituitary. Early experimental work on primates revealed the dependence of gonadotropin secretion on GnRH frequency by showing that pulsatile but not constant delivery of exogenous GnRH can restore gonadotropin secretion in animals with hypothalamic lesions [58]. It is now clear that gonadotropin secretion is suppressed when the GnRH frequency is

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either too high or too low, and this effect is mediated through complex signalling networks that allow cells to regulate the synthesis of LH and FSH differentially in response to GnRH frequency [59,60]. Several mathematical models related to GnRH signalling have been proposed [61] and a mechanistic model of the pathway has shown that the nonlinear relationship between gonadotropin secretion and GnRH pulse frequency is most likely due to the convergent feed-forward architecture of the network [61,62]. The model suggests that frequency decoding is primarily achieved due to the synergistic effect of multiple signalling pathways (e.g., the extracellular signal regulated kinase (ERK) pathway and the nuclear factor of activated T cells (NFAT) pathway) on the expression of gonadotropin-related genes. This contrasts with upstream negative feedback interactions (e.g., due to agonist-induced receptor internalization) that were previously thought to play a crucial role in frequency decoding. Instead, the model shows that feedback plays a different role, allowing the pituitary system to cope with cell–cell heterogeneity and process GnRH information more reliably [63].

At the level of the hypothalamus, a coarse-grained neuronal population model has advanced our understanding of how GnRH pulsatility is sustained and regulated [64]. The model draws on recent experimental work, which demonstrates the pivotal role of neuropeptide signalling within the arcuate nucleus kisspeptin population for GnRH pulse generation [65,66]. The model supports the idea that the kisspeptin population drives GnRH pulses by postulating that it operates as a relaxation oscillator due to neuropeptideergic negative and positive feedback interactions mediated by neurokinin B and dynorphin, respectively. Furthermore, the model predicts that pulsatile dynamics depend on basal activity levels in the kisspeptin population and highlights the tipping-point behaviour of the system as basal activity increases. Using optogenetics, these model predictions were confirmed in vivo, showing that pulses can be directly controlled in oestrous mice by selectively exciting kisspeptin neurons in the arcuate nucleus with continuous low-frequency (1 Hz and 5 Hz) light stimulation [64]. Thus, this is yet another example of how even simple phenomenological models can lead to useful and experimentally testable insights.

Mathematical modelling has also been employed to understand the macroscopic processes involved in follicular development [67]. Although gonadotropins are known to control the development of ovarian follicles and their secretory activity, little attention has been given to...
sex steroid secretion and how it feeds back to upstream components of the HPG axis modulating GnRH and gonadotropin secretion. These feedback interactions underpin the ovarian cycle and have a critical role in women’s physiology and reproductive health, thus representing a unique opportunity for experimental physiologists, clinicians, and mathematical modellers alike [68].
Hybrid Systems: A New Paradigm to Establish How the Parts Contribute to the Whole

Like alpha and beta cells in the pancreas, the five endocrine cell types of the anterior pituitary generate electrical activity in the form of spikes and bursts [69]. Electrical activity brings Ca\textsuperscript{2+} into the cells through ion channels, which triggers hormone secretion and stimulates vesicle refilling. In the absence of hypothalamic signals, pituitary gonadotrophs fire sharp spikes at a slow rate, releasing very little hormone. By contrast, lactotrophs and somatotrophs fire in bursts, which are longer electrical events than spikes. Bursts provide more time for Ca\textsuperscript{2+} to enter cells, so lactotrophs and somatotrophs have a high basal rate of hormone release [70]. While pituitary cells have similar amounts of most voltage- and Ca\textsuperscript{2+}-activated channels, they differ in the amount of large-conductance potassium (BK) channels. Lactotrophs and somatotrophs have a high density of BK channels, while gonadotrophs have very few [71]. This is paradoxical because BK channels are repolarising channels (in neurons and other cell types). BK channels typically open quickly during an action potential, reducing its duration. However, in pituitary cells like somatotrophs and lactotrophs, BK channels seem to increase event duration, turning spikes into bursts. This prompts the question of whether gonadotrophs would burst if they expressed BK channels.

A mathematical model predicted that assimilation of BK channels into gonadotroph electrical activity can switch its firing dynamics from spiking to bursting. By opening quickly at the beginning of an action potential, BK channels limit the activation of other, slower K\textsuperscript{+} channels, which in turn prevents these channels from repolarising the cell [72]. Analysis of the model suggests that this effect is robust to changes in the expression of other ion channels [73] but leaves open the question of whether fast BK current activation promotes bursting in real cells. This problem was elegantly solved through the dynamic clamp technique (Figure 4). Bursting lactosomatotroph cells first had their BK channels blocked by a channel antagonist, resulting in a switch from bursting to spiking in most cells. Then, a BK current calculated in real time from a mathematical model was added back to the cells via a computer-assisted dynamic clamp. This made the cells switch back to bursting and, importantly, it occurred only if the modelled BK current was fast enough, demonstrating that the mechanism identified by the mathematical model was correct. Finally, use of the dynamic clamp to add a model BK current into spiking gonadotrophs made these cells switch to bursting, demonstrating that the difference between electrical activity patterns in lactosomatotroph and gonadotroph cells could be explained by the presence or absence of BK channels [73].

The dynamic clamp was instrumental in establishing the role of BK channels, by linking a model-based mathematical mechanism to real pituitary cells. It illustrates the power of hybrid systems to combine experiments and modelling. Another elegant example of such a system was developed by Dhumpa et al. [74] to show that islets of Langerhans can synchronise their insulin secretion through feedback from the liver. To do so, they introduced islets loaded with a fluorescent Ca\textsuperscript{2+} indicator into a microfluidic chamber and interfaced the global Ca\textsuperscript{2+} signal from the islet population with a mathematical model of glucose release by the liver in response to insulin. The modelled glucose level was then delivered back to the islet chamber. Without liver feedback, the islets produced independent oscillations. As soon as the feedback was turned on, however, the islets began to synchronise, as evidenced by the resulting global Ca\textsuperscript{2+} oscillation, out of phase with the resulting glucose oscillation. This demonstrated that the liver might act as a coordinator of activity in the islet population and enabled testing of the effectiveness of this coordination as the speed of the liver feedback was varied. Thus, hybrid systems allow us to determine the role played by components of a biological system, by controlling key parameters of such components, particularly the timescale on which they operate.
Concluding Remarks and Future Perspectives

The complexity of endocrine systems is evidenced by the number of molecular interactions occurring at multiple levels of organisation that are necessary to achieve robust control of hormone secretion. Strikingly, many endocrine axes exhibit the same control strategies to regulate hormone levels within a homeostatic range: feedback loops, network organisation of components, and collective behaviour that cannot be explained solely by investigating the dynamics of individual cells. Here, we have reviewed recent examples from three major

Outstanding Questions

Hormone pulsatility in endocrine systems is ubiquitous, but whether this is an optimal solution compared with constitutive secretion needs to be investigated. Is there an energetic advantage in the reduced amount of hormone needed for pulsatile signalling?

Compared with constant hormone levels, rhythmic hormone secretion contains more information in the form of, for example, amplitude and frequency. How is this information encoded and how do different tissues read the same blood signalling message in different ways?

Hormone rhythms are known to affect gene expression at different timescales and in a context-dependent way. At a cellular level, how are chromatin responses determined by the pattern of nuclear receptor activation?

How does chronodisruption, especially between different local and central pacemakers, cause disease?

Can our increased knowledge of hormone dynamics be used in therapeutics that improve patient care?
endocrine axes where mathematical models have delivered insight about dynamic behaviour that was difficult to interpret solely by looking at the experimental data. The relevance and timeliness of using mathematical tools to understand these control strategies is largely driven by the urgency of understanding their dysregulation in reproductive, metabolic, and stress-related conditions, including complex psycho-immunoneuroendocrine disorders [6,75–77].

The key role of hormone dynamics in health and disease has suggested future research avenues at the interface of mathematical modelling and experimental neuroendocrinology. In this sense, frequency encoding and decoding mechanisms underlying pulsatile hormone secretion remain an understudied area [77]. For instance, ultradian hormone stimulation is known to induce glucocorticoid receptor-mediated pulses of gene transcription [78], and there exists a growing realisation that understanding how the dynamics of glucocorticoid signalling affects gene regulation is key to the design of effective chronotherapies [79,80]. The development of such understanding will be likely to involve modelling the role of hormone pulsatility on continuous dynamic equilibration and stochastic dynamic interactions at the level of DNA binding [75].

Another burgeoning area of research is the crosstalk interactions between endocrine axes. For instance, hypercortisolism induced by chronic stress, Cushing’s syndrome, or medication is a known risk factor for the development of diabetes. This has prompted investigations on the links between glucocorticoid dynamics and insulin secretion and resistance [81–83]. Similarly, a mathematical model linking the HPA and metabolic axes describes a way in which circadian glucocorticoid oscillations regulate a transcriptional circuit underlying adipocyte differentiation [84], suggesting mechanisms by which conditions that disrupt pulsatile glucocorticoid secretion could lead to obesity. By contrast, insulin-induced hypoglycaemia is an acute stressor that both significantly activates the HPA axis and inhibits pulsatile LH secretion in rats [85], evidencing crosstalk interactions between the metabolic, stress, and reproductive axes. Gender differences in endocrine regulation are also being investigated via mathematical methods, as suggested by a model exploring the effects of testosterone on the HPA axis response to stress [86].

While in most experimental research it is sufficient to ‘let data speak for itself’, existing experimental protocols as applied to complex endocrine phenomena often struggle to combine data at different levels of organisation. As a result, the mutual interactions between factors underlying endocrine regulation and the different timescales at which they occur are often ignored. This is where mathematical models offer a solution to interpret the data and gain insight on the underlying dynamics. Moreover, models help us think beyond the limits of ‘what we can do’ at the laboratory bench and start asking ‘what if’ questions. This not only stimulates creative interdisciplinary collaborations but also advances the field by replacing a static, snapshot view of endocrine function with one where complex, multiscale regulation underpins hormone dynamics (see Outstanding Questions).

Acknowledgments

This work was funded by the Medical Research Council (MRC) through MRC Fellowship MR/P014747/1 (to E.Z.), MRC Fellowship MR/P01478X/1 (to K.C.A.W.), and MRC Grant MR/J008893/1 (to F.S., E.Z., and S.L.L.), Engineering and Physical Sciences Research Council (EPSRC) Grant EP/N014391/1 (to M.V., S.L.L., and K.T.A.), and Wellcome Trust Grant WT105618MA (to K.C.A.W. and K.T.A.). This study did not generate any new data.

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