Multiple-scale neuroendocrine signals connect brain and pituitary hormone rhythms

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Small assemblies of hypothalamic parvocellular neurons release their neuroendocrine signals at the median eminence to control long-lasting pituitary hormone rhythms essential for homeostasis. How such rapid hypothalamic neurotransmission leads to slowly-evolving hormonal signals remains unknown. Here, we show that the temporal organization of dopamine release events in freely-behaving animals relies on a set of characteristic features that are adapted to the dynamic dopaminergic control of pituitary prolactin secretion, a key reproductive hormone. First, locally generated dopamine release signals are organized over more than four orders of magnitude (0.001 Hz - 10 Hz). Second, these dopamine events are finely-tuned within and between frequency domains as building blocks that recur over days to weeks. Third, an integration time window is detected across the median eminence, and consists of high-frequency dopamine discharges that are coordinated within the minutes range. Thus, a hierarchical combination of time-scaled neuroendocrine signals displays local-global integration to connect brain-pituitary rhythms and pace hormone secretion.

A remarkable function of the brain is its capability to integrate temporal information with complex physiological responses. This has been well established for behavioral responses such as non-rapid eye movement (NREM) sleep, where three neuronal oscillations with distinct frequency bands support information transfer (1). Yet the neuronal mechanisms that orchestrate the dialog between the brain and other basic functions like reproduction, lactation and growth remain largely unknown (2-5). They depend on the fine tuning of pituitary hormone pulses by small assemblies of hypothalamic neuroendocrine or parvocellular neurons, which release specific secretagogues at the median eminence (ME) (4, 6).

Here, we took advantage of the anatomical organisation of the ME to investigate how the tuberoinfundibular (TIDA) neuronal population (7, 8) releases dopamine (DA) to negatively regulate pituitary secretion of prolactin (PRL), a key reproductive hormone (2). To do so, miniaturised amperometric carbon fiber implants were used to detect DA release events (9) for days to weeks in freely-behaving mice. Using this approach, we uncovered a hierarchically-organized delivery of release events over four orders of magnitude (from <0.1 sec to several hours), which correlate with the dynamics of PRL in the bloodstream.

RESULTS

Frequency-coding of DA release events in vivo

To characterize the release dynamics of TIDA nerve terminals in vivo, we employed long-term constant voltage amperometry in awake mice using thin (30 μm tip diameter) carbon fibers implanted into the ME (Fig. 1A). Voltage was clamped at -700 mV to allow detection of DA released from TIDA neurons. DA amperometry was performed continuously during several days, and the relationship with PRL secretion was assessed using tail blood micro-sampling for high-sensitivity mPRL ELISA developed in-house (10) (Fig. 1A). Single carbon fiber electrode recordings revealed robust DA currents (median 325 nA, IQR: 127 to 822 nA) due to oxidation of DA to dopamine-o-quinone (Fig. 1A), and these could be robustly detected over the long term (Fig. 1B) (n = 7 virgin female mice). We then used DA as a relevant readout to explore the dynamics of TIDA neuron population function in freely-behaving animals. DA currents at the ME level discharged over different timescales (Fig. 1B, C) and more frequently during the night than day (Fig. 1D) (mean counts/h from ZT 0, in 6 hour blocks: 18.7, 27.2, 28.6, 30.4), implying that the strength of TIDA neuron excitability is likely modular around the day/night switch. DA release events were often grouped and interspaced by long-lasting (dozens of minutes to several hours) silent periods, suggesting nested relations between high and low frequency output patterns (Fig. 1B,C). No clear association between DA current density and estrus cycle stage was detected (Fig. 1C and Fig. S1).

Analysis of inter-event intervals (IEI) for DA release unveiled a wide range of time intervals, from less than 100 ms to a few hours, with two principal frequencies of 1.5 Hz and 12 Hz (Fig. 1E). A delay of several minutes between decreasing PRL levels and the onset of DA release was often observed. The neuronal mechanisms that or-ganize nested relations between high and low frequency output patterns (Fig. 1B,C). No clear association between DA current density and estrus cycle stage was detected (Fig. 1C and Fig. S1).

The hypothalamo-pituitary axis controls a wide-range of homeostatic processes including growth, stress and reproduction. Despite this, the hypothalamic neuron firing patterns that lead to slowly-evolving pituitary hormone rhythms remain enigmatic. Here, we employed in vivo amperometric recordings in freely-behaving mice to investigate how tuberoinfundibular neurons release dopamine (DA) at the median eminence (ME) to control pituitary prolactin secretion. Using this approach, we show that DA release occurs as multiple locally-generated and time-scaled secretory events, which are integrated over a range of minutes across the ME. These results provide a broad physiological mechanism for the dialog that occurs between the brain and pituitary to dictate hormone rhythms over multiple timescales, from ultradian to seasonal.

Significance

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We next examined whether these sub-second DA release events possess a secondary/tertiary organization at the local level. In vivo monitoring of DA release events at the median eminence level. (A) Electrodes were implanted at the median eminence (ME) of mice and dopamine (DA) was detected using constant voltage amperometry. Serial blood micro-sampling was performed from the tail vein. (B) Representative 24 hour recording of DA release, with zoom of a 10-minute sequence (bottom). (C) Representative 11 day recording from a female mouse. Each vertical line corresponds to a single secretion event. The stage of the estrus cycle is indicated on the left for each day (M, metestrus; D, diestrus; P, proestrus, E, estrus). (D) Mean distribution of DA release events during the day (n = 80 days from 7 female mice). (E) Histogram of inter-event intervals (IEIs): two prominent frequencies are apparent at 1.5 and 12 Hz (≥ 1 Hz) were not synchronized (Fig. 3A), DA secretory response to a i.p. injection of 1μg ovine PRL (PRL injected at time 0)(from 5 animals, 7 injections). (F) Relation between DA and PRL. Average normalized PRL levels occurring around a DA event (n = 501 DA events, from six 1 hour long sessions) (black, mean; blue, SEM). (G) DA secretory response to a i.p. injection of 1μg ovine PRL (PRL injected at time 0)(from 5 animals, 7 injections). (H) Distribution of the IEIs of DA events induced by i.p. injection of PRL. (I) Example of simultaneous recording of PRL levels (red) and DA release events (black). In all cases, bar graphs show the mean ± SEM.

Long-range organization of DA release events at the local ME level

We next examined whether these sub-second DA release events possess a secondary/tertiary organization at the local level i.e. in the close vicinity of carbon fiber tips. Using cluster analysis to group DA currents on the basis of their shape, and bootstrapping to identify temporal series of events appearing with a higher-than-chance frequency during the recording period, a specific pattern of DA release events was found, with stereotypical features remaining consistent between different animals recorded on different days (Fig. 2A, Fig. S4).

Further analyses demonstrated that these stereotyped patterns of DA release events were not randomly-distributed, but rather appeared as chains of sequential events within the same group and/or between groups (Fig. 2 B-E). These recurrent motifs of DA release events were scaled from the millisecond (Fig. 2B-D) right up to the hour (Fig. 2E) range, and could even be detected over days (Fig. 2F-I). Thus, the mechanisms controlling TIDA neuron activities appear to be inherently robust.

Local-global integration of DA release events across the median eminence

A long-standing question regarding parvocellular neuron function is how nerve terminals discharge their neurohormones across the ME to sculpt pituitary output (2-4, 6). Given that TIDA nerve terminals abut over the whole ME (7, 8), dual-carbon fiber recordings were carried out 500 μm apart rostro-caudally, spanning the population (n = 3 animals). While distant DA events at high frequencies (≥1 Hz) were not synchronized (Fig. 3A), DA events were coordinated with IEIs in the minutes range during lactation. Further analyses demonstrated that these stereotyped patterns of DA release events were not randomly-distributed, but rather appeared as chains of sequential events within the same group and/or between groups (Fig. 2 B-E). These recurrent motifs of DA release events were scaled from the millisecond (Fig. 2B-D) right up to the hour (Fig. 2E) range, and could even be detected over days (Fig. 2F-I). Thus, the mechanisms controlling TIDA neuron activities appear to be inherently robust.

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This organization occurs more than four orders of magnitude of frequency, from infra-slow vessels in a freely-behaving mouse model. DA release events are repeated over weeks as network-driven rhythms that cover (E-F) Example of temporal patterns on DA release. Each dot represents a single DA release event, colored depending on the subgroup (as in figure 2A). Each line shows one repetition of the sequence during the recording; five examples of repetition are shown for each pattern. (F-I) Frequency of the four temporal patterns during 8 days of recording.

**DISCUSSION**

Our results show how an ensemble of parvocellular TIDA neurons delivers its neuroendocrine products towards ME portal vessels in a freely-behaving mouse model. DA release events are repeated over weeks as network-driven rhythms that cover more than four orders of magnitude of frequency, from infra-slow (<0.001 Hz) to fast rhythms (1-10Hz). This organization occurs not only locally within, but also across the TIDA neuron assembly, as DA release events are scaled over the minute-range throughout the ME (Fig. 4).

Specifically, the use of miniaturized carbon fibers stereotaxically-implanted into the ME allowed us to detect and discriminate DA-related currents in vivo, which were far more complex, but also more organized than spike firing activities recorded in parvocellular neurons from either brain slices (9, 14-17) or anesthetized animals (18). Even though the small tip of the carbon fiber was likely able to detect DA release from only a few TIDA neurons, we observed a variety of rhythms. First, high frequency (about 1 and 10 Hz) events were prominent locally but not synchronized globally. As the site of recording is variable and these rhythms were observed in all animals, a large number of local DA release processes presumably originate from TIDA neurons capable of secreting at high rates. The latter would be considered as "executive" in the top-down control of pituitary PRL rhythms by hypothalamic DA inputs, since they coincided with drops in pituitary PRL secretion. Second, slower rhythms of DA release (with time periods of minutes to hours) were detectable locally due to the ability of small carbon fibers to measure DA events over days to weeks with no noticeable deleterious effects. Strikingly, these were not distinguishable from high frequency DA events with which a hierarchical combination occurred regarding both the specific frequencies generated and how they organize in time as temporal motifs. Since the local-global integration of high frequency DA
neuronal neighbours within the arcuate nucleus (19). As D1 and D2 receptors are expressed in the ME (20), this raises the possibility that DA release events at the ME, even those organized over slow rhythms, may also contribute to the regulation of other neuropeptides, such as those underlying circadian locomotor- and growth-hormone pulses (20, 21).

The discovery of a multiple-time-scale integration of DA delivery at the neurohemal space provides a hitherto unknown element concerning how the brain dialogues with peripheral organs via a neuroendocrine connection. Such hierarchical organization of rhythms has been observed in other brain regions where multiple oscillations co-occur, with the slower oscillation generally driving local, faster oscillations (1). A similar multiple-time-scale neurohemal code may plausibly be shared by other assemblies of hypothalamic parvocellular neurons. Notably, the ME is capable of delivering hormone changes over a wide-range of timescales, from ultradian to seasonal (22, 23). This neurohemal structure may thus provide a model system for investigating how parvocellular outputs are translated into slowly-evolving endocrine outcomes.

MATERIALS AND METHODS

Detailed methods are provided in SI Materials and Methods. Briefly, carbon fiber microelectrodes were fabricated using a single 30 µm thread of carbon fiber, coated in Nation and connected to a gold-plated pin. C57BL6 female mice were stereotaxically implanted with carbon fiber microelectrodes at the level of the median eminence (stereotaxic coordinates relative to Bregma -1.3 mm rostro-caudal; 0 mm medio-lateral; 6.1 mm ventral). After recovery, mice were transferred to recording cages, connected to an electrical sleeve to allow free movement, and carbon fibers were held at 700 mV throughout the recording to detect secretion of DA. Repeated tail blood microsampling was performed to measure blood PRL levels, using a home-made ELISA. All statistical analysis was performed with R software.

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