



Macroscale intrinsic network architecture of the hypothalamus

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Control of multiple life-critical physiological and behavioral functions requires the hypothalamus. Here, we provide a comprehensive description and rigorous analysis of mammalian intrahypothalamic network architecture. To achieve this at the gray matter region (macroscale) level, macroscale connection (macroconnection) data for the rat hypothalamus were extracted from the primary literature. The dataset indicated the existence of 7,982 (of 16,770 possible) intrahypothalamic macroconnections. Network analysis revealed that the intrahypothalamic macroconnection network (its macroscale subconnectome) is divided into two identical top-level subsystems (or subnetworks), each composed of two nested second-level subsystems. At the top-level, this suggests a deeply integrated network; however, regional grouping of the two second-level subsystems suggested a partial separation between control of physiological functions and behavioral functions. Furthermore, inclusion of four candidate hubs (dominant network nodes) in the second-level subsystem that is associated prominently with physiological control suggests network primacy with respect to this function. In addition, comparison of network analysis with expression of gene markers associated with inhibitory (GAD65) and excitatory (VGLUT2) neurotransmission revealed a significant positive correlation between measures of network centrality (dominance) and the inhibitory marker. We discuss these results in relation to previous understandings of hypothalamic organization and provide, and selectively interrogate, an updated hypothalamus structure–function network model to encourage future hypothesis-driven investigations of identified hypothalamic subsystems.

hypothalamus | mammal | neuronal connections | neurome | neuroinformatics

The renowned theoretical physicist Richard P. Feynman, whose birth centenary was in 2018, also explored far afield (1) and valued modeling problems to gain understanding—a philosophy alluded to in two statements he wrote on his office chalkboard that were present there on his last day of life (Fig. 1A): “What I cannot create, I do not understand,” and “Know how to solve every problem that has been solved.” The first statement supports using models to gain understanding; the second recognizes the value to future progress in understanding how solutions (leading to models) are arrived at. In biology, these tenets are exemplified by the discovery of the structure of DNA (2), with empirical evidence from X-ray crystallography experiments (3) (Fig. 1B) leading to a structural model (Fig. 1C). Evidence-based models are also a mainstay of systems neuroscience, in which the system being modeled—the nervous system—is generally considered to be the most complex biological system (4). A central goal, supported by models, is to understand the connective organization of the nervous system both intrinsically (its connectome) (5, 6) and in relation to the body (the neurome; Fig. 1E) (6, 7) at different scales of granularity. These scales range from gray matter region (macroscale), to neuron type (mesoscale), to single neuron (microscale), to synaptic (nanoscale) (6). Macroscale neuronal connection data are typically obtained from pathway-tracing experiments (Fig. 1D).

Using data-driven approaches, we recently investigated the macroscale network of the cerebral hemispheres (8) and their principle parts: the cerebral cortex (9) and cerebral nuclei (10). This led to novel network models for two of the four major divisions of the forebrain that play an essential role in the cognitive control of behavior. However, cerebral hemisphere function also requires ancillary neuronal networks that enable cognitively directed motor actions to occur in concert with sensory cues and behavioral state. Prominent supporting networks include those in the thalamus and hypothalamus (the two other main divisions of the forebrain). The thalamus plays a major role in supporting cognition by processing sensory information en route to the cerebral cortex, whereas the hypothalamus is vital for subcognitive control of fundamental physiological processes and survival behaviors (11).

Experimental evidence acquired over more than a century has established the necessity of the hypothalamus for the control of those behavioral and physiological functions of the body that are essential for survival and reproduction (for review, see refs. 12–14).

Significance

Control of multiple life-critical physiological and fundamental behavioral functions requires the hypothalamus. Here, we provide a comprehensive description and analysis of mammalian intrahypothalamic network organization at the level of gray matter regions (macroscale). Network analysis revealed deep top-level network integration, but regional organization of two second-level subsystems (or subnetworks) indicated partial separation between control of physiological functions and behavioral functions; furthermore, inclusion of dominant network nodes in the subnetwork associated prominently with physiological functions suggests network primacy for physiological control. Comparing network organization to inhibitory (GAD65) and excitatory (VGLUT2) neurotransmission-associated gene markers revealed a significant positive correlation between network centrality (dominance) and the inhibitory marker. We introduce a novel intrahypothalamic network model to guide future hypothesis-driven investigations into hypothalamic structure and function.

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Data deposition: An implementation of the network analysis method of multiresolution consensus clustering that was used in the present work is available at <https://github.com/Leueb/HierarchicalConsensus>. All connection reports used for this study are provided as supporting information (SI) in spreadsheet format (Microsoft Excel) in *SI Appendix*. The connection reports (including associated methodological information) are also deposited as a searchable resource at The Neurome Project (<http://www.neuromeproject.org>).

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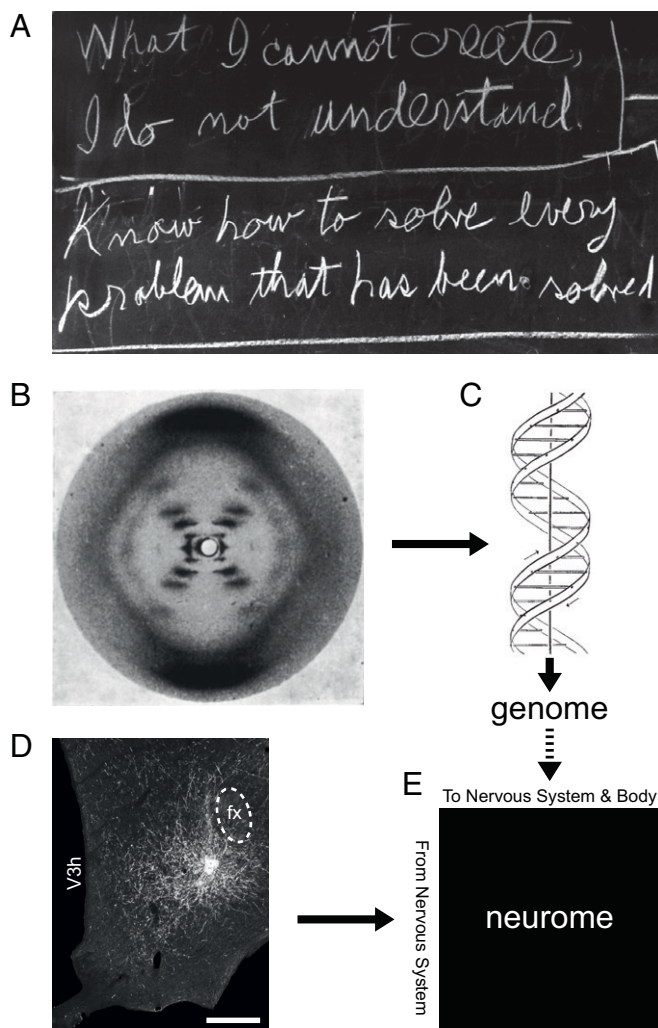


Fig. 1. Empirical models advance understanding. (A) Interrelated statements by Richard Feynman conveying how knowledge-based modeling can advance scientific understanding. (B) The X-ray diffraction pattern of DNA obtained by Rosalind Franklin (B) was instrumental for determining the double helical structure of DNA by James Watson and Francis Crick (C), enabling genome determination. (D) Visualized injection site of the anterograde neuronal pathway tracer *Phaseolus vulgaris*-leucoagglutinin in the rat hypothalamus. fx, fornix; V3h, third ventricle, hypothalamic part. (Scale bar: 250 μm .) (E) Data obtained from pathway-tracing experiments can be used to construct a network model for the complete nervous system that describes connections between all parts of the nervous system and between the nervous system and the rest of the body—a neurome (7); genome structure is a fundamental determinant of neurome structure (dashed arrow). (A) Image courtesy of the Archives, California Institute of Technology. (B) Reproduced by permission from ref. 3, Springer Nature: *Nature*, copyright (1953). (C) Reproduced by permission from ref. 2, Springer Nature: *Nature*, copyright (1953). (D) Reproduced from ref. 18.

Hypothalamic involvement in this control is diverse and includes all three nervous system divisions for motor output: neuroendocrine, autonomic, and somatomotor. This is illustrated by (i) hypothalamic neuroendocrine control of the pituitary gland, (ii) autonomic control of the cardiovascular system and abdominal viscera via preautonomic (sympathetic and parasympathetic preganglionic) connections, and (iii) (via other polysynaptic connections) hypothalamic somatomotor control of motivated behaviors that are fundamental to survival (defensive and aggressive, ingestive, reproductive, and exploratory behaviors) (11, 12).

Hypothalamic functional diversity reflects its structural and connective complexity, from embryonic development to adult. A systematic study of mammalian brain development in the mid-1990s concluded that embryological differentiation of the rat hypothalamus was “an unusually complex, little understood process” (15). More recent investigations have advanced our understanding of the underlying genetics, but (as noted in a recent review), “the development of the hypothalamus remains poorly understood, with large and obvious gaps in the literature at every developmental stage” (16). Hypothalamic cytoarchitecture is highly differentiated, generally more so than other central nervous system (CNS) divisions (17). Illustrative of hypothalamic connective complexity, one of its major subdivisions—the lateral hypothalamic area (LHA)—contains the most highly connected CNS regions identified to date [in terms of macroscale connections (macroconnections)] (18–20).

Challenges notwithstanding, some general organizing principles have emerged. A classical cytoarchitecturally based structural description of the hypothalamus divides it into three longitudinal zones: periventricular, medial, and lateral (21); and four transverse rostral-to-caudal levels: preoptic, supraoptic (latterly referred to as anterior), tuberal, and mammillary (22) (for review, see refs. 12 and 13). Developmental analysis to some extent supports an outside-in sequence with respect to the differentiation of the three longitudinal zones (lateral to periventricular), and differentiation of the four rostral-to-caudal levels is understood primarily in relation to adjacent structures (15).

Synthesis of structural and functional data has engendered various models of hypothalamic participation in the control of different fundamental behaviors, including ingestive (23), defensive (24), reproductive (25), and exploratory (26). However, despite persistent efforts, these models remain quite rudimentary, partly due to a lack of basic data and partly due to fragmentary synthesis of the available data. Nevertheless, it is pertinent to mention a general model for hypothalamic involvement in patterned motor actions, which places it at the top of a motor control hierarchy that includes successive motor pattern initiators, generators, and finally motoneuron pools—with “control” being used in the sense of “a certain level of endogenous activity, (perhaps some form of ‘set-point’)” (see figure 9 and text of ref. 11). The model is supported by loss/preservation-of-function experiments and is exemplified by control of locomotion. However, general applicability is suggested, providing a basis for understanding hypothalamic organization in relation to the three categories of motor output (neuroendocrine, autonomic, and somatic) that enable and sustain diverse behavioral expression (11, 27).

Given the diverse and complex nature of the hypothalamus, a holistic model of its intrinsic connection network would provide a foundation for future hypothesis-driven investigations into how hypothalamic circuits relate not only to specific sensory, cognitive, behavioral state, and motor functions, but also to the overall function of the hypothalamus as it relates to the interdependent motivations of individual and species survival. Accordingly, here, we investigated the organizing principles of the mammalian intrahypothalamic network based on rat macroconnection data that were obtained from pathway-tracing experiments and published in the primary literature. This is complemented with comparative analysis of gene markers associated with inhibitory [glutamic acid decarboxylase 65 (GAD65)] and excitatory [vesicular glutamate transporter 2 (VGLUT2)] neurotransmission. Network analysis is based on a weighted and directed connection matrix (for all 130 hypothalamic gray matter regions; 65 on each side), which is a macroscale intrahypothalamic subconnectome, and follows a strategy we have employed previously (8–10).

Results

Analysis Framework. The analysis is based on macroconnections reported in the primary literature between all regions of the adult rat hypothalamus, including connections originating and terminating

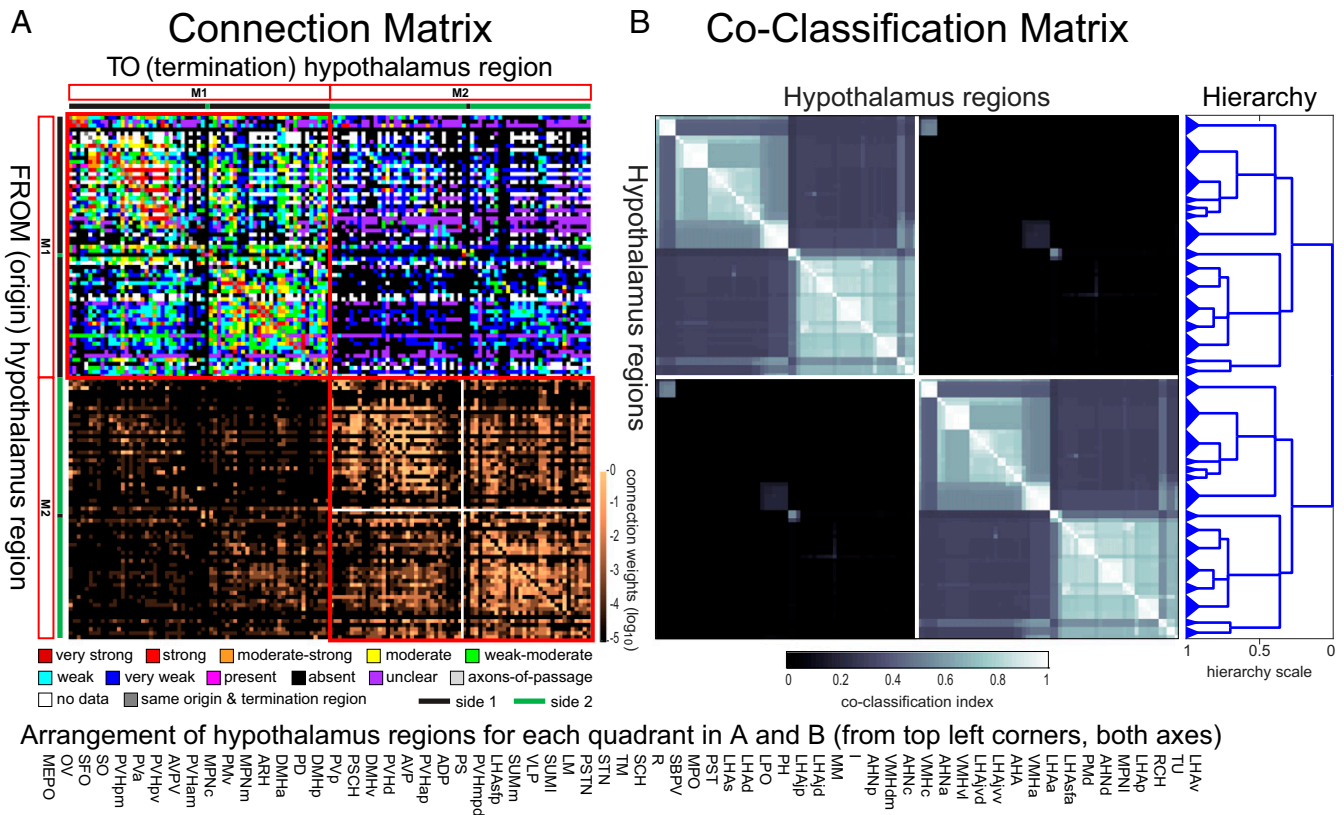


Fig. 2. (A and B, Top) Connection and coclassification network matrices for the bilateral intrahypothalamic subconnectome (HY2). (Bottom) Arrangement of hypothalamus regions for each quadrant in A and B. (A) Directed and weighted monosynaptic macroconnection matrix for the rat hypothalamus, with gray matter region sequence in a subsystem arrangement derived from MRCC analysis (shown in B). Connection weights are represented by descriptive values (Upper half, color key shown Below), and on a log₁₀ scale (Lower half, scale shown at right edge); sides 1 and 2 are indicated by green and black bars, respectively. Two bilateral top-level subsystems (M1 and M2) are outlined in red; two second-level subsystems are delineated by a white cross (shown only for M2 for clarity but applies to both M1 and M2). (B) Complete coclassification matrix obtained from MRCC (as in A) for the 130 regions (65 per side) of the hypothalamus. A linearly scaled coclassification index (shown Below) gives a range between 0 (no coclassification at any resolution) and 1 (perfect coclassification across all resolutions). Ordering and hierarchical arrangement are determined after building a hierarchy (Right) of nested solutions that recursively partition each cluster (i.e., subsystem), starting with the two top-level subsystems. The 30 subsystems obtained for the finest partition are indicated on the left edge of the dendrogram, while the two identical top-level subsystems (corresponding to M1 and M2) appear at the root of the tree (far right edge). A total of 21 distinct hierarchical levels are present, as determined by the sum of vertical cuts through each unique set of branches. The length of each distinct set of branches represents a distance between adjacent solutions in the hierarchical tree that may be interpreted as its persistence along the entire spectrum; dominant solutions extend longer branches, while fleeting or unstable solutions extend shorter branches. All solutions plotted in the tree survive the statistical significance level of $\alpha = 0.05$. Abbreviations are defined in Dataset S2.

hierarchy of nested partitions with 30 bottom-level subsystems (15 on each side of the brain) (Fig. 2B). For HY1, the coclassification matrix comprises a 12-level hierarchy of nested partitions with 17 bottom-level subsystems (SI Appendix, Fig. S2B).

A flatmap representation (Fig. 3) facilitates comparison of the top-level modules of HY1 and HY2 (and the latter's two second-level subsystems). A first point to note is that the bilateral structure of each top-level HY2 module is the result of the predominantly crossed connections of a single outlying region—the retina. Second, it is apparent when comparing HY1 to HY2 that regions in HY1 module (M)1 or M3 assign respectively and exclusively to HY2 M1/2 second-level subsystems 1.1 (HY1 M1) or 1.2 (HY1 M3), whereas regions in HY1 M2 assign to one or the other of these HY2 subsystems (Fig. 3). Considering region number/module for HY1, M1 and M3 are the largest and each contains a similar number of regions (27 for HY1 M1 and 29 for HY1 M3) compared with nine regions for HY1 M2. Given that regions in HY1 M1 and HY1 M3 assign exclusively to either subsystem 1.1 or 1.2 of HY2 M1/2, while those in HY1 M2 split between these HY2 subsystems, the number of regions in each is also similar (33 for HY2 M1/2 subsystem 1.1, and 32 for HY2 M1/2 subsystem 1.2). Third, a medial–lateral topological division

is apparent: regions in HY1 M1 and HY2 M1/2 subsystem 1.1 are mostly medial, whereas regions in HY1 M3 and HY2 M1/2 subsystem 1.2 are mostly lateral; by comparison, longitudinal axis region subsystem assignment is evenly distributed (Fig. 3). However, the nine regions of HY1 M2 are an exception to this general pattern (they are mostly caudal and include both medial and lateral components).

Centrality Metrics and Additional Network Attributes. In addition to MRCC analysis, we investigated hypothalamic network properties using four common network centrality metrics (measures that indicate the dominance/“importance” of each node in the network): degree, strength, betweenness, and closeness (8–10). The centrality metric of degree measures the number of input (in-degree) or output (out-degree) connections for each network node (here, each gray matter region); strength represents the total weight of each node's macroconnections; and the related centrality measures of betweenness and closeness take account of the shortest path between nodes and are considered to provide an indication of node centrality with respect to information flow.

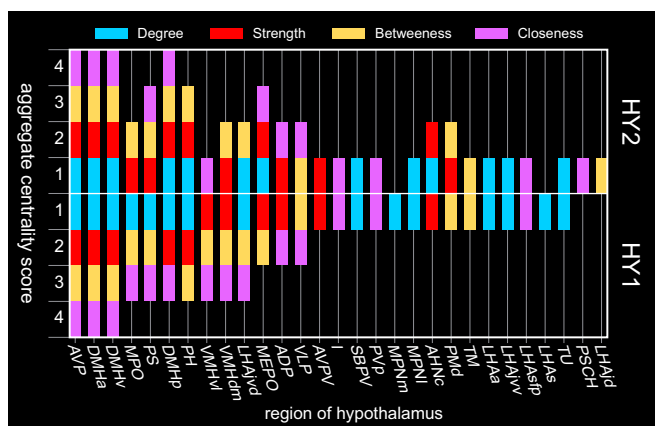


Fig. 4. Central nodes of the intrahypothalamic network. Identification of candidate hub regions (and others with high network centrality) for the bilateral (HY2) and unilateral (HY1) hypothalamic subconnectomes. Regions are assigned a score of 0 to 4 according to the number of times they fall within the top 20th percentile for each of four measures of centrality (degree, strength, betweenness, and closeness) and are arranged from left to right by HY1 descending aggregate centrality and topographically (17). Regions with a centrality score of 4 are considered candidate hubs. For individual-region centrality values for each measure of centrality (for HY2), see *SI Appendix, Fig. S3*. Note that aggregate centrality scores are modulated between HY1 and HY2, indicative of the relevance of HY2 contralateral connections to the overall structure of the network. Abbreviations are defined in *Dataset S2*.

length: 240.0 ± 2.9 and 241.4 ± 5.1]. These metrics are only weakly indicative of small-world organization, mainly due to only modest levels of clustering relative to the null model. With respect to rich-club attributes, neither HY1 nor HY2 was found to contain a densely connected subgraph of hubs. Analysis of the HY1 network revealed a 48-node subgraph with significantly greater density compared with 1,000 randomized networks; however, the excess in density (over the null model) was less than 7% and the subgraph included almost three-fourths of the entire network, thus excluding it from consideration as a rich-club (no rich-club was found in HY2).

Comparison of Network Centrality with Markers of Excitatory and Inhibitory Neurotransmission. A critical determinant of neuronal network function is whether it is inhibitory or excitatory. To investigate this property, we mapped gene markers associated with inhibitory (GAD65) (30), and excitatory (VGLUT2) (31) synaptic neurotransmission. The levels of mRNA for GAD65 and VGLUT2 were analyzed for 64 of 65 hypothalamic regions—the retina was not included, but it was previously reported that its output neurons express VGLUT2 (32).

The results indicate that GAD65 and VGLUT2 are both highly and heterogeneously expressed in the hypothalamus, consistent with and confirmatory to previous work (33, 34). GAD65 and/or VGLUT2 mRNA was detected in all regions, and most regions (81.5%, 53/65) express both markers (*Dataset S3*). However, one region expressed GAD65, but not VGLUT2 (i.e., anterior hypothalamic nucleus dorsal region), and six regions (seven if including the retina) expressed VGLUT2, but not GAD65 [i.e., supraoptic nucleus, PMd, medial mammillary nucleus, lateral mammillary nucleus (LM), PSTN, and STN]. Moreover, in regions that expressed both GAD65 and VGLUT2, there was considerable inter- and intraregion variation in expression levels, and equal expression levels of both markers in the same region was rare (nine regions, 13.8%). Nevertheless, aggregate regional expression levels for the hypothalamus were approximately equal [expression levels of GAD65 were just 2%

higher than those of VGLUT2 (*Dataset S3*)—a slim margin further reduced by inclusion of the retina].

To relate expression of GAD65 and VGLUT2 to the results of the network analysis, their regional expression levels were compared with the aggregate centrality scores for each hypothalamic region in HY2 (Fig. 5). This comparative analysis revealed a significant positive correlation between GAD65 and all four measures of centrality (degree: $\rho = 0.256$, $P = 0.04$; strength: $\rho = 0.317$, $P = 0.01$; betweenness: $\rho = 0.343$, $P = 0.005$; and closeness: $\rho = 0.400$, $P = 0.001$; Spearman’s rank order correlation). In contrast, only closeness presented a significant (negative) correlation with VGLUT2 ($\rho = -0.276$, $P = 0.02$). (*SI Appendix, Fig. S4*).

Discussion

Analysis of macroconnection data extracted from the primary literature for the rat intrahypothalamic subconnectome (HY2) indicated the existence of 7,982 of 16,770 possible connections (from 87.7% data coverage), a connection density of 47.6%. Comparable analysis recently applied to the endbrain (EB) and its principle divisions—cerebral cortex (CTX) and cerebral nuclei (CNU)—indicated connection densities of 17.9% (EB), 22.8% (CNU), and 24.5% (CTX) (8). Evidently, HY2 is markedly more connection dense than the intrinsic networks (subconnectomes) for these other divisions of the forebrain. Connection density differences for the component ipsilateral and contralateral subconnectomes are even greater. For example, the connection density of the ipsilateral intra-CTX subconnectome is 37.7% compared with 62.6% for the hypothalamus (a 66% increase); for the contralateral subconnectome, it is 10.2% for the CTX compared with 32.5% for the hypothalamus (a 219% increase) (8). The comparatively high hypothalamic connection density accounts for the marginal expression of the small-world network attribute (characterized by simultaneous high clustering and short path length) compared with relatively robust small-worldness exhibited by EB2, CTX2, and CNU2 (*SI Appendix, Fig. S5*). This organization may reflect reduced wiring cost afforded by the greater spatial compactness of hypothalamic regions compared with CNU and CTX regions (17). Similar considerations may underlie the absence of intrahypothalamic rich-club expression (a feature of networks whose most highly connected nodes are highly interconnected).

Comparing the current MRCC analysis to an earlier-alluded-to hypothalamic structure–function model (11, 35) facilitates exploration of possible functional interpretations. To elaborate, the earlier model identifies subgroupings based on structural and functional properties, giving five divisions: (i) a neuroendocrine motor zone associated with pituitary gland control; (ii) medial zone nuclei forming part of a putative behavior control column; (iii) a highly interconnected group of five rostrally located regions and the three regions of the DMH that together are considered to form a theoretical “visceromotor network” (35) for generating patterned autonomic and neuroendocrine motor output; (iv) a periventricular region related to (and possibly expanding) the visceromotor network, defined essentially by what remains of the periventricular and medial zones after subtracting the neuroendocrine motor zone, medial zone nuclei, and visceromotor network; and (v) a lateral zone associated prominently with the behavior control column, which is supported by more recent data (18–20, 36).

The arrangement of regions in the two second-level HY2 subsystems largely follows the five divisions of the earlier model, with HY2 M1/2 subsystem 1.1 including (and mostly composed of) all regions of the neuroendocrine motor zone and the visceromotor network, and with HY2 M1/2 subsystem 1.2 mostly composed of medial zone nuclei (behavior control column), the periventricular region, and the lateral zone (Fig. 6). This grouping of regions supports an updated structure–function model for the hypothalamus comprising two longitudinal divisions: one that is

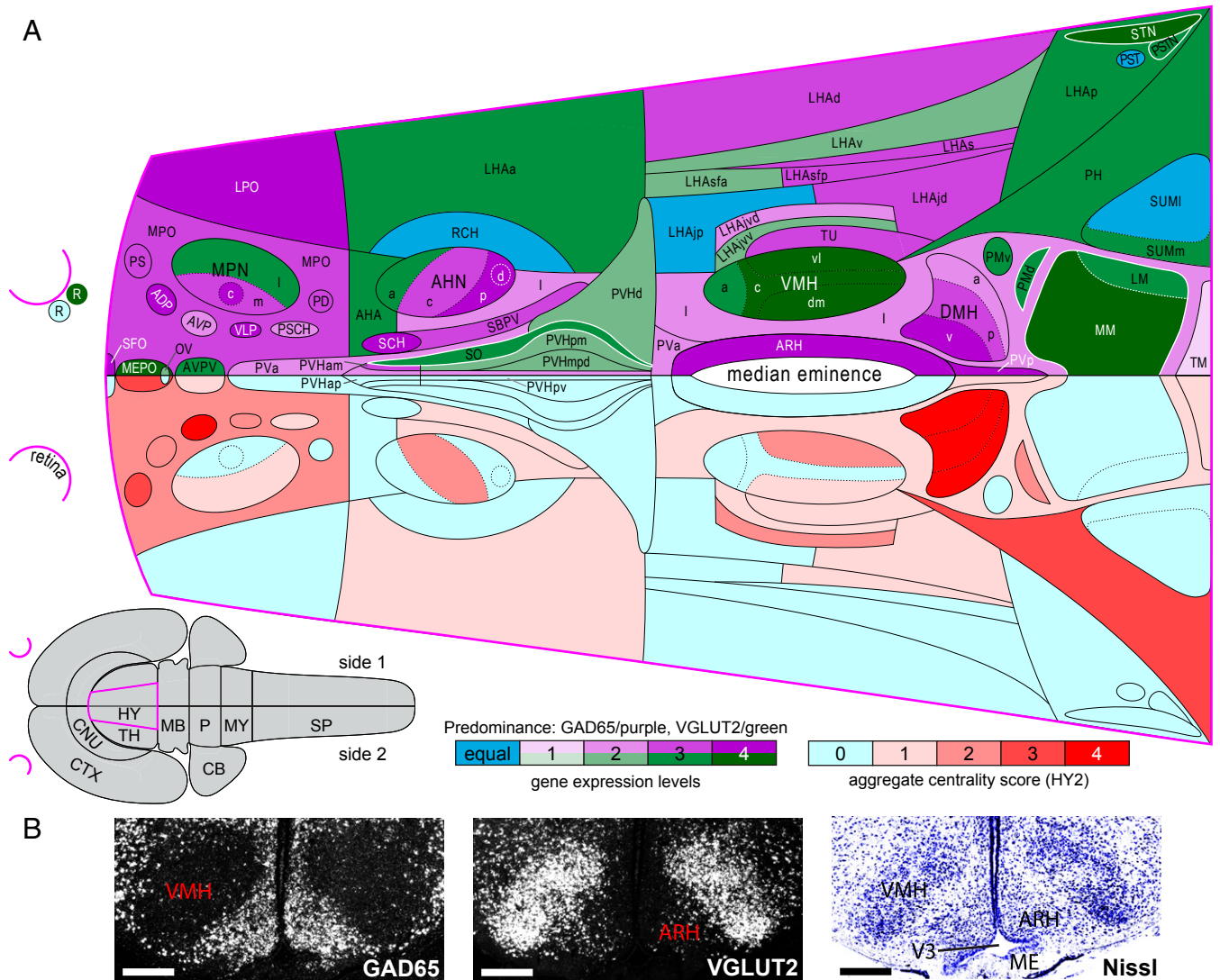


Fig. 5. Comparison of intrahypothalamic network centrality with GAD65 and VGLUT2 mRNA expression. (A, Lower half of flatmap) Regional distribution of aggregate centrality measures for the bilateral hypothalamus macroscale subconnectome (HY2) (color key shown *Below Right*). (A, Upper half of flatmap) mRNA expression levels of GAD65 and VGLUT2 (color key shown *Below Middle*). For comparison, each bilateral dataset is represented on one side of a rat hypothalamus flatmap. The hypothalamus (magenta) and its spatial relation to the CNS is represented on the gray flatmap at *Lower Left*. A five-point index is used for both centrality and mRNA expression levels: for centrality, regions are assigned a score of 0 to 4 according to the number of times they fall within the top 20th percentile for each of four measures of centrality (degree, strength, betweenness, and closeness); for gene expression, regions are assigned a score of 1 to 4 according to binned data for their average expression levels ranging from 0 to 7 (1, 0 to 1.75; 2, >1.75 to 3.5; 3, >3.5 to 5.25; and 4, >5.25 to 7). Flatmap coloration for aggregate centrality follows the corresponding color scale (candidate hubs have a score of 4 and are bright red). Flatmap coloration for mRNA expression indicates region predominance (GAD65/purple, VGLUT2/green) and expression level (1 to 4); regions with equal expression levels of both genes are blue (these regions had a binned score of 2); regions outlined in white express either GAD65 or VGLUT2. (B) Representative darkfield photomicrographs of mRNA expression for GAD65 (*Left*) and VGLUT2 (*Middle*) at a midrostrocaudal level of the VMH nucleus (shown in corresponding Nissl-stained section image, *Right*). Note the inverse relationship between GAD65 and VGLUT in the VMH and ARH. (Scale bars: 500 μ m.) ME, median eminence; V3, third ventricle. Abbreviations for the hypothalamus flatmap are defined in [Dataset S2](#) and those for the gray flatmap are defined in [Fig. 3](#). Maps adapted from [ref. 17](#).

mostly periventricular and associated prominently with physiological control, and the other mostly medial–lateral and associated prominently with behavioral control ([Fig. 7](#)). The suggested divisions are supported further by inclusion in HY2 M1/2 subsystem 1.1 of neuroendocrine-related hypothalamic circumventricular organs (organum vasculosum of the lamina terminalis and subfornical organ) as well as four candidate hub regions that all are members of the visceromotor network: AVP, DMHa, DMHp, and DMHv. However, the existence of only two bilateral top-level modules for HY2 underscores integration of the complete intrahypothalamic network—congruent with the functional understanding that behavioral and physiological homeostatic processes are

mutually supportive and that both support the overarching function of the hypothalamus to support survival and reproduction ([11](#)) ([Fig. 7](#)).

Whichever level of network resolution is interrogated, the general comparative (and hypothesis-generating) approach is to consider how a novel network structure compares to existing models. We have followed this approach at a high level (above discussion) and, in the remaining discussion, consider selected examples at lower nested levels of the network hierarchy.

In the earlier model (see [figure 12](#) of [ref. 11](#)), regions of the behavior control column division are subdivided by structural connectivity and by association with different categories of

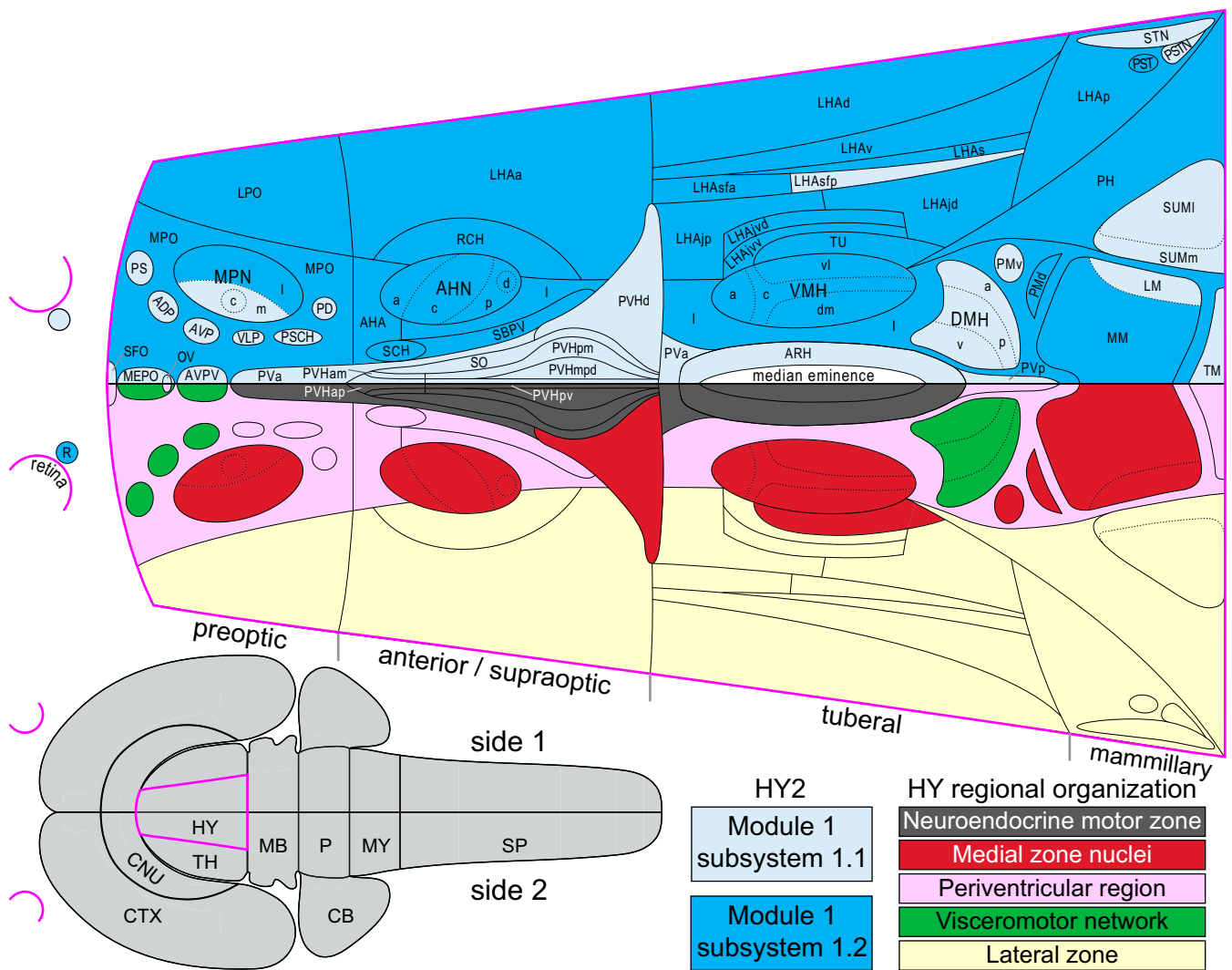


Fig. 6. Comparison of current (network) and previous (cell group-based) models of hypothalamic organization. An established cell group-based model of hypothalamic organization (*Lower* half of flatmap) divides it into four transverse levels (preoptic, anterior/supraoptic, tuberal, mammillary) and three longitudinal zones (periventricular, medial, lateral). Refinements to this model include the identification of a neuroendocrine motor zone (gray), medial zone nuclei (red) considered to form the rostral end of a behavior control column, and a periventricular region (pink) containing a putative visceromotor network (green), leaving the remaining lateral zone (yellow). The *Upper* half of the flatmap shows the two top-level partitions of the bilateral hypothalamus macroscale subconnectome (HY2). For comparison, each organization schema is presented on a single side. Most HY2 subsystem 1.1 regions (lighter blue) are in the periventricular zone and include all regions of the neuroendocrine motor zone and a putative visceromotor network, whereas most HY2 subsystem 1.2 regions (darker blue) are in the lateral and medial zones. The hypothalamus (magenta) and its spatial relation to the CNS is represented on the gray flatmap at *Lower Left*.

goal-directed behavior. For example, reproductive behavior involves the medial preoptic nucleus (MPN) lateral part, the ventromedial hypothalamic nucleus (VMH) ventrolateral part, and the PMv; defensive behavior involves the anterior hypothalamic nucleus, the VMH dorsomedial part, and the PMd. More recent evidence indicates defensive behavior may also involve the VMH central part (24) and the LHAjd (37). With one exception (PMv), all these regions are in HY2 M1/2 subsystem 1.2, consistent with an updated model (Fig. 7).

Additional comparison with lower-level HY2 subsystems reveals novel associations. For example, the PMv previously appeared to be an outlier, but is grouped in a third-level subsystem that includes several other regions associated with reproductive function, including the two other MPN divisions (central and lateral) and especially the anteroventral periventricular nucleus and ARH involved in critical neuroendocrine reproductive control (12). In fact, while the PMv is associated strongly with both neuroendocrine and somatomotor control

(12, 24, 27), the current model emphasizes the former. This illustrates further the highly integrated intrahypothalamic network structure and also suggests that theoretical functional interpretation of an MRCC nested network hierarchy is informed by its consensus basis, analogous to a multilayered functional heatmap. With respect to the third-level PMv-included HY2 subsystem, it also suggests that other regions in this subsystem not clearly associated with reproductive function merit reevaluation. This strategy could also be applied to other regions and subsystems, such as the DMH candidate hub that is indicated to integrate behavioral and behavioral-state control (Fig. 7) (38).

As a second example, note that HY1 has six top-level modules (i.e., M1 to M3, each side) and that one of these modules, HY1 M2, splits between HY2 M1/2 subsystems 1.1 and 1.2 (unlike HY1 M1 and HY1 M3 that transpose directly to HY2 M1/2 subsystems 1.1 and 1.2, respectively) (Figs. 3 and 7). This distinguishes HY1 M2 from HY1 M1 and HY1 M3, and HY1 M2 is further distinguished by having only nine regions compared with

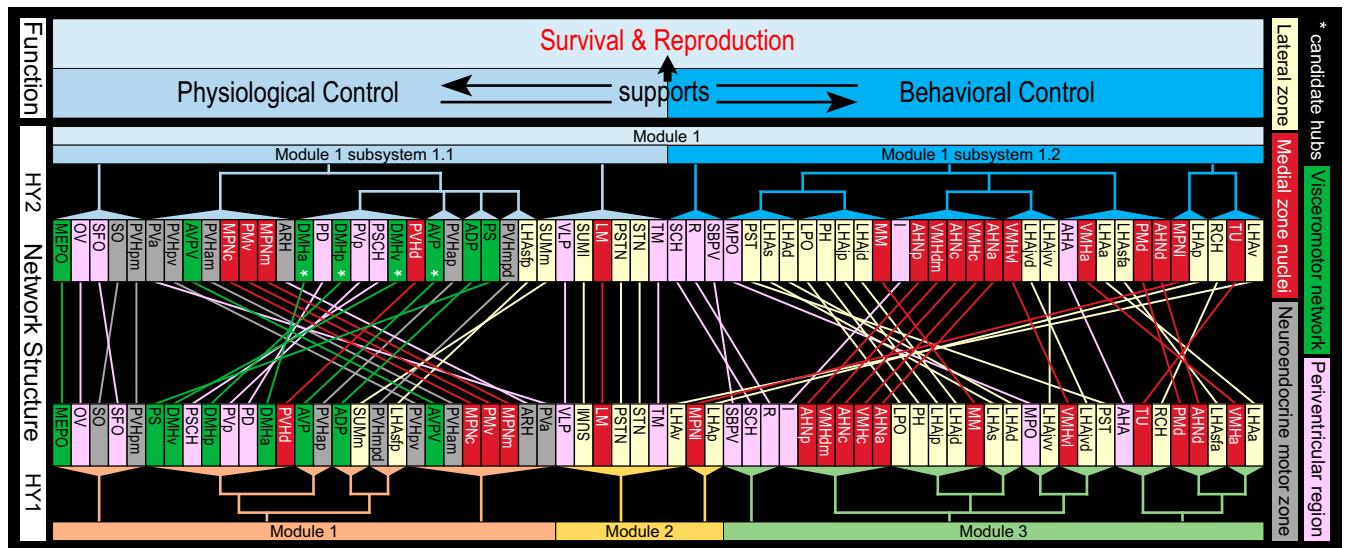


Fig. 7. Hypothalamus structure–function network model. Network structures emerging from MRCC analysis of the bilateral (HY2) and unilateral (HY1) hypothalamus macroscale subconnectomes (colored dendrograms) are compared with one another and with an earlier model of hypothalamic cell group and network organization (11, 35). Colored lines connecting regions for HY1 and HY2 show correspondence between subsystem assignment determined by MRCC. Two-thirds of HY2 M1/2 subsystem 1.1 regions are in the periventricular zone, which includes the periventricular region (pink); included in this partition are all regions of a putative hypothalamic visceromotor pattern generator network (green), all regions of the neuroendocrine motor zone (gray), and both hypothalamic circumventricular organs [organum vasculosum of the lamina terminalis (OV) and subfornical organ (SFO)]; in contrast, four-fifths of HY2 M1/2 subsystem 1.2 regions are in the medial (dark red) and lateral (light yellow) hypothalamic zones. Overall, this organization suggests a prominent functional association for HY2 M1/2 subsystem 1.1 with physiological control (especially neuroendocrine signaling), and for subsystem 1.2 with behavioral control (especially somatomotor signaling). Moreover, primacy of physiological control is suggested by inclusion of all four candidate hubs in HY2 M1/2 subsystem 1.1 (white asterisks). However, both HY2 top-level subsystems 1.1 and 1.2 include regions involved in autonomic and behavioral-state control, and the existence of only two bilateral HY2 top-level modules underscores deep intrahypothalamic integration. Communication between HY2 M1/2 subsystems 1.1 and 1.2 may be considered mutually supportive, and both support the prime function of the hypothalamus to support survival and sexual reproduction. Abbreviations are defined in [Dataset S2](#).

27 and 29 for HY1 M1 and HY1 M3, respectively. At the second hierarchical level, HY1 M2 has two subsystems of seven and two regions. All but one region in the seven-region subsystem are in HY2 M1/2 subsystem 1.1, with the remaining region and the two-region subsystem in HY2 M1/2 subsystem 1.2 (Fig. 7). At face value, this indicates that HY1 M2 is less robust than either HY1 M1 or HY1 M3. Given also the suggested general differentiation between physiological and behavioral control for HY2 M1/2 subsystems 1.1 and 1.2, it is salient to consider functional associations for the HY1 M2 regions that assign differently to these HY2 subsystems. We consider here the six regions of HY1 M2 that are in HY2 M1/2 subsystem 1.1: VLP, supramammillary nucleus (SUM) lateral part (SUMl), LM, TM, PSTN, and STN. In addition, in HY2, these six regions are in a partition that includes a seventh region: the SUM medial part (SUMm) (Fig. 7).

The VLP and TM are implicated (oppositely) in behavioral-state control (VLP in sleep state; TM in awake state) (39). The LM has an established role in processing directional heading information during locomotion (40). The SUMm and SUMl are also associated with locomotion via their generative role for the hippocampal theta rhythm (41) present during locomotion (42); further SUM association is with REM sleep, which is also positively correlated to the hippocampal theta rhythm (43). The PSTN is indicated to play a broad role that includes coordination of parasympathetic responses associated with cardiovascular function and ingestive behavior, and may also influence central relay of sensory information relating to the latter (44, 45). Lastly, the STN is implicated in somatomotor control, notably of orofacial and limb movements (46). With respect to limb movements, STN lesion is associated with the rare movement-disorder ballism, characterized by uncontrollable “throwing” of the limbs (47). The STN has also received attention as a therapeutic target site for other movement disorders, including Parkinson’s disease

(47, 48). In sum, these regions are associated with goal-directed behavior, locomotion necessary to obtain a goal, and behavioral state that determines when these are active (broad functions germane to multiple specific behaviors). Given indicated STN involvement in several diseases affecting locomotor control and its use as a therapeutic target, correlative investigation of other regions in the same HY2 third-level subsystem appears relevant.

Our analysis of the macroscale intrinsic network architecture of the hypothalamus reveals structure–function relations that tend to increase in specificity with increasing network resolution, but the whole network is richly integrated. More generally, we have demonstrated how data-driven network modeling approaches can be employed as hypothesis-generating tools, with selected examples provided by interrogation of an updated model of the intrahypothalamic network, and we hope this encourages further investigation of the multiple intrahypothalamic subnetworks described here. Thus far, we have investigated macroscale subconnectomes for the cerebral hemispheres (8–10) and for the hypothalamus (this study). Future investigations will be aided by a more comprehensive understanding of the network architecture of the nervous system.

Materials and Methods

Network Analysis. All network analysis methods used here follow those described previously (8–10), including a recently introduced method for MRCC analysis (8, 29). All macroconnection data obtained from the primary literature were interpreted in relation to the current version of the only available standard, hierarchically organized, annotated parcellation and nomenclature for the rat brain (17). Within- and between-sides connection reports were assigned ranked qualitative connection weights (reported values) according to their description; an ordinal scale [1 (very weak) to 7 (very strong)] was used. Connection report data and annotations are provided in a Microsoft Excel spreadsheet ([Dataset S1](#)), as are the data extracted from these reports to construct connection matrices ([Dataset S2](#)). To facilitate access to, and use of,

the connection-report data, they are freely available as a searchable resource at The Neurome Project (<http://www.neuromeproject.org>). For weighted network analysis, an exponential scale was applied to the ordinal weight categories. As in previous work (8–10), the scale spanned 4 orders of magnitude and is consistent with quantitative pathway-tracing data in rats (7). Network analyses were carried out on the directed and log-weighted rat intrahypothalamic macroconnection matrix (Dataset S2, worksheet “HY topographic bins”) using tools collected in the Brain Connectivity Toolbox (www.brain-connectivity-toolbox.net/).

In Situ Hybridization for GAD65 and VGLUT2. Methods and controls for isotopic in situ hybridization for detection of GAD65 and VGLUT2 mRNA were fully described previously (49). Tissue sections were obtained from an adult male Sprague-Dawley rat (all procedures were approved by the Institutional Animal Care and Use Committee at the University of Southern California). cDNA probes used to generate ³⁵S-UTP-labeled cRNA probes for in situ hybridization were obtained from the following sources: GAD65 from M. G. Erlander, University of California, Los Angeles, CA (50), and VGLUT2 from D. R. Ziegler, University of Pikeville, Pikeville, KY (34). For analysis of VGLUT2 and GAD65 mRNA expression, an ordinal value ranging from 1 (very weak) to 7 (very strong) was recorded that qualitatively reflected signal strength (silver grains visible under darkfield microscopic illumination) relative to the overall range

of signal observed independently for each gene marker (absence of signal was denoted by a 0). Data acquisition was aided with the use of a specialized Microsoft Excel template (Axiome C, created by J.D.H.) that facilitated data entry across multiple levels for each hypothalamic gray matter region as described in a rat brain reference atlas (17). Accordingly (excepting the retina), data were acquired for all hypothalamic regions across 22 transverse levels of the hypothalamus, with individual values recorded for each region at atlas-level resolution (average of four atlas levels per region) (Dataset S3). To negate the effects of interanimal variability, series of sections from the same rat brain were analyzed for each mRNA sequence and matched with an adjacent series of Nissl-stained sections to enable data transposition to the reference atlas. Analysis encompassed expression of markers on either side of the brain (no appreciable difference in signal between sides was observed for any region).

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