Neurons in the Rat Lateral Hypothalamic Area Integrate Information from the Gastric Vagal Nerves and the Cerebellar Interpositus Nucleus

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\textbf{Key Words}
Gastric vagal nerves \cdot Cerebellar interpositus nucleus \cdot Lateral hypothalamic area \cdot Glycemia-sensitive neuron \cdot Cerebellohypothalamic projections \cdot Feeding control

\textbf{Abstract}
Previous investigations have demonstrated that the neuronal activity in the lateral hypothalamic area (LHA) is respectively modulated by afferent inputs from the gastric vagal nerves innervating the upper gastrointestinal tract, as well as the cerebellar interpositus nucleus (IN). The aim of this study was to examine whether the gastric vagal and cerebellar IN inputs converge onto single LHA neurons in rats, especially those sensitive to glycemia. Of the 114 LHA neurons recorded, 60 (52.6\%) and 51 (44.7\%) responded to gastric vagal and cerebellar IN stimulation, respectively. Of the 60 LHA neurons responsive to gastric vagal stimulation, 30 also responded to the cerebellar IN stimulus, indicating a convergence of gastric vagal and cerebellar inputs onto single hypothalamic cells. When the gastric vagal nerves and cerebellar IN were stimulated simultaneously, a summation of the responses was observed in all 6 neurons tested. Moreover, of 24 neurons that responded to both the gastric vagal and cerebellar IN stimuli, 15 (62.5\%) were identified as glycemia-sensitive. These results demonstrate that the visceral information transmitted by the gastric vagal nerves and the somatic information forwarded by the cerebellar IN converge onto single LHA neurons, especially those sensitive to glycemia. The findings also suggest that integration of somatic-visceral responses related to short-term feeding regulation may take place in the LHA.

\textbf{Introduction}

The hypothalamus is a central area where many peripheral signals and neural pathways that relate to autonomic functions and motivated behaviors converge \cite{1, 2}. The lateral hypothalamic area (LHA) is closely associated with the initiation and regulation of feeding \cite{3–8}, and it is well known that neurons in the LHA can sense the blood glucose concentration \cite{9, 10}. A fall in blood glucose may induce hunger and initiate feeding \cite{9, 11}, and a growing body of evidence suggests that those neurons sensitive to glycemia integrate food and water intake, various forms of social behavior and neuroendocrine signals involved in the regulation of homeostasis of blood glucose, feeding and other physiological functions \cite{12, 13}.
Previous electrophysiological studies in the cat have demonstrated that gastric vagal afferent inputs reach the LHA [14, 15], suggesting that meal-related visceral signals induced by food intake and digestion can be transmitted to the LHA via the gastric vagal nerves [16]. However, these studies have not indicated whether the neurons receiving the gastrointestinal signals relayed by the gastric vagal nerves are glycemia-sensitive or not. More recently, the role of the cerebellum in regulating non-somatic visceral processes, feeding behavior included, has received increasing attention [17–25]. Stimulation of the rat cerebellar fastigial nucleus is known to evoke post-synaptic responses and to modulate the discharge of LHA neurons, including the glucose-sensitive/glycemia-sensitive neurons [20, 24]. Other studies have revealed that glucose-sensitive and glucose-insensitive neurons in the LHA respond to stimulation of the cerebellar interpositus nucleus (IN) in cats [21, 22]. These results suggest that the cerebellum may forward some movement-related information to the LHA through the cerebellohypothalamic projections [26], and may be involved in the regulation of additional non-somatic functions such as food intake.

The above-mentioned findings also suggest that LHA neurons may play a role in the integration of visceral and somatic information conveyed via gastric vagal and cerebellorubal pathways, respectively. However, it is unclear if inputs from the gastric vagal nerves and cerebellar IN converge onto single LHA neurons. One of the purposes of this study is to explore this issue and to examine whether the inputs from these two sources interact with each other. Since feeding behavior is closely related to fluctuation in the blood glucose level [9, 11, 27, 28], the sensitivity of the recorded neurons to intravenous injection of glucose was also investigated. Our results demonstrate a convergence of gastric vagal and cerebellar IN afferent inputs to single LHA neurons, including cells sensitive to glycemia.

Materials and Methods

Animals and Surgery

Experiments were performed on 34 adult Sprague-Dawley rats of either sex weighing 250–300 g. The rats were anesthetized with urethane (0.8 g/kg) plus α-chloralose (65 mg/kg) given intraperitoneally. A supplemental dose of anesthetic was given usually every 5–6 h. A jugular catheter was placed for intravenous administration of supplemental anesthetic, glucose, normal saline and mannitol. Rectal temperature was maintained at 37 ± 0.5°C by keeping the animals on a heating pad. The ECG was continuously monitored on an oscilloscope. A left femoral catheter filled with normal saline containing heparin (500 IU/ml) was connected to a self-made blood pressure amplifier through a transducer (BLPR, WPI, USA). Arterial pressure was measured throughout the experimental sessions, and no obvious fluctuation was found during and after stimulation of the gastric vagal nerves or cerebellar IN [23–25].

The abdomen was opened by a midline incision and the gastric branches of the dorsal, and ventral vagal trunks were carefully dissected from the surrounding tissues to expose the nerve bundles and were then placed over a bipolar hook electrode. After closing the abdominal incision, the animal was mounted in a stereotaxic frame (1404, David Kopf Instruments, USA). The cisterna magna was opened so that there was no accumulation of cerebrospinal fluid. The scalp was incised and two small circular holes were made in the skull above the LHA and cerebellar IN. After the dura mater was removed, the exposed brain surfaces were covered with 2% Ringer agar. A concentric bipolar stainless steel electrode (ID 0.1 mm, OD 0.4 mm, tip exposure 0.2 mm) was used to stimulate the cerebellar IN. The electrode was stereotaxically placed in the nucleus at A –11.6, L 1.5–2.5 and H 4.5–5.0 according to the atlas of Paxinos and Watson [29].

Stimulation, Recording, Data Acquisition and Analysis

Double-negative rectangular pulses were used to stimulate the gastric vagal nerves (intensity 300–500 μA, duration 0.5 ms, interval 10 ms) [14, 15] and cerebellar IN (intensity 50–200 μA, duration 0.4 ms, interval 10 ms) [24, 25]. The paired pulses were applied to the gastric vagal nerves or cerebellar IN every 15 s up to 100 trials for constructing a peri-stimulus time histogram (PSTH; see below for details). Single neuronal discharges were recorded extradurally from the LHA (A –2.3 to –2.8, L 1.5–2.0, H 7.5–9.0) [29], which is contralateral to the stimulated cerebellar IN, using a glass electrode filled with 1% solution of pontamine sky blue in 0.5 M sodium acetate (DC resistance 5–10 MΩ). The LHA neuronal discharges were conventionally amplified and monitored on an oscilloscope, and simultaneously fed into a window discriminator. The standard rectangular pulses (5 V, 1.0 ms) triggered from the spikes were sent through an A/D interface (1404 Plus, CED, UK) to a laboratory computer which was used to analyze the data online by the software Spike 2 (CED, UK). If the recorded neuron responded to the gastric vagal stimulation and/or cerebellar IN stimulation, the sensitivity of the cell to intravenous glucose administration (0.4 M, 0.5 ml/kg), which modified blood glucose in small amounts to simulate normal physiological glycemic fluctuations associated with feeding [9, 10, 28], was tested further. Cells that showed a specific and significant inhibition in response to the injection of glucose but not to the administration of normal saline (0.9% NaCl, 0.5 ml/kg, used as an osmotic and non-glycemic control) and mannitol (0.4 M, 0.5 ml/kg, used as a volumetric control) were considered to be glycemia-sensitive neurons [10, 21, 22, 24, 25, 28]. In order to examine the possible convergence of gastric vagal and cerebellar IN afferent inputs on the glycemia-sensitive neurons of LHA, in some cells that responded to both the gastric vagal nerves and cerebellar IN stimulation and showed an inhibitory response to glucose administration, the effect of simultaneous stimulation of gastric vagal nerves and cerebellar IN was observed further.

PSTHs of the neuronal discharges were generated by the computer to assess the effects of gastric vagal and cerebellar IN stimulations (sampling interval = 2 ms, sampling length = 1,600 ms, accumulated in 100 trials) and the effects of glucose, saline and mannitol injections (sampling interval = 1 s, sampling length = 900 s, single trial). The response patterns of LHA neurons to the gastric vagal, cerebellar IN and glucose stimulations were determined by
comparing the discharge rate of cells recorded in the post-stimulation response window with the basal firing rate of the cell in the pre-stimulation control window. According to the previously established criteria [20, 21, 23–25, 30], inhibition due to stimulation was considered to be significant if the discharge rate in the response window decreased at least to 30% of the basal rate in the control window, while excitation was considered to be significant if the discharge rate increased to 200% of the basal rate. Then, the difference between the firing rate in the response window and the firing rate in the control window was statistically evaluated by using Student’s t test, and the difference was considered to be significant if the p values were $<0.05$. For all the cells recorded, the raw data were simultaneously fed into the computer through the A/D interface and collected as records in the form of oscilloscopic traces with a sampling rate of 10 KHz.

**Histology**

At the end of each experiment, pontamine sky blue was injected into the recording site (20 $\mu$A, 10 min) and DC current (10 $\mu$A, 20 s) was passed through the stimulation electrode to deposit iron at the site of stimulation. The brain was then removed and fixed with 10% formaldehyde containing 1% potassium ferrocyanide. A week later, frozen coronal sections (80 $\mu$m thickness) of the brain were prepared to identify the recording and stimulation sites. If the blue dots that indicated the recording and/or stimulating sites were outside the target nuclei, the data were excluded from further analyses.

**Results**

**LHA Neuronal Responses to Gastric Vagal Stimulation**

A total of 114 LHA neurons were recorded in this study, 60 of which (52.6%) responded to the stimulation of gastric vagal nerves (table 1). Of the 60 responsive neurons, 53 responded phasically (53/60, 88.3%) to the stimulation. This type of response includes a transient decrease or increase in discharge rate within a relatively constant latency (40–126 ms) following the gastric vagal stimulation (fig. 1a, b). The remaining 7 responsive neurons (7/60, 11.7%) showed a change in their firing pattern to the stimulation (fig. 1c, d).
Among the 53 neurons that responded to the gastric vagal stimulation with a phasic response, 39 (73.6%) were inhibited (including inhibitory-excitatory responses, \( p < 0.05 \); fig. 1a), and 14 (26.4%) were excited (including excitatory-inhibitory responses, \( p < 0.05 \); fig. 1b). The mean latencies of the inhibitory and excitatory responses were 89.7 ± 20.1 and 97.9 ± 21.8 ms (mean ± SD), respectively. There was no significant difference in the mean latencies of the inhibitory and excitatory responses (\( p > 0.05 \)). On 5 phasically responsive LHA neurons, the relationship between the stimulation intensity and the response magnitude was further examined. The results showed that these neurons exhibited stimulation intensity-dependent responses to the gastric vagal stimulation, revealing the existence of recruitment of the gastric vagal afferent inputs to the LHA neurons (fig. 2a–c).

In the remaining 7 LHA neurons whose firing patterns were modified in response to stimulation of gastric vagal nerves, 5 responded with a long-lasting decrease (\( n = 3 \)) or increase (\( n = 2 \)) in their firing rate (fig. 1c) and 2 changed their firing pattern from single and discrete spiking to burst discharging consisting of 2–5 spikes (fig. 1d). Once the stimulus was stopped or the intensity of stimulation was reduced to subthreshold, the changed firing patterns of these neurons reverted to their original discharge manner (fig. 1c, d), suggesting that the responses were stimulus-related. These results indicate that neurons in the LHA may show different response characteristics to the feeding-related visceral signals transmitted by the gastric vagal nerves.

We successfully completed the glycemia-sensitivity tests on 45 of the 60 LHA neurons that responded to the gastric vagal stimulation, although the remaining 15 cells were lost during the test session. On the basis of the test-

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**Table 1.** The responses of 114 LHA neurons to gastric vagal and cerebellar IN stimulation

<table>
<thead>
<tr>
<th>Neurons responding to gastric vagal stimulation</th>
<th>Neurons responding to cerebellar IN stimulation</th>
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<tbody>
<tr>
<td>Phasic responses 53 (53/60, 88.3%)</td>
<td>51 (51/114, 44.7%)</td>
</tr>
<tr>
<td>Inhibition 39 (39/53, 73.6%)</td>
<td>36 (36/51, 70.6%)</td>
</tr>
<tr>
<td>Excitation 14 (14/53, 26.4%)</td>
<td>15 (15/51, 29.4%)</td>
</tr>
<tr>
<td>Tonic responses 7 (7/60, 11.7%)</td>
<td>–</td>
</tr>
<tr>
<td>No responses 54</td>
<td>63</td>
</tr>
</tbody>
</table>

\(^1\) The total number of LHA neurons responding to gastric vagal stimulation, including phasic and tonic responsive cells, was 60 (60/114, 52.6%).

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**Fig. 2.** An LHA neuron that responded to the gastric vagal stimulation showing inhibition to the intravenous glucose administration. **a–c** PSTHs show the intensity-dependent excitatory responses of the LHA neuron to gastric vagal stimulations at different intensities (\( p < 0.05 \)). The stimulation intensity used for each experiment is given at the top right corner of each corresponding PSTH. The cell exhibited a specific inhibitory response to glucose administration (\( p < 0.05 \); d), but had no response to the saline and mannitol injection (\( p > 0.05 \); e, f), suggesting that the cell was a glycemia-sensitive neuron. Arrows under PSTHs (a–c) indicate the stimulation artifact, and the bars in PSTHs (d–f) show the glucose, saline and mannitol injection, which lasted for 30 s.
ed neurons showing a specific and significant inhibitory response ($p < 0.01$) to the intravenous injection of glucose but no response to the administration of normal saline and mannitol, 27 (27/45, 60%) were found to be glycemia-sensitive neurons. One such case is illustrated in figure 2d–f. The result demonstrates that glycemia-sensitive neurons in the LHA receive inputs from gastric vagal nerves.

**LHA Neuronal Responses to Cerebellar IN Stimulation**

Among the 114 LHA neurons recorded in this study, 51 (51/114, 44.7%) responded to the contralateral cerebellar IN stimulation (table 1). Of the 51 LHA neurons, 36 (36/51, 70.6%) showed inhibitory responses (including inhibitory-excitatory responses) and 15 (15/51, 29.4%) exhibited excitatory responses (including excitatory-inhibitory responses; fig. 3). Moreover, 33 of the 36 inhibited neurons (91.7%) showed a short latency (range 2–14 ms) response to the IN stimulation with a mean latency of 9.1 ± 2.8 ms, and the other 3 inhibited neurons (8.3%) responded to the IN stimulation with a long latency of >20 ms. Fourteen of the 15 excited neurons had a short latency excitatory response of 4–18 ms and the mean latency was 12.6 ± 3.2 ms, and another one showed a long latency of 50 ms to the stimulation. The characteristics of the responses of these responsive LHA neurons to the cerebellar IN stimulation were consistent with previous studies in cats [21, 22]. Furthermore, the results from 5 tested LHA neurons showed that the cells responded to the cerebellar IN stimulation in a stimulus intensity-dependent manner (fig. 4a–c).

We also conducted glycemia-sensitivity tests on 36 of the 51 LHA neurons that responded to the cerebellar IN stimulation. The results showed that 21 of the examined cells (21/36, 58.3%) were sensitive to glycemia (fig. 4d–f).

**LHA Neuronal Responses to Simultaneous Stimulation of the Gastric Vagal Nerves and Cerebellar IN**

Interestingly, in this study 30 of the 60 LHA neurons responsive to gastric vagal stimulation also responded to cerebellar IN stimulation. Twenty-one (70%) of these 30 neurons responded in the same direction to gastric vagal and cerebellar IN stimulation (i.e., the neurons showed either inhibitions or excitations to both of the stimulations); the remaining 9 cells responded in a different direction to the gastric vagal and cerebellar IN stimulation (9/30, 30%; i.e., inhibition/excitation or excitation/inhibition). These results suggest the possibility that convergence of gastric vagal and cerebellar IN inputs may happen in the same LHA neuron. To examine the possibility, the effect of simultaneous stimulation of gastric vagal nerves and cerebellar IN was observed in 6 cells. The results demonstrated that simultaneously stimulating these
Fig. 4. An LHA neuron that responded to the cerebellar IN stimulation showing inhibition to the intravenous glucose administration. The cell exhibited an intensity-dependent excitatory responses to the IN stimulations at different intensities (p < 0.05; a–c). In addition, in company with the increase in stimulation intensity (200 μA), the cell also showed inhibition following the excitatory response (p < 0.05; c). The stimulation intensity is given at the top right corner of each corresponding PSTH. The glycemia-sensitivity test showed that the neuron responded to the glucose injection with a specific inhibition (p < 0.05; d), but not to the saline and mannitol application (p > 0.05; e, f), suggesting that the cell was glycemia-sensitive. The layout of this figure is the same as figure 2.

Fig. 5. An LHA neuron that received convergent inputs from gastric vagal nerves and cerebellar IN showing inhibition following an intravenous glucose administration. a Stimulation of the gastric vagal nerves resulted in the recorded LHA neuron an excitation (p < 0.05). b Stimulation of the cerebellar IN induced an inhibitory response in the neuron (p < 0.05). c Simultaneous stimulation of the gastric vagal nerves and cerebellar IN did not elicit a significant cell response (p > 0.05), suggesting the existence of summation of the gastric vagal- and cerebellar IN-induced responses on the LHA cell. The test of intravenously injecting glucose (d), saline (e) and mannitol (f) revealed that the cell was a glycemia-sensitive neuron. a–c Arrows indicate the stimulation artifact. d–f Bars show the glucose, saline and mannitol injections, which lasted for 30 s.
two sites caused a summation of responses in all 6 LHA neurons tested, i.e., the responses of these neurons to simultaneous stimulation of the gastric vagal and cerebellar IN were attenuated in comparison with either stimulating gastric vagal or cerebellar IN alone if the responses of the neurons to stimulation of vagal and cerebellar IN were not in the same response direction (fig. 5); while the responses induced by simultaneously stimulating the two sites were more pronounced if the original responses were in the same direction (data not shown).

To further examine whether the gastric vagal and cerebellar IN inputs converge onto the same single LHA glycemia-sensitive neuron, we conducted a glycemia-sensitive test on 24 of 30 LHA neurons that responded to both the gastric vagal and cerebellar IN stimulation. The results showed that 15 of 24 cells (62.5%) were glycemia-

Table 2. Glycemia sensitivity of the LHA neurons responding to gastric vagal or/cerebellar IN stimulation

<table>
<thead>
<tr>
<th>Type of neurons</th>
<th>Neurons responding to gastric vagal stimulation (n = 45)</th>
<th>Neurons responding to cerebellar IN stimulation (n = 36)</th>
<th>Neurons responding to stimulations of gastric vagal nerves and cerebellar IN (n = 24)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycemia-sensitive neurons</td>
<td>27 (60%)</td>
<td>21 (58.3%)</td>
<td>15 (62.6%)</td>
</tr>
<tr>
<td>Glycemia-insensitive neurons</td>
<td>18 (40%)</td>
<td>15 (41.7%)</td>
<td>9 (37.4%)</td>
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</table>

Fig. 6. The distribution of neurons that responded to both gastric vagal and cerebellar IN stimulation in the LHA. 

- **a** A transverse section of the cerebellum (80 μm in thickness) showing the stimulation site in the IN (indicated by the arrowhead) and a transverse section of the hypothalamus (80 μm in thickness) presenting the recording site (indicated by the arrowhead).
- **b** Histological reconstruction showing the sites of neurons that responded to gastric vagal and cerebellar IN stimulation in the LHA. Three transverse sections of the hypothalamus corresponding to the planes Bregma –2.30 mm, –2.56 mm and –2.80 mm of the atlas of Paxinos and Watson [29] are presented, respectively. The recorded cells were either glycemia-sensitive (▲) or glycemia-insensitive (○) neurons.
sensitive neurons (fig. 5d–f), demonstrating that the afferent inputs from the gastric vagal nerves and cerebellar IN converge onto a single glycemia-sensitive neuron of the LHA, and that most LHA neurons which responded to gastric vagal and/or cerebellar IN stimulation were sensitive to glycemia (table 2).

Distribution of Responsive Neurons in the LHA

Histologically verified recording sites for the 30 LHA neurons that responded to both the gastric vagal nerves and cerebellar IN are reconstructed in figure 6. Fifteen of them were glycemia-sensitive neurons. It can be seen from figure 6 that the neurons responsive to both vagal and cerebellar IN stimulation and the glycemia-sensitive neurons were scattered over the LHA and showed no specific regional difference.

Discussion

It has been demonstrated that the gastric vagal fibers innervating the upper gastrointestinal tract can transmit three classes of meal-related information to the brain: mechanical distention of the lumen or gut contraction, chemical properties of luminal contents and gut peptides. Each of them reduces the meal size [16, 31]. This fact strongly suggests that gastrointestinal vagal afferent fibers may convey the negative-feedback signals to central areas, such as the nucleus tractus solitarius and hypothalamus, which play important roles in the control of food intake. However, little is known about the exact relationship between the peripherally activated vagal signals and the hypothalamic control of feeding behavior. The present study revealed that 52.6% of the recorded LHA neurons responded to gastric vagal stimulation and that most of the cells were inhibited. These results obtained from experiments on rats are comparable with previous findings from cats [14, 15]. Although the sub-diaphragmatic vagal trunks contain afferent and efferent components, a histological and functional study has well documented that at least 80% of the vagal fibers are afferent [32]. Besides, our previous study [25] revealed that vagotomy abolished the similar vagal-induced neuronal responses of the ventromedial hypothalamic nucleus (VMN), indicating that the stimulation may not activate any other spinal afferent pathways to the hypothalamus. In this study, the long response latencies of the LHA neuronal responses to gastric vagal stimulation suggested that inputs from the gastrointestinal tract to the hypothalamus are polysynaptic. Yuan and Barber [14, 15] demonstrated that they were relayed by the nucleus tractus solitarius. In addition, data from this study also showed that the responses of LHA neurons to gastric vagal stimulation were diverse, i.e., besides the transient phasic response, some LHA cells responded to gastric vagal stimulation with a change in their firing pattern. The result indicates that LHA neurons may employ various spiking patterns to encode different meal-related information that has arisen from the gastrointestinal tract for accurate transmission and processing of visceral sensory inputs in feeding control. It is thus suggested that the inputs from the gastrointestinal tract to LHA neurons, conveyed by gastric vagal afferent fibers, may be an important component of the negative feedback regulation of feeding.

Feeding behavior is closely linked to fluctuation in the blood glucose level [9–11, 27], and it is well documented that about 40% of the neurons in the LHA are glycemia-sensitive neurons [11, 33] which may be involved in meal onset triggered by a blood glucose drop [28]. In this study, the correlation between the gastric vagal afferent inputs and the glycemia-sensitive neurons in the LHA was further documented. Among 45 LHA neurons that responded to the gastric vagal stimulation, 27 (60%) were found to be sensitive to glycemia. The results prove that the gastrointestinal inputs conveyed by the gastric vagal afferent fibers mostly reach those glycemia-sensitive neurons of the LHA. Therefore, the modulatory effect of the gastric vagal afferent inputs on the LHA glycemia-sensitive neurons may play an important role in the short-term regulation of feeding behavior.

On the other hand, the cerebellar influence on feeding control has also been revealed successively during the past decade. Behavioral studies have demonstrated that animals with lesions of cerebellar cortex showed an alteration in food intake behavior, a disturbance in nutritional utilization, as well as a decrease in body weight [18, 34, 35]. A recent study using the technique of functional magnetic resonance imaging provided more direct evidence of cerebellar involvement in feeding control, indicating that the cerebellum was indeed activated by glucose intake in human subjects without any accompanying motor behavior [17]. Although the connections between the cerebellar fastigial nucleus and the glucose-sensitive/glycemia-sensitive neurons in the LHA have been disclosed [20, 22, 24], the cerebellar IN, which developed later in vertebrate phylogeny for adapting to the need for more precise control of distal limb muscles and seems to be more important in reaching and grasping food than fastigial control of axial and facial musculatures [36], has been neglected in research on cerebellohypothalamic pro-
The cell's sensitivity to the glucose administration (i.v.) on those LHA neurons that responded to both gastric vagal and cerebellar IN stimulation was also examined in the present study. Among 24 tested LHA neurons, the majority (15/24, 62.5%) was identified to be sensitive to glycemia (table 2). Because the ratio of glycemia-sensitive neurons in the LHA is less than 50% [11, 33], the present result provides strong evidence that gastrointestinal inputs and cerebellar IN inputs not only converge onto single LHA neurons but also mostly converge onto those LHA cells sensitive to glycemia. On the other hand, some studies also showed a strong relationship between the LHA glycemia-insensitive neurons and water- or food-seeking behavior [37–39]. Therefore, it can be inferred that the convergent inputs from the gastric vagal nerves and the cerebellar IN may also be involved in the formation of feeding motion by their effects on the glycemia-sensitive neurons in the LHA. On the basis of the above facts and the present data, we suggest that neurons in the LHA, especially those glycemia-sensitive neurons receiving both gastric vagal inputs and cerebellar IN inputs may integrate the meal-initiating signals (such as hypoglycemia), meal-limiting information (conveyed via gastric vagal afferents), and meal-related somatic information (forwarded by cerebellar IN through cerebellohypothalamic projections), and consequently trigger an appropriate behavioral response related to food intake. Combined with our previous findings that the cerebellar IN and gastric vagal afferent inputs reach and converge onto VMN glycemia-sensitive neurons [25] and the well-established functional reciprocal interaction between LHA and VMN on feeding control, we propose that this hypothalamic (including LHA and VMN) integration of the somatic-visceral (cerebellar IN, gastric vagal and glycemia) response may play a critical role in the short-term regulation of feeding behavior.

Acknowledgements

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