EXPERIMENTAL STUDY

Effect of the GH secretagogue L-163,255 and restricted feeding time on GH pulsatility in the rat

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Abstract

Objective: To determine the effect of repeated treatments with the growth hormone secretagogue (GHS) L-163,255 on the pulsatile release of GH when administered in meal-fed rats before and after feeding.

Design: The first group of rats (AL, n = 6) had food available ad libitum. The second (restricted, R, n = 6), third (GHSB, n = 6), and fourth (GHSA, n = 6) groups were fed from 1100 to 1400 h. Groups GHSB and GHSA were given GHS by gavage, 3.0 mg/kg L-163,255, at 1000 h (before feeding, B) and at 1500 h (after feeding, A) respectively. Three weeks after the initiation of the treatment, blood samples were collected at 10-min intervals over 6 h, and GH levels were determined.

Results: In group R, the concentrations of GH were higher before feeding than during feeding; P < 0.05. The average concentrations of the peak in response to GHS were higher when GHS was administered before (121.70 ± 33.68 ng/ml) than after (49.67 ± 17.87 ng/ml) feeding. The mass of GH, as calculated by deconvolution analysis was also higher in the GHSB group than in the GHSA group (251.6 ± 64.1 ng/ml per min vs 85.3 ± 22.9 ng/ml per min respectively, P < 0.05).

Conclusion: L-163,255 is effective in inducing GH release after repeated oral administration in rats. The effectiveness is greater when GHS is administered before rather than after feeding in meal-fed animals.

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Introduction

Growth hormone (GH) is a peptide hormone, produced and secreted from somatotrophs in the anterior pituitary gland. It exerts its action by binding to GH receptor in target tissues. GH is released in episodic bursts in a pulsatile manner, regulated by episodic changes in the concentration of two other hormones, growth hormone releasing hormone (GHRH) and somatostatin (SS). GHRH stimulates secretion whereas SS inhibits GH secretion (1). Growth hormone secretion can also be induced by synthetic ligands. Several such GH releasing substances, different from the naturally occurring GHRH, have been synthesized. These substances are called growth hormone secretagogues (GHS). The GHS are synthetic peptidyl and non-peptidyl molecules that exhibit strong GH releasing activity in vivo and in vitro. GHS induces GH secretion in a dose-dependent manner when administered i.v. or orally (2–5).

The GH secretagogues act through the growth hormone secretagogue receptor (GHS-R), a G protein coupled receptor (6). Ghrelin, an acylated stomach peptide, was recently reported as the first known endogenous ligand for this receptor (7). More endogenous ligands have been reported subsequently (8, 9). The GHS receptor cannot be activated by GHRH or somatostatin and the GH secretagogues affect GH secretion via a mechanism that is different from that of GHRH. Interestingly, the secretagogues and the peptidergic GH releasing hormone are synergistic with respect to GH release (4–6, 9).

The oral bioavailability of the first known peptidyl GHS was limited. Recently available GH secretagogues now include small-molecule, non-peptide secretagogues that can be administered orally to stimulate significant GH secretion (2).

The particular secretagogue used in this study is the spiropiperidine, L-163,255, developed by Merck and Co. Inc. It has been found to be very effective at stimulating
the secretion of GH, and subsequent elevation of circulating levels of insulin-like growth factor-I (10).

It is well known that GH is secreted in a pulsatile manner in episodic bursts (1). Studies in steers have shown a specific pattern of GH secretion that is related to feeding (11–13). Animals were ‘meal-fed’, such that they received food for only a certain period of time. In this case, the animals were fed from 0730–0930 h (12) or from 1000 to 1200 h (13). Under these conditions, GH was secreted in relatively high levels 1–3 h before the feeding period. During the feeding period GH was secreted at a very low level. After feeding, GH levels were relatively low, but started to rise again. Intravenous injection of bovine GHRH induced a rise in GH concentration in meal-fed steers. This rise was significantly higher when GHRH was injected 20 min before feeding than when it was injected 20 min after feeding (13).

There are no data available concerning the effect of GHS L-163,255 on the pulsatile release of GH, when administered to meal-fed rats. The present study was designed to determine the possible effects of GHS L-163,255 on the pulsatile release of GH when administered orally in meal-fed rats before and after feeding.

Repeated treatment with growth hormone secretagogues results, in some cases, in a desensitisation and a reduced GH release in response to these substances (2). A second objective of this study was to test whether the effect of GHS L-163,255 on GH release in rats is sustained after repeated oral treatment.

Materials and methods

All animal procedures were approved by the Institutional Animal Care and Use Committee (IACUC) of Virginia Commonwealth University (VCU).

Male Sprague–Dawley rats (Charles River Laboratories, Wilmington, MA, USA) were obtained at 125–150 g body weight (approximately 38–42 days of age). The animals were kept in individual steel cages in animal quarters controlled for temperature (21–23 °C) and light (14 h light:10 h darkness).

All rats had free access to water. Animals were fed a high protein diet consisting of 35.3% casein (Harlan Teklad, Madison, WI, USA), 34.7% sucrose, 15% cornstarch, 5.0% fibre-celulfil (Harlan Teklad), 5.0% corn oil, 3.5% mineral mix (AIN93, Harlan Teklad), 1.0% vitamin mix (AIN93, Harlan Teklad), 0.3% dl-methionine (Sigma Chemical Co., St Louis, MO, USA) and 0.2% choline bitartrate (Sigma Chemical Co.).

The animals were divided into four weight-matched groups. The first group (AL, n = 6) was fed the high protein diet which was available ad libitum (AL). The second group (R, n = 6) was ‘meal-fed’ such that the availability of the high protein diet was restricted to a period from 1100 to 1400 h. The third group (GHSB, n = 6) was food restricted with L-163,255 given one hour before (B) feeding (i.e. at 1000 h). The GHS was given by gavage at a dose of 3.0 mg/kg body weight. The frequency of administration was twice a week. Due to the limitations in the numbers of animals that could be sampled on one day (3), the animals received L-163,255 either on Monday and Friday or on Tuesday and Saturday. The fourth group (GHSA, n = 6) was food restricted and L-163,255 was given 1 h after (A) the end of the feeding period (i.e. at 1500 h). Route, dose and intervals of administration were the same as for the GHSB group. The two control groups (AL and R) were gavaged with saline at the same time intervals as the GHSB animals.

Body weight and food consumption were measured daily. Calculation of food intake was made by dividing the amount of food consumed per 100 g body weight (g food/100 g body weight). Calculation of food efficiency was made by dividing the amount of weight gained by the amount of food consumed (g weight gained/g food consumed).

For the blood sampling, polyvinyl catheters (Dural Plastics Ltd, Auburn, Australia) containing 50 IU/ml heparin, were implanted into the right atrium of all rats through the right external jugular vein. After testing, the catheters were plugged with a stainless steel plug. The operation was performed under ether anaesthesia. After recovering on a heating pad the rats were allowed to return to their cages. This operation was performed three days before sampling in order to allow the rats to adapt to the catheters.

Blood sampling began three weeks after the initiation of the GHS treatment for the determination of GH in plasma. Thirty minutes before sampling, a 50-cm polyvinyl extension was attached to the catheter with the use of a small piece of stainless steel tubing. Blood samples were collected at 10-min intervals for a period of 6 h. Sampling started 30 min before oral administration of saline or GHS by gavage. For groups AL, R and GHSB sampling lasted from 0930 h to 1530 h and for group GHSA it lasted from 1430 h to 2030 h.

Blood samples (0.25 ml) were collected with heparinized 1.0 ml syringes via the polyvinyl catheters and transferred into 1 ml heparinized conical tubes. Plasma was immediately separated with centrifugation and stored at −20 °C until assayed for GH by RIA. The red blood cells were re-suspended up to the initial volume in 0.9% w/v sodium chloride solution, and re-injected into the animal via the catheter, after the subsequent sample.

Assay for GH was performed with a radioimmunoassay (RIA) kit containing rat GH RP-2 as reference preparation, provided by Dr A F Parlow and the National Hormone and Pituitary Program of the NIDDK. The intra-assay coefficient of variation was 6.6% at 1.0 ng/tube and the interassay coefficient of variation was 10.5% at 0.625 ng/tube. Assay sensitivity was 4.0 ng/ml.
The pulsatility of the GH secretion was calculated with the multiple parameter deconvolution method of Veldhuis et al. (14) and Veldhuis and Johnson (15). This method uses the plasma concentrations of GH for the calculation of the mean GH secretory burst area (mass) for the specific GH burst that followed GHS treatment. It also allowed for the calculation of other parameters of GH pulsatile secretion over the 6-h period. These other parameters included: basal secretion, half-duration, half-life, number of bursts, interburst interval, burst amplitude, pulsatile production rate, and total production rate.

Intergroup comparisons were carried out with analysis of variance (ANOVA) followed by Duncan’s Multiple Range test in case of significance. All results are presented as means ± S.E.M.

Results

In general, the dose of L-163,255 used in these experiments caused no significant changes in food intake, weight gain, or food efficiency.

Figure 1 illustrates the mean plasma GH concentrations during the 6-h sampling period. In the AL group, the normal pattern for the GH secretion was observed. In the food-restricted group, R, the concentrations of GH were significantly higher before feeding (17.6±2.4 ng/ml from 0930 h to 1100 h) than during feeding (11.2±1.2 ng/ml from 1110 h to 1400 h, P < 0.05). After the feeding period, the GH levels were 11.0±2.9 ng/ml from 1410 to 1530 h.

A marked increase in circulating GH levels was observed in groups GHSA and GHSB with a maximum at 30 min after the administration of the GHS L-163,255. All six rats of the GHSB group responded to the GHS with a GH peak after 30–40 min (peak height range 38–248 ng/ml). Five out of six rats of the GHSA group responded to the GHS with a GH peak (peak height range 46–105 ng/ml).

The mean values for GH 30 min after saline or GHS administration by gavage were 14.32±4.10 ng/ml, 23.09±8.31 ng/ml, 121.70±33.68 ng/ml and 49.67±17.87 ng/ml in groups AL, R, GHSB and GHSA respectively. The GH concentration at that time was significantly higher in the GHSB group than in the saline-treated groups (AL vs GHSB P < 0.05, R vs GHSB P < 0.05). In the GHSA group, a peak in response to GHS was also observed. However, the difference between groups GHSA and R was not significant despite the clear GH peak in the GHSA group (R vs GHSA P = 0.2). Comparison of the peaks of GH 30 min after GHS administration showed that the administration of GHS before feeding stimulated a higher peak of GH than when GHS was given after feeding (P < 0.05).

Figure 2 shows the results of deconvolution analyses of the individual burst mass immediately after GHS administration. The mass of GH secreted by the pituitary gland in response to GHS before feeding (GHSB) was significantly greater than that stimulated by GHS after feeding (GHSA) (251.6±64.1 ng/ml per min vs 85.3±22.9 ng/ml per min respectively, P < 0.05). In contrast to these significant differences, there was no significant difference in any of the overall parameters of GH pulsatility during the 6-h period of sampling.

Table 1 shows the average values for weight gain, food intake and food efficiency (means ± S.E.M.) in the
four groups of rats. Group AL had higher weight gain and food intake. No significant differences were observed between groups R, GHSB and GHSA.

**Discussion**

Growth hormone is released by the pituitary gland in episodic pulses in the rat, human, and other mammals (1). Feeding is known to affect GH secretion (11–13). The GH pulses are known to increase before feeding, and to decrease during feeding and for up to 2–3 h afterwards (11–13). Consistent with these results, the rats of the restricted (R) group in our experiment had lower average GH concentrations during the feeding period from 11.10 h to 1400 h than during the period before feeding.

The GHS L-163,255 has been found to be effective in stimulating GH secretion in pigs (10, 16), horses (17) and chickens (18). This is the first observation about the effect of this GHS in rats. In the GHS-treated animals a clear peak in response to GHS was observed 30 min after gavage.

In a previous study, reduction of the serum GH response to an injection of growth hormone releasing factor (GRF) was observed when steers, fed once a day at 1400 h, were injected with GRF after feeding. The best response to GRF was observed when the GRF was injected 2 h before feeding (12). In another experiment, injection of bovine GHRH to meal-fed steers before feeding caused a significantly higher increase in circulating GH than after feeding (13).

The possible effect of restricted feeding on the response to GHS is not known. Our data show that there is a significantly higher impact on circulating GH when the GHS is administered before feeding. The administration of GHS caused a significantly higher release of GH when it was given before feeding than after feeding. The mass of GH that was secreted after the GHS stimulus, as calculated with deconvolution analysis, was significantly higher in GHSB than in GHSA (251.6±64.1 ng/ml vs 85.3±22.9 ng/ml, \( P < 0.05 \)). Likewise, the average peak height 30 min after GHS was also higher in the GHSB than in the GHSA group.

With respect to the mechanism by which GHS, or other GH-releasing stimuli, might be more effective before, in contrast to after, a restricted period of feeding, we would hypothesize an important role for the level of somatostatin tone from the hypothalamus. McMahon et al. (13) reported a relatively low level of activity of somatostatin neurons during the period before restricted feeding in steers. We would suspect that a low level of somatostatin coming from the hypothalamus would allow for a greater response to GH-releasing stimuli. On the other hand, a relatively high level of somatostatin tone after feeding might be important to a lower level of GH stimulation.

Christensen et al. (17) administered the GHS L-163,255, i.v., to mares at doses of 1.0 and 5.0 mg/kg body weight. Those animals were fed twice daily, at 0800 h and at 1500 h and the GHS was injected at 1300 h. A GH peak in response to GHS was observed 34 min after injection. There was no difference in GH release between the 1.0 and the 5.0 mg/kg dose. In our experiment, the GH release in response to oral administration of L-163,255 in a dose of 3.0 mg/kg body weight was observed after a time period of 30 min. This suggests that the GHS L-163,255 is quickly absorbed through the intestinal system and acts quickly. Thus, oral administration is

**Table 1** Weight gain, food intake and food efficiency in the four groups of rats studied. Results are means ± S.E.M.

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Weight gain (g/day)</th>
<th>Food intake (g/100 g body weight)</th>
<th>Food efficiency (g of food intake/g weight gained)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AL</td>
<td>6</td>
<td>8.38 ± 2.57</td>
<td>9.94 ± 0.53</td>
<td>0.55 ± 0.19</td>
</tr>
<tr>
<td>R</td>
<td>6</td>
<td>5.39 ± 2.82</td>
<td>7.65 ± 1.05</td>
<td>0.46 ± 0.21</td>
</tr>
<tr>
<td>GHSB</td>
<td>6</td>
<td>4.83 ± 2.70</td>
<td>7.45 ± 1.00</td>
<td>0.45 ± 0.22</td>
</tr>
<tr>
<td>GHSA</td>
<td>6</td>
<td>4.99 ± 2.80</td>
<td>7.42 ± 1.02</td>
<td>0.45 ± 0.19</td>
</tr>
</tbody>
</table>

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Effect of GHS and restricted feeding in rats

This is the first observation, to our knowledge, about the effect of GHS L-163,255 in rats. Our results confirm the observation in other species of animals that the GHS L-163,255 can cause significant growth hormone release (10, 16–18). In the rat there is a maximum release about 30 min after the oral administration of GHS.

A desensitization and sustained amplification of pulsatile profile of GH release has been observed after repeated treatment with GH secretagogues (2). Repeated administration of the MK-0677 GHS to dogs resulted in a decrease in the GH response by 41–77% (19). In our experiment, the GH release in response to the GHS was observed after a 3-week treatment with GHS in bi-weekly intervals. Although the design of our experiment did not allow the measurement of the response to L-163,255 after first time administration for comparison, our data suggest that this secretagogue is capable of effectively stimulating GH secretion even after repeated administration.

In contrast to the significant acute stimulation of GH release within about 30 min after treatment, GHS did not exert a significant effect on the overall parameters of GH pulsatility during the total sampling period of 6 h. These results lead to the probability that GHS can acutely affect GH secretion without necessarily disrupting its overall, long-term, pulsatile pattern.

Growth hormone secretion is known to be significantly related to ‘fasted’ and ‘fed’ states. In humans, GH is secreted at significantly higher levels during the fasted state (20). We had not initially considered this study to be related to ‘fasted’ versus ‘fed’ states. However, one could consider the time immediately preceding a 3-h/day feeding period to be a relatively ‘fasted’ state. Likewise, the period after feeding would be a ‘fed’ state. Despite the typical incongruence of GH pulsatility in rats to that of humans with respect to fasting, i.e. rats typically show decreased GH secretion during fasting (21), the current data indicate that rats are very likely to be more responsive to GH-releasing stimuli during a period in which they have certainly been without food for a relatively long period of time. It is possible that, although the pituitary gland may be more responsive to GH-releasing stimuli, those stimuli do not come down from the hypothalamus at that time.

We have actually thought of the current studies as being designed around the ‘anticipation of feeding’. In that sense they have not necessarily been studies of the ‘fasted’ state per se. Studies of potential differences between ‘fasting’ and ‘anticipation of feeding’ might be useful for a better understanding of GH physiology and growth.

Despite the clear acute effect on GH release, no significant difference in growth was observed in the GHSB and GHSA groups compared with the restricted group. This may be due to the meal feeding itself, or to the dose of GHS which was not sufficient, or to some feedback effect. Tests with higher doses of GHS should be conducted.

We conclude that L-163,255 is effective in inducing GH release after repeated oral administration at a dose of 3 mg/kg in rats. The effectiveness of this GHS is greater when it is administered before feeding than after feeding in meal-fed animals. This observation may be useful for the optimization of the effectiveness of this secretagogue.

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