Clinical Research Article

Corroboration of Height Velocity Prediction Markers for rhGH With an Oral GH Secretagogue Treatment in Children With GHD

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Abstract

Context: Recombinant human growth hormone (rhGH) is approved for treatment of pediatric growth hormone deficiency (GHD), with greatest growth responses observed in those with severe GHD. Orally administered GH secretagogues (GHS) may be useful treatment in patients with moderate GHD. Distinguishing children with severe vs moderate GHD could identify children who would be better treated with rhGH or GHS.

Objectives: Evaluate baseline insulin-like growth factor-I (IGF-I) and stimulated peak GH response as predictors of 12-month height velocity (HV) in children with GHD.

Design: Data on children with GHD were analyzed in a legacy data base (GeNeSIS data).

Participants: 514 naïve to rhGH-treatment, prepubertal children with idiopathic isolated GHD for whom stimulated GH, baseline serum IGF-I, and first-year HV during rhGH treatment data are available.

Outcome Measures: Children with severe or moderate GHD were categorized based on GH and IGF-I data and evaluated based on baseline auxologic and hormone profiles and first-year growth response to rhGH.

Results: Cohorts of severe and moderate GHD were 81/514 (15.8%) and 433/514 (84.2%). Cohorts differed significantly with regard to indicators of GHD [eg, baseline height SD score (SDS), height SDS minus target height SDS, HV, HV SDS, and change in height SDS during rhGH treatment]. Multiple regression analysis showed IGF-I and stimulated GH were significant predictors of HV independent of other known variables. Expected first-year HV in moderate GHD was 8.3 cm/y.

Conclusions: The combination of peak GH to GH stimulation testing and baseline IGF-I concentration are predictive enrichment markers for annualized HV responses to rhGH therapy.
Growth hormone deficiency (GHD) in children is a well-recognized medical condition leading to physical disability and psychological distress associated with short stature starting during childhood and persisting throughout adult life [1-4]. The amount of growth hormone (GH) secreted by children with GHD covers a range, from zero in the most severe cases to measurable but subnormal quantities in milder disease [5,6]. The condition is rare and is therefore defined as an “orphan” disease by Food and Drug Administration, European Medicines Agency, and other regulatory authorities. The only currently approved therapy for GHD is recombinant human growth hormone (rhGH), which must be given by daily subcutaneous injection. Nonadherence to this therapy is a widely recognized issue [7-9]. Ibutamoren (LUM-201, Lumos, Austin, TX, USA), previously MK-0677 (Merck and Company, Rahway, NJ, USA) is an orally bioavailable small molecule that was instrumental in the cloning of the GH secretagogue receptor [10,11]. The GH secretagogue receptor was used to identify ghrelin as the natural ligand for this receptor [12]. Ibutamoren increases the release of endogenous GH in adults [13] and in children more mildly affected by GHD [14,15] and has the potential to stimulate childhood growth over the longer term, attenuating the physical and psychological symptoms associated with short stature. Internal feedback systems limit the secretion of GH upon administration of such a GH-releasing agent [16], potentially preventing safety issues associated with exposure to excessive levels of GH. Because it is administered orally rather than by daily injection, ibutamoren treatment could increase adherence to a regimen of growth-promoting therapy, easing the burden of the disease on patients and families. Results of a previous study found no statistically significant difference in annualized height velocity (HV) after 6 months of treatment with ibutamoren or daily rhGH injections when moderate GHD was defined as a baseline insulin-like growth factor-I (IGF-I) > 30 µg/L and a peak GH to single dose ibutamoren ≥5 µg/L [17]. Consequently, baseline serum IGF-I and GH response to a single dose of the investigative drug were used to identify a subset of patients with moderate GHD. In addition, that study showed a peak GH response to a single dose of ibutamoren >5 µg/L was equivalent to a GH response in a standard stimulation test of >2 µg/L [17]. In planning for a comparative trial of different doses of ibutamoren vs standard doses of injected rhGH, data illustrating the effect of rhGH in a large population of children who participated in the Genetics and Neuroendocrinology of Short Stature International Study (GeNeSIS; Eli Lilly and Company, Indianapolis, IN, USA) were evaluated. Since theoretically GH secretagogues require a functionally intact pituitary gland, the analysis focused on patients with idiopathic isolated GHD (IGHD) excluding those with organic GHD or combined pituitary hormone deficiency. The objective of this analysis was (i) to corroborate the criteria for moderate GHD determined from the previous study [17] in an independent cohort and (ii) to obtain an average value of first-year annualized HV attained with standard rhGH therapy in this subset as a comparator for ibutamoren treatment. Such an approach would strengthen the evidence required by providers, patient families, and others to evaluate the effectiveness of the oral GH secretagogue in comparison with that of currently standard rhGH therapy.

**Patients and Methods**

**Source of data**

Data from GeNeSIS (Clinical Trial Registry Number: NCT01088412) were analyzed to serve as the independent cohort. Anonymized data of individual participants were made available by Lilly on the platform www.clinicalstudydatarequest.com after approval by an independent review committee.

**Patients and data collection**

GeNeSIS was a prospective, open-label, observational research program conducted in 30 countries at more than 800 study sites between 1999 and 2015. The main objective was to investigate safety and effectiveness of rhGH treatment (Humatrope®, Eli Lilly and Company, Indianapolis, IN, USA) in pediatric patients with growth failure. Data were collected by investigators according to their standard practices and entered on case report forms. Data included etiology of GHD; demographic, auxologic, and biochemical data; status of other pituitary hormones; and information on abnormal brain or pituitary imaging [18,19]. Bone age was mostly determined according to Greulich-Pyle, and data obtained by the Tanner-Whitehouse method were converted as described before [20]. Baseline data were collected before initiation of GH treatment, and follow-up data were typically collected at 6-month intervals. The plausibility of reported data was reviewed using scatter plots, and sites were queried to comment on or to correct implausible data. The diagnosis of GHD in an individual patient was

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**Key Words:** recombinant human growth hormone, rhGH, GeNeSIS, GH deficiency, GHD, insulin-like growth factor-I, IGF-I
patients with idiopathic GHD without additional growth-limiting disorders, further selection steps followed: IGHD, GH-naïve at baseline, GH-treated, chronological age ≥4 y and ≤10 y at baseline, pubertal stage 1 (B1 in girls, G1 in boys) at baseline, first-year annualized HV (cm/y) available and plausible, stimulated maximum GH and baseline serum IGF-I available (Fig. 1).

Statistics

Age- and sex-based SDS values for height [24], annualized HV [25], body mass index (BMI) [26], target height according to Tanner, bone age, and serum IGF-I were calculated as described previously [18, 19]. Results are presented as mean ± SD unless otherwise stated. Comparisons among groups were performed by 1-way analysis of variance for continuous variables and by chi-square test for categorical variables. For factors influencing response to rhGH, multiple linear regression analysis was performed with first-year annualized HV as the response variable and varied sets of exploratory variables. The baseline characteristics examined included chronological age, sex, weight SDS, BMI SDS, height SDS, difference between height and target height SDS, dose per week, serum IGF-I, and maximum GH (continuous or categorical by the cutoff). Sex was not a significant predictor and was excluded from subsequent modeling. Weight SDS was also removed from the models because of high collinearity with height SDS, and BMI SDS was selected instead. Model diagnostics were carried out by examining residuals, leverages, and Cook's distances; 3 influential outliers were thus excluded from the analyses. Both standardized and unstandardized regression parameter estimates were reported. All analyses were conducted using SAS 9.4 (Cary, NC, USA). A P-value of less than 0.05 was considered statistically significant.

Results

Patients

The independent cohort that was analyzed after the filtering process identified patients considered a priori the most suitable candidates for ibutamoren therapy (Fig. 1). The authors believe that the diagnosis of idiopathic IGHD was the most important diagnostic category for identifying the patients of interest, accounting for 73.3% of all patients with GHD (9022 of 12 315 patients). Only rhGH treatment-naïve patients at baseline who actually received treatment were eligible (72.4% of idiopathic IGHD and 53.0% of all GHD). To avoid the interference caused by patients undergoing a pubertal growth spurt and the high HV of patients at a very young age, the study population was further limited to patients with
baseline chronological age between 4 and 10 years and prepubertal stage (Tanner 1), which further reduced the cohort (30.5% of rhGH-treated naive idiopathic IGHD, 16.2% of all GHD). Because the primary endpoint was first-year annualized HV, only patients for whom this variable was available and plausible were included (27.2% of rhGH-treated naive idiopathic IGHD patients, 14.4% of all GHD patients). To test the hypothesis that stimulated maximum GH and baseline serum IGF-I are suitable markers for determining the severity of GHD, we focused finally on the cohort where both variables were available and plausible (N = 514, 7.9% of rhGH-treated naive idiopathic IGHD patients, 4.2% of all GHD patients). Based on a previous study [17], we hypothesized that either stimulated maximum GH < 2 µg/L or baseline serum IGF-I ≤ 30 µg/L indicate severe GHD in contrast to moderate GHD predicted by the combined markers GH ≥ 2 µg/L and IGF-I > 30 µg/L. Cohort sizes of these 2 subgroups were 81/514 (15.8%) with severe GHD and 433/514 (84.2%) with moderate GHD.

### Demographics at baseline

Baseline characteristics before start of rhGH treatment for all patients and for the proposed subgroups are shown in Table 1. The total number of boys [313/514 (61%)] was greater than that of girls [201/514 (39%)]. The individual GHD subgroups were also predominantly male; however, the difference was less marked in the moderate GHD cohort. Chronological age was similar for the subgroups. Bone age was significantly more retarded in the severe vs the moderate GHD group when adjusted for chronological age. Height SDS was more diminished in the severe GHD cohort (−3.01 vs −2.58; P < 0.0001), while target height SDS was less diminished (−0.52 vs −0.75; P = 0.0146); consequently, the difference between the patients’ height SDS and their target (mid-parental) height SDS was, on average, greater (−2.49 vs −1.82; P < 0.0001). Pretreatment annualized HV was not different between the 2 cohorts. BMI SDS was significantly lower in the severe GHD group (−1.36 vs −0.37; P < 0.0001). The stimulated maximum GH peaks and baseline serum IGF-I values (concentrations or means of SDS) were different between both groups, as expected. The values complied with the predefined cutoffs of these parameters. The mean GH dose was slightly greater in the severe GHD group (0.21 mg/kg/week vs 0.20 mg/kg/week; P < 0.0200), which lay well in the recommended dose ranges of 0.18 to 0.25 mg/kg/week (Europe) and 0.18 to 0.3 mg/kg/week (USA).

### Changes During rhGH treatment

Mean height SDS at 1 year of rhGH treatment (Table 2) was not significantly different between the severe and moderate GHD cohorts due to a greater mean increase in the severe GHD group (0.86 vs 0.61; P < 0.0001). This finding corresponds with the greater mean first-year HV and HV SDS of the severe GHD group (−1.36 vs −0.37; P < 0.0001). The stimulated maximum GH peaks and baseline serum IGF-I values (concentrations or means of SDS) were different between both groups, as expected. The values complied with the predefined cutoffs of these parameters. The mean GH dose was slightly greater in the severe GHD group (0.21 mg/kg/week vs 0.20 mg/kg/week; P < 0.0200), which lay well in the recommended dose ranges of 0.18 to 0.25 mg/kg/week (Europe) and 0.18 to 0.3 mg/kg/week (USA).
### Table 1. Baseline characteristics of the final study group and the cohorts with severe or moderate GH deficiency

| Variable                                      | All          | Severe GHD IGF-I ≤ 30µg/L or GH < 2 µg/L | Moderate GHD IGF-I > 30µg/L and GH ≥ 2 µg/L | P       |
|----------------------------------------------|--------------|------------------------------------------|------------------------------------------|---------|
| Male/female (%)                              | 514 61/39    | 81 72/28                                 | 433 59/41                                | 0.0349* |
| Chronological age (year)                     | 514 7.05 1.7 | 81 6.80 1.7                               | 433 7.10 1.7                             | 0.1535  |
| Bone age (year)                              | 380 5.04 2.0 | 60 4.39 2.3                               | 320 5.17 1.9                             | 0.0049  |
| Bone age—chronological age (year)            | 380 -2.14 1.2 | 60 -2.52 1.6                            | 320 -2.07 1.1                           | 0.0072  |
| Bone age SDS                                 | 380 -2.67 1.5 | 60 -3.15 1.8                           | 320 -2.57 1.4                           | 0.0044  |
| Height SDS                                   | 514 -2.65 0.8 | 81 -3.01 1.0                           | 433 -2.58 0.7                           | <0.0001 |
| Target height SDS                            | 492 -0.71 0.8 | 80 -0.52 0.8                           | 412 -0.75 0.8                           | 0.0146  |
| Height SDS – target height SDS               | 492 -1.93 1.0 | 80 -2.49 1.1                           | 412 -1.82 0.9                           | <0.0001 |
| Pretreatment HV (cm/y)                       | 363 4.80 1.3 | 54 4.99 1.6                               | 309 4.76 1.3                             | 0.2352  |
| Pretreatment HV SDS                          | 363 -1.29 1.4 | 54 -1.09 1.7                           | 309 -1.32 1.3                           | 0.2562  |
| Body mass index SDS                          | 510 -0.53 1.4 | 80 -1.36 1.7                           | 430 -0.37 1.3                           | <0.0001 |
| Maximum GH peak (µg/L)*                     | 514 7.40 4.8; 10.5 | 81 5.10 1.4; 9.3                     | 433 7.73 5.4; 10.8                       | <0.0001 |
| IGF-I (µg/L)*                                | 514 80 49; 109 | 81 22 11; 43                           | 433 86 60; 114                         | <0.0001 |
| IGF-I SDS                                    | 509 -1.72 1.7 | 79 -4.13 2.3                           | 430 -1.28 1.2                           | <0.0001 |
| rhGH dosage weekly (mg/kg/wk)                | 507 0.20 0.045 | 80 0.21 0.053                         | 427 0.20 0.043                         | 0.0200  |

*P* values for comparing severe and moderate GHD groups were calculated by one-way ANOVA unless specified otherwise.

Abbreviations: ANOVA, analysis of variance; GH, growth hormone; GHD, growth hormone deficiency; IGF-I, insulin-like growth factor-I; HV, height velocity; rhGH, recombinant human growth hormone; SDS, SD score.

*P*-value by chi-square test.

*Because of skewed distribution of serum GH and IGF-I concentrations, data are shown as median, first quartile, and third quartile. *P*-value calculated using Wilcoxon rank sum test.
and ended up at greater concentrations. The change in IGF-I SDS was greater in the severe GHD group (3.01 vs 1.35, \( P < 0.0001 \)).

**GH, IGF-I, and height velocity**

To further confirm the combined cutoffs for stimulated maximum GH and serum IGF-I concentration as positive enrichment markers, we divided the population into 2 stimulated GH categories (<2 µg/L or ≥2 µg/L) and varied IGF-I cutoffs from 0 to 120 µg/L. First-year annualized HV was chosen as the dependent response variable (Table 3). Evidently, low GH or low IGF-I result in a better growth response with a maximum at GH < 2 µg/L and IGF-I < 30 µg/L (mean HV = 12.6 cm/y). In patients with low GH (<2 µg/L) and IGF-I above the cutoff, mean annualized HV was clearly smaller and decreased further with increasing baseline IGF-I cutoff, but to a smaller degree. In patients with greater GH (≥2 µg/L) but IGF-I below the cutoffs (0-120 µg/L), mean HV was greater and decreased slightly from 10.6 to 8.6 cm/y. Mean annualized HV was smallest in patients with greater GH (GH ≥ 2 µg/L) and decreased slightly with IGF-I above increasing cutoffs (8.5-7.8 cm/y). At GH ≥ 2 µg/L and IGF-I cutoff >30 µg/L, the first-year HV was 8.3 ± 1.6 cm/y. When serum IGF-I was divided into 2 categories (≤30 µg/L or >30 µg/L) and cutoffs for stimulated GH maximum varied between 0 and 10 µg/L, similar picture was obtained (Table 4). Mean HV was greatest in the low IGF-I group (≤30 µg/L) and decreased with increasing stimulated GH cutoff values. In the high IGF-I group (>30 µg/L), mean HV was practically independent of the GH cutoff values; for GH greater than the increasing cutoffs, mean annualized HV remained around 8.3 cm/y. A visual impression of the dependence of the mean first-year HV on the choice of cutoff values for stimulated GH maximum and baseline serum IGF-I is given in Figures 2 and 3.

**Multiple regression analysis**

HV during rhGH treatment in patients with GHD is influenced by a number of parameters. To assess the value of stimulated maximum GH and serum IGF-I concentration as markers for rhGH response in comparison with other potential markers, we conducted a series of multiple linear regression analyses with first-year HV as the response variable and a variety of exploratory variables including stimulated maximum GH and baseline IGF-I (continuous or categorical according to the cutoffs), sex, baseline chronological age, baseline bone age, baseline height or height SDS, baseline BMI SDS, target height or target...
height SDS, and GH dose. In general, the models were significant (Pr(|t|<|0.0001)) and of acceptable quality. Depending on the set of exploratory variables, chronological age, BMI SDS, and rhGH dose contributed significantly to growth prediction. However, IGF-I (absolute, SDS, or categorical) was the most powerful predictor of first-year HV in all models, with an inverse relationship to HV. Relevant parameters of the best model are shown in Table 5. The single most important predictor was the categorical variable of baseline IGF-I > 30 µg/L with a standardized estimate of -0.2982. The second most important predictor was baseline chronological age, followed by GH dose, baseline BMI SDS, baseline height SDS minus target height SDS, and maximum GH ≥ 2 µg/L. Baseline height SDS was not a significant predictor. The significance of the combined markers stimulated GH ≥ 2 µg/L and baseline IGF-I > 30 µg/L was verified by this multivariate analysis (Table 5).

### Discussion

Examination of the GeNeSIS data corroborates the use of baseline IGF-I concentration and stimulated GH as meaningful indicators of the severity of GH and the response to treatment. Using cut-off values comparable to those determined in a companion study [17], baseline IGF-I concentration ≤30 ng/mL and stimulated GH < 2 are independent indicators of good annualized HV to rhGH in children with GHD. In the IGF-I cut-off groups, subjects with lower IGF-I concentration had higher annualized HV than those with higher IGF-I; annualized HV decreases as the IGF-I cut-off is increased. In the GH cut-off groups, annualized HV is highest in the low GH group and, again, HV decreases with increasing IGF-I. The IGF-I and GH cutoffs identified in this study also serve to segregate the study population into severe and moderate GHD groups. Those with baseline IGF-I concentration ≤30 ng/ml or GH < 2 have more severe GHD and are found to have lower height.
SDS, greater differences in chronological and bone age, greater differences in height SDS and target height SDS, and lower pretreatment height velocities. Moreover, the group with severe GHD has greater first-year annualized HV when treated with daily injections of rhGH.

Ibutamoren is a GHS receptor agonist that was studied in the late 1990s as a potential therapy for pediatric GHD. At the time, study results suggested a lack of efficacy when compared to rhGH [14,27]. Re-examination of the data in 2018 confirmed the findings from these studies. When data from all children studied were combined, the growth data during the first 6 months of rhGH or ibutamoren therapy showed that daily injected rhGH was superior to once-daily oral ibutamoren therapy. However, when only the data from children with severe GHD were considered, the efficacy of rhGH treatment was even more pronounced. In contrast, among children with moderate GHD, the difference in 6-month HV between the ibutamoren- and

Table 4. First-year height velocity (cm/y) during rhGH treatment in low (≤30 µg/L) and high (>30 µg/L) IGF-I groups according to different stimulated GH peak concentrations (N = 514)

| GH Cutoff (µg/L) | Baseline GH < cutoff | Baseline GH ≥ cutoff |
|------------------|----------------------|----------------------|
|                  | AHV Mean (cm/y) | AHV SD (cm/y) | n   | AHV Mean (cm/y) | AHV SD (cm/y) | n   |
| IGF-I ≤ 30       |                    |                    |      |                    |                    |      |
| 0                | —                   | —                   | —    | 10.19              | 2.51              | 59   |
| 1                | 13.24               | 1.98               | 7    | 9.78               | 2.29              | 52   |
| 2                | 12.63               | 2.31               | 9    | 9.75               | 2.30              | 50   |
| 3                | 12.57               | 2.04               | 13   | 9.52               | 2.21              | 46   |
| 4                | 12.52               | 1.97               | 14   | 9.47               | 2.21              | 45   |
| 5                | 11.69               | 2.47               | 18   | 9.53               | 2.25              | 41   |
| 6                | 11.21               | 2.52               | 25   | 9.44               | 2.24              | 34   |
| 7                | 11.09               | 2.49               | 31   | 9.20               | 2.16              | 28   |
| 8                | 11.03               | 2.36               | 36   | 8.88               | 2.18              | 23   |
| 9                | 10.89               | 2.38               | 38   | 8.93               | 2.28              | 21   |
| 10               | 10.64               | 2.43               | 44   | 8.90               | 2.35              | 15   |
| IGF-I > 30       |                    |                    |      |                    |                    |      |
| 0                | —                   | —                   | —    | 8.28               | 1.58              | 455  |
| 1                | 9.85                | 0.96               | 5    | 8.27               | 1.58              | 450  |
| 2                | 8.09                | 2.06               | 22   | 8.29               | 1.56              | 433  |
| 3                | 8.40                | 1.89               | 36   | 8.27               | 1.56              | 419  |
| 4                | 8.42                | 1.77               | 68   | 8.26               | 1.55              | 387  |
| 5                | 8.46                | 2.07               | 118  | 8.22               | 1.37              | 337  |
| 6                | 8.40                | 1.93               | 167  | 8.21               | 1.34              | 288  |
| 7                | 8.38                | 1.83               | 213  | 8.20               | 1.33              | 242  |
| 8                | 8.33                | 1.78               | 246  | 8.23               | 1.31              | 209  |
| 9                | 8.29                | 1.71               | 291  | 8.28               | 1.33              | 164  |
| 10               | 8.34                | 1.69               | 334  | 8.13               | 1.22              | 121  |

Abbreviations: AHV, annualized height velocity; GH, growth hormone; IGF-I, insulin-like growth factor-I.

Figure 2. Dependence of first-year HV on stimulated maximum GH (<2 µg/L vs ≥2 µg/L) and varying baseline IGF-I cutoffs.

Figure 3. Dependence of first-year HV on baseline IGF-I (≤30 µg/L vs >30 µg/L) and varying baseline GH cutoff.
rhGH-treated patients was not statistically significant [17]. Based on this observation, the data were retrospectively examined, and 2 criteria were selected to evaluate the effect of treatment based on the severity of GHD in individual children. We hypothesized that a peak GH response to a single test dose of 0.8 mg/kg ibutamoren orally and baseline serum IGF-I could distinguish between the children who had moderate GHD and those who were severely deficient [17]. As there were no data on the response to a test dose of ibutamoren in the GeNeSIS data base, we also performed a receiver operator curve analysis from the data of acute response to ibutamoren and compared the results to those of either serum IGF-I or standard GH stimulation tests. We determined that a cutoff of 5 µg/L in the ibutamoren stimulation test was equivalent to a peak response to a standard GH stimulation test of 2 µg/L [17]. For serum IGF-I, a conservative cutoff of 30 µg/L was tested as a way to distinguish between moderate and severe GHD.

Since this was a retrospective analysis and the number of children in the study [17] was limited, we wanted to confirm this approach, in particular the GH and IGF-I cutoffs, in an independent large cohort by studying typical indicators of the severity of GHD in children treated with rhGH. When the GeNeSIS data base was made accessible to investigators, courtesy of Eli Lilly and Company through clinicalstudydatarequest.com, we were given permission to interrogate the data base. This paper presents the results of that interrogation. Theoretically, responsiveness to an oral GH secretagogue requires functionally intact somatotrophs in the pituitary. It is unlikely that this prerequisite is fulfilled in patients with an organic etiology of GHD such as genetic defect, abnormal pituitary development, intracranial tumor, certain clinical syndromes, central nervous system malformations, irradiation, and other causes. Most of these etiologies cause combined pituitary hormone deficiencies. Therefore, we focused on patients with idiopathic IGHD, which accounts for the great majority of patients with GHD [4,28,29].

In this study, severe GHD was defined by either IGF-I ≤ 30 µg/L or maximum GH < 2 µg/L, while moderate GHD was defined by IGF-I > 30 µg/L and GH ≥ 2 µg/L. Both cohorts differed significantly at baseline in height SDS and height SDS minus target height SDS, which are considered typical indicators of the severity of GHD [30]. As expected by definition, stimulated GH and baseline serum IGF-I were also significantly different between both groups. P-values were provided, because in the severe GHD cohort the logical connector for GH and IGF-I is “or,” which may allow for substantial overlap of IGF-I or GH values with the moderate GHD cohort. Growth responses to rhGH treatment such as annualized HV, HV SDS, and change in height SDS were significantly greater in the severe GHD cohort, consistent with reports from other studies [31-34].

Using first-year annualized HV as a surrogate marker of rhGH responsiveness, it was evident that low stimulated maximum GH and low baseline serum IGF-I indicated good response to rhGH treatment, although the response decreased with increasing values of either parameter. This is consistent with reports from other studies [32-36]. In conclusion, these data confirm that the combined use of stimulated GH peak and baseline IGF-I with the applied cutoffs allows for identifying children with moderate GHD who may be candidates for ibutamoren testing and potentially treatment.

A relevant question is whether stimulated maximum GH and serum IGF-I are still significant indicators of severe or moderate GHD, if other contributing variables are included. A series of multivariate analyses of annualized HV from

### Table 5. Multiple regression analysis: final model for prediction of first-year height velocity

| Variable                                              | Parameter estimate | Standard error | Pr>|t| | Standardized estimate |
|-------------------------------------------------------|-------------------|---------------|---|----------------------|
| Intercept                                             | 10.6721           | 0.6349        | <0.0001 | 0                   |
| BL IGF-I > 30 µg/L (= 1)                               | −1.5835           | 0.2326        | <0.0001 | −0.2982              |
| Maximum GH ≥ 2 µg/L (= 1)                              | −0.7749           | 0.2881        | 0.0074  | −0.1117              |
| BL chronological age (y)                               | −0.2112           | 0.0406        | <0.0001 | −0.2131              |
| BL height SDS                                         | −0.1238           | 0.1250        | 0.3225  | −0.0520              |
| BL height SDS − target height SDS                     | −0.2404           | 0.0962        | 0.0128  | −0.1322              |
| BL body mass index SDS                                 | 0.2114            | 0.0525        | <0.0001 | 0.1730               |
| rhGH dose (mg/kg/wk)                                  | 7.0316            | 1.6539        | 0.0001  | 0.1763               |

Note that baseline serum IGF-I and stimulated maximum GH concentrations are included as categorical variables defined by the stated cutoffs.

Abbreviations: BL, baseline; GH, growth hormone; IGF-I, insulin-like growth factor-I; rhGH, recombinant human growth hormone; SDS, SD score.

*BL = baseline, at start of rhGH treatment.
514 children treated with rhGH indicated that age, rhGH dose, the difference between patient and target height SDS, BMI SDS, GH stimulation test result, and baseline IGF-I concentration were all independent indicators. These findings are consistent with reports from other studies in prepubertal children with idiopathic GHD [31,33-39]. Based on our analysis, prepubertal children aged 4 to 10 years with moderate idiopathic IGHD (stimulated GH ≥ 2 µg/L and IGF-I > 30 µg/L) have the potential to grow on average 8.3 cm/y in the first year of rhGH treatment. Data from the KIGS registry of patients with idiopathic GHD, not necessarily IGHD, including children at younger ages (1-10 years) and a GH cutoff of 5 µg/L have a similar mean first-year HV (8.6 cm/y) [36]. We hypothesize that this HV will be true both for treatment with daily rhGH injections or with once daily oral ibutamoren treatment.

An unexpected finding from this analysis was that catch-up growth decreased from >12 cm/y for patients with severe GHD to 8.3 cm/y for patients with moderate GHD, where moderate GHD was defined by peak GH to a stimulation test of 2 µg/L and serum IGF-I of 30 µg/L or greater.

A weakness of the GeNeSIS data was that the diagnosis of GHD was not subject to a stringent protocol: GH stimulation tests were not standardized, and GH measurements were not performed by the same assay in a central laboratory. This is typical for a “real-world” observational study. On the other hand, a strength of these data is that IGF-I measurements were performed in central laboratories or data were converted to central laboratory data. Regarding the primary objective of the current study, corroborating the peak GH and IGF-I cutoff values of the twin study [17], another weakness may be that the mean rhGH doses were different. The dose-dependence of GH treatment has in fact been demonstrated in various studies [40,41], but its role may only be moderate [42]. The mean dose in the twin study was 0.30 mg/kg/week (maximum approved US dose) whereas in the current study it was 0.20 mg/kg/week. This difference may cause slightly different growth responses. In this context, however, it should be kept in mind that dose-response relationships in biology commonly follow logit laws resulting in S-shaped curves (effect linear vs dose logarithmic). Therefore, a 30% decline in dose may have only a moderate effect and is unlikely to affect the study objective.

In conclusion, baseline IGF-I and stimulated GH alone or together are significant indicators of the degree of pediatric GHD, independent of other markers. In children with idiopathic IGHD, their combined use with cutoffs >30 µg/L for IGF-I and ≥2 µg/L for GH are predictive enrichment markers to segregate HV responses to rhGH therapy and can identify patients with moderate GHD who qualify for oral GH secretagogue testing and treatment. Notably, children with the most severe GHD grow at highest HV in response to rhGH treatment. Our working hypothesis is that children with moderate GHD will grow at similar rates in response to either daily injections of rhGH or oral ibutamoren.

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**Additional Information**

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**Data Availability:** Restrictions apply to the availability of some or all data generated or analyzed during this study to preserve patient confidentiality or because they were used under license. The corresponding author will on request detail the restrictions and any conditions under which access to some data may be provided.

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