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Identification of factors associated with good response to growth hormone therapy in children with short stature: results from the ANSWER Program®

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Abstract

Objective: To identify factors associated with growth in children on growth hormone (GH) therapy using data from the American Norditropin Studies: Web-enabled Research (ANSWER) Program® registry.

Methods: GH-naïve children with GH deficiency, multiple pituitary hormone deficiency, idiopathic short stature, Turner syndrome, or a history of small for gestational age were eligible (N = 1,002). Using a longitudinal statistical approach, predictive factors were identified in patients with GHD for change from baseline in height standard deviation score (ΔHSDS) following 2 years of treatment.

Results: Gradual increases in ΔHSDS over time were observed for all diagnostic categories. Significant predictive factors of ΔHSDS, ranked by significance were: height velocity (HV) at 4 months > baseline age > baseline HSDS > baseline body mass index (BMI) SDS > baseline insulin-like growth factor I (IGF-I) SDS; gender was not significant. HV at 4 months and baseline BMI SDS were positively correlated, whereas baseline age, HSDS, and IGF-I SDS were negatively correlated with ΔHSDS.

Conclusions: These results may help guide GH therapy based on pretreatment characteristics and early growth response.

Introduction

Treatment with exogenous growth hormone (GH) has become a well-accepted therapeutic option for children with growth failure. Since the availability of recombinant human GH (rhGH) in 1985, a wide range of conditions associated with decreased growth, including GH deficiency (GHD), Turner syndrome (TS), Noonan syndrome (NS), children born small for gestational age (SGA), Prader-Willi syndrome (PWS), idiopathic short stature (ISS), and SHOX (short stature homeobox) gene haploinsufficiency have been approved by the United States Food and Drug Administration (FDA) for treatment [1-4].

Treatment with GH has been demonstrated to increase both short-term growth and adult height in pediatric patients with a variety of different growth disorders [5-8]. However, considerable variability in response to this treatment has been reported across and within different diagnostic categories [9-11]. Such variability makes decisions about whether to treat with GH, when to begin treatment, and what dosing to use more difficult [12].

Reports from clinical trials and analyses suggest multiple factors that influence the response to GH treatment. Variables associated with better responses to GH treatment in patients with ISS include first-year growth response, younger age at start of treatment, the difference in height at the start of treatment from target height SD score (HSDS), and GH dose [13,14]. Additional factors may include underlying genetic conditions,
The maximal stimulated serum GH concentration was also recorded. Height and BMI SDS (z scores) were calculated according to the standard formulas provided by the Center for Disease Control and Prevention [32]. IGF-I SDS scores were calculated using a standard algorithm and reference values provided by Diagnostic Systems Laboratories, Inc. (Webster, TX, USA). Data were collected at baseline and at 4 months, 1 year, and 2 years of GH treatment. Data at 4 months were collected within a 1-month window and data at 1 and 2 years were collected within a 3-month window. To eliminate the potential of erroneous data having been entered, the following rules were used to remove patients from the analysis: lack of height information at baseline, 4 months, 1 year, or 2 years; baseline age 0 or > 18 years; baseline HSDS less than -5 or greater than +2; and baseline height < 35 cm or > 200 cm. Also, patients were excluded when key variables from baseline or subsequent values were deemed physically or chronologically implausible (3.77% of potential subjects were excluded according to this criteria).

**Regression model development**

In this study, a longitudinal statistical approach was used to identify factors that have significant predictive value for change in HSDS from baseline (AHSDS) in a regression model. AHSDS data collected following the first and second years of treatment were included in the model. A smoothing procedure was applied for the corresponding mean value curves for first-year HV and baseline age. Due to the limited number of patients in the MPHD, TS, SGA, and ISS diagnostic categories, regression analysis was only performed for patients with GHD. The curves presented were built using polynomial regression. The quadratic polynomial regression, under the assumption that the height SD is not a function of baseline age, provided a sufficient fitting, while higher terms (for example, cubic and fourth degree) were not statistically significant.

**Results**

**Baseline demographics**

The ANSWER Program registry (as of November 30, 2009) contained information for over 9,000 patients, of which 1,002 GH treatment-naive patients from selected diagnostic categories (GHD, MPHD, TS, SGA, and ISS) met the criteria for inclusion in this analysis. Baseline demographic characteristics for the subjects in this study by specific diagnostic category are summarized in Table 1. The study included longitudinal data for 698 patients with GHD, 71 with MPHD, 60 with TS, 50 with SGA, and 123 with ISS. Mean baseline ages were lower for MPHD (6.4 years), SGA (7.1 years), and TS (8.5 years) groups compared to those for GHD (10.9 years) or ISS (11.2 years). Baseline mean peak GH levels were lowest for patients with GHD and MPHD (5.5 and
3.6 ng/mL, respectively). Baseline mean GH dose (μg/kg/day) for the different diagnostic categories was the lowest for MPHD patients, consistent with their apparently greater degree of GH deficiency and associated GH sensitivity. For all diagnostic categories, the mean and median GH dose did not increase more than 0.007 mg/kg/day over the two years, indicating a very narrow GH dose change over this period.

### Height outcomes

The effects of GH treatment on ΔHSDS over 2 years of treatment are shown in Table 2. Gradual increases in ΔHSDS were observed over time and ranged between 0.15 (ISS) to 0.37 (MPHD) at 4 months, and 0.82 (TS) to 1.20 (MPHD) at 2 years, with the largest ΔHSDS observed at year 1 and year 2 in patients with MPHD and SGA. Annualized HV at 4 months was 13.6 cm/year for MPHD, and between 8.33 (TS) and 9.96 (SGA) cm/year for the other indications (Figure 1). Within each diagnostic category, mean annualized HV was the greater during the first year, and generally decreased during the second year. Mean annualized HV at 1 year was greatest for MPHD at 10.74 cm/year, and ranged between 7.97 (TS) and 9.57 (GHD) for the other indications.

### Regression analysis

Linear regression was performed on HSDS data for patients with GHD (Table 3). Variables significantly associated with ΔHSDS 1 and 2 years included HV at 4 months, and baseline age, HSDS, BMI SDS, and IGF-I SDS. The ranking of importance of predictive factors as related to ΔHSDS (as determined by the F value, the higher the more influential) was as follows: HV at 4 months > baseline age > baseline HSDS > baseline BMI SDS > baseline IGF-I SDS. HV at 4 months and baseline BMI SDS were positively correlated with ΔHSDS, while baseline age, HSDS, and IGF-I SDS were negatively correlated with ΔHSDS. Gender was less influential than the above factors (Table 3) and was not detected as statistically significant in this analysis.

### Table 1 Baseline demographics by diagnostic category

|        | GHD | MPHD | Turner | SGA | ISS |
|--------|-----|------|--------|-----|-----|
| n      | Mean (SD) | n | Mean (SD) | n | Mean (SD) | n | Mean (SD) |
| Gender |     |     |        |     |     |     |     |     |
| Male   | 543 | -3 | 53 | -3 | 0 | -3 | 33 | -3 | 91 | -3 |
| Female | 155 | -18 | 18 | -18 | 60 | -17 | 17 | -17 | 32 | -17 |
| Age    | 698 | 10.9 (3.46) | 71 | 6.4 (5.23) | 60 | 8.5 (4.17) | 50 | 7.1 (4.41) | 123 | 11.2 (2.88) |
| HSDS   | 698 | -2.2 (0.86) | 71 | -2.0 (1.36) | 60 | -2.5 (0.77) | 50 | -2.8 (0.97) | 123 | -2.3 (0.68) |
| IGF-I SDS | 605 | -2.5 (1.26) | 31 | -3.2 (1.54) | 34 | -2.0 (1.43) | 32 | -2.1 (1.53) | 114 | -2.2 (1.11) |
| BMI SDS | 681 | -0.1 (1.38) | 49 | 0.5 (1.84) | 55 | 0.5 (0.97) | 49 | -0.8 (1.35) | 123 | -0.8 (3.38) |
| Bone Age, yrs | 616 | 9.4 (3.31) | 39 | 8.0 (4.58) | 43 | 7.8 (3.45) | 44 | 6.1 (3.36) | 115 | 9.7 (2.93) |
| Peak GH, ng/mL | 606 | 5.5 (2.69) | 40 | 3.6 (3.03) | 5 | 12.5 (8.25) | 17 | 13.8 (10.95) | 97 | 15.2 (8.10) |
| GH dose, μg/kg/day | 697 | 45.9 (10.1) | 71 | 40.6 (11.2) | 60 | 51.9 (9.0) | 50 | 49.9 (13.5) | 123 | 46.1 (8.6) |

BMI, body mass index; GH, growth hormone; GHD, growth hormone deficiency; HSDS, height standard deviation score; IGF-I, insulin-like growth factor I; ISS, idiopathic short stature; MPHD, multiple pituitary hormone deficiency; SD, standard deviation; SDS, standard deviation score; SGA, small for gestational age.

### Table 2 HSDS and ΔHSDS by diagnostic category

|        | HSDS | ΔHSDS |
|--------|------|-------|
| HSDS   | n | Mean (SD) | n | Mean (SD) |
| GHD    | 698 | -2.22 (0.86) | - | - |
| Year 1 | 698 | -1.61 (0.83) | 698 | 0.61 (0.49) |
| Year 2 | 697 | -1.17 (0.88) | 697 | 1.06 (0.64) |
| MPHD   | 71 | -1.98 (1.36) | - | - |
| Year 1 | 70 | -1.13 (1.04) | 70 | 0.85 (0.76) |
| Year 2 | 70 | -0.79 (1.04) | 70 | 1.20 (1.04) |
| Turner | 60 | -2.49 (0.77) | - | - |
| Year 1 | 59 | -2.32 (0.82) | 59 | 0.18 (0.20) |
| Year 2 | 60 | -1.99 (0.86) | 60 | 0.50 (0.31) |
| SGA    | 50 | -2.76 (0.97) | - | - |
| Year 1 | 50 | -1.96 (0.93) | 50 | 0.80 (0.59) |
| Year 2 | 50 | -1.59 (1.00) | 50 | 1.18 (0.65) |
| ISS    | 123 | -2.31 (0.68) | - | - |
| Year 1 | 123 | -1.77 (0.69) | 123 | 0.54 (0.38) |
| Year 2 | 123 | -1.41 (0.79) | 123 | 0.90 (0.59) |

GHD, growth hormone deficiency; HSDS, height standard deviation score; ISS, idiopathic short stature; MPHD, multiple pituitary hormone deficiency; SD, standard deviation; SDS, standard deviation score; SGA, small for gestational age.
Analysis of the mean values was used to build smoothed curves for demonstration of the relationship between first-year ΔHSDS and baseline age (Figure 2A and 2B), and between first-year HV and baseline age (Figure 2C and 2D) in male and female patients with GHD. Generally, the curves demonstrate that younger baseline age is associated with greater ΔHSDS and HV in these patients. Similar curves were observed with male and female patients for both ΔHSDS and first-year HV.

Discussion
In this longitudinal study of GH treatment in patients with GHD, MPHD, TS, SGA, and ISS, HSDS improved over time. For patients with GHD, several variables were identified that correlated with growth response during the first and second years of GH treatment. HV at 4 months was the most significant predictor of ΔHSDS observed in the first 2 years of GH treatment. This observation that 4-month HV was such a strong predictor is a novel finding, since many studies do not consistently report growth this early in the treatment cycle. Additional factors that were influential in predicting HSDS outcomes were ranked in order of importance: younger baseline age > lower baseline HSDS > higher baseline BMI SDS > lower baseline IGF-I SDS.

For the GHD patient population, age and baseline HSDS were important determinants of the response to GH treatment, as previously demonstrated [18,20]. However, other reports have also indicated additional significant factors, such as birth weight SDS and GH dose [20]. The present results also indicated that higher baseline BMI was positively correlated with the growth response to GH for patients with GHD. Birth weight SDS and weight SDS were shown to be correlated with growth response to GH in the Pharmacia Kabi International Growth Study, suggesting that the heavier the child was, the greater the expected growth response to GH treatment [20]. The impact of BMI in this study might reflect, at least in part, the importance of nutrition for optimization of outcomes in patients receiving GH [1,33].

In general, the results from this analysis are consistent with previously published results for specific patient populations. A prior prediction study in patients with TS indicated that first-year growth response to GH was

Table 3 Regression model for longitudinal ΔHSDS at year 1 and year 2 for patients with GHD (n = 698).

| Characteristic      | β Estimate | F Value | P Value |
|---------------------|------------|---------|---------|
| Height velocity at 4 months | 0.0319     | 214.31  | < .0001 |
| Baseline age        | -0.0439    | 74.17   | < .0001 |
| Baseline HSDS       | -0.0776    | 29.11   | < .0001 |
| Baseline BMI SDS    | 0.0398     | 20.62   | < .0001 |
| Baseline IGF-I SDS  | -0.0245    | 5.74    | 0.0169  |
| Gender              | -0.0438    | 2.46    | 0.1175  |

BMI, body mass index; GHD, growth hormone deficiency; HSDS, height standard deviation score; IGF-I, insulin-like growth factor I; SDS, standard deviation score.
significantly influenced by weekly GH dose, chronological age, HSDS, body weight SDS, number of GH injections per week, and adjunctive oxandrolone treatment [26]. Predictors of the growth response over a longer duration of treatment (2-4 years) included HV during previous years, weekly GH dose, weight SDS, age, and oxandrolone treatment [26]. In SGA patients, results from one study found that first-year growth response to GH treatment was the most important predictor of second-year growth response [21]. Other variables that were significantly correlated with the growth response to GH included GH dose, weight and age at the start of treatment, and midparental HSDS [21]. Studies in the ISS patient population have identified additional factors as predictors of longer-term responses to GH, including baseline HSDS, GH dose, weight at the start of treatment, and first-year growth response [13,14]. It is important to recognize that this category may be the most heterogenous, with growth failure being a consequence of many different etiologies.

Specific results from other studies that are consistent with the present analysis, include the lack of gender effect on response to GH treatment. Analysis of results from the Pfizer Kabi International Growth Study database found no significant gender-related differences in effects of GH in HV or HSDS over 2 or 3 years of treatment [34]. In 8,018 patients with ISS in the National Cooperative Growth Study there was no significant effect of gender on first-year HV or first-year change from baseline in height SDS [35]. In a recent report, a large cohort of male and female patients with GHD, MPHD, TS, SGA, NS, and ISS from the ANSWER Program registry was used to assess gender-related differences in \( \Delta \)HSDS following 2 years of GH treatment. Results demonstrated increased \( \Delta \)HSDS in all patients, however, clinically relevant gender differences were not observed [36]. The importance of early timing for initiation of treatment from the present analysis is also consistent with previous findings. A National Registry of Growth Hormone Treatment report in the Netherlands that included 342 patients (diagnosis of GHD or a maximal GH response during provocation tests of less than 11 ng/mL) indicated that initiation of treatment before puberty resulted in a change from baseline in height SDS of 0.71 vs 0.58 for those who initiated treatment after puberty [19]. Results from the French registry of 2,852 patients with idiopathic GHD also indicated that prepubertal initiation of GH treatment was associated with significantly greater adult height gain [37].

Although in this study it is not known what proportion of patients
across the different diagnostic categories may have been in puberty, the mean baseline chronological and bone ages are consistent with the majority of patients being prepubertal, and this likely lessens the impact of puberty on the observed growth responses.

The different correlations between baseline age, HSDS, BMI SDS, and IGF-I SDS, with growth response over 2 years of treatment with GH, carry implications for clinical practice. The correlation of baseline age with ΔHSDS and HV in the patients with GHD further supports the initiation of GH at as young an age as possible to promote optimal growth. This concept is supported by results from another study that demonstrated a relationship between baseline age and first-year HV for patients with GHD, MPHD, and TS [15]. Several consensus statements endorse the use of GH treatment as soon as a diagnosis is made or growth failure is demonstrated for patients from several diagnostic categories [38-41]. The inverse relationship observed between baseline IGF-I and the two-year change in HSDS is consistent with an increased sensitivity to the effects of GH in patients who have a greater degree of GHD. In this non-interventional observational study, serum IGF-I was measured at a number of commercial laboratories reflecting routine clinical practice. IGF-I SDS was calculated using one formula which provided consistency to the analysis. This is also reflected in the finding that mean baseline IGF-I SDS in both the GHD and MPHD populations was lower than that observed in non-GHD patients. The positive correlation observed between baseline BMI SDS and ΔHSDS may emphasize the importance of nutrition in patients with growth failure [33,40]. An abnormally low BMI in pediatric patients may be a sign of malnutrition, which can also be associated with growth disturbances. In the end, the role of baseline age, HSDS, BMI SDS, and IGF-I SDS in the response of individual patients to GH therapy should all be considered for optimal management of short stature or growth failure.

Conclusion
The present results from a large patient cohort enrolled in the ANSWER Program registry demonstrate gradual increases in ΔHSDS over time for all diagnostic categories. For patients with GHD, greater HV at 4 months is the most significant predictor of ΔHSDS over the first 2 years of GH treatment, while gender did not have any influence.

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References
1. Grumbach MM, Bin-Abbas BS, Kaplan SL: The growth hormone cascade: progress and long-term results of growth hormone treatment in growth hormone deficiency. Horm Res 1998, 49(suppl 2):1-57.
2. Mazzanti L, Tamburroto F, Bergamaschi R, Scarano E, Montanari F, Torella M, Ballarinie C, Cicognani A: Developmental syndromes: growth hormone deficiency and treatment. Endocr Dev 2009, 14:114-134.
3. Harris M, Hofman PL, Cutfield WS: Growth hormone treatment in children: review of safety and efficacy. Paediatr Drugs 2004, 6:99-106.
4. Richmond E, Ragol AD: Current indications for growth hormone therapy for children and adolescents. Endocr Rev 2010, 18:92-108.
5. Bryant J, Baxter L, Cave CB, Milne R, Recombinant growth hormone for idiopathic short stature in children and adolescents. Cochrane Database Syst Rev 2007, CD004440.
6. Baxter L, Bryant J, Cave CB, Milne R: Recombinant growth hormone for children and adolescents with Turner syndrome. Cochrane Database Syst Rev 2007, CD003887.
7. Maiorana A, Cianfarani S: Impact of growth hormone therapy on adult height of children born small for gestational age. Pediatrics 2009, 124 e519-e531.
8. Crabe R, von HM, Engrand P, Chatelan P: Recombinant human growth hormone for children born small for gestational age: meta-analysis confirms the consistent dose-effect relationship on catch-up growth. J Endocrinol Invest 2008, 31:346-351.
9. Pasquono AM, Passeri F, Municchi G, Segni M, Pucarelli I, Larizza D, Bossi G, Severi F, Galasso C: Final height in Turner syndrome patients treated with growth hormone. Horm Res 1996, 46:269-272.
10. Ranke MB, Lindberg A: KIGS International Board: Height at start, first-year growth response and cause of shortness at birth are major determinants of adult height outcomes of short children born small for gestational age and Silver-Russell syndrome treated with growth hormone: analysis of data from KIGS. Horm Res Paediatr 2010, 74:259-266.
11. Sas TC, de Ridder MA, Wit JM, Rotteveel J, Oostdijk W, Reeser HM, Otten BJ, de Munck Koozer-Schrama SM: Adult height in children with growth hormone deficiency: a randomized, controlled, growth hormone dose-response trial. Horm Res Paediatr 2010, 74:172-181.
12. Hintz RL: Growth hormone treatment of idiopathic short stature: clinical studies. Growth Horm IGF Res 2005, 15(suppl A):S5-58.
13. Zucchinì S: Growth hormone use in the treatment of idiopathic short stature. Curr Opin Investig Drugs 2008, 9:396-401.
14. Ranke MB, Lindberg A, Price DA, Darendellì F, Bertsson-Wikland K, Wilton P, Reiter EO: Age at growth hormone therapy start and first-year responsiveness to growth hormone are major determinants of height outcome in idiopathic short stature. Horm Res 2007, 68:53-62.
Ranke MB: New paradigms for growth hormone treatment in the 21st century: prediction models.