Abstract

Objectives: Our aim was to analyze a large cohort of childhood onset GH deficiency (CO-GHD) adults from a unique adult center, in order to analyze their clinical management and to study the metabolic and bone status in relation to GHD and to the other pituitary deficits, and to evaluate these parameters during the long-term follow-up.

Design and methods: Observational retrospective cohort study on 112 consecutive CO-GHD adults transferred to our unit from 1st January 1994 to 1st March 2012. Evaluation of GHD in pediatrics and after transition was conducted following consensus guidelines. Data recorded from pediatric and adult files were GH doses, pituitary magnetic resonance imaging and function, and metabolic and bone status.

Results: Most patients presented with severe CO-GHD (64%) associated with other pituitary deficits (66%). CO-GHD was acquired in 56%, congenital in 33%, and idiopathic in 11% cases. Most patients (83%) stopped GH before transfer, at 16.3 years (median), despite persistence of GHD. Median age at transfer was 19.4 years. After transfer, GHD persisted in 101 patients and four of the 11 resolutive GHD were non idiopathic. IGF1 level was $<-2$ SDS in 70% of treated patients at transfer and in 34% of them after 3 years of treatment. Follow-up showed improvement in lipid profile and bone mineral density in severely persistent GHD patients under GH therapy. In multivariate analysis, the associated pituitary deficits seemed stronger determinant factors of metabolic and bone status than GHD.

Conclusions: This study raises concern about discontinuation of GH replacement therapy in pediatrics in severely persistent GHD patients and about the often insufficient dose of GH in the treatment of adult patients.

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Introduction

The optimal management of GH deficiency (GHD) during childhood, through adolescence, and all along adulthood is a challenge to pediatric and adult endocrinologists. In childhood, reduced linear growth is the primary consequence of untreated GHD and historically GH replacement therapy (GHRT) was stopped once final height was achieved. During the last decade, there has been a growing body of literature describing the negative impact of GHD on body composition, bone health, metabolic profile, cardiovascular risk, and quality of life, in older adolescents and in adults, pleading for a maintenance of GHRT in young adults with persistent GHD (1, 2). All of our knowledge on the long-term management of childhood onset GHD (CO-GHD) patients, once they become adults, relate to either large but multicentric studies (3, 4, 5, 6, 7, 8) or to monocentric studies on small cohorts (9, 10, 11). There is a lack of large monocentric studies addressing this issue.

Our adult endocrinological department is involved in the management of patients with rare endocrine disorders. We decided to study the consecutive CO-GHD patients transferred to our unit by all our collaborating pediatric departments, from 1994 to 2012. Our aim was to analyze the first large cohort of CO-GHD adults from a unique adult center, in order to evaluate their clinical management and to study their metabolic and bone status in relation to GHD and to the other pituitary deficits, and the evaluation of these parameters during long-term follow-up.
Subjects and methods

Patients

This observational retrospective cohort study took place in the adult department of endocrinology and reproductive medicine in Pitie-Salpetriere Hospital, Paris, France. The identities of all patients with CO-GHD transferred to our unit have been listed prospectively in a database since 2006. At that time, we retrieved, from the common database system of our hospital, the identities of the patients with CO-GHD transferred to us since 1994, and retrospectively added them to our database.

In our study, we included CO-GHD patients, transferred from any pediatric department to our unit, from 1st January 1994 to 1st March 2012, and who had at least one evaluation in our department.

CO-GHD was defined following the consensus guidelines (12). A peak GH concentration below 20 mIU/l after a provocative test was the threshold for defining GHD. Severe GHD was considered for peak GH concentration below 10 mIU/l and partial GHD for peak GH concentration between 10 and 20 mIU/l.

In pediatrics, provocative GH tests performed werearginine + insulin (n = 36), ornithine (n = 15), glucagon + betaxolol (n = 13), insulin tolerance test (ITT; n = 8), clonidine (n = 2), or l-dopa (n = 2). During childhood, most patients with a suspected isolated GHD had two provocative tests and most patients with a CNS pathology, a history of irradiation, or multiple pituitary deficits had one provocative test. We recorded in our database only the results from one stimulation test in pediatrics. For some patients (n = 16), the type of stimulation test used was unknown, but we had a pediatric report with the peak value of GH. In the other cases (n = 20), we had a pediatric report concluding to a GHD but we did not have the test results.

After transfer to our adult department, GHD was re-evaluated after a discontinuation of GH therapy of at least 1 month. The provocative test used was the ITT. During the transition period, a peak GH concentration below 15 mIU/l after ITT was considered as severe GHD (partial if peak GH between 15 and 20 mIU/l), following the 2005 consensus (13).

Other pituitary functions were also evaluated. Gonadotrophic deficiency was defined by an absence of spontaneous pubertal development at the physiologic age for puberty, and low estradiol (for girls) or testosterone (for boys) associated with low gonadotrophins (FSH and LH) after pubertal induction. Thyrotropic deficiency was defined by free thyroxine (FT_{3}) below normal range with TSH under upper normal range. Corticotropic deficiency was defined by plasma cortisol response to ITT or Synacthene 0.25 mg below 415 nmol/l, with normal or low ACTH. Lactotrophic deficiency was defined by prolactin under normal range, after puberty. The presence of a diabetes insipidus was assessed by a water restriction test of at least 12 h.

In all patients, a pituitary magnetic resonance imaging (MRI) was carried out to search for malformation, tumor, or infiltration of the hypothalamic–pituitary region.

Pediatric data

We went to the different pediatric departments in order to have direct access to the pediatric files. Files were reviewed for age, gender, results of GH stimulation tests and insulin-like growth factor 1 (IGF1) level, assessment of other anterior and posterior pituitary deficits, and MRI results of the pituitary. For patients for whom we could retrieve from the pediatric files, the information about GH treatment during childhood, age and dose at initiation, and pattern and duration of treatment was noted. Final and target heights (cm and SDS) were recorded.

Transition data

Age at last consult in the pediatric department and age at first consult in the adult department were noted. The same data as in childhood were recorded concerning the evaluation of pituitary function. If GH was resumed, dose and age were noted. If GH was stopped, reasons were specified.

Adult data

Data recorded in the adult department were pituitary function (IGF1 level and evaluation of other pituitary deficits), treatment with GH (dose, observance, and side effects), anthropometric measurements (weight, height, and BMI), metabolic parameters (lipid profile, fasting glycemia (FG), HbA1c, and body composition assessment by dual-energy X-ray absorptiometry (DEXA)), and bone metabolism parameters (osteocalcin and bone mineral density (BMD) assessment by DEXA).

We recorded GH doses and IGF1 levels at each visit (consultation or hospitalization). For all the other parameters, we recorded the results from the evaluations (EVAL) performed during hospitalizations. By definition, all patients had at least one evaluation during hospitalization in our department.

For patients with severe persistent GHD, at the first evaluation in our department (EVAL1), we compared the parameters of EVAL1 with those recorded at the second evaluation (EVAL2), the closest to 3 years after EVAL1. The patients with severe persistent GHD at transition, who have undergone EVAL1 and EVAL2, were divided into two groups: group 1 being the patients who received GH therapy (either continuously or discontinuously) between EVAL1 and EVAL2, and group 2 being the patients who did not receive GH therapy between EVAL1 and EVAL2.

We analyzed the recorded parameters based on IGF1 SDS values, and compared patients with IGF1 < −2
SDS with patients with IGF1 $\geq -2$ SDS. We also studied the influence, of all the recorded parameters, of the pituitary deficits associated with the GHD.

Hormonal assays

The level of IGF1 was measured using immunoradiometric assay (CIS Bio, Gif sur Yvette, France). Results were expressed in nanomoles per liter and in Z-score (SDS). As the IGF1 values were not normally distributed, IGF1 SDS calculation was done after log transformation (ln).

Statistical analysis

Descriptive statistics used numbers and percentages for qualitative variables and means and standard deviations or medians and inter-quartile intervals (Q1-Q3) for quantitative ones. Distributions of actual following times and maximum potential following times were estimated by the Kaplan–Meier method. Differences between two groups were tested using chi-square tests or Fisher’s exact tests for qualitative variables and two-sample Wilcoxon tests for quantitative ones. Paired comparisons between EVAL1 and EVAL2 distributions were tested using McNemar tests for qualitative variables and signed-rank tests for quantitative ones. Relationships between two quantitative variables were assessed using Spearman’s rank correlation coefficient tests. Multivariate analyses aimed at finding the relationships between each biological variable and the types of deficit, and were performed using linear models estimated using stepwise regression. All tests were two-sided, with a $P$ value $<0.05$ considered as significant. Computations were performed using the SAS V9 statistical package.

Results

Description of the cohort and pediatric data

We reviewed the files of 112 consecutive patients with CO-GHD transferred to our unit, with at least one evaluation during hospitalization. We did not have direct access to 20 pediatric files, but did not exclude them because we obtained the reports of pediatric hospitalization that concluded the presence of CO-GHD (all of these 20 patients were re-evaluated in the adult department and they had persistent GHD).

Patients (62% men and 38% women) mostly had severe GHD, usually associated with other pituitary deficits and due in a majority of cases to an acquired or a congenital cause (Table 1). Seventy-six percent of the congenital GHDs were due to pituitary malformation, and 48% of the acquired GHD were secondary to craniopharyngioma (data not shown).

Information about GH doses, age at initiation, and discontinuation, in the pediatric department, were available only for 96 patients (86%). Median age at initiation of GH was 10.0 years (6.3–13.9), median initial GH doses were $34.7 \mu g/kg per day (27.6–38.8)$ (after 5 years $31.4 \mu g/kg per day (24.4–34.5))

Eighty of the 96 patients (83%) stopped GH before being transferred to the adult department, at a median age of 16.3 years (14.7–17.3). Among those 80 patients, 39 had a dynamic re-evaluation of the somatotropic axis in the pediatric department, showing persistent GHD in 36 of them.

Median target height was 0.8 SDS ($-0.8$ to 0.7). Median final height was $-0.2$ SDS ($-0.9$ to 0.4). Median difference between median target and final heights was $-0.17$ SDS ($-1.0$ to 0.6). Final height was $< -2$ SDS in three women and four men. There was no difference in final height in either sex according to the etiology of the GHD.

Transition period

Median age at the last visit in the pediatric department was 19.1 years (16.1–27.3) and median age at first visit in the adult department was 19.4 years (18.4–21.3). Median time between last visit in pediatrics and first visit in the adult department was 4.6 months (2.6–6.7). Sex, etiology of GHD, year of transition, age at transfer, and time lapse between last visit in the pediatric department and first visit in the adult department were not correlated (data not shown).

Median time between the end of GH treatment in pediatrics department and first evaluation (EVAL1) in the adult department was 3.4 years (1.7–5.8) and all patients

Table 1 Degree of GHD and its association with other pituitary deficits, according to the etiology, at evaluation in the pediatric department.

<table>
<thead>
<tr>
<th>CO-GHD</th>
<th>Whole cohort ($n=112$)</th>
<th>Acquired ($n=63$)</th>
<th>Congenital ($n=37$)</th>
<th>Idiopathic ($n=12$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Degree of GHD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Severe</td>
<td>64% (72/112)</td>
<td>64% (40/63)</td>
<td>73% (27/37)</td>
<td>42% (5/12)</td>
</tr>
<tr>
<td>Partial</td>
<td>18% (20/112)</td>
<td>14% (9/63)</td>
<td>16% (6/37)</td>
<td>42% (5/12)</td>
</tr>
<tr>
<td>Missing data</td>
<td>18% (20/112)</td>
<td>22% (14/63)</td>
<td>11% (4/37)</td>
<td>16% (2/12)</td>
</tr>
<tr>
<td>Pituitary deficits</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Multiple deficits*</td>
<td>66% (74/112)</td>
<td>65% (41/63)</td>
<td>76% (28/37)</td>
<td>42% (5/12)</td>
</tr>
<tr>
<td>Isolated GHD</td>
<td>22% (25/112)</td>
<td>19% (12/63)</td>
<td>22% (8/37)</td>
<td>42% (5/12)</td>
</tr>
<tr>
<td>Missing data</td>
<td>12% (13/112)</td>
<td>16% (10/63)</td>
<td>2% (1/37)</td>
<td>16% (2/12)</td>
</tr>
</tbody>
</table>
who stopped GH in childhood did so for more than 6 months when they arrived in the adult department. Patients who did not discontinue GH before transfer were re-evaluated between 1 and 2 months after GH withdrawal, according to consensus guidelines.

**Follow-up in the adult department**

**GH status at EVAL1** During transition, at their first evaluation (EVAL1) in the adult department, 90 patients had an ITT to re-evaluate the somatotropic axis (Fig. 1). The 22 others did not: they had organic disease with at least three additional pituitary hormone deficiencies and IGF1 level $< -2$ SDS.

Among the 90 re-tested patients, 79 (88%) had a persistent GHD (71 patients had a peak GH below 15 mU/l, considered as severe GHD during the transition period, and eight patients had a peak GH between 15 and 20 mU/l, considered as partial GHD during the transition period). The remaining 11 patients had a restored normal response of GH secretion in response to ITT (GH peak $> 20$ mU/l): seven with an idiopathic GHD, two with an ectopic neurohypophysis (one with interrupted stalk), one with an empty sella, and one with Langerhans cell histiocytosis.

In all, we considered that 101 ($= 79 + 22$) patients had persistent GHD at the first evaluation in our department, comprising 93 ($= 71 + 22$) patients with an indication to resume GHRT according to consensus guidelines. The number of patients who received GH at some point during adult follow-up was 49.

At EVAL1, 33 patients with persistent GHD were treated with GH (currently or with a withdrawal of $<1$ month): 16 had not stopped since pediatrics and the other ones had restarted at first consult in the adult department, before first evaluation. In treated patients, median GH dose was 0.60 mg/day (0.4–1). IGF1 level was $<-2$ SDS in 23 (70%) of the treated patients (median IGF1 $= -3.3$ SDS ($-4.2$ to $-2.9$)) with median GH doses of 0.60 mg/day (0.50 to 0.85). IGF1 level was $>-2$ SDS in the other 10 (30%) patients treated (median IGF1 $= -0.8$ SDS ($-1.1$ to 0.9)) with the median GH doses of 0.6 mg/day (0.6 to 0.8). There was no significant difference between the doses of GH in patients with IGF1 $< -2$ SDS compared with patients with IGF1 $> -2$ SDS ($P = 0.6$). However, it is important to note that among the 13/33 patients with the highest GH doses ($> 1$ mg/day), six of them had very poor observance. In the 20/33 remaining patients (with GH doses $< 1$ mg/day), only three had observance issues.

At EVAL1, 97% of patients declared no side effect of GH treatment, but the information was lacking in 70% of the files. Fifteen patients had a history of seizures, and only one reported seizure aggravated under GH therapy.

Among the 68 GHD patients who were not GH treated at EVAL1, IGF1 was $<-2$ SDS in 75%, with a median IGF1 of $-3.9$ SDS ($-4.3$ to $-3.4$). IGF1 was $> -2$ SDS in the other 25%, with a median IGF1 of $-1.1$ SDS ($-1.7$ to $-0.6$).

**Evaluation of recorded parameters according to the treatment group** Sixty-three patients with severe persistent GHD (peak GH $< 15$ mU/l) at the first evaluation, at transition in the adult unit, had more than one evaluation in our department.

Median time between EVAL1 and EVAL2 was 2.5 years (1.6–3.3). For 39/63 patients, EVAL2 took place within 2–4 years after the transfer.

Patients in group 1 ($n = 45$) were treated with GH either continuously or discontinuously between EVAL1 and EVAL2. Patients treated discontinuously had a median time under GH between both evaluations of 40%, meaning that the median time under GH between

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**Figure 1** GH treatment and reevaluation of GH axis after transfer. Proportion of patients who stopped GH treatment in pediatrics and number of patients who had a dynamic reevaluation of the somatotropic function at their first evaluation in the adult department.

* Dynamic reevaluation, insulin tolerance test.
** Number of patients with a dynamic reevaluation of the somatotropic function after transfer.
*** None under GH therapy at the first evaluation in the adult department.
EVAL1 and EVAL2 was a little less than half of the time between the two evaluations. Patients in group 2 \((n=18)\) had an indication for GHRT, but did not wish to resume this treatment.

Table 2 shows the clinical practice of how many patients of each group had each of the recorded parameters at both the evaluations. We compared the two groups at EVAL1, at EVAL2, and the evaluation of all parameters within and between each group between EVAL1 and EVAL2. Results are shown in Table 3.

**GH status.** Doses of GH were obviously different between the two groups, at each hospitalization. In group 1, there was no significant difference in doses between EVAL1 and EVAL2; however, doses tended to decrease.

IGF1 SDS values were not significantly different between group 1 and group 2 at EVAL1; however, they were significantly different at EVAL2 \((P=0.005)\). In group 1, there was a rise in median IGF1 of 1.8 SDS, which was significant \((P=10^{-4})\). In group 2, IGF1 SDS values were not significantly different between EVAL1 and EVAL2.

At EVAL1, IGF1 value was \(<2\) SDS in 70% of patients in group 1 and in 73% of patients in group 2 \((P=1)\). At EVAL2, IGF1 was <2 SDS in 34% of patients in group 1 and in 69% of patients in group 2 \((P=0.02)\). The difference in the proportion of patients with IGF1 <2 SDS in group 1, between EVAL1 (70%) and EVAL2 (34%) was strongly significant \((P=7 \times 10^{-4})\).

**Anthropometric parameters.** Patients in group 1 were significantly younger and taller than patients in group 2 at each evaluation. No one significantly grew between EVAL1 and EVAL2, but in group 1 patients tended to grow \((P=0.055)\). Weight and BMI were not different between the groups at each hospitalization; however, they rose significantly in both groups between EVAL1 and EVAL2. Although patients in group 2 tended to put on more weight than in group 1, the difference was not significant.

**Metabolic parameters.** Evaluation of total cholesterol (TC) showed a significantly difference between the two groups (it dropped by a median of \(-0.15\) mmol/l in group 1 and raised by a median of 0.3 mmol/l in group 2, \(P<0.05\)).

The same trend was observed for triglycerides (TG) and LDL-c, without reaching statistical significance, once adjusted for sex and BMI. No significant difference was found regarding the body composition once adjusted for sex and height (data not shown).

FG was higher in group 2 at each time, and difference between the two groups was almost significant at EVAL1. FG rose between EVAL1 and EVAL2 in both the groups, but it was not significant once adjusted for BMI. HbA1c was lower in group 1 than that in group 2 at EVAL2, but it lost significance once adjusted for BMI.

**Bone metabolism.** We observed a significant rise in spine and hip BMD in group 1 between EVAL1 and EVAL2. However, at each hospitalization there was no difference between the two groups and the modification of the BMD between the two hospitalizations was not different between the two groups. No statistical difference was found for osteocalcin (data not shown).

Other comparisons. We also compared all the parameters between EVAL1 and EVAL2, by dividing the patients with severe persistent GHD at transition and who had a second evaluation \((n=63)\) differently. On one hand, we compared patients always under GH vs patients never under GH vs patients taking GH discontinuously. On the other hand, we compared patients always under GH vs patients taking GH discontinuously pooled with patients never under GH. We found no significant difference for any of the studied parameters for any of these comparisons (data not shown).

**Comparison of the recorded parameters according to IGF1.** We compared all the parameters at EVAL1 and EVAL2 according to IGF1 SDS values, whether they were \(<-2\) or \(\geq -2\) SDS (data not shown). Surprisingly, doses of GH were not significantly related to IGF1 SDS values. None of the recorded parameters were significantly linked to IGF1 SDS values, apart from osteocalcin level at EVAL2, which was significantly lower in patients with low IGF1 (median: 22 ng/ml \((7–47)\) vs 35 ng/ml \((29–55)\), \(P=0.04\)). The sex ratio of patients with IGF1 <2 SDS was 1 at both evaluations.

**Influence of the associated pituitary deficits.** We studied the correlations of all the quantitative variables to the number of pituitary deficits associated to the GHD. Significant results are presented in Table 4. The number of pituitary deficits was strongly correlated between EVAL1 and EVAL2 \((r=0.92, P<10^{-4})\). At EVAL1, TC, TG, LDL-c, weight, BMI, and percentage of fat mass were positively correlated to the number of pituitary deficits.
insulin-like growth factor 1 (IGF1) at Eval 2 (P < 0.001). The corticotropic deficiency was found significantly linked to FGF at Eval 1 (P = 0.02), fat mass percentage at Eval 1 (P = 0.01), hip Z-score (P = 0.006, even though it was in the opposite direction than expected), and IGF1 at Eval 2 (P = 0.004). Thyreotropic deficiency was significantly linked to LDL-c at Eval 1 (P = 0.02); and diabetes insipidus was significantly linked to fat mass percentage at Eval 1 (P = 0.04).

Comparison of recorded parameters according to the etiology of GHD Among the 63 patients with severe persistent GHD, who had at least two evaluations, 44 had an acquired GHD and 15 a congenital

Table 4 Correlations of all the studied quantitative variables to the number of pituitary deficits associated with the GHD.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Eval 1</th>
<th>Eval 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>r</td>
<td>P</td>
</tr>
<tr>
<td>IGF1 SDS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>Weight</td>
<td>0.41</td>
<td>0.001</td>
</tr>
<tr>
<td>BMI</td>
<td>0.42</td>
<td>5 × 10^{-3}</td>
</tr>
<tr>
<td>Fat mass (%)</td>
<td>0.81</td>
<td>10^{-4}</td>
</tr>
<tr>
<td>TC</td>
<td>0.30</td>
<td>0.03</td>
</tr>
<tr>
<td>TG</td>
<td>0.34</td>
<td>0.01</td>
</tr>
<tr>
<td>LDL-c</td>
<td>0.39</td>
<td>0.02</td>
</tr>
<tr>
<td>HDL-c</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>FGF</td>
<td>−0.35</td>
<td>0.01</td>
</tr>
<tr>
<td>Z-score hip</td>
<td>0.58</td>
<td>0.02</td>
</tr>
</tbody>
</table>

TC, total cholesterol; TG, triglycerides; FG, fasting glycemia; r, correlation; NS, non significant.

Table 3 Comparison of recorded parameters according to the treatment group at both evaluations. For each studied parameter, only when the data were not available for all patients, we indicated the number of patients for whom they were available.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Eval 1</th>
<th>Eval 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Group 1</td>
<td>Group 2</td>
</tr>
<tr>
<td></td>
<td>Median (Q1–Q3) (n = 45)</td>
<td>Median (Q1–Q3) (n = 18)</td>
</tr>
<tr>
<td>GH dose (mg/day)</td>
<td>0.60 (0.4 to 1) (n = 15)</td>
<td>0 (n = 15)</td>
</tr>
<tr>
<td>IGF1 (SDS)</td>
<td>−3 (−3.8 to −1.5) (n = 15)</td>
<td>−3 (−4.2 to −1.9) (n = 15)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>19.3 (18.4 to 21.2) (n = 45)</td>
<td>21.8 (19.4 to 22.6) (n = 18)</td>
</tr>
<tr>
<td>Height (SDS)</td>
<td>0.2 (−0.8 to −0.5) (n = 15)</td>
<td>−0.8 (−2.2 to 0.2) (n = 15)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>72.8 (62 to 91) (n = 15)</td>
<td>65 (59 to 72) (n = 15)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>0.81 (0.6 to 1.49) (n = 41)</td>
<td>0.86 (0.59 to 1.80) (n = 15)</td>
</tr>
<tr>
<td>TG (mmol/l)</td>
<td>3.19 (2.5 to 3.66) (n = 31)</td>
<td>2.56 (2.29 to 3.88) (n = 15)</td>
</tr>
<tr>
<td>LDL-c (mmol/l)</td>
<td>3.90 (3.7 to 4.5) (n = 43)</td>
<td>4.45 (4 to 4.95) (n = 16)</td>
</tr>
<tr>
<td>FG (mmol/l)</td>
<td>5.3 (5.1 to 5.5) (n = 21)</td>
<td>5.3 (5.2 to 5.6) (n = 15)</td>
</tr>
<tr>
<td>HbA1c (%)</td>
<td>0.96 (0.84 to 1.08) (n = 32)</td>
<td>0.93 (0.84 to 1.0) (n = 15)</td>
</tr>
<tr>
<td>BMD spine (g/cm²)</td>
<td>0.94 (0.83 to 1.04) (n = 32)</td>
<td>0.83 (0.81 to 0.87) (n = 15)</td>
</tr>
<tr>
<td>BMD hip (g/cm²)</td>
<td>0.94 (0.83 to 1.04) (n = 32)</td>
<td>0.83 (0.81 to 0.87) (n = 15)</td>
</tr>
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</table>

EVAL1, first evaluation in the adult department (after transfer from the pediatric department); EVAL2, second evaluation in the adult department (the closest possible to 3 years after transfer); Group 1, patients who received GH between Eval 1 and Eval 2; Group 2, patients who never had GH between Eval 1 and Eval 2; TC, total cholesterol; TG, triglycerides; FG, fasting glycemia; BMD, bone mineral density. This table compares the studied parameters between treated (group 1) and untreated (group 2) patients at first evaluation after transfer (Eval 1) and at second evaluation (Eval 2) in the adult department. It also compares the evolution of these parameters between Eval 1 and Eval 2 in each group. The symbols indicate a significant difference (P < 0.05) for each comparison: *significant difference between group 1 and group 2 at Eval 1; †significant difference between group 1 and group 2 at Eval 2; ‡significant difference between Eval 1 and Eval 2 for group 1; and §significant difference between Eval 1 and Eval 2 for group 2.

aAdjusted on sex and BMI.

bAdjusted on BMI.
GHD. We compared all the qualitative and quantitative parameters at EVAL1 and at EVAL2, and their evaluation between EVAL1 and EVAL2, in acquired vs congenital GHD patients. The parameters which reached significance were the presence of a diabetes insipidus being more frequent in acquired GHD (none was observed in congenital GHD: \( P=10^{-5} \)), LDL-c being higher at EVAL1 in acquired than congenital GHD (median 3.22 (2.60–3.96) vs 2.29 (1.86–2.54), \( P=0.02 \)), and FG being higher at EVAL2 in congenital than in acquired GHD (median 4.7 (43–5.3) vs 4.2 (3.95–4.55), \( P=0.02 \)).

Follow-up During their follow-up in the adult department, 63% of patients came at least once a year and 76% of patients visited our unit within the last 2 years. Seventeen percent of patients were lost to follow-up 1 year after the transfer. Up to 10 years after the transfer, less than one-third of the patients were lost to follow-up (data not shown).

Discussion

This study shows how 112 consecutive CO-GHD adults are taken care of, around and after the transition period. The advantage of this study is that all patients have been transferred and followed-up in a unique adult department, though they came from different pediatric units.

Most patients (two-thirds) had severe GHD in childhood, associated with other pituitary deficits, which is in a close proportion to what has been observed in other studies (12). Idiopathic GHD is less frequent in our study than that in others (5, 11), but it is not surprising because in most cases these patients recover a normal somatotropic function (14) and thus are not transferred to an adult department.

Our patients were transferred around the age 19, which seems appropriate. Indeed, it is suggested that transfer should be organized at the time of another change in the adolescent life, for example, the end of school (15). This age is close to what was observed in other studies of transitioning CO-GHD patients (5, 11) and to the age considered as ideal by patients in an auto-questionnaire (16). In this study, age at transfer is independent from etiology, sex, or year of transfer. Patients treated with GH during young adulthood (group 1) are transferred at a younger age than patients with persistent GHD who are not treated with GH during adulthood (group 2), regardless of the etiology. This suggests that the pediatric team organizes transfer sooner for patients who do not discontinue GH in pediatrics or who agree to the idea of resumption.

Median time between last consult in the pediatric department and first consult in the adult department was short (4.6 months), showing that when transition was prepared, there was a continuity in care around transfer. After transfer, most patients were regularly followed, with less than one-third of patients lost to follow-up. The success of transition was mandatory for patients with chronic conditions, such as CO-GHD, in order to optimize long-term follow-up and medical outcome (17, 18).

There has been a lot of concern about the impact of discontinuation of GH during transition, even after reaching adult height, leading to adverse effects on lipid profile, body composition, other cardiovascular risk factors, and acquisition of peak bone mass (4, 10, 19). Several studies have shown benefit from not interrupting GHRT at this point in time (4, 6, 7, 9, 19, 20). In our cohort of CO-GHD patients, GHRT was discontinued before transfer in most patients, even though they had persistent GHD, raising concern about their full bone and body composition maturation. Some of them continued to be followed in pediatrics several years before transfer. These observations could be taken into account by pediatricians when deciding whether to stop GH at the end of adolescence or not, and when considering the most adequate time for transfer. However, we noticed differences in pediatric management according to the various departments from which our patients came.

Transition period is an appropriate time for reassessment of GH status, unless there is high likelihood of persistent GHD (1). Four patients with non-idiopathic GHD have recovered a normal response of GH secretion in response to stimulation, two of which having an etopic neurohypophysis, a finding which has already been described (21).

Patients under GHRT at first evaluation in the adult department (EVAL1) had a GH dose about 50% lower than the recommended pediatric dose (25 μg/kg per day) and close to the recommended doses around transition (between 12.5 and 25 μg/kg per day or even lower, depending on authors) (13, 14, 22, 23). The currently recommended doses during transition are possibly insufficient. Indeed, in our study 70% of treated patients at EVAL1 had IGF1 values \(<-2\) SDS, independent of the sex. However, most of the patients with the highest doses of GH showed poor observance. It is most surprising that patients under GHRT with IGF1 values \(<-2\) SDS have a dose of GH similar to those of patients with IGF1 values \(\geq-2\) SDS, but this can be explained by the fact that three patients with low IGF1 have the highest doses prescribed and severe observance issues.

In patients with severe persistent GHD, IGF1 SDS values are significantly higher in the treated group (group 1) than in the non-treated group (group 2) at EVAL2, which is consistent with the fact that IGF1 level is correlated to GH treatment (6, 24), but this is not the case at EVAL1. The rise in IGF1 SDS values in the treated group between both evaluations is significant (\( P=0.01 \)), and there are significantly fewer patients with IGF1 values \(<-2\) SDS in the treated group at EVAL2 than at
EVAL.1. There is an improvement in the efficiency of the treatment in time, probably linked to the fact that doses of GH were more appropriate in older patients, and to the fact that patients who continue GHRT for years during adulthood are motivated and show better observance than in late adolescence.

In CO-GHD adult patients with persistent GHD, we observe a rise in weight and BMI with time, whether they resumed GHRT or not. Previous studies have found an effect of GH treatment on body composition in patients with CO-GHD or adult onset-GHD, but not on weight or BMI (25). However, we have not found a significant modification of body composition with time, but there is an important lack of data, especially in the untreated group.

This study made us aware of the number of missing explorations during adulthood in patients with persistent GHD from childhood. For instance, although all patients were followed in the same adult department, some of them did not have an evaluation of body composition or BMD by DEXA, especially patients who did not resume GHRT. Even more data were missing concerning the cardiovascular explorations (cardiac sonography, supra-aortic root Doppler), making it impossible to study those parameters. Metabolic parameters were more uniformly explored.

We observe an improvement in lipid profile over time only in GHD patients who receive GHRT during adulthood (group 1), which is in agreement with previously published studies (26, 27, 28). We did not find a difference in lipid profile according to IGF1 SDS values. Even though results are controversial, others studies have found that a rise in IGF1 under GHRT during transition is associated with a drop in TC and LDL-c (29). An explanation for the fact that we have found a significant variation in some parameters according to GH treatment status, but not according to IGF1 status (< -2 or ≥ -2 SDS), could lie in our methodology. Indeed, groups of treatment were elaborated using all the visits (consult and hospitalization), as the IGF1 status groups were determined from a unique dosage on day of evaluation, which might not be adequately representative of its long-term level.

FG tends to rise in GHD patients receiving GHRT during adulthood (group 1), but it also does in untreated GHD patients (group 2). On the other hand, HbA1c tends to be lower in group 1 than that in group 2 after a few years of the follow-up (EVAL2). The net effect of GHRT on insulin resistance is difficult to predict. GHRT lowers fat mass, and increasing IGF1 improves insulin sensitivity (30). However, GH also has direct insulin antagonistic effects in the liver and other tissues (31). A meta-analysis of placebo-controlled studies showed that GHRT was associated with a slight rise in both fasting glucose and fasting insulin levels (32).

The variations observed in the metabolic parameters are not explained by the etiology of the GHD, and in particular, no significant difference is found between patients with a history of craniopharyngioma and the other patients.

We observe a significant rise in spine and hip BMD in GHD adult patients under GHRT (group 1), which is in agreement with previously published data (8, 24, 33). GHRT has an overall anabolic effect on bone, but its effects are complex and the results are biphasic (6). Levels of osteocalcin are higher in patients with IGF1 values ≥ -2 SDS than that in patients with IGF1 < -2 SDS, which is consistent with the results from another study (4).

There is a strong correlation between the number of deficits associated with GHD, lipid profile, weight, BMI, and percentage of fat mass. A positive correlation has been found between time off GHRT during transition and TC, LDL-c, and TG in patients with non-idiopathic CO-GHD, who are more inclined to have associated deficits (4). It is possible that the presence of associated deficits increases the deleterious effects of GHD on lipid profile. A study compared patients with isolated and associated CO-GHD, and concluded that in GHD patients, associated deficits aggravate BMI, waist circumference, and lower IGF1 levels, but GHRT ameliorates these parameters in the same extent in isolated and associated GHD (34).

In multivariate analysis, the only variable significantly associated to lipid profile, weight, and BMI, is the gonadotrophic deficit. Deleterious effects on BMI or lipid profile in patients with either insufficient treatment for congenital hypogonadotropic hypogonadism or with anti-androgenic drugs for prostate cancer have been found (35, 36). Besides, estrogen substitution in women with gonadotropic and somatotropic deficits leads to hepatic resistance to GH action and thus to a decrease in IGF1 level compared with women without estrogen (37). In our study, women with gonadotrophic deficit received oral estrogen substitution therapy, thus potentially have lower levels of IGF1, explaining higher levels of TC and LDL-c. We also observe a significant association between corticotrophic deficit and a higher FG, as well as a higher fat mass percentage. We could hypothesize that, as previously reported, glucocorticoid replacement therapy increases morning insulin resistance and abdominal fat mass and weight (38). Finally, we found a positive correlation between thyreotropic deficit and LDL-c. Effects of hypothyroidism on lipid profile are well known. However, in our study, most patients with thyreotropic deficit are adequately supplemented in i-T4. We could hypothesize that because the conversion of T4 into T3 is reduced in untreated GHD patients (39), it is possible that some patients with normal FT4 levels are nevertheless insufficiently supplemented.

In conclusion, important reflections in pediatric and adult endocrinologists concerning the management of CO-GHD patients around and after transition are necessary because real-life practice is far from published recommendations. Impact of GHD on metabolism, body,
and bone compositions seems inferior to that of other pituitary deficits. More studies, particularly prospective, are needed to further address the issue of the care of CO-GHD patients during and after transition.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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