CLINICAL STUDY

Tumour recurrence and enlargement in patients with craniopharyngioma with and without GH replacement therapy during more than 10 years of follow-up

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Abstract

Objective: Most patients who have been treated for craniopharyngioma (CP) are GH deficient (GHD). GH replacement therapy (GHRT) may stimulate tumour regrowth; and one of the concerns with long-term GHRT is the risk of tumour progression. Therefore, the objective was to study tumour progression in CP patients on long-term GHRT.

Design: Case–control study.

Patients and methods: The criteria for inclusion of cases were: i) GHD caused by CP; ii) GHRT ≥3 years; and iii) regular imaging. This resulted in 56 patients (mean age at diagnosis 25 ± 16 years) with a mean duration of GHRT of 13.6 ± 5.0 years. As controls, 70 CP patients who had not received GHRT were sampled with regard to follow-up, gender, age at diagnosis and initial radiation therapy (RT).

Results: The 10-year tumour progression-free survival rate (PFSR) for the entire population was 72%. There was an association (hazard ratio, P value) between PFSR and initial RT (0.13, p < 0.001) and residual tumour (3.2, p < 0.001). The 10-year PFSR was 88% for the GHRT group and 57% for the control group. Substitution with GHRT resulted in the following associations to PFSR: GHRT (0.57, 0.17), initial RT (0.16, p < 0.001), residual tumour (2.6, < 0.01) and gender (0.57, 0.10). Adjusted for these factors, the 10-year PFSR was 85% for the GHRT group and 65% for the control group.

Conclusions: In patients with CP, the most important prognostic factors for the PFSR were initial RT and residual tumour after initial treatment. Long-term GHRT did not affect the PFSR in patients with CP.

Introduction

Craniopharyngiomas (CPs) are histologically benign suprasellar tumours with a strong tendency towards tumour progression and a capacity to invade the surrounding tissues (1). The primary tumour treatment for CP includes surgery and, in selected cases, radiation therapy (RT). In many cases, the tumour’s volume effect and/or the tumour treatment result in panhypopituitarism (2, 3).

Studies on hypopituitary CP patients have shown an increased standard mortality rate (SMR) (2.9–9.3), which was significantly higher than that in other groups of patients with hypopituitarism (4, 5, 6). The main causes of the increased SMR were cerebro- and cardiovascular accidents. Before the introduction of GH replacement therapy (GHRT), hypopituitarism in patients with a variety of causes including pituitary tumours was associated with a reduced quality of life (7, 8), a reduced bone mass (9) and an increased mortality, especially from vascular diseases (4, 5, 10). GHRT was introduced in the early 1990s. Studies have shown an improvement in quality of life, bone mass and a reduction in several vascular risk factors in patients with hypopituitarism receiving GHRT (11, 12, 13).

It has been shown that CP patients with GH deficiency (GHD) benefit to the same extent from GHRT as non-functioning pituitary adenoma (NFP A) patients with GHD (14). The more severe hypopituitarism and the higher mortality rate in CP patients compared with NFP A patients emphasise the need for optimal replacement therapy in CP patients.

Because of the known mitogenic effects of GH, it has been feared that GHRT may increase tumour recurrence or tumour enlargement in patients with a history of pituitary tumours (15, 16). Recent findings indicate that this is not the case in patients with NFP A (17, 18). It is well known that CPs have a high tumour progression rate – in fact after 10 years of follow-up around half of all CP patients who have not received RT develop tumour progression (1). The aim of this study

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was therefore to address the safety issue of whether long-term GHRT is associated with an increased frequency of recurrence or enlargement of CPs. A secondary aim was to obtain new long-term follow-up data on tumour progression in patients with CP.

**Subjects and methods**

The study was performed as an open prospective longitudinal case–control study of consecutive CP patients with hypopituitarism and GHD in order to study the effects of long-term GHRT at the Department of Endocrinology, Centre for Endocrinology and Metabolism at the Sahlgrenska University Hospital, Gothenburg, Sweden. Controls were sampled among CP patients not treated with GHRT at the Erlangen-Nuremberg University Hospital, Germany.

**Patients treated with GHRT**

Patients treated with GHRT were selected based on three inclusion criteria: i) hypopituitarism and GHD caused by CP; ii) GHRT for at least 3 years; and iii) imaging performed before commencing GHRT and after at least 2 years of treatment. A total of 56 patients with CP were eligible for inclusion in the study. The primary treatment of the tumours performed between 1969 and 2005 (the vast majority in the 1980s and 1990s) is presented in Table 1. The surgical approach was transcranial in 47 cases (89%) and transsphenoidal in six cases (11%). The percentage of patients receiving RT, either as part of the primary treatment (30 patients) or as treatment due to tumour progression (five patients), was 63%. When RT was a part of the primary treatment, the method used was conventional fractionated external radiotherapy except in two patients who were treated with stereotactic radiotherapy (Table 1).

The average observation period on GHRT was 13.6 ± 5.0 years. At final imaging, 43 patients (77%) were examined with magnetic resonance imaging (MRI) and 13 patients (23%) with computed tomography (CT). GHD was diagnosed using established criteria (11). The GH dose was titrated individually in order to maintain serum insulin-like growth factor 1 (IGF1) levels within the age-related reference range (19, 20). The average GH dose for the enrolled patients at the end of the observation period was 0.45 ± 0.12 mg/day (mean ± s.d.) for men and 0.72 ± 0.44 mg/day for women. At the end of the observational period, all patients had reached adulthood. The percentage of patients who had zero, one, two or three additional deficiencies beside GHD was 2, 5, 7 and 86%, respectively. Diabetes insipidus was present in 50 patients (89%). All patients were monitored and treated for additional hormonal deficiencies besides GHD.

Informed written consent was obtained from all patients. The study was approved by the Regional Ethics Review Board at the Sahlgrenska Academy, University of Gothenburg, Sweden.

**Patients not treated with GHRT**

A group of 70 CP patients not treated with GHRT was sampled from the hospital database at the Erlangen-Nuremberg University Hospital. The sampling process involved three mandatory criteria (diagnosis, type of initial treatment and status of GHRT) and four dispositive criteria (gender, age, age at diagnosis and duration of follow-up). The results of the sampling process are shown in Table 1.

The primary treatment of the CP tumours is presented in Table 1. The surgical approach was transcranial in 45 cases (71%) and transsphenoidal in 18 cases (29%). The percentage of patients for whom RT was part of the tumour treatment, either as part of the primary treatment (23 patients) or as treatment due to tumour progression (nine patients), was 46%. When RT was a part of the primary treatment, the method used was conventional fractionated external radiotherapy (Table 1).

**Table 1** Details of enrolled patients: gender distribution, age, age at diagnosis, observation period, primary treatment, and proportion of patients with a residual tumour after primary treatment.

<table>
<thead>
<tr>
<th></th>
<th>CP treated with GHRT (n=56)</th>
<th>CP not treated with GHRT (n=70)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender (M/F; %)</td>
<td>57/43</td>
<td>49/51</td>
</tr>
<tr>
<td>Agea (years)</td>
<td>46.6±16.1 (22–75)</td>
<td>45.7±16.2 (6–76)</td>
</tr>
<tr>
<td>Age at diagnosisa (years)</td>
<td>25.1±16.4 (4–62)</td>
<td>32.3±16.9 (2–65)</td>
</tr>
<tr>
<td>Observational perioda (years)</td>
<td>13.6±5.0 (3–28)</td>
<td>13.4±7.8 (3–34)</td>
</tr>
<tr>
<td>Primary treatment n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Surgery alone</td>
<td>25 (45)</td>
<td>45 (64)</td>
</tr>
<tr>
<td>Surgery combined with RT</td>
<td>28 (50)</td>
<td>18 (26)</td>
</tr>
<tr>
<td>Cyst puncture alone</td>
<td>1 (2)</td>
<td>2 (3)</td>
</tr>
<tr>
<td>Cyst puncture combined with RT</td>
<td>2 (4)</td>
<td>4 (6)</td>
</tr>
<tr>
<td>RT alone</td>
<td>0 (0)</td>
<td>1 (1)</td>
</tr>
<tr>
<td>Residual tumour after primary treatment n (%)</td>
<td>16 (29)</td>
<td>33 (47)</td>
</tr>
</tbody>
</table>

CP, craniopharyngioma; GHRT, GH replacement therapy; RT, radiation therapy.

*Values are presented as mean ± s.d. (range).
The average follow-up period from the primary treatment to the last imaging examination was 13.4±7.8 years. At the final imaging, 52 patients (74%) were examined with MRI and 18 patients (26%) with CT. The percentage of patients who had zero, one, two or three other deficiencies than GHD was 13, 17, 34 and 36% respectively. Diabetes insipidus was present in 33 patients (47%).

Informed written consent was obtained from all patients.

**Methods: evaluation of tumour progression with and without GHRT**

Patients treated with GHRT were divided into two groups on the basis of their tumour status before commencement of GHRT. Patients with a known residual tumour were categorised at the end of the observational period as to the presence or absence of tumour enlargement. Patients without any visible residual tumour at baseline were categorised either as having a recurrence or no recurrence at the end of the observation period. Tumour status was evaluated by regular imaging of the sella turcica and its surroundings throughout the follow-up period. Tumour recurrence or enlargement was determined by comparison of the imaging at the end of the observation period compared with baseline, before commencement of GHRT. All tumour recurrence or enlargement, regardless of size or clinical relevance, resulted in categorisation as recurrence or enlargement at the time it was first detected on imaging. Tumour progression was defined as either tumour recurrence or tumour enlargement. Recurrence and enlargement were classified as being of clinical significance if any change in the clinical management was necessary. Change in clinical management was defined as intervention with cyst puncture, surgery or radiotherapy. All imaging was performed as part of a routine clinical surveillance programme, which was individualised based on age and tumour characteristics. Serum IGF1 concentration was determined by hydrochloric acid ethanol extraction RIA. IGF1 SDS were calculated as described previously (21).

Patients not treated with GHRT were evaluated in the same way as patients treated with GHRT, with the exception of the definition of the baseline, which was set to the time of the first tumour treatment.

**Statistical analysis**

All descriptive statistical analyses are presented as mean values and s.d. The mean of IGF1 values and SDS were compared between GHRT patients with and without a tumour progression by Student’s t-test after ascertaining the assumption of normality. Kaplan–Meier survival curves were obtained to describe the progression-free survival (PFS) of patients in Table 2. The log-rank test was used to compare tumour progression times between subgroups. A proportional hazard model was used to analyse the impact of the potentially influential variables of gender (male ( =1) vs female ( =0)), age at diagnosis, residual tumour after primary treatment, initial RT and GHRT. Hazard ratios (HRs), 95% confidence intervals (CIs) and P values are presented. A SAS macro (22) was used to calculate the adjusted survival probabilities including a 95% CI, stratified by GHRT. The level of significance was set at P < 0.05 (two-sided test). Statistical analyses were performed using SPSS 18.0 (Chicago, IL, USA), SAS 9.2 (Cary, NC, USA) and Matlab 7.9.0 (Natick, MA, USA).

**Results**

**All patients: long-term follow-up**

Of 126 CP patients, 39 (31%) developed tumour progression during the follow-up period. The PFS rates (PFSRs) at 10 and 15 years were 72 and 67% respectively (Table 2). Tumour progression was observed after primary tumour treatment with surgery alone in 32 patients, surgery combined with RT in four patients, cyst puncture alone in one patient and combined cyst puncture and RT in two patients. The
patients with tumour progression were subsequently treated with surgery in 18 cases, combined surgery and RT in 12 cases, RT alone in one case, combined cyst puncture and RT in two cases, cyst puncture alone in one case and expectation was considered adequate in five cases (Table 3).

The tumour progression outcomes were analysed using Cox regression, which showed that initial RT and the absence of residual tumour after primary treatment were independently associated with a significantly lower frequency of tumour progression whereas age at diagnosis and gender had no significant effect on the frequency of tumour progression. The results of the Cox regressions are shown in Table 4.

All patients were divided into four groups depending on whether they were primarily treated with RT (yes/no) and whether they had a residual tumour after primary treatment (yes/no). The 5-, 10- and 15-year PFSRs are presented in Table 2. In all patients who had received initial RT, the 10-year PFSR was 90% compared with 58% of those without RT as part of their primary treatment. The statistical analysis showed a significant difference in PFSR between the patients in the four subgroups (Fig. 1). The best long-term PFSR was seen in patients treated with initial RT without a residual tumour whereas patients not treated with initial RT but who had a residual tumour had the worst outcome.

When the progression rate for the subgroup of patients without initial RT and with no visible residual tumour was used as baseline, the estimated HRs (95% CI for HR; P value) for tumour progression were (adjusted for gender) 3.0 (1.5–6.1; <0.005) for patients without initial RT and a residual tumour; 0.1 (0.01–0.79; 0.03) for patients with initial RT and no residual tumour; and 0.4 (0.16–1.1; 0.08) for patients with initial RT and a residual tumour (Fig. 1B).

Table 3 Treatment targeting tumour progression in patients with and without GHRT depending on initial RT and the presence of residual tumour.

<table>
<thead>
<tr>
<th>Treatment of tumour progression (no. of patients with tumour progression/total no. of patients)</th>
<th>Surgery</th>
<th>Surgery combined with RT</th>
<th>RT</th>
<th>Cyst puncture</th>
<th>Cyst puncture combined with RT</th>
<th>Clinically insignificant tumour progression</th>
</tr>
</thead>
<tbody>
<tr>
<td>All GHRT patients (9/56)</td>
<td>4</td>
<td>4</td>
<td>1</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>No initial RT and no residual tumour (4/21)</td>
<td>3</td>
<td>1</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>No initial RT and residual tumour (4/5)</td>
<td>–</td>
<td>3</td>
<td>1</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Initial RT and no residual tumour (1/19)</td>
<td>1</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Initial RT and residual tumour (0/11)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>All non-GHRT patients (30/70)</td>
<td>14</td>
<td>8</td>
<td>–</td>
<td>1</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>No initial RT and no residual tumour (16/35)</td>
<td>10</td>
<td>5</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>No initial RT and residual tumour (9/12)</td>
<td>2</td>
<td>3</td>
<td>–</td>
<td>–</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Initial RT and no residual tumour (0/2)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Initial RT and residual tumour (5/21)</td>
<td>2</td>
<td>–</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>2</td>
</tr>
</tbody>
</table>

CP, craniopharyngioma; GHRT, GH replacement therapy; RT, radiation therapy.

*Clinically significant tumour progression was defined as tumour progression resulting in additional tumour treatment.

Table 4 Cox regressions analysing possible factors influencing the progression-free survival rates in craniopharyngioma patients.

<table>
<thead>
<tr>
<th>Cox regression analysis of possible factors influencing tumour progression</th>
<th>HR</th>
<th>95% CI</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All patients (n = 126)</td>
<td>0.13</td>
<td>0.05–0.33</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Residual tumour after primary treatment</td>
<td>3.2</td>
<td>1.6–6.2</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Age at diagnosis</td>
<td>1.0</td>
<td>1.0–1.0</td>
<td>0.40</td>
</tr>
<tr>
<td>Gender*</td>
<td>0.55</td>
<td>0.28–1.1</td>
<td>0.08</td>
</tr>
<tr>
<td>Evaluation of GHRT</td>
<td>0.57</td>
<td>0.26–1.3</td>
<td>0.17</td>
</tr>
<tr>
<td>Treated with GHRT</td>
<td>0.16</td>
<td>0.06–0.40</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Residual tumour after primary treatment</td>
<td>2.6</td>
<td>1.3–5.3</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Gender*</td>
<td>0.57</td>
<td>0.29–1.1</td>
<td>0.10</td>
</tr>
</tbody>
</table>

| Evaluation of GHRT by clinically significant tumour progression | 0.65 | 0.29–1.5 | 0.30 |
| Treated with GHRT | 0.12 | 0.04–0.36 | <0.001 |
| Residual tumour after primary treatment | 2.2 | 1.1–4.7 | <0.04 |
| Gender* | 0.72 | 0.35–1.4 | 0.35 |

GHRT, GH replacement therapy; RT, radiation therapy; HR, hazard ratio.

*Male coded 1 and female coded 0.

*Clinically significant tumour progression was defined as tumour progression resulting in additional tumour treatment.
Tumour progression in patients treated with GHRT

Of the 56 patients treated with GHRT, nine (16%) developed tumour progression. In patients treated with GHRT, the estimated 10-year PFSR was 88% (Table 2). All tumour recurrences or enlargements were clinically significant. In the nine cases, where tumour progression occurred, eight patients had initially undergone surgery alone whereas one patient had undergone surgery and RT (Table 5).

The influence of initial RT on tumour PFSR is presented in Table 5. Tumour treatments targeting tumour progression divided depending on initial therapy are presented in Table 3.

In all patients with GHRT, the mean of the last serum IGF1 concentration was $222\pm107$ mg/l. There was no significant difference in the serum IGF1 level ($P=0.35$) or the SDS ($P=0.15$) between patients with tumour progression ($266\pm138$ mg/l; IGF1 SDS $1.23\pm2.3$) and patients without tumour progression ($214\pm100$ mg/l; IGF1 SDS $0.10\pm2.1$).

Tumour progression in patients not treated with GHRT

Of the 70 patients not receiving GHRT, 30 (43%) developed tumour progression. The 10-year PFSR was 57% for patients not treated with GHRT (Table 2). Five out of 30 patients developed clinically insignificant tumour progression. In the cases where tumour progression occurred, the initial tumour treatment was surgery alone in 24 patients, combined surgery and RT in three patients, cyst puncture in one patient and combined cyst puncture and RT in two patients (Table 5).

The effect of initial RT on tumour PFSR for patients not treated with GHRT is shown in Table 5. Tumour treatments targeting tumour progression divided depending on initial therapy are presented in Table 3.

Comparison of tumour progression in patients with and without GHRT

The 10-year PFSRs in patients treated with and without GHRT were 88 and 57% respectively. A Cox regression analysis showed that initial RT and absence of residual tumour after primary treatment were significantly associated with higher PFSR whereas treatment with GHRT and gender did not affect the PFSR (Table 4). Adjusted for these uneven distributed factors, the 10-year PFSRs were 85% in patients treated with GHRT and 65% in patients not treated with GHRT (Fig. 2).

When considering only patients with a clinically significant tumour progression (i.e. needing further tumour treatment), the 10-year PFSRs were 88% for the patients treated with GHRT and 63% for patients not treated with GHRT. Cox regression analysis showed that initial RT and absence of residual tumour after primary
treatment were associated with significantly higher clinical PFSRs where there was no residual tumour (23, 24). Adjusted for these uneven distributed factors, the 10-year clinical PFSRs were 85% in GHRT patients and 69% in non-GHRT patients.

No associations between tumour progression or clinically significant tumour progression and GHRT were found.

**Discussion**

This study on 126 CP patients with and without GHRT, who had a mean follow-up of over 13 years, provides new long-term follow-up data and allows for an evaluation of GHRT effect on tumour progression rate. In the current study, the overall 10- and 15-year PFSRs were 72 and 67% respectively. The strongest predictors of tumour PFS were initial RT and tumour status after the primary tumour treatment, whereas long-term GHRT had no significant impact on tumour progression.

The tumour progression rate in patients with CPs is high but differs widely in earlier studies mainly depending on the extent of surgical removal and if initial RT was used. Reports with follow-ups of 10 years or more in CP patients are scarce, but earlier papers have reported a 10-year PFSR of 41% in CP patients not treated with initial RT who had a residual tumour or 42–81% where there was no residual tumour (23, 24, 25, 26). In the current study, the 10-year PFSRs in CP patients not primarily treated with RT and with or without a residual tumour after the initial treatment were 32% and 66% respectively. The addition of RT to the primary treatment has been reported to increase the 10-year PFSRs in CP patients with and without a residual tumour after the initial tumour treatment to 84–90% and almost 100% respectively (23, 24, 27). The corresponding results in our study were 82 and 100%.

The two most important predictive factors, according to this study, for tumour PFSR were initial RT and the absence of a residual tumour after primary tumour treatment. Although the tumour progression rate is considerably lower in patients in whom total tumour removal has been ensured, there is a risk that an aggressive surgical approach would cause extensive hypothalamic injury with a poorer metabolic and neuropsychological outcome (28). Therefore, in many patients, the goal is not to achieve a radical tumour removal but to perform tumour debulking and, if needed, combine with adjuvant RT in order to minimise tumour progression.

Initial treatment with RT was the single most important factor for preventing tumour progression. However, RT is also associated with an increased risk of developing additional pituitary hormone deficiencies (29, 30), an increased risk of optic neuropathy (29, 30), secondary brain tumours (31) and cerebrovascular accidents (32, 33). In many centres, the practice of RT has therefore been used more restrictively during later years, although its use is still frequent in CPs due to their high risk of tumour progression. This study does not add any information on the risk–benefit of RT except for the aspects related to long-term tumour progression rates.

No correlation between age at diagnosis of CP and tumour progression was found in this study, which is in agreement with earlier studies (23, 27, 34). Thus, our results are in line with the previous reports and add 2 Cox regression of progression-free survival rates adjusted for initial RT, residual tumour and gender.  

*Clinically insignificant tumour progression was defined as tumour progression not resulting in additional tumour treatment.
further to the limited available data on the long-term follow-up of tumour progression in CP patients.

Our study does not lend support to the concern that long-term GHRT increases the risk of tumour recurrence or tumour enlargement in CP patients. There are only a few previous studies comparing the tumour progression rate in CP patients with and without GHRT. Karavitaki et al. (35) reported no influence of 6 years of GHRT on tumour progression rate in 32 GH-treated patients. The study had, however, a large difference in the frequency of initial RT between the GH-treated and non-GH-treated patients. In a recent study from Müller et al. (36), there was no difference in tumour progression between 54 CP patients treated with GHRT and 60 CP patients without GHRT after a mean follow-up of 2.8 years. In the current study, after adjustment for initial RT, residual tumour after primary treatment and gender, the 10-year PFSRs were 85% for the GHRT-treated patients and 65% for patients not treated with GHRT. Furthermore, circulating levels of IGF1 in the GHRT group, which are closely correlated with the biological effect of GH, were not different in CP patients with and without tumour progression. Our results, therefore, support the previous reports that physiological doses of GH as replacement therapy does not influence the rate of tumour progression in patients with CP.

Selection bias is a possible limitation of this trial as all GHRT patients were treated and followed at one university clinic, whereas the patients without GHRT were treated at another centre. The distributions in age at diagnosis and in the year of diagnosis were similar for both centres. A potential selection bias would occur if CP patients with GHD from the Gothenburg clinic were excluded from GHRT because of a residual tumour with a high risk of tumour progression. This is unlikely as we only found two patients with GHD who had not been included in the original GHD cohort from the clinic in Gothenburg, due to lack of medical records and due to their being lost to follow-up. Childhood-onset and adult-onset CPs have not been analysed separately, as age at diagnosis did not affect the rate of tumour progression in this study or in earlier studies (23, 27, 34) (see Table 4).

The management of patients with CP is a clinical challenge both in terms of the best choice of primary tumour treatment and the long-term management of the endocrine and metabolic consequences of the tumour itself and its treatment. In summary, our study with 13 years of follow-up highlights the effect of initial RT and the absence of a residual tumour on tumour progression rates in CP patients. In addition, long-term GHRT and serum IGF1 concentrations did not influence the risk of tumour recurrence or tumour enlargement. Based on our results from this large study with long-term follow-up, we conclude that GHRT does not increase the risk of tumour recurrence or tumour enlargement in CP patients.

Declaration of interest

Authors D S Olsson, M Buchfelder, K Wiendieck, N Kremenevskaja, B-A Bengtsson, K-E Jakobsson, M Jarfelt and A G Nilsson have nothing to declare. G Johannsson has received grant support from NovoNordisk and honorarium from NovoNordisk, Pfizer, Eli Lilly and Merck Serono.

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