Abstract

Objective: Investigate whether intervention with GH after tibial fracture enhances fracture healing.

Design: Randomised, double-blind, placebo-controlled study in 406 patients (93 women, 313 men, age: 18–64 years) with tibial fracture.

Methods: Patients were stratified by tibial fracture (open or closed) and allocated to placebo or GH treatment (15, 30 or 60 μg/kg daily, until clinically assessed healing or until 16 weeks post-surgery). Primary outcome was time from surgery until fracture healing and assessment of healing was done centrally and observer blinded. Patients reported for evaluation every 4 weeks until 24 weeks, and at 9 and 12 months.

Results: GH did not accelerate time to healing in the combined group of open and closed fractures. When separately analysing the closed and open fractures, a significant difference in time to healing was observed between treatment groups, exclusively in the closed fractures (P<0.05; subgroup analysis revealed that the 60 μg/kg group was significantly different from placebo). The relative risk of fracture healing for 60 μg/kg versus placebo during the 12 month was: all fractures, 1.16; 95% CI: (0.86; 1.57) (ns); closed fractures, 1.44; 95% CI: (1.01; 2.05; P<0.05); open fractures, 0.75; 95% CI: (0.42; 1.31) (ns). The estimated median number of days before fracture healing in closed fractures was 95 with 60 μg/kg versus 129 with placebo (95% CI: (94; 129) and (94; 249)) corresponding to approximately 26% decrease in healing time.

Conclusions: In the overall group of open and closed tibial fractures, no significant enhancement of fracture healing was observed with GH, whereas in closed tibial fractures, GH accelerated healing significantly.

Introduction

Growth hormone (GH) plays a crucial role in the maintenance of bone mass by regulating bone resorption and formation. A few studies on accidental hip fracture patients have looked potentially promising (1–3), and there is evidence from animal models and from in vitro studies that GH stimulates fracture healing (4–10). Thus, GH may have potential as a therapeutic option in enhancement of the healing of these fractures; however, no published data from clinical trials exist. The tibia is the most commonly fractured long bone and a fracture that tends to heal more slowly than other long bones (11, 12). Tibial fractures are frequently associated with complications, such as delayed unions, non-unions and infections. Despite improvements in surgical techniques (13), treatment of these fractures remains a challenge in orthopaedic surgery. The prevalence of impaired healing depends on multiple factors and ranges from <5 to 100% (13–15). A reduction in healing time or complications would relieve some of the burden that tibial fractures impose on patients and society (16–19).

GH plays a central role for several metabolic functions including remodelling of the skeleton (20, 21). In adults, bone remodelling is regulated by a complex interaction of circulating GH, insulin-like growth factors (IGFs), IGFBP-proteins (IGFBPs), locally produced IGFs, IGFBPs and other growth factors, acting in an autocrine and paracrine way (22–24). Stimulation of osteoblasts is believed to be the main mechanism of action (22), but angiogenesis, preservation of muscle tissue and function and improved wound healing may also be involved (25, 26).

The purpose of this study was to investigate the efficacy and safety of GH in the treatment of tibial fractures. The primary objective was to investigate the efficacy of GH as compared with placebo on fracture healing in tibia fractures treated surgically with...
intramedullary nailing and a secondary objective was to investigate the safety of GH in patients with tibia fractures.

**Patients and methods**

**Patients**

In order to obtain 340 evaluable outcomes, it was planned to randomise 400 patients. The study population comprised all tibial fracture patients suitable for intramedullary nailing according to the standard of treatment of the medical centres participating in this study and fulfilling the inclusion criteria of the study protocol. The main inclusion criteria comprised: male and females aged ≥18 and <65 years, and intramedullary nailing surgery of a tibial fracture categorised as Tscherne type C1, C2, C3 (closed fractures) or Gustilo type I, II, IIIa (open fractures) (27, 28). Informed consent was obtained from each patient prior to study inclusion.

Exclusion criteria included acromegaly, open growth plate, chronic endocrine or metabolic disease including diabetes and severe obesity (body mass index (BMI) > 32.0), chronic liver disease, known or suspected malignant disease, glucosuria, pre-existing bone and/or soft tissue infection, severe head injury (Glasgow Coma Scale <8), patients in need of mechanical ventilation (except during surgery) or circulatory support, or any condition that the investigator believed would interfere with trial participation or evaluation. Patients above 65 years of age were excluded in order to reduce variability in this first RCT in tibial fracture patients.

The study was conducted in accordance with the guidelines of the Helsinki Declaration on human experimentation. The protocol was approved by local Institutional Review Boards and by local and national Ethics Committees, as appropriate, and conducted in accordance with the International Conference on Harmonisation guidelines for good clinical practice.

**Study design**

The patients were stratified by fracture (open or closed) and randomly allocated to either GH (Norditropin SimpleXx, Novo Nordisk, Gentofte, Denmark: 10 mg/cartridge) or placebo (3:1), at one of the three dose levels: 15, 30 or 60 µg/kg body weight. All GH doses administered were at a pharmacological level. The highest dose level was chosen from clinical experience showing this dose increases IGF-I levels distinctively and the lowest dose level was chosen to be significantly higher than the approximate average physiological daily production of GH in normal adults (29).

Investigators were instructed to use the lowest available patient number from the randomisation list generated by the sponsor (in blocks of 12 for each stratum). Treatment was double-blinded with regard to GH versus placebo. For each patient number, a sealed code was kept at the trial site, at the sponsor’s local office and at the sponsor’s international product safety office.

Patients reported for evaluation every 4 weeks until 24 weeks post-surgery, and at 9 and 12 months to document safety and efficacy.

**Treatment**

GH or placebo treatment was initiated within 3 days after intramedullary nailing surgery and continued either until clinically assessed healing or until 16-weeks post-surgery, whichever occurred first. Trial product was administered daily by s.c. injections. In order to minimise adverse events such as water retention, the patients were titrated up to their target dose over three weekly steps. In case of events during treatment, a dose reduction in steps of 25% was considered. A drug accountability check was performed by the study monitor to ensure that each patient had at least used 75% of the required number of cartridges and had been exposed to trial drug during at least 75% of the intended treatment period.

**Radiological and clinical assessments**

The primary endpoint was time from surgery until fracture has healed and radiographic assessment of healing status was performed centrally by a panel of three experienced, observer-blinded, orthopaedic specialists (Synarc, San Francisco, CA, USA). Each observer was blinded to the treatment regimen, the clinical status of the patients, and to the results of the other radiological readings. Fracture healing was defined by disappearance of the fracture lines and/or cortical bridging in three out of the four cortices viewed on the anteroposterior and lateral radiographs. X-rays of the injured tibia were taken pre- and post-surgery and at each visit.

The secondary endpoints comprised individual investigator (clinically) assessed healing, the number of secondary procedures related to the fracture (e.g. exchange nail and bone graft), serum IGF-I and IGFBP-3, routine haematology and other safety assessments (see below). Clinically, assessed healing was based on the individual investigator’s radiographic evaluation and physical examination (full weight bearing and no pain or movement at the fracture site).

**Hormone assays**

Blood collections were performed at baseline prior to surgery and at 4, 8, 12, 16, 20 and 24 weeks post-surgery. Each subject had blood samples collected at the same time of day. Serum IGF-I levels were determined by
commercially available IRMAs (Immunotech, Marseille, France). Intra- and inter-assay coefficients of variation were 6.3 and 6.8% respectively.

Serum IGFBP-3 was determined by commercially available IRMAs (Immunotech, Marseille, France). Intra- and inter-assay coefficients of variation were 6.0 and 9.5% respectively. Osteocalcin (OST) was measured using IRMA Osteo-Riact kit from Scherring (Gi1-sur-Yvette, France). Intra- and inter-assay coefficients of variation were 2.8 and 5.2% respectively. Serum CrossLaps (CTX) was measured by a one-step ELISA assay using the commercially available serum CrossLaps kit (Nordic Biosciences Diagnostics, Herlev, Denmark). Intra- and inter-assay coefficients of variation were 5.1 and 6.5% respectively. All biochemical analyses were performed at a central laboratory (MDS Pharma Services, Paris, France).

Safety assessment
At each trial visit, plasma glucose, blood pressure and weight were measured, while glycosylated haemoglobin (HbA1c) was recorded at baseline (before GH treatment), at the end of treatment visit (at completion of GH treatment) and finally at week 24 (≥8 weeks after completion of GH treatment). Plasma glucose and HbA1c were measured by routine automated methods. All adverse events during the trial period were recorded, evaluated and followed up until the patient had recovered, stabilised, recovered with sequelae or died. An independent Data Safety Monitoring Board (DSMB) evaluated safety data during and after the study.

Statistical analysis
All statistical analyses were based on the intention-to-treat population, defined as all randomised patients exposed to trial product (n=406). The primary endpoint ‘time from surgery until fracture has healed’ was interval censored and the primary analysis takes this into account using a semi-parametric stratified proportional hazards regression model for interval-censored data (30). The effect of treatment on healing time was evaluated by estimation of the relative risk corresponding to healing, comparing active treatment groups with placebo. The primary analysis was based on all data points from the entire 12-month study period. The exact time (number of days) since surgery was derived from the recorded visit dates. Relative risk >1 corresponds to acceleration of fracture healing in comparison with placebo. The test of no difference between treatment groups was performed using a likelihood ratio test. Clinically, assessed healing was analysed similarly.

The influence on treatment effect of age at baseline, country, gender, fracture stratum, type of fracture and nail insertion technique was investigated in the primary analysis model, by including the investigated variable and a treatment group by variable interaction term, and test for no interaction. The influence of secondary procedures was investigated by censoring at the time of the first secondary procedure in a separate analysis. All other analyses were conducted on data with no censoring at time of first secondary procedure. The frequencies of secondary procedures were analysed using a χ² test.

In closed fractures, the median numbers of days before fracture healing, based on the proportional hazards regression model, were estimated post hoc using the information from the estimated survival functions and point-wise confidence bands for the placebo and the 60 μg/kg group. Wald’s test was used for direct comparisons between the 60 μg/kg group and placebo. No correction for multiple testing was performed.

The estimated sample size was based on the assumption that two-thirds of the fractures would be closed and one-third open. Furthermore, it was assumed that approximately 60% of open fractures and 80% of closed fractures would be healed at week 24. With these assumptions, the aim was to detect difference at week 24 in open fractures between placebo (60% healed) and GH (80% healed) and in closed fractures between placebo (80% healed) and GH (95% healed). Using a two-group χ² test of equal proportions with a 0.05 two-sided significance level, 80% power for detecting an increase in healed fractures from 60 to 80% was reached, when the sample size in each group was 82 patients. Similarly, 80% power for detecting an increase in healed fractures from 80 to 95% was reached, when the size in each group was 88 patients. This sample size calculation is considered conservative, as the primary efficacy analysis is not based on the proportions of healed fractures at week 24 alone, but on all information from the interval-censored data during the entire 12-month study period.

Results
Characteristics of the patients
An overview of the trial is shown in Fig. 1. Four hundred and twelve patients were screened, of whom 406 were randomised and exposed to trial drug, and 368 (90%) completed the trial. Baseline characteristics across treatment groups were similar (Table 1). The population comprised patients aged between 18 and 64 years; 75% were males. Twenty-nine percent (29%) of the patients had an open tibial fracture.

Treatment
Little variation across visits was seen with regard to the mean dose prescribed for the patients at each dose level (15 μg/kg GH, 1.0–1.1 mg; 30 μg/kg GH, 2.1–2.2 mg; 60 μg/kg GH, 4.1–4.2 mg; the three placebo dose levels combined, 2.5–2.6 mg/kg). A successful dosage differentiation between treatment groups is indicated by the IGF-I and IGFBP-3 data.

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IGF-I levels increased markedly above physiological levels with GH treatment, especially with 60 µg/kg. OST and CTX increased significantly, exceeding the effect of bone fracture alone distinctively (Table 2). Eighty-five percent of the patients fulfilled the criteria for treatment compliance, as judged from the drug accountability check.

Primary outcome

The primary endpoint was time from surgery until fracture has healed and was assessed by radiographic evaluation until 12 months post-surgery. When analysing the 12-month study period (primary analysis including all primary endpoint data in closed and open fractures), no

![Patient flow diagram. All patients were included in the intention to treat (ITT) analysis, except one patient who withdrew before receiving any trial drug (placebo group). Protocol violators mainly comprised of patients who did not fulfil the drug accountability check. Withdrawn patients were included in the analyses with all completed visits.](image)

**Table 1** Patient characteristics at baseline.

<table>
<thead>
<tr>
<th></th>
<th>Placebo</th>
<th>15 µg GH/kg</th>
<th>30 µg GH/kg</th>
<th>60 µg GH/kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients randomiseda</td>
<td>100 (100%)</td>
<td>99 (100%)</td>
<td>99 (100%)</td>
<td>108 (100%)</td>
</tr>
<tr>
<td>Patients completed</td>
<td>91 (90%)</td>
<td>93 (94%)</td>
<td>85 (86%)</td>
<td>99 (92%)</td>
</tr>
<tr>
<td>Fracture stratum</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Open fractures</td>
<td>29 (29%)</td>
<td>23 (23%)</td>
<td>27 (27%)</td>
<td>37 (34%)</td>
</tr>
<tr>
<td>Closed fractures</td>
<td>71 (71%)</td>
<td>76 (77%)</td>
<td>72 (73%)</td>
<td>71 (66%)</td>
</tr>
<tr>
<td>Age (mean years (s.d.))</td>
<td>38 (12)</td>
<td>36 (12)</td>
<td>39 (12)</td>
<td>38 (12)</td>
</tr>
<tr>
<td>Weight (mean kg (s.d.))</td>
<td>76 (13)</td>
<td>72 (14)</td>
<td>71 (12)</td>
<td>73 (14)</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>18 (18%)</td>
<td>25 (25%)</td>
<td>29 (29%)</td>
<td>21 (19%)</td>
</tr>
<tr>
<td>Male</td>
<td>82 (82%)</td>
<td>74 (75%)</td>
<td>70 (71%)</td>
<td>87 (81%)</td>
</tr>
<tr>
<td>Smoker</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>48 (48%)</td>
<td>47 (47%)</td>
<td>43 (43%)</td>
<td>53 (49%)</td>
</tr>
<tr>
<td>Yes</td>
<td>52 (52%)</td>
<td>52 (53%)</td>
<td>56 (57%)</td>
<td>55 (51%)</td>
</tr>
</tbody>
</table>

*aExcluding one patient not exposed to trial product.*
significant difference between treatment groups was seen (*P = 0.47; Table 3). In the pre-planned supplementary analysis of primary endpoint, however, a significant interaction with the treatment effect of GH was seen for fracture stratum (open or closed; *P = 0.02) and furthermore, a significant difference between treatment groups was seen in closed fractures (*P = 0.03), with the greatest acceleration of fracture healing in the 60 µg/kg group. In contrast to closed fractures, no significant difference was seen between the treatment groups in open fractures (*P = 0.35). The relative risk of fracture healing for 60 µg/kg versus placebo during the 12-month study period was: all fractures, 1.16; 95% CI: (0.86; 1.57) (ns); closed fractures, 1.44; 95% CI: (1.01; 2.05; *P = 0.045); open fractures, 1.05; 95% CI: (0.42; 2.81) (ns). The proportion of healed fractures across all visits is shown in Fig. 2. A significant interaction with the treatment effect of GH was also seen for gender (*P = 0.01; the treatment seeming to be more efficacious in females in the 60 µg/kg group). No significant interactions were seen between treatment and age, nail insertion technique or country.

### Additional analyses on primary outcome

The *post hoc* analysis revealed that estimated median number of days before fracture healing in closed fractures was 95 with 60 µg/kg versus 129 with placebo (95% CI: (94; 129) and (94; 249)) corresponding to approximately 26% decrease in healing time.

### Secondary outcomes

#### Clinically assessed healing
No statistically significant difference in clinically (individual investigator) assessed healing was seen between any of the treatment groups in

Table 3 shows the relative risk of radiologically assessed fracture healing (GH/placebo) during the 12-month study period.
Figure 2 Radiologically assessed healing across visits in all tibial fractures, closed tibial fractures and open tibia fractures. The duration of GH/placebo treatment lasted until clinically assessed healing or 16-week post-surgery, whichever occurred first.
the analysis model and no significant interaction was seen with fracture stratum. However, a trend towards enhancement of healing during GH treatment (60 μg/kg) as compared with placebo was noticed in both open and closed fractures during the GH treatment period (Fig. 3). The individual investigator assessed healing occurred at an earlier time point compared with the centrally performed radiological assessment (Fig. 3). The median number of days from surgery until clinically assessed healing was 108 with placebo versus 87 with 60 μg/kg GH, indicating that full weight bearing was achieved earlier with 60 μg/kg GH.

**Secondary surgical procedures** A total of 102 secondary procedures planned after randomisation, categorised as most invasive (e.g. bone graft and exchange of nail) or less invasive (e.g. dynamisation and removal of screw), were performed during the trial: 21, 21, 25 and 35 procedures with placebo, 15, 30 and 60 μg/kg GH respectively. The frequency of these secondary procedures did not differ significantly between the treatment groups ($P = 0.24$). Half were performed in open fractures and the majority were categorised as less invasive. Additionally, 43 secondary procedures planned prior to randomisation were performed. No significant influence was seen by secondary procedures on the primary analysis when censoring at the time of the first secondary procedure, i.e. the relative risk for fracture healing was highest with 60 μg/kg GH, but no statistical significant difference was seen between any of the treatment groups (Table 3).

**Adverse events** Treatment emergent adverse events (TEAE) are summarised in Table 4. The proportion of patients experiencing any TEAE was lowest with placebo (35%) and highest with 60 μg/kg (58%). A TEAE was defined as any adverse event occurring from initial trial product administration until 7 days after last trial product administration. The most frequent TEAEs were infections and events related to fluid retention. The dose was reduced in 14 patients: 1, 3, 3 and 7 patients with placebo, 15, 30 and 60 μg/kg GH respectively. Dose reduction was done by steps of 25%.

The relation to trial drug was by the individual investigator rated as possible/probable in six TEAEs: one event in the 15 μg/kg group (extraskeletal ossification); one event in the 30 μg/kg group (allergic dermatitis) and four events in the 60 μg/kg group (pyrexia, acute cholecystitis, increased blood glucose and non-ketotic hyperglycaemic hyperosmolar coma). Two patients died during the follow-up period. Cause of death was pulmonary tuberculosis (60 μg/kg) and lung squamous cell carcinoma with metastasis (15 μg/kg; on re-examination, this carcinoma was detectable prior to the trial). A trial drug relation was considered unlikely by the investigator, the sponsor and the DSMB for both deaths.

Random plasma glucose and HbA1c values were similar across treatment groups and visits (Table 5). A total of 20 patients experienced increased HbA1c (>6.4%) and/or increased non-fasting plasma glucose (>11.0 mmol/l). Subsequent analyses revealed that five of these patients had increased levels at baseline. The increases were transient in all patients except for two patients in the placebo group and one patient in the 30 μg/kg group, the latter with a normal non-fasting plasma glucose (5.2 mmol/l) and a slightly increased HbA1c (6.9%) at the last visit.

No clinically relevant safety finding was observed with regard to blood pressure. Mean values were similar across visits and treatment groups (range: 78.1–80.7 mmHg (diastolic) and 123.0–129.9 mmHg (systolic)).
Discussion

This is the first report of a randomised, double-blind, placebo-controlled clinical trial investigating the effect of systemic GH treatment on the healing of tibial fractures. The main result from the study is that in the overall group of open and closed tibial fractures, no effect of GH was seen for the primary endpoint (time from surgery until fracture healed) based on data from the entire 12-month study period. However, when separately analysing the closed fractures, a significant difference in time to healing was observed between treatment groups, with the largest effect in the 60 mg/kg group. Tibia fractures treated with intramedullary nailing was included in this study since it is now the standard treatment of the majority of these fractures in most countries and is a technique which is being increasingly used as the gold standard of treatment in order to provide union earlier – making healing time the most important outcome. The estimated median number of days from surgery to fracture healing in closed fractures was 95 days in the 60 mg/kg GH group versus 129 days in the placebo group corresponding to a 26% decrease in healing time.

In the individual investigator assessment of healing, no statistically significant difference was observed between any of the treatment groups. However, a tendency towards enhancement of healing during GH treatment (60 mg/kg) as compared with placebo was noticed in both open and closed fractures during the GH treatment period and this pattern of accelerated healing during GH treatment with 60 mg/kg was consistent with the radiological assessment of healing (Fig. 3). It is speculated that a larger variability in the individual assessment of fracture healing may impact the probability of detecting significant differences compared to the centrally, observer-blinded radiographic evaluation of fracture healing. The investigators assessed healing at an earlier time point than the centrally performed radiographic evaluation in accordance with the previous results from a single-blinded trial on tibial fracture healing (31).

GH has crucial effects on longitudinal growth in childhood and adolescence and GH deficiency (GHD) in

Table 4 Treatment emergent adverse events (TEAEs).

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Placebo n (%)</th>
<th>15 μg hGH/kg n (%)</th>
<th>30 μg hGH/kg n (%)</th>
<th>60 μg hGH/kg n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients randomised</td>
<td>100</td>
<td>99</td>
<td>99</td>
<td>108</td>
</tr>
<tr>
<td>All TEAEs</td>
<td>35 (35%)</td>
<td>38 (38%)</td>
<td>40 (40%)</td>
<td>53 (58%)</td>
</tr>
<tr>
<td>Oedema lower limb</td>
<td>–</td>
<td>1 (1%)</td>
<td>1 (1%)</td>
<td>1 (1%)</td>
</tr>
<tr>
<td>Oedema upper limb</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Oedema peripheral</td>
<td>–</td>
<td>–</td>
<td>1 (1%)</td>
<td>1 (1%)</td>
</tr>
<tr>
<td>Oedema</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Peripheral swelling</td>
<td>–</td>
<td>2 (2%)</td>
<td>1 (1%)</td>
<td>6 (6%)</td>
</tr>
<tr>
<td>Swelling</td>
<td>–</td>
<td>–</td>
<td>1 (1%)</td>
<td>–</td>
</tr>
<tr>
<td>Abscess soft tissue</td>
<td>1 (1%)</td>
<td>1 (1%)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Localised infection</td>
<td>–</td>
<td>–</td>
<td>1 (1%)</td>
<td>–</td>
</tr>
<tr>
<td>Post-operative infection</td>
<td>–</td>
<td>–</td>
<td>1 (1%)</td>
<td>–</td>
</tr>
<tr>
<td>Post-operative wound infection</td>
<td>1 (1%)</td>
<td>2 (2%)</td>
<td>2 (2%)</td>
<td>2 (2%)</td>
</tr>
<tr>
<td>Wound infection</td>
<td>1 (1%)</td>
<td>2 (2%)</td>
<td>2 (2%)</td>
<td>3 (3%)</td>
</tr>
<tr>
<td>Wound sepsis</td>
<td>–</td>
<td>1 (1%)</td>
<td>1 (1%)</td>
<td>5 (5%)</td>
</tr>
<tr>
<td>Arthralgia</td>
<td>–</td>
<td>3 (3%)</td>
<td>3 (3%)</td>
<td>3 (3%)</td>
</tr>
</tbody>
</table>

Table 5 Summary of HbA1c and random plasma glucose levels during the study.

<table>
<thead>
<tr>
<th>Treatment (N/N)</th>
<th>Placebo mean (s.d.)</th>
<th>15 μg GH/kg mean (s.d.)</th>
<th>30 μg GH/kg mean (s.d.)</th>
<th>60 μg GH/kg mean (s.d.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline (N/N)</td>
<td>97/100</td>
<td>95/99</td>
<td>96/95</td>
<td>103/107</td>
</tr>
<tr>
<td>HbA1c (%)</td>
<td>5.23 (0.40)</td>
<td>5.17 (0.37)</td>
<td>5.15 (0.39)</td>
<td>5.17 (0.41)</td>
</tr>
<tr>
<td>Glucose (mmol/l)</td>
<td>6.45 (1.31)</td>
<td>6.38 (1.30)</td>
<td>6.35 (1.15)</td>
<td>6.63 (1.34)</td>
</tr>
<tr>
<td>End of treatment (N/N)</td>
<td>75/95</td>
<td>73/95</td>
<td>70/94</td>
<td>85/100</td>
</tr>
<tr>
<td>HbA1c (%)</td>
<td>5.34 (0.42)</td>
<td>5.38 (0.43)</td>
<td>5.40 (0.52)</td>
<td>5.41 (0.71)</td>
</tr>
<tr>
<td>Glucose (mmol/l)</td>
<td>5.51 (1.23)</td>
<td>5.49 (0.95)</td>
<td>5.69 (1.38)</td>
<td>5.86 (2.16)</td>
</tr>
<tr>
<td>Post-treatment (N/N)</td>
<td>85/93</td>
<td>93/96</td>
<td>90/94</td>
<td>94/100</td>
</tr>
<tr>
<td>HbA1c (%)</td>
<td>5.36 (0.51)</td>
<td>5.32 (0.40)</td>
<td>5.33 (0.46)</td>
<td>5.27 (0.43)</td>
</tr>
<tr>
<td>Glucose (mmol/l)</td>
<td>5.40 (0.89)</td>
<td>5.37 (0.74)</td>
<td>5.27 (0.90)</td>
<td>5.33 (0.76)</td>
</tr>
</tbody>
</table>

| Glucose > 11.0 mmol/l or HbA1c > 6.4% at any time (N) | 3 | 3 | 6 | 8 |

*Number of patient with HbA1c data/number of patients with plasma glucose data.

*Post-treatment HbA1c and plasma glucose data were measured at week 24 (≥8 after completion of GH treatment).
adults has been associated with bone loss and increased fracture risk (32, 33), whereas bone mass has been reported to be increased and fracture risk decreased in acromegaly (34, 35). It has therefore been hypothesised that GH may have a role in enhancement of fracture healing and a number of animal studies have demonstrated that GH stimulates bone formation (4–6), whereas others have shown no such response (36–38). This discrepancy has been explained by a variety of experimental designs used, the GH dosage and the animal species. However, in recent animal studies, it has been shown that systemically administered species-specific GH treatment induces a pronounced effect on bone formation. This was observed during both intramembranous bone formation and during secondary fracture healing, where bone formation is predominantly achieved by endochondral ossification (7–9, 39). Further, it has been demonstrated that local injection of GH enhances bone formation and mechanical strength of tibia fractures (10). Thus, the GH enhancement of closed tibia fracture healing observed in the present study is in agreement with recent findings from experimental animal studies.

An apparent difference in enhancement of tibia fracture healing between open and closed fractures was observed in the present study. However, the open fractures comprised a relatively small group in which healing time was more variable and as expected, the healing time was longer in open than in closed fractures. Thus, the GH treatment period in open fractures comprised a relatively shorter part of the full healing period than in closed fractures. It could be speculated that this might explain why GH seemed to accelerate healing of open fractures only in the early part of the study. Hence, it is hypothesised that GH treatment may accelerate the healing in open fractures similar to that observed for closed fractures if being administered during the entire period of healing and recovery. The apparent lack of effect in open fractures in the current study could also be related to an impeded opportunity for GH to reach osteoblasts when extensive soft tissue damage is present. Thus, further research is needed with an extended treatment period to reveal if GH may be beneficial in the more complicated open fractures.

A previous study reported a very moderate effect in recovery of elderly patients with hip fractures receiving GH treatment, and this moderate effect might be ascribed to the short treatment duration (6 weeks) and the relatively low GH dose (i.e. 20 μg/kg per day) administered (1). In the present study, no significant enhancement of fracture healing effect was observed in open or closed fractures with 15 or 30 μg/kg, although an increase was observed at these dose levels with regard to levels of IGF-I and markers of bone remodelling. Resistance to GH action has previously been described for catabolic and surgical trauma patients (40), and it is likely that GH resistance also is present in patients with tibial fractures. Apart from fracture healing, the effect of the 60 μg/kg dose was also distinctively more pronounced on levels of IGF-I and OST, as compared with the lower dose groups, and we therefore anticipate that 60 μg/kg per day is the lowest effective dose to enhance tibial fracture healing.

In addition to fracture stratum, a significant treatment interaction effect was observed for gender. It is well known that IGF-I levels are highly dependent on age, gender and sex hormone status. However, in the present study, no gender difference in circulating IGF-I levels was observed in response to the same doses of GH per kg body weight (data not shown). Clinical experience with GH administration in healthy adults suggests that a possible gender interaction would be expected to have been in favour of males and not females, as GH have been shown to induce a lower IGF-I response in females compared with males (41). In this study, the number of females was relatively low and thus it is not possible to draw any conclusion regarding gender interaction with treatment response. However, a possible gender dimorphism with regard to healing of tibial fractures cannot be ruled out.

Considering the pharmacological GH dose levels administered to the adult tibial fracture population in this study, few adverse events were observed and no significant safety issues emerged in this study. Further to the safety evaluation, the frequency of the secondary surgical procedures did not differ between the treatment groups. The proportion of patients experiencing any TEAE was lowest with placebo and highest with 60 μg/kg. However, the most frequent adverse events were consistent with those observed in patients with tibial fractures (infections) or observed during GH treatment in adults (transient events of swelling, oedema and arthralgia). This is of importance as previous studies investigating the effect of high doses of GH on the outcome of critically ill adults reported increased relative risk of death (42). A likely reason why no serious safety issues emerged in this study is that the patients were relatively healthy except for having a tibial fracture, and they were in no need of mechanical ventilation or circulatory support. Thus, no new GH safety issues were identified in the current study. However, based on our experience and in accordance with generally accepted guidelines (43), it seems prudent to monitor markers of glucose homoeostasis in all patients treated with GH – in particular those vulnerable to induction of pre-diabetes, e.g. patients with increased baseline values of measures of glucose homoeostasis and elderly.

In conclusion, the results demonstrate that in the overall group of open and closed tibial fractures, no significant effect of GH on time to healing was identified, whereas in closed tibial fractures comprising the vast majority of tibial fractures, GH accelerated healing significantly during the 12-month study period. The GH-induced enhancement of healing may be of potential benefit in patients with closed tibial fractures.
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