Growth hormone treatment in Prader-Willi syndrome patients: systematic review and meta-analysis

Caroline de Gouveia Buff Passone 1,2, Ruth Rocha Franco,3 Simone Sakura Ito,3 Evelinda Trindade,3 Michel Polak,2 Durval Damiani,3 Wanderley Marques Bernardo3

ABSTRACT

Background Growth hormone (GH) treatment is currently recommended in Prader-Willi syndrome (PWS) patients. Objectives To evaluate the impact (efficacy and safety) of the use of recombinant human GH (rhGH) as a treatment for PWS. Method We performed a systematic review and, where possible, meta-analysis for the following outcomes: growth, body mass index, body composition, cognitive function, quality of life, head circumference, motor development/strategy, behaviour and adverse effects. We included all PWS patients, with all types of genetic defects and with or without GH deficiency, who participated in rhGH studies performed in infancy, childhood and adolescence, that were either randomised controlled trials (RCTs) (double-blinded or not) or non-randomised controlled trials (NRCTs) (cohort and before and after studies). The databases used were MEDLINE, Embase and Cochrane Central. Results In 16 RCTs and 20 NRCTs selected, the treated group had an improvement in height (1.67 SD scores (SDS); 1.54 to 1.81); body mass index z-scores (−0.67 SDS; −0.87 to −0.47) and fat mass proportion (−6.5% SDS; −8.46 to −4.54) compared with the control group. Data about cognition could not be aggregated. Conclusion Based on high quality evidence, rhGH treatment favoured an improvement of stature, body composition and body mass index, modifying the disease’s natural history; rhGH treatment may also be implicated in improved cognition and motor development in PWS patients at a young age. Ethics and dissemination The current review was approved by the ethical committee of our institution. The results will be disseminated through conference presentations and publications in peer-reviewed journals. PROSPERO registration number CRD42019140295

INTRODUCTION

Prader-Willi syndrome (PWS) is a rare genetic disorder resulting from the loss of imprinted gene expression within the paternal chromosome 15q11-q13. It has a prevalence rate of 1/10 000 to 30 000 and is characterised by endocrine abnormalities due to hypothalamic-pituitary insufficiency and complex physical, behavioural and intellectual impairment.

PWS is the main genetic cause of severe obesity in children. An early team approach to patient management allows improved quality of care and provides these individuals with a better quality of life.1 During early childhood, delayed motor and language development are observed; average milestones are achieved at approximately double the expected age. Intellectual and learning disabilities are variable and generally evident by the time the child reaches school age. Children with PWS can also develop psychiatric disorders throughout their lives.2

Growth hormone (GH) deficiency can occur in 40% to 100% of PWS patients, depending on GH stimulated test.3 The Pfizer International Growth Study (KIGS), with a cohort of 424 children with PWS, showed...
a prevalence for growth hormone deficiency of 74%.\textsuperscript{4} In the absence of GH replacement, patients with PWS usually present with short stature from the second decade of life, but it may be noticed earlier in childhood. It has been shown that insulin-like growth factor 1 and insulin-like growth factor binding protein 3 serum levels are low in patients with PWS.\textsuperscript{4,5} The association of GH deficiency, severe obesity and high fat mass indexes served as the foundation for the first studies to introduce the treatment with somatropin/recombinant human GH (rhGH) in these patients. Several studies trying to understand the real impact of the treatment with rhGH in PWS have since emerged.\textsuperscript{5–9}

The use of rhGH is currently recommended in PWS patients with and without GH deficiency to improve short stature and metabolism;\textsuperscript{8–10} its use may also have beneficial effects on reducing weight gain and possibly on appetite regulation.\textsuperscript{9,10} GH acts as an anabolic agent that increases lean body mass and reduces fat mass.\textsuperscript{7,11} Once the natural history of PWS and its association with the progressive increase of fat mass with age was established, the positive effect of rhGH on body composition

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**Figure 1** Preferred Reporting Items for Systematic Reviews and Meta-Analyses flow chart detailing selection of studies included in the meta-analysis. NRCTs; non-RCTs; RCTs, randomised controlled trials.

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**Figure 2** Difference between rhGH treated group related to Z-stature in RCTs (RevMan 5.3). (A) Random sequence generation (selection bias). (B) Allocation concealment (selection bias). (C) Blinding of participants and personnel (performance bias). (D) Blinding of outcomes assessment (detection bias). (E) Incompleted outcome data (attrition bias), selective reporting (reporting bias). (G) Intention-to-treat. (H) Sample size determination. (I) Outcome. (J) Early interruption. (K) Prognostic characteristics. GH, growth hormone; RCTs, randomised controlled trials; rhGH, recombinant human GH.
motivated its use in PWS patients as well as its approval by the US Food and Drug Administration since 2000.8 12

Another very important aspect of the use of rhGH therapy in PWS patients concerns the improvement in motor development, since hypotonia is a major problem in infants with PWS, may improve behaviours, since there is a high incidence of psychiatric problems, and may result in a better quality of life.13–15 Moreover, rhGH therapy may improve motor development, since hypotonia is a major problem.13–15 In addition, some concerns about the increased risk of death in these patients have emerged, especially regarding an increased incidence of sleep apnoea.19–22

The purpose of this systematic review is to provide a concise summary of the current knowledge regarding the use of rhGH treatment in patients with PWS, including recommendations, outcomes and implications for future research and clinical practice.

**OBJECTIVE**

To evaluate the impact (efficacy and safety) of the use of rhGH as a treatment for PWS. We performed a systematic

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**Table 1** Characteristics of the randomised controlled trials studies that evaluated the use of growth hormone in patients with Prader-Willi syndrome

<table>
<thead>
<tr>
<th>Author/year</th>
<th>Age</th>
<th>N rhGH/ control</th>
<th>Dose of rhGH</th>
<th>Period (years)</th>
<th>Country</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bakker NE et al16</td>
<td>6–14 years</td>
<td>12/16</td>
<td>1 mg/m2/day</td>
<td>2</td>
<td>Netherlands</td>
<td>Z-stature and body composition (%fat mass and LBM-SDS)</td>
</tr>
<tr>
<td>de Lind van Wijngaarden RFA et al13</td>
<td>4, 9±3.0 years (&lt;3 years and &gt;3 years)</td>
<td>29/21</td>
<td>1 mg/m2/day</td>
<td>1;2</td>
<td>Netherlands</td>
<td>Z-stature, BMI-SDS, body composition (%fat)</td>
</tr>
<tr>
<td>Böhm B et al13</td>
<td>3–11 years</td>
<td>19</td>
<td>0.033 mg/kg/ day</td>
<td>2</td>
<td>Sweden</td>
<td>Behaviour (specific test for the study) and cognition (Raven’s Standard Progressive Matrices test, Arthur’s adaptation of Leiter’s Performance Scale; verbal comprehension test (SPIQ test), Terman’s Scale of Intelligence, Bender Gestalt test, Harris Draw-a-Man)</td>
</tr>
<tr>
<td>Lo ST et al14</td>
<td>3, 5–14 years</td>
<td>24/18</td>
<td>1 mg/m2/day</td>
<td>1;2</td>
<td>Netherlands</td>
<td>Behaviour (Developmental Behaviour Checklist of Children with intellectual disability and Children Social Behaviour Questionnaire)</td>
</tr>
<tr>
<td>Reus L et al17</td>
<td>12.9±7.1 months</td>
<td>10/12</td>
<td>1 mg/m2/day</td>
<td>2</td>
<td>Netherlands</td>
<td>Muscular thickness with ultrasound, muscle strength (Infant Muscle Strength metre) and motor performance (Gross Motor Function Measure)</td>
</tr>
<tr>
<td>Siemensma EPC et al114</td>
<td>3, 5–14 (mean 7,4 years)</td>
<td>29/21</td>
<td>1 mg/m2/day</td>
<td>2</td>
<td>Netherlands</td>
<td>WISC-R (vocabulary, similarities, block design and picture arrangement) and Wechsler Preschool and Primary Scale of Intelligence-Revised Dutch version</td>
</tr>
<tr>
<td>Festen DAM et al7</td>
<td>3–14 years</td>
<td>25/25</td>
<td>1 mg/m2/day</td>
<td>2</td>
<td>Netherlands</td>
<td>Z-stature, body composition (%fat, LBM-SDS), head circumference (z)</td>
</tr>
<tr>
<td>Festen DAM et al16</td>
<td>Grupo I: 2–3 years C: 1–5 years</td>
<td>20/22</td>
<td>1 mg/m2/day</td>
<td>1</td>
<td>Netherlands</td>
<td>Motor and mental development (BSID-II),</td>
</tr>
<tr>
<td>Myers SE et al14</td>
<td>4–37 months</td>
<td>12/11</td>
<td>1 mg/m2/day</td>
<td>1</td>
<td>USA</td>
<td>Z-stature, head circumference, body composition (%fat, LBM-kg), language/cognition (age of first spoken word and Capute scales of infant’s language), motor development (age of walking and Toddler and Infant Motor Evaluation)</td>
</tr>
<tr>
<td>Camrel AL et al18</td>
<td>4–37 months</td>
<td>15/14</td>
<td>1 mg/m2/day</td>
<td>1</td>
<td>USA</td>
<td>Growth velocity, body composition and motor development (Toddler Infant Motor Evaluation)</td>
</tr>
<tr>
<td>Whitman BY et al17</td>
<td>4–16 years</td>
<td>35/19</td>
<td>1 mg/m2/day</td>
<td>2</td>
<td>USA</td>
<td>Behaviour (Oxford Survey Diagnostic Instrument and Family Inventory of Life Events)</td>
</tr>
<tr>
<td>Camrel AL et al6</td>
<td>4–16 years</td>
<td>35/19</td>
<td>1 mg/m2/day</td>
<td>1</td>
<td>USA</td>
<td>Body composition (%fat, LBM-kg), growth velocity, BMI-kg, muscular strength and agility (agility run, broad jump, sit-ups in 20 secs and upper extremity strength)</td>
</tr>
<tr>
<td>Lindgren AC et al6</td>
<td>3–12 years</td>
<td>15/12</td>
<td>0.1 IU/kg/day</td>
<td>1</td>
<td>Sweden</td>
<td>Z-stature, BMI-SDS, body composition (%fat, LBM-kg), growth velocity.</td>
</tr>
</tbody>
</table>

BMI, body mass index; LBM, lean body mass; rhGH, recombinant human GH; SDS, SD scores.
### Results of cognitive function, behaviour, motor development and quality of life

<table>
<thead>
<tr>
<th>Author</th>
<th>Test type</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cognitive function</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dykens EM et al</td>
<td>Cognition: KBIT-2 (verbal IQ, non-verbal IQ, composite IQ) VABS-II (communication and daily live skills, socialisation, adaptive composite)</td>
<td>Children receiving GHT had significantly higher verbal and composite IQs, and adaptive communication and daily living skills. Those who started before 12 months of age had higher non-verbal and composite IQs.</td>
</tr>
<tr>
<td>Donze SH et al</td>
<td>Psychomotor development (Bayley Scales of Infants II - BSID-II - mental and motor development)</td>
<td>During 3 years of GH, mental development increased from 58.1% (2.8) at baseline to 79.6% (3.7) (p&lt;0.01). A lower baseline psychomotor development and a younger age at start of GH treatment were associated with a higher increase in mental development (p&lt;0.01). No control group for this analysis.</td>
</tr>
<tr>
<td>Siemensma EPC et al</td>
<td>WISC-R (Vocabulary, similarities, block design and picture arrangement) and Wechsler Preschool and Primary Scale of Intelligence-Revised Dutch version</td>
<td>In short-term, mGH prevented deterioration of certain cognitive skills (similarities and vocabulary domain) and significantly improved abstract reasoning and visuospatial skills during 4 years of GH treatment. Children with lower cognitive functioning at baseline, GH treatment had a greater effect on abstract verbal reasoning and visuospatial skills in comparison with control group.</td>
</tr>
<tr>
<td>Festen DAM et al</td>
<td>Mental development Bayley Scales of Infant Development II (BSID-II)</td>
<td>Mental development improved significantly during the first year of study in the GH group versus the control group: median (IQR) change was +9.3% (-5.3 to 13.3) versus -2.9% (8.1 to 4.9) (p&lt;0.05) in mental development.</td>
</tr>
<tr>
<td>Myers SE</td>
<td>Language/cognition (age of first spoken word and Capute scales of Infants language)</td>
<td>GH-treated PWS group progressed significantly more during the first year in both language and cognitive development than the PWS control group; First words were spoken at 14 months, in 12 subjects treated before their first birthday and 17.2 in late-treated subjects, compared with reported data for global PWS of 21–53 months.</td>
</tr>
<tr>
<td>Böhm B et al</td>
<td>Cognition (Raven’s Standard, Progressive Matrices test, Arthur’s adaptation of Leiter’s Performance Scale; SPIQ test, Terman’s Scale of Intelligence, Harris Draw-a-Man)</td>
<td>No difference was found in these patients (mean age 6.5 years) regarding cognitive function.</td>
</tr>
<tr>
<td><strong>Behaviour outcome</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dykens EM et al</td>
<td>Repetitive Behaviour Scale and Adaptive Behaviour</td>
<td>mGH treated versus treatment naive 4 to 21 years old children with PWS had significantly higher adaptive behaviour standard scores, but no differences in repetitive behaviour scales.</td>
</tr>
<tr>
<td>Lo ST et al</td>
<td>Behaviour (Developmental Behaviour Checklist of Children with intellectual disability and Children Social Behaviour Questionnaire)</td>
<td>No difference reported.</td>
</tr>
<tr>
<td>Whitman BY et al</td>
<td>Behaviour (Offord Survey Diagnostic Instrument and Family Inventory of Life Events)</td>
<td>No differences between groups; however, a significant positive effect (reduction of depressive symptoms) was noted for the treated group regarding cognitive function.</td>
</tr>
<tr>
<td>Böhm B et al</td>
<td>Behaviour (specific test for this study)</td>
<td>No difference between groups. Successively worsened behaviour after discontinuing GH treatment.</td>
</tr>
<tr>
<td><strong>Motor development/ muscle strength</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Donze SH et al</td>
<td>Psychomotor development (BSID-II - mental and motor development)</td>
<td>During 3 years of GH, mean motor development increased from 41.9% (2.9) to 78.2% (3.9) (p&lt;0.01). A lower baseline psychomotor development and a younger age at start of GH treatment were associated with a higher increase in motor development (p&lt;0.01).</td>
</tr>
<tr>
<td>Reus L et al</td>
<td>Muscular thickness by ultrasound</td>
<td>GH has a positive effect on muscle thickness (biceps; forearm flexor, quadriceps and tibialis anterior) in PWS infants.</td>
</tr>
<tr>
<td>Reus L et al</td>
<td>Muscle strength (Infant Muscle Strength Meter - AIMMS) and Motor function (Gross Motor Function Measure - GMFM); Bayley Scales of Infants II- BSID-II</td>
<td>The AIMMS and GMFM clearly revealed a significant positive effect of GH on motor development; the child’s maximum motor potential increased with GH treatment, thereby resulting in a clear functional improvement. BSID-II revealed no effect of GH on motor development.</td>
</tr>
<tr>
<td>Festen DAM et al</td>
<td>Motor and mental development, Bayley Scales of Infant Development II (BSID-II)</td>
<td>Significant improvement in motor development (+11.2% vs –18.5%) at the first year of treatment in the treated group versus control group from the baseline.</td>
</tr>
<tr>
<td>Myers SE et al</td>
<td>Motor development (age of walking and Toddler and Infant Motor Evaluation)</td>
<td>A trend towards improved mobility and stability percentile rankings was noted with GH therapy, however, wide variability among PWS subjects was seen at all time points. Eleven PWS subjects treated before their first birthday walked independently at a mean age of 23.5 months. Five late-treated PWS subjects walked independently at a mean age of 24.3 months.</td>
</tr>
<tr>
<td>Carrel AL et al</td>
<td>Muscular strength</td>
<td>Documented changes in physical function (strength and agility testing) in PWS children treated with GH in first 2 years therapy, but no change in the following 2 years.</td>
</tr>
<tr>
<td>Carrel AL et al</td>
<td>Muscular strength and agility (agility run, broad jump, sit-ups in 20 s and upper extremity strength)</td>
<td>Agility run (faster by 2.3±0.5s), broad jump (farther by 3.3±1.9 inches), abdominal strength (an improvement of 3.0±2.1 sit-ups/20 s) and upper extremity strength (increase of 2.5±1.8 weight-lift repetitions/30 s); compared with baseline performance</td>
</tr>
<tr>
<td><strong>Quality of life</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bakker NE et al</td>
<td>DUX25 and DUXPW questionnaires for children and parents</td>
<td>GH-treated children showed a significant improvement in health-related quality of life during the 2-year RCT in the physical subdomain of the DUX25 and the DUXPW, compared with the untreated ones. During long-term GH treatment (11 year), questionnaires scores remained stable. Social subdomain was higher in children with a deletion than in children with an uniparental disomy or imprinting defect, according to parents</td>
</tr>
<tr>
<td>Sipilä I et al</td>
<td>16D (generic 16-dimension health-related quality of life) instrument for adolescents</td>
<td>The effect of rGH therapy remains unclear because of a lack of untreated control group with PWS and lack of comparable baseline measurements for evaluating changes over time.</td>
</tr>
</tbody>
</table>

GH, growth hormone; GHT, GH therapy; IQs, intelligent quotients; PWS, Prader-Willi syndrome; RCT, randomised controlled trial; rGH, recombinant human GH.
Table 2 Characteristics of the non-randomised studies that evaluated the use of growth hormone in patients with Prader-Willi syndrome

<table>
<thead>
<tr>
<th>Author</th>
<th>Type</th>
<th>Age</th>
<th>N rhGH</th>
<th>Dose of rhGH</th>
<th>Period (years)</th>
<th>Country</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Donzé SH et al</td>
<td>BEFORE AND AFTER</td>
<td>6 months-3 years</td>
<td>29</td>
<td>1 mg/m²/day</td>
<td>3</td>
<td>Netherlands</td>
<td>Psychomotor development (BSID-II - mental and motor development), head circumference</td>
</tr>
<tr>
<td>Dykens EM et al</td>
<td>(three separated studies)</td>
<td>4-21 years</td>
<td>96; 127; 168</td>
<td>1 mg/m²/day</td>
<td>6, 2</td>
<td>USA</td>
<td>BMI-SDS, Cognition: KBIT-2 (verbal IQ, non-verbal IQ, composite IQ) VABS-II (communication and daily life skills, socialisation, adaptive composite), Hyperphagia Questionnaire; Repetitive Behaviour Scale</td>
</tr>
<tr>
<td>Bakker NE et al</td>
<td>BEFORE AND AFTER</td>
<td>6-4.3 years</td>
<td>1566</td>
<td>0.23 mg/kg/week</td>
<td>3</td>
<td>KIGS database - worldwide</td>
<td>Z-stature, BMI-SDS, adverse effects</td>
</tr>
<tr>
<td>Bakker NE et al</td>
<td>BEFORE AND AFTER</td>
<td>6-14 years</td>
<td>76</td>
<td>1 mg/m²/day</td>
<td>11</td>
<td>Netherlands</td>
<td>Quality of life (DUX25 and DUXPW questionnaires)</td>
</tr>
<tr>
<td>Scheermeyer E et al</td>
<td>BEFORE AND AFTER</td>
<td>13-24 months 2-12 years</td>
<td>72</td>
<td>1 mg/m²/day</td>
<td>3</td>
<td>Australia</td>
<td>Z-stature, BMI-SDS and adverse effects</td>
</tr>
<tr>
<td>Lo ST 2015</td>
<td>COHORT</td>
<td>3.5-14 years</td>
<td>24</td>
<td>1 mg/m²/day</td>
<td>2;8</td>
<td>Netherlands</td>
<td>Behaviour, Developmental Behaviour Checklist of Children with intellectual disability (DBC) and Children's Social Behaviour Questionnaire (CSBQ)</td>
</tr>
<tr>
<td>Lo ST et al</td>
<td>COHORT</td>
<td>3.5-14 years</td>
<td>53</td>
<td>1 mg/m²/day</td>
<td>2;7</td>
<td>Netherlands</td>
<td>VABS-II, Cognitive function: Bayley Scales of Infants II - NL (BSID-II), Wechsler Preschool and Primary Scale of Intelligence-Revised Dutch version</td>
</tr>
<tr>
<td>Bakker NE et al</td>
<td>BEFORE AND AFTER</td>
<td>3-7 years (mean: 5.49)</td>
<td>60</td>
<td>1 mg/m²/day</td>
<td>8</td>
<td>Netherlands</td>
<td>Z-stature, BMI-SDS, lean body mass-SDS, % fat mass-SDS, head circumference</td>
</tr>
<tr>
<td>Meinhardt U et al</td>
<td>COHORT</td>
<td>0.4-12.2 years (mean:3.8)</td>
<td>41</td>
<td>0.03 mg/kg/day</td>
<td>1; 6</td>
<td>Switzerland, Denmark, Germany</td>
<td>Z-stature, body composition and adverse effects</td>
</tr>
<tr>
<td>Siemensma EP et al</td>
<td>COHORT</td>
<td>3.5-14 years</td>
<td>29</td>
<td>1 mg/m²/day</td>
<td>2; 4</td>
<td>Netherlands</td>
<td>WISC-R (vocabulary, similarities, block design and picture arrangement) and Wechsler Preschool and Primary Scale of Intelligence-Revised Dutch version - less 7 years old</td>
</tr>
<tr>
<td>Colmenares A et al</td>
<td>COHORT</td>
<td>1-15 years (mean: 6±3.7)</td>
<td>36</td>
<td>0.03 mg/kg/day</td>
<td>3</td>
<td>France</td>
<td>Z-stature, BMI-SDS and body composition- lean mass (kg), %fat mass</td>
</tr>
<tr>
<td>Sipilä I et al</td>
<td>BEFORE AND AFTER</td>
<td>2.0-10.3 years (mean: 6.6)</td>
<td>20</td>
<td>0.033 mg/kg/</td>
<td>1; 10</td>
<td>Finland</td>
<td>Z-stature, BMI-SDS, body composition (%fat), and quality of life (HRQOL)</td>
</tr>
<tr>
<td>Carrel AL et al</td>
<td>COHORT</td>
<td>4-32 months/. 6-9 years</td>
<td>46</td>
<td>1 mg/m²/day</td>
<td>6</td>
<td>USA</td>
<td>Z-stature, body composition and muscular strength</td>
</tr>
<tr>
<td>de Lind van Wijngaarden RFA et al</td>
<td>COHORT</td>
<td>5.9±3.2 years</td>
<td>55</td>
<td>1 mg/m²/day</td>
<td>4</td>
<td>Netherlands</td>
<td>Z-stature, BMI-SDS, body composition (%fat-SDS and lean body mass-SDS), head circumference</td>
</tr>
<tr>
<td>Nyunt O et al</td>
<td>COHORT</td>
<td>&lt;3 years and &gt;3 years</td>
<td>54</td>
<td>0.6 mg/m²/day</td>
<td>4</td>
<td>Australia</td>
<td>Z-stature and BMI-SDS</td>
</tr>
<tr>
<td>Lindgren AC et al</td>
<td>BEFORE AND AFTER</td>
<td>4.9-11.3 years (mean: 6.9)</td>
<td>22</td>
<td>0.03 mg/kg/d</td>
<td>10</td>
<td>Sweden</td>
<td>Z-stature, BMI-SDS and body composition</td>
</tr>
<tr>
<td>Lin H-Y et al</td>
<td>BEFORE AND AFTER</td>
<td>1.3-13.5 years</td>
<td>46</td>
<td>0.01U/kg/day</td>
<td>5</td>
<td>Taiwan</td>
<td>Z-stature, BMI-SDS and adverse effects</td>
</tr>
<tr>
<td>Angulo MA et al</td>
<td>COHORT</td>
<td>8.3±2.7 years</td>
<td>21</td>
<td>0.25 mg/kg/week</td>
<td>7.9</td>
<td>USA</td>
<td>Z-stature and BMI-SDS</td>
</tr>
<tr>
<td>Carrel AL et al</td>
<td>BEFORE AND AFTER</td>
<td>12 years</td>
<td>48</td>
<td>1 mg/m²/day</td>
<td>4</td>
<td>USA</td>
<td>Z-stature, body composition, muscular strength, adverse effects</td>
</tr>
<tr>
<td>Tauber M et al</td>
<td>BEFORE AND AFTER</td>
<td>8.4±3.2 years</td>
<td>14</td>
<td>0.5 U/kg/week</td>
<td>3.6</td>
<td>France</td>
<td>Z- stature SDS, BMI-SDS, growth velocity</td>
</tr>
</tbody>
</table>

BMI, body mass index; rhGH, recombinant human GH; SDS, SD scores.

review followed where possible by meta-analysis for the following outcomes:

- Primary outcomes: growth, body mass index, body composition, cognitive function.
- Secondary outcomes: quality of life, head circumference, motor development/muscle strength, behaviour and adverse effects.

METHODS
The systematic review and meta-analysis were performed according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines (online supplementary file 1) and it was registered with PROSPERO (International Prospective Register of Systematic Reviews). We included all PWS patients, with all types of genetic defects, with or without GH deficiency, who participated in GH studies performed in infancy, childhood and adolescence, that were either randomised controlled trials (RCTs) (double-blinded or not) or non-randomised controlled trials (NRCTs) (cohort and before and after studies). Narrative reviews, systematic review protocols, case reports, abstracts, data from unpublished research or incomplete articles were...
Figure 3  
Association between rhGH treatment and Z-stature in the NRCTS (RevMan 5.3).  
(A) Inclusion criteria.  
(B) Condition and measure.  
(C) Diagnostic method.  
(D) Consecutive inclusion.  
(E) Loss of participants.  
(F) Group characteristics.  
(G) Clinical information.  
(H) Outcomes.  
(J) Demographics. NRCTs, non-randomised controlled trials; rhGH, recombinant human growth hormone.

Figure 4  
Association between rhGH-treated group and the no treatment group according to growth velocity in RCTs (RevMan 5.3).  
(A) Random sequence generation (selection bias).  
(B) Allocation concealment (selection bias).  
(C) Blinding of participants and personnel (performance bias).  
(D) Blinding of outcomes assessment (detection bias).  
(E) Incomplete outcome data (attrition bias).  
(F) Selective reporting (reporting bias).  
(G) Intention-to-treat.  
(H) Sample size determination.  
(I) Outcome.  
(J) Early interruption.  
(K) Prognostic characteristics. RCTs, randomised controlled trials; rhGH, recombinant human growth hormone.

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Motor development/muscle strength: evaluated by different types of measures.

- Adverse effects: sleep abnormalities, initial headache, peripheral oedema, scoliosis, diabetes and mortality.

Electronic searches

The search for evidence on the efficacy of somatropin (rhGH) on PWS was performed through a systematic review based on the PRISMA checklist. There were no restrictions on language or period. The databases used were MEDLINE (via PubMed), Embase and Cochrane Central with the following search strategies.

MEDLINE (via PubMed)

(Prader-Willi syndrome OR Prader-Labhart-Willi syndrome OR Labhart Willi Prader Fanconi syndrome OR Royer syndrome) AND (human growth hormone OR hGH OR somatotropin OR somatropin OR Serostim OR Zomacton OR Cryo-Tropin OR recombinant human growth hormone AND (mammalian) OR r-hGH-M OR r-hGH AND (m) OR Humatrope OR Umatrope OR Maxomat OR Norditropin OR Nutropin OR Omnitrope OR Saizen OR Genotropin OR Genotonorm)

Embaze

(Prader-Willi Syndrome OR Prader-Labhart-Willi Syndrome OR Labhart Willi Prader Fanconi Syndrome OR Royer Syndrome) AND (human growth hormone OR hGH OR somatotropin OR somatropin OR Serostim OR Zomacton OR Cryo-Tropin OR recombinant human

excluded. Studies with less than 1 year of rhGH use were also excluded.

The rhGH group was compared with a matching placebo group or patients who received no drugs during the 1 year follow-up in RCTs. For NRCTs we evaluated patients at the start and at the end of the treatment. All studies that had patients who received a subcutaneous application of rhGH at any dosage were included.

We analysed the following outcome measures:

- Growth: Z-stature SD scores (SDS) according to WHO growth charts, Centers for Disease Control and Prevention (CDC) growth charts or by PWS curve; growth velocity: centimetres/year or growth velocity SDS.23–25

- Body mass index (m²/kg): BMI-SDS according to WHO growth charts and CDC growth charts.23–25

- Body composition (lean mass and fat mass): measure by dual-energy X-ray absorptiometry (DXA-type Lunar Prodigy, GE Healthcare-Chalfont St Giles, UK, or Madison, Wisconsin). We evaluated the proportion of total body/trunk fat mass (kilograms) or SDS of lean mass.

- Cognitive function: all types of validated questionnaires.

- Quality of life: evaluated by different validated instruments.

- Behaviour: evaluated by different validated instruments.

- Head circumference: Z-score according to CDC growth charts and Netherlands charts.26

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growth hormone AND (mammalian) OR rhGH-M OR rhGH AND (m) OR Humatrope OR Umatrope OR Maxomar OR Norditropin OR Nutropin OR Omnitrope OR Saizen OR Genotonorm OR Genotropin OR Genotonorm)

Cochrane
(Prader-Willi syndrome OR Prader-Labhart-Willi syndrome OR Labhart Willi Prader Fanconi syndrome OR Royer syndrome) AND (human growth hormone)

All PWS articles were evaluated independently by two reviewers. Following the eligibility criteria, studies were evaluated first by title and then by abstract. The full-text of selected articles was assessed for data extraction and bias evaluation. A characterisation matrix was constructed with the following variables: author, publication year, study design, country, age, period, dose and outcomes. When the data were not clear in the paper, we contacted the authors by email. Grey literature (thesis, book chapters) and references of selected articles were reviewed.

Selection of studies and data extraction
Identification, screening and eligibility assessments were performed independently in an unblinded, standardised manner by two reviewers. Disagreements between reviewers were discussed with a third researcher. Extraction of dichotomous (adverse effects) and continuous variables (all other outcomes) was independently performed.

Assessment of risk of bias in the included studies
Risk of bias was evaluated separately by two reviewers. The Cochrane Collaboration Risk of Bias tool was used for randomised trials, the New Castle Ottawa Scale was used for cohort studies and the Joanna Briggs Institute Instrument for used for before and after studies and for cohort studies when only one arm of the cohort was analysed.27 28

Measures of treatment effect and units of analyses
We performed separated meta-analyses of RCTs and NRCTs. Associations were demonstrated in relative and absolute risks, difference in means, SD, SEs and 95% CIs. Different follow-up lengths were standardised by using patient-years of follow-up. When different measures or instruments were used in the studies, SEs were used to estimate outcomes. P values lower $p=0.05$ were considered statistically significant to evaluate treatment effect. Studies that did not report data in terms of means and their respective SD were recalculated or not included in the analysis. Studies that mentioned medians, 25th and 75th percentiles and CIs were included after transforming these statistics into SD by Hozo et al method.29 When the data were not clear in the paper or were missing, we contacted the authors by email.

Subjects were evaluated in two subgroups: infants (under 3.5 years old) and children and adolescents. We selected this younger group to look for differences in the early start of rhGH and the cut-off was based in the majority of articles found.
The software used was Cochrane Review Manager (V.5.3; The Nordic Cochrane Centre). The possibility of publication bias was assessed by evaluating a funnel plot (Beggar’s test) for asymmetry. We used GRADE considerations (study limitations, consistency of effect, imprecision, indirectness and publication bias) to assess the data quality for each outcome and to draw conclusions about the quality of the evidence within the text of the review.

**Patient and public involvement**

Patients and/or the public were not involved in the design, or conduct, or reporting or dissemination plans of this research.

**RESULTS**

The electronic search yielded 463 titles from the databases until September 2019, and after text selection, this review included 16 completed RCTs and 20 NRCTs (figure 1). The excluded studies after full-text analysis (n=18) are presented in a separate table (see online supplementary file 2).

**Randomised control studies**

We performed the meta-analysis for the evaluation of growth hormone treatment versus no treatment in 9 of the 16 included RCT (table 1). No study used placebo due to ethical issues. The variables analysed were height z-scores (Z,E), growth velocity (cm), BMI z-scores (Z-BMI), proportion of fat mass (%), lean mass z-scores (Z and kg) and head circumference z-scores (Z).

The seven meta-analysis excluded studies contain outcomes related to cognitive function, behaviour, quality of life and motor development. For the evaluation of cognitive function in patients with PWS, different questionnaires were used; therefore, meta-analysis was not possible.33-35 The same occurred with the behaviour outcome.17 33 36 Regarding quality of life, only one study was found.18 Five studies with different measures were found for muscular strength/motor development.5 16 35 37 38 All these studies are described in table 1 and analysed at the end of results section.

**Non-randomised studies**

Regarding the non-randomised studies, 20 were selected: 11 before and after studies and 9 cohort studies (table 2). Of these, only 14 could be evaluated regarding the following outcomes: height z-scores (Z,E), Z-BMI, proportion of fat mass (%) and adverse effects. The patient-year (number of patients x years of follow-up) unit was used for long-term follow-up studies.

There were only two studies that evaluated quality of life,18 39 two that evaluated behaviour with different questionnaires13 36 and four that evaluated cognitive function, also with different evaluation methods (table 2).13-15 94 We found two studies for muscular strength/motor development.15 40 They are analysed at the end of results section. For the meta-analysis process and for a better approach to all data, we transformed all cohort studies into before and after studies. Therefore, the study of Carrel 2010 was excluded from all analysis because it had no baseline data.
For all the studies included, the obtained results are reported according to each outcome.

**Growth: z-stature and growth velocity**
Among the nine RCTs with a total of 328 patients, the treated group had an improvement in height of 1.67 (95% CI 1.54 to 1.81) SDS compared with that in the control group. In the younger group (under 3.5 years), the improvement was 1.08 (95% CI 0.81 to 0.35) SDS, and in the older group, it was 1.87 (95% CI 1.71 to 2.02) SDS. The Z-stature meta-analysis quality of evidence was moderate when all groups were included and high when the two age groups were analysed separately, due to decreased heterogeneity (figure 2, online supplementary file 3).

In the NRCT studies, the treated group had a significant improvement in stature (1.52 (95% CI 0.86 to 2.16) SDS) during the 2909 patients-years of follow-up (figure 3), this meta-analysis, however, was based in low quality evidence (online supplementary file 4).

The group receiving rhGH treatment had a significant improvement in growth velocity of 5.44 cm (95% CI 3.27 to 7.61) compared with that in the control group among the three RCTs (figure 4), but the quality of evidence was also low (online supplementary file 3). We chose to report the growth velocity measure in centimetres and due to that one article that used only SDS measure was excluded.

**Z-BMI**
The group receiving rhGH treatment had a Z-BMI decrease of −0.67 SDS (95% CI −0.87 to −0.47) compared with that in the control group among the three RCTs (figure 5). The quality of evidence was high (online supplementary file 3). The study of Bakker 2015 was excluded after examining the funnel plot.18

The NRCT studies showed no difference during the 2237 patients-years of follow-up after rhGH treatment (figure 6). It is important to note that no change in BMI during the follow-up of these patients was considered the opposite of the disease natural history. The quality of this evidence was very low (online supplementary file 4).

**%Fat mass**
In six of the included RCTs, which evaluated a total of 204 patients, the group treated with rhGH had a significant reduction in the fat mass proportion of −6.5 (95% CI −8.46 to −4.54) % compared with that in the control group. These data have high quality of evidence (figure 7) (online supplementary file 3). All studies used Lunar DXA. We excluded the de Lind van Wijngaarden RFA 2010 because they used a different type of measure for body fat mass (fat mass SDS according to Dutch patients).41

In the NRCT studies, we also found a significant reduction in the proportion of fat mass (−7.04%; 95% CI −8.39 to −5.7)) in 198 patients-year of follow-up with a low
quality of evidence (figure 8) (online supplementary file 4). The data in the de Lind van Wijngaarden RFA 2009 study were reported in SDS-fat mass and were not included in the analysis.9

Lean mass
The group receiving rhGH treatment had an SDS-lean mass improvement of 2.03 (95% CI 1.34 to 2.71) SDS compared with that of the control group among two randomised controlled trials with moderate quality of evidence (figure 9, online supplementary file 3). The studies by Lindgren 1998 and Carrel 1999 used another variable and showed an increase of 4.28 (95% CI 2.26 to 6.30) kg of lean mass with high quality of evidence (online supplementary file 3).5,6

Head circumference
The group receiving rhGH treatment had a head circumference improvement of 0.55 (95% CI 0.25 to 0.86) cm compared with that of the control group among the three RCTs including a total of 114 patients. The quality of the evidence was moderate (figure 10, online supplementary file 3).

Cognitive function, behaviour, motor development/muscular strength and quality of life
All the results from RCTs and NRCTs were described in table 3.

Adverse effects
In the RCT studies, there were few reports related to the adverse effects of rhGH use. Carrel 1999 reported one case of cerebral pseudotumour, and Myers reported one case of scoliosis.5,6 For a better analysis, we used ‘before and after studies’, which evaluated more patients and had a longer follow-up. Side effects related to rhGH in patients/year of follow-up were 227. We only included the following side effects for this analysis that are according the authors are related to the treatment: sleep apnoea (n=35), initial headache (n=9), peripheral oedema (n=7), type 2 diabetes mellitus (n=11), scoliosis (n=64). This last side effect is cited and counted, but it is difficult to establish a relationship between its rhGH progression and treatment. The quality of this evidence was moderate. Twelve deaths were reported in all these studies. (figure 11, online supplementary file 4).

DISCUSSION
PWS is a challenging and rare disease with few treatment options. To the best of our knowledge, this is the first meta-analysis that evaluated the use of rhGH in a large number of PWS patients while assessing different domains, allowing an overall vision of the impact of rhGH treatment.

Our study showed that rhGH use during infancy promoted an improvement of stature, body composition (%fat mass and lean body mass) and growth velocity. Indeed, data about final height confirmed this evidence.3 24 42 43 The impact of +1.67SD in stature and the decrease of 6.5% to 7.0% in fat mass is not negligible in PWS daily life and can contribute with their fight against obesity and their complications. The short-term impact on BMI, in association with diet, was good; in the long run, rhGH controlled the weight gain that is part of the natural history of PWS, but there was a great deal of variability among the studies.

The confidence level for each effect changed according to the quality of the evidence. We found a high quality of evidence for stature, BMI and %fat mass, and we found a moderate level for the other parameters in the RCT studies. Furthermore, a reasonable number of non-RCT studies confirmed our findings even with a low quality of evidence.

Although the effects of rhGH on growth and body composition were more objectively reported, cognitive function studies were quite difficult to evaluate. Most cognitive function studies reported improvements in the different measures used, especially in verbal and non-verbal IQ and vocabulary domain.13 44 Children that started the treatment at a young age (less than 1 year old) reported better adaptive communication skills in long-term studies.13 44 Indeed, in Bayley Scales for Infants two
studies showed differences in mental development.\textsuperscript{15} 16 Unfortunately, these data could not be aggregated.

The actions of rhGH on cognitive function could be explained by the presence of brain GH receptors, especially at a younger age, during which brain development is taking place at a more intense pace.\textsuperscript{14} Moreover, GH can increase the muscle tone, as we will discuss in reviewed articles, and stimulate physical activity. Therefore, it can promote articulation and motor control for optimal development of communication skills, which additionally promotes social interaction and physical tasks. These effects might be cumulative over time. Also, the largest effect of GH treatment on adaptive functioning was reached if GH treatment was started at a younger age during infancy.\textsuperscript{15}

Muscular strength and motor development seemed to improve in treated patients according to selected articles.\textsuperscript{5} 15 16 37 40 44 Multilevel regression models controlled for age and baseline muscle thickness identified a statistically significant rhGH dependency in the Reus article.\textsuperscript{37} The authors discuss a major response in the first years of treatment and at a young age.\textsuperscript{16} 40

The impact on behaviour in patients with PWS was not clear. Most of the articles did not show any difference. Positive results are cited in Böhm \textit{et al} that found worse behaviour in PWS patients, as evaluated by a parental questionnaire, after cessation of hGH treatment and in Dykens \textit{et al} that showed better adaptive behaviour scores.\textsuperscript{13} 27 Regarding quality of life, there is only one study with a specific PWS questionnaire that showed an improvement at the start of the treatment, but not in the long run.\textsuperscript{18}

The low number of adverse effects in RCTs are generally poorly reported in literature, as we found in our analysis.\textsuperscript{46} In NRCTs, the number of reported adverse events indicated a low risk of mortality and few complications during long-term follow-up in studies with moderate quality of evidence. Sleep apnoea still needs to be evaluated with polysomnography. Even in cohort studies, we have some difficulties to access all information regarding adverse effects, so we cannot fully comment on safety.

The different measures adopted, especially regarding the body composition analysis (related to the Dutch SDS for all parameters) and the cognitive function evaluation, were a limitation for this study. Many articles could not be used. It was not possible to analyse the impact in mental development and IQ, an important initial endpoint for this study, due to the lack of universal questionnaires in this field. We are convinced that the elaboration of a specific questionnaire to evaluate the syndrome at each age would add great value for this analysis in future studies. Additionally, more studies evaluating cognitive function and quality of life are needed in PWS patients.

Another barrier was the difficulty in finding RCTs that studied the use of rhGH in patients with PWS. The main reason is that professionals who treat PWS feel uncomfortable and unethical withholding rhGH treatment in a group of their patients with PWS. At the same time, many countries still do not have access to rhGH treatment. Potential biases could be related to exclusion criteria in clinical studies, where severe cases or adherence difficulties (social and economic problems) could be excluded. Therefore, larger registry is still important to evaluate long-term follow-up with specific emphasis on the covariates of PWS to extrapolate our results. The genetic variability of PWS, with different implicated genes, is a great challenge for patients’ comparison, especially regarding cognition. Differences according to the dose response and genetic subtypes could not be analysed in this review but may be very important to ascertain conclusive disease knowledge and its prognosis following treatment.

**CONCLUSION**

Growth hormone treatment can promote an improvement of stature, body composition and BMI, modifying the disease’s natural history; growth hormone treatment may also be implicated in improving cognition and motor development at a young age.

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**Contributors**

All authors contributed to the focus of the systematic review. CP and WB drafted the initial protocol, with guidance and feedback at all stages from SI, RF, ET, DD and MP. CP, SI and RF collected data from the articles. CP, WB DD and MP interpreted the results. All authors have revised and reviewed the results and have approved the final manuscript.

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**Competing interests**

No, there are no competing interests.

**Patient consent for publication**

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**Provenance and peer review**

Not commissioned; externally peer reviewed.

**Data availability statement**

Data are available upon reasonable request. All data relevant to the study are included in the article or uploaded as supplementary information. Data from published studies were analysed.

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