Liraglutide pharmacokinetics and exposure-response in adolescents with obesity

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Summary
Background: Obesity in adolescence presents a major public health challenge, often leading to obesity in adulthood with associated chronic disease.
Objectives: This study aimed to perform a population pharmacokinetic and exposure-response analysis of liraglutide by meta-analysis of data from trials conducted in children, adolescents and adults with obesity.
Methods: The population pharmacokinetic analysis investigated the effect of covariates body weight, age group (children, adolescents and adults) and sex on liraglutide exposure in adolescents compared with previous results in adults. The exposure-response relationship of liraglutide for the change from baseline in body mass index standard deviation score (BMI SDS) was evaluated in adolescents and compared to that in adults.
Results: Body weight was the main covariate affecting liraglutide exposure, with lower exposures at higher body weights, whereas age group was of no importance and sex was of little importance. An exposure-response relationship was demonstrated for liraglutide in both adolescents and adults as the decrease in BMI SDS from baseline increased in an exposure-dependent manner with increasing liraglutide exposure.
Conclusions: The population pharmacokinetic analysis supported similar liraglutide exposures in adolescents and adults; body weight was the most important covariate affecting exposure. An exposure-response relationship was established for liraglutide.

KEYWORDS
adolescents, clinical trial, GLP-1, liraglutide, paediatric, pharmacokinetics

1 | INTRODUCTION

The prevalence of obesity over the past 30 years has more than doubled in children and tripled in adolescents, reaching epidemic proportions in the United States.1,2 While the overall prevalence of obesity is lower in children than in adults, rates of increase in childhood obesity are now higher than those seen in adults in many countries.3 Childhood obesity presents a major public health challenge since obesity in childhood and adolescence often leads to obesity in adulthood.4-6 Paediatric obesity is associated with a wide range of chronic diseases, including type 2 diabetes, hypertension, hyperlipidaemia, polycystic ovary syndrome and sleep apnea.7-10 Prolonged obesity continuing into adulthood may lead to complications such as cardiovascular disease,11,12 which was responsible for more than two-thirds of adult deaths associated with high body mass index (BMI) worldwide in 2015.3 Childhood and adolescent obesity may also...
adversely affect quality of life, resulting in adverse psychosocial problems such as low self-esteem, depression and reduced educational achievement. The economic consequences of obesity and its related complications are also high.

Current treatments for children and adolescents with obesity tend to favour lifestyle modifications that target diet and exercise; however, such programs often have limited effect on reducing BMI in adolescents because weight loss and maintenance of weight loss are hard to achieve. Bariatric surgery may be an effective alternative for individuals with morbid obesity, though there remains a treatment gap for patients who do not meet criteria for surgery and those who struggle to lose weight with lifestyle interventions alone. Few approved pharmacotherapies are available for the treatment of obesity in the paediatric population. In the United States, liraglutide 3.0 mg was recently approved for chronic weight management by the Food and Drug Administration (FDA) for use in patients aged ≥12 years weighing >60 kg. Both orlistat (patients ≥12 years) and phentermine (patients >16 years) are also approved for weight management, and setmelanotide is approved for the treatment of certain genetic causes of obesity (patients ≥6 years). Liraglutide 3.0 mg is also approved for use in adolescents in Brazil and Saudi Arabia and orlistat is approved in Switzerland for adolescent use by the Swiss Regulatory Agency, Swissmedic.

Glucagon-like peptide-1 (GLP-1), an incretin hormone secreted by intestinal L cells in response to food intake, stimulates insulin secretion and inhibits glucagon secretion in a glucose-dependent manner. Liraglutide is a GLP-1 receptor agonist that promotes weight loss through reduced appetite and a subsequent reduction in energy intake. As an adjunct to a reduced-calorie diet and increased physical activity, liraglutide 3.0 mg (Saxenda) once daily is approved in the United States, European Union and elsewhere (65 countries) for chronic weight management in adults with obesity (BMI ≥30 kg/m²), or overweight (≥27 to <30 kg/m²) in the presence of at least one weight-related comorbidity.

The efficacy and safety of liraglutide in adults were evaluated in the Satiey and Clinical Adiposity – Liraglutide Evidence (SCALE) program. The safety, tolerability, pharmacokinetics and pharmacodynamics of liraglutide in adolescents aged 12-17 years and in children aged 7-11 years have previously been investigated in short-term (<10 weeks) randomized, controlled trials. In phase 3 randomized, controlled 56-week trial investigating the effects of liraglutide for weight management in pubertal adolescents with obesity, liraglutide, in addition to lifestyle therapy, significantly reduced the BMI standard-deviation score (SDS) compared with placebo (estimated treatment difference: −0.22; 95% confidence interval [CI], −0.37, −0.08; P = 0.002).

A previous population pharmacokinetic analysis performed with liraglutide using adult data from the 56-week phase 3a SCALE Obesity and Prediabetes trial (1839) and SCALE Diabetes trial (1922) trials found that sex and body weight were the only intrinsic factors that influenced the exposure (ie, the average steady-state plasma concentration) of liraglutide 3.0 mg. Moreover, in a previous exposure-response analysis using data from the aforementioned SCALE trials and a phase 2, 20-week trial (1807), greater weight loss was observed with higher liraglutide exposures. This report aimed to perform population pharmacokinetic and exposure-response analyses of data from the recent 56-week trial in adolescents (trial 4180) in a meta-analysis including data from previous trials conducted in children, adolescents and adults with obesity, to support the 3.0 mg liraglutide dose for weight management in an adolescent population. The population pharmacokinetic analysis will provide additional information including adolescent data from a long-term (52-week) phase 3 trial, to support previous findings. The exposure-response analysis is the first to evaluate phase 3 data from both adults and adolescents together.

METHODS

2.1 Population pharmacokinetic analysis

The objectives of the population pharmacokinetic analysis were (1) to investigate body weight, age group (children aged 7-11 years, adolescents aged 12-17 years and adults aged ≥18 years) and sex as covariates to determine whether their impact on liraglutide exposure was in accordance with previous results in adults and (2) to compare the drug exposure for adolescents in the 56-week trial 4180 to previous results from shorter trials in adolescents, children and adults. As the focus of the analysis was a comparison of different age populations, age was prespecified to be included initially as a categorical variable. If proven to be an important covariate, further exploration of this relationship was possible if relevant; however, this was not the case.

Data from trials 4180 and 3967 (in adolescents), 4181 (children) and 3630 (adults) were included in the pharmacokinetic assessment of liraglutide (Table 1). In trial 4180, Tanner staging (stages 2-5) of adolescents was assessed by site staff trained in pubertal assessments. For females, assessments of breast and pubic hair development were made. For males, assessments of testicular volume (by orchidometer), penis development and pubic hair development were made. Prepubertal patients were not permitted in the trial. In all trials, liraglutide concentration in plasma was determined using a validated enzyme-linked immuno-sorbent assay (ELISA) with a lower limit of quantification (LLOQ) of 0.03 nmol/L. The ELISA was a sandwich immunoassay with two monoclonal antibodies directed against different epitopes on liraglutide and was according to guidance regarding recovery, accuracy, precision, sensitivity and stability.

A standard one-compartment model with first-order absorption and elimination was the starting point for the description of liraglutide pharmacokinetics, and the model was developed and validated according to the US FDA and European Medicines Agency guidelines. The structural model was parameterized in terms of the following parameters:

- kₚ (absorption rate constant)
- CL/F (apparent clearance)
- V/F (apparent volume of distribution)
The first-order conditional estimation with interaction (FOCE-I) method was used for the population pharmacokinetic analysis, implemented in the non-linear mixed effects modelling (NONMEM) software. Due to pharmacokinetic sampling in steady-state conditions, exposure levels were expected to be much higher than the LLOQ (mean values 500- to 1000-fold above LLOQ). Observed values close to or below the LLOQ were therefore believed to be due to missed doses. Data records with missing concentration values or concentration values below the LLOQ were excluded from the population pharmacokinetic analysis in order not to overestimate CL/F. Stricter exclusion criteria were tested in sensitivity analyses but were found not to be relevant.

### TABLE 1 Trial design and baseline characteristics of the trials included in the population PK analysis

| Trial design | Trial 4180\textsuperscript{23} (NCT02918279) Phase 3a trial in adolescents N = 121 | Trial 4181\textsuperscript{22} (NCT02696148) Phase 1 trial in children N = 13 | Trial 3967\textsuperscript{21} (NCT01789086) Phase 1 trial in adolescents N = 13 | Trial 3630\textsuperscript{20} (NCT00978393) Phase 1 trial in adults N = 29 |
|---------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------|
| Weekly dose escalation steps (mg/day) | 0.6, 1.2, 1.8, 2.4, 3.0 | 0.3, 0.6, 0.9, 1.2, 1.8, 2.4, 3.0 | 0.6, 1.2, 1.8, 2.4, 3.0 | 0.6, 1.2, 1.8, 2.4, 3.0 |
| Maintenance doses (mg/day) | 0.6 (n = 1); 1.2 (n = 1); 1.8 (n = 2); 2.4 (n = 10); 3.0 (n = 107) | 2.4 (n = 1); 3.0 (n = 12) | 2.4 (n = 1); 3.0 (n = 12) | 3.0 |
| Treatment duration\textsuperscript{a} | 56 weeks | 7 weeks | 5-6 weeks | 35 days |
| Completed trial (\% liraglutide vs placebo) | 81%; 79% | 88%; 75% | 93%; 100% | 90% across groups |
| Sparse PK sampling weeks | 8, 12, 16, 30, 42, 56 | NA | NA | NA |
| Number of pre-dose (trough) PK samples during dose escalation | NA | 7\textsuperscript{b} | 4 | NA |
| Number of PK samples after last dose | NA | 5 | 6 | 12 |
| Nominal timing of PK sampling after the last dose | NA | Pre-dose, 1, 2, 3, 24, 72 hours | Varying according to assigned sequence | Pre-dose, 2, 4, 1, 13, 15, 18, 20, 24, 36, 48, 60 hours |
| Demographics | | | | |
| Sex | | | | |
| Female | 67 (55.4%) | 7 (53.8%) | 10 (76.9%) | 11 (37.9%) |
| Male | 54 (44.6%) | 6 (46.2%) | 3 (23.1%) | 18 (62.1%) |
| Race | | | | |
| White | 102 (84.3%) | 6 (46.2%) | 12 (92.3%) | 25 (86.2%) |
| Black or African American | 13 (10.7%) | 7 (53.8%) | - | - |
| Asian | 2 (1.7%) | - | - | - |
| Other | 4 (3.3%) | - | 1 (7.7%) | 4 (13.8%) |
| Ethnicity | | | | |
| Not Hispanic or Latino | 92 (76.0%) | 10 (76.9%) | 13 (100%) | 26 (89.7%) |
| Hispanic or Latino | 29 (24.0%) | 3 (23.1%) | - | 3 (10.3%) |
| Age, years (mean [SD]) | 14.6 (1.6) | 9.8 (0.9) | 15.1 (1) | 47.8 (13.8) |
| Range | 12-17 | 8-11 | 13-16 | 20-72 |
| Body weight, kg (mean [SD]) | 99.4 (19.7) | 69.1 (10.8) | 102.1 (12.2) | 102.3 (15.6) |
| Range | 62.1-178.2 | 53.9-86.8 | 79.9-119.2 | 74.2-131.6 |

Note: Data for demographics are presented as number and percentage of participants, unless otherwise stated. Abbreviations: N, number of participants in the population PK analysis (those on liraglutide); NA, not applicable; PK, pharmacokinetic; SD, standard deviation.

\textsuperscript{a}Including dose-escalation.

\textsuperscript{b}Including one trough sample before last dose.
An analysis of the influence of covariates on exposure was carried out including all tested covariates in one step, using a confirmatory approach,\textsuperscript{32} and disregarding any interactions between age group and sex. Model development involved estimation of a base model without covariates and a full model including all predefined covariates. If any covariates without statistically significant effect could be excluded, the reduced model would be considered the final population pharmacokinetic model. The covariates were prespecified and based on earlier findings in phase 3 trials with adults as well as prior knowledge from smaller trials in children and adolescents.

The covariate effects of baseline body weight, sex and age group were investigated for CL/F. Due to the sparseness of data, as well as the focus on average exposure, only the effect of baseline body weight was investigated with respect to V/F. The CL/F and V/F were parameterized as follows for the $i^{th}$ subject (shown for CL/F only):

$$\text{CL}/F_i = TVCL \cdot E_{\text{body weight,CL}} \cdot E_{\text{sex}} \cdot E_{\text{age group}} \cdot \exp(\eta_{CL,i})$$

(1)

$$E_{\text{body weight,CL}} = \frac{\text{body weight}}{100 \text{ kg}}^{\theta_{\text{body weight,CL}}}$$

(2)

$$E_{\text{sex}} = (\theta_{\text{male}})^{\text{male}}$$

(3)

$$E_{\text{age group}} = (\theta_{\text{child}})^{\text{child}} \cdot (\theta_{\text{adolescent}})^{\text{adolescent}}$$

(4)

$$V/F_i = TVV \cdot E_{\text{body weight,V}} \cdot \exp(\eta_{V,i})$$

(5)

where $TVCL$ and $TVV$ are “typical values (TV)” of apparent clearance (CL/F) and volume of distribution (V/F), respectively, for a reference subject (an adult female with a body weight of 100 kg) and the $\theta$ values are the estimated covariate effect parameters.

Specific steady-state exposures for individual participants were based on the full model, including all covariates, and derived from the subject-specific post-hoc CL/F estimates, the maintenance dose and the dose interval (24 hours):

$$C_{\text{avg}} = \frac{\text{Dose}}{(\text{CL}/F \cdot 24 \text{ hours})}$$

(6)

The CIs for full model parameters were estimated using bootstrapping. For each investigated covariate, differences were considered relevant if the 90% CI of the estimated mean of the relative exposure fell outside the standard bioequivalence limits (0.80-1.25).

Between-subject variability was included for CL/F and V/F, assuming log-normal distributions without correlation between parameters. Furthermore, CL/F and V/F were estimated using a full variance-covariance matrix. No between-subject variability was included for $k_e$. Within-subject variability (residual) was described by a proportional error model. Standard graphical quality analyses, including goodness-of-fit plots (Figure S1), were made during qualification of the pharmacokinetic model.

### 2.2 Exposure-response analysis

The objectives of the exposure-response analysis were: (1) to investigate the exposure-response relationship of liraglutide in adolescents with respect to the change from baseline in BMI SDS and (2) to determine whether the exposure-response relationship for the change from baseline in BMI SDS was similar in adults and adolescents.

An evaluation of the exposure-response of liraglutide in adults has been published previously using data from the phase 2 trial (1807), the phase 3a SCALE Obesity and Prediabetes trial (1839) and the phase 3a SCALE Diabetes trial (1922).\textsuperscript{28} Data from the phase 3a trial 4180 (in adolescents) were additionally included in the present exposure-response assessment of liraglutide (Table 2). The analyses were conducted using response data at week 20 (phase 2 trial 1807), week 50 (phase 3a trial 1922) or week 56 (phase 3a trials 4180 and 1839) to align with the original analysis in adults. The appropriateness of using different trial lengths for the historical adult trials was evaluated by graphically exploring the weight loss longitudinally. This was deemed appropriate as the maximal effect on weight reduction was reached after 20 weeks of treatment and remained stable thereafter for the trials of longer duration.

The exposure variable used for the analysis was individual model-based average liraglutide concentration ($C_{\text{avg}}$) estimates from the population pharmacokinetic analyses of the four trials (for trial 4180, LLOQ samples were included in the exposure-response data set to better reflect actual exposure - accounting for potential non-compliance). For the primary exposure-response analysis of change from baseline in BMI SDS, BMI SDS was calculated according to the World Health Organization (WHO).\textsuperscript{33} For adults, the growth charts for 19-year-old females or males were used for the calculation. The following response variables were evaluated as supportive analyses: changes from baseline in body weight (%), BMI (%), waist circumference (cm) and the proportion of participants achieving a 5% reduction in body weight.

A modified exposure-response model based on the previous meta-analysis\textsuperscript{28} was used in the development of the present exposure-response model. The starting models included the effect of all investigated covariates on the placebo effect $E_0$, and of selected covariates on the $E_{\text{max}}$, as identified based on previous analyses.\textsuperscript{28} The covariate selection was made as follows. Covariates for $E_0$ were removed in a stepwise manner, via backwards elimination based on a likelihood ratio test, keeping the covariates that were both statistically significant ($P < 0.05$) and clear (ie, clearly needed to describe a potential difference in observed exposure-response when stratified by the covariate). Based on the reduced model from the previous step, covariates for $E_{\text{max}}$ were investigated via forwards inclusion. Based on plots of exposure-response stratified by covariates, each covariate factor was considered. If the data indicated differences in the treatment effect, the covariates were investigated by forwards inclusion and included if statistically significant ($P < 0.05$). If the covariates appeared to be relevant based on the plots, they were tested in the following order: predefined covariates for $E_{\text{max}}$ were investigated first, then other clear covariates for $E_{\text{max}}$.

Standard goodness-of-fit plots were made for checking and evaluating the model.
| Trial design | Participants on liraglutide treatment | Participants on placebo treatment | Participants with normoglycemia | Participants with prediabetes | Participants with type 2 diabetes | Weekly dose escalation steps (mg/day) | Maintenance doses (mg/day) | Treatment duration | Completed trial (% liraglutide vs placebo) |
|-------------|-------------------------------------|----------------------------------|---------------------------------|--------------------------------|----------------------------------|-----------------------------------|--------------------------|----------------|-----------------------------------------|
| Trial 4180\(^{23}\) (NCT02918279) Phase 3a trial in adolescents N = 247 | 121 | 331 | 126 | 183 | 62 | 2 | 0.6, 1.2, 1.8, 2.4, 3.0 | 0.6 (n = 1); 1.2 (n = 1); 1.8 (n = 2); 2.4 (n = 10); 3.0 (n = 107) | 56 weeks | 81%; 79% |
| Trial 1807\(^{27}\) (NCT00422058) Phase 2 trial in adults N = 415 | 331 | 2339 | 84 | 205 | 210 | 0 | 0.6, 1.2, 1.8, 2.4, 3.0 | 1.2, 1.8, 2.4, 3.0 | 20 weeks | 85%; 81% |
| Trial 1839\(^{24}\) (NCT01272219) Phase 3a trial in adults N = 3250 | 2339 | 123 | 911 | 715 | 2000 | 0 | 0.6, 1.2, 1.8, 2.4, 3.0 | 3.0 | 56 weeks | 72%; 64% |
| Trial 1922\(^{25}\) (NCT01272232) Phase 3a trial in adults N = 707 | 123 | 1250 | 205 | 1250 | 707 | 707 | 1.8, 3.0 | 1.8, 3.0 | 56 weeks | 77%; 66% |

**Demographics**

**Sex**

| Female | 145 (58.7%) | 313 (75.4%) | 2535 (78%) | 352 (49.8%) |
| Male | 102 (41.3%) | 102 (24.6%) | 715 (22%) | 355 (50.2%) |

**Race**

| White | 217 (87.9%) | 413 (99.5%) | 3072 (94.5%) | 669 (94.6%) |
| Asian | 2 (0.8%) | - | 118 (3.6%) | 16 (2.3%) |
| Black or African American | 19 (7.7%) | - | - | - |
| American Indian or Alaska Native | 1 (0.4%) | - | 7 (0.2%) | 4 (0.6%) |
| Native Hawaiian or other Pacific Islander | - | - | 2 (0.1%) | - |
| Other | 8 (3.2%) | 2 (0.5%) | 51 (1.6%) | 18 (2.5%) |

**Ethnicity**

| Not Hispanic or Latino | 194 (78.5%) | 415 (100%) | 2913 (89.6%) | 668 (90.2%) |
| Hispanic or Latino | 53 (21.5%) | - | 337 (10.4%) | 69 (9.8%) |

**Age, years (mean [SD])**

| 14.5 (1.6) | 46.4 (10.4) | 45.3 (11.9) | 54.8 (10.2) |

**Body weight, kg (mean [SD])**

| 12-17 | 18-65 | 18-78 | 24-82 |
| 100.8 (20.7) | 97.7 (12.9) | 106.7 (21.4) | 106.0 (21.2) |
| 62.1-178.2 | 69.2-141.2 | 63.0-244.0 | 60.1-193.3 |

**BMI, kg/m\(^2\) (mean [SD])**

| 35.6 (5.4) | 34.4 (2.8) | 38.4 (6.3) | 37.2 (6.8) |
| 26.6-58.8 | 29.1-41.0 | 27.0-77.2 | 27.0-67.6 |

**BMI SDS\(^{b}\) (mean [SD])**

| 3.2 (0.7) | 2.8 (0.4) | 3.4 (1.0) | 3.2 (1.1) |
| 2.1-6.5 | 1.9-3.7 | 1.4-9.3 | 1.4-8.4 |

Note: Data for demographics are presented as number and percentage of participants, unless otherwise stated. Abbreviations: BMI, body mass index; BMI SDS, body mass index standard deviation score; N, number of participants in the exposure-response analysis; NA, not applicable; PK, pharmacokinetic; SD, standard deviation. \(^a\)Including dose-escalation. \(^b\)BMI SDS represents the number of SDs from a reference standard population mean BMI.
The final models were parameterized as follows:

\[
\text{BMI SDS}_{\text{CFB}} = E_0 + E_{\text{max}} \frac{C_{\text{avg}}}{C_{\text{avg}} + EC_{50}} + E_{\text{cov}} + e,
\]

where CFB denoted the “change from baseline” in BMI SDS. \(E_0\) was the placebo effect, \(E_{\text{max}}\) was the maximal drug effect, \(EC_{50}\) the exposure leading to half-maximum effect, \(E_{\text{cov}}\) covariate effects on the overall response (ie, placebo and treatment-related responses) and \(e\) the normal distributed residual error. Sex, baseline response parameter, age group and trial factors were considered as covariates. Baseline BMI SDS and age group were included as covariates in the final model.

The exposure-response evaluation for changes from baseline in body weight, BMI and waist circumference were evaluated using similar models as for the BMI SDS:

\[
\text{Body weight}_{\text{CFB}} = E_0 + E_{\text{max}} \left(1 + \left(\text{male}\right) \gamma \right) \frac{C_{\text{avg}}}{C_{\text{avg}} + EC_{50}} + E_{\text{cov}} + e
\]

\[
\text{BMI}_{\text{CFB}} = E_0 + E_{\text{max}} \left(1 + \left(\text{male}\right) \gamma \right) \frac{C_{\text{avg}}}{C_{\text{avg}} + EC_{50}} + E_{\text{cov}} + e
\]

\[
\text{Waist}_{\text{CFB}} = E_0 + E_{\text{max}} \frac{C_{\text{avg}}}{C_{\text{avg}} + EC_{50}} + E_{\text{cov}} + e
\]

\(1_{\text{adolescent}}\) was the covariate representing a lower \(E_{\text{max}}\) in adolescents compared to adults, where adolescent was an indicator variable for adolescents. \(1_{\text{male}}\) was the covariate representing a lower \(E_{\text{max}}\) in males compared to females, where \(\gamma\) was an indicator variable for males. \(\gamma\) was the Hill coefficient. In the final models, \(E_{\text{cov}}\) covariate effects for body weight and BMI included age group, while for waist circumference the covariate effects were baseline waist circumference (cm), age group and diabetic state (ie, diabetes vs non-diabetes). The “non-diabetes” also included individuals with prediabetes. The other expressions were as defined above.

The exposure-response relationship for the proportion of participants with at least 5% reduction in body weight (body weight responder rate [BW,RR]) was estimated using logistic regression analyses parameterized as:

\[
\text{BW,RR} \sim \text{invlogit} \left(\frac{E_{\text{max}} \left(1 + 1_{\text{male}}\right) \gamma \frac{C_{\text{avg}}}{C_{\text{avg}} + EC_{50}} + E_0 + E_{\text{cov}}}{E_{\text{max}} \left(1 + 1_{\text{male}}\right) \gamma \frac{C_{\text{avg}}}{C_{\text{avg}} + EC_{50}} + E_0 + E_{\text{cov}}}\right)
\]

The expressions were as defined above. In the final model, age group was included as a covariate effect.

### 2.3 | Data analysis software

The software program NONMEM (ICON Development Solutions, Ellicott City, MD, USA) version 7.3 was used for the population pharmacokinetic analysis. R version 3.2.3 (R Foundation, Revolution Analytics, Mountain View, CA, USA) was used for data file processing, explorative data analysis and plotting. Exposure-response analyses were implemented in R.

### 3 | RESULTS

#### 3.1 | Population pharmacokinetic analysis

##### 3.1.1 | Model development and qualification

A base model without covariates was developed based on prior knowledge of liraglutide pharmacokinetics. Subsequently, a full model, which was the base model with all investigated covariates included, was estimated for the covariate analysis. All investigated covariates were significant, except for age group. The full model could therefore not be reduced and, therefore, the full model was adopted as the final model.

A one-compartment model successfully described the pharmacokinetics of liraglutide. The parameter estimates from the base and final models are presented in Table S1 and Table 3, respectively.

##### 3.1.2 | Demographics and datasets

A total of 176 individuals was included in the analysis: 13 children from trial 4181, 121 adolescents from trial 4180, 13 adolescents from trial 3967 and 29 adults from trial 3630 (Table 1). The majority of participants (82.4%) were white (11.4% were black or African American) and non-Hispanic (80.1%).

For trial 4180, the final dataset comprised 646 pharmacokinetic observations from 121 adolescents (94 observations [14.6% of the final dataset] were below the LLOQ; 22 samples from two individuals were excluded during data cleaning due to timing being missing or an inadequate dosing history. There were also many lower-than-expected individual liraglutide values in trial 4180 (Figure S2). Data cleaning for trials 4181, 3967 and 3630 was conducted similarly to previously reported analyses. The number of observations included for each trial as well as the number of samples below the LLOQ and excluded is presented by trial in the Supporting Information.

##### 3.1.3 | Covariate analysis

The effects of intrinsic covariates on liraglutide exposure are shown in Figure 1. All covariates were tested simultaneously; thus for a given covariate effect, the effects of the other covariates were accounted for. In accordance with previous findings in adults, body weight was the main intrinsic covariate affecting liraglutide exposure, with lower exposure at higher body weights. Age group was of no importance and sex was of little importance.

The inverse relationship between body weight and exposure was apparent across trials and age groups (Figure 2).
I n d i v i d u a l  a n d  m e a n  Cavg values appeared to be similar in adolescents and adults when adjusted to the 3.0 mg dose, whereas children had slightly higher liraglutide concentrations (Figure 3A). When adjusted for differences in body weight, however, exposures were similar across all age groups (Figure 3B).

### 3.2 | Exposure-response analysis

#### 3.2.1 | Demographics and datasets

In the exposure-response full dataset, there were more females than males in most of the trials, except for trial 1922 in which sex was evenly distributed (Table 2). Otherwise the demographic characteristics were generally similar across trials. Several race and ethnicity categories were represented for both adolescents and adults, with 7.7% of participants black or African American in trial 4180 (Table 2). The mean baseline body weight was 100.8 kg in adolescents from trial 4180 and ranged from 97.7 kg in the phase 2 trial in adults (trial 1807) to 106.7 kg in the largest phase 3 trial in adults (trial 1839). Baseline body weights and BMI SDS values were similar between adolescents and adults (Figure S3).

The exposure-response dataset comprised placebo and treatment data from a total of 4619 individuals: 4372 adults from trials 1807, 1839 and 1922 and 247 adolescents from trial 4180. Last-observation-carried-forward (LOCF) imputation was used if the observation...
was missing at the end of treatment, consistent with the original analysis. Data for waist circumference were not obtained for all adult trial participants, so this dataset included a total of 3163 individuals.

### 3.2.2 Liraglutide exposure-response relationship in adolescents and adults

An exposure-response relationship was demonstrated for liraglutide in both adolescents and adults with a greater decrease from baseline in BMI SDS as liraglutide exposure increased (Figure 4A). Similarly, an exposure-response relationship, with larger responses at higher exposures, was identified for changes from baseline in BMI (%) (Figure 4B), body weight (%) (Figure 4C), waist circumference (Figure 4D) as well as the proportion of participants achieving at least a 5% weight loss (Figure 4E).

For all response variables, similar exposure-response relationships were seen in both adolescents and adults. The two age groups differed in the placebo group response, with lower mean responses in adolescents; this difference was consistent across the exposure range as well. There was a tendency for a steeper exposure-response in adolescents for all response variables, leading to overlapping effects with adults at the highest liraglutide exposures. However, the effect of age group on the maximal effect in the model (E_{max}) was not significant in these analyses. The exposure-response for adolescents in trial 4180 who gained height of at least 0.5 cm during the trial was not significantly different from the exposure-response for those who had reached their final height. Approximately 64% of the adolescents in the exposure-response analysis grew more than 0.5 cm during the trial. Of these, some received liraglutide and some received placebo.

For those receiving placebo treatment, the average growth was 1.8 cm, while for those receiving liraglutide treatment, the average growth was 1.4 cm (data not shown).

In accordance with the similar demographic characteristics of adults and adolescents included in the exposure-response analyses, the exposures of liraglutide were overlapping in the two populations, as shown for all participants with exposures adjusted to the liraglutide 3.0 mg dose (Figure S4). In trial 4180, liraglutide exposures were also similar in adolescents in earlier stages of pubertal development (Tanner stages 2 and 3) as compared with those in later stages (Tanner stages 4 and 5) (Figure S5).

### 4 DISCUSSION

The present population pharmacokinetic analysis demonstrates that the principal covariate affecting the exposure of liraglutide 3.0 mg is body weight, with lower exposures as body weight increases. Age group was not an important covariate and, as the body weight ranges were similar between adolescent trial participants and historical adult trial participants, the exposure ranges in the two populations were comparable. An exposure-response relationship was demonstrated for liraglutide in both adolescents and adults, with larger responses at higher exposures. Thus, as liraglutide exposures increased, the mean changes from baseline in BMI SDS, BMI (%), body weight (%), waist circumference and the proportion of participants achieving at least a 5% weight loss increased in an exposure-dependent manner.

The recommended 3.0 mg liraglutide dose was shown to provide similar exposures in adolescents and adults, even without adjusting for baseline body weight, which supports using the same dose as approved in adults for weight management in adolescents. The added benefits of increasing liraglutide exposures, in terms of a greater reduction in BMI SDS and greater changes in other response variables, were similar in adolescents with obesity and adults with overweight or obesity, thus further supporting the 3.0 mg treatment dose. While individual and mean exposures appeared to be slightly higher in children than in adolescents or adults, exposures were similar across age groups after adjusting for body weight.

Although body weight was found to be the most important covariate affecting exposure and an exposure-response relationship was demonstrated, the maintenance dose for liraglutide 3.0 mg is not dose-adjusted according to body weight. All phase 3 trials in the
liraglutide development program evaluated a 3.0 mg maintenance dose in both adults and paediatric patients without weight-adjusted dosing. Furthermore, weight loss has been shown to vary according to initial body weight, and a clinically relevant weight loss is achieved across baseline BMI categories, supporting that weight-adjusted dosing is not necessary.

A previous population pharmacokinetic analysis of liraglutide dosed up to 3.0 mg also identified body weight as a key covariate affecting exposure in adults with overweight or obesity with or without type 2 diabetes mellitus; sex was additionally identified as a key covariate in this large phase 3 dataset. The effect of sex was less clear in the present smaller adolescent dataset.

The primary response variable used for the present exposure-response analysis of liraglutide for weight management was BMI SDS, which is a measure of the number of SDs from the population mean BMI, matched for age and sex. The exposure-response relationship

**FIGURE 4** Liraglutide exposure-response relationships in adults and adolescents. Figures show the weight-related outcomes for liraglutide exposure in adolescents and adults with respect to A: BMI SDS, B: BMI (%), C: body weight (%), D: waist circumference (cm) and E: the proportion of individuals achieving 5% weight reduction. Data points with error bars are arithmetic means with 95% CIs for each of 3 quantiles of Cavg for liraglutide and one quantile for placebo (at Cavg of 0 nmol/L). Lines are covariate-adjusted, model derived relationships. Data are included from trial 1807 after 20 weeks of treatment, trials 4180 and 1839 after 56 weeks of treatment and trial 1922 after 50 weeks of treatment. BMI, body mass index, BMI SDS, body mass index SD score; Cavg, average liraglutide concentration; CI, confidence interval; N, number of participants; PK, pharmacokinetic.
with respect to change from baseline in BMI SDS was successfully described by an $E_{\text{max}}$ model with baseline BMI SDS and age group as covariates on the placebo response. As the BMI SDS tends to be skewed at higher BMI SDS values,\textsuperscript{33} it was important to evaluate results together with other weight-related parameters. Supportive exposure-response analyses were therefore also conducted for the percentage change from baseline in body weight and BMI, change from baseline in waist circumference as well as for the proportion of participants achieving at least 5% weight loss. The exposure-response relationships for the additional response variables were all successfully described by $E_{\text{max}}$ models of exposure-response.

The difference in the weight-loss responses observed between adults and adolescents in the placebo group, where adults achieved a greater mean weight loss than adolescents, could have been due to lower adherence to active treatment and/or the diet and exercise program in some adolescents, although linear growth (height increase) in a part of the adolescent population might also have contributed. There were several lower-than-expected individual liraglutide concentration measurements in trial 4180 in adolescents, which could also potentially indicate a reduced adherence to the trial medication for some trial participants. Low liraglutide concentrations were also observed in four children in trial 4181, although the reason for this was unknown.\textsuperscript{22} Lower adherence in adolescents compared with adults is also a significant concern in diabetes management.\textsuperscript{39} Clinically, there is an ongoing focus to reinforce adherence to both medication and behavioural strategies such as diet and exercise in both diabetes and obesity management.\textsuperscript{39\textendash}41 In contrast, for adolescents with the highest liraglutide exposures, there was a tendency for weight-loss responses being comparable to the adult responses. Although the tendency for a steeper exposure-response in adolescents was not significant in this small data set, the exposure estimates in trial 4180 were based on sparse pharmacokinetic sampling over a long time period, and the estimates could be influenced by samples obtained during periods with reduced or varying adherence to treatment. Thus, the individual exposure estimates also could reflect the adherence level.

Limitations of the present exposure-response analysis include a relatively low number of paediatric participants as well as some variability in the populations included in terms of inclusion criteria and background lifestyle therapy. Trial design differed between the trials and included counselling in healthy nutrition and physical activity for all participants, but for the adult participants in trials 1839 and 1922 a calorie-restricted diet was also included. Moreover, the reduced adherence to treatment in adolescents compared with adults as speculated above could not be confirmed in practice. In the exposure-response analysis, the value for a 19-year-old female or male was used as the adult value for the BMI SDS response variable,\textsuperscript{73} assuming that these growth charts would be representative also for older adults. Strengths of the analysis include an overall high rate of completion among participants across trials. Long-term exposure data were available through sparse pharmacokinetic sampling, and the availability of data in participants receiving placebo allowed for the described exposure-response analyses. Finally, the availability of large datasets in adults allowed for prior model development, joint analyses and outcome comparisons.

In summary, the population pharmacokinetic analysis indicated similar exposures in adolescents and adults, with similar body weight ranges between the studied adolescents and adults. Body weight was identified as the most important covariate affecting exposure. An exposure-response relationship with a larger response with increasing liraglutide exposure was established for BMI SDS change from baseline as well as the percentage change from baseline in body weight and BMI, change from baseline in waist circumference, and for the proportion of participants with at least 5% weight loss. While a lower mean response was observed in adolescents compared to adults, consistent with the placebo response across the exposure range, the added benefits of increasing liraglutide exposures were similar in adolescent and adults with overweight or obesity.

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KCCP and PMH planned the trial and the modelling analysis. DH was involved in the conduct of the trial and planning of the modelling analysis. LDM was involved in trial conduct as investigator. KCCP conducted the modelling analysis. All authors were involved in interpreting the analysis results and writing the paper and had final approval of the submitted and published versions. The authors wish to thank Angela Stocks, PhD (Larix A/S, Copenhagen, Denmark) for editorial and medical writing services, which were funded by Novo Nordisk.

**CONFLICT OF INTEREST**

KCCP, PMH, DH and NR are employees of Novo Nordisk and KCCP, PMH and DH own stocks in Novo Nordisk. LDM has received grant support paid to the University at Buffalo from AstraZeneca and Novo Nordisk and fees for serving as a healthcare professional consultant from Novo Nordisk.

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**REFERENCES**

1. Sanyaolu A, Okorie C, Qi X, Locke J, Rehman S. Childhood and adolescent obesity in the United States: a public health concern. *Glob Pediatr Health*. 2019;6:1-11.
2. Hales CM, Carroll MD, Fryar CD, Ogden CL. Prevalence of obesity among adults and youth: United States, 2015-2016. *NCHS Data Brief*. 2017;288:1-8.
3. Afshin A, Forouzanfar MH, Reitsma MB, et al. Health effects of overweight and obesity in 195 countries over 25 years. *N Engl J Med*. 2017;377(1):13-27.
4. NCD Risk Factor Collaboration (NCD-RisC). Worldwide trends in body-mass index, underweight, overweight, and obesity from 1975 to 2016: a pooled analysis of 2416 population-based measurement studies in 128·9 million children, adolescents, and adults. *Lancet*. 2017;390(10113):2627-2642.
5. Singh AS, Mulder C, Twisk JW, van Mechelen W, Chinapaw MJ. Tracking of childhood overweight into adulthood: a systematic review of the literature. *Obes Rev*. 2008;9(5):474-488.
6. Ward ZJ, Long MW, Resch SC, Giles CM, Cradock AL, Gotmarker SL. Simulation of growth trajectories of childhood obesity into adulthood. *N Engl J Med*. 2017;377(22):2145-2153.
7. Wabitsch M. Overweight and obesity in European children: definition and diagnostic procedures, risk factors and consequences for later health outcome. Eur J Pediatr. 2000;159(Suppl 1):S8-513.

8. Lakshman R, Elks CE, Ong KK. Childhood obesity. Circulation. 2012;126(14):1770-1779.

9. Di Cesare M, Sorić M, Bovet P, et al. The epidemiological burden of obesity in childhood: a worldwide epidemic requiring urgent action. BMC Med. 2019;17(1):212.

10. Nittari G, Sciri S, Petrelli F, Pirillo I, di Luca NM, Grappasonni I. Fighting obesity in children from European World Health Organization member states. Epidemiological data, medical-social aspects, and prevention programs. Clin Ter. 2019;170(3):e223-e230.

11. Gungor N, Thompson T, Sutton-Tyrrell K, Janosky J, Arslanian S. Early signs of cardiovascular disease in youth with obesity and type 2 diabetes. Diabetes Care. 2005;28(5):1219-1221.

12. Singh GM, Danaei G, Farzadfar F, et al. The age-specific quantitative effects of metabolic risk factors on cardiovascular diseases and diabetes: a pooled analysis. PLoS One. 2013;8(7):e65174.

13. Li Q, Blume SW, Huang JC, Hammer M, Graf TR. The economic burden of obesity by glycemic stage in the United States. Pharmacoeconomics. 2015;33(7):735-748.

14. Chu DT, Minh Nguyet NT, Dinh TC, et al. An update on physical activity for the prevention of obesity in children, adolescents, and young adults. Int J Obes. 2015;39(10):1521-1526.

15. MacLean PS, Higgins JA, Giles ED, Sherk VD, Jackman MR. The role for adipose tissue in weight regain after weight loss. Obes Rev. 2015;16(Suppl 1):45-54.

16. Czepiel KS, Perez NP, Campoverde Reyes KJ, Sabharwal S, Stanford FC. Pharmacotherapy for the treatment of overweight and obesity in children, adolescents, and young adults in a large health system in the US. Front Endocrinol. 2020;11:290.

17. Lagorros YT, Rössner S. Obesity management: what brings success? Therap Adv Gastroenterol. 2013;6(1):77-88.

18. Holst JJ. The physiology of glucagon-like peptide 1. Physiol Rev. 2007;87(4):1409-1439.

19. Drucker DJ, Nauck MA. The incretin system: glucagon-like peptide-1 receptor agonists and dipeptidyl peptidase-4 inhibitors in type 2 diabetes. Lancet. 2006;368(9548):1696-1705.

20. van Can J, Sloth B, Jensen CB, Flint A, Blaak EE, Saris WH. Effects of the once-daily GLP-1 analog liraglutide on gastric emptying, glycemic parameters, appetite and energy metabolism in obese, non-diabetic adults. Int J Obes. 2014;38(6):784-793.

21. Danne T, Biester T, Kapitzke K, et al. Liraglutide in an adolescent population with obesity: a randomized, double-blind, placebo-controlled 5-week trial to assess safety, tolerability, and pharmacokinetics of liraglutide in adolescents aged 12–17 years. J Pediatr. 2017;181:146-153.e3.

22. Mastrandrea LD, Witten L, Carlsson Petri KC, Hale PM, Hedman HK, Riesenber RA. Liraglutide effects in a paediatric (7-11 y) population with obesity: a randomized, double-blind, placebo-controlled, short-term trial to assess safety, tolerability, and pharmacokinetics of liraglutide in adolescents aged 12–17 years. J Pediatr. 2017;181:146-153.e3.

23. Kelly AS, Auerbach P, Barrientos-Perez M, et al. A randomized, controlled trial of liraglutide for adolescents with obesity. N Engl J Med. 2020;382(22):2117-2128.

24. Pi-Sunyer X, Astrup A, Fujioka K, et al. A randomized, controlled trial of 3.0 mg of liraglutide in weight management. N Eng J Med. 2015;373(1):11-22.

25. Davies MJ, Bergentst R, Bode B, et al. Efficacy of liraglutide for weight loss among patients with type 2 diabetes: the scale diabetes randomized clinical trial. JAMA. 2015;314(7):687-699.

26. Overgaard RV, Petri KC, Jacobsen LV, Jensen CB. Liraglutide 3.0 mg for weight management: a population pharmacokinetic analysis. Clin Pharmacokinet. 2016;55(11):1413-1422.

27. Astrup A, Rössner S, Van Gaal L, et al. Effects of liraglutide in the treatment of obesity: a randomised, double-blind, placebo-controlled study. Lancet. 2009;374(9701):1606-1616.

28. Wilding JP, Overgaard RV, Jacobsen LV, Jensen CB, le Roux CW. Exposure-response analyses of liraglutide 3.0 mg for weight management. Diabetes Obes Metab. 2016;18(5):491-499.

29. Office of Clinical Pharmacology Review. NDA for Saxenda. 2013. https://www.accessdata.fda.gov/drugsatfda_docs/nda/2014/206321Orig1s000ClinPharmRPdf.pdf. Accessed January 2021.

30. Food and Drug Administration. Guidance for industry: population pharmacokinetics. 2019. https://www.fda.gov/media/128793/download. Accessed August 2020.

31. European Medicines Agency. Guideline on the investigation of drug interactions. 2012. https://www.ema.europa.eu/en/documents/scientific-guideline/guideline-investigation-drug-interactions-revision-1_en.pdf. Accessed August 2020.

32. Hu C, Zhang J, Zhou H. Confirmatory analysis for phase III population pharmacokinetics. Pharm Stat. 2013;10(1):14-26.

33. de Onis M, Onyango AW, Borghi E, Siyam A, Nishida C, Siekmann J. Development of a WHO growth reference for school-aged children and adolescents. Bull World Health Organ. 2007;85(9):660-667.

34. Food and Drug Administration. Prescribing information for Saxenda. 2014. https://www.accessdata.fda.gov/drugsatfda_docs/label/2014/206321Orig1s000lbl.pdf. Accessed January 2021.

35. European Medicines Agency. Summary of product characteristics for Saxenda. 2015. https://www.ema.europa.eu/en/documents/product-information/saxenda-epar-product-information_en.pdf. Accessed January 2021.

36. Blackman A, Foster GD, Zammit G, et al. Effect of liraglutide 3.0 mg in individuals with obesity and moderate or severe obstructive sleep apnea: the SCALE sleep apnea randomized clinical trial. Int J Obes. 2016;40(8):1310-1319.

37. Wadden TA, Hollander P, Klein S, et al. Weight maintenance and additional weight loss with liraglutide after low-calorie-diet-induced weight loss: the SCALE maintenance randomized study. Int J Obes. 2013;37(11):1443-1451.

38. le Roux C, Arøda V, Hemmingsson J, Cancino AP, Christensen R, Pi-Sunyer X. Comparison of efficacy and safety of liraglutide 3.0 mg in individuals with BMI above and below 35 kg/m2: a post-hoc analysis. Obes Facts. 2017;10(6):531-544.

39. Cox L, Hunt J. Factors that affect adolescents’ adherence to diabetes treatment. Nurs Child Young People. 2015;27(1):16-21.

40. Gandhi K, Vu BK, Eshtehardi SS, Wasserman RM, Hilliard ME. Adherence to medications for assessment in research and practice. Diabetes Manag. 2015;5(6):485-498.

41. Bean MK, Mazzeo SE, Stern M, Bowen D, Ingersoll K. A values-based motivational interviewing (MI) intervention for pediatric obesity: study design and methods for MI values. Contemp Clin Trials. 2011;32(5):667-674.

**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of this article.