PEX-168 improves insulin resistance, inflammatory response and adipokines in simple obese mice: a mechanistic exploration

Zeyuan Guo¹, Yuting Wu¹, Lihua Zhu², Yong Wang³, Daorong Wang²,³,⁴ and Xiaofang Sun¹,²*

Abstract

Background: Polyethylene glycol loxenatide (PEX-168) is a new antidiabetic drug; as such, there are not yet any reports on its weight loss effect. Therefore, this trial was designed to investigate the effect of PEX-168 on simple obese mice.

Methods: Thirty healthy male C57BL/6 mice were randomly selected and divided into a control group (NC) and an obesity model group. The high-fat diet-induced simple obesity mice were divided into a model control group (HF) and three intervention groups. The intervention groups were injected with different doses of PEX-168 intraperitoneally once a week for 12 weeks (low (LD), medium (MD) and high (HD)). Fasting blood glucose (FBG), body weight and food intake were measured from 1 to 12 weeks after PEX-168 injection. The serum insulin (INS), C-reactive protein (CRP), chemerin and omentin levels were measured after 12 weeks.

Results: Compared with the HF group, the low dose of PEX-168 reduced the body weight of the mice in a short period of time (8 weeks), and the mice in the MD and HD groups showed a significant decrease in body weight (P < 0.05). The low dose of PEX-168 could effectively improve the blood glucose and homeostasis model assessment of insulin resistance (Homa-IR) of the mice (FBG P < 0.05 INS, Homa-IR P < 0.001), but there was no significant difference between different doses (P > 0.05). CRP levels in the MD and HD groups were significantly improved (P < 0.05). The levels of serum chemerin and omentin in the intervention groups were also significantly improved (P < 0.01), but there was no significant difference between the different doses (P > 0.05).

Conclusions: PEX-168 significantly reduced the body weight of simple obese mice and improved the insulin resistance. PEX-168 may regulate the expression of chemerin and omentin through its hypoglycaemic effect, and the weight-reducing effect of PEX-168 is unlikely to be the reason for the changes in both.

Keywords: GLP-1, weight loss, chemerin, omentin, Homa-IR, CRP
Introduction
The incidence rate of obesity has doubled in recent decades, which has caused health challenges among the world population [1]. Obesity is not only a risk factor of cardiovascular disease and type 2 diabetes mellitus [2], but also increases the financial burden of medical care [3]. Lifestyle (e.g., diet and exercise) and behaviour changes are the cornerstones of obesity management [4], but they are difficult for patients to comply with and maintain, and effective noninvasive treatments are limited[5]. The risks of bariatric surgery are high, and the surgery is only suitable for a small group of patients [6]. Considering the huge financial burden caused by obesity, the cost of interventions that can sustain weight loss is also critical. Therefore, new treatment options are of great significance for overweight and obese patients without type 2 diabetes to better manage their weight, and prevent or improve insulin resistance and inflammation, thereby reducing the risk of diabetes and improving their quality of life [7].

GLP-1 is a peptide substance that has a series of physiological functions such as promoting insulin release, promoting the proliferation and differentiation of pancreatic β-cells, increasing satiety and reducing food intake, and has a low risk of hypoglycaemia [8]. Many studies have demonstrated that overweight or obese people with type 2 diabetes have a better tolerance to the GLP-1 receptor agonist liraglutide, which can reduce food intake without increasing energy consumption, to achieve a good weight loss goal [9, 10]. However, due to the action of dipeptidyl peptidase IV, the half-life of GLP-1 in vivo is short, and several repeated doses are required to achieve therapeutic effects. Compared with natural GLP-1, although liraglutide is not easily and rapidly degraded by dipeptidyl peptidease IV in vivo, its half-life is also only 13 h, and it still needs to be injected subcutaneously once a day, which affects patient compliance. PEX-168 is modified by amino acids and pegylation of the chemical structure of Exenatide, which is the first long-acting GLP-1 receptor agonist (GLP-1 RA) developed independently in China, and has entered the Chinese health insurance system. Its introduction for the treatment of obesity can significantly reduce medical costs. PEX-168 becomes a long-acting formulation through the modification of polyethylene glycol, and only needs to be administered once a week, thus increasing patient compliance.

However, research in Jessica and Tina pointed out that further research is still needed to determine the effect of GLP-1 receptor agonists in overweight or obese patients without type 2 diabetes [11, 12]. Studies have proven the efficacy and safety of PEX-168 in the treatment of patients with T2DM [13, 14], but the effect of PEX-168 on simple obesity and on the regulation of chemerin and omentin expression has not been reported. This study therefore selected six indicators to evaluate the effect of PEX-168 according to prior research and the drug characteristics. Serum omentin and chemerin are newly discovered novel adipokines. As an inflammatory chemokine and adipokine, chemerin is considered to be a common factor between obesity and T2DM [15]. Recent studies have shown that plasma levels and gene expression of omentin-1 are negatively associated with obesity and insulin resistance [16]. Both are closely related to obesity and disorders of glycolipid metabolism and can reflect the hyperlipidaemia and inflammatory state of patients. Obesity is a significant determinant of insulin resistance, and adipose tissue plays a key role in insulin resistance [17]; insulin resistance is a common pathophysiological condition of obesity and diabetes [18]. Diet-induced obesity is associated with chronic low-grade inflammation of the hypothalamus. Metabolic inflammation is also a significant feature of obesity. Obesity is associated with an increased risk of cardiovascular disease (CVD), and the risk factor is modifiable [19]. CRP is now one of the most powerful predictors of inflammatory and lipid markers of cardiovascular events [20]. Obesity, inflammation and insulin resistance are closely related; obese patients without dyslipidaemia, insulin resistance, inflammatory state, and impaired fasting blood sugar have a very low risk of developing diabetes [21].

Therefore, the aim of this study was to observe the effects of different doses of PEX-168 on body weight, intake, Homa-IR, CRP, and the adipokines chemerin and omentin in simple obese mice induced by a high-fat diet, to observe whether there was variability between different doses and to provide a basis for subsequent clinical studies.

Materials and methods
Reagents and materials
Long-acting glucagon-like peptide-1 analogue polyethylene glycol losenatide (PEX-168) was provided by Hausen Pharmaceutical Co., Ltd (Jiangsu, China), lot number: H20190024, specification: 0.5 ML (0.1 mg) per bottle, stored at 4–8°C, Elisa kit was purchased from Liko Biotechnology Co., Ltd (Nanjing, China), precision electronic balance was purchased from Shuangjie Electronics Co., Ltd (Jiangsu, China). Automatic biochemical analyzer (Hitachi, Japan), Glucometer (Roche, Germany).

The basic diet and high-fat diet were purchased from Yangzhou University College of Veterinary Medicine (Jiangsu, China) and Medison Biomedical Co., Ltd (Jiangsu, China) respectively. The basic diet was composed of 22.8% protein, 13.8% fat, and 63.4% carbohydrate, the high-fat diet was composed of 26.2% protein, 34.9% fat, and 26.3% carbohydrate. The caloric density
was 3656 kcal/kg for the basic diet and 5240 kcal/kg for the high-fat diet.

Animals

Animal model and test methods

Thirty-six-week-old SPF-grade male C57BL/6 mice were purchased from the College of Agriculture, Yangzhou University, and housed in cages at room temperature between 18 and 24 °C. They were first fed normal chow for one week for acclimation and exposed to light for 12 H daily, with unlimited access to chow and water. This experiment was approved by the Animal Ethics Committee of Yangzhou University. Starting from the age of seven weeks, they were randomly divided into five groups, and one group (NC group, n = 6) was randomly selected separate from the five groups as a control group and continued to be fed with ordinary feed, and the rest of the groups were fed high-fat feed to establish an obesity model until the weight of the obesity model group exceeded 20% of the control group, and there was no difference in body weight between the obesity groups, meanings that the model is successful. After successful modeling, the four obesity model groups were randomly divided into three intervention groups: low dose (0.03 mg/kg) group (LD group, n = 6), medium dose (0.1 mg/kg) group (MD group, n = 6), high dose (0.33 mg/kg) group (HD group, n = 6) and an obese control group (HF group, n = 6).

After grouping, the three intervention groups and the obese control group transitioned from a high-fat diet to a normal diet after one week. The intervention started at the end of the dietary transition period and lasted for 12 weeks. The intervention groups were injected intraperitoneally with different doses of PEX-168, according to their specific subgroup, once a week at 8 a.m. on Tuesday; the HF and NC groups were injected intraperitoneally with the same volume of saline. Before the intervention, the mean FBG of mice in the NC group was 7.08 mmol/L and that of obese mice in the four groups was 7.89 mmol/L. The FBG was within the normal range, indicating that the mice were all non-diabetic simple obese mice. The body weight and food intake of the mice were measured regularly every day. At the end of the animal experiment (Day 7 after the twelfth intervention), all mice were fasted overnight for 12 h, blood was collected from the eyes, and the supernatant was collected after centrifugation at 5000 r/min for 15 min.

Indicators and measurement

(1) General condition: daily observation of the mice’s activities, including observation of their appearance, urine, stool, activity, gait, spirit, appetite, and any abnormal conditions.

(2) Determination of body weight, food intake and blood glucose: The body weight of mice was measured regularly every day, the appropriate amount of food was placed in the enclosures and weighed before putting in, and the weight of the remaining feed was weighed the next day (the large pieces of residue in the cage box of each group were weighed and recorded, and the residue crumbs were ignored), and the daily food intake was calculated. Mice were fasted for 12 h each week without the restriction of water intake, and their fasting blood glucose was measured regularly by tail-tip cutting and hand-held glucometer using the next morning for 12 weeks.

(3) Serum biochemical analysis: At the end of 12 weeks intervention, all mice were fasted overnight for 12 H, and they were executed by cervical dislocation after eye blood collection, and the supernatant was preserved after centrifugation at 5000r/min for 15 min. The levels of FBG, INS and CRP were detected, and the level of Homa-IR was calculated according to the steady-state model method formula: Homa-IR index = FBG (mmol/L) x INS (mIU/L) / 22.5. The serum chemerin and omentin levels were determined in mice according to the kit instructions.

Statistical analysis

Analysis was performed using SPSS 22.0. The measurement data were presented as the means ± standard deviation (x ± s); statistical comparisons among groups were performed using one-way analysis of variance (ANOVA) and t-tests; Pearson correlation analysis was used for correlation; P < 0.05 was considered statistically significant.

Results

General conditions

All three groups of mice in the intervention group had a significant decrease in activity and water intake after being injected with PEX-168 during the first two weeks, especially in the three days after the injection. The inhibitory effect gradually improved with the extension of the intervention time, and the LD group recovered the fastest. During the intervention period, all mice had dark and shiny coats.

Effect on the intake of mice

Food intake in the three intervention groups had a significant decrease during the first three weeks. The inhibition of food intake was strongest on the first day of each intervention week, and gradually improved in the second six days. In a word, the food intake suppression effect gradually weakened with the duration of the intervention (Fig. 1). Compared with the HF group, the food intake of the three intervention groups was significantly suppressed on the first day of injection, and the food intake of the LD and MD groups was significantly lower than that of the NC group at the end of the intervention period.
reduced in the intervention period, and the difference was statistically significant (LD $P < 0.05$, MD $P = 0.004 < 0.01$, HD $P = 0.002 < 0.01$). In the late intervention period, the difference in food intake between the HF and LD groups was not statistically significant ($P > 0.05$).

Effects on body weight, blood glucose, and insulin in mice

The mean weight of the mice in the control group before the start of the experiment was 21.89 g. There was no significant difference in weight between all the obese mice groups ($P > 0.05$). During the intervention period, the body weight of mice in the HF group was relatively stable. After the intervention, the body weight of the mice in the three groups was still higher than that in the NC group, and the difference was statistically significant ($P < 0.001$). Compared with the HF group, the weight of the mice in the three groups had a downward trend, among which the body weight of mice in MD and HD groups decreased significantly and the difference was statistically significant (MD $P = 0.009 < 0.01$, HD $P = 0.025 < 0.05$). The weight loss of mice in the LD group was statistically significant in the first eight weeks ($P < 0.05$) (Tables 1 and 2).

After the intervention, the levels of FBG, INS, and Homa-IR of mice in all four groups were significantly higher than those in the NC group, with statistically significant differences ($P < 0.001$). When compared with the HF group, the levels of FBG, INS and Homa-IR of obese mice in the three intervention groups were significantly lower, with statistically significant differences (FBG $P < 0.05$, INS Homa-IR $P < 0.001$). There were no significant differences in the improvement of FBG, INS and Homa-IR in mice between the three intervention groups. In the 10th week of the intervention, a transient hypoglycaemic event occurred in the MD and HD groups, with four and three mice having FBG below 5 mmol/L, with

![Fig. 1 The food intake of each group was compared for 12 weeks. Data are means ± s.d. Values were expressed as a box and whisker with minimum and maximal value. The figures showed the average daily food intake per week in each group.](image)

### Table 1

| Groups | NC       | HF       | LD       | MD       | HD       |
|--------|----------|----------|----------|----------|----------|
| Week 0 | 21.88±0.46 g | 31.33±1.53 g | 30.29±1.48 g | 31.97±2.88 g | 29.90±2.23 g |
| Week 4 | 22.38±0.68 g | 29.64±1.38 g | 28.02±1.47 g | 27.00±1.13 g | 27.16±1.33 g |
| Week 8 | 22.39±0.85 g | 31.07±2.40 g | 28.42±1.18 g | 27.86±1.26 g | 28.56±1.24 g |
| Week 12| 23.52±0.81 g | 31.57±2.08 g | 31.95±1.97 g | 28.67±0.83 g | 29.04±1.67 g |

Data are means ± s.d
incidences of 67% and 50%, respectively, followed by a return to normal levels.

Effects on CRP, and the adipokines chemerin and omentin in mice
As shown in Table 3, compared with the NC group, there were significant differences in the levels of CRP, chemerin and omentin in the four groups (chemerin and omentin $P < 0.001$, CRP $P < 0.05$). CRP was significantly lower in the MD and HD groups than that in the HF group, and the differences were statistically significant ($P < 0.05$). There was no difference in the improvement between the MD and HD groups ($P > 0.05$). The serum chemerin levels of mice in all three intervention groups were lower than those in the HF group, and the difference was statistically significant ($P < 0.01$). The serum omentin levels of mice in all three intervention groups were higher than those in the HF group, and the difference was statistically significant ($P < 0.001$). There was no significant difference in the improvement of serum chemerin and omentin levels between the 3 groups of mice receiving different doses of PEX-168 ($P > 0.05$).

Insulin resistance in mice was negatively correlated with serum omentin levels ($r = -0.836$, $P < 0.001$) and positively correlated with serum chemerin levels ($r = 0.828$, $P < 0.001$).

Discussion

Body weight and food intake
In the present study, low doses of PEX-168 inhibited the progression of obesity in the short term, and medium to high concentrations of PEX-168 inhibited the development of obesity throughout the intervention. The corresponding changes in food intake and body weight also demonstrated that the weight-reducing effect of PEX-168, like other GLP-1 analogues, was mainly through inhibition of intake, which is also consistent with animal experiments with liraglutide. However, the decrease in food intake was unstable during the intervention and was gradually normalized by the mice as the intervention progressed and the inhibition of food intake diminished. This may be because there are other factors that inhibit food intake besides just the feeling of fullness, such as gastrointestinal discomfort. The most common adverse reaction of GLP-1 RA is gastrointestinal reactions, which gradually decrease with a longer dosing time and are significantly dose-dependent[22]. GLP-1 also has a profound inhibitory effect on gastric emptying. The delayed gastric emptying induced by GLP-1 is influenced by a rapid response at the level of vagal activation, and the delay in gastric emptying by GLP-1 is significantly attenuated after long-term administration of GLP-1 analogues[23], which was also consistent with the changes in intake of the intervention groups in this experiment. In addition, PEX-168 is a long-acting preparation with a more sustained hypoglycaemic effect and less effect on gastric emptying, leading to a gradual weakening of the inhibitory effect on food intake.

Insulin resistance
In this study, the hypoglycaemic effect of PEX-168 on simple obese mice manifested at low doses, and all three doses of PEX-168 inhibited the development of prediabetes and improved insulin resistance in nondiabetic mice.

Table 2  Comparison of weight、FBG、FINS、Homa-IR in each group

| Groups | Quantity | Weight (g) | FBG (mmol/L) | FINS (mIU/L) | Homa-IR |
|--------|----------|------------|--------------|--------------|---------|
| NC     | n=5      | 23.52±0.90a | 7.40±0.12a   | 7.70±0.05a   | 2.59±0.05a |
| HF     | n=6      | 31.57±2.28  | 8.33±0.15    | 10.01±0.10   | 3.79±0.08 |
| LD     | n=5      | 31.95±2.20b | 8.02±0.15ab  | 8.50±0.28ab  | 3.1±0.14ab |
| MD     | n=6      | 28.67±0.91ab| 8.08±0.17ab  | 8.62±0.24ab  | 3.17±0.13ab |
| HD     | n=5      | 29.04±1.87ab| 7.94±0.21ab  | 8.65±0.29ab  | 3.12±0.07ab |

*a P < 0.05 vs. the HF group; *b P < 0.05 vs. the NC group. Data are means ± s.d.

Table 3  Comparison of CRP、Chemerin and Omentin levels in each group

| Groups | CRP (ug/L) | Chemerin (ng/L) | Omentin (ng/L) |
|--------|------------|-----------------|----------------|
| NC     | 1327.39±27.05a | 50.85±1.19a     | 147.76±1.62a   |
| HF     | 1410.48±25.45b | 56.79±0.85b     | 131.14±3.14b   |
| LD     | 1403.08±35.26b| 54.65±1.47ab    | 139.67±3.88ab  |
| MD     | 1370.17±31.32ab| 54.30±1.33ab    | 139.10±3.22ab  |
| HD     | 1370.44±33.90ab| 54.66±1.02ab    | 137.58±2.83ab  |

*a P < 0.05 vs. the HF group; *b P < 0.05 vs. the NC group. Data are means ± s.d.
simple obese mice. In Khound's animal study[24], it was shown that elevated GLP-1 prevented the overproduction of VLDL and improved insulin resistance induced by a high-fat diet in mice, which is consistent with the results of the present study.

However, clinical trials have confirmed the safety of PEX-168 in regulating blood glucose in T2DM patients, and the risk of hypoglycaemia is very low with monotherapy. In this study, PEX-168 was used in nondiabetic simple obese mice, and hypoglycaemic events occurred in the middle- and high-dose groups, indicating that there is a certain risk of hypoglycaemia when PEX-168 intervenes in simple obese mice with normal blood glucose, and the low dose should be considered the starting dose.

**Inflammatory reaction**

C-reactive protein, synthesized by the liver, is a sensitive marker of systemic inflammation. It is a nonspecific acute phase reactant that has traditionally been used to detect acute injury, infection and inflammation[25]. Recent studies have shown that diabetes, obesity and elevated levels of CRP, TNF-alpha and leptin are closely associated[26].

In the present study, the level of CRP was significantly elevated in obese mice compared to NC mice, which is consistent with previous reports. The association between obesity and elevated serum CRP levels has been well explained by pathophysiological mechanisms. In this experiment, only the MD and HD groups showed significant improvement in inflammation, and both body weight and the level of CRP in the LD group were not significantly different from those in the HF group, indicating that PEX-168 at medium doses and above significantly reduces body weight in simple obese mice, thereby improving the inflammatory reaction and reducing cardiovascular risk.

**Adipose factor**

In this study, the level of serum chemerin was significantly higher and the level of serum omentin was significantly lower in obese mice than in the NC group. The concentration of chemerin was associated with BMI, adipocyte volume and number, and in adult obese patients, body weight was significantly and positively correlated with circulating chemerin levels[27]. Batista assessed the concentration of omentin in obese patients[28], and normal weight subjects showed higher levels of omentin than overweight and obese patients, which is also consistent with the findings of the study.

Interestingly, although the LD group did not lose weight at the end of the intervention, the levels of FBG, INS, Homa-IR, chemerin and omentin were improved, and the improvements were equal across doses. In Yang's study[29], it was suggested that the GLP-1 analogue liraglutide could improve insulin resistance in high-fat diet-induced obese mice by improving endoplasmic reticulum stress, thereby reducing chemerin levels. K. Tan and Yan pointed out that[16, 30]insulin and glucose could significantly and dose-dependently reduce omentin-1 mRNA and protein levels, and the plasma level of omentin-1 was independently and negatively correlated with fasting insulin and HOMA-IR. K. Tan also pointed out that BMI or WHR is unlikely to be responsible for the decrease in omentin-1 mRNA expression and protein levels in female PCOS patients. In this study, insulin resistance was highly and significantly correlated with both chemerin and omentin, so it can be inferred that the regulation of chemerin and omentin expression by PEX-168 may be related to the hypoglycaemic effect of PEX-168, and the weight-reducing effect of PEX-168 is unlikely to be the reason for the changes in both. This conclusion needs to be demonstrated by further experimental studies.

**Study strengths and limitations**

In this study, the weight reduction effect of PEX-168 was first studied and the effects of it on adipokines chemerin and omentin were further explored. PEX-168 has good potential in treating obesity and preventing the development of diabetes. PEX-168 significantly improves the adipokines through its hypoglycaemic effects. These results provide further mechanistic insight into the action of PEX-168 in the treatment of obesity and diabetes.

However, it has been suggested that[11]non-type 2 diabetic patients lose more weight than type 2 diabetic patients when treated with GLP-1 agonists for weight loss, and this could not be confirmed in this experimental design without the inclusion of a diabetic obesity model group.

Due to our initial experimental design, the sample comprised male mice. The results in this animal study pertain only to the male sex. Since it is a novel hypoglycaemic and anti-obesity drug, we did not set a drug control group in this experiment, but to explore the appropriate concentration to achieve positive effects. However, the addition of a drug control group (like liraglutide) would have made the results more visual and clinically meaningful. We call for the inclusion of the drug control group in the further experiment.

The intervention period was not long enough. In clinical trials, it was noted that PEX-168 blood levels reached stability after four weeks of intervention, but in this study the measurements of the mice only started to
stabilize at a later stage, so the intervention period should be extended.

Conclusions
In summary, the data suggest that in nondiabetic simple obese mice, the antidiabetic drug PEX-168 can effectively reduce body weight, improve insulin resistance, reduce the inflammatory reaction, reduce chemerin and increase omentin levels, and prevent the development of prediabetes. PEX-168 regulates the expression of chemerin and omentin probably through its hypoglycaemic effects, and the weight-reducing effect of PEX-168 is unlikely to be the reason for the changes in both. This study may contribute to the guidance of clinically relevant drugs, broaden the treatment of obesity and contribute to the further exploration of the exact mechanism by which PEX-168 regulates adipokines chemerin and omentin. We will improve the experimental design and conduct further prospective clinical studies in the future.

Abbreviations
PEX-168: Polyethylene glycol loxenatide; GLP-1: Glucagon-like peptide-1; GLP-1 RA: Glucagon-like peptide-1 receptor agonist; WHR: Waist-to-hip ratio; Homa-IR: Homeostasis model assessment of insulin resistance; FBG: Fasting blood glucose; INS: Insulin; CRP: C-reactive protein; BMI: Body mass index; PCOS: Polycystic ovarian syndrome

Acknowledgements
The authors thank Yangzhou University and Northern Jiangsu People’s Hospital for their support of this study.

Authors’ contributions
XS has made substantial contributions to conception and design, ZG and YW have carried out all trials and conducted statistical analysis. ZG has drafted the manuscript and YW has helped with the drafting. LZ, YW and YW have carried out all trials and conducted statistical analysis. ZG has helped with the drafting. All authors read and approved the final manuscript.

Funding
Not applicable.

Availability of data and materials
Data and material would be supplied based on reasonable request. If someone wants to request the data, please email 13,665,278,170@163.com.

Declarations
Ethics approval and consent to participate
The study was carried out in compliance with the ARRIVE guidelines and the Basel Declaration. The study was approved by The Ethic Committee of Northern Jiangsu People’s Hospital.

Consent for publication
All authors have seen the manuscript and agreed to submit it to the journal.

Competing interests
The authors declare that they have no competing interests.

Author details
1College of Nursing, Yangzhou University, Yangzhou, China. 2Northern Jiangsu People’s Hospital, Yangzhou, China. 3General Surgery Institute of Yangzhou, Yangzhou University, Yangzhou, China. 4Clinical Medical College of Yangzhou University, Yangzhou, China.

References
1. Andolfi C, Fischella PM. Epidemiology of Obesity and Associated Comorbidities. Journal of laparoendoscopic & advanced surgical techniques Part A. 2018;28(8):919–24.
2. Wang S, Peng D. Regulation of adipocyte autophagy—the potential anti-obesity mechanism of high density lipoprotein and Apolipoprotein-A-I. Lipids in health and disease. 2012;11:131
3. Liang Y, Qi Y. Developmental trajectories of adolescent overweight/obesity in China: socio-economic status correlates and health consequences. Public health. 2020;185:246–53.
4. Rock CL, Flatt SW, Paliz B, Quintana EL, Heath DD, Rana BK, et al. Effects of diet composition on weight loss, metabolic factors and biomarkers in a 1-year weight loss intervention in obese women examined by baseline insulin resistance status. Metabolism: clinical and experimental. 2016;65(11):1605–13.
5. Kim DD, Basu A. Estimating the Medical Care Costs of Obesity in the United States: Systematic Review, Meta-Analysis, and Empirical Analysis. Value in health: the journal of the International Society for Pharmacoeconomics and Outcomes Research. 2016;19(5):602–13.
6. Tewksbury C, Crowley N, Parrott JM, Andromalos L, Isom KA, Smith E, et al. Weight Loss Prior to Bariatric Surgery and 30-Day Mortality, Readmission, Recuperation, and Intervention: an MBSAQIP Analysis of 349,016 Cases. Obesity surgery. 2019;29(11):3622–8.
7. Neeland JI, Turer AT, Ayers CR, Powell-Wiley TM, Vega GL, Farzaneh-Far R, et al. Dysfunctional adiposity and the risk of prediabetes and type 2 diabetes in obese adults. Jama. 2012;308(11):1150–9.
8. Klinger S, Poussin C, Debril MB, Dolci W, Haillan PA, Thorens B. Increasing GLP-1-induced beta-cell proliferation by silencing the negative regulators of signaling CAMP response element modulator-alpha and DUSP14. Diabetes. 2008;57(3):584–93.
9. van Can J, Sloth B, Jensen CB, Flint A, Blaak EE, Saris WH. Effects of the once-daily GLP-1 analog lixioglitazone on gastric emptying, glycemic parameters, appetite and energy metabolism in obese, non-diabetic adults. International journal of obesity (2005). 2014;38(6):784-93.
10. Tronieri JS, Waddell TA, Walsh OA, Berkowitz RI, Alamuddin N, Gruber K, et al. Effects of lixioglitazone plus phentermine in adults with obesity following 1 year of treatment by lixioglitazone alone: A randomized placebo-controlled pilot trial. Metabolism: clinical and experimental. 2019;66:83–91.
11. Vilsboll T, Christensen M, Junker AE, Knop FK, Gluud LL. Effects of glucagon-like peptide-1 receptor agonists on weight loss: systematic review and meta-analyses of randomised controlled trials. BMJ (Clinical research ed). 2012;344:d7771.
12. Potts JE, Gray LJ, Brady EM, Khunti K, Davies MJ, Bodicoat DH. The Effect of Glucagon-Like Peptide 1 Receptor Agonists on Weight Loss in Type 2 Diabetes: A Systematic Review and Mixed Treatment Comparison Meta-Analysis. PloS one. 2015;10(6):e0126769.
13. Shuai Y, Yang G, Zhang Q, Li W, Luo Y, Ma J, et al. Efficacy and safety of polyethylene glycol loxenatide monotherapy in type 2 diabetes patients: A multicentre, randomized, double-blind, placebo-controlled phase 3a clinical trial. Diabetes, obesity & metabolism. 2021;23(1):116–24.
14. Gao F, Xu X, Mo Z, Ma J, Zhang Q, Yang G, et al. Efficacy and safety of polyethylene glycol loxenatide as add-on to metformin in patients with type 2 diabetes: A multicentre, randomized, double-blind, placebo-controlled, phase 3b trial. Diabetes, obesity & metabolism. 2020;22(12):2375–83.
15. Ernst MC, Sinal CJ. Chemerin: at the crossroads of inflammation and obesity. Trends in endocrinology and metabolism. TEM. 2012;21(11):660–7.
16. Yan P, Li L, Yang M, Liu D, Liu H, Boden G, et al. Effects of the long-acting human glucagon-like peptide-1 analog lixioglitazone on plasma omentin-1 levels in patients with type 2 diabetes mellitus. Diabetes research and clinical practice. 2011;92(3):368–74.
17. Miao Z, Alvarez M, Ko A, Bhagat Y, Rahmani E, Jew B, et al. The causal effect of obesity on prediabetes and insulin resistance reveals the important role of adipose tissue in insulin resistance. PLoS genetics. 2020;16(9):e1000918.
18. Gao C, Rao M, Huang W, Wan Q, Yan P, Long Y, et al. Resistant starch ameliorated insulin resistant in patients of type 2 diabetes with obesity: a systematic review and meta-analysis. Lipids in health and disease. 2019;18(1):205.
19. Shabana, Shahid SU, Sarwar S. The abnormal lipid profile in obesity and coronary heart disease (CHD) in Pakistani subjects. Lipids in health and disease. 2020;19(1):73.

20. Szmitko PE, Verma S. C-reactive protein and the metabolic syndrome: useful addition to the cardiovascular risk profile? Journal of the cardiometabolic syndrome. 2006;1(1):66–9; quiz 70–1.

21. Sung KC, Lee MY, Kim YH, Huh JH, Kim JY, Wild SH, et al. Obesity and incidence of diabetes: Effect of absence of metabolic syndrome, insulin resistance, inflammation and fatty liver. Atherosclerosis. 2018;275:50–7.

22. Brown E, Cuthbertson DJ, Wilding JP. Newer GLP-1 receptor agonists and obesity-diabetes. Peptides. 2018;100:61–7.

23. Nauck MA, Kemmeries G, Holst JJ, Meier JJ. Rapid tachyphylaxis of the glucagon-like peptide 1-induced deceleration of gastric emptying in humans. Diabetes. 2011;60(5):1561–5.

24. Khound R, Tahir J, Baker C, Adeli K, Su Q. GLP-1 Elicits an Intrinsic Gut-Liver Metabolic Signal to Ameliorate Diet-Induced VLDL Overproduction and Insulin Resistance. Arteriosclerosis, thrombosis, and vascular biology. 2017;37(12):2252–9.

25. Backes JM, Howard PA, Moriarty PM. Role of C-reactive protein in cardiovascular disease. The Annals of pharmacotherapy. 2004;38(1):110–8.

26. Mirza S, Hossain M, Mathews C, Martinez P, Pino P, Gay JL, et al. Type 2-diabetes is associated with elevated levels of TNF-alpha, IL-6 and adiponectin and low levels of leptin in a population of Mexican Americans: a cross-sectional study. Cytokine. 2012;57(1):136–42.

27. Rowicka G, Dyląg H, Chełchowska M, Weker H, Ambroszkiewicz J. Serum Calprotectin and Chemerin Concentrations as Markers of Low-Grade Inflammation in Prepubertal Children with Obesity. International journal of environmental research and public health. 2020;17(20).

28. de Souza Batista CM, Yang RZ, Lee MJ, Glynn NM, Yu DZ, Pray J, et al. Omentin plasma levels and gene expression are decreased in obesity. Diabetes. 2007;56(6):1655–61.

29. Yang J, Ao N, Du J, Wang X, He Y. Protective effect of liraglutide against ER stress in the liver of high-fat diet-induced insulin-resistant rats. Endocrine. 2015;49(1):106–18.

30. Tan BK, Adya R, Farhatullah S, Lewandowski KC, O’Hare P, Lehnert H, et al. Omentin-1, a novel adipokine, is decreased in overweight insulin-resistant women with polycystic ovary syndrome: ex vivo and in vivo regulation of omentin-1 by insulin and glucose. Diabetes. 2008;57(4):801–8.

**Publisher’s Note**
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.