Determination of Ghrelin’s Role in the Pathogenesis of Pregnancy Induced Hypertension

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Abstract

Introduction

Pregnancy induced hypertension (PIH) is a disease of an unclear etiology that complicates 2-3 % of pregnancies. PIH is a critical cause of perinatal mortality of pregnant women and neonates and a major cause of intraterine growth retardation and of iatrogenic prematurity [1]. PIH is a multiorgan complication; its main causes result from the interaction of the mother’s immunological system with trophoblast antigens. The ethiopathogenesis of pregnancy induced hypertension remains to be elucidated.

A crucial risk factor of pregnancy induced hypertension is obesity. Relationships between hypertension and obesity are well documented [2], ultimately involving excessive retention of sodium by the kidneys, insulin resistance, and chronic stimulation of the sympathetic system, leading to vessel remodelling. Leptin levels are increased in obese patients, stimulating the proliferation of smooth muscles cells of arterial vessels and influencing Renin–Angiotensin–Aldosterone System activity. Pregnant women experience excessive appetite, which leads to overall body mass increase [3].

Ghrelin, a hormone mainly produced in the digestive tract, is involved in releasing growth hormone, increasing appetite, and inducing obesity. Ghrelin stimulates the secretion of prolactin and adrenocorticotrophic hormone and has a crucial impact on steroidogenesis and carbohydrate metabolism [4]. Disorders surrounding adrenocorticotrophic hormone secretion are observed in obese individuals. Results regarding changes in ghrelin secretion in various types of hypertension, however, are equivocal. Ghrelin is known to play a crucial role in the intraterine growth of the fetus; blood ghrelin concentration in intraterine growth retardation fetuses has been reported to be higher than in healthy ones [5].

Based on the possible relationship between hypertension and obesity in pregnant women together with the change in ghrelin secretion in obese individuals, we sought to discern whether ghrelin governs related secretion disorders and plays a role in the ethiopathogenesis of pregnancy induced hypertension.

We sought to answer the following questions:

It is evident that our research has both scientific and practical aspects. Showing a relationship between ghrelin secretion and hypertension and/or obesity would prove that ghrelin analysis is a useful tool in evaluating the fetal condition and/or pregnancy prognosis.

Materials and Methods

Our study included 61 pregnant women. The first group consisted of 20 healthy pregnant (HP), the second of 20 hypertensive pregnant women (PIH), and the third of 21 pregnant women with hypertension and obesity (OPIH). The mean ages of the groups were: 28.1 ± 1.1, 25.7 ± 1.2 and 27.5 ± 1.1 years respectively. Differences between the mean plasma level as compared with NP (812 ± 83, 716 ± 66, and 717 ± 52 vs 1057 ± 74, respectively). We found no significant ghrelin reduction after eating in obese hypertensive pregnant women. The blood ghrelin concentration in pregnant women negatively correlated with mean arterial pressure. In spite of the differences in ghrelin concentration in women, whom a decrease in ghrelin concentration after a standard meal was observed compared to women, who didn’t reduce of ghrelin secretion no differences were found in mean arterial pressure.

Conclusion: The relationship between arterial hypertension and ghrelin secretion disorders during hypertensive pregnancy is uncertain and requires further examination.

Keywords: Ghrelin; Pregnancy; Hypertension; Obesity

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ages were not significant. All women gave birth to healthy children with Apgar scores between 7 and 10 points. The control group, consisting of 19 healthy non-pregnant women at the age of 20. 2 ± 0.1 (NP), was also examined. We received a statement on informed consent from the study participants. The study protocol was approved by the Bioethical Commission of the Medical University of Silesia. Pregnant women (unifetal pregnancies) with both hypertension and other systemic diseases (including diabetes) were not included to the study. Blood from each woman was drawn from the basilic vein after a fasting period 2-3 days before delivery. Haemoglobin, glucose, protein, creatinine, and uric acid concentrations were determined. Plasma ghrelin concentration was assessed before and one hour after the intake of a standard breakfast. All patients had had their arterial blood pressure measured before blood was drawn. 

The mean arterial blood pressure (MAP) was determined according to the formula:

\[
\text{MAP} = \frac{\text{systolic pressure} + (\text{diastolic pressure} - \text{diastolic pressure})}{3}
\]

Body Mass Index (BMI) was calculated according to the formula:

\[
\text{BMI} = \frac{\text{body mass}}{\text{height}^2} \text{ (kg/m}^2\text{)}
\]

A radio immunological method was used to determine ghrelin concentration using the Total Ghrelin Ria Kit. The standard breakfast included: 40 g of cornflakes, 250 ml of milk, 100 ml of grapefruit juice, and 5 g of sugar, totaling an energetic value of 369.6 kcal (1545 kJ).

Results are presented as mean ± SEM. The distribution normality was assessed using the Kolmogorov–Smirnov test. A student's t-test was employed to determine the mean for dependent and independent variables. The U-Mann Whitney test was used to analyze ghrelin concentration data. Occurrence correlations were evaluated using the Pearson's linear correlation and Kendall test (for Ghrelin). Differences were considered statistically significant at p<0.05.

Results

Hypertensive groups of pregnant women expressed significantly increase MAP compared both, to values at the beginning of pregnancy – PIH: 114 ± 2 vs. 96 ± 1 mmHg and OPIH: 119 ± 3 vs. 99.8 ± 1 mmHg - p<0.01 and to healthy pregnant ones HP: 92 ± 1 vs. 90 ± 1 mmHg - p<0.01, respectively. Clinical and laboratory characteristics of studied populations are presented in Table 1. Our results revealed significantly lower plasma ghrelin concentrations in all pregnant women as compared to non pregnant ones. Ghrelin concentration significantly decreased after a standard meal in normal pregnant women; however, this trend did not occur in OPIH group members (Table 2).

Based on the Kendall correlation coefficient, a negative correlation was found between the random variables BMI and ghrelin in the population of healthy pregnant women (HP) for the assumed level of significance \(\alpha\), p<0.05.

As a result of the statistical analysis for the Pearson’s linear correlation testa positive correlations between BMI and MAP and BMI and glucose were found in hypertensive pregnant women (PIH+OPIH) p=0.036 and p<0.001 respectively.

Based on Pearson's linear correlation test, a positive correlations were found between both the random variables BMI and MAP as well as BMI and glucose in the whole population of pregnant women (HP+PIH+OPIH) for assumed levels of significance \(\alpha\), p<0.001. Additionally, as a result of the statistical analysis for the Kendall correlation coefficient, a negative correlation between the random variables MAP and ghrelin was found in this population for the assumed relevance level \(\alpha\), p=0.043. The calculated correlation coefficient \(\tau\) is -0.26, which means that the dependence level between the analyzed statistical characteristics in the probe taken from the subject general population was low.

In pregnant women, whom a decrease in ghrelin concentration after a standard meal was observed (n=43) – 846.4 ± 44. 8 ng/ml compared to pregnant women, who didn't reduce of ghrelin secretion after standard breakfast (n=18) – 513. 5 ± 40. 5 ng/ml, no differences were found in mean arterial pressure 109 ± 2 vs 109 ± 1 mmHg.

Results Summary

Our results shows a standard clinical and biochemical picture of pregnancy characterized by an increase in body mass and BMI, a decrease in hemoglobin level, and a decrease in creatinine and protein concentrations in all pregnant subjects as compared to non pregnant ones. Hypertensive pregnant women (PIH and OPIH) exhibited higher blood pressure during pregnancy. Our results revealed significantly lower plasma ghrelin concentrations in all pregnant women in comparison with non pregnant ones.

PIH differed in women with uncomplicated pregnancies with no negative correlation between ghrelin concentration and BMI. Additionally, no difference in glucose, creatinine, and protein concentration was noted. OPIH differed in women with uncomplicated pregnancies by the significance in glycaemia and uricaemia. No significant ghrelin decrease after a standard meal was observed. The OPIH women differed from the PIH women by higher BMI and glycaemia, but not by blood pressure.

A negative correlation between mean arterial pressure and ghrelin concentration was reported in all pregnant women together. In spite of the aforementioned differences in ghrelin concentration in women, whom a decrease in ghrelin concentration after a standard meal was observed compared to women, who didn't reduce of ghrelin secretion no differences were found in mean arterial pressure.

Discussion

Hypertension is a frequent complication of pregnancy, and the etiopathogenesis is multifactorial. Generalized blood vessel contraction, which induces vascular resistance, is the basic pathophysiological phenomenon. These contractions can have destructive impact on vessels. Hypertension in pregnancy is associated with blood vessel sensitivity increases to substances of pressure activity [3].

Many pathogenetic foci are listed, including genetically determined parameters that also may contribute to blood pressure increases during pregnancy. Important contributors to developing arterial hypertension during pregnancy are environmental factors [6]. The impacts of body mass increases during pregnancy and insufficient nutrients for the mother and fetus, including arterial hypertension, contribute to etiopathogenesis. It is important than women control body mass gain because an excess can result in hypertension [7]. The impact of obesity in the etiopathogenesis of arterial hypertension is undeniable [8]. Both obesity and arterial hypertension in pregnant women are known to lead to uteroplacental blood flow impairment [9], resulting in changes in placenta morphology, ultimately decreasing the active surface of the organ.

Ghrelin influences the energy regulation of the body. Ghrelin is...
The reason for these lower ghrelin levels remains to be determined. The lack of significant differences in ghrelin concentrations in hypertensive groups does not exclude it as influencing the etiopathogenesis of hypertension. The central and peripheral activities of ghrelin include stimulating both hunger feelings and motoric activity of the alimentary tract, and initiating eating. An increase in blood ghrelin concentration precedes a meal. Limiting food intake and following a low protein diet leads to greater ghrelin secretion. Conversely, a high-fat and carbohydrate-rich diet hampers its secretion. Kawamura et al. [14] showed that ghrelin levels significantly rise during starvation. Additionally, ghrelin levels in slim individuals increases progressively before each meal and decreases just one hour after it [15]. Firczyk characterized hemodialysed patients by higher ghrelin concentrations than the control group [16]. We report a significant decrease in ghrelin concentration after a standard meal in pregnant women. This normal trend, however, fails to appear in pregnant women with arterial hypertension and obesity.

During pregnancy, a negative correlation between BMI and ghrelin concentration has been observed. Again, we failed to find this correlation in pregnant women with hypertension, both in the PIH and OPIH groups. We conclude that arterial hypertension in pregnant women complicates these relationships.

In patients with polycystic ovary syndrome, the ghrelin levels correlate negatively with BMI, a trend opposing that in healthy women [17,18].
Jolda-Mysłowska [19] determined ghrelin concentrations in patients with idiopathic arterial hypertension, finding that concentrations in blood serum in these patients were lower than control. Makino et al. [5] obtained different results, reporting that patients with pregnancy induced hypertension exhibited higher ghrelin levels. Additionally, they noted a positive correlation between ghrelin concentration and arterial pressure in these patients, and a positive relationship between ghrelin and blood pressure in healthy pregnant women, which was not observed in non pregnant ones.

The relationship between ghrelin, insulinemia, and insulin resistance in obese hypertension patients was examined by Dytfeld et al. [20]. Decreased concentrations of ghrelin in blood serum were observed in obese individuals with hypertension. Body mass reduction was associated with an increase in ghrelin concentration [17].

Prerequisites exist regarding the extent of ghrelin secretion disorders in the etiopathogenesis of arterial hypertension. Although BMI and MAP as well as BMI and glucose concentrations were correlated, ghrelin concentration was only found to correlate with mean arterial pressure. Thus, it cannot be unequivocally presumed whether ghrelin deficiency is responsible for arterial hypertension or vice versa (that the occurrence of hypertension modifies ghrelin secretion). Our reported negative relationship between mean arterial pressure and ghrelin concentration in all pregnant subjects suggests an etiopathogenic link between pregnancy induced hypertension and ghrelin secretion.

Makino et al. [5] claimed that ghrelin levels are lower in obese patients, and a negative correlation between ghrelin concentration and BMI exists. A ghrelin level decline in the third trimester of pregnancy was described by Fuglsang et al. [21], which supported Makino’s result, showing that women reach their greatest body mass at the end of the pregnancy period.

Considering the impact of ghrelin on glucose metabolism, its influence on insulin resistance stimulation, obesity, and GH secretion, which by its anti-insulin activity increases glucose concentration in blood serum, should be highlighted. This peripheral activity of ghrelin in glucose metabolism has been confirmed in the literature [22].

Standard meal consumption allowed for monitoring hormonal mechanisms, including ghrelin secretion disorders, which can potentially influence arterial pressure. During a standard pregnancy, a standard meal causes ghrelin concentration to significantly decrease; the lack of such a decrease in hypertensive pregnant women with obesity can foster etiopathogenetic prerequisites for hormone secretion. Whether this hormone configuration is typical for the obese in the hypertensive group of pregnant women requires further investigation. Notably, the lack of the aforementioned correlation in the hypertensive groups (PIH, and OPIH), stands against a connection between arterial hypertension and ghrelin secretion disorders.

No mean arterial pressure difference between groups exhibiting decreased ghrelin concentration was seen after a standard meal. This data also argue against a close dependence of ghrelin secretion changes and arterial hypertension.

It can thus be stated that the relationship between arterial pressure values and ghrelin secretion disorders in hypertensive pregnant women is uncertain and requires further investigation.

Hypertension during pregnancy remains problematic. Due to complications of hypertension, relevant investigations revolve around various scientific centers. Suggested procedure algorithms prove that a current, constant search for optimal diagnostics and therapy are underway. The discovery of novel hormone activity will enable recognition of a pregnancy-induced hypertension mechanism.

Conclusions

1. Ghrelin concentration in the blood serum of pregnant women is lower than in non-pregnant women.

2. Women with pregnancy induced hypertension and obesity differ from healthy pregnant women in food intake-induced change of ghrelin secretion.

3. The relationship between arterial hypertension and ghrelin secretion disorders during hypertensive pregnancy is uncertain and requires further examination.

Disclosure Statement

The authors have nothing to disclose.

References


