Human relaxins (RLNH1, RLNH2), their receptor (RXFP1) and fetoplacental growth

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Abstract

Relaxin, a systemic and placental hormone, has potential roles in fetoplacental growth. Human placenta expresses two RLN genes, RLNH1 and RLNH2. Maternal obesity is common and is associated with abnormal fetal growth. Our aims were to relate systemic and cord blood RLNH2, placental RLNs and their receptor (RXFP1) with fetoplacental growth in context of maternal body mass index, and associations with insulin-like growth factor 2 (IGF2) and vascular endothelial growth factor A (VEGFA) in the same placentas. Systemic, cord blood and placental samples were collected prior to term labor, divided by prepregnancy body mass index: underweight/normal (N=25) and overweight/obese (N=44). Blood RLNH2 was measured by ELISA; placental RLNH2, RLNH1, RXFP1, IGF2 and VEGFA were measured by quantitative immunohistochemistry and mRNAs were measured by quantitative reverse transcription PCR. Birthweight increased with systemic RLNH2 only in underweight/normal women (P=0.036). Syncytiotrophoblast RLNH2 was increased in overweight/obese patients (P=0.017) and was associated with placental weight in all subjects (P=0.038). RLNH1 had no associations with birthweight or placental weight, but was associated with increased trophoblast and endothelial IGF2 and VEGFA, due to female fetal sex. Thus, while systemic RLNH2 may be involved in birthweight regulation in underweight/normal women, placental RLNH2 in all subjects may be involved in placental weight. A strong association of trophoblast IGF2 with birthweight and placental weight in overweight/obese women suggests its importance. However, an association of only RLNH1 with placental IGF2 and VEGFA was dependent upon female fetal sex. These results suggest that both systemic and placental RLNs may be associated with fetoplacental growth.

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Introduction

Fetal growth is an important predictor of perinatal morbidity and mortality as well as adult-onset cardiometabolic disease (Barker 2006, Unterscheider et al. 2014, Smith et al. 2016). Despite its importance, the physiology underlying normal fetal growth remains incompletely understood. Maternal obesity is associated with both excessive and restricted fetal growth (Higgins et al. 2011, Magann et al. 2013). Thus, the need to understand how fetal growth is controlled is escalating as rates of maternal obesity continue to increase in the United States (Fisher et al. 2013). Increased maternal body mass index (BMI) in the early second trimester is also associated with increased placental weight (Roland et al. 2014), and placental weight itself is independently associated with adult-onset disease (Barker et al. 2010).

Relaxin (RLN) is known for its role in remodeling of reproductive tissues (Hisaw 1926) and as a growth factor (McMurtry et al. 1980, Millar et al. 2003). Three RLN genes, RLN1, RLN2 and RLN3 are present in humans and great apes, whereas RLN in other monkeys and mammals have only a single gene analogous to human RLN2 (Hayes 2004). RLN1 and RLN2 are expressed in human decidua, placenta and prostate (Hansell et al. 1991), and RLN3 is primarily expressed in the nervous system of rodents and macaques (Ma et al. 2009). Maternal systemic RLN is relaxin H2 (RLNH2) and secreted from the corpus luteum during gestation, as shown by concurrent ovarian vein and peripheral blood sampling in pregnant women (Weiss et al. 1976). The placenta does not contribute to this circulating RLN, as women with ovum-donated pregnancies, where there is no functional corpus luteum, have undetectable systemic RLN (Johnson et al. 1991). The RLN receptor was first identified as a leucine-rich repeat G-protein-coupled...
receptor (LGR7) (Hsu et al. 2002) currently known as the RLN family peptide receptor 1 (RXFP1) and is expressed in placenta (Lowndes et al. 2006). Although the function of RLNH1 remains unknown (Bathgate et al. 2013), it shares the same receptor as RLNH2, but binds with a fivefold lower affinity than RLNH2, suggesting preferential RLNH2 binding (Garibay-Tupas et al. 1995).

RLN and RXFP1 are expressed by the endothelium and vascular smooth muscle of the systemic vasculature in rodents (Jelinic et al. 2014) and RXFP1 in umbilical endothelium and vascular smooth muscle in humans (Sarwar et al. 2015). Evidence for roles in both vasodilation and angiogenesis have been obtained primarily from animal models, but potential contributions of systemic RLN to human maternal gestational vascular adaptations are likely (Conrad 2011). RLNH1 and RLNH2 as local hormones can potentially target fetal-placental vessels to increase placental perfusion, with important implications for its involvement in fetal growth. However, little is known about the roles of RLN within the placenta and its potential effects on placental vasculature or in placental growth.

Fetal growth may be partially regulated by systemic and/or placental RLN either directly or indirectly by induction of other fetalplacental growth factors such as insulin-like growth factor 2 (IGF2) (Millar et al. 2003) and vascular endothelial growth factor A (VEGFA) (Unemori et al. 1999). Growth-restricted fetuses exhibit decreased placental IGF2 expression (McMinn et al. 2006, Demetriou et al. 2014), and IGF2 dysregulation is associated with Beckwith–Wiedemann syndrome, a fetal overgrowth condition (Shuman et al. 1993–2016). RLNH2 increases transcription of IGF2 in human fetal membranes (Millar et al. 2003). VEGFA promotes vascular endothelial cell growth and vasodilation and contributes to placental angiogenesis (Andraweera et al. 2012), and fetal growth restriction is associated with decreased maternal serum VEGF (Savvidou et al. 2006). RLNH2 has been shown to induce the expression of VEGF in endometrial cells and monocytes (Unemori et al. 1999, 2000). However, little is known about the relationships between RLNH2, RLNH1 and either IGF2 or VEGFA in the human placenta.

A better understanding of fetal growth is integral to the goal of optimizing the intrauterine environment and improving the immediate and lifelong health of the newborn. The aims of this study were to relate maternal and fetal systemic RLNH2 and placental RLNH1, RLNH2 and RXFP1 in the same patients at term and to show whether there were any associations with fetoplacental growth and levels of placental IGF2 or VEGFA in the context of maternal BMI.

Materials and methods

Study participants and sample collection

The sample set was used previously (Tsai et al. 2015a, b) and approved by the Western Institutional Review Board with all participants providing written informed consent. Women at greater than or equal to 37 weeks 0 days gestation with a singleton pregnancy were recruited at Kapiolani Medical Center for Women and Children (Honolulu, HI, USA). Potential participants were limited to those undergoing prelabor cesarean delivery to avoid potentially confounding changes related to labor. Patients with premature rupture of membranes, pregestational diabetes, acute infection at the time of delivery, maternal inflammatory/connective tissue/autoimmune disease and steroid use were excluded. In addition, patients with gestational diabetes were excluded due to its association with increased fetal growth (Langer et al. 2005). A placental pathologist examined all tissues, and those with histologic signs of inflammation/infection were discarded. Patients were divided by prepregnancy body mass index (BMI) into underweight/normal (Uw/N) (BMI: 16.0–24.9 kg/m², n=25) and overweight/obese (Ow/Ob) (BMI ≥25.0 kg/m², n=44). Fasting maternal blood was collected prior to delivery and venous cord blood within 15 min of delivery. Plasma was aliquoted and stored at −80°C. Two full-thickness placental samples, excluding basal plate, were collected at the cord insertion site within 15 min of delivery. One sample was fixed in neutral-buffered formaldehyde for 72 h, and the other was immediately stored at −80°C.

Maternal plasma analysis

Maternal plasma interleukin-6 (IL-6) is increased in obesity (Eder et al. 2009) and was measured by High-Sensitivity Quantikine ELISA (R&D Systems), sensitivity 0.016 pg/mL and intra-assay coefficient of variation 7.8%. Maternal and fetal plasma RLN were measured by ELISA (Abipco, Salem, NH, USA), sensitivity 4 pg/mL and intra-assay coefficient of variation 5.2%. This ELISA was not reported to be specific for RLNH2, but prior work has demonstrated that only RLNH2 is systemic (Johnson et al. 1991). Maternal plasma samples were unavailable for all subjects; therefore, for RLNH2, Uw/N (n=24) and Ow/Ob (n=43) and for IL-6, Uw/N (n=22) and Ow/Ob (n=39) were used. All samples were run in duplicate.

Placental immunohistochemistry and quantitation

Immunohistochemistry was performed for quantitative assessment (Taylor & Levenson 2006). Serial sections (5 μm) were heated at 95°C for 30 min in sodium citrate buffer (10 mM, pH 6.0) for antigen retrieval for all antigens except VEGFA, which was heated in 10 mM Tris/1 mM EDTA, pH 9.0, for 20 min. The Vectastain Elite kit (Vector Labs, Burlingame, CA, USA) was used according to the manufacturer’s protocol. Nonspecific binding was blocked with normal goat serum for 30 min prior to incubation for 60 min with either rabbit monoclonal to RLNH2 (Abcam ab 183505, 0.06 pg/μL), mouse monoclonals to RLNH1 from both Abcam, (ab 14329) and R&D Systems (MAB 3257, 3 pg/μL), RXFP1 (Sigma-Aldrich, 0.06 pg/μL), and IGF2 (Novus Biologicals, Littleton, CO, USA, 0.5 pg/μL) or rabbit polyclonal to VEGFA (Santa Cruz Biotechnology, 0.4 pg/μL). Negative controls were species-specific non-immune IgG at equivalent concentrations with 3,3-diaminobenzidine substrate for visualization.
Slides were rinsed, counterstained with hematoxylin, cleared and mounted.

Imaging was performed with an Olympus BX51 microscope (Olympus America) and a CRI Nuance spectral analyzer (Caliper Life Sciences, Hopkinton, MA, USA). A series of brightfield images were acquired between 420 and 700 nm wavelengths at 20-nm intervals. These images were stacked to create a 3-dimensional image cube. The Inform software (version 2.0.2; PerkinElmer) segmented tissues into cell types, unmixed the spectral components and quantitated staining levels (Stack et al. 2014). Average signal intensity per pixel from 5 different fields per patient was obtained as mean optical density units (ODU). Signal intensities were quantitated at 400x magnification. Quantitation of villous vascular endothelium was by manual identification of vessels on each image. Laboratory personnel were blinded to clinical information.

**RNA isolation and quantitative reverse transcription PCR (qRT-PCR)**

Total RNA was extracted from 30 mg placental tissue using the RNeasy Fibrous MiniKit (Qiagen) and quality was assessed by Agilent Bioanalyzer 2100 (Agilent Technologies). Total RNA (400 ng) was reverse transcribed using Multiscribe reverse transcriptase (Applied Biosystems), incubating at 25°C for 10 min, 37°C for 120 min and 95°C for 5 min. For RT-PCR, gene-specific TaqMan Assays on Demand probe sets for RLN1 (Hs00366417_m1), RLN2 (Hs00762019_s1) and two reference genes SDHA (Hs00188166_m1) and YWHAZ (Hs00237047_m1) were selected for placenta as recommended (Meller et al. 2005: 1). Reactions in triplicate were cycle at 95°C for 1 min, 1 cycle at 95°C for 10 min, 40 cycles at 95°C for 15 s and 60°C for 1 min. Results were normalized to the geometric mean of SDHA and YWHAZ expression in each sample and reported as the mean ± s.d.

*Statistical analyses*

Data were analyzed with SAS statistical software, version 9.3 (SAS Institute Inc, Cary, NC) and SPSS statistics for Macintosh, version 23 (IBM). Demographic and clinical data are compared by Student’s t-tests and chi-square or Fisher’s exact test for continuous and categorical variables respectively. Immunostaining intensities between Uw/N and Ow/Ob patients were compared with Student’s t-tests. Relationships between RLNH1 and RLNH2, RXFP1, IGF2, VEGFA, with fetal and placental growth and fetal sex were compared for all samples and stratified by BMI group using linear regression models. P values of ≤0.05 were considered statistically significant.

**Results**

**Demographic and clinical data**

Of 84 patients, 6 were excluded for chorioamnionitis or placental histologic inflammation and 9 for gestational diabetes, leaving 69 patients in this study. Demographic and clinical data are shown in Table 1. Of these, 3 were underweight (Uw), 22 normal weight (N), 21 overweight (Ow) and 23 obese (Ob), which were combined into 2 groups Uw/N (n = 25) and Ow/Ob (n = 44) with mean pregestational BMIs of 21.3 and 32.0 respectively (P < 0.001). The Uw/N group was significantly older than the Ow/Ob group (P = 0.034), but otherwise there were no differences. Maternal plasma IL-6 and placental weight in relation to prepregnancy BMI are shown in Fig. 1A and B respectively. Maternal IL-6 (P < 0.001) (Fig. 1A) and placental weight (P = 0.025) (Fig. 1B) both increased linearly with prepregnancy BMI, validating our sample set as these observations were consistent with previous data (Eder et al. 2009, Roland et al. 2014).

**Maternal and cord plasma RLNH2**

Birthweight increased as maternal plasma RLNH2 increased for Uw/N (P = 0.036) but not for Ow/Ob patients (P = 0.553) (Fig. 1C). There was no association between maternal plasma RLNH2 and placental weight (data not shown). The mean ± s.d. of maternal and cord plasma RLNH2 levels for all patients were 260 ± 158.8 pg/mL and 22 ± 12.5 pg/mL respectively, with cord plasma RLNH2 significantly lower than maternal RLNH2 (P < 0.001). Data for all patients showedcord plasma

**Table 1 Demographic and clinical characteristics of underweight/normal and overweight/obese patients.**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Uw/N (n = 25)</th>
<th>Ow/Ob (n = 44)</th>
<th>Significance (P)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal age (years)</td>
<td>33 ± 5.9</td>
<td>30 ± 5.0</td>
<td>0.034*</td>
</tr>
<tr>
<td>Primiparous</td>
<td>6 (24.0)</td>
<td>7 (15.9)</td>
<td>0.524</td>
</tr>
<tr>
<td>Prepregestational BMI (kg/m²)</td>
<td>21.3 ± 2.2</td>
<td>32.0 ± 6.3</td>
<td>&lt;0.001*</td>
</tr>
<tr>
<td>Male newborn</td>
<td>8 (32.0)</td>
<td>25 (56.8)</td>
<td>0.083</td>
</tr>
<tr>
<td>Gestational age at delivery (weeks)</td>
<td>38.9 ± 0.7</td>
<td>39.0 ± 0.5</td>
<td>0.237</td>
</tr>
<tr>
<td>Gestational weight gain (pounds)</td>
<td>13.2 ± 5.13</td>
<td>14.9 ± 7.5</td>
<td>0.570</td>
</tr>
<tr>
<td>Smoking at delivery</td>
<td>0 (0.0)</td>
<td>3 (6.8)</td>
<td>0.160</td>
</tr>
<tr>
<td>Hypertension</td>
<td>0 (0.0)</td>
<td>2 (4.5)</td>
<td>0.531</td>
</tr>
<tr>
<td>Birthweight (g)</td>
<td>3259 ± 418</td>
<td>3445 ± 483</td>
<td>0.096</td>
</tr>
<tr>
<td>Placental weight (g)</td>
<td>475 ± 92</td>
<td>535 ± 102</td>
<td>0.011*</td>
</tr>
</tbody>
</table>

Results are expressed as mean ± s.d. or frequency (%).

*P < 0.05.

Ow/Ob, overweight/obese; Uw/N, underweight/normal.
levels increased as maternal plasma RLNH2 increased ($P<0.001$, Fig. 1D), but there were no associations of cord plasma RLNH2 with either birthweight or placental weight. From these data, the levels of maternal systemic RLNH2 may be more important for birthweight in Uw/N than those in Ow/Ob women.

Placental RLNH2: protein and gene expression

In the placenta, RLNH2 immunolocalized strongly to the syncytiotrophoblast and the few remaining cytotrophoblasts, with lighter staining in the fetal vascular endothelial cells. Immunostaining was also lighter in Uw/N compared to Ow/Ob samples, examples shown (Fig. 2A and B) with a negative control (Fig. 2C). Quantitation with all subjects showed that placental weight increased as syncytiotrophoblast RLNH2 increased ($P=0.038$) (Fig. 2D), with no increase in placental weight with RLNH2 in the vasculature. However, there were no relationships between syncytiotrophoblast ($P=0.185$) or vascular ($P=0.942$) RLNH2 and birthweight. Subdivision of patients into two groups, showed that syncytiotrophoblast RLNH2 was significantly greater in Ow/Ob patients compared to Uw/N patients ($P=0.017$) (Fig. 2E and F). In contrast, for the two groups of patients, there was no significant difference in RLNH2 in the placental vasculature (not shown). Placental RLNH2 mRNA in Ow/Ob patients (Fig. 2G) showed a similar pattern with placental weight as its protein (Fig. 2F), as well as with birthweight (not shown), neither were significant. However, the RLNH2 mRNA for Uw/N subjects differed from that for Ow/Ob subjects and was significantly but inversely related to both placental weight ($P=0.018$) (Fig. 2G) and birthweight ($P=0.017$) (not shown).

Placental RXFP1

The relaxin receptor (RXFP1) was immunolocalized in the same placentas and examples are shown for Uw/N (Fig. 3A) and Ow/Ob (Fig. 3B) with a negative control (Fig. 3C). The staining intensity was very good in the syncytiotrophoblast and vascular endothelial cells, with additional light and intermittent staining in the vascular smooth muscle as previously shown (Jelinic et al. 2014). Separate quantitation of these was not possible; therefore, the results for the vasculature were a combination of both endothelial and smooth muscle staining. Quantitation showed significantly more RXFP1 in the syncytiotrophoblast of Ow/Ob compared to the Uw/N patients ($P=0.054$) (Fig. 3D) with no difference in the vasculature ($P=0.664$) (Fig. 3E). When plotted for individual subjects, placental weight was not associated with RXFP1 in either the syncytiotrophoblast ($P=0.977$) (Fig. 3F) or vasculature ($P=0.093$) (Fig. 3G).

Summary: maternal and placental RLNH2 and RXFP1, according to prepregnancy maternal BMI

A diagram summarizing these data is shown in Fig. 4. Maternal systemic RLNH2, a product of the corpus luteum, was increased as birthweight increased in Uw/N patients only. There was no association with placental weight. However, for all subjects, syncytiotrophoblast RLNH2 was related to placental weight, as shown. In the Ow/Ob patients, who had heavier placentas, there were also relatively higher levels of syncytiotrophoblast RLNH2 and RXFP1 compared to the Uw/N patients. These data suggest that in Uw/N patients, maternal plasma RLNH2 may be associated with birthweight, whereas placental RLNH2 in all subjects may be associated with placental weight.

Placental RLNH2 and associations with IGF2 and VEGFA

RLNH2 has been shown to increase IGF2 (Millar et al. 2003) and VEGFA (Unemori et al. 1999) in different cell types. Although any direct relationship has not been shown in placental tissue or cells, we sought associations between RLNH2 and these important placental growth factors. Therefore, we immunolocalized both proteins in the placentas of our patients and sought associations with the concomitant expression of RLNH2. IGF2 localized to the remaining cytotrophoblast and vascular
endothelium, examples are shown for Uw/N (Fig. 5A) and Ow/Ob (Fig. 5B), with a negative control (Fig. 5C). Placental weight ($P < 0.001$) and birthweight ($P = 0.002$) (Fig. 5D and E respectively) both significantly increased as cytотrophoblast IGF2 increased in the Ow/Ob patients, but not in the Uw/N patients.

VEGFA immunolocalized to the syncytiotrophoblast and remaining cytотrophoblast, with slightly stronger staining in the latter. It also immunolocalized to the vascular endothelium. These results agree with results from Clark and coworkers (1996) and therefore are not shown here. There were no significant associations between placental RLNH2 and either IGF2 or VEGFA in the trophoblast or placental vasculature.

**Placental RLNH1 and associations with RXFP1, IGF2 and VEGFA**

The negative results showing lack of any association of RLNH2 with the growth factors, IGF2 and VEGFA were unexpected. We therefore considered the homologous hormone RLNH1, also expressed in the placenta (Hansell et al. 1991). The recent availability of antibodies to RLNH1 allowed us explore this in our samples. Initially, we used an antibody from the same source as the RLNH2 antibody used here, but this was unsuitable for immunohistochemistry in the placenta, giving high background staining. We therefore used another recently available antibody to RLNH1 (see ‘Materials and methods’ section). This showed specific localization to the syncytiotrophoblast, remaining cytотrophoblast and fetal endothelium, examples are shown for Uw/N (Fig. 6A) and Ow/Ob (Fig. 6B), with a negative control (Fig. 6C). There were no differences in staining intensity in the two patient groups. Quantitation of RLNH1 according to placental weight showed different results from those of RLNH2, with no significant changes in the groups of patients, although the levels in Ow/Ob patients diverged from those of the Uw/N group (Fig. 6D). However, levels of mRNA for RLNH1 according to placental weight (Fig. 6E) showed no decline in the Uw/N patients as did mRNA for RLNH2 (Fig. 2G); indeed, placental weight tended to increase as RLNH1 mRNA increased ($P = 0.07$).

The associations of placental RLNH1 and RLNH2 proteins with placental expression of IGF2 and VEGFA
are due to their actions on the same receptor (RXFP1) (Garibay-Tupas et al. 1995) as well as the ability of RLN to increase expression of RXFP1. This has been demonstrated in vitro, in isolated decidual cells (Mazella et al. 2004) and in full-thickness fetal membrane explants (Lowndes et al. 2006). In this study, in the syncytiotrophoblast, we found a significant positive association of RLNH2 with syncytiotrophoblast RXFP1 ($P < 0.001$), and of RLNH1 with RXFP1 ($P = 0.005$) in the same tissues. Similarly, within the vascular endothelium, RLNH2 ($P = 0.009$) and RLNH1 ($P = 0.058$) were positively associated with RXFP1. Thus, both RLNH1 and RLNH2 were likely to have positively affected the expression of RXFP1.

In spite of this, there was a markedly different association of RLNH1 and RLNH2 with IGF2 and VEGFA in the trophoblast and vasculature (Table 2). This shows a highly significant association of RLNH1 with IGF2 (trophoblast $P < 0.001$, vasculature $P = 0.002$) and VEGFA (trophoblast $P = 0.001$, vasculature $P = 0.043$), and no significant associations for RLNH2 with either IGF2 or VEGFA.

Although syncytiotrophoblast RLNH1 levels showed no differences between women with male or female fetuses ($P = 0.537$), its associations with IGF2 and VEGFA differed significantly by fetal sex (Table 3). IGF2 in the trophoblast increased as trophoblast RLNH1 increased in female ($P < 0.001$) but not in male ($P = 0.6$) fetuses.
Similarly, vascular IGF2 increased as trophoblast RLNH1 increased for female ($P = 0.005$), but not for male fetuses ($P = 0.4$). Trophoblast VEGFA was also more closely associated with trophoblast RLNH1 in female ($P = 0.002$) compared to male fetuses ($P = 0.3$), whereas vascular VEGFA was also significantly associated with RLNH1 in female ($P < 0.001$) but not male fetuses ($P = 0.1$).

**Discussion**

This study is the first to suggest the possible roles for maternal systemic RLNH2 as well as placental RLNH2 and RLNH1 in fetoplacental growth. We have shown associations at term between maternal plasma RLNH2 with birthweight, syncytiotrophoblast RLNH2 with placental weight and RLNH1 with placental IGF2 and VEGFA in women with a female fetus. In addition, the relationship between maternal systemic RLNH2 and cord plasma levels suggests the possible transfer of RLNH2 from maternal blood. We have also demonstrated strong immunostaining of RXFP1 in the endothelium of placental vessels, suggesting that RLNs may act as vasodilators. Thus, these results are consistent with potential roles for RLNs in increasing placental

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*Figure 5* Placental IGF2 and fetoplacental growth. Examples of IGF2 immunostaining in (A) Uw/N, (B) Ow/Ob and (C) negative control. Anti-IGF2 only immunostained the remaining cytotrophoblast (T) and villous vascular endothelial cells (EC). (D) Quantitation showed that placental weight increased as IGF2 increased for Ow/Ob ($P < 0.001$) (open squares) but not Uw/N subjects (closed circles). (E) Birthweight increased as IGF2 increased for Ow/Ob subjects ($P = 0.002$) (open squares) and not Uw/N subjects (closed circles).

*Figure 6* Placental RLNH1 and placental weight. Examples of RLNH1 immunostaining in (A) Uw/N, (B) Ow/Ob and (C) negative control. Anti-RLNH1 immunostained the syncytiotrophoblast (ST) and villous vascular endothelial cells (EC). (D) Quantitation showed that syncytiotrophoblast RLNH1 was unrelated to placental weight for either Uw/N ($P = 0.605$) or Ow/Ob ($P = 0.202$) patients. (E) Placental RLNH1 mRNA showed a tendency to increase as placental weight increased ($P = 0.070$) in Uw/N subjects (closed circles) with no relationship in Ow/Ob patients ($P = 0.354$) (open squares).
perfusion, in addition to possibly affecting local growth factor production. Together, these could increase fetoplacental growth.

Maternal systemic RLN is only a product of the corpus luteum (Weiss et al. 1976, Johnson et al. 1991). A progressive relationship between increasing obesity and decreasing reproductive hormone secretion with evidence for decreased corpus luteum function in obese women has suggested that obesity plays a role in relative infertility (Rochester et al. 2009). It is possible that a healthier corpus luteum in the Uw/N subjects resulted in the significant relationship between maternal RLNH2 and birthweight. The absence of any relationship between maternal RLNH2 and birthweight in the Ow/Ob patients is also consistent with previous work showing decreased vascular responsiveness to systemic RLNH2 in overweight non-pregnant rats (van Drongelen et al. 2012). We have also shown no association between maternal RLNH2 and placental weight, suggesting that systemic RLNH2 is probably not involved in the regulation of placental growth, and that this is more likely to be controlled locally within the placenta. Thus, syncytiotrophoblast RLNH2 and RXFP1 were both increased in Ow/Ob women compared to the Uw/N group, which may be related to their heavier placentas. Interestingly, although increasing syncytiotrophoblast RLNH2 was associated with increasing placental weight, regardless of maternal BMI, increasing RLNH2 mRNA was associated with decreasing birthweight and placental weight in the Uw/N women, possibly reflecting an acute compensatory effect or negative feedback.

Table 2  Associations of placental RLNH1 and RLNH2 with placental expression of IGF2 and VEGFA in the trophoblast and vasculature.

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Fetal-placental growth factor</th>
<th>RLNH1*</th>
<th>RLNH2*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trophoblast</td>
<td>IGF2</td>
<td>&lt;0.001</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td>VEGFA</td>
<td>0.001</td>
<td>0.1</td>
</tr>
<tr>
<td>Vasculature</td>
<td>IGF2</td>
<td>0.002</td>
<td>0.9</td>
</tr>
<tr>
<td></td>
<td>VEGFA</td>
<td>0.043</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Results expressed as P-value of association. 
*Positive relationship for all associations.
IGF2, insulin-like growth factor 2; RLNH1, relaxin H1; RLNH2, relaxin H2; VEGFA, vascular endothelial growth factor A.

For this study, we have used some novel and newly developed commercial reagents for measurement of both RLNH1 and RLNH2 mRNAs and proteins. The high level of homology for the RLNs at both the RNA and protein levels makes specific identification of one in the presence of the other difficult. Over a period of years, we have reported the measurement of RNAs for both hormones using different strategies (Hansell et al. 1991, Garibay-Tupas et al. 1995), as well as raising antisera to synthetic peptides of RLNH1 and RLNH2 (Tashima et al. 2002). While it is normal to report methods of production of reagents, for those commercially available this can be proprietary. However, information provided on the reagents used here stated that the RLNH1 primers and probe mapped within a single exon, but genomic DNA may be detected. We therefore used only high-quality RNA devoid of DNA contamination. For the antibodies, we tried to use equivalent reagents for RLNH1 and RLNH2, but this was not possible because the equivalent RLNH1 antibody was unsuitable for immunohistochemistry. Therefore, although our anti-RLNH2 was directed to a sequence in the B-chain, the alternative anti-RLNH1 used, was to human prorelaxin-1. However, this antibody had no cross-reactivity with recombinant human prorelaxin-2. We acknowledge that these differences may be important, but our use for intracellular detection by immunohistochemistry, rather than for solution assays like Western blotting or ELISA, may have also been advantageous. Our results for RLNH1 and RLNH2 were appropriately similar for such highly homologous molecules, while in some instances, they were markedly different. This lends credence to the view that the reagents used here allowed specific detection of the two intracellular hormones. At the same time, we also recognize the need for confirmation of our results with different reagents and different methodologies.

Unfortunately, it was not possible to confirm our results by Western blotting. The low levels of RLNH1 and RLNH2 expression in the placenta, together with their adsorptive properties preclude this methodology. However, it is now possible to make quantitative immunohistochemistry like an immunoassay carried out on a tissue section, but rigorous multiple precautions need to be taken into account (Taylor & Levenson 2006, Stack et al. 2014). In addition, quantitation needs to be performed with state-of-the-art software allowing specific intracellular measurements (Fiore et al. 2012). Just as the recent availability of reagents for RLNH1, which for the first time has allowed this hormone to be studied concurrently with RLNH2, similarly, new methodologies are becoming available for the study of low levels of local hormones in tissues such as the placenta.

The two human RLNs, RLNH1 and RLNH2, both signal through RXFP1 by G-protein coupling and adenylate cyclase signaling to increase intracellular cAMP levels (Halls et al. 2009). We have shown here strong associations between the levels of both RLNs and that of RXFP1, with
RLNH2 having a stronger relationship than RLNH1 in both the syncytiotrophoblast and placental vasculature. A considerable body of work has been directed to understanding the action of RLNH2 on RXFP1, which has been shown to be complex and dependent upon the cell type (Halls et al. 2009). Activation of this receptor by RLNH2 uses G-protein signaling, but also tyrosine kinase as another signaling mechanism (Bartsch et al. 2001). Our results on the associations of RLNH1 and RLNH2 with IGF2 and VEGFA were markedly different, suggesting that different signaling mechanisms may be induced by activation of RXFP1 on the syncytiotrophoblast by these two hormones. Further work is required in order to understand how they differentially regulate these two growth factors.

We have shown that RXFP1 is well expressed in late gestation by both the syncytiotrophoblast and vascular endothelium. Vascular responsiveness to RLNH2 is associated with RXFP1 expression in the endothelium (Jelinic et al. 2014) involving the endothelin-1/nitric oxide vasodilation pathway (Novak et al. 2006). Our results are therefore consistent with a role for RLN in increasing placental perfusion, and as a consequence, it could affect the fetoplacental growth. At the same time, RLNH2 increases VEGF (Unemori et al. 1999, 2000) thereby increasing angiogenesis and influencing growth. IGF2 is a well-recognized fetoplacental growth factor (Fowden 2003, Kappil et al. 2015). The results presented here are the first to suggest a possible maternal BMI-dependent relationship between placental IGF2 and fetoplacental growth, with stronger effects in Ow/Ob women. The IGF axis in cord blood is sensitive to maternal BMI, with BMI-dependent differences in IGF-binding proteins (Ferraro et al. 2012). Further work is needed to see if altered IGF2 in maternal obesity is related to increased body fat and insulin resistance in the newborns of obese mothers (Catalano et al. 2009). Although RLNH2 has been shown to induce IGF2 in the fetal membranes (Millar et al. 2003), our results showed a stronger association with RLNH1 in the placenta, which was similar for VEGFA. As RLNH1 is limited to primates, its functional role as a placental hormone may be important. Although RLNH1 is not a product of the corpus luteum, it is produced in low levels by the syncytiotrophoblast and is perfectly positioned to target syncytiotrophoblast RXFP1 to increase growth factor production. Although this study is preliminary and we have only shown associations, it would be important in the future to show a direct linkage between RLNH1 binding to RXFP1 in the syncytiotrophoblast and the increased production of IGF2 and VEGFA.

The dogma in the RLN field is that it is of greater importance in female compared to male reproduction. We show here that the relationships between placental RLNH1 and both IGF2 and VEGFA were highly specific to female fetal sex. Human data as well as animal models demonstrate sex-specific fetal growth and metabolic adaptations to environmental insults, with female fetuses generally exhibiting greater responsiveness and resilience than male fetuses (Clifton 2005, Gonzalez-Bulnes et al. 2015, Rosenfeld 2015), with improved neonatal outcomes (Simchen et al. 2014, Wallace et al. 2016). Thus, further evaluation of RLNH1 in pathologic settings in pregnancy could assess its contribution to this well-known disparity.

Our data show associations between the RLNs and fetoplacental growth and fetoplacental growth factors, but do not prove causation. We also recognize the need for these results to be confirmed and extended using other methodologies. In addition, our data are also limited to term gestation, while fetoplacental growth at delivery is the cumulative result of processes spanning the whole pregnancy. Studies in earlier gestation are therefore needed, although it would be difficult to obtain normal human samples. In conclusion, these preliminary results suggest that maternal RLNH2 and placental RLNH1 and RLNH2 may have maternal BMI- and fetal sex-dependent associations with human fetoplacental growth.

**Declaration of interest**

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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