Role of ghrelin in fertilization, early embryo development, and implantation periods

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Abstract

In order to clarify the physiological role of ghrelin in gestation, we evaluated the effects of administration of exogenous ghrelin (2 or 4 nmol/animal per day) or its antagonist (6 nmol/animal per day of (D-Lys3)GHRP6) on fertilization, early embryo development, and implantation periods in mice. Three experiments were performed, treating female mice with ghrelin or its antagonist: i) starting from 1 week before copulation to 12 h after copulation, mice were killed at day 18 of gestation; ii) since ovulation induction until 80 h later, when we retrieved the embryos from oviducts/uterus, and iii) starting from days 3 to 7 of gestation (peri-implantation), mice were killed at day 18. In experiments 1 and 3, the antagonist and/or the highest dose of ghrelin significantly increased the percentage of atrophied fetuses and that of females exhibiting this finding or a higher amount of corpora lutea compared with fetuses (nCL/nF) (experiment 3: higher nCL/nF-atrophied fetuses: ghrelin 4, 71.4–71.4% and antagonist, 75.0–62.5% vs ghrelin 2, 46.2–15.4% and control, 10–0.0%; n=7–13 females/group; P<0.01). In experiment 2, the antagonist diminished the fertilization rate, and both, ghrelin and the antagonist, delayed embryo development (blastocysts: ghrelin 2, 62.5%; ghrelin 4, 50.6%; and antagonist, 61.0% vs control 78.4%; n=82–102 embryos/treatment; P<0.0001). In experiment 3, additionally, ghrelin (4 nmol/day) and the antagonist significantly diminished the weight gain of fetuses and dams during pregnancy. Our results indicate that not only hyperghrelinemia but also the inhibition of the endogenous ghrelin effects exerts negative effects on the fertilization, implantation, and embryo/fetal development periods, supporting the hypothesis that ghrelin (in ‘adequate’ concentrations) has a physiological role in early gestational events.

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Introduction

Ghrelin was originally identified in 1999 as the endogenous ligand of the growth hormone secretagogue receptor (GHSR1a; Kojima et al. 1999, 2001, Barreiro & Tena-Sempere 2004, Fernández-Fernández et al. 2005). This substance is a 28-amino acid peptide, mainly secreted by the stomach and the hypothalamus. Although it essentially stimulates GH secretion and food intake, several studies have linked ghrelin to the reproductive physiology, mainly with the hypothalamic–hypophyseal–gonadal axis modulation (Kojima et al. 1999, 2001, Barreiro et al. 2002, Kawamura et al. 2003, Fernández-Fernández et al. 2006, García et al. 2007, Zhang et al. 2007, Tena-Sempere 2008a,b). Moreover, as plasma ghrelin concentrations dramatically increase during fasting or undernourishment (Kojima & Kangawa 2005), this peptide has been proposed as an inhibitory signal for reproductive physiology and/or behavior during food scarcity periods (Fernández-Fernández et al. 2006, Tena-Sempere 2008a,b), especially in females (Schneider 2004, Bertoldi et al. 2011).

Nevertheless, it is well known that ghrelin plasma level in mothers and fetuses increases during pregnancy in several mammalian species, suggesting that this peptide exerts some physiological roles in gestation. Nonetheless, there is still consensus neither about the stage of gestation in which ghrelin peaks (Gualillo et al. 2001, Fernández-Fernández et al. 2004, Shibata et al. 2004, Fuglsang et al. 2005, Govoni et al. 2007, Harrison et al. 2007, Palík et al. 2007) nor about its origin, with the placenta, corpora lutea, decidual endometrium, and/or fetal pancreas postulated as possible candidates (Gualillo et al. 2001, Caminos et al. 2003, Tanaka et al. 2003, Harrison et al. 2007). Moreover, as GHSR1a has been detected in mammalian placentae, endometria, and fallopian tubes (Gaytán et al. 2005, Harrison et al. 2007), it has been proposed that ghrelin might be one of the numerous peptides that regulate embryo implantation (Tanaka et al. 2003). Concordantly, in
humans, in vitro experiments have demonstrated a decidualization-stimulating effect of ghrelin upon endometrial stromal cells in cultures (Tanaka et al. 2003). Finally, the ghrelin receptor has been identified in mammalian oocytes and embryos (Kawamura et al. 2003, Du et al. 2010), and ghrelin secretion has been linked to early embryo development. In fact, while some authors have claimed that this peptide has deleterious effects on the progress of mouse embryo to blastocyst (Kawamura et al. 2003), other authors have demonstrated that it has stimulatory dose-dependent effects in sheep or pigs (Wang et al. 2007, Zhang et al. 2007).

Given the above-mentioned data, it is evident that although ghrelin is supposed to physiologically participate in early gestational events, there is a lack of knowledge about this presumable role, due to fragmentary evidences mostly obtained from in vitro designs. Therefore, the objective of this study is to investigate, by means of an in vivo approach, the effects of exogenous ghrelin administration (2 or 4 nmol/animal per day) or endogenous ghrelin inhibition (by 6 nmol/animal per day of (D-Lys3)GHRP6) on fertilization, early embryo development, and implantation periods in pre/gestating mice. For these purposes, three experiments were performed treating female mice with two doses of ghrelin or an antagonist: i) starting from 1 week before copulation to 12 h after copulation (gestation day 1), mice were killed at day 18 of gestation; ii) starting from ovulation induction to 80 h after copulation (gestation day 1), mice were killed at day 18 of gestation; and iii) starting from days 3 to 7 of gestation (peri-implantation period), mice were killed at day 18.

Materials and methods

Animals

Experiments were conducted in accordance with the Guide for the Care and Use of Laboratory Animals from the Medicine School of the Cordoba National University (UNC-RHCS 674/09). This Animal Ethical Committee approved the protocols used in this study.

We used adult female (60–80 days) inbred Albino Swiss mice (N:NIH) maintained on a 14 h light:10 h darkness cycle at 22 ± 2 °C with water provision ad libitum. In order to avoid an increase in the body weight associated with possible orexigenic/anorexigenic effects of ghrelin or its antagonist, treated females received the same amount of food (Grupo Pilar-Gepsa, Cordoba, Argentina) as that consumed by control females at the same gestational period. Nevertheless, it is important to remark that, in previous experiments, we determined that the doses of ghrelin or its antagonist used in this study do not significantly modify daily food intake (P Torres, E M Luque & A C Martini 2013, unpublished observations).

Chemicals

Ghrelin (Pi-Proteomics, Huntsville, AL, USA) and its antagonist, (D-Lys3)GHRP6 (Sigma–Aldrich), were both dissolved in isotonic solution (0.9% ClNa solution) and injected subcutaneously, administering twice a day half the daily dose (at 0900 and 1700 h). Control animals received the vehicle in the same regimen. The doses of ghrelin used in this study were selected based on its ability to increase GH secretion from a dose–response curve performed previously and published by our group (Bertoldi et al. 2011). Furthermore, similar doses were used by other authors, with comparable results in GH secretion (Wei et al. 2006, Sun et al. 2007). Concordantly, the selected dose of the antagonist had been previously demonstrated to inhibit the effects of endogenous hyperghrelinemia or exogenously administered ghrelin, which makes this option an effective ghrelin antagonist protocol (Bertoldi et al. 2011).

Experimental groups

All the experiments were performed on female mice subjected to one of the following treatments:

i) ghrelin 2: administration of 2 nmol/animal per day of ghrelin dissolved in 0.2 ml isotonic solution.
ii) Ghrelin 4: administration of 4 nmol/animal per day of ghrelin dissolved in 0.2 ml isotonic solution.
iii) Ant: administration of 6 nmol/animal per day of (D-Lys3)-GHRP6 (ghrelin antagonist) dissolved in 0.2 ml isotonic solution.
iv) Control: administration of 0.2 ml isotonic solution/animal per day.

Experiment 1

In order to evaluate the role of ghrelin in the peri-fertilization period, we injected ghrelin or its antagonist into adult female mice starting from 1 week before the copulation confirmation day, inclusive. By the afternoon of the seventh day of treatment, a male mouse was introduced into the cage in which the female was housed, and evidences of copulation were evaluated every morning from this day onwards. All experimental females were receptive to males within the first 3 days, which implies that ghrelin and the antagonist were injected at a time period ranging from 7 to 9 days before copulation.

The day that copulation was confirmed was considered as day 1 of gestation. Females were killed on day 18 by cervical dislocation, in order to obtain ovaries and gravid uterus and to quantify the number of corpora lutea (nCL), litter size, viable pups’ weight, atrophied fetuses, and the nCL/number of fetuses (nF). In previous studies, we considered the nCL as a sign of ovulation index and we have also considered that a higher nCL/nF, depending on the treatment period, may reflect alterations in reproductive processes such as fertilization, embryo development, and/or implantation (Puechagut et al. 2012).

Experiment 2

With the objective to evaluate the role of ghrelin in early embryo development, adult females were injected with ghrelin or the antagonist starting from the day of pharmacologically induced ovulation (with 5 IU pregnant mares’ serum and,
48 h later, 10 IU human chorionic gonadotropin (hCG)) to the third day of pregnancy. By the afternoon of the hCG administration, male mice were introduced into the cages in which females were housed and copulation was confirmed the subsequent morning. After 2 days (80 h after the estimated ovulation time), females were killed in order to obtain reproductive organs and to quantify the fertilization rate and embryo development.

**Experiment 3**

In order to evaluate the role of ghrelin in the implantation process, pregnant females were injected with ghrelin or the antagonist starting from days 3 to 7 of gestation (peri-implantation period). At day 18, females were killed and again the nCL in the ovaries as well as the number and weight of the fetuses found in the uterus was evaluated. The percentage of atrophied fetuses and nCL/nF were also recorded. The initial body weight of females (pregnancy day 1) and weight gain during pregnancy were assessed, weighing dams at gestation days 7, 14, and 18. Moreover, after killing and gravid uterus removal on day 18, dams weight was assessed in order to subtract the litter weight from that of the mother.

**Confirmation of copulation**

The morning(s) after the introduction of an adult male mouse into the cage housing females, vaginal smears were collected and examined, and copulation was considered positive when a vaginal plug or spermatozoa in the vaginal smear were detected.

**Number of corpora lutea**

After females were killed at day 18, both ovaries were collected and placed in a capsule containing 2 ml modified Tyrode’s medium and the number of fresh corpora lutea/female was recorded under a stereoscopic magnifying glass. These corpora lutea are usually seen as notoriously round red-orange vascularized structures that protrude from the ovary surface. In previous experiments performed in our laboratory, we found that under control conditions, more than 80% of the females showed an equal nCL and nF. That is why we consider this parameter as an acceptable sign of ovulation (Puechagut et al. 2012).

**Atrophied fetuses**

After females were killed at day 18 of gestation, the uterus was isolated and placed in a capsule containing 2 ml modified Tyrode’s medium. Under an inverted microscope, the number of oocytes or embryos, the percentage of fertilization (the number of oocytes with respect to the total amount of oocytes and embryos recovered), and the developmental status of the embryos were recorded. The results are expressed as the percentage of non-fertilized gametes and the percentage of embryos in each one of the two developmental stages: i) we considered as ‘embryos developed up to the morula stage’, to the sum of embryos of 2, 4, or more cells plus those in the morula stage and ii) ‘blastocysts’, to the sum of compact plus expanded plus hatched blastocysts (Luque et al. 2010).

**Embryonic developmental status**

Oviducts and uteri were obtained after 80 h of estimated ovulation from those females for which copulation had been confirmed and flushed in different capsules with 1.5 ml modified Tyrode’s medium. Under an inverted microscope, the number of oocytes or embryos, the percentage of fertilization, percentage of embryos developed up to the morula stage, and percentage of blastocysts – represented in Figs 2 and 3, and 4) were analyzed by the χ² test and those expressed as mean ± S.E.M. (body weight gain during pregnancy, nCL, litter size, weight of viable pups, nCL/nF, and atrophied fetuses – represented in Tables 1 and 2, Fig. 5) were analyzed using a one-way ANOVA with Fisher’s LSD test as post-hoc comparison analysis. In all cases, P values was calculated using the following formula: 100 – (nF × 100/nCL). When we found a higher nCL/nF, we considered that some alterations had occurred in reproductive processes.

**Statistical analysis**

The results are expressed as percentage or mean ± S.E.M. Those represented as percentage (percentage of females with a higher nCL/nF, percentage of females with one or more atrophied fetuses, percentage of fertilization, percentage of embryos developed up to the morula stage, and percentage of blastocysts – represented in Figs 2 and 3, and 4) were analyzed by the χ² test and those expressed as mean ± S.E.M. (body weight gain during pregnancy, nCL, litter size, weight of viable pups, nCL/nF, and atrophied fetuses – represented in Tables 1 and 2, Fig. 5) were analyzed using a one-way ANOVA with Fisher’s LSD test as post-hoc comparison analysis. In all cases, P values was calculated using the following formula: 100 – (nF × 100/nCL). When we found a higher nCL/nF, we considered that some alterations had occurred in reproductive processes.

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The morning(s) after the introduction of an adult male mouse into the cage housing females, vaginal smears were collected and examined, and copulation was considered positive when a vaginal plug or spermatozoa in the vaginal smear were detected.
under 0.05 were considered statistically significant. Statistical analyses were performed using Infostat 2000 (Infostat version 1.1, Group Infostat, FCA-UNC, Córdoba, Argentina).

Results

Experiment 1

When female mice received treatment starting from 7 to 9 days before copulation to 12 h after copulation, we found that on day 18 of gestation, neither ghrelin (in any dose) nor the antagonist modified the nCL, litter size, or weight of viable pups (Table 1). In those females treated with ghrelin (both doses) or the antagonist, we did find an increase, although not significant, in the percentage of nCL/nF with respect to control. A similar increasing tendency was detected in the percentage of atrophied fetuses, reaching statistical difference only in the group treated with the antagonist vs control. On the other hand, we detected an important increase in the percentages of treated females exhibiting a higher nCL/nF or some degree of fetal atrophy with respect to controls (Fig. 2). These differences reached statistical significance for a higher nCL/nF in females injected with ghrelin and for atrophied fetuses in mice treated with the antagonist.

Experiment 2

When treating females starting from the day of pharmacological induction of ovulation to the third day of pregnancy, we observed that the antagonist significantly diminished the fertilization index compared with the other groups. Additionally, both doses of ghrelin (in a dose-dependent manner) and the antagonist significantly delayed embryo development; this was demonstrated by the increase in the percentage of embryos that developed up to the morula stage and a decrease in the proportion of blastocysts (20–35% reduction; Fig. 3).

It is important to mention that, as expected for the time elapsed from estimated ovulation to embryo retrieval, only 6.4% of total oocytes/embryos were obtained from oviductal flushing (93.6% were obtained from uterus). As we did not find any differences in the developmental stage of the embryos retrieved from oviducts vs uterus or in the proportion of females exhibiting oocytes/embryos in the oviducts between experimental groups (results not shown), for statistical purposes, all retrieved oocytes/embryos, independent of their origin, were considered together in each group.

Experiment 3

In females treated with ghrelin or the antagonist during the peri-implantation period (days 3–7 of pregnancy) and killed at day 18, we did not observe any differences in the litter size (including atrophied fetuses), but found a significant decrease in the weight of viable pups, in all treatments vs control. This reduction showed higher values in females treated with the highest dose of ghrelin.
Atrophied fetuses were those remarkably smaller than the normal ones at gestation day 18. Ghrelin or its antagonist was injected subcutaneously twice a day (half of the daily dose in each injection). Control females were injected with the vehicle (isotonic solution) in the same regimen. The results are expressed as the percentage of females showing each characteristic. Fetal atrophy was considered as the presence of one or more fetuses remarkably smaller than the normal ones. The number of females in each group: control = 10, ghrelin (2 nmol/animal per day) = 13, ghrelin (4 nmol/animal per day) = 7, and (D-Lys₃)GHRP₆ = 8. a, b, c and d, P<0.01 vs control.

or with the antagonist (Table 2). In the former two groups, we also found an increase in the percentage of nCL/nF or that of atrophied fetuses, although these differences reached statistical significance only for fetal atrophy (Table 2). Concordantly, when these two parameters were expressed in relation to the proportion of females with a higher nCL/nF or some degree of fetal atrophy, we found that ghrelin (4 nmol/animal per day) and the antagonist significantly increased the percentage of females with both features when compared with control ones (Fig. 4).

Finally, when evaluating the increase in the weight of dams during pregnancy, we found that the highest dose of ghrelin and the antagonist significantly decreased the weight gain in females during pregnancy with respect to control or ghrelin (2 nmol/animal per day; Fig. 5). This phenomenon may be attributed to the smaller weight of the viable pups plus the increase in the percentage of atrophied fetuses (fetuses very small for their gestational age), as we found no differences in the body weight of females (at day 18) after gravid uterus removal (‘dams weight gain’, Fig. 5).

### Discussion

The main objective of our study is to explore the physiological role of ghrelin in early gestational events, by means of an in vivo approach of hyperghrelinemia or inhibiting endogenous ghrelin effects. The results of this experimental design support the hypothesis that ghrelin modulates several gestational processes, including fertilization, early embryo development, and possibly implantation, exerting detrimental effects at high concentrations, at which this peptide may function as a food scarcity signal. Moreover, the administration of the antagonist provoked deleterious effects on these processes as well, suggesting that the physiological increase in plasma ghrelin concentrations during gestation would be necessary for a normal/optimal pregnancy progress. A number of new experiments are being currently developed in our laboratory in order to elucidate the cellular/molecular basis of such actions.

One of the major findings of our study was that not only ghrelin (in its highest dose) but also its antagonist significantly increased the percentage of nCL/nF or that of atrophied fetuses and/or the percentages of females exhibiting a higher nCL/nF or fetal atrophy in comparison to dams injected with the vehicle (experiments 1 and 3). As the first parameter was calculated as the amount of fetuses with respect to the nCL (suggestive of ovulation index), a smaller number of conceptuses may reflect alterations at diverse times of the process, i.e. at fertilization, early embryo development, and/or implantation. As females were treated during the peri-implantation period in the third experiment (in which the percentage of nCL/nF was more evident), the impairment of the fertilization process by ghrelin or the antagonist, at least in this experiment, is not a suitable explanation. A negative effect of the different treatments upon implantation and/or placenta formation/function seems to be more feasible; the results detected in experiment 3 for atrophied fetuses support

### Table 1

Reproductive success of female mice injected with exogenous ghrelin or with (D-Lys₃)GHRP₆ (ghrelin antagonist) starting from 1 week before copulation.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Control (n=9)</th>
<th>Ghrelin (2 nmol/animal per day) (n=9)</th>
<th>Ghrelin (4 nmol/animal per day) (n=11)</th>
<th>(D-Lys₃)GHRP₆ (6 nmol/animal per day) (n=8)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of corpora lutea</td>
<td>11.9±0.6</td>
<td>12.7±0.8</td>
<td>12.6±0.6</td>
<td>13.3±0.7</td>
</tr>
<tr>
<td>Litter size</td>
<td>11.7±0.7</td>
<td>11.7±1.1</td>
<td>11.3±0.7</td>
<td>12.3±0.6</td>
</tr>
<tr>
<td>Weight of viable pups (g)</td>
<td>1.0±0.1</td>
<td>1.1±0.0</td>
<td>0.9±0.1</td>
<td>1.0±0.1</td>
</tr>
<tr>
<td>nCL/nF (%)</td>
<td>2.2±2.2</td>
<td>7.3±6.7</td>
<td>9.8±5.4</td>
<td>6.8±3.8</td>
</tr>
<tr>
<td>Atrophied fetuses (%)</td>
<td>0.1±0.1</td>
<td>0.7±0.4</td>
<td>0.6±0.3</td>
<td>1.9±0.5*</td>
</tr>
</tbody>
</table>

Ghrelin or its antagonist was injected subcutaneously twice a day (half of the daily dose in each injection). Control females were injected with the vehicle (isotonic solution) in the same regimen. The results are expressed as mean±S.E.M. nCL/nF, number of corpora lutea/number of fetuses, calculated as 100−(nF at gestation day 18×100/nCL). When positive, this index was considered as a sign of alterations in reproductive processes. Atrophied fetuses were those remarkably smaller than the normal ones at gestation day 18. n, number of females. *P=0.0244 vs control.
Table 2 Reproductive success of female mice injected with exogenous ghrelin or with (D-Lys3)GHRP6 (ghrelin antagonist) starting from days 3 to 7 of pregnancy (peri-implantation period).

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Control (n = 12)</th>
<th>Ghrelin (2 nmol/animal per day) (n = 14)</th>
<th>Ghrelin (4 nmol/animal per day) (n = 9)</th>
<th>(D-Lys3)GHRP6 (6 nmol/animal per day) (n = 11)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of corpora lutea</td>
<td>12.0±0.3</td>
<td>13.5±0.8</td>
<td>13.4±0.7</td>
<td>13.1±0.7</td>
</tr>
<tr>
<td>Litter size</td>
<td>11.5±0.6</td>
<td>12.2±1</td>
<td>10.0±1.2</td>
<td>12.3±0.9</td>
</tr>
<tr>
<td>Weight of viable pups (g)</td>
<td>1.0±0.3</td>
<td>0.9±0.1*</td>
<td>0.8±0.1†</td>
<td>0.8±0.1‡</td>
</tr>
<tr>
<td>Atrophied fetuses (%)</td>
<td>0.0±0.0</td>
<td>0.2±0.2</td>
<td>1.3±0.4†</td>
<td>0.9±0.3</td>
</tr>
</tbody>
</table>

Ghrelin or its antagonist was injected subcutaneously twice a day (half of the daily dose in each injection). Control females were injected with the vehicle (isotonic solution) in the same regimen. The results are expressed as mean±S.E.M. nCL/nF, number of corpora lutea/number of fetuses, calculated as 100–(nF at gestation day 18×100/nCL). When positive, this index was considered as a sign of alterations in reproductive processes. Atrophied fetuses were considered as those remarkably smaller than the normal ones at gestation day 18. n, number of females. *P=0.0488 vs control; †P=0.0020 (ghrelin 4 and antagonist vs control); and ‡P=0.0020 (ghrelin 4 vs ghrelin (2 nmol)).

This hypothesis. In fact, these two parameters may be a reflection of altered placenta formation and/or function.

With regard to this fact, ghrelin has been proposed as one of the multiple signals involved in the crosstalk between the invading placenta and the endometrium (Tanaka et al. 2003, Tawadros et al. 2007). Tanaka et al. suggested that ghrelin may be involved in endometrial embryo receptivity and that it may influence pre-implantation embryo development, acting as a paracrine/autocrine factor. These authors performed an immunohistochemical analysis demonstrating that the strongest signal for ghrelin was found in the extravillous trophoblast cells from the first trimester placenta, which constitutes the frontier of invasion into the maternal endometrium. Moreover, they found that the in vitro addition of ghrelin to the culture medium stimulated the decidualization of human endometrial stromal cells obtained in the luteal phase (Tanaka et al. 2003). Similarly, other authors have reported that the in vitro culture of the placenta cell line JEG-3 with ghrelin (100–1000 pg/ml) for 48 h significantly stimulated cell proliferation and decreased apoptosis (Rak-Mardyla & Gregoraszczuk 2010). These evidences are consistent with a dramatic increase in the peptide mRNA expression in the endometrium during early pregnancy and with the endometrial expression of the GHSR (GHSR1a) throughout the normal menstrual cycle and early pregnancy, demonstrated in vivo by Tanaka et al. (2003). Interestingly, these same authors found that in patients with ectopic pregnancies, ghrelin mRNA levels did not increase and ghrelin immunoreactivity was not detected in decidual cells.

All these evidences support the hypothesis that the increase in ghrelin secretion typical of early pregnancy contributes to endometrial decidualization and placenta formation. Therefore, it is possible that the alterations in this expected ghrelin raise (i.e. over increase or inhibition) exert deleterious effects on placenta.

Currently, investigations are being developed in our laboratory on the association between ghrelin and nitric oxide (NO) synthesis during the peri-implantation period and its effect on placental development. The placenta is a fast growing tissue that requires active blood supply and rapid vascular development endotherial NO synthases (iNOS and eNOS) are known to be involved in implantation regulation, placental formation, and placental nutrient transport to fetuses (Sladek et al. 1997, Kaufmann et al. 2003, Maul et al. 2003, Khan et al. 2012, Kulandalevu et al. 2012, Kusinski et al. 2012). On the other hand, in non-reproductive models, it has been demonstrated that ghrelin enhances NOS activity and consequently NO synthesis (Rodriguez-Pacheco et al. 2008, Morley et al. 2011, Sudar et al. 2011, Wang et al. 2012). All these evidences suggest that the detrimental effects on the implantation ratio observed in our study may be linked to an alteration in the NO synthesis exerted by abnormal ghrelin concentrations. Moreover, the significant decrease in fetal body weight at gestational day 18 may be related to a reduced nutrient transport ability of the placenta.

Figure 5 Weight gain of dams of females injected with ghrelin (2 or 4 nmol/animal per day) or ghrelin antagonist (D-Lys3)GHRP6 (6 nmol/animal per day) starting from days 3 to 7 of gestation. Ghrelin and the antagonist were injected subcutaneously twice a day (half of the daily dose at each injection). Control females were injected with the vehicle (isotonic solution) in the same regimen. The results are expressed as the percentage of weight gain with respect to day 1 of gestation, calculated using the formula: (body weight×100/body weight at day 1)–100. In addition, after killing and gravid uterus removal on day 18, the weight of dams was assessed in order to subtract the litter weight from that of the mother (‘dams weight gain’). The number of females in each group: control = 12, ghrelin (2 nmol) = 14, ghrelin (4 nmol) = 9, and (D-Lys3) GHRP6 = 11.

a, P=0.0392 vs control and ghrelin; b, P=0.0292 vs control and ghrelin 2; c and d, P=0.0222 vs control and ghrelin 2.
A special comment is worthy on the increase in the percentage of females exhibiting a higher nCL/nF or atrophied fetuses observed in experiment 1, which included treatment with ghrelin or the antagonist starting from 1 week before the day of copulation, inclusive. This treatment did not comprise the implantation phase; in fact, it includes ovulation and presumably fertilization, as well as a fraction of the early embryo development period. In conclusion, the alterations found in this experiment at the implantation and/or fetal development processes may be linked to defects of other parameters such as oocyte quality or progesterone secretion by corpus luteum (Li et al. 2011, Rak-Mardyla et al. 2012). Regarding this issue, it has been reported that the corpora lutea secrete ghrelin, especially in the luteal phase and that this peptide significantly inhibits 3β-hydroxysteroid dehydrogenase activity and progesterone secretion (Rak-Mardyla et al. 2012). In agreement with these findings, results in human obtained in an assisted reproduction center have demonstrated that the follicular fluid concentrations of ghrelin negatively correlated with progesterone values (Li et al. 2011). More importantly, these authors informed that follicular fluid ghrelin negatively correlated with the embryo cleavage rate and number of viable embryos at day 3, explained by alterations in oocyte quality (Li et al. 2011). Some experiments are being currently performed in our laboratory, destined to evaluate, in our experimental conditions, oocyte morphology, maturation, and spontaneous activation.

In our study, we found that the treatment of pregnant females with ghrelin or its antagonist during the early embryo developmental period (starting from hCG injection to 80 h after estimated ovulation) significantly delayed embryo development to blastocyst in a dose-dependent manner. A study of the in vitro effects of different doses of ghrelin on ovine oocyte maturation and embryo development has been recently published (Wang et al. 2013). While the smaller doses of ghrelin (50 ng/ml) increased blastocyst rates and the total number of cells per blastocyst (compared with 0 or 10 ng/ml), a higher dose (250 ng/ml) decreased these parameters, suggesting that there is an ‘appropriate’ concentration of the peptide that promotes blastocyst formation (Wang et al. 2013). This is consistent with the results obtained in our study, in which, the addition of not only ghrelin but also the antagonist altered embryo progress. Other authors have also performed in vitro experiments and observed that ghrelin significantly diminished the progression of two-cell embryos to blastocysts, and that coinubcation with ghrelin and the same antagonist used in our study reversed these deleterious effects (Kawamura et al. 2003). To our knowledge, this is the first study that evaluates the in vitro effects of administration of ghrelin and its antagonist on embryo development.

In our study, in this same experiment, the antagonist significantly diminished the fertilization index. To our knowledge, such an effect has not been informed in any of the previous studies. In vitro experiments are being currently performed in our laboratory in order to elucidate these aspects.

It is important to notice that the results obtained in the former experiment were for females with pharmacologically induced superovulation. It has been reported that this treatment may alter the oocyte quality/maturity and consequently embryo development (Erzegd & Storeng 2001). With this in mind, the actual effects of ghrelin or the antagonist during early embryo development in naturally ovulated females are still unknown.

As mentioned before, in experiment 3, on day 18, we found a significant decrease in the weight of fetuses obtained from dams treated with ghrelin or the antagonist during the peri-implantation period. This may explain the reduction in the weight gain of dams during pregnancy (in groups ghrelin (4 nmol) and antagonist), as there were no differences in the body weight of these females after removal of the gravid uterus. Indeed, the reduction in the weight gain of dams was maximal in ghrelin (4 nmol) group, which is the one that presented the smallest litter size and weight of pups. Concordantly, in previous studies carried out in our laboratory with a ghrelin analog called hexarelin, we found that this substance, when administered during the first week of pregnancy, significantly increased the percentage of nCL/nF and tended to reduce the litter size (Luque et al. 2010, Puechagut et al. 2012).

It has been reported that ghrelin administered to pregnant rats easily crosses through the placenta to the fetus blood and amniotic fluid and that this substance stimulates epithelial cell proliferation in cultures (Nakahara et al. 2006). Besides, several authors have informed that the administration of ghrelin to pregnant rats (starting from pregnancy days 1 to 11 or from day 14 or 15 to delivery) increases the weight of pups at delivery (Hayashida et al. 2002, Fernández-Fernández et al. 2005, Nakahara et al. 2006). This fact has been confirmed by our team in mice treated with 4 nmol ghrelin during the whole gestation period (P Torres, E Luque and A C Martini, unpublished observations). Nevertheless, in this study, ghrelin was administered only during the peri-implantation period, in which it exerted detrimental effects on fetal weight. Nonetheless, not only ghrelin but also the antagonist exerted these negative actions, suggesting that the peri-implantation period is a critical stage in which ‘adequate’ ghrelin concentrations would be necessary for placenta formation and/or functioning. As mentioned above, it is well known that a functionally altered placenta may provoke a reduction in fetus development and weight gain (Kusinski et al. 2012); this is probably the explanation of our results, as the percentages of nCL/nF and atrophied fetuses were higher in the three experimental groups, especially when using 4 nmol ghrelin.

Finally, although so far descriptive, our in vivo study supports the hypothesis that during early gestation,
‘adequate’ ghrelin concentrations would be necessary to guarantee normal/optimal pregnancy progression. Certainly, additional experiments will provide further information about the cellular/molecular basis of these results.

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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Author contribution statement

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