Insulin-like growth factor (IGF)-I, IGF-II, IGF-binding protein-2 and pregnancy-associated glycoprotein mRNA in pigs with somatotropin-enhanced fetal growth

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Abstract

Fetal growth is increased when pregnant gilts are treated with recombinant porcine somatotropin. The mechanism for increased fetal growth was examined by measuring the expression of IGF-I and -II and IGF-binding protein-2 (IGFBP-2) mRNA in liver and reproductive tissues of somatotropin- and saline-treated pregnant gilts.

Twenty-four pregnant gilts received daily injections of either saline (control; n=12) or 5 mg recombinant porcine somatotropin (n=12) from day 30 to day 43 of gestation. Gilts were slaughtered on day 44 of gestation and liver, ovary, placenta, placental uterus (uterus with adjacent placental tissue) and non-placental uterus (region of the necrotic tip) were collected. The mRNAs for somatotropin receptor, IGFs -I and -II, IGFBP-2 and pregnancy-associated glycoprotein (a marker of trophoblast tissue) were analyzed by Northern blotting or ribonuclease protection assay.

Gilts treated with somatotropin had heavier fetuses and placentas. The concentration of mRNA for the components of the IGF system was tissue-dependent. The uterine IGF-I mRNA concentration was greater in non-placental than in placental uterus. The greatest IGF-II mRNA concentration was observed in placenta, and adjacent uterine tissue expressed IGFBP-2 mRNA intensely. In non-placental uterus, IGFBP-2 mRNA was nearly undetectable. Somatotropin-dependent regulation of IGF-I was only observed in liver, where the greatest somatotropin receptor mRNA concentration was found. In the pregnant uterus, somatotropin failed to change the concentration of IGF or IGFBP-2 mRNA. Pregnancy-associated glycoprotein mRNA concentration was decreased by somatotropin.

In summary, increased fetal growth in somatotropin-treated pregnant pigs was not associated with changes in IGF or IGFBP-2 mRNA concentration in reproductive tissues. Other mechanisms, therefore, lead to enhanced fetal growth in somatotropin-treated pregnant pigs.

Introduction

Fetal growth is a complex process that depends on the transfer of maternal nutrients across the placenta, and the expression of fetal and placental growth factors and morphogens that control development (D’Ercole & Underwood 1981, Han & Hill 1992). The rate of fetal growth is partially dependent on the insulin–like growth factor (IGF) system that co-ordinates several aspects of uterine, placental, and fetal development (Owens 1991). The IGF system consists of two related growth factors (IGF–I and –II) and a group of IGF–binding proteins (IGFBPs; Jones & Clemmons 1995). The IGFBPs can modify IGF activity by binding IGFs and preventing IGF receptor activation. It is also possible that IGFBPs have actions that are independent of their roles as proteins that bind IGFs (Rechler 1997).

Uterine tissues express components of the IGF system in a tissue- and developmentally specific pattern. In the pig, most IGF-I mRNA is found within the myometrium (Song et al. 1996), but endometrial IGF-I is steroidogenically regulated during early pregnancy (Simmen et al. 1990). The placenta is the primary location of fetal IGF-II production. Later in porcine pregnancy, endometrial IGF-I decreases (Simmen et al. 1992), but uterine IGFBP-2 (the predominant IGFBP in porcine uterus during pregnancy) increases within the endometrium (Song et al. 1996). The importance of IGF-I and IGF-II for fetal development was demonstrated in gene knockout experiments in which mice with deletion of either gene were born at 60% of normal weight (Baker et al. 1993). The conclusion was that both IGF-I and IGF-II are required for fetal growth.

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In addition to the locally acting IGFs (either uterine IGF-I or placental IGF-II), maternal blood IGF-I concentrations appear to play a role in fetal development. In humans, farm animals, and laboratory animals, maternal IGF-I concentrations are correlated with the size of the fetus and birth weight (Hall et al. 1986, Gluckman & Barry 1988, Lassarre et al. 1991, Gluckman et al. 1992). It may be possible, therefore, to increase fetal development and birth weight by increasing maternal IGF-I concentrations during pregnancy. Indeed, fetal or neonatal weight may be increased when recombinant porcine somatotropin (rpST) was administered and maternal IGF-I increased during pregnancy. Indeed, fetal or neonatal weight and birth weight by increasing maternal IGF-I concentration may be possible, therefore, to increase fetal development directly in response to somatotropin are undefined. Any method to increase birth weight and improve neonatal survival.

The molecular mechanisms leading to increased fetal growth in response to somatotropin are undefined. Any direct effects of rpST on the pregnant uterus should depend on somatotropin receptor location. The objectives of this study were, therefore, to determine the tissue location of somatotropin receptor in the pregnant pig and to assay liver (the traditional site for somatotropin action) as well as reproductive tissues for changes in IGF-I, IGF-II, and IGFBP-2 gene expression in response to rpST during pregnancy. Pregnancy-associated glycoprotein (PAG) was also measured as a marker of placental development.

**Materials and Methods**

**Animals, treatments and collection of tissues**

Twenty-four crossbred (Yorkshire × Landrace) gilts (Sus scrofa, domestica) were used. Gilts were artificially inseminated at their second post-pubertal estrus (6–8 months of age). Approximately 21 days following insemination, gilts were checked with a boar for return to estrus, and blood was analyzed for progesterone concentration. Gilts that failed to express estrus and had elevated progesterone (>10 ng/ml) were classified as pregnant.

Pregnant gilts were assigned to the experiment. The experiment contained three replicates. Replicates 1 and 2 consisted of 6 gilts each (12 gilts total; 6 rpST and 6 control). Replicate 3 consisted of 12 gilts (6 rpST and 6 control). Gilts in replicate 3 had one uterine horn ligated to give a 2:1 ratio of uterine space per potential conceptus between the crowded and uncrowded pregnant horns (Dziuk 1968, Sterle et al. 1995). The uterine crowding was done to satisfy the objectives of a related experiment (Sterle et al. 1995) and did not compromise the objective of the present experiment, because treatment (rpST or saline) was balanced within the replicate.

Gilts were randomly assigned within each replicate to receive daily intramuscular injections of 1 ml saline (control) or 5 mg rpST (Monsanto, St Louis, MO, USA) in 1 ml injectable water from day 30 to day 43 of gestation. Gilts were grouped until day 44 of gestation, when they were slaughtered by electrocution and exsanguination according to University of Missouri Animal Care and Use Committee guidelines.

The uterus was recovered within 5 min of exsanguination and individual fetuses were weighed. Immediately after removal of the fetuses, samples of ovary (corpora lutea (CL) and follicles isolated and frozen separately or frozen together as whole ovary), uterus, placenta and liver were placed in separate plastic vials and frozen in liquid nitrogen. Uterine tissues were collected from two locations. The first location was the uterus adjacent to the placenta (placental uterus). The placenta was stripped from the uterus, weighed, and uterine and placental tissue sampled and frozen separately. The placenta and placental uterus were collected from the second fetus of a uterine horn (the first fetus was adjacent to the tip of uterus). A second uterine sample was taken from the region between the second and third placenta, in the region of the necrotic tips. This was termed non-placental uterus. Uterine samples were collected from the crowded uterine horn in replicate 3. In four gilts (two rpST and two control), endometrium and myometrium were isolated from a second placental location. After freezing in liquid nitrogen, all samples were stored at −80 °C until RNA extraction.

**Isolation of RNA**

Unless otherwise noted, chemicals and reagents were purchased from the Sigma Chemical Company (St Louis, MO, USA). Frozen tissue was removed from storage at −80 °C and mortared to a fine powder under liquid nitrogen. The powder was then homogenized in 4 M guanidinium thiocyanate, and RNA isolated according to Chomczynski & Sacchi (1987). Integrity of rRNA was monitored by electrophoresis of an aliquot of the RNA preparation through 1% agarose in Tris–borate/EDTA (0·09 M Tris–borate, 0·002 M EDTA) with 1·25 µM ethidium bromide, followed by ultraviolet illumination. Purified RNA was dissolved in H2O, quantified based on A260 measurement, and stored at −80 °C.

**Northern blotting**

Five of six gilts from replicates 1 and 2 were randomly selected from each treatment for analyses of somatotropin...
receptor, IGF-II, IGFBP-2 and PAG mRNA and 28S rRNA (control) by Northern blotting. Analysis of RNA by Northern blotting was performed as previously described (Lucy et al. 1993). Total cellular RNA (25 μg) was electrophoresed after denaturation with 1 M glyoxal and 50% dimethyl sulfoxide in a 1:2% agarose gel in 10 mM sodium phosphate (pH 6.5). A reduced amount of RNA (2.5 μg) was used for IGF-II and PAG blots for placenta samples. The RNA was transferred to nitrocellulose by capillary transfer for 15 h in 20-strength SSPE (3 M NaCl, 0·2 M NaH₂PO₄, 0·2 M EDTA, pH 7·0) and the nitrocellulose was prehybridized for 2 h at 42°C in 50% (v/v) formamide, 5-strength SSPE, 5-strength Denhardt’s solution, 0·1% (w/v) SDS and 100 μg/ml tRNA. Following prehybridization, the filter was incubated in the prehybridization solution for 24 h at 42°C with a ³²P-labelled cDNA probe.

The cDNA probes were bovine somatotropin receptor (Hauser et al. 1990), porcine IGF-II (Yuan et al. 1996a), bovine IGFBP-2 (Kirby et al. 1996), porcine PAG (a mixture of porcine PAG1 and PAG2; Szafranska et al. 1996) and porcine 28S rRNA (Yuan et al. 1996b). Labeling of the cDNA probes was by random priming (Multiprime DNA Labeling Systems, Amersham International, Amersham, Bucks, UK) using [α-³²P]dCTP.

Northern blot washing conditions were: 2 × 15 min: 1-strength SSC (0·15 M NaCl, 0·017 M sodium citrate, pH 7·0), 0·1% SDS, 50°C; followed by 10 min: 0·1-strength SSC, 0·1% SDS, 50°C. Autoradiography was performed using XOMAT-AR film (Eastman Kodak, Rochester, NY, USA) for 12–72 h at −80°C with an intensifying screen. As a positive control for RNA loading and transfer, blots were stripped of labeled probe by incubating in boiling H₂O for 30 min and hybridized with a 28S rRNA probe using the conditions described above. Some blots were stripped and hybridized several times with several different cDNA probes.

**Analysis of RNA by ribonuclease protection assay**

The porcine IGF-I ribonuclease protection assay was used for analyses in replicates 1, 2 and 3. Additional gilts in replicate 3 were used because IGF-I mRNA showed greater between-animal variability than the other mRNA. Therefore, to increase the sensitivity of the statistical analyses, replicate 3 gilts were randomly selected from each treatment and analyzed for IGF-I mRNA. The porcine IGF-I cDNA fragment used for production of the complementary ribonucleotide probe has been described previously (Yuan et al. 1996a). Antisense ribonucleotide probes were generated using a Riboprobe Gemini II Core system (Promega Corporation, Madison, WI, USA). Linearized plasmid (200 ng) was incubated with T7 polymerase (Promega), [α-³²P]dCTP (New England Nuclear, Boston, MA, USA) and appropriate buffers to yield an antisense ribonucleotide probe. Before use in nuclease protection assays, ribonucleotide probes were extracted with phenol:chloroform (1:1) and then chloroform. Unincorporated [α-³²P]dCTP was removed by centrifugation through a G50 Sephadex spin column (Boehringer-Mannheim, Indianapolis, IN, USA). Ribonuclease protection assays (Lee & Costlow 1987) were performed on 25 μg total cellular RNA by using the RPA II kit (Ambion Inc., Austin, TX, USA). Protected mRNA fragments were identified by their electrophoretic mobility through 8% acrylamide, 8 M urea gels (Acryl-A-Mix 8, Promega). Gels were dried and autoradiography was performed using XOMAT-AR film for 24 h at −80°C with intensifying screens.

**Statistical analyses**

Relative amounts of mRNA were measured by scanning densitometry of autoradiographs. The GP Tools version 3·0 (BioPhotonics Corporation, Ann Arbor, MI, USA) program was used to determine density. The 28S ribosomal band from either a Northern blot (samples analyzed by Northern blotting) or an ethidium bromide-stained gel (samples analyzed by ribonuclease protection assay) was used to normalize the signal for each mRNA by calculating a ratio (arbitrary units) of mRNA to 28S RNA. Data were analyzed by least squares analysis of variance (SAS Institute 1987). A statistical model that included both factors (i.e. tissue, and the main effect of treatment). Tissue means were separated by Duncan’s multiple range test. A statistical model that included both factors (i.e. tissue, and the main effect of treatment). Tissue means were separated by Duncan’s multiple range test. A statistical model that included the main effect of tissue was used to determine differences in mRNA expression across different locations (i.e. liver versus ovary versus uterus etc.). Tissue means were separated by Duncan’s multiple range test. A statistical model that included the main effect of treatment (rpST or control) was used to evaluate the response of a single tissue treatment. The effect of replicate (1 to 3) was included for the analyses of IGF-I mRNA concentration. The statistical model included the effects of treatment, replicate, and treatment by replicate. Data are presented as least squares means ± standard errors of the least squares mean. Unless stated otherwise, data were considered statistically significant at P<0.05.

**Results**

**Fetal and placental development**

Maternal and fetal blood IGF-I and IGF-II, as well as gross weights and measurements of the uterus and fetuses, were reported in a previous publication (Sterle et al. 1995). The rpST-treated pigs had greater maternal blood IGF-I (268±14 versus 78±14 ng/ml on day 43; rpST versus control; P<0.01), heavier placentas (71±2±3·5 versus 58±4±3·4; P<0.02), and tended to have greater fetal blood IGF-I (14·2±1·4 versus 10·0±1·5 ng/ml;
Number of fetuses per pregnancy, embryonic survival (day 44), uterine weight, and fetal blood IGF-II concentrations were similar for rpST and control animals. Maternal blood IGF-II concentrations decreased during rpST treatment (191 ± 14 versus 283 ± 14 ng/ml on day 43; rpST versus control; \( P < 0.05 \)).

**Somatotropin receptor mRNA**

The somatotropin receptor mRNA was analyzed in two pigs (one rpST and one control; the effect of treatment on somatotropin receptor mRNA was not tested) to determine the relative concentrations of somatotropin receptor in liver and reproductive tissues. The somatotropin receptor mRNA was detected as 4·0 kb mRNA on Northern blots (Fig. 1; mRNA from one of two pigs is shown). There was an effect of tissue on somatotropin receptor mRNA concentration (\( P < 0.01 \)). Liver (28·5 ± 3·6 arbitrary units) expressed at least seven-fold more somatotropin receptor mRNA than reproductive tissues (ovary (mixture of corpus luteum, follicles and stroma), corpus luteum (CL), ovarian follicles (Fol), placental uterus (Plac ut), myometrium (Myo), endometrium (Endo), non-placental uterus (n-Plac ut) and placenta (Plac)). The concentration of somatotropin receptor mRNA was similar within reproductive tissues (\( P > 0.05 \)).

**IGF-I mRNA**

A subset of samples was analyzed within a single ribonuclease protection assay for a tissue comparison of IGF-I mRNA concentration (Fig. 2; placenta was not analyzed because our preliminary Northern blot analyses failed to detect IGF-I mRNA in placenta). There was a tissue by treatment interaction (\( P < 0.05 \)) for IGF-I mRNA concentration (Fig. 2). Non-placental uterus (4·9 ± 0·7 arbitrary units) had a greater IGF-I mRNA concentration than placental uterus (1·7 ± 0·7) or ovary (2·1 ± 0·7), and the concentration of IGF-I mRNA in these tissues was not changed by rpST. The IGF-I mRNA concentration in liver, however, was increased in rpST-treated pigs (6·1 ± 1·0 versus 0·4 ± 1·0; rpST versus control respectively). The analyses of additional samples confirmed that IGF-I mRNA concentration was increased in liver by rpST (\( P < 0.01 \)), whereas ovarian or uterine IGF-I mRNA concentrations were similar for rpST and control (Table 1). An effect of replicate was detected for uterus because the crowded uterus (replicate 3) had greater IGF-I mRNA concentrations in placental uterus (0·3 ± 0·1, 0·4 ± 0·1 and 0·8 ± 0·1 for replicates 1, 2 and 3 respectively; \( P < 0.10 \)) and non-placental uterus (0·1 ± 0·1, 0·2 ± 0·1 and 0·4 ± 0·1; replicates 1, 2 and 3 respectively; \( P < 0.05 \)). There was no effect of replicate on ovarian IGF-I mRNA concentration (\( P > 0.05 \)).

**IGF-II mRNA**

The mRNA for IGF-II was detected as a 2·5 kb band on Northern blots (Fig. 3A). A tissue comparison, a subset of mRNA samples (\( n = 24 \)) was analyzed for IGF-II mRNA on a single Northern blot (six tissues from two rpST and two control pigs). Greatest expression of mRNA for IGF-II was found in placenta (1·5 ± 0·1 arbitrary units; Fig. 3A; tissue \( P < 0.001 \)). Lower concentrations of mRNA for IGF-II were found in liver (0·6 ± 0·1), CL (0·5 ± 0·1), follicle (0·7 ± 0·1), ovary (0·5 ± 0·1) and placental uterus (0·7 ± 0·1). The presence of IGF-II mRNA in placental uterus may have been associated with placental tissue interdigitated into the uterine wall, because PAG mRNA was detected within placental uterus samples (see below). These interdigitated placental fragments were not removed by manual stripping of the placenta from the uterus. Statistical analyses of the subset of mRNA samples did not detect a treatment or treatment by tissue interaction. Analyses of a greater number of samples for IGF-II mRNA resulted in similar conclusions (i.e. similar IGF-II mRNA concentrations for rpST and control groups within the liver, reproductive tissues and placenta (Fig. 3B; Table 1)).
IGFBP-2 mRNA

The mRNA for IGFBP-2 was detected as a 1.5 kb band on Northern blots. There was an effect of tissue (P<0.001) on IGFBP-2 mRNA concentration. The placental uterus and non-placental uterus had distinctly different IGFBP-2 mRNA concentrations. The expression of IGFBP-2 was greatest within the endometrial layer of the placental uterus (26.0±1.1 arbitrary units; Fig. 4A). The second greatest mRNA concentration was in placental uterus (sample included endometrium and myometrium; 15.0±1.1). Other tissues (including non-placental uterus) contained less IGFBP-2 mRNA than placental endometrium or placental uterus (liver (1.1±1.1), CL (1.9±1.1), ovarian follicle (2.1±1.1), whole ovary (mixture of follicles, CL and stroma; 1.6±1.1), myometrium (1.0±1.1), non-placental uterus (2.5±1.1) and placenta (4.6±1.1)). A greater number of samples was analyzed to determine the effects of treatment on IGFBP-2 mRNA concentration. The IGFBP-2 mRNA concentrations were similar within liver, ovary, uterus and placenta for rpST-treated and control pigs (Table 1; Fig. 4B).

PAG mRNA

The mRNA for PAG was detected as a 1.4 kb mRNA within the porcine placenta. When analyzed within

Table 1

<table>
<thead>
<tr>
<th>Tissue</th>
<th>IGF-I</th>
<th>IGF-II</th>
<th>IGFBP-2</th>
<th>PAG</th>
</tr>
</thead>
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<tr>
<td></td>
<td>rpST</td>
<td>SAL</td>
<td>S.E.</td>
<td>rpST</td>
</tr>
<tr>
<td>Liver</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
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<td>n=7</td>
<td>n=7</td>
<td>n=5</td>
<td>n=5</td>
</tr>
<tr>
<td>Ovary</td>
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<td>0.6</td>
<td>0.3</td>
<td>0.3</td>
</tr>
<tr>
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<td>n=8</td>
<td>n=5</td>
<td>n=5</td>
<td>n=5</td>
</tr>
<tr>
<td>Placental uterus</td>
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<td>0.6</td>
<td>0.1</td>
<td>0.6</td>
</tr>
<tr>
<td>n=8</td>
<td>n=9</td>
<td>n=5</td>
<td>n=5</td>
<td>n=5</td>
</tr>
<tr>
<td>Non-placental uterus</td>
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<td>0.2</td>
<td>0.1</td>
<td>0.2</td>
</tr>
<tr>
<td>n=8</td>
<td>n=8</td>
<td>n=5</td>
<td>n=5</td>
<td>n=5</td>
</tr>
<tr>
<td>Placenta</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.3</td>
</tr>
<tr>
<td></td>
<td>n=5</td>
<td>n=5</td>
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<td>n=5</td>
</tr>
</tbody>
</table>

**P<0.01 versus control.
reproductive tissue and liver, the PAG mRNA was also detected in placental uterus and endometrium (collected adjacent to the placenta; Fig. 5A). The presence of PAG mRNA in the placental uterus and endometrium suggested that residual placental tissue contaminates uterine tissue when the placenta is manually stripped from the uterus. The concentration of PAG mRNA was decreased in placenta (Table 1; Fig. 5B; \( P < 0.01 \)) for rpST-treated compared with control pigs.

Discussion

Administration of rpST to pregnant pigs increased maternal blood IGF-I and the weight of fetuses and placentas (Sterle et al. 1995). Maternal liver responded to rpST with an increased IGF-I mRNA concentration (Fig. 2). The change in IGF-I gene expression agrees with the increase in blood IGF-I that occurs after rpST administration (Buonomo et al. 1995, Sterle et al. 1995). Similar changes in IGF-I gene expression were not observed...
within reproductive tissues. The difference may reflect the relative concentrations of somatotropin receptor within liver compared with reproductive tissues (Fig. 1). The small concentration of somatotropin receptor in reproductive tissues may preclude a local response to rpST. Our inability to detect differences in IGF-I mRNA concentration within reproductive tissues in response to somatotropin treatment agrees with data from laboratory and ruminant species, where somatotropin failed to increase uterine IGF-I (Mathews et al. 1989, Kirby et al. 1996). A uterine IGF-I response to somatotropin was found in the ovariectomized, hypophysectomized rat but the response was small compared with the IGF-I response to estradiol (Murphy & Ghalary 1990). Skeletal muscle also had an undetectable IGF-I response to somatotropin (Coleman et al. 1994), perhaps because somatotropin receptors are not highly expressed in muscle (Florini et al. 1996). One study of pregnant pigs at similar stages of gestation showed increased IGF-I mRNA in the endometrium after rpST treatment (Kelley et al. 1995). Our study failed to confirm this previous observation, perhaps because we tested whole uterus, where the bulk of IGF-I mRNA arises from the myometrium (Song et al. 1996). Our conclusion is that somatotropin is not a dominant regulator of total uterine IGF-I mRNA concentration in pregnant pigs.

A third replicate was included for the IGF-I mRNA analyses. This was done because a definitive answer for the IGF-I response was not obtained after our analyses of replicates 1 and 2. The third replicate included surgically induced uterine crowding. The uterine IGF-I mRNA concentration in pigs from replicate 3 was greater than those of pigs in replicates 1 and 2. This was only observed for uterine tissues (i.e. ovarian tissues were similar for IGF-I mRNA concentration across replicates). The induction of uterine IGF-I mRNA in response to a crowded uterus may stimulate uterine or placental growth and development. The present experiment was not designed with a contemporary control for uterine crowding within replicate 3. The response of IGF-I mRNA concentration to a crowded uterus will, therefore, require additional studies.

The failure of a local (uterine) response to rpST demonstrates that enhanced fetal growth in rpST-treated pigs is not associated with increased IGF expression within the reproductive tract. Other mechanisms, therefore, must lead to increased fetal growth in rpST-treated gilts. One possibility is that increased IGF-I from the maternal liver acts as an endocrine growth factor for greater fetal growth and development. In previous studies, maternal IGF-I concentrations were correlated with the size of the fetus and birth weight (Hall et al. 1986, Gluckman & Barry 1988, Lassarre et al. 1991, Gluckman et al. 1992). The greater maternal IGF-I in rpST-treated gilts may act through similar mechanisms. A second possibility is that rpST causes nutrient partitioning that favors fetal development. Injection of pigs with rpST increased plasma glucose and free fatty acid concentrations (Gopinath & Etherton 1989, Dunshea et al. 1992, Buonomo et al. 1995, Hansen et al. 1997). The greater blood nutrient amounts may act as substrates for fetal and placental growth (Liu et al. 1994). Endocrine IGF-I and nutrient partitioning are only two of several mechanisms that may lead to enhanced...
fetal development. We did not examine changes in the expression of other fetal or placental growth factors in response to rpST. Furthermore, somatotropin receptors are widely distributed within maternal and fetal tissues. Therefore, the effects of rpST may have been mediated directly at multiple sites.

An unexpected observation from this study was the unique distribution of IGF-I, IGF-II and IGFBP-2 mRNA within the pregnant uterus. The IGF-I and IGFBP-2 gene expression within the uterus adjacent to the placenta (placental uterus) was different from the IGF-I and IGFBP-2 gene expression within the uterus adjacent to the necrotic tip, where two neighboring placentas meet (non-placental uterus). The concentration of IGFBP-2 mRNA was increased several-fold in placental uterus compared with non-placental uterus (Fig. 4A). The presence of IGFBP-2 mRNA in placental uterus was not caused by contaminating placental tissue, because little IGFBP-2 mRNA was detected within placenta (Fig. 4A). Not all genes were up-regulated in placental uterus, however, because IGF-I mRNA concentration was lower in placental compared with non-placental locations (Fig. 2). The intense expression of IGFBP-2 in the region of the placental uterus suggests a specific induction of IGFBP-2 by molecules produced by the placenta. One hypothesis, which needs to be tested, is that the intense expression of IGF-II within the placenta (Fig. 3A) induces the expression of IGFBP-2 by the uterine endometrium. A similar relationship exists in the ovine uterus, in which placental cells expressing IGF-II are closely apposed by endometrial cells expressing IGFBP-2 (Delhanty & Han 1993, Reynolds et al. 1997). The IGFBP-2 may act as a barrier to the movement of placental IGF-II from the pregnant uterus. In the non-placental uterus, IGFBP-2 may not be expressed because of inadequate placental signal from the necrotic tip.

A second unexpected finding was the relative expression of IGF-I in placental compared with non-placental uterus. We detected a decreased IGF-I mRNA concentration in the placental uterus compared with the non-placental uterus. The difference in IGF-I mRNA between the two uterine locations probably reflects the local influence of the conceptus. In the pregnant porcine and ovine uterus, IGF-I mRNA decreases during pregnancy (Song et al. 1996, Reynolds et al. 1997). The placenta is an active site of IGF-II mRNA synthesis in both pig (Simmen et al. 1992) and sheep (Reynolds et al. 1997), and IGF-II synthesis by the placenta may inhibit the synthesis of IGF-I by the uterus through a local IGF negative feedback loop. Infusion of IGF-I into fetal sheep caused a decrease in IGF-I mRNA within fetal liver (Kind et al. 1996). It is possible, therefore, that the synthesis of IGF-I within a tissue is partially dependent on circulating concentrations of IGFs. Within the pregnant uterus, IGF-II synthesis by the placenta may feedback negatively on IGF-I synthesis by the uterus. The absence of IGF-II synthesis within the necrotic tip may lead to greater IGF-I mRNA synthesis in adjacent uterine locations.

PAG was measured because it is a major placental protein that could be used as a marker for chorionic tissue. Our initial concern was that the presence of IGF-II mRNA within placental uterus was caused by residual placental tissue that was not removed by manual stripping of the placenta from the uterus. Indeed, we detected PAG mRNA in our placental uterus samples and believe that some of the IGF-II mRNA in placental uterus arises from placental contamination. Manually stripping placenta from pregnant pig uterus (a routine technique in studies of pregnant pig endometrium; e.g. Simmen et al. 1992) does not necessarily remove placental tissues that are deeply interdigitated within the uterus. Studies of pregnant endometrium should be interpreted with caution, therefore, when placentas are manually stripped and the studied mRNA (e.g. IGF-II) is also expressed by placenta.

As expected, mRNA for PAG was abundant in placenta. An unexpected finding, however, was that rpST decreased PAG mRNA concentration. Although PAGs are intensely expressed within the placentas of sheep (Xie et al. 1996), cattle (Xie et al. 1994) and pigs (Szafranska et al. 1995), a specific function for PAG has not been identified. The reason for somatotropin-induced PAG down-regulation is also unknown. The response may be secondary to structural changes in the placenta or may reflect rpST- or IGF-I-dependent aspects of PAG physiology. Although the observation was impossible to measure, we experienced greater difficulty separating uterus from placenta in rpST-treated pigs. Samples for histological analyses were not collected in the present experiment. Subsequent histological studies of rpST-treated pregnant pigs, however, suggest greater attachment by more extensive interdigitation of the placenta and the maternal epithelium (J A Sterle, unpublished observations). The importance of decreased PAG mRNA concentration in the process is not known.

In conclusion, rpST did not cause a change in IGF-I, IGF-II or IGFBP-2 mRNA concentration within the pregnant uterus of the pig. Changes in IGF-I mRNA concentration were observed in maternal liver, where greatest somatotropin receptor mRNA concentration was detected. The pregnant pig uterus demonstrated regions of gene expression where placental and non-placental uterus have distinctly different IGF-I and IGFBP-2 mRNA concentrations. Increased fetal growth may result from a somatotropin- or IGF-I-mediated endocrine mechanism that may stimulate fetal growth or change nutrient availability to the fetus. Other mechanisms are also possible, including altered expression of other fetal or placental growth factors. The importance of each possibility to fetal growth will require further study.
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