Developmental Programming: Prenatal Testosterone Excess and Insulin Signaling Disruptions in Female Sheep

Chunxia Lu, Rodolfo C. Cardoso, Muraly Puttabyatappa, and Vasantha Padmanabhan

Department of Pediatrics, University of Michigan, Ann Arbor, Michigan

ABSTRACT

Women with polycystic ovary syndrome often manifest insulin resistance. Using a sheep model of polycystic ovary syndrome-like phenotype, we explored the contribution of androgen and insulin in programming and maintaining disruptions in insulin signaling in metabolic tissues. Phosphorylation of AKT, ERK, GSK3beta, mTOR, and p70S6K was examined in the liver, muscle, and adipose tissue of control and prenatal testosterone (T)-, prenatal T plus androgen receptor antagonist (flutamide)-, and prenatal T plus insulin sensitizer (rosiglitazone)-treated fetuses as well as 2-yr-old females. Insulin-stimulated phospho (p)-AKT was evaluated in control and prenatal T-, prenatal T plus postnatal flutamide-, and prenatal T plus postnatal rosiglitazone-treated females at 3 yr of age. GLUT4 expression was evaluated in the muscle at all time points. Prenatal T treatment increased mTOR, p-p70S6K, and p-GSK3beta levels in the fetal liver with both androgen antagonist and insulin sensitizer preventing the mTOR increase. Both interventions had partial effect in preventing the increase in p-GSK3beta. In the fetal muscle, prenatal T excess decreased p-GSK3beta and GLUT4. The decrease in muscle p-GSK3beta was partially prevented by insulin sensitizer cotreatment. Both interventions partially prevented the decrease in GLUT4. Prenatal T treatment had no effect on basal expression of any of the markers in 2-yr-old females. At 3 yr of age, prenatal T treatment prevented the insulin-stimulated increase in p-AKT in liver and muscle, but not in adipose tissue, and neither postnatal intervention restored p-AKT response to insulin stimulation. Our findings provide evidence that prenatal exposure to excess testosterone (T) leads to both androgens and insulin may be involved in the programming of these metabolic disruptions.

androgen antagonist, insulin resistance, insulin sensitizer, PCOS

INTRODUCTION

Polycystic ovary syndrome (PCOS) is one of the most common infertility disorders in women of reproductive age [1]. The diagnostic features of PCOS, as defined by the Rotterdam consensus criteria [2, 3] and accepted at an evidence-based conference at the National Institutes of Health [4, 5], include ovulatory and menstrual dysfunctions, hyperandrogenism, and polycystic ovaries. While these are the cardinal features of PCOS, insulin resistance is the main metabolic abnormality observed in the majority of these patients [6, 7]. The mechanisms leading to the development and maintenance of hyperandrogenemia and insulin resistance and the extent to which insulin and androgens contribute to this metabolic perturbation have been the topic of intense investigation. Studies addressing insulin resistance in women with PCOS show that insulin sensitizers reduce hyperandrogenism [8] and attenuation of androgen actions improves insulin sensitivity [9], pointing to a reciprocal role of androgens and insulin in the maintenance of hyperandrogenism and insulin resistance in women with PCOS.

Because the diagnosis of PCOS is often made only after the establishment of reproductive competence, early perturbations leading to the development of insulin resistance in women with PCOS are unknown. While there is evidence for a genetic basis for PCOS [10], prenatal origins of this syndrome are supported by reports that children born small for their gestational age [11] for PCOS [10], prenatal origins of this syndrome are supported by reports that children born small for their gestational age [11] have an increased propensity to develop PCOS. Animal models of PCOS provide unique opportunities to identify early disruptions and investigate the developmental progression of the PCOS phenotype. Studies in several animal models have shown that prenatal exposure to excess testosterone (T) leads to reproductive and metabolic dysfunctions during adult life that resemble those seen in women with PCOS [12-15]. More specifically, prenatal T treatment in sheep, a precocial large animal model, results in oligo-/anovulation, functional hyperandrogenism, polycystic ovarian morphology, and peripheral insulin resistance associated with altered insulin signaling in metabolic tissues [14, 16]. While the adult phenotype of this sheep model has been extensively characterized, early perturbations that lead to these defects remain unexplored. Investigation of the developmental changes in animal models of PCOS phenotype and their reversal with interventions may help identify preventive strategies in humans.

The mechanisms by which prenatal T treatment leads to PCOS phenotype in the female sheep are unclear. Similar to the hormonal changes observed in PCOS women during pregnancy [17, 18], gestational T treatment in sheep increases not only maternal circulating concentrations of T but also insulin levels (Fig. 1A) [19]. The relative contribution of sex steroids and insulin in programming insulin resistance in the offspring as well as the early tissue-specific disruptions in metabolic tissues induced by prenatal T excess are unknown. Pharmacological interventions with steroid receptor antagonists or insulin sensitizers during pregnancy and after establishment of the PCOS phenotype can provide insights into the mechanisms and pathways involved in the development and maintenance of insulin resistance.

The objectives of this study are twofold: 1) to determine the ontogeny of tissue-specific disruptions in insulin signaling in prenatal T-treated sheep (Fig. 1A) and 2) to utilize pharma-
cological agents during pregnancy and after the establishment of the PCOS-like phenotype to determine the relative contributions of sex steroids and insulin in the development and/or maintenance of the pathology. We hypothesize that prenatal T excess, either by its androgenic action or via perturbed maternal insulin homeostasis, alters the activity level of key members of the insulin-signaling pathway in a tissue- and development-specific manner.

MATERIALS AND METHODS

Breeding and Maintenance

All animal procedures used in these studies were approved by the Animal Care and Use Committee of the University of Michigan and are consistent with the National Institutes of Health Guide for Use and Care of Animals. All studies were conducted at the University of Michigan Research Facility using multiparous Suffolk sheep. Details regarding animal maintenance, breeding, and lambing have been described previously [20]. In brief, starting 2–3 wk before and continuing until the time of breeding, ewes were group fed daily with 0.5 kg shelled corn and 1.0–1.5 kg alfalfa hay/ewe. Ewes were mated to raddled Suffolk rams, and mating was confirmed by the presence of paint on the back of the females. Once mated, females were randomly assigned to the different treatment groups, housed under a natural photoperiod in the pasture, and group fed with a daily maintenance diet of 1.25 kg alfalfa/brome mix hay/ewe. After weaning at ~8 wk of age, female lambs were maintained outdoors and fed a pelleted diet (Shur-Gain; Nutreco Canada Inc., Guelph, ON) consisting of 3.6 MCal/kg digestible energy and 18% crude protein. When twins or triplets were involved, only one female offspring from each mother was randomly selected.

Experimental Design

**Study 1:** early disruptions in insulin-signaling pathway induced by prenatal T excess. To address early perturbations induced by exposure to excess T and investigate the relative contributions of androgen and insulin in mediating these effects, four groups of female fetuses were studied at Gestational Day 90 (GD90): 1) control (C), 2) gestational T treatment (T), 3)
gestational cotreatment with T and flutamide (TF), an androgen antagonist, and 4) gestational cotreatment with T and rosiglitazone (TR), an insulin sensitizer. Sample size for all markers in liver and adipose tissue were C = 9, T = 6, TF = 6, and TR = 6. Due to the high variability in expression patterns, sample sizes were increased by adding animals from a second cohort (this cohort did not have a TR group) to arrive at a final sample size of C = 16, T = 14, TF = 15, and TR = 6. Gestational T treatment consisted of twice weekly intramuscular administration of 100 mg T propionate (1.2 mg/kg; Sigma-Aldrich, St. Louis, MO) suspended in 2 ml corn oil from GD30 to GD90. Flutamide (10 mg/kg) or rosiglitazone (30 mg/kg) was packed into capsules (Profine Capsule Filling System; Torpac, Fairfield, NJ), individually weighed, and administered daily. The daily dose of flutamide administered to pregnant ewes in this study has been previously shown to block the effects of both exogenous and endogenous androgens on phenotypic virilization in males and prenatal T-treated female sheep [21]. Rosiglitazone (~0.11 mg/kg) (Avandia; GlaxoSmithKline, Research Triangle Park, NC) was also administered orally on a daily basis. The treatment dose of rosiglitazone is within the range used to treat women with PCOS [22, 23]. Basal expression of phospho (p)-Akt, in the liver, skeletal muscle, and subcutaneous and visceral adipose tissue and GLUT4 in the muscle were evaluated in C, T, T+F, and T+R females. In addition, changes in p-Akt were also measured following in vitro insulin stimulation in all tissues. Figure 1C illustrates the sequence of experimental procedures in study 3.

**Tissue Harvesting and Processing**

All animals were fasted for 48 hr prior to tissue collection. For studies 1 and 2, liver (obtained from the tip of the left lobe), skeletal muscle (obtained from the vastus lateralis), and visceral adipose tissue samples (obtained from the mesenteric fat surrounding the ventral sac of the rumen) were collected from GD90 fetuses (study 1) and 2-yr-old females (study 2). Tissues were flash-frozen and stored at −80°C until processed for studying changes in basal expression of members of the insulin-signaling cascade. For study 3, the protocol for assessment of insulin-stimulation of Akt in metabolic tissues was similar to the procedures described previously [32–34]. Briefly, liver, muscle, and abdominal and subcutaneous adipose tissues were minced, pre-incubated with saline at 37°C for 10 min, and then incubated in 1 ml saline containing 100 nM insulin (Sigma-Aldrich) at 37°C for 2 or 5 min. The tissue designated as the treatment control was incubated similarly but without insulin. After treatment, the samples were snap-frozen in dry ice and stored at −80°C until further processing.

**Western Blot Analysis**

Tissue samples were homogenized in radio-immunoprecipitation assay buffer (Pierce RIPA Buffer; Thermo Scientific, Rockford, IL) containing protease inhibitors (Complete Mini; Roche Diagnostics, Indianapolis, IN) and phosphatase inhibitors (PhosSTOP; Roche Diagnostics). Insulin homogenates were centrifuged at 10,000 x g for 15 min at 4°C and the whole-cell protein extract was used for the analysis. Equal amounts of protein (20–50 μg) were resolved on SDS-PAGE and transferred into a nitrocellulose membrane. Levels of phosphorylated and total forms of each protein, as well as the corresponding loadings in different groups, were determined using ImageJ software. For studies 1 and 2, the ratio of band densities for phospho-/total protein or total protein/loading control from the treatment groups were expressed relative to the controls from the respective autoriograms, which were set at 100%. For study 3, samples at 0, 2 and 5 min time points from any given animal were blotted at the same time, and the ratio of phospho-/total protein was calculated as the insulin-stimulation effect was expressed relative to time point 0, which was set at 100%. In addition, samples at time point 0 from all treatment groups were run and blotted together to evaluate changes in the basal levels of p-Akt. The ratio of p-Akt/Akt at time point 0 of the treatment groups were expressed relative to the controls, which was set at 100%. Anti-p-AktSer473, Akt, mTORSer2448, mTOR, p-GSK3βSer9, p-GSK3αSer9, p-ERKThr202/Tyr204, p70S6K, p-p70S6KThr421/Ser424, glyceraldehyde-3-phosphate dehydrogenase (GAPDH), and β-actin antibodies were purchased from Cell Signaling Technology (Cell Signaling Technology, Inc., Beverly, MA). Anti-Glut4 was purchased from Abcam (Cambridge, MA) (Supplemental Table S1, available online at www.biolreprod.org). Because of technical issues in detecting β-actin in some series, GAPDH was replaced as a loading control. The specificity of the antibodies was confirmed by visualization of protein bands of the correct size.
RESULTS

Study 1: Early Disruptions in Insulin-Signaling Pathway Induced by Prenatal T Excess

Figure 2 summarizes the results of AKT and GSK3β protein expression. At GD90, no differences were observed among treatment groups for the ratio of p-AKT/AKT in any of the three tissues studied (top panel). In contrast, prenatal T treatment altered the ratio of p-GSK3β/GSK3β in a tissue-specific manner and was manifested as an increase in the liver, a decrease in the muscle, and no change in the adipose tissue (bottom panel). This increase in liver p-GSK3β/GSK3β was prevented partly by both interventions (TF and TR not different from C or T). Similarly, the decrease in muscle GSK3β/GSK3β was also partially prevented in the TR group. Consistent with the decreased p-GSK3β in the muscle, a trend (P = 0.07) for reduced GLUT4 expression was also observed in prenatal T-treated fetuses. Both interventions partially prevented this decrease in GLUT4 expression (Fig. 2).

Investigation of the upstream regulators of GSK3β, namely p-mTOR and p-p70S6K, found that the levels of p-mTOR in the liver and muscle did not differ significantly between C and T fetuses (Fig. 3, left column, top and middle rows); however, total mTOR was increased in the liver, suggestive of increased mTOR activity. Consistent with this, the levels of p-p70S6K (while p-p85S6K was not detectable in our samples), a downstream molecule of the mTOR pathway and an upstream regulator of GSK3β, was significantly increased in the liver of prenatal T-treated fetuses. This increase in p-p70S6K was also observed in the TR group but partially prevented in the TF group (Fig. 3, middle column, top row). In contrast, gestational T treatment did not alter the levels of p-p70S6K in the muscle (Fig. 3, middle column, middle row). In the adipose tissue, although p-mTOR was significantly increased in T fetuses, the expression of total mTOR was significantly lower. In addition, p-p70S6K did not differ between T and C females (Fig. 3, bottom row). The increase in p-mTOR levels in adipose tissue was prevented by cotreatment with either androgen antagonist.

FIG. 2. Changes in expression levels of total and phosphorylated AKT and GSK3β in the liver (left column), muscle (middle column), and adipose tissue (right column) as well as GLUT4 expression in the muscle from GD90 fetuses. Representative Western blots (n = 5/group) are shown along with mean ± SEM of the p-AKT/AKT, AKT/actin, p-GSK3β/GSK3β, GSK3β/actin, and GLUT4/GAPDH ratios. C, control; T, prenatal T; TF, prenatal T plus flutamide; and TR, prenatal T plus rosiglitazone. Different superscripts indicate statistically significant changes; b#, P = 0.07, C versus T. Note statistical outcomes of analyses of protein expression patterns in the muscle before and after addition of the second cohort are similar (composite shown).
or insulin sensitizer. Gestational T treatment did not alter the levels of p-ERK in all the three tissues studied (Fig. 3, right column).

Study 2: Contributions of Androgen and Insulin in Programming Adult Disruptions in Members of the Insulin-Signaling Pathway in Prenatal T-Treated Sheep

As opposed to the changes evidenced in GD90 fetuses, prenatal T treatment did not affect the basal levels of total or phosphorylated AKT, GSK3β, or ERK (Fig. 4) in the liver, muscle, and adipose tissue at 2 yr of age. A decrease in p-AKT and p-ERK levels in the liver (Fig. 4, top row, left and right columns) and a tendency for an increase (P = 0.07) in p-GSK3β expression in the adipose tissue (Fig. 4, bottom row, left column) were evident in the TR group compared to controls. In addition, no changes were observed in basal GLUT4 expression in the muscle across all experimental groups (data not shown).

Study 3: Impact of Prenatal T Excess on AKT Response to Insulin Stimulation

The ratio of p-AKT/AKT in metabolic tissues did not differ between groups prior to insulin stimulation (Fig. 5). In addition, no changes were observed in basal GLUT4 expression in the muscle among treatment groups (data not shown). Prenatal T treatment decreased the p-AKT/AKT ratio following insulin stimulation in the muscle and liver but not in visceral and subcutaneous adipose tissues (Fig. 5). Levels of p-AKT increased significantly (~150%) in the liver of C females after 2 min of insulin stimulation (Fig. 5, top panel, left). This increase, however, was not evident in the liver obtained from prenatal T-treated females. Similarly, p-AKT levels increased after 2 (~240%) and 5 min (~220%) of insulin stimulation in the skeletal muscle of C animals (Fig. 5, top panel, right), but this increase was not evident in the muscle of prenatal T-treated females. Postnatal androgen antagonist or insulin sensitizer treatment failed to restore normal insulin stimulation of AKT phosphorylation in the muscle and liver. Insulin stimulated the phosphorylation of AKT in visceral and subcutaneous adipose tissue to a similar degree in both C and T animals (Fig. 5, bottom panel). An increase in the insulin-stimulated p-AKT level was observed in both the visceral and subcutaneous adipose depots of T+F females and only in the subcutaneous compartment of T+F when compared to C females.
DISCUSSION

Findings from this study extend our previous observations on the mRNA expression level [16] to a functional level and indicate that prenatal exposure to excess T disrupts the normal activation (phosphorylation) of proteins involved in the regulation of glucose homeostasis in the female fetus, culminating in tissue-specific changes in the insulin-signaling cascade during adulthood. Changes in the phosphoprotein level of key insulin-signaling molecules in the female fetus are in the direction of increased insulin sensitivity in the liver, decreased sensitivity in the muscle, and unaltered sensitivity in the adipose tissue. The adult metabolic consequences from prenatal exposure to T excess were manifested as reduced insulin-stimulated levels of p-AKT in both the liver and muscle, and unaltered p-AKT levels in visceral and subcutaneous adipose tissues. The direction of these changes is consistent with our previous work on the mRNA level [16], showing that the liver and skeletal muscle, but not the adipose tissue, are likely insulin resistant in adult sheep prenatally exposed to T excess. These findings and the involvement of androgenic and metabolic pathways in mediating these alterations are discussed below.

Androgenic Versus Metabolic Programming of Insulin Signaling in the Liver

The liver plays a key role in maintaining glucose levels in the blood [35, 36]. The increased total mTOR coupled with augmented p-p70S6K, a downstream molecule of the mTOR pathway, are suggestive of the activation of the mTOR pathway by gestational T excess in female sheep during fetal life. Consistent with this, phosphorylation at Ser-9 of GSK3β, a downstream target of p70S6K was also increased. Although Ser-9 phosphorylation can be regulated through both AKT and ERK pathways, the absence of changes in the phosphorylation status of AKT and ERK indicates that the increased GSK3β phosphorylation in prenatal T-treated fetuses is predominantly regulated through the mTOR pathway (Fig. 6, left panel).

Considering that the gestational T treatment extended through Day 90 of fetal life, changes observed in GD90 fetuses are likely to be the direct result of activation of the androgen receptor. Interactions between androgen and mTOR pathways have been demonstrated in prostate cancer cells with high levels of T stimulating mTOR activity [37, 38]. Androgens have also been shown to upregulate mTOR activity...
in the ovary [39]. The finding that the androgen antagonist flutamide prevented the increase in total mTOR also supports a predominantly androgenic mediation in the activation of the mTOR pathway (Fig. 6, left panel). The inability of insulin sensitizer cotreatment to prevent the prenatal T-induced increase in p-p70S6K in concert with partial reversal of p-GSK3β increase in the T group by insulin sensitizer cotreatment (TR group not different from C or T) is supportive of the potential for metabolic mediation independent of p70S6K.

In contrast to the activation of the mTOR pathway in the fetal liver, changes in basal phosphorylation of AKT, ERK, and GSK3β were not evident in 2-yr-old, prenatal T-treated sheep. However, a reduction in insulin-stimulated phosphorylation of AKT was evident in 3-yr-old females, supportive of the liver being insulin resistant at this developmental stage (later adulthood). The finding that postnatal androgen antagonist treatment failed to prevent this reduction suggests that the negative effect of gestational T on adult hepatic insulin sensitivity is likely independent of postnatal activation of the androgen receptor. The fact that rosiglitazone, a high-affinity ligand for the transcription factor peroxisome proliferator–activated receptor-γ (PPAR-γ) [40] that alters the expression of genes involved in glucose metabolism [41, 42], also failed to prevent this reduction may relate to the differential effects of rosiglitazone in different tissues [43]. Rosiglitazone has been shown to exacerbate hepatic insulin resistance associated with increased triglyceride and fatty acyl-CoA contents [44] and to reduce phosphorylation of AKT in human hepatocarcinoma cells [45].

Together, our results indicate that prenatal T treatment, at least partially via the androgenic pathway, alters insulin signaling in the fetal liver consistent with increased insulin sensitivity. The findings of decreased hepatic responsiveness to insulin in adult females are suggestive of organizational (permanent) effects of prenatal T excess on liver function. Developmental differences in the effect of prenatal T excess, such as enhanced insulin sensitivity in fetal liver versus reduced sensitivity in adult life, are consistent with compensatory mechanisms being in place to prevent the development of pathology during early life and failing to do so during later life.

**Androgenic Versus Metabolic Programming of Insulin Signaling in the Muscle**

The decrease in phosphorylation of GSK3β at Ser-9 in the fetal muscle is suggestive of reduced insulin sensitivity in this early developmental stage [46, 47]. Consistent with this
premise, prenatal T excess was found to decrease the expression of GLUT4 in the fetal muscle, where it is primarily expressed [48]. Previous studies have found GLUT4 levels to be reflective of glucose uptake [49, 50]. The lack of changes in the mTOR, AKT, and ERK pathways together with the failure of androgen antagonist intervention in preventing the decrease in p-GSK3\(\beta\) suggests that GSK3\(\beta\) phosphorylation in the muscle during fetal life is likely regulated through other signaling pathways, such as WNT and protein kinase C [51, 52] (Fig. 6, middle panel). Further investigation is required to clarify this possibility.

Failure of insulin to stimulate p-AKT in the skeletal muscle from prenatal T-treated sheep at 3 yr of age is consistent with gestational T excess reducing insulin sensitivity in the muscle not only during fetal life but also during adulthood. The finding that postnatal treatment with insulin sensitizer did not restore AKT stimulation by insulin in the muscle of T females is in agreement with a previous study [53] that found rosiglitazone to improve glucose uptake without altering the insulin signaling in human skeletal muscle. These findings suggest a weak positive relationship between insulin activation of the IRS-1/PI3-kinase/AKT pathway in the muscle and peripheral insulin sensitivity. Thus, the previously reported [24] improvement in peripheral insulin sensitivity with rosiglitazone treatment in prenatal T-treated sheep may be independent of the activation of this pathway in the muscle. Moreover, observations that prenatal intervention with either androgen antagonist or insulin sensitizer failed to prevent the changes in insulin signaling in the skeletal muscle indicate that the effects of prenatal T are likely mediated by mechanisms independent of either androgen or insulin receptors, such as aromatization of T into estradiol.

Paradoxically, p-mTOR levels were increased in both liver and muscle tissues from T females (Fig. 3). Although the exact mechanism is unclear, this increase in p-mTOR may be due to a predominantly estrogenic milieu in T females (facilitated by aromatization of T into estradiol). Estrogen has been shown to upregulate the mTOR pathway in human breast cancer cell lines [54, 55]. Unfortunately, because we found no effects of flutamide treatment alone on peripheral insulin sensitivity in a prior study (Padmanabhan et al., unpublished results), this group was not generated with this cohort, precluding us from investigating direct effects of flutamide on metabolic tissues. Although a rosiglitazone-only group was generated as part of this investigation, due to the absence of its effect on peripheral insulin sensitivity (Padmanabhan et al., unpublished results), this group was not included in the extensive tissue-specific analyses.

Androgenic Versus Metabolic Programming of Insulin Signaling in the Adipose Tissue

Contrary to changes observed in the fetal liver and muscle, prenatal T excess did not alter the phosphorylation of GSK3\(\beta\) and p70S6K in the fetal adipose tissue (Fig. 6, right panel). These data suggest that the increase in mTOR phosphorylation compensates for the reduced total mTOR expression, culminating in lack of change in the net mTOR activity. These findings coupled with the lack of changes in insulin-stimulated levels of p-AKT in the visceral and subcutaneous adipose tissues in 3-yr-old females indicate that prenatal T treatment had no effect on the insulin sensitivity in the adipose tissue. In vitro studies with human subcutaneous adipocytes have also found no effects of chronic T treatment on IRS-1-associated PI3-kinase and its downstream AKT activities following...
insulin stimulation [56]. Whether changes in insulin sensitivity in the adipose tissue are evident at later ages remains to be determined.

Interestingly, postnatal rosiglitazone treatment augmented the p-AKT response to insulin-stimulation in the adipose tissue. This observation is in agreement with previous reports that rosiglitazone improves insulin sensitivity in the adipose tissue by activating the IRS-1/PI3-kinase/AKT pathway [57]. The trend for increased basal level of GSK3β in the adipose tissue of TR females at 2 yr of age is consistent with the positive effect of rosiglitazone on insulin sensitivity in the adipose tissue. Collectively, our observations indicate that prenatal T treatment does not alter insulin sensitivity in the adipose tissue in female sheep during the developmental stages investigated.

Relationship Between Tissue-Specific Changes in Insulin Signaling and Peripheral Insulin Resistance in Prenatal T-Treated Sheep

In addition to alterations in reproductive function [14], our previous longitudinal studies have identified marked fluctuations in insulin sensitivity throughout development in prenatal T-treated female sheep [31, 58, 59]. More specifically, prenatal T-treated females manifest insulin resistance relative to controls during the infantile [58] and early juvenile life [31], a similar degree of insulin sensitivity as controls during the prepubertal and postpubertal periods when controls also become insulin resistant [59], and re-emergence of insulin resistance compared to controls during adulthood [31]. Assessment of peripheral insulin sensitivity at early developmental time points in a subset of females from study 2 confirmed a similar pattern of fluctuation, with T females manifesting insulin resistance at ~6 wk of age (which was prevented by rosiglitazone treatment) and normal insulin sensitivity at ~20 mo of age compared to C females [26]. While the experimental paradigm used here is the same as in our previous studies investigating peripheral glucose-insulin homeostasis, one limitation of the present study is the lack of assessment of peripheral insulin sensitivity in these animals just prior to tissue collection. Whether tissue-specific alterations precede the re-emergence of peripheral insulin sensitivity at later ages is unclear. Considering that the tissue specific studies in protein expression point to downstream effects on glycogen synthesis, a second limitation of the study is the lack of assessment of glycogen content. Because glycogen content in tissue can be affected by multiple factors, including fasting and storage for extended period [60], glycogen content could not be assessed in this study. In spite of these limitations, the current findings provide valuable insights on the potential mechanisms linking prenatal exposure to T excess and the development of tissue-specific insulin resistance in female sheep.

When comparing fetal versus adult responses (Fig. 7), findings in the liver of increased insulin sensitivity during fetal life in contrast to reduced sensitivity during adulthood support the notion that mechanisms are developed by the liver to compensate for the early insults. Such adaptive mechanisms appear to be organ-specific, having different trajectories for different organ systems and dependent on critical periods of differentiation. For instance, the direction of changes in insulin sensitivity in the muscle, namely reduced sensitivity in both fetal and adult females, differs from the trajectory seen in the liver. It is still possible that compensatory mechanisms similar to those seen in the liver during fetal life may have occurred in the muscle later during postnatal life. More detailed time course studies would be necessary to investigate this possibility.

Translational Relevance

In addition to the similarities in peripheral insulin resistance [14], our observations indicate that prenatal T treatment in sheep leads to tissue-specific changes in insulin signaling during adult life that parallel those reported in women with PCOS. Disruptions in skeletal muscle in prenatal T-treated sheep are suggestive of reduced insulin sensitivity and are consistent with those seen in women with PCOS [61]. However, similar to the present findings in prenatal T-treated sheep (study 2), such disruptions appear not to be manifested as changes in the basal AKT activity in women with PCOS [62]. Attenuated p-AKT response in skeletal muscle collected from prenatal T-treated sheep (study 3) and PCOS women [62] becomes evident only upon insulin stimulation. At the adipose tissue level, the finding of normal responsiveness to insulin in prenatal T-treated females is also consistent with observations in PCOS women of no differences in the abundance of several insulin-signaling molecules at basal levels [63] and after insulin stimulation of isolated adipocytes [7, 64]. Unfortunately, to our
knowledge, studies investigating insulin signaling in the liver of PCOS women are not available for comparison. For obvious reasons, tissue-specific changes occurring during fetal or early postnatal life that contribute to the development of insulin resistance in women with PCOS are difficult to determine. While extrapolation of findings in this sheep model to human pathology should be done cautiously, observations that PCOS may have developmental origins [11, 65–68] and prenatal T treatment recapitulates the metabolic phenotype of PCOS in female sheep [14, 31] suggest that animal models, such as prenatally T-treated sheep, may provide a means to identify early targets for the development of intervention strategies.

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