Review Article

The Role of Metformin in Metabolic Disturbances during Pregnancy: Polycystic Ovary Syndrome and Gestational Diabetes Mellitus

Joselyn Rojas, Mervin Chávez-Castillo, and Valmore Bermúdez

Endocrine and Metabolic Diseases Research Center, School of Medicine, University of Zulia, 20th Avenue, Maracaibo 4004, Venezuela

Correspondence should be addressed to Joselyn Rojas; rojas.joselyn@gmail.com

Received 21 June 2014; Revised 7 November 2014; Accepted 19 November 2014; Published 8 December 2014

Abstract

Maintenance of gestation implicates complex function of multiple endocrine mechanisms, and disruptions of the global metabolic environment prompt profound consequences on fetomaternal well-being during pregnancy and postpartum. Polycystic Ovary Syndrome (PCOS) and gestational diabetes mellitus (GDM) are very frequent conditions which increase risk for pregnancy complications, including early pregnancy loss, pregnancy-induced hypertensive disorders, and preterm labor, among many others. Insulin resistance (IR) plays a pivotal role in the pathogenesis of both PCOS and GDM, representing an important therapeutic target, with metformin being the most widely prescribed insulin-sensitizing antidiabetic drug. Although traditional views neglect use of oral antidiabetic agents during pregnancy, increasing evidence of safety during gestation has led to metformin now being recognized as a valuable tool in prevention of IR-related pregnancy complications and management of GDM. Metformin has been demonstrated to reduce rates of early pregnancy loss and onset of GDM in women with PCOS, and it appears to offer better metabolic control than insulin and other oral antidiabetic drugs during pregnancy. This review aims to summarize key aspects of current evidence concerning molecular and epidemiological knowledge on metformin use during pregnancy in the setting of PCOS and GDM.

1. Introduction

Infertility currently affects approximately 48.5 million of women aged 20–44 years around the world [1], with severe implications in their physical and mental well-being [2]. Female fertility entails a complex array of endocrine mechanisms surrounding the integrity of the hypothalamus-pituitary-ovary (HPO) axis, which are especially important in maintenance of a healthy pregnancy, particularly due to the demands of the growing fetus [3]. Many conditions may disrupt this environment, and Polycystic Ovary Syndrome (PCOS)—an endocrine-metabolic disease that encompasses multiple hormonal alterations related to female infertility—stands out mainly due to its high prevalence, affecting 6-7% of women aged 12–45 years [4], with a worrisome 70% of women estimated to remain undiagnosed [5].

The hallmarks of this gynecoendocrine disease are disruption of ovarian steroidogenesis, giving rise to hyperandrogenemia and insulin resistance (IR) [6]. A complex IR-hyperinsulinemia-hyperandrogenemia cycle involved in the endocrine disruptions in PCOS [7] leads not only to the typical clinical picture of PCOS—featuring oligoanovulation and hyperandrogenic manifestations—but also to diverse cardiometabolic comorbidities, such as impaired glucose tolerance [8], dyslipidemia [9], hypertension [10], central obesity [11,12], accelerated atherosclerosis [13], and metabolic syndrome [14], which can appear as a myriad of distinct metabolic phenotypes [15] including mild, moderate, and severe forms of PCOS.

Insulin resistance is an important component in the etiopathogenesis of PCOS, being associated with obesity, acanthosis nigricans, hirsutism [16], and early pregnancy loss [17] in these women. In addition, utilizing the HOMA-IR index as a surrogate for IR quantification, Huidobro et al. [18] reported this condition to be associated with gestational diabetes mellitus (GMD), which supports the notion that this pregnancy-related metabolic disorder may be part of...
2. Insulin Resistance as the Key Endocrine Disruption in Polycystic Ovary Syndrome

The etiology of PCOS is complex and multifactorial, including several endocrine disturbances, such as (a) increased pulsatile secretion of gonadotropin-releasing hormone (GnRH) and luteinizing hormone (LH), prompting theca cell hyperstimulation and androgen hypersecretion [27]; (b) nonselection of a dominant ovarian follicle, mediated by intrinsic and extrinsic ovary factors, with follicular cells hyperplasia [28]; (c) genetic predisposition to hyperandrogenemia, linked to abnormal in utero androgenic exposure [29]; and (d) genetic predisposition to hyperinsulinemia, also linked to prenatal androgen exposure and pancreatic β-cell dysfunction [30]. Although it is difficult to establish the relative importance or chronology of these and subsequent alterations, PCOS is characterized by an IR-hyperinsulinemia-hyperandrogenemia positive feedback circuit (Figure 1), where the latter component determines the majority of clinical manifestations and the diagnostic criteria for this condition (Table 1). Moreover, obesity is a very common feature in females with PCOS, which appears to magnify all previous pathophysiologic mechanisms [7].

Insulin resistance, defined as a decrease in cellular responsiveness to insulin signaling [31], triggers increased insulin secretion, a phenomenon termed "compensatory hyperinsulinemia" [32]. Although this mechanism attempts to maintain lipid, carbohydrate, and protein metabolism homeostasis, it contributes to multiple aggregate consequences, such as the cardiovascular PCOS comorbidities [33], and favors hyperandrogenemia through various pathways. In this respect, disruption of the HPO is particularly relevant: insulin has been shown to elevate GnRH and LH secretion both dose- and time-dependently [34, 35], potentially mediated through the MAPK pathway [36]. This results in increased frequency and amplitude of GnRH and LH pulse secretion, with increased LH/FSH ratio, potentiating ovarian steroidogenic alterations [6]. Other features frequently found in women with PCOS act in synergy with insulin towards enhancing LH release, including hyperleptinemia via AgRP/NPY neural pathways and kisspeptidergic signaling [37], and decreased opioidergic tone, which appears to sensitize pituitary LH-secreting cells to GnRH signaling [38]. Hyperinsulinemia has also been associated with diminished Sex Hormone-Binding Globulin (SHBG) levels, although insulin appears to be unable to directly inhibit shbg expression; instead, this effect depends on hyperglycemia-mediated Hepatocyte Nuclear Factor 4-α downregulation [39]. Lower SHBG synthesis results in increased sex hormone availability, exacerbating androgenic signaling [40].

Lastly, PCOS is also characterized by selective IR in ovarian tissue, wherein mitogenic pathways are favored while...
Figure 1: The insulin resistance-hyperinsulinemia-hyperandrogenemia cycle in Polycystic Ovary Syndrome. PCOS is dominated by three major endocrine disruptions: insulin resistance, hyperinsulinemia, and hyperandrogenemia. Although it is difficult to establish which disturbance develops first in any given case, these components are interconnected by many reinforcing mechanisms, constituting a positive feedback cycle. Furthermore, obesity and chronic inflammatory states—present in both obese and lean women with PCOS—amplify pathophysiologic pathways linked to all elements in this triad. The cycle leads to the manifestations of PCOS and infertility, complications during pregnancy, and chronic cardiometabolic comorbidities.

metabolic signaling is absent, yielding follicular cell hyperplasia and potentiation of steroidogenesis [41]. Several theories surround this concept, including cAMP-dependent activation of PKA with subsequent activation of Steroidogenic Acute Regulatory (StAR) protein [42], increased PI3K/Akt activity via serine phosphorylation by a hypothetical kinase in theca cells [43], and inositol phosphoglycan signaling, which appears to deviate from insulin-dependent pathways aside from being activated by the insulin receptor itself [44]. At any rate, IR-hyperinsulinemia activity leads to hyperandrogenemia, which in turn induces pro-IR structural and functional modifications in key insulin target tissues, including decreased amount of more oxidative, insulin-sensitive type I muscle fibers, and increased amount of more glycolytic, less sensitive type II fibers [40], as well as elevated lipolysis in adipocytes, favoring free fatty acid- (FFA-) mediated IR [45], perpetuating the IR-hyperinsulinemia-hyperandrogenemia feedback [7].

Although physical activity and lower caloric intake are considered fundamental lifestyle interventions [46], insulin-sensitizing agents are also a hallmark of PCOS management, with metformin being the most frequently used molecule [47]. Metformin has been described to offer significant improvement of several parameters, including Body Mass Index (BMI), LH, androstenedione, testosterone [48], DHEAS, blood pressure [49], menstrual cyclicity, fasting insulin [50], IR, dyslipidemia, oxidative stress, endothelial dysfunction [51], and several inflammatory markers [52]. This biguanide has also been reported to improve other features such as anovulation rate and acne [53] as well as BMI and LH [54] in non-IR women with PCOS. Moreover, it appears to be beneficial in both obese and lean women with PCOS [53], which may explain the persistent benefits of metformin even with several different metabolotypes.

The subset of lean women with PCOS is particularly interesting. Although all PCOS phenotypes tend towards a more “apple-like” adipose distribution [55], lean subjects usually have less visceral fat [56]. Likewise, in these individuals, IR and hyperandrogenemia are predominantly related to low SHBG levels [57], with increased risk for elevated inflammation markers [58] and early vascular disease [59]. Although both lean and obese PCOS women tend to exhibit higher oxidative stress [60], they appear to behave differently regarding aging and risk of developing type 2 diabetes mellitus (DM2), which seems to be less frequent in lean women with PCOS [61]. Indeed, women who are able to maintain normal weight with aging appear to boast a healthier metabolic profile than those who do not [62]. These differences may influence the impact of metformin in each group [63]: whereas reproductive benefits are observed in both obese and lean PCOS women [64], metabolic advantages, such as lowering of proinsulin and insulin levels, are seen predominantly in the obese and overweight subset [65].

Other antidiabetic drugs have been evaluated to be applied in PCOS, particularly thiazolidinediones (TZD). Despite reports indicating these agents to be more effective than metformin at reducing IR in subjects with PCOS [66], their use remains less widespread, due to concerns of increased cardiovascular risk [67]. Indeed, despite significantly ameliorating IR, glucose homeostasis, hyperandrogenic ovarian response, and systemic inflammation [68, 69], TZD appear to induce several deleterious modifications in
cardiac tissue transcriptomes, including upregulation of metalloproteinases implicated in atheromatous plaque rupture, potassium channels required for action potential generation, and genes involved in sphingolipid and ceramide metabolism [70]. Beyond these molecular findings, the impact of TZD on cardiovascular risk is also reflected in epidemiologic findings, with a higher risk of congestive heart failure in prediabetic and diabetic subjects (RR = 1.72, 95% CI: 1.21–2.42, P = 0.002) [71].

3. Exacerbation of Physiologic Insulin Resistance as the Fundament of Gestational Diabetes Mellitus

Insulin resistance is a physiologic state during gestation, driven by several maternal hormones such as estrogen, progesterone, cortisol, and particularly human placental lactogen (hPL) [72]. Target cell modifications include defective tyrosine phosphorylation of the β subunit of the insulin receptor [73] and decreased expression of IRS-1 [74], whereas expression of the p85α subunit of phosphoinositol 3-kinase is increased, which interferes with heterodimeric conformation of this enzyme and thus prevents further insulin signaling [72]. Similarly, GLUT4 expression has been noted to be decreased in adipose tissue of pregnant females, significantly hindering insulin responsiveness [75]. Although the elevated serum levels of free fatty acids triggered by IR represent an important adaptive mechanism in order to increase the glucose offer for fetal metabolism, they also serve as a self-reinforcing pathway for IR (Figure 2) [76].

These pro-IR phenomena are counterbalanced by several pancreatic function-enhancing signals, which allow for the typical over twofold increase in insulin secretion during the second and third trimesters of gestation [77]. These signals include hPL, prolactin, and estrogens, all of which rise progressively and prominently throughout pregnancy [78], associated with increases in pancreatic β-cell mass and insulin transcription, and improve glucose-stimulated insulin secretion by promoting glucokinase and GLUT-2 expression, as well as raising glucose utilization and oxidation in pancreatic β cells [78]. These compensatory pathways are valuable, as they aim to maintain adequate glucose metabolism whilst allowing for increased FFA production [77]. Nonetheless, these mechanisms may be intrinsically defective or insufficient in some women, leading to the development of GDM, defined as glucose intolerance of onset or first recognition during pregnancy [79].

To this end, obesity is an important risk factor for GDM, with an OR = 2.6; 95% CI: 2.1–3.4; P < 0.05 [80]. Aside from
enhancing all previously described pro-IR mechanisms [72], obesity favors the development of a systemic inflammatory state, with elevated levels of mediators such as TNF [81]. This cytokine is implicated in IR by allowing IRS-1 serine phosphorylation via activation of JNK and NF-κB pathways [82]. Likewise, states of nutrient excess have been linked to upregulation of p70 S6K1, an IRS-1 serine kinase which induces degradation of this protein and may contribute to IRS-1 deficiency in GDM [72]. Similarly, both obesity and PCOS are associated with decreased expression of GLUT4 [83].

Another important factor is adiponectin, a proteic hormone with insulin-sensitizing activity, whose levels are decreased in obesity [84]. Although adipocytes are the primary site for adiponectin synthesis, placental production of adiponectin appears to be a paramount regulator of metabolism homeostasis during gestation [85]. Moreover, cytokines such as TNF, IFNγ, IL-6, and leptin have been found to modulate adiponectin and adiponectin receptor expression in women with GDM [86], harmonizing with reports associating hypoadiponectinemia with postpartum IR, β-cell dysfunction, and dysglycemia [87]. Expression of PPARγ is also diminished, leading to subdued lipogenic pathways, favoring greater FFA release [88] and disturbance of proper lipid partition, which would enhance lipid deposition in nonprofessional tissues such as skeletal muscle, enhancing the IR cycle [7]. Other related metabolic markers have been independently associated with higher risk for GDM: the Coronary Artery Risk Development in Young Adults (CARDIA) Study [89] reported that impaired fasting glucose (OR = 4.74; 95% CI: 2.14–10.51; P < 0.01), hyperinsulinemia (OR = 2.36; 95% CI: 1.20–4.63; P < 0.01), and low levels of HDL-C (OR = 3.07; 95% CI: 1.62–5.84; P < 0.01) are associated with GDM risk after adjusting for race, age, parity, and birth order.

4. Implications of Gestational Diabetes Mellitus on Fetomaternal Health

Gestational diabetes mellitus has been noted to prevail in females with predisposition to metabolic disturbances, with pregnancy acting as stress test on endocrine physiology [90], reflected on both obesity and PCOS representing independent risk factors for GDM, as previously discussed [20, 80]. This condition entails several consequences on both mother and offspring well-being. Maternal implications consist principally of higher risk for development of DM2 after pregnancy, with approximately 10% of women diagnosed with DM2 shortly after delivery and up to 40% after 10-year follow-up [91]. Indeed, gestation may reveal or worsen preexisting defects in β-cell function, accelerating onset of DM2 and other related conditions [90]. This influence is present even in nonobese women with GDM, with findings of endothelial dysfunction and chronic inflammation markers—both associated with the pathogenesis of DM2, cardiovascular disease, and metabolic syndrome—in this population [92]. HOMA-IR assessment boasts promising results as predictor of postpartum β-cell dysfunction [93].

On the other hand, the Hyperglycemia and Adverse Pregnancy Outcome (HAPO) study [94] has demonstrated that hyperglycemia during pregnancy—even in nondiabetic ranges—is associated with increased birth weight and elevated cord blood C-peptide serum levels. GDM is related to greater risk of macrosomia, shoulder dystocia, birth injuries, neonatal hypoglycemia, hypocalcemia, hyperbilirubinemia, respiratory distress syndrome, and polycythemia [95], as well as teratogenesis, particularly in obese subjects [96]. Furthermore, elevated cord-blood insulin concentrations are linked to glucose intolerance in offspring, and children exposed to GDM appear to display various metabolic disturbances well into childhood, including higher blood pressure and lower HDL-C [97].

These epidemiological data obey profound disruptions in embryonic and fetal metabolism, and numerous hypotheses attempt to explain this panorama. The theory of fuel-induced teratogenesis was first outlined by Freinkel [98], who proposed fuel excess and overgrowth to be the pathogenic basis of maternal hyperglycemia. This notion is founded on findings of maternal hyperglycemia-induced enhancing fetal insulin secretion, potentiating tissue growth—macrosomia—via fetal IGF-1 [99]. Alternatively, Hales and Barker [100] have propelled the thrifty phenotype theory, suggesting in utero malnutrition to bear a strong influence on postnatal risk of obesity, cardiovascular disease, and DM2, and even risk of PCOS and future pregnancy complications [101]. These premises are complemented by the concept of metabolic memory, related to endocrine-metabolic reprogramming of offspring amidst the diabetic environment during pregnancy [102]. This notion encompasses fetal inflammation, blunted myogenesis, oxidative stress, and disruption of immune system tolerance, among various other alterations [103]. Likewise, fetal exposure to diabetes appears to modify hypothalamic functionality in animal models, associated with hyperphagic behavior and obesity-proneness after birth [104].

AMP-dependent kinase (AMPK), a classic target of metformin action, may be an important mediator in this context [105], as it intervenes in processes such as lipogenesis via inhibition of acetyl-CoA carboxylase [106], myogenesis through the modulation of myocyte enhancer factor 2 [107], cell cycle [108], and appetite pathways [109]. Animal models have shown that metformin-induced AMPK activation yields beneficial effects over embryonic implantation [110], fetal inflammation [111], maternal liver function [112], and pregnancy outcomes [113]. Notwithstanding that these and other molecular pathways remain under research and certain aspects require further characterization, metformin has proven to beat the test of time, standing as a promising recourse in many circumstances, including GDM.

5. Metformin Pharmacokinetics during Pregnancy

Uptake and distribution of metformin towards the circulatory system requires the participation of bidirectional transporters located in the intestine and liver [114, 115]; see Figure 3. In the apical membrane of enterocytes, PMAT (Plasma Membrane Monoamine Transporter) and OCT3 (Organic Cation Transporters) mediate absorption. Mobilization of the drug
6 International Journal of Reproductive Medicine

Figure 3: Absorption and distribution of metformin during pregnancy.

Towards the liver requires OCT1, OCT2, and OCT3, while OCT2 is needed in order to reach the bloodstream, kidneys, and excretion [116]. Renal clearance of metformin increases during mid (723 ± 243 mL/min, \( P < 0.01 \)) and late pregnancy (625 ± 130 mL/min, \( P < 0.01 \)) [116], relating to a concentration of the drug in umbilical cord blood at time of birth between undetectable levels and 1263 ng/mL. Placental tissue expresses OCT2 transporter, yet under strict epigenetic control [117, 118], underlying ample interindividual differences in this aspect. However, other transporters are also involved in drug efflux through the placenta. Reflecting the high protectiveness of the human syncytiotrophoblast regarding the fetus, this tissue has been described to express a series of transporters in the apical membrane, such as P-glycoprotein (P-gp), Multidrug Resistance-Associated Protein 1 (MRP1), and Breast Cancer Resistance Protein (BCRP) [119–122], with metformin being transported mainly via P-gp (58% ± 20%) and BCRP (25% ± 14%) [119]. Competition between this biguanide and other drugs can also limit the exposure of the fetus, further limiting the presence of toxic concentrations during pregnancy.

Animal studies using dosages up to 600 mg/kg daily have failed to report evidence of teratogenic effects [123] and extremely high dosages between 900 and 1500 mg/kg daily failed to induce carcinogenicity [124]. Furthermore, in 2003 Gutzin et al. [125] reported their results concerning first trimester exposure, ascertaining no higher rates of major malformations with an OR of 1.05 (95% CI: 0.65–1.70), while neonatal death rendered an OR of 1.16 (95% CI: 0.67–2.00). Likewise, Gilbert et al. [126] conducted a meta-analysis on 8 studies concerning fetal malformations associated with metformin use during pregnancy, indicating this drug to yield an OR of 0.50 (95% CI: 0.15–1.60)—rendering a minor protective effect. Finally, the pooling analysis showed that the control group had a malformation rate of 7.2%, compared to 1.7% in the metformin group [126], strongly supporting metformin’s safety during pregnancy.

Concerning breast milk-related exposure [127], it has been confirmed that metformin can be detected at ranges between 0.13 and 0.28 mg/mL, equivalent to <0.5% of the mother’s weight-adjusted dosage [106]. Other reports have quantified metformin in breast milk at 0.28–1.08% [128] and 0.18–0.21% [129] of maternal dose. Placental partition coefficient for metformin has been calculated at 36.3%, with a cord plasma concentration of 0.1–2.9 mg/L during labor [130]. Such findings confirm that neonatal exposure to metformin is actually quite insignificant, and it is not related to glucose abnormality in infants, granting safe use before, during, and after pregnancy [128–130].

6. Metformin Use in Pregnant Women with Polycystic Ovary Syndrome: Different Outcomes, Different Efficacy

Because infertility is one of the main consequences of female reproduction in patients with PCOS [4, 5], ovulation induction remains the most common intervention during fertility counseling. Current guidelines heavily promote lifestyle modifications and support clomiphene as the first-line agent for ovulation induction, while recognizing that complementation with metformin improves ovulation and pregnancy success [131], as reported by Lord et al. [22] in their meta-analysis concerning effectiveness of this antidiabetic drug in achievement of ovulation in 15 trials involving 543 participants. This yielded an OR of 3.88 (95% CI: 2.25–6.69) for metformin alone and 4.41 (95% CI: 2.37–8.22) for metformin combined with clomiphene. In addition, the results from Khorram et al. [132] showed that two-week treatment with insulin reduced insulin levels and IR while improving SHBG levels and clomiphene-induced ovulation. In regards to metformin and gonadotropin use, Palomba et al. [133] reported that the biguanide improved live birth rates (OR = 1.95; 95% CI: 1.10–3.44; \( P = 0.020 \)) and pregnancy success (OR = 2.25; 95% CI: 1.50–3.38; \( P < 0.0001 \)).

Early pregnancy loss (EPL) is defined as the interruption of pregnancy before the 20th week of gestation [134]. Although chromosomal abnormalities are the principal cause
of EPL [135], they are uncommonly reported in women with PCOS [136]. It has been proposed that endocrine disruptions may play a role in EPL, with elevated androgens being associated with EPL in women with PCOS, and with recurrent EPL in women with and without PCOS [21]. Additionally, several endometrial molecular alterations have been described during implantation in PCOS: (a) androgen-dependent suppression of glycodelin [137], a cell-adhesion molecule involved in endometrial receptivity [138]; (b) IR-hyperinsulinemia can also diminish glycodelin expression, alongside IGFBP-1, key molecules for endometrial preimplantation maturation [139]; and (c) a hypofibrinolytic state due to increased synthesis of plasminogen activator inhibitor-1 (PAI-1), which has been found to be an independent risk factor for EPL in PCOS [140]. In this context, PCOS patients prescribed with metformin have lower pooled odds ratios for EPL (OR = 0.32, 95% CI: 0.19–0.55) and preterm birth (OR = 0.30, 95% CI: 0.13–0.68) [141], suggesting that this treatment can reverse the impact of PCOS on implantation success observed in this gynecoenocrine disease.

Other benefits have been attributed to metformin throughout gestation in women with PCOS, but perhaps one of the most important ones, is the 40% reduction of new-onset diabetes in high risk individuals as reported by Salpeter et al. [142]. In their meta-analysis using 31 trials and 4,570 subjects, the resulting pooled OR was 0.6 (95% CI: 0.5–0.8), with an absolute risk reduction of 6% (95% CI: 4–8) during a period of treatment of 1.8 years [142]. On the other hand, Nawaz et al. [143] have described decreased prevalence of fetal growth restriction and increased live birth rates, as well as an absence of intrauterine deaths or stillbirths, in women taking metformin during pregnancy, in line with claims of metformin being unrelated to teratogenicity [144].

Nevertheless, metformin during pregnancy appears unable to significantly reduce rates of preeclampsia and preterm birth in subjects with PCOS. A randomized, placebo-controlled, double-blind, multicenter study by Vanky et al. [145] found that preeclampsia prevalence was 7.4% in the metformin group and 3.7% in the placebo group (3.7%; 95% CI: 1.7–9.2; \( P = 0.18 \)), whereas preterm birth prevalence was 3.7% in the metformin group and 8.2% in the placebo group (4.4%; 95% CI: −10.1–1.2; \( P = 0.12 \)); the inefficacy of metformin at preventing preeclampsia may be due to the complex etiopathogenesis of this disease. Data from Stridsklev et al. [146] support this phenomenon, in which reporting metformin treatment did not affect uterine artery flow during gestation, while also describing an association between uterine artery flow and androgens, highlighting the complexity of the mechanisms underlying placentation, conservation of uterine artery flow, and vessel compliance [147, 148].

Indeed, despite several mechanisms related to IR-hyperinsulinemia being involved in the etiopathogenesis of preeclampsia—chronic systemic inflammation, increased sympathetic tone, and vascular smooth muscle growth [149]—metformin may be unable to effectively modify the pathogenic root of this disease, which is faulty placentation [150]. Similarly, although metformin's effects may aid in prevention of preterm birth by ameliorating oxidative stress and chronic inflammation [151], various elements underlying preterm labor may escape the reach of metformin's activity, including the most common factors associated with this condition—defective placentation, intrauterine infection, and maternal immunologic receptivity [152].

Still, metformin seems to offer other benefits to offspring of women with PCOS even in the postnatal period. In this scenario, metformin throughout pregnancy has been associated with diminished neonatal hypoglycemia [153], as well as normal growth and motor-social development in the first 18 months of life [154]. Likewise, the growth and motor-social skills of breast-fed children of women with PCOS taking metformin have been demonstrated to be similar to those of formula-fed infants, with no abnormalities [155].

### 7. Metformin in Pregnant Women with Gestational Diabetes Mellitus: Challenging Insulin as the Go-To Therapy

Although insulin therapy has been considered the best management option for GDM, recent evidence diverges from this precept. The first major trial concerning the use of metformin and/or insulin during pregnancies complicated with GDM was the metformin in gestational diabetes (MiG) [156], whose goal was to determine the effects of either drug on prevention of fetal hyperinsulinemia and promotion of lower maternal glycermia. This research group ascertained metformin (500–2500 mg/day) with or without supplemental insulin not to be associated with higher perinatal complications, in comparison to insulin alone [157], findings later corroborated by Silva et al. [158]. Furthermore, patients tend to prefer metformin over insulin as treatment schemes and would rather be prescribed such drug if possible [156]. Likewise, metformin use during pregnancy failed to adversely affect maternal lipid parameters, C-reactive protein levels, or birth weight [159].

After this emblematic trial, several other studies have supported the effectiveness of metformin in GDM. Nironmanesh et al. [160] conducted a randomized controlled trial with 160 pregnant patients with GDM, 80 of them treated with metformin (500–2500 mg) and the rest with insulin NPH (0.2 U/kg bedtime) and regular (1 U per 10 mg/dL over). Results revealed metformin to reduce rates of macrosomia and maternal weight gain. Additionally, Rowan et al. [161] also ascertained a decline in macrosomia and preeclampsia rates and suggested glycemic goals in GDM should be more rigorous. Metformin in GDM has also been described to lower incidence of surgical delivery [162]. Notably, these effects are observed even in spite of lowering of vitamin B12 [163], a recognized side effect of the drug [164].

Although various oral hypoglycemic agents—aside from metformin—are known to confer adequate metabolic control during pregnancy compared to insulin [165], metformin seems to be the superior choice, offering better control than glyburide, as reported by Silva et al. [158]. This research group has also reported newborns from mothers treated with metformin to obtain lower weight (3193 g versus 3387 g; \( P = 0.01 \)) and ponderal index results (2.87 versus 2.96; \( P = 0.05 \)) as
well as less maternal weight gain, in women with GDM, when compared to those treated with glyburide (10.3 kg versus 7.6 kg; \( P = 0.02 \)) [166], possibly reducing probabilities of other weight-related complications, such as preeclampsia. On the other hand, data on TZD use during GDM is relatively scarce, and trials conducted to date are considered insufficient to definitively establish these drugs as safe during pregnancy [25]. In this context, PPARγ has been noted to be key in embryonic development [167], and TZD administration during pregnancy has been associated with impaired fetal development [168], with this drug class remaining within the FDA Pregnancy Category C [169]. Therefore, further research is needed to explore the role of TZD in pregnancy and GDM.

Beyond evidence supporting metformin use in GDM, a key issue regarding pharmacological management of this disease is the prediction and selection of the best suited alternative (insulin alone, metformin alone, or both combined) for each specific patient. Insulin remains the most recommended option in mild cases of GDM [170] and in women with elevated BMI [171]. Indeed, in women with GDM, HOMA-IR values 1.29–2.89—interpreted as decreased insulin secretion—have been proposed to indicate a requirement of insulin therapy, whereas values >2.89 are thought to underline insufficient compensation of IR, rendering insulin-sensitizing agents more adequate [172]. Likewise, women with GDM and a fasting glucose result from oral glucose tolerance test below 93.3 mg/dL have displayed a probability of favorable pharmacological response of 93% to metformin [173]. On the other hand, early detection of GDM is a predictor for supplemental insulin treatment in women initially treated with metformin [174], as well as older age and elevated serum fructosamine concentration [175].

8. Concluding Remarks

Pregnancies complicated with GDM or with history of PCOS are a challenge for both obstetricians and endocrinologists, representing a halfway point where these specialties merge and highlighting the importance of multidisciplinary prenatal management. In our experience, we have observed that patients with PCOS who continue with metformin treatment throughout pregnancy and those who receive this drug as a pharmacological intervention in GDM yield better pregnancy outcomes and a better postpartum metabolic prognosis for both mothers and their offspring.

Nevertheless, further studies are needed to uncover and elucidate the benefits and shortcomings of metformin in this context, in both molecular and epidemiological fields. Ongoing studies concerning these issues include the Metformin to Prevent Late Miscarriage and Preterm Delivery in Women With Polycystic Ovary Syndrome Trial (PregMet2) [176] and the Metformin Treatment in Gestational Diabetes and Non-insulin Dependent Diabetes in Pregnancy in a Developing Country Trial (migdm&tdm) [177] as well as additional data from the MiG trial, among many others. Indeed, the future appears compelling and exciting in this aspect, with these sources promising valuable information which may reshape and refine views on metformin use during pregnancy.

Conflict of Interests

There are no financial or other contractual agreements that might cause conflict of interests.

Acknowledgments

This work was supported by Research Grant no. CC-0437-10-21-09-10 from CONDES, University of Zulfit, and Research Grant no. FZ-0058-2007 from Fundacite-Zulia.

References

[1] H. Teede, A. Deeks, and L. Moran, “Polycystic ovary syndrome: a complex condition with psychological, reproductive and metabolic manifestations that impacts on health across the lifespan,” BMC Medicine, vol. 8, article 41, 2010.
[2] M. T. Sheehan, “Polycystic ovarian syndrome: diagnosis and management,” Clinical Medicine & Research, vol. 2, no. 1, pp. 13–27, 2004.
[3] C. B. Kallen, “Steroid hormone synthesis in pregnancy,” Obstetrics and Gynecology Clinics of North America, vol. 31, no. 4, pp. 795–816, 2004.
[4] J. Vrbikova and V. Hainer, “Obesity and polycystic ovary syndrome,” Obesity Facts, vol. 2, no. 1, pp. 26–35, 2009.
[5] M. N. Mascarenhas, S. R. Flaxman, T. Boerma, S. Vanderpoel, and G. A. Stevens, “National, regional, and global trends in infertility prevalence since 1990: a systematic analysis of 277 health surveys,” PLoS Medicine, vol. 9, no. 12, Article ID e1001356, 2012.
[6] E. Diamanti-Kandarakis, “Polycystic ovarian syndrome: pathophysiology, molecular aspects and clinical implications,” Expert Reviews in Molecular Medicine, vol. 10, p. e3, 2008.
[7] J. Rojas, M. Chávez, L. Olivar et al., “Polycystic ovary syndrome, insulin resistance, and obesity: navigating the pathophysiologic labyrinth,” International Journal of Reproductive Medicine, vol. 2014, Article ID 719050, 17 pages, 2014.
[8] S. A. Arslanian, V. D. Lewy, and K. Danadian, “Glucose intolerance in obese adolescents with polycystic ovary syndrome: roles of insulin resistance and β-cell dysfunction and risk of cardiovascular disease,” The Journal of Clinical Endocrinology and Metabolism, vol. 86, no. 1, pp. 66–71, 2001.
[9] S. Robinson, A. D. Henderson, S. V. Gelding et al., “Dyslipidaemia is associated with insulin resistance in women with polycystic ovaries,” Clinical Endocrinology, vol. 44, no. 3, pp. 277–284, 1996.
[10] M. W. Elting, T. J. M. Korsen, P. D. Bezemer, and J. Schoemaker, “Prevalence of diabetes mellitus, hypertension and cardiac complaints in a follow-up study of a Dutch PCOS population,” Human Reproduction, vol. 16, no. 3, pp. 556–560, 2001.
[11] Z. H. Huang, B. Manickam, V. Ryvkin et al., “PCOS is associated with increased CD11c expression and crown-like structures in adipose tissue and increased central abdominal fat depots independent of obesity,” The Journal of Clinical Endocrinology & Metabolism, vol. 98, no. 1, pp. E17–E24, 2013.
[12] S. Borruei, E. Fernández-Durán, M. Alpañés et al., “Global adiposity and thickness of intraperitoneal and mesenteric adipose tissue depots are increased in women with polycystic ovary syndrome (PCOS),” Journal of Clinical Endocrinology and Metabolism, vol. 98, no. 3, pp. 1254–1263, 2013.
[13] R. Shroff, A. Kerchner, M. Maifeld, E. J. R. van Beek, D. Jagasia, and A. Dokras, “Young obese women with polycystic ovary
syndrome have evidence of early coronary atherosclerosis,” The Journal of Clinical Endocrinology & Metabolism, vol. 92, no. 12, pp. 4609–4614, 2007.

[14] A. J. Cussons, B. G. A. Stuckey, and G. F. Watts, “Metabolic syndrome and cardiometabolic risk in PCOS,” Current Diabetes Reports, vol. 7, no. 1, pp. 66–73, 2007.

[15] M. C. Amato, V. Guarnotta, D. Forti, M. Donatelli, S. Dolcimacolo, and C. Giordano, “Metabolically healthy polycystic ovary syndrome (MH-PCOS) and metabolically unhealthy polycystic ovary syndrome (MU-PCOS): a comparative analysis of four simple methods useful for metabolic assessment,” Human Reproduction, vol. 28, no. 7, pp. 1919–1928, 2013.

[16] E. Mor, A. Zograbyan, P. Saadat et al., “The insulin resistant subphenotype of polycystic ovary syndrome: clinical parameters and pathogenesis,” The American Journal of Obstetrics and Gynecology, vol. 190, no. 6, pp. 1654–1660, 2004.

[17] L. B. Craig, R. W. Ke, and W. H. Kutteh, “Increased prevalence of insulin resistance in women with a history of recurrent pregnancy loss,” Fertility and Sterility, vol. 78, no. 3, pp. 487–490, 2002.

[18] A. Huidobro, A. M. Prentice, A. J. C. Fulford, and J. Rozowski, “Antropometría como predictor de diabetes gestacional: estudio de cohorte,” Revista Médica de Chile, vol. 138, pp. 1373–1377, 2010.

[19] C. M. Clark Jr., C. Qiu, B. Amerman et al., “Gestational diabetes: should it be added to the syndrome of insulin resistance?” Diabetes Care, vol. 20, no. 5, pp. 867–871, 1997.

[20] C. M. Boomsma, M. J. C. Eijkemans, E. G. Hughes, G. H. A. Visser, B. C. J. M. Fauser, and N. S. Macklon, “A meta-analysis of pregnancy outcomes in women with polycystic ovary syndrome,” Human Reproduction Update, vol. 12, no. 6, pp. 673–683, 2006.

[21] S. Kamalanathan, J. P. Sahoo, and T. Sathypalan, “Pregnancy in polycystic ovary syndrome,” Indian Journal of Endocrinology and Metabolism, vol. 17, pp. 37–43, 2013.

[22] J. M. Lord, I. H. Flight, and R. J. Norman, “Insulin-sensitising drugs (metformin, troglitazone, rosiglitazone, pioglitazone, D-chiro-inositol) for polycystic ovary syndrome,” Cochrane Database of Systematic Reviews, vol. 3, Article ID CD003053, 2003.

[23] H. C. Zisser, “Polycystic ovary syndrome and pregnancy: is metformin the magic bullet?” Diabetes Spectrum, vol. 20, no. 2, pp. 85–89, 2007.

[24] M.-E. Lautzis, D. G. Goulis, and M. Vrontakis, “Efficacy and safety of metformin during pregnancy in women with gestational diabetes mellitus or polycystic ovary syndrome: a systematic review,” Metabolism: Clinical and Experimental, vol. 62, no. 11, pp. 1522–1534, 2013.

[25] D. S. Feig, G. G. Briggs, and G. Koren, “Oral antidiabetic agents in pregnancy and lactation: a paradigm shift?” Annals of Pharmacotherapy, vol. 41, no. 7–8, pp. 1174–1180, 2007.

[26] Package Insert for Glucophage, http://www.glucophagexr.com/pages/default.aspx.

[27] C. R. McCartney, C. A. Eagleson, and J. C. Marshall, “Regulation of gonadotropin secretion: implications for polycystic ovary syndrome,” Seminars in Reproductive Medicine, vol. 20, no. 4, pp. 317–325, 2002.

[28] M. Karoshi and S. O. Okolo, “Commentary: Polycystic ovarian disease (PCOD): a misnomer, looking for a new name,” International Journal of Fertility and Women’s Medicine, vol. 49, no. 4, pp. 191–192, 2004.

[29] D. H. Abbott, D. A. Dumesic, and S. Franks, “Developmental origin of polycystic ovary syndrome—a hypothesis,” Journal of Endocrinology, vol. 174, no. 1, pp. 1–5, 2002.

[30] D. A. Dumesic, D. H. Abbott, and V. Padmanabhan, “Polycystic ovary syndrome and its developmental origins,” Reviews in Endocrine and Metabolic Disorders, vol. 8, no. 2, pp. 127–141, 2007.

[31] M. H. Shanik, Y. Xu, J. Skrha, R. Dankner, Y. Zick, and J. Roth, “Insulin resistance and hyperinsulinemia: hyperinsulinemia the cart or the horse?” Diabetes Care, vol. 31, supplement 2, pp. S262–S268, 2008.

[32] G. M. Reaven, “Compensatory hyperinsulinemia and the development of an atherogenic lipoprotein profile: the price paid to maintain glucose homeostasis in insulin-resistant individuals,” Endocrinology and Metabolism Clinics of North America, vol. 34, no. 1, pp. 49–62, 2005.

[33] J. Rojas, V. Bermúdez, E. Leal et al., “Insulinorresistencia e hiperinsulinemia como factores de riesgo para enfermedad cardiovascular,” AVFT, vol. 27, pp. 29–39, 2008.

[34] N. Sekar, J. C. Garney, and J. D. Veldhuis, “Mechanisms underlying the steroidogenic synergy of insulin and luteinizing hormone in porcine granulosa cells: joint amplification of pivotal sterol-regulatory genes encoding the low-density lipoprotein (LDL) receptor, steroidogenic acute regulatory (stAR) protein and cytochrome P450 side-chain cleavage (P450scc) enzyme,” Molecular and Cellular Endocrinology, vol. 159, no. 1-2, pp. 25–35, 2000.

[35] E. Y. Adashi, A. J. W. Hsueh, and S. S. C. Yen, “Insulin enhancement of luteinizing hormone and follicle-stimulating hormone release by cultured pituitary cells,” Endocrinology, vol. 108, no. 4, pp. 1441–1449, 1981.

[36] R. Salvi, E. Castillo, M.-J. Voirl et al., “Gonadotropin-releasing hormone-expressing neurons immortalized conditionally are activated by insulin: implication of the mitogen-activated protein kinase pathway,” Endocrinology, vol. 147, no. 2, pp. 816–826, 2006.

[37] J. W. Hill, J. K. Elmquist, and C. F. Elias, “Hypothalamic pathways linking energy balance and reproduction,” American Journal of Physiology: Endocrinology and Metabolism, vol. 294, no. 5, pp. E827–E832, 2008.

[38] A. D. Eyvazzadeh, K. P. Pennington, R. Pop-Busui, M. Sowers, J.-K. Zubieta, and Y. R. Smith, “The role of the endogenous opioid system in polycystic ovary syndrome,” Fertility and Sterility, vol. 92, no. 1, pp. 1–12, 2009.

[39] D. M. Selva, K. N. Hogeveen, S. M. Innis, and G. L. Hammond, “Monosaccharide-induced lipogenesis regulates the human hepatic sex hormone-binding globulin gene,” Journal of Clinical Investigation, vol. 117, no. 12, pp. 3979–3987, 2007.

[40] A. Gambineri, C. Pelusi, V. Vicennati, U. Pagotto, and R. Pasquali, “Obesity and the polycystic ovary syndrome,” International Journal of Obesity, vol. 26, no. 7, pp. 883–896, 2002.

[41] C.-B. Book and A. Dunaif, “Selective insulin resistance in the polycystic ovary syndrome,” Journal of Clinical Endocrinology and Metabolism, vol. 84, no. 9, pp. 3110–3116, 1999.

[42] E. Méndez, N. Montserrat, and J. V. Planas, “Modulation of the steroidogenic activity of luteinizing hormone by insulin and insulin-like growth factor-I through interaction with the cAMP-dependent protein kinase signaling pathway in the trout ovary,” Molecular and Cellular Endocrinology, vol. 229, no. 1-2, pp. 49–56, 2005.

[43] A. Dunaif, J. Xia, C.-B. Book, E. Schenker, and Z. Tang, “Excessive insulin receptor serine phosphorylation in cultured
fibroblasts and in skeletal muscle: a potential mechanism for insulin resistance in the polycystic ovary syndrome,” *Journal of Clinical Investigation*, vol. 96, no. 2, pp. 801–810, 1995.

[44] J. E. Nestler, D. J. Jakubowicz, and M. J. Iuorno, “Role of inositolphosphoglycan mediators of insulin action in the polycystic ovary syndrome,” *Journal of Pediatric Endocrinology & Metabolism*, vol. 13, supplement 5, pp. 1295–1298, 2000.

[45] P. Arner, “Effects of testosterone on fat cell lipolysis. Species differences and possible role in polycystic ovarian syndrome,” *Biochimie*, vol. 87, no. 1, pp. 39–43, 2005.

[46] L. J. Moran, R. Pasquali, H. J. Teede, K. M. Hoeger, and R. J. Norman, “Treatment of obesity in polycystic ovary syndrome: a position statement of the Androgen Excess and Polycystic Ovary Syndrome Society,” *Fertility and Sterility*, vol. 92, no. 6, pp. 1966–1982, 2009.

[47] L. Radosh, “Drug treatments for polycystic ovary syndrome,” *American Family Physician*, vol. 79, no. 8, pp. 671–676, 2009.

[48] A. D. Genazzani, E. Chierchia, E. Rattighieri et al., “Metformin administration restores allopregnalone response to adrenocorticotropic hormone (ACTH) stimulation in overweight hyperinsulinemic patients with PCOS,” *Gynecological Endocrinology*, vol. 26, no. 9, pp. 684–689, 2010.

[49] E. M. Velazquez, S. Mendoza, T. Hamer, F. Sosa, and C. J. Glueck, “Metformin therapy in polycystic ovary syndrome reduces hyperinsulinemia, insulin resistance, hyperandrogenemia, and systolic blood pressure, while facilitating normal menses and pregnancy,” *Metabolism: Clinical and Experimental*, vol. 43, no. 5, pp. 647–654, 1994.

[50] A. Kriplani and N. Agarwal, “Effects of metformin on clinical and biochemical parameters in polycystic ovary syndrome,” *The Journal of Reproductive Medicine*, vol. 49, no. 5, pp. 361–367, 2004.

[51] D. Kocer, F. Bayram, and H. Diri, “The effects of metformin on endothelial dysfunction, lipid metabolism and oxidative stress in women with polycystic ovary syndrome,” *Gynecological Endocrinology*, vol. 30, no. 5, pp. 367–371, 2014.

[52] E. Diamanti-Kandarakis, T. Paterakis, and H. A. Kandarakis, “Indices of low-grade inflammation in polycystic ovary syndrome,” *Annals of the New York Academy of Sciences*, vol. 1092, pp. 175–186, 2006.

[53] S. Tan, S. Hahn, S. Benson et al., “Metformin improves polycystic ovary syndrome symptoms irrespective of pre-treatment insulin resistance,” *European Journal of Endocrinology*, vol. 157, no. 5, pp. 669–676, 2007.

[54] J. Nawrocka and A. Starczewski, “Effects of metformin treatment in women with polycystic ovary syndrome depends on insulin resistance,” *Gynecological Endocrinology*, vol. 23, no. 4, pp. 231–237, 2007.

[55] R. Horejsi, R. Möller, S. Rackl et al., “Android subcutaneous adipose tissue topography in lean and obese women suffering from PCOS: comparison with type 2 diabetic women,” *The American Journal of Physical Anthropology*, vol. 124, no. 3, pp. 275–281, 2004.

[56] J. G. Dolfing, C. M. Stassen, P. M. M. Van Haard, B. H. R. Wolfenbuttel, and D. H. Schweitzer, “Comparison of MRI-assessed body fat content between lean women with polycystic ovary syndrome (PCOS) and matched controls: less visceral fat with PCOS,” *Human Reproduction*, vol. 26, no. 6, pp. 1495–1500, 2011.

[57] J.-P. Baillargeon and A. Carpentier, “Role of insulin in the hyperandrogenemia of lean women with polycystic ovary syndrome and normal insulin sensitivity,” *Fertility and Sterility*, vol. 88, no. 4, pp. 886–893, 2007.

[58] R. Keskin Kurt, A. G. Okyay, A. U. Hakverdi et al., “The effect of obesity on inflammatory markers in patients with PCOS: a BMI-matched case-control study,” *Archives of Gynecology and Obstetrics*, vol. 290, no. 2, pp. 315–319, 2014.

[59] C. Celik, E. Bastu, R. Abali et al., “The relationship between copper, homocysteine and early vascular disease in lean women with polycystic ovary syndrome,” *Gynecological Endocrinology*, vol. 29, no. 5, pp. 488–491, 2013.

[60] S. A. Blair, T. Kyaw-Tun, I. S. Young, N. A. Phelan, J. Gibney, and J. McEneny, “Oxidative stress and inflammation in lean and obese subjects with polycystic ovary syndrome,” *Journal of Reproductive Medicine*, vol. 58, no. 3–4, pp. 107–114, 2013.

[61] D. W. Stovall, A. P. Bailey, and L. M. Pastore, “Assessment of insulin resistance and impaired glucose tolerance in lean women with polycystic ovary syndrome,” *Journal of Women's Health*, vol. 20, no. 1, pp. 37–43, 2011.

[62] S. Livadas, A. Kollias, D. Panidis, and E. Diamanti-Kandarakis, “Diverse impacts of aging on insulin resistance in lean and obese women with polycystic ovary syndrome: evidence from 1345 women with the syndrome,” *European Journal of Endocrinology*, vol. 171, no. 3, pp. 301–309, 2014.

[63] G. Önalan, U. Goktolga, T. Ceyhan, T. Bagis, R. Onalan, and R. Pabayçu, “Predictive value of glucose—insulin ratio in PCOS and profile of women who will benefit from metformin therapy: obese, lean, hyper or normoinsulinemic?” *European Journal of Obstetrics Gynecology and Reproductive Biology*, vol. 123, no. 2, pp. 204–211, 2005.

[64] A. S. Kumari, A. Haq, R. Jayasundaram, L. O. Abdel-Wareth, S. A. al Haija, and M. Alvares, “Metformin monotherapy in lean women with polycystic ovary syndrome,” *Reproductive BioMedicine Online*, vol. 10, no. 1, pp. 100–104, 2005.

[65] A. Kruszynska, J. Sliwinska-Szrednicka, W. Jeske, and W. Zgliczyński, “Proinsulin, adiponectin and hsCRP in reproductive age women with polycystic ovary syndrome (PCOS)—the effect of metformin treatment,” *Endokrynologia Polska*, vol. 65, no. 1, pp. 2–10, 2014.

[66] Q. Du, Y. J. Wang, S. Yang, B. Wu, P. Han, and Y. Y. Zhao, “A systematic review and meta-analysis of randomized controlled trials comparing pioglitazone versus the treatment of polycystic ovary syndrome,” *Current Medical Research and Opinion*, vol. 28, pp. 723–730, 2012.

[67] A. Ziaee, S. Oveis, A. Abedini, S. Hashemipour, T. Karimzadeh, and A. Ghorbani, “Effect of metformin and pioglitazone treatment on cardiovascular risk profile in polycystic ovary syndrome,” *Acta Medica Indonesiana*, vol. 44, no. 1, pp. 16–22, 2012.

[68] V. R. Aroda, T. P. Ciaraldi, P. Burke et al., “Metabolic and hormonal changes induced by pioglitazone in polycystic ovary syndrome (PCOS)—the effect of metformin treatment,” *Journal of Clinical Endocrinology and Metabolism*, vol. 94, no. 2, pp. 469–476, 2009.

[69] T. P. Ciaraldi, V. Aroda, S. R. Mudaliai, and R. R. Henry, “Inflammatory cytokines and chemokines, skeletal muscle and oxidative stress, in women with polycystic ovary syndrome: a randomized, placebo-controlled clinical trial,” *Journal of Clinical Endocrinology and Metabolism*, vol. 62, no. 11, pp. 1587–1596, 2013.

[70] K. D. Wilson, Z. Li, R. Wagner et al., “Transcriptome alteration in the diabetic heart by rosiglitazone: implications for cardiovascular mortality,” *PLoS ONE*, vol. 3, no. 7, Article ID e2609, 2008.

[71] R. M. Lago, P. P. Singh, and R. W. Nesto, “Congestive heart failure and cardiovascular death in patients with prediabetes and type 2 diabetes given thiazolidinediones: a meta-analysis
of randomised clinical trials,” *The Lancet*, vol. 370, no. 9593, pp. 1129–1136, 2007.

[72] L. A. Barbour, C. E. McCurdy, T. L. Hernandez, J. P. Kirwan, P. M. Catalano, and J. E. Friedman, “Cellular mechanisms for insulin resistance in normal pregnancy and gestational diabetes,” *Diabetes Care*, vol. 30, no. 2, pp. S112–S119, 2007.

[73] J. Shao, P. M. Catalano, H. Yamashita et al., “Decreased insulin receptor tyrosine kinase activity and plasma cell membrane glycoprotein-1 overexpression in skeletal muscle from obese women with gestational diabetes mellitus (GDM): evidence for increased serine/threonine phosphorylation in pregnancy and GDM,” *Diabetes*, vol. 49, no. 4, pp. 603–610, 2000.

[74] P. M. Catalano, S. E. Nizielski, J. Shao, L. Preston, L. Qiao, and J. E. Friedman, “Downregulated IRS-1 and PPARγ in obese women with gestational diabetes: relationship to FFA during pregnancy,” *American Journal of Physiology: Endocrinology and Metabolism*, vol. 282, no. 3, pp. E522–E533, 2002.

[75] S. Okuno, S. Akazawa, I. Yasushi et al., “Decreased expression of the GLUT4 glucose transporter protein in adipose tissue during pregnancy,” *Hormone and Metabolic Research*, vol. 27, no. 5, pp. 231–234, 1995.

[76] E. Sivan and G. Boden, “Free fatty acids, insulin resistance, and pregnancy,” *Current Diabetes Reports*, vol. 3, no. 4, pp. 319–322, 2003.

[77] P. M. Catalano, L. Huston, S. B. Amini, and S. C. Kalhan, “Longoitudinal changes in glucose metabolism during pregnancy in obese women with normal glucose tolerance and gestational diabetes mellitus,” *The American Journal of Obstetrics and Gynecology*, vol. 180, no. 4, pp. 903–916, 1999.

[78] A. Nadal, P. Alonso-Magdalena, S. Soriano, A. B. Ropero, and I. Quesada, “The role of oestrogens in the adaptation of islets to insulin resistance,” *The Journal of Physiology*, vol. 587, no. 21, pp. 5031–5037, 2009.

[79] R. Kaaja and T. Rönnemaa, “Gestational diabetes: pathogenesis and consequences to mother and offspring,” *Review of Diabetic Studies*, vol. 5, no. 4, pp. 194–202, 2008.

[80] J. L. Weiss, F. D. Malone, D. Emig et al., “Obesity, obstetric complications and cesarean delivery rate—a population-based screening study,” *American Journal of Obstetrics & Gynecology*, vol. 190, no. 4, pp. 1091–1097, 2004.

[81] T. Tzanavari, P. Giannogonas, and K. P. Karalis, “TNF-α and obesity,” *Current Directions in Autoimmunity*, vol. 11, pp. 145–156, 2010.

[82] S. E. Shoelson, J. Lee, and A. B. Goldfine, “Inflammation and insulin resistance,” *The Journal of Clinical Investigation*, vol. 116, no. 7, pp. 1793–1801, 2006.

[83] D. Rosenbaum, R. S. Haber, and A. Dunaf, “Insulin resistance in polycystic ovary syndrome: decreased expression of GLUT-4 glucose transporters in adipocytes,” *The American Journal of Physiology—Endocrinology and Metabolism*, vol. 264, no. 2, pp. E197–E202, 1993.

[84] J. Kawano and R. Arora, “The role of adiponectin in obesity, diabetes, and cardiovascular disease,” *Journal of the Carbohydrate Metabolic Syndrome*, vol. 4, no. 1, pp. 44–49, 2009.

[85] I. L. M. H. Aye, T. L. Powell, and T. Jansson, “Review: adiponectin—the missing link between maternal adiposity, placental transport and fetal growth?” *Placenta*, vol. 34, pp. S40–S45, 2013.

[86] J. Chen, B. Tan, E. Karteris et al., “Secretion of adiponectin by human placenta: differential modulation of adiponectin and its receptors by cytokines,” *Diabetologia*, vol. 49, no. 6, pp. 1292–1302, 2006.

[87] R. Retnakaran, Y. Qi, P. W. Connelly, M. Sermer, A. J. Hanley, and B. Zinman, “Low adiponectin concentration during pregnancy predicts postpartum insulin resistance, beta cell dysfunction and fasting glycaemia,” *Diabetologia*, vol. 53, no. 2, pp. 268–276, 2010.

[88] T. A. Buchanan and A. H. Xiang, “Gestational diabetes mellitus,” *The Journal of Clinical Investigation*, vol. 115, no. 3, pp. 485–491, 2005.

[89] P. M. Catalano, L. Preston, L. Qiao, and J. E. Friedman, “Downregulated IRS-1 and PPARγ in obese women with gestational diabetes: relationship to FFA during pregnancy,” *American Journal of Physiology: Endocrinology and Metabolism*, vol. 282, no. 3, pp. E522–E533, 2002.

[90] S. Okuno, S. Akazawa, I. Yasushi et al., “Decreased expression of the GLUT4 glucose transporter protein in adipose tissue during pregnancy,” *Hormone and Metabolic Research*, vol. 27, no. 5, pp. 231–234, 1995.

[91] E. Sivan and G. Boden, “Free fatty acids, insulin resistance, and pregnancy,” *Current Diabetes Reports*, vol. 3, no. 4, pp. 319–322, 2003.

[92] P. M. Catalano, L. Huston, S. B. Amini, and S. C. Kalhan, “Longitudinal changes in glucose metabolism during pregnancy in obese women with normal glucose tolerance and gestational diabetes mellitus,” *The American Journal of Obstetrics and Gynecology*, vol. 180, no. 4, pp. 903–916, 1999.

[93] A. Nadal, P. Alonso-Magdalena, S. Soriano, A. B. Ropero, and I. Quesada, “The role of oestrogens in the adaptation of islets to insulin resistance,” *The Journal of Physiology*, vol. 587, no. 21, pp. 5031–5037, 2009.

[94] R. Kaaja and T. Rönnemaa, “Gestational diabetes: pathogenesis and consequences to mother and offspring,” *Review of Diabetic Studies*, vol. 5, no. 4, pp. 194–202, 2008.

[95] J. L. Weiss, F. D. Malone, D. Emig et al., “Obesity, obstetric complications and cesarean delivery rate—a population-based screening study,” *American Journal of Obstetrics & Gynecology*, vol. 190, no. 4, pp. 1091–1097, 2004.

[96] T. Tzanavari, P. Giannogonas, and K. P. Karalis, “TNF-α and obesity,” *Current Directions in Autoimmunity*, vol. 11, pp. 145–156, 2010.

[97] S. E. Shoelson, J. Lee, and A. B. Goldfine, “Inflammation and insulin resistance,” *The Journal of Clinical Investigation*, vol. 116, no. 7, pp. 1793–1801, 2006.

[98] D. Rosenbaum, R. S. Haber, and A. Dunaf, “Insulin resistance in polycystic ovary syndrome: decreased expression of GLUT-4 glucose transporters in adipocytes,” *The American Journal of Physiology—Endocrinology and Metabolism*, vol. 264, no. 2, pp. E197–E202, 1993.

[99] J. Kawano and R. Arora, “The role of adiponectin in obesity, diabetes, and cardiovascular disease,” *Journal of the Carbohydrate Metabolic Syndrome*, vol. 4, no. 1, pp. 44–49, 2009.

[100] I. L. M. H. Aye, T. L. Powell, and T. Jansson, “Review: adiponectin—the missing link between maternal adiposity, placental transport and fetal growth?” *Placenta*, vol. 34, pp. S40–S45, 2013.

[101] J. Chen, B. Tan, E. Karteris et al., “Secretion of adiponectin by human placenta: differential modulation of adiponectin and its receptors by cytokines,” *Diabetologia*, vol. 49, no. 6, pp. 1292–1302, 2006.
K. Franke, T. Harder, L. Aerts et al., “Programming” of orexigenic and anorexigenic hypothalamic neurons in offspring of treated and untreated diabetic mother rats,” Brain Research, vol. 1031, no. 2, pp. 276–283, 2005.

J. Rojas, N. Arraiz, M. Aguirre, M. Velasco, and V. Bermúdez, “AMPK as target for intervention in childhood and adolescent obesity,” Journal of Obesity, vol. 2011, Article ID 252817, 19 pages, 2011.

R. W. Brownsey, A. N. Boone, J. E. Elliott, J. E. Kulpa, and W. M. Mikhailov and R. J. Shaw, “The AMPK signalizing pathway coordinates cell growth, autophagy and metabolism,” Nature Cell Biology, vol. 13, no. 9, pp. 1016–1023, 2011.

R. Stark, S. E. Ashley, and Z. B. Andrews, “AMPK and the neuroendocrine regulation of appetite and energy expenditure,” Molecular and Cellular Endocrinology, vol. 366, no. 2, pp. 215–223, 2013.

P. Banerjee, R. R. Bhonde, and R. Pal, “Diverse roles of metformin during peri-implantation development: revisiting novel molecular mechanisms underlying clinical implications,” Stem Cells and Development, vol. 22, no. 22, pp. 2927–2934, 2013.

N. Desai, A. Roman, B. Rochelson et al., “Maternal metformin treatment decreases fetal inflammation in a rat model of obesity and metabolic syndrome,” American Journal of Obstetrics and Gynecology, vol. 209, no. 2, pp. 136-e1–136-e9, 2013.

H.-Y. Lee, D. Wei, and M. R. Loeken, “Lack of metformin effect on mouse embryo AMPK activity: implications for metformin treatment during pregnancy,” Diabetes/Metabolism Research and Reviews, vol. 30, no. 1, pp. 23–30, 2014.

G. S. Eng, R. A. Sheridan, A. Wyman et al., “AMP kinase activation increases glucose uptake, decreases apoptosis, and improves pregnancy outcome in embryos exposed to high IGF-1 concentrations,” Diabetes, vol. 56, no. 9, pp. 2228–2234, 2007.

J. J. Marin, “Plasma membrane transporters in modern liver pharmacology,” Scientific, vol. 2012, Article ID 428139, 15 pages, 2012.

L. Gong, S. Goswami, K. M. Giacomini, R. B. Altman, and T. E. Klein, “Metformin pathways: pharmacokinetics and pharmacodynamics,” Pharmacogenetics and Genomics, vol. 22, no. II, pp. 820–827, 2012.

J. W. Jonker and A. H. Schinkel, “Pharmacological and physiological functions of the polyspecific organic cation transporters: OCT1, 2, and 3 (SLC22A1-3),” Journal of Pharmacology and Experimental Therapeutics, vol. 308, no. 1, pp. 2–9, 2004.

S. Eyal, T. R. Easterling, D. Carr et al., “Pharmacokinetics of metformin during pregnancy,” Drug Metabolism and Disposition, vol. 38, no. 5, pp. 833–840, 2010.

J. Saito, T. Hirota, N. Kikunaga, K. Otsubo, and I. Ieiri, “Inter-individual differences in placental expression of the SLC22A2 (OCT2) gene: relationship to epigenetic variations in the 5’-upstream regulatory region,” Journal of Pharmaceutical Sciences, vol. 100, no. 9, pp. 3875–3883, 2011.

M. Kovo, N. Kogman, O. Ovadia, I. Nakash, A. Golan, and A. Hoffman, “Carrier-mediated transport of metformin across the human placenta determined by using the ex vivo perfusion of the placental cotyledon model,” Prenatal Diagnosis, vol. 28, no. 6, pp. 544–548, 2008.

M. Maliepaard, G. L. Scheffer, I. F. Faneyte et al., “Subcellular localization and distribution of the breast cancer resistance protein transporter in normal human tissues,” Cancer Research, vol. 61, no. 8, pp. 3458–3464, 2001.

M. V. St.-Pierre, M. A. Serrano, R. I. R. Macias et al., “Expression of members of the multidrug resistance protein family in human term placenta,” The American Journal of Physiology—Regulatory Integrative and Comparative Physiology, vol. 279, no. 4, pp. R1495–R1503, 2000.

V. Ganapathy, P. D. Prasad, M. E. Ganapathy, and F. H. Leibach, “Placental transporters relevant to drug distribution across the maternal-fetal interface,” Journal of Pharmacology and Experimental Therapeutics, vol. 294, no. 2, pp. 413–420, 2000.

G. G. Briggs, R. K. Freeman, and S. J. Yaffe, Drugs in Pregnancy and Lactation, Lippincott Williams & Wilkins, Philadelphia, PA, USA, 2002.

Package Insert: Metformin Hydrochloride, Bristol-Myers Squibb, New York, NY, USA, 2009.

S. J. Gutzin, E. Kozer, L. A. Magee, D. S. Feig, and G. Koren, “The safety of oral hypoglycemic agents in the first trimester of pregnancy: a meta-analysis,” Canadian Journal of Clinical Pharmacology, vol. 10, no. 4, pp. 179–183, 2003.

C. Gilbert, M. Valois, and G. Koren, “Pregnancy outcome after first-trimester exposure to metformin: a meta-analysis,” Fertility and Sterility, vol. 86, no. 3, pp. 658–663, 2006.

G. G. Briggs, P. J. Ambrose, M. P. Nageotte, G. Padilla, and S. Wan, “Excretion of metformin into breast milk and the effect on nursing infants,” Obstetrics and Gynecology, vol. 105, no. 6, pp. 1437–1441, 2005.

C. J. Glueck and P. Wang, “Metformin before and during pregnancy and lactation in polycystic ovary syndrome,” Expert Opinion on Drug Safety, vol. 6, no. 2, pp. 191–198, 2007.

S. J. Gardiner, C. M. J. Kirkpatrick, E. J. Begg, M. Zhang, M. Peter Moore, and D. J. Saville, “Transfer of metformin into human milk,” Clinical Pharmacology and Therapeutics, vol. 73, no. 1, pp. 71–77, 2003.

B. Charles, R. Norris, X. Xiao, and W. Hague, “Population pharmacokinetics of metformin in late pregnancy,” Therapeutic Drug Monitoring, vol. 28, no. 1, pp. 67–72, 2006.

T. D. R. Vause, A. P. Cheung, S. Sierra et al., “Ovulation induction in polycystic ovary syndrome,” Journal of Obstetrics and Gynaecology Canada, vol. 32, no. 5, pp. 495–502, 2010.

O. Khorrarn, J. P. Hellwell, S. Katz, C. M. Bonpane, and L. Jaramillo, “Two weeks of metformin improves clomiphene citrate-induced ovulation and metabolic profiles in women with polycystic ovary syndrome,” Fertility and Sterility, vol. 85, no. 5, pp. 1448–1451, 2006.

S. Palomba, A. Falbo, and G. B. La Sala, “Metformin and gonadotropins for ovulation induction in patients with polycystic ovary syndrome: a systematic review with meta-analysis of randomized controlled trials,” Reproductive Biology and Endocrinology, vol. 12, article 3, 2014.

R. G. Farquharson, E. Jauniaux, and N. Exalto, “Updated opinion on drug safety,” Human Reproduction, vol. 20, no. 11, pp. 3008–3011, 2005.

H. B. Ford and D. J. Schust, “Recurrent pregnancy loss: etiology, diagnosis, and therapy,” Reviews in Obstetrics and Gynecology, vol. 2, pp. 76–83, 2009.
[165] J. S. Dhulkotia, B. Ola, R. Fraser, and T. Farrell, “Oral hypoglycemic agents vs insulin in management of gestational diabetes: a systematic review and metaanalysis,” *American Journal of Obstetrics & Gynecology*, vol. 203, no. 5, pp. 457.e1–457.e9, 2010.

[166] J. C. Silva, C. Pacheco, J. Bizato, B. V. De Souza, T. E. Ribeiro, and A. M. Bertini, “Metformin compared with glyburide for the management of gestational diabetes,” *International Journal of Gynecology and Obstetrics*, vol. 111, no. 1, pp. 37–40, 2010.

[167] Y. Barak, M. C. Nelson, E. S. Ong et al., “PPARγ is required for placental, cardiac, and adipose tissue development,” *Molecular Cell*, vol. 4, no. 4, pp. 585–595, 1999.

[168] J. Sevillano, I. C. López-Peréz, E. Herrera, M. Del Pilar Ramos, and C. Bocos, “Englitazone administration to late pregnant rats produces delayed body growth and insulin resistance in their fetuses and neonates,” *Biochemical Journal*, vol. 389, no. 3, pp. 913–918, 2005.

[169] P. Froment and P. Touraine, “Thiazolidinediones and fertility in polycystic ovary syndrome (PCOS),” *PPAR Research*, vol. 2006, Article ID 73986, 8 pages, 2006.

[170] J. Gui, Q. Liu, and L. Feng, “Metformin vs insulin in the management of gestational diabetes: a meta-analysis,” *PLoS ONE*, vol. 8, no. 5, Article ID e64585, 2013.

[171] V. W. Wong and B. Jalaludin, “Gestational diabetes mellitus: who requires insulin therapy?” *Australian and New Zealand Journal of Obstetrics and Gynaecology*, vol. 51, no. 5, pp. 432–436, 2011.

[172] A. Sokup, B. Ruszkowska-Ciastek, K. Góralczyk, M. Walentowicz, M. Szymański, and D. Rośc, “Insulin resistance as estimated by the homeostatic method at diagnosis of gestational diabetes: estimation of disease severity and therapeutic needs in a population-based study,” *BMC Endocrine Disorders*, vol. 13, article 21, 2013.

[173] A. Corbould, F. Swinton, A. Radford, J. Campbell, S. McBeath, and A. Dennis, “Fasting blood glucose predicts response to extended-release metformin in gestational diabetes mellitus,” *Australian and New Zealand Journal of Obstetrics and Gynaecology*, vol. 53, pp. 125–129, 2013.

[174] C. P. Spaulonci, L. S. Bernarides, T. C. Trindade, M. Zugaib, and R. P. Francisco, “Randomized trial of metformin vs insulin in the management of gestational diabetes,” *American Journal of Obstetrics & Gynecology*, vol. 209, no. 1, pp. 34.e1–34.e7, 2013.

[175] K. Tertti, U. Ekblad, P. Koskinen, T. Vahlberg, and T. Rönnemaa, “Metformin vs. insulin in gestational diabetes. A randomized study characterizing metformin patients needing additional insulin,” *Diabetes, Obesity and Metabolism*, vol. 15, no. 3, pp. 246–251, 2013.

[176] Identifier NCT01587378, Metformin to Prevent Late Miscarriage and Preterm Delivery in Women with Polycystic Ovary Syndrome Trial (PregMet2), ClinicalTrials.gov, Bethesda, Md, USA, National Library of Medicine, 2012, http://clinicaltrials.gov/show/NCT01855763.

[177] National Library of Medicine, “Metformin treatment in gestational diabetes and noninsulin dependent diabetes in pregnancy in a developing country (migdm&t2dm),” ClinicalTrials.gov NCT01855763, National Library of Medicine, Bethesda, Md, USA, 2013, http://clinicaltrials.gov/show/NCT01855763.