What have we learned from monogenic forms of severe insulin resistance associated with PCOS/HAIRAN?

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Insulin resistance and polycystic ovary syndrome (PCOS) have important but complex pathophysiological relationships. Both conditions are associated with type 2 diabetes and cardiovascular risks. However, obesity, which is an important contributor to these diseases, is a frequent confounding factor.

Severe insulin resistant states are observed in rare but very diverse diseases, of genetic or acquired origin, ranging from major dysfunctions of insulin receptor to abnormalities in lipid droplets formation and syndromes of body fat redistribution, in the absence of obesity. Hyperandrogenemia and PCOS are common findings in women with severe insulin resistance syndromes, whatever their causes.

Reports of the ovarian phenotypes in women with monogenic forms of severe insulin resistance syndromes have stimulated many studies about the roles of the ovarian insulin-insulin growth factor (IGF) system. In addition, PCOS features could be the referring signs leading to the diagnosis of several syndromes of severe insulin resistance.

**Primary dysfunctions of insulin receptors and PCOS**

The first clinical evidence for an important role of insulin in regulating ovarian functions came from the identification by R. Kahn of severe insulin resistance syndromes with acanthosis nigricans and virilization in 1976 (1). The patients were
described with hirsutism, polycystic ovaries, clitoral enlargement, and for some of them accelerated early growth and coarse figures (1). Therefore, it was proposed that insulin could enhance androgen production by the ovary. This hypothesis was reinforced later by the demonstration of insulin’s ability to stimulate steroidogenesis in ovarian cells in vitro, and the identification of insulin receptors in stromal and follicular compartments of the human ovary (for review, see (2)). However, defects in the insulin signalling are located at the receptor level in these diseases, leading to insulin resistance and diabetes. Indeed, insulin receptor gene mutations or auto-antibodies have then been shown to be responsible for type A and type B insulin resistance syndromes, respectively (3, 4). The ovary expressing not only insulin receptors, but also type I and type II IGF receptors (2), the most likely hypothesis is that the major hyperinsulinemia observed in type A insulin resistance syndromes acts on the ovary through the stimulation of IGF receptors. In addition, in severe insulin resistance syndrome, hyperinsulinemia is responsible for a decrease in the liver production of sex hormone binding globulin (SHBG) and IGF1 binding protein-1 (IGFBP-1), further enhancing the increase in free testosterone and IGF1 levels.

Concerning type B insulin resistance syndromes, auto-antibodies directed against insulin receptors could have complex effects, with inhibition and/or activation of the different insulin signalling pathways (5), that could differently modulate insulin-mediated effects on the ovary.

**Lipodystrophic syndromes and PCOS**

Other syndromes of severe insulin resistance are associated with PCOS features. It is the case of lipodystrophic syndromes, either generalized, partial, genetic, or acquired (6). Recent studies have evidenced new molecular defects involved in these diseases (7-16), in favour of a primary role of adipose tissue development, and in particular of adipocyte lipid droplets formation, in generalized congenital lipoatrophies. Insulin resistance is thought to be secondary to the defective lipid storage in lipodystrophic adipose tissue, with post-receptor defects in insulin-mediated signalisation events. In accordance, acromegaloid features (17) and liver steatosis (18) associated with severe insulin resistance syndromes have been explained by partial insulin resistance affecting glucose-lowering effects of insulin, but not insulin-activated mitogenic signals nor hepatic de novo lipogenesis,
respectively. In accordance, a recent studies revealed an association between polymorphisms in two components of the insulin signalling pathways, AKT2 and GSK3β, and PCOS (19). In addition, post-binding defects in insulin receptor signalling have been evidenced in common forms of PCOS (20, 21).

Whatever the origin of severe insulin resistance (primary insulin receptors defects or lipodystrophies), in vivo hyperinsulinemia has been clearly shown to promote ovarian growth and androgen synthesis independently of gonadotropins (22, 23).

**Endocrine defects of adipose tissue and PCOS**

In lipodystrophic syndromes, the endocrine deficiency of adipose tissue has been shown to play important pathophysiological roles in metabolic alterations. In particular, adiponectin and leptin are severely decreased, contributing to the ectopic lipid storage in non-adipose cells which inhibits insulin signalling (lipotoxicity). Leptin replacement therapy has been shown to restore menstrual functions in lipodystrophic women, which could be due to improvements in both insulin sensitivity and LH pulsatility (24, 25). However, the morphologic polycystic ovarian pattern did not change after leptin treatment. Conversely, the leptin replacement therapy of young hypoleptinemic children with congenital generalized lipodystrophic syndromes did not modify the onset of puberty (26). In addition, adiponectin deficiency probably does not play an important primary role in PCOS associated with severe insulin resistance syndromes. Indeed, adiponectin is decreased in lipodystrophies, but is significantly increased in patients with primary defects of insulin receptors (mutations or auto-antibodies) as compared to controls, although the mechanisms are not known precisely (27, 28).

Finally, PCOS features are not always present in insulin resistance syndromes (29-31). This is in favour of an aggravating, but not a primary role of insulin resistance on ovary dysfunctions.

Conversely, although the PCOS phenotype show a considerable heterogeneity in metabolic features, hirsutism and oligomenorrhea were shown to be the referring signs in several cases of severe insulin resistance syndromes (32, 33).
Conclusion
Monogenic forms of severe insulin resistance associated with PCOS have highlighted the involvement of hyperinsulinemia in the ovarian pathophysiology. Although the pathogenic mechanisms are probably multifactorial, these diseases show the major contributor role for insulin resistance in ovarian dysfunction leading to PCOS. Conversely, at the clinical level, the careful characterization of patients with PCOS features can lead to diagnosis of rare severe insulin resistance syndromes, with and without lipodystrophy.

References

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