Steroidogenic profiles of males with congenital hypogonadotropic hypogonadism on hCG/rFSH and on testosterone replacement

Julia Rohayem1, Paul-Martin Holterhus2, Sandra Laurentino1, Sabine Kliesch2, Eberhard Nieschlag1, Michael Zitzmann1, Alexandra Kulle2
1Center of Reproductive Medicine and Andrology, Department of Clinical and Surgical Andrology, University of Münster, Germany
2Children’s Hospital Kiel, Department of Pediatric Endocrinology and Diabetes, University of Schleswig-Holstein, Germany

INTRODUCTION

In males with congenital hypogonadotropic hypogonadism (CHH), LH/rFSH stimulation of gonads is deficient. Two hormone replacement strategies are employed to induce and maintain virilisation: substitution of testosterone and gonadotropin replacement with hCG/rFSH.

AIMS

We aimed to delineate the role of gonadotropins in pathways of male androgen biosynthesis

PATIENTS AND METHODS

In 25 males with CHH, first undergoing hCG/rFSH treatment and then testosterone replacement, serum steroid hormone profiles (precursors of testosterone and its metabolites) were analyzed, using liquid chromatography-tandem mass spectrometry (LC-MS/MS). Data were compared to those of healthy controls, matched for age, BMI and serum testosterone (T) concentration.

RESULTS

Combined treatment of CHH males with hCG and rFSH resulted in steroid hormone profiles similar to those of healthy men, but this was not the case, while a regimen based on exogenous testosterone was used:

While CHH patients were on T substitution, decreased serum concentrations of some members of the classical Δ4 pathway of androgen biosynthesis (progesterone (p=0.0104), 17-OH-progesterone (17 OHP) (p=0.0031)) and of the alternative T pathway steroid androstenediol (p=0.0044) were observed, compared to controls. The marker steroid of the backdoor DHT pathway androstenediol (p=0.025), was slightly increased. Some Δ4 pathway steroids (androstenedione, (A4)), the testosterone metabolites DHT and 17-β estradiol (E2), the Δ5 pathway steroid dehydroepiandrosterone sulfate (DHEAS) and all measured 11-oxygenated C19 androgens (11-keto-testosterone (11 K T), 11-keto-dihydro-testosterone (11 K DHT) and 11-keto-androstenedione (11 K A4)) were comparable to those of controls.

By contrast, on hCG/rFSH replacement, steroid profiles resembled those of healthy male controls, regarding the Δ 4 pathway of androgen biosynthesis (17-OHP, A4) and the metabolite DHT, the marker steroid of the alternative T pathway androstenediol, the Δ5 pathway steroid DHEAS and all aforementioned 11-oxygenated C19 androgens (11 K A4, 11 K T, 11 K DHT). Serum progesterone (Δ 4 pathway) was slightly decreased (p=0.0104); the testosterone metabolite E2 and the backdoor DHT pathway steroid androstenediol were increased (both p<0.0001).

CONCLUSIONS

In males with CHH, a replacement with hCG/rFSH mimics physiologic steroid hormone profiles better than a substitution with testosterone. Gonadotropins induce Δ4 classic pathway steroid production and co-activate the alternative pathway of T biosynthesis. The backdoor pathway of DHT, synthesis of DHEAS and of 11-oxygenated C19 steroids are activated independently of gonadotropins. The documented differences in replacement strategies may impact on long term male health.