Gonadotropin-Dependent Pubertal Disorders are Common in Patients with Virilizing Adrenocortical Tumors in Childhood

Authors:

Monica F. Stecchini¹, Zilda Braid¹, Candy B. More¹, Davi C. Aragon¹, Margaret Castro², Ayrton C. Moreira², Sonir R. Antonini³

Authors’ Affiliations:

Departments of ¹Pediatrics and ²Internal Medicine, Ribeirao Preto Medical School, University of Sao Paulo. Ribeirao Preto – SP, Brazil.

Corresponding Author: Sonir R. Antonini, MD, PhD. Departamento de Pediatria, Faculdade de Medicina de Ribeirao Preto, Universidade de Sao Paulo. Avenida Bandeirantes, 3900 Monte Alegre, CEP 14049-900, Ribeirao Preto, Sao Paulo, Brazil.
E-mail: antonini@fmrp.usp.br

Abbreviated Title: Precocious Puberty in Children with Adrenocortical Tumors

Keywords: adrenocortical tumor, gonadotropin-dependent precocious puberty, early puberty, final height.

Word count: 3,981
ABSTRACT

OBJECTIVE: To investigate the impact of early exposure to androgen excess on gonadotropin-dependent puberty (GDP) and final height (FH) of patients with androgen-secreting adrenocortical tumors (ACT) in childhood. METHODS: Retrospective cohort study. Occurrence of GDP and achievement of FH were evaluated. Central precocious puberty (CPP) and early fast puberty (EFP) were considered pubertal disorders. Patients with normal puberty and pubertal disorders were compared. RESULTS: The study included 63 patients (44F), followed in a single institution from 1975 until 2017. At diagnosis of ACT, median age was 25.8 months; duration of signs, six months; stature SDS, 0.5 (-3.6 – 3.9); and bone age advancement, 14.7 months (-27.9 – 85.4). To date, 37 patients developed GDP: 26 had normal puberty; one, precocious thelarche; seven, CPP; and three, EFP. GnRHa effectively treated CPP/EFP. Tall stature and older age at diagnosis of ACT were associated with risk of CPP alone [RR 4.17 (95% CI 1.17 – 14.80)] and CPP/EFP [RR 3.0 (95% CI 1.04 – 8.65)]. Recurrence/metastasis during follow-up were associated with risk of CPP alone [RR 4.17 (95% CI 1.17 – 14.80)] and CPP/EFP [RR 3.0 (95% CI 1.12 – 8.02)]. Among the 19 patients that reached FH, stature SDS dropped from 1.4 to -0.02 since diagnosis of ACT (p=0.01). Seventeen achieved normal FH. There was no difference in FH SDS between patients with normal puberty and pubertal disorders (p=0.75). CONCLUSIONS: Gonadotropin-dependent pubertal disorders are common in patients with androgen-secreting ACT in childhood. FH is usually not impaired. The study reinforces the importance of close follow-up after surgery to identify and treat consequences of early exposure to androgen excess.
INTRODUCTION

Gonadotropin-dependent precocious puberty or central precocious puberty (CPP) is caused by re-activation of the hypothalamic-pituitary-gonadal axis (HPG), while gonadotropin-independent precocious puberty or peripheral precocious puberty (PPP) is caused by autonomous secretion of sex steroids from ovaries, testicles or adrenals, before the age of eight years in girls and nine years in boys (1). In some circumstances, PPP can trigger CPP through mechanisms that are not well understood, but that may be related to the skeletal maturation induced by androgen excess (2).

The development of secondary CPP has been extensively reported in patients with lately diagnosed or poorly controlled congenital adrenal hyperplasia (CAH), and other conditions associated with androgen excess, usually after initiation of treatment of the underlying disease (3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16). In this context, androgen-secreting adrenocortical tumors (ACTs), which usually present with virilizing features, linear growth acceleration and bone age (BA) advancement (17), could also lead to secondary CPP. However, this outcome has been considered uncommon in ACTs (18).

ACTs are rare in children worldwide, with an overall incidence of 0.2 – 0.4 cases per million per year (17). However, their incidence is 10 – 15 times higher in Southern and Southeastern Brazil (19), where 75 – 97% of the patients harbor a specific P53 mutation. The P53 p.R337H is the most significant and well characterized genetic risk for ACT in these regions; it is widespread due to a founder effect (20, 21, 22, 23, 24). This mutation was initially described in association with ACT in Brazilian children (21), but it has been shown to be associated with the development of other types of cancer in adulthood, in the context of Li-Fraumeni syndrome (25, 26).
Pediatric ACTs are more commonly diagnosed in early childhood (< 4 years) and predominantly affect females (19, 23). The majority of these tumors secrete hormones, especially androgens and cortisol, and the most common clinical presentation is a virilizing syndrome – isolated or associated with Cushing syndrome (17). The diagnosis of ACT is suggested by clinical, laboratory and imaging features, and confirmed by pathological findings after tumor resection (17). In the pediatric population, imaging criteria and histologic classification systems used to distinguish benign from malignant tumors for adult patients do not reflect the future tumor behavior and the patient’s outcome. Therefore, the term “ACT” is preferred and represents the sum of adenomas and carcinomas (17, 27, 28). Open adrenalectomy is the treatment of choice, which may be complemented with mitotane and chemotherapy, based on tumor staging (29).

Although morbidity (metastasis, recurrence and complications of ACT treatment) and mortality in patients with ACTs are well documented (17, 19, 23, 24), the late effects of early exposure to androgen excess on gonadotropin-dependent puberty and on growth have been scarcely presented in case reports (30, 31, 32, 33), in small series of patients (18, 34) or in studies with other purposes (19, 24).

We hypothesized that children with androgen-secreting ACTs may be at increased risk of developing early activation of the HPG axis, and that final height (FH) may not be impaired if adequate treatment is used. Therefore, the objective of this study was to investigate the impact of early exposure to androgen excess on gonadotropin-dependent pubertal development and on FH of patients treated for androgen-secreting ACTs during childhood, in a single reference center, over the period of four decades.

SUBJECTS AND METHODS
Study design and Patients

This was a retrospective cohort study, performed at the Hospital of Ribeirao Preto Medical School from the University of Sao Paulo. The study was approved by this institution’s ethics committee and conducted in compliance with the Declaration of Helsinki.

Records from patients followed at this institution, from September 1975 until September 2017, were reviewed. Inclusion criteria was: surgical treatment of an androgen-secreting ACT before the age of 18 years. Exclusion criteria was: presence of other conditions that also influence growth and pubertal development.

Epidemiological, clinical, anthropometrical, hormonal, radiological, histopathological and genetic data were collected from the eligible patients’ medical records.

Descriptive data

The age at diagnosis of ACT and the age of tumor resection were considered the same – and presented as “age at diagnosis of ACT” – because surgery was performed shortly after the clinical / biochemical / radiological diagnosis of ACT in all cases.

The duration of ACT signs reflected the period between the family’s perception of clinical signs until the diagnosis of ACT.

The main clinical presentation was divided as: asymptomatic; unspecific signs – weight loss, fever, abdominal mass or pain; virilizing syndrome – precocious pubarche, clitoris or penile enlargement, axillary hair and odor, acne, muscular hypertrophy, aggressiveness – denoting androgen excess; and Cushing syndrome – weight gain, central obesity, facial plethora, violet striae – denoting cortisol excess (17).

Height and body mass index (BMI) were expressed in standard deviation scores (SDS), according to the World Health Organization (WHO) curves for children of 0 – 5 years
(2006) and 5 – 19 years (2007), using the statistical software LMSgrowth 2002-12 Medical Research Council, UK. Short stature was defined as stature < -2 SDS and tall stature, as stature > +2 SDS (35). FH was considered when BA was 15 and 17 years for females and males, respectively. Target height and range were calculated for girls as \([\text{mother’s stature} + \text{father’s stature} – 13]/2 \pm 9\) and for boys as \([\text{father’s stature} + \text{mother’s stature} + 13]/2 \pm 10\), in centimeters (36). Obesity was defined as BMI > +3 SDS in children from 0 – 5 years or > +2 SDS in children from 5 – 19 years, while overweight was defined as BMI > +2 SDS in children from 0 – 5 years or > +1 SDS in children from 5 – 19 years.

Pubertal stage was described according to Marshall and Tanner classification (37, 38). BA was determined by radiologists and confirmed by the attending endocrinologists, based on the Greulich-Pyle method (39). It was presented as delta \((\Delta)\) BA (difference between BA and chronological age) and classified as advanced when \(\Delta\) BA was > +1 SDS for sex and chronological age (1).

In terms of hormone production, ACTs were categorized as pure androgen secreting tumors (virilizing tumors or V-ACT), when androgen excess was isolated, or androgen and cortisol secreting tumors (mixed tumors or M-ACT), when androgen excess was combined with autonomous cortisol secretion, confirmed by a dexamethasone suppression test (17). Plasma concentration of dehydroepiandrosterone sulfate (DHEA-S), androstenedione, testosterone and 17-hydroxyprogesterone (17-OHP), as well as cortisol (17), was determined by immunoassays.

The diagnosis of ACT was suggested by clinical, laboratory and imaging features, and confirmed by pathological findings after tumor resection.

Tumor stage was established according to the International Pediatric Adrenocortical Tumor Registry (IPACTR) criteria, as I, II, III or IV (29).
Sanger sequencing of TP53 exon 10 was routinely performed for patients with the diagnosis of ACT to investigate the presence of the P53 p.R337H mutation (17).

**Outcomes and Predictors**

The development of gonadotropin-dependent pubertal disorders was defined as the primary outcome of this study. CPP was defined as the presence of breast in girls (Tanner stage 2 – thelarche) before eight years and of testicular enlargement (≥ 4 cm³) in boys before nine years, with progression of pubertal signs, secondary to activation of the HPG axis. Early puberty (EP) – although within the normal pubertal range – was defined as the presence of breast between 8 and 9 years in girls and of testicular enlargement between 9 and 10 years in boys. Fast puberty (FP) was defined as progression from one pubertal stage to another in less than 6 months (1). The combination of EP and FP was named early fast puberty (EFP) (40, 41). Precocious menarche was defined as cyclic vaginal bleeding before 9 years (1). CPP and EFP were considered pubertal disorders.

Sex, age, duration of ACT signs, Tanner stage for pubic hair, stature SDS, Δ BA, tumor type, DHEA-S and testosterone levels, and tumor stage at diagnosis of ACT, recurrence and/or metastasis during follow-up, as well as FH, were compared between patients with normal puberty and with pubertal disorders.

**Statistical analysis**

Quantitative variables were expressed as median and range, if continuous, or as absolute number and fractions, if discrete.

Gonadotropin-dependent pubertal development was analyzed based on the comparison of two groups: normal puberty (normal onset + EP without FP) and pubertal
disorders (CPP + EFP). Relative risk (RR) was used to investigate associations, based on categorical variables: sex (female or male), age (≥ 4 or < 4 years), duration of ACT signs (≥ 6 or < 6 months), Tanner stage for pubic hair (> 2 or ≤ 2), stature SDS (> 2 or ≤ 2), bone age (advanced or not advanced), tumor type (V-ACT or M-ACT), tumor stage (III/IV or I/II), DHEA-S and testosterone levels (both: ≥ 300 or < 300 ng/dL) at diagnosis of ACT, and recurrence/metastasis during follow-up (present or absent). The definition of these categories was based on the assumption that longer and more severe exposure to androgen excess would increase the risk of pubertal disorders. The results were presented as RR, with 95% confidence intervals (CI).

Wilcoxon rank-sum test was used to compare continuous variables between groups. Wilcoxon signed-rank test was used to compare initial and final stature SDS of patients that reached FH. For the hypothesis tests, significance level of $\alpha = 0.05$ was used.

The software STATA 15.1 was used for statistical analysis.

RESULTS

Subjects

From September 1975 until September 2017, 70 patients with the diagnosis of ACT before the age of 18 years were followed at the Hospital of Ribeirao Preto Medical School, University of Sao Paulo – which is located in the Southeast region of Brazil. Of these, the records of six female patients were not included due to: lack of surgical treatment (n=1), lack of proper hormonal assessment before diagnosis of ACT (n=3) and presence of isolated cortisol excess (n=2). Additionally, the record of one female patient was excluded due to the presence of an overgrowth syndrome – Beckwith-Wiedeman syndrome (n=1). Therefore,
the records of 63 patients with the diagnosis of an androgen-secreting ACT, both V-ACT and M-ACT, before the age of 18 years were eligible for this study (Figure 1).

**General characteristics**

Table 1 shows the general features of the 63 patients included in the study. The majority were female (44/63) and white (58/63), with median age at diagnosis of ACT of 25.8 months (2.1 – 192.2). Median duration of ACT signs was six months (0 – 60): < 6 months for 30 patients, between 6 and 12 months for 19 patients, between 12 and 24 months for 9 patients, and ≥ 24 months for 5 patients. Family history of ACT was present in 8/63 and of other types of cancer in 1/3 of the patients.

The majority of patients (54/63) presented clinically with a virilizing syndrome, seven also had Cushing syndrome and two patients were asymptomatic. In these two patients, the diagnosis was made because of significant family history of ACT and presence of the P53 p.R337H mutation – which motivated hormone assessment – in one case, or because of altered 17-OHP in the neonatal screening for CAH – which is an unusual form of diagnosis – in the other. However, hormonal assessment revealed that the majority of the tumors (55/63) was actually M-ACT.

Median SDS for stature and for BMI at diagnosis of ACT was 0.50 (-3.55 – 3.89) and 1.38 (-1.88 – 6.33), respectively. Eleven patients had tall stature and two, short stature. Twelve patients had obesity and nine, overweight. There was no difference regarding stature SDS (p=0.38) or BMI SDS (p=0.18) between patients with V-ACT [stature SDS: 1.24 (-0.08 – 1.97); BMI SDS: 0.55 (-0.63 – 2.45)] and patients with M-ACT [stature SDS: 0.48 (-3.55 – 3.89); BMI SDS: 1.41 (-1.88 – 6.33)].
Median DHEA-S was 748.0 µg/dL (32.0 – 4950.0) [19.5 (0.8 – 128.7) µmol/L] and median testosterone was 242.5 ng/dL (33.6 – 1800.0) [8.4 (1.2 – 62.4) nmol/L].

BA was available for 52 out of the 63 patients: it was advanced in 33, adequate for chronological age in 17 and delayed in two patients. In general, BA was 14.7 months advanced (-27.9 – 85.4). Median Δ BA was 17.5 months (0.4 – 49.1) for patients with V-ACT and 11 months (-27.9 – 85.4) for patients with M-ACT (p=0.84).

Tumor stage I was the most frequent (38/63), followed by III (11/63), II (8/63) and IV (6/63). The mutational status was available for 52 out of the 63 patients: 48/52 harbored the P53 p.R337H mutation.

Median duration of follow-up was 73.7 months (0.2 – 295.4). During this period, 16 patients had recurrence and/or metastasis, 17 were treated with chemotherapy (mitotane; combination of other chemotherapy agents – mainly cisplatin, doxorubicin, etoposide; or mitotane associated with these agents) and only three patients received radiotherapy (doses were not available). Fourteen patients died (all deaths were related to ACT complications), 16 lost follow-up and five were discharged after 10 years. Of note, the patients that were discharged after 10 years had the diagnosis of ACT in 1984 – 1993. At the time, the risk of developing other types of cancer in adulthood was less known and there was no specific long-term follow-up protocol for these patients and families.

**Pubertal outcomes**

*Figure 2* shows the pubertal outcomes among the 63 study patients. To date, 37 patients developed gonadotropin-dependent puberty: 26 developed puberty within normal age range, one (F) had isolated precocious thelarche, seven (4F: 3M) had CPP, and three (F) had EFP. Median time and range (in months) between diagnosis of ACT and pubertal onset...
were: for normal puberty, 83.8 (-8.8 – 103.5) in females and 102.4 (64.4 – 113.6) in males; for CPP, 34.9 (3.8 – 73.8) in females and 2.7 (-2.1 – 3.1) in males; and for EFP, 40.0 (0.5 – 94.7). One boy and one girl had developed normal puberty before diagnosis of ACT, but the accurate age was not available. None of the patients had delayed puberty.

Of the 26 patients that did not develop gonadotropin dependent puberty: 12 are still on follow-up (current age: 43.5 months [19 – 99]), while eight died (age: 65.5 months [12 – 146]) and six lost follow-up (age: 46 months [23 – 93]) before developing gonadotropin-dependent puberty.

Regarding gonadotropin-releasing hormone analog (GnRHa) treatment, among the three CPP female patients who were treated, one had menarche previously to treatment, at 8y5mo, and died before the end of GnRHa treatment, due to ACT advanced disease stage. Another had menarche at 12y4mo, and the last one is currently nine years old, is still on medication and had no menarche yet. One CPP female patient received no treatment and had menarche at 10y3mo. One boy with CPP was treated for eight years, until age 12. The other two boys with CPP received no treatment, because of multiple ACT-related complications. All three girls with EFP were treated with GnRHa. Menarche occurred in one at 12y5mo, one was lost to follow up, and one had no menarche yet.

Additionally, one girl had isolated precocious thelarche at 7y3mo, but had no rapidly progressive puberty and achieved menarche at 13y1mo. Of note, one girl, currently 7y11mo, did not develop puberty yet, but had an isolated precocious vaginal bleeding at age 7y3mo.

Table 2 presents the general features of the 10 patients with pubertal disorders and the 26 with normal puberty. The patient with isolated precocious thelarche was not included in the comparisons. Patients with CPP and EFP tended be older (p=0.07) and have
more advanced bone age ($p=0.05$) at diagnosis of ACT, but there was no difference regarding duration of ACT signs ($p=0.86$), stature SDS ($p=0.59$), or DHEA-S and testosterone levels ($p=0.89$ and $p=0.43$, respectively).

At diagnosis of ACT, tall stature was associated with increased risk of developing CPP [RR 4.17 (95%CI 1.17 – 14.80)], and older age, with increased risk of developing pubertal disorders [RR 3.0 (95%CI 1.04 – 8.65)]. During follow-up, the presence of recurrence and/or metastasis was associated with the development of CPP alone [RR 4.17 (95%CI 1.17 – 14.80)] and also with pubertal disorders in general [RR 3.0 (95%CI 1.12 – 8.02)]. There was no association regarding duration of ACT signs, Tanner stage for pubic hair, BA advancement, tumor type, androgen levels, or tumor stage at diagnosis of ACT (Tables 3 and 4).

**Final height**

FH was reached in 19 out of 63 patients (8M: 11F). Among these, two patients had adult short stature, with no available target height; none of them had had pubertal disorders. All other patients achieved normal stature; the 10 (5M: 5F) for whom target height was available reached their genetic potential. Among these 19 patients, stature SDS dropped from 1.42 at diagnosis of ACT to -0.02 at FH ($p=0.013$) (Figure 2).

Of these 19 patients, 14 had normal puberty, three had CPP, one had EFP, and one had precocious thelarche. There was no difference in FH SDS between patients with normal puberty and pubertal disorders ($p=0.75$) (Figure 2).

**DISCUSSION**
There are scanty data on the late consequences of early exposure to high androgen levels on pubertal development and FH of patients with childhood ACTs. This retrospective cohort study, performed in a single reference center, clearly shows that gonadotropin-dependent pubertal disorders are common in patients who previously had androgen-secreting ACTs during childhood. Additionally, it confirms that FH is usually not impaired in this group of patients.

The general features of the patients from this cohort were similar to previously published series (19, 24, 34, 42, 43). The majority were female (44/63), had the diagnosis of ACT before the age of 4 years (44/63), with clinical presentation of a virilizing syndrome (54/63), and harbored the P53 p.R337H mutation (48/52). Despite the predominance of clinical signs of androgen excess, routine assessment of cortisol autonomous secretion (17) revealed that most ACTs (55/63) had a mixed pattern of hormone secretion. All patients underwent open surgery, and some had adjuvant chemotherapy, based on tumor stage, as it is preconized (29). Of note, radiotherapy is not routinely recommended for ACT patients, but it was used for three patients from 1986 – 1988, when mitotane and other chemotherapy agents were not easily available.

The development of CPP after prolonged androgen exposure, especially after its withdrawal, has been well documented in CAH (3, 4, 5, 6), Leydig cell tumors (8, 9, 10, 11, 12, 13, 14, 15), familial male-limited precocious puberty (FMPP) (7) and even in interpersonal transfer of testosterone gel (16). The mechanisms that trigger secondary HPG axis activation are not well understood, but the temporal correlation between skeletal maturation and HPG axis maturation may be a clue (44). A study on male patients with CAH or FMPP showed an association between the most severe skeletal advancement and the earliest age of gonadotropin-dependent puberty (44). In another study, CAH patients with...
more advanced BA, more rapid linear growth and taller stature during the first two years of life also developed CPP (5). The occurrence of gonadotropin-dependent pubertal disorders mainly after treatment of the underlying disorder may be explained by the withdraw of a peripheral inhibitor of HPG axis, such as testosterone, in a patient with advanced BA (31). It is known that the decrease in testosterone levels during therapy may lead to secondary CPP in boys with FMPP, for example (7).

In ACTs, the development of CPP has been considered rare (18). There are only scarce studies reporting gonadotropin-dependent pubertal disorders after treatment of ACT. These studies are either small series (18, 31, 33, 34) or larger series in which the number of patients that reached pubertal age is not stated; therefore, the proportion of pubertal disorders cannot not be accurately calculated (19, 24). In the present study, this issue was addressed in a straight line. Indeed, 37 out of 63 patients developed gonadotropin-dependent puberty, and, among them, 10/37 had pubertal disorders. Thus, for the first time, a great number of ACT patients were followed until puberty, allowing us to clearly demonstrate that gonadotropin-dependent pubertal disorders are common in patients that had androgen-secreting ACTs during childhood.

In the present study, tall stature at diagnosis of ACT was associated with the development of CPP, and more patients with pubertal disorders had advanced BA at diagnosis of ACT, compared with those with normal puberty. In accordance to these data, the patients that developed CPP or EFP from previous reports (18, 31, 33, 34) had tall stature and advanced BA at diagnosis of ACT. Although the duration of ACT signs before the diagnosis of ACT was not associated with the development of pubertal disorders, this study demonstrated that recurrence and/or metastasis increased the risk of CPP, as well as of pubertal disorders in general. These conditions may lengthen the period of androgen
exposure, contributing to the skeletal maturation and further HPG axis activation. At last, there was an association between older age at diagnosis of ACT and the development of pubertal disorders, which may be explained by the proximity to the actual pubertal age. Indeed, a study of hypothalamic stimulation with clomiphene citrate in six children with CAH and one boy with ACT showed that luteinizing hormone response increases with chronological age and, more particularly, with BA (45).

GnRHa have been shown to be effective for the treatment of secondary CPP, and their indications include prevention of precocious menarche in girls and adult short stature, as well as psychosocial maladjustment, in both sexes (1, 2). In the present study, GnRHa effectively treated the patients with gonadotropin-dependent pubertal disorders. Menarche occurred at an appropriate age for the treated patients, and the one with an episode of vaginal bleeding before GnRHa treatment presented no recurrence. Additionally, treated patients reached normal FH, within the genetic potential. However, due to the small number of patients with pubertal disorders that reached FH, comparison between treated and not treated patients was not possible. Treatment with GnRHa or cyproterone acetate had only been reported in three cases of CPP in patients with ACT, but there is no information on menarche or FH of these patients (18, 31, 33).

It is well known that hyperfunction of the adrenal cortex in childhood, with elevated secretion of androgens and glucocorticoids, may cause significant disturbance of linear growth. While androgen excess advances growth and BA, glucocorticoid excess impairs skeletal maturation (46). In the presence of both, however, the androgen effects appear to dominate the clinical features (47). This predominance may explain why the majority of patients in this cohort had normal (79.3%) or tall stature (17.5%) and advanced BA (63.5%) at diagnosis of ACT, although most had M-ACT. This finding is in accordance with previous
reports (18, 19, 24, 30, 31, 32, 34, 42, 43, 47). In addition, the combination of androgen and glucocorticoid effects may have accounted for the lack of difference in initial stature SDS, BMI SDS and ∆ BA between V-ACT and M-ACT. Despite the acceleration of linear growth and skeletal maturation at diagnosis of ACT, there was a reduction of stature SDS from initial to final height, and the majority of patients (17 out of 19) reached normal FH. These results reinforce that growth tends to return to the patient’s predetermined channel after successful removal of a virilizing ACT (47), with good prognosis on FH (18, 19, 30, 32).

The retrospective character of this study imposed some limitations. Missing data may have weakened the statistical analysis. The duration of follow-up was different among the study subjects and many did not develop gonadotropin-dependent puberty and / or reach FH because they were still young, died or lost follow-up at the time of the study. Additionally, BA radiographies were not reviewed in a blinded fashion by a single reader, since many of the images were unavailable. However, to our knowledge, this is the largest cohort of ACT patients from a single center evaluated for pubertal and FH outcomes.

In conclusion, this study clearly shows that gonadotropin-dependent pubertal disorders are more common than previously expected in patients with childhood virilizing ACTs. Additionally, it confirms that FH is usually not impaired, reiterating the good prognosis for FH in these patients. At last, it reinforces the importance of close and prolonged endocrinology follow-up after surgery, not only to detect ACT-related complications, but also to promptly identify and treat consequences of early exposure to androgen excess.

DECLARATION OF INTEREST

The authors report no conflict of interest in this work.
**FUNDING**

This work was supported by FAPESP (Sao Paulo State Research Agency; Grants #2015/19663-5 and #2014/03989-6). M.F.S. and C.B.M. are recipients of research fellowships from CAPES and FAPESP – Brazil, respectively.

**ACKNOWLEDGMENTS**

The authors are grateful to all medical, health care, laboratory and administrative personnel who were involved in the care of these patients over the last four decades in our institution. In addition, we thank all patients and their families for their trust in our service.

We dedicate this work to Professor Romulo Sandrini (Federal University of Parana), whose work with pediatric ACT patients has been an inspiration. In addition, we dedicate this work to the memory of Professor Jose T. Verissimo, who first diagnosed and treated pediatric ACT patients in our institution in the early 1960's.
REFERENCES

1. Brito V, Spinola-Castro A, Kochi C, Kopacek C, Silva P, & Guerra-Júnior G. Central precocious puberty: revisiting the diagnosis and therapeutic management. *Archives of Endocrinology and Metabolism* 2016, 60, 163–172. (doi:10.1590/2359-3997000000144)

2. Fuqua JS. Treatment and outcomes of precocious puberty: An update. *Journal of Clinical Endocrinology and Metabolism* 2013, 98, 2198–2207. (doi:10.1210/jc.2013-1024)

3. Dacou-Voutetakis C & Karidis N. Congenital adrenal hyperplasia complicated by central precocious puberty: treatment with LHRH-agonist. *Ann N Y Acad Sci* 1993, 687, 250–254.

4. Speiser P. Transient central precocious puberty in non-classic 21-hydroxylase deficiency. *The Journal of Pediatric Endocrinology and Metabolism* 1995, 8, 287–289.

5. Soliman AT, Ailamki M, Aisalmi I, & Asfour M. Congenital Adrenal Hyperplasia Complicated by Central Precocious Puberty: Linear Growth During Infancy and Treatment With Gonadotropin-Releasing Hormone Analog. *Metabolism* 1997, 46, 513–517.

6. Guven A, Cebeci AN, & Hancili S. Gonadotropin releasing hormone analog treatment in children with congenital adrenal hyperplasia complicated by central precocious puberty. *Hormones* 2015, 14, 265–271. (doi:10.14310/horm.2002.1555)

7. Almeida MQ, Brito VN, Lins TSS, Guerra-Junior G, Castro M De, Antonini SR, Arnhold IJP, Mendonca BB, & Latronico AC. Long-term treatment of familial male-limited precocious puberty (testotoxicosis) with cyproterone acetate or ketoconazole. *Clinical Endocrinology* 2008, 69, 93–98. (doi:10.1111/j.1365-2265.2007.03160.x)
8. Mengel W & Knorr D. Leydig cell tumours in childhood. *Prog Pediatr Surg* 1983 **16** 133–138.

9. Criscoulo T, Sinisi A, Perrone L, Graziani M, Bellastella A, & Faggiano M. Isosexual precocious pseudopuberty secondary to a testosterone-secreting Leydig cell testicular tumour: true isosexual development early after surgery. *Andrologia* 1986 **18** 175–183.

10. Ghazi A, Rahimi F, Ahadi M, & Sadeghi-Nejad A. Development of true precocious puberty following treatment of a Leydig cell tumor of the testis. *The Journal of Pediatric Endocrinology and Metabolism* 2001 **14** 1679–1681.

11. Kiepe D, Annette RU, Autschbach F, Kessler M, Schenk JP, & Bettendorf M. Sexual Pseudo-Precocity Caused by a Somatic Activating Mutation of the LH Receptor Preceding True Sexual Precocity. *Hormone Research in Paediatrics* 2008 **70** 249–253. (doi:10.1159/000151598)

12. Lignitz S, Partsch CJ, Wudy SA, Hartmann MF, & Pohlenz J. Clinical and metabolic findings in a 6-year-old boy with a Leydig cell tumour. *Acta Paediatrica* 2011 **280–282.** (doi:10.1111/j.1651-2227.2011.02338.x)

13. Olivier P, Simoneau-roy J, & Francoeur D. Leydig Cell Tumors in Children: Contrasting Clinical, Hormonal, Anatomical, and Molecular Characteristics in Boys and Girls. *The Journal of Pediatrics* 2012 **161** 1147–1152. (doi:10.1016/j.jpeds.2012.05.039)

14. Santos-Silva R, Bonito-Vítor A, Campos M, & Fontoura M. Gonadotropin-Dependent Precocious Puberty in an 8-Year-Old Boy with Leydig Cell Testicular Tumor. *Hormone Research in Paediatrics* 2014 **82** 133–137. (doi:10.1159/000358084)

15. Verrotti A, Penta L, Zenzeri L, Lucchetti L, Giovenali P, & Feo P De. True Precocious Puberty Following Treatment of a Leydig Cell Tumor: two case reports and literature
16. Brachet C & Heinrichs C. Central precocious puberty after interpersonal transfer of testosterone gel: just a coincidence? *The Journal of Pediatric Endocrinology and Metabolism* 2012 **25** 757–760. (doi:10.1515/jpem-2012-0067)

17. Antonini SR, Leal LF, & Cavalcanti MM. Pediatric adrenocortical tumors: diagnosis, management and advancements in the understanding of the genetic basis and therapeutic implications. *Expert Review of Endocrinology and Metabolism* 2014 **9** 445–464.

18. Salt A, Savage M, & Grant D. Growth patterns after surgery for virilising adrenocortical adenoma. *Arch Dis Child* 1992 **67** 234–236.

19. Sandrini R, Ribeiro RC, & Lacerda L De. Childhood Adrenocortical Tumors. *The Journal of Clinical Endocrinology & Metabolism* 1997 **82** 2027–2031.

20. Latronico AC, Pinto EM, Domenice S, Candida M, Villares B, Martin RM, Zerbini MC, Lucon AM, & Mendonca BB. An Inherited Mutation Outside the Highly Conserved DNA-Binding Domain of the p53 Tumor Suppressor Protein in Children and Adults with Sporadic Adrenocortical Tumors. *JCEM* 2001 **86** 4970–4973. (doi:10.1210/jcem.86.10.7957)

21. Ribeiro RC, Sandrini F, Figueiredo B, Zambetti GP, & Michalkiewicz E. An inherited p53 mutation that contributes in a tissue-specific manner to pediatric adrenal. *PNAS* 2001 **98** 9330–9335. (doi:10.1073/pnas.161479898)

22. Sandrini F, Villani DP, Tucci S, Moreira AC, Castro M, & Elias LLK. Inheritance of R337H p53 Gene Mutation in Children with Sporadic Adrenocortical Tumor. *Horm Metab Res* 2005 . (doi:10.1055/s-2005-861373)

23. Leal LF, Mermejo LM, Ramalho LZ, Martinelli CE, Yunes A, Seidinger AL, Jose M,
Cardinalli IA, Brandalise SR, Moreira AC, Tone LG, Scrideli CA, Castro M, & Antonini SR. Wnt/beta-Catenin Pathway Deregulation in Childhood Adrenocortical Tumors. *JCEM* 2011 96 3106–3114. (doi:10.1210/jc.2011-0363)

24. Mastellaro MJ, Ribeiro RC, Oliveira-filho AG, Seidinger AL, Cardinalli IA, Miranda ECM, Aguiar SS, Brandalise SR, Yunes JA, & Barros-Filho AA. Adrenocortical tumors associated with the TP53 p.R337H germline mutation can be identified during childhood. *Jornal de Pediatria* 2017 550 1–8. (doi:10.1016/j.jped.2017.06.009)

25. Isabel M, Achatz W, Olivier M, Le F, Ashton-prolla P, Giugliani R, Inez E, Casali C, Rocha D, Luiz A, Hainaut P, & Vargas R. The TP53 mutation, R337H, is associated with Li-Fraumeni and Li-Fraumeni-like syndromes in Brazilian families. *Cancer Letters* 2007 245 96–102. (doi:10.1016/j.canlet.2005.12.039)

26. Kratz CP, Achatz MI, Brugi L, Frebourg T, Garber JE, Greer M lousie C, Hansford JR, Janeway KA, Kohlmann WK, Mcgee R, Mullighan CG, Onel K, Pajtler KW, Stefan MP, & Savage SA. Cancer Screening Recommendations for Individuals with Li-Fraumeni Syndrome. *Clinical Cancer Research* 2017 23 38–46. (doi:10.1158/1078-0432.CCR-17-0408)

27. Lau SK & Weiss LM. The Weiss system for evaluating adrenocortical neoplasms: 25 years later. *Human Pathology* 2009 40 757–768. (doi:10.1016/j.humpath.2009.03.010)

28. León MIM, Chaparro SR, Lara BW, & Pinos MDD. Tumores corticosuprarrenales pediátricos: imagen de adenomas y carcinomas. *Radiologia* 2012 54 342–349. (doi:10.1016/j.rx.2011.02.005)

29. Ribeiro R, Pinto E, Zambetti G, & Rodriguez-Galindo C. The International Pediatric Adrenocortical Tumor Registry initiative: Contributions to clinical, biological, and
treatment advances in pediatric adrenocortical tumors. Molecular and Cellular Endocrinology 2012 351 37–43. (doi:10.1016/j.mce.2011.10.015)

30. Valerio G, Spagnuolo M, Muzzi G, Buono P, Lombardi F, Palmieri R, & Franzese A. Adrenocortical Tumor in a Boy: Final Height is Not Impaired Despite a Severe Advancement of Bone Age. Journal of Pediatric Endocrinology & Metabolism 2003 16 1061–1063.

31. Kim MS, Yang EJ, Cho DH, Hwang PH, & Lee D yeol. Virilizing Adrenocortical Carcinoma Advancing to Central Precocious Puberty after Surgery. Korean Journal of Family Medicine 2015 150–153. (doi:10.4082/kjfm.2015.36.3.150)

32. Breidbart E. Pubertal outcome in a female with virilizing adrenocortical carcinoma. J Pediatr Endocrinol Metab 2016 29 503–509. (doi:10.1515/jpem-2015-0123.)

33. Ersoy B, Kizilay D, Cayirli H, Temiz P, & Gunsar C. Central Precocious Puberty Secondary to Adrenocortical Adenoma in a Female Child: Case Report and Review of the Literature. Journal of Pediatric and Adolescent Gynecology 2017 30 591–594. (doi:10.1016/j.jpag.2017.05.009)

34. Lee P, Winter R, & Green O. Virilizing Adrenocortical Tumors in Childhood: Eight Cases and a Review of the Literature. Pediatrics 1985 76 437–444.

35. Barstow C & Rerucha C. Evaluation of Short and Tall Stature in Children. American Family Physician 2015 92 43–50.

36. Tanner J. The use and abuse of growth standards. In Human Growth, edn 2nd, pp 95–112. Eds Falkner F & Tanner J. 1986.

37. Marshall W & Tanner J. Variations in Pattern of Pubertal Changes in Girls. Archives of Disease in Childhood 1969 44 291–303. (doi:10.1136/adc.44.235.291)

38. Marshall W & Tanner J. Variations in the Pattern of Pubertal Changes in Boys.
39. Greulich WW & Pyle Sl. Radiographic Atlas of Skeletal Development of the Hand and Wrist. edn 2nd, Stanford University Press, Palo Alto, CA, 1959.

40. Lazar L, Kauli R, Pertzelan A, & Phillip M. Gonadotropin-Suppressive Therapy in Girls with Early and Fast Puberty Affects the Pace of Puberty but Not Total Pubertal Growth or Final Height. *J Clin Endocr Metab* 2002 **87** 2090–2094.

41. Stagi S, Bindi G, Galluzzi F, Cauza F la, & Salti R. Precocious, early and fast puberty in males with Chiari I malformation. *J Pediatr Endocrinol Metab* 2004 **17** 1137–1140.

42. Ribeiro R, Neto R, Schell M, Lacerda L, Sambaio G, & Cat I. Adrenocortical Carcinoma in Children: a Study of 40 Cases. *Journal of Clinical Oncology* 1990 **8** 67–74.

43. Wolthers O, Cameron F, Scheimberg I, Honour J, Hindmarsh P, Savage M, Stanhope R, & Brook C. Androgen secreting adrenocortical tumours. *Arch Dis Child* 1999 **80** 46–50.

44. Flor-cisneros A, Leschek EW, Merke DP, Barnes KM, Coco M, Cutler Jr GB, & Baron J. In Boys with Abnormal Developmental Tempo, Maturation of the Skeleton and the Hypothalamic-Pituitary-Gonadal Axis Remains Synchronous. *The Journal of Clinical Endocrinology & Metabolism* 2004 **89** 236–241. (doi:10.1210/jc.2002-021954)

45. Moreira A, Veríssimo J, Foss M, Iazigi N, Maciel L, Pimenta W, Rodrigues J, & Santoro J. Pubertal Maturation of the LH Stimulatory Response to Clomiphene Citrate in Congenital Virilizing Adrenal Hyperplasia. *Clinical Endocrinology* 1982 **17** 441–447.

46. Savage M, Scommegna S, Carroll P, Ho J, Monson J, Besser G, & Grossman A. Growth in Disorders of Adrenal Hyperfunction. *Horm Res* 2002 **58** 39–43. (doi:10.1159/000064767)

47. Hauffa BP, Roll C, Mühlenberg R, & Havers W. Growth in children with adrenocortical tumors. *Klin Pädiatr* 1991 **203** 83–87.
FIGURE LEGENDS

Figure 1. Selection of patients with diagnosis of adrenocortical tumor in childhood according inclusion and exclusion criteria.

Figure 2. Pubertal outcomes in 63 patients with diagnosis of adrenocortical tumor in childhood.

Figure 3. Initial and final height according to pubertal development in patients with diagnosis of adrenocortical tumor in childhood.
70 patients with ACT < 18 years (1975 – 2017)

64 patients included in the study

63 eligible patients

6 patients not included

1 patient without surgery

1 patient with diagnosis and treatment of ACT at another hospital

2 patients without hormonal assessment

2 patients with pure cortisol secreting ACT

1 patient excluded

1 patient with an overgrowth syndrome (Beckwith-Wiedeman)
63 patients with ACT < 18 years

37 patients already developed gonadotropin-dependent puberty

- 10 patients had pubertal disorders
  - 7 patients with CPP
  - 3 patients with EFP
- 1 patient had isolated precocious thelarche
- 26 patients had normal puberty
  - 9 patients with EP

26 patients haven’t developed gonadotropin-dependent puberty

- 1 patient had precocious isolated vaginal bleeding without other signs of puberty
Table 1. General features of the 63 patients with adrenocortical tumors < 18 years from 1975 to 2017.

| Features                                      | Total n = 63 |
|-----------------------------------------------|--------------|
| **At diagnosis of ACT**                       |              |
| **Sex**                                       |              |
| Female                                        | 44           |
| Male                                          | 19           |
| **Skin color**                                |              |
| White                                         | 58           |
| Non-white                                     | 5            |
| **Age (months)**                              | 25.8 (2.1 – 192.2) |
| **Duration of ACT signs (months)**             | 6 (0 – 60)   |
| **Family history of ACT**                     | 8            |
| **Family history of other types of cancer**    | 21           |
| **Clinical features**                         |              |
| Pubarche                                      | 51           |
| Macrogenitosomy                               | 14/19        |
| Clitorimegaly                                  | 31/44        |
| Acne                                          | 34           |
| Axillary odor                                  | 15           |
| Muscular hypertrophy                          | 21           |
| Deep voice                                    | 14           |
| Agressiveness                                 | 15           |
| Irritability                                  | 9            |
| Hypertension                                  |              |
| Present                                       | 35           |
| Absent                                        | 16           |
| Data not available                            | 12           |
| Weight gain                                   | 10           |
| Facial pletora                                | 5            |
| Abdominal mass                                | 7            |
| Abdominal pain                                | 3            |
| **Tanner stage for pubic hair**               |              |
| PH1                                           | 8            |
| PH2                                           | 22           |
| PH3                                           | 22           |
| PH4                                           | 6            |
| PH5                                           | 5            |
| **Stature SDS**                               | 0.5 (-3.5 – 3.9) |
| **BMI SDS**                                   | 1.4 (-1.9 – 6.3) |
| **DHEA-S** *(µg/dL)*                          | 748 (32 – 4950) |
| **Testosterone** **(ng/dL)**                  | 242.5 (33.6 – 1800) |
| **∆ bone age (months)**                       | 14.7 (27.9 – 85.4) |
| **Tumor hormone profile**                     |              |
| Androgen secreting ACT (Virilizing)           | 8            |
| Androgen and cortisol secreting ACT (Mixed)   | 55           |
| **Tumor stage (IPACTR)**                      |              |
| I                                             | 38           |
| II                                            | 8            |
| III                                           | 11           |
| IV                                            | 6            |
| **PS3 p.R337H mutation**                      |              |
| Present                                       | 48           |
| Absent                                        | 4            |
| Data not available                            | 11           |
| **During follow-up**                          |              |
| **Chemotherapy**                              | 17           |
| **Recurrence / metastasis**                   | 16           |
| **Death**                                     | 14           |
| **Loss of follow-up**                         | 16           |
| **Discharge after 10 years**                  | 5            |
| **Duration of follow-up (months)**             | 73.7 (0.2 – 295.4) |

* n=55; ** n=54; *** n=52

ACT: adrenocortical tumor; PH: pubic hair; SDS: stature standard deviation; BMI: body mass index; DHEAS: dehydroepiandrosterone sulfate; ∆: delta (difference between bone age and chronological age); IPACTR: International Pediatric Adrenocortical Tumor Registry.
Table 2. General features of the 36# patients with adrenocortical tumor that developed gonadotropin-dependent puberty during follow-up.

| Features | Pubertal disorders (n = 10) | Normal puberty (n = 26) |
|----------|----------------------------|------------------------|
| **At diagnosis of ACT** | | |
| **Sex** | | |
| Female | 7 | 15 |
| Male | 3 | 11 |
| **Skin color** | | |
| White | 10 | 26 |
| **Age (months)** | 67.7 (5.3 – 95.5) | 24.7 (4.7 – 192.2) |
| **Duration of ACT signs (months)** | 6 (1 – 12) | 6 (1.5 – 18) |
| **Tanner stage for pubic hair** | | |
| PH1 | 1 | 3 |
| PH2 | 4 | 9 |
| PH3 | 4 | 10 |
| PH4 | 0 | 2 |
| PH5 | 1 | 2 |
| **Stature SDS** | 1.63 (-1.96 – 2.44) | 1.29 (-3.55 – 3.29) |
| **BMI SDS** | 1.07 (-0.31 – 5.22) | 1.44 (-0.63 – 3.22) |
| **DHEA-S (µg/dL)** | 735 (33 – 3968) | 763 (32 – 4678)* |
| **Testosterone (ng/dL)** | 381 (58 – 900) | 235 (33.6 – 1728)* |
| **Δ bone age (months)** | 37.7 (-1.5 – 79.7)** | 19.7 (-5.6 – 49.1)*** |
| **Tumor Hormone Profile** | | |
| Virilizing-ACT | 0 | 6 |
| Mixed-ACT | 10 | 20 |
| **Tumor stage (IPACTR)** | | |
| I | 5 | 18 |
| II | 1 | 4 |
| III | 4 | 2 |
| IV | 0 | 2 |
| **P53 p.R337H mutation** | | |
| Present | 8 | 20 |
| Absent | 1 | 2 |
| Data not available | 1 | 4 |
| **During follow-up** | | |
| Chemotherapy | 6 | 5 |
| Recurrence / metastasis | 5 | 4 |
| Death | 4 | 2 |
| Loss of follow-up | 0 | 9 |
| Discharge after 10 years | 1 | 3 |
| **Duration of follow-up (months)** | 78.8 (4 – 247.6) | 150.3 (8.2 – 295.4) |

*The patient with precocious thelarche was not included in the analysis. *n = 23; **n = 8; ***n = 22
ACT: adrenocortical tumor; PH: pubic hair; SDS: stature standard deviation; BMI: body mass index; DHEAS: dehydroepiandrosterone sulfate; Δ: delta (difference between bone age and chronological age); IPACTR: International Pediatric Adrenocortical Tumor Registry.
Table 3. Relative risk for the development of central precocious puberty in pediatric patients with history of adrenocortical tumors.

| FEATURES AT DIAGNOSIS OF ACT | CPP | NP | RR | CI 95% |
|------------------------------|-----|----|----|--------|
| N                            | 7   | 26 |    |        |
| SEX                          |     |    |    |        |
| F                            | 4   | 15 | 0.98 | (0.26; 3.71) |
| M                            | 3   | 11 |    |        |
| AGE AT DIAGNOSIS             |     |    |    |        |
| ≥ 48 months                 | 4   | 6  | 3.07 | (0.84; 11.25) |
| < 48 months                  | 3   | 20 |    |        |
| DURATION OF ACT SIGNS        |     |    |    |        |
| ≥ 6 months                  | 4   | 14 | 1.11 | (0.29; 4.21) |
| < 6 months                  | 3   | 12 |    |        |
| TANNER STAGE – PUBIC HAIR   |     |    |    |        |
| > PH2                       | 4   | 15 | 0.98 | (0.26; 3.71) |
| ≤ PH2                       | 3   | 11 |    |        |
| STATURE SDS                 |     |    |    |        |
| > 2                         | 4   | 4  | 4.17 | (1.17; 14.8) |
| ≤ 2                         | 3   | 22 |    |        |
| BONE AGE                    |     |    |    |        |
| Advanced                    | 5   | 13 | 2.78 | (0.37; 20.59) |
| Not advanced                | 1   | 9  |    |        |
| DHEA-S (µg/dL)              |     |    |    |        |
| ≥ 300                       | 5   | 17 | 0.91 | (0.22; 3.78) |
| < 300                       | 2   | 6  |    |        |
| TESTOSTERONE (ng/dL)        |     |    |    |        |
| ≥ 300                       | 4   | 8  | 2.00 | (0.54; 7.39) |
| < 300                       | 3   | 15 |    |        |
| TUMOR HORMONE PROFILE       |     |    |    |        |
| V-ACT                       | 0   | 6  | *   | *      |
| M-ACT                       | 7   | 20 |    |        |
| TUMOR STAGE                 |     |    |    |        |
| III and IV                  | 3   | 4  | 2.78 | (0.80; 9.65) |
| I and II                    | 4   | 22 |    |        |
| RECURRENCE / METASTASIS     |     |    |    |        |
| (during follow-up)          |     |    |    |        |
| Present                     | 4   | 4  | 4.17 | (1.17; 14.8) |
| Absent                      | 3   | 22 |    |        |

* Not done.

ACT: adrenocortical tumor; CPP: central precocious puberty; NP: normal puberty; RR: relative risk; SDS: stature standard deviation; V-ACT: androgen secreting ACT (virilizing ACT); M-ACT: androgen and cortisol secreting ACT (mixed ACT); DHEAS: dehydroepiandrosterone sulfate; Δ: delta (difference between bone age and chronological age).
Table 4. Relative risk for the development of central precocious puberty or early fast puberty in pediatric patients with history of adrenocortical tumors.

| FEATURES AT DIAGNOSIS OF ACT | CPP/EFP | NP | RR     | CI 95%          |
|------------------------------|---------|----|--------|-----------------|
| N                            | 10      | 26 |        |                 |
| SEX                          |         |    |        |                 |
| F                            | 7       | 15 | 1.48   | (0.46; 4.81)    |
| M                            | 3       | 11 |        |                 |
| AGE AT DIAGNOSIS             |         |    |        |                 |
| ≥ 48 months                 | 6       | 6  | 3.00   | (1.04; 8.65)    |
| < 48 months                 | 4       | 20 |        |                 |
| DURATION OF ACT SIGNS        |         |    |        |                 |
| ≥ 6 months                  | 5       | 14 | 0.90   | (0.31; 2.56)    |
| < 6 months                  | 5       | 12 |        |                 |
| TANNER STAGE – PUBLIC HAIR  |         |    |        |                 |
| ≤ PH2                       | 6       | 15 | 1.07   | (0.36; 3.15)    |
| > 2                         | 4       | 11 |        |                 |
| STATURE SDS                 |         |    |        |                 |
| ≤ 2                         | 6       | 22 |        |                 |
| > 2                         | 4       | 4  | 2.33   | (0.27; 6.29)    |
| BONE AGE                    |         |    |        |                 |
| Advanced                    | 6       | 13 | 1.74   | (0.42; 7.17)    |
| Not advanced                | 2       | 9  |        |                 |
| DHEA-S (µg/dL)              |         |    |        |                 |
| ≥ 300                       | 8       | 17 | 1.28   | (0.34; 4.84)    |
| < 300                       | 2       | 6  |        |                 |
| TESTOSTERONE (ng/dL)        |         |    |        |                 |
| ≥ 300                       | 6       | 8  | 2.03   | (0.71; 5.88)    |
| < 300                       | 4       | 15 |        |                 |
| TUMOR HORMONE PROFILE       |         |    |        |                 |
| M-ACT                       | 10      | 20 |        |                 |
| V-ACT                       | 0       | 6  | *      | *               |
| TUMOR STAGE                 |         |    |        |                 |
| I / II                      | 6       | 22 |        |                 |
| RECURRENT / METASTASIS      |         |    |        |                 |
| (during follow-up)          |         |    |        |                 |
| Present                     | 5       | 4  | 3.00   | (1.12; 8.02)    |
| Absent                      | 5       | 22 |        |                 |

*Not done.

ACT: adrenocortical tumor; CPP: central precocious puberty; NP: normal puberty; RR: relative risk; SDS: stature standard deviation; V-ACT: androgen secreting ACT (virilizing ACT); M-ACT: androgen and cortisol secreting ACT (mixed ACT); DHEAS: dehydroepiandrosterone sulfate; ∆: delta (difference between bone age and chronological age).