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## Gonadal Function in Adult Male Patients with Congenital Adrenal Hyperplasia

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#### 1 Gonadal Function in Adult Male Patients with Congenital Adrenal Hyperplasia

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55 Abstract

56	Context: Current knowledge on gonadal function in Congenital Adrenal Hyperplasia (CAH) is mostly
57	limited to single center/country studies enrolling small patient numbers. Overall data indicate that
58	<mark>gonadal function can be compromised in men with CAH.</mark> Gonadal function can be compromised in
59	male patients with congenital adrenal hyperplasia (CAH), however previous studies have been
60	limited to reports from a single center/country or small patient numbers.
61	<b>Objective:</b> To determine gonadal function in men with CAH within the European "dsd-LIFE" cohort.
62	Design: Cross-sectional clinical outcome study, including retrospective data from medical records.
63	Methods: Fourteen academic hospitals included 121 men with CAH aged 16-68 years. Main outcome
64	measures were serum hormone concentrations, semen parameters, and imaging data of the testes.
65	<b>Results:</b> At the time of assessment, <del>19/83</del> 14/69 patients had a serum testosterone <mark>concentration</mark>
66	<del>level</del> below the reference range; <mark>8</mark> <mark>7</mark> of those were hypogonadotropic, <del>10</del> <mark>6</mark> normogonadotropic, and
67	1 hypergonadotropic. In contrast, <del>in the presence of</del> among the patients with normal serum
68	testosterone ( <del>64/83</del> <mark>55/69</mark> ), <del>5</del> <mark>4</mark> <del>patients</del> were hypogonadotropic, <del>50</del> <mark>44</mark> normogonadotropic, and <del>9</del> <mark>7</mark>
69	hypergonadotropic. The association of decreased testosterone with reduced gonadotropin
70	concentrations (Odds Ratio (OR)= <del>8.0 [2.2-29.6]</del> 12.8 [2.9-57.3]) was weaker than the association
71	between <mark>serum</mark> androstenedione/testosterone ratio ≥ 1 and reduced gonadotropin concentrations
72	(OR= <del>16.8 [2.0-142.5]</del> <mark>39.3 [2.1-732.4]</mark> ). <mark>Evaluation of sperm quality revealed decreased</mark>
73	sperm concentration <del>s</del> (15/39), <del>decreased</del> motility (13/37), and abnormal morphology (4/28) <del>were</del>
74	also observed. Testicular adrenal rest tumor (TART)s were present in 39/80 patients, with a higher
75	prevalence in patients with the most severe genotype (14/18), and in patients with increased current
76	17-hydroxyprogesterone ( <del>12/18</del> <mark>20/35</mark> ) or androstenedione ( <del>16/26</del> <mark>12/18</mark> ) <mark>serum</mark> concentrations.
77	Forty-three children were fathered by 26/113 patients.

- 78 **Conclusions:** Men with CAH have a high risk of developing hypothalamic-pituitary-gonadal
- 79 disturbances and spermatogenic abnormalities. Regular assessment of endocrine gonadal function
- 80 and of imaging for TART development by imaging are recommended, in addition to measures for
- 81 fertility protection.

#### 82 Introduction

Congenital adrenal hyperplasia (CAH) is an autosomal recessive disorder resulting in impaired
 adrenocortical steroid synthesis by several enzyme deficiencies. The most common form (>95%) is
 21-hydroxylase deficiency (210HD) with an incidence of 1:15 000, leading to glucocorticoid and often
 also mineralocorticoid deficiency in combination with androgen excess ies <sup>1, 2</sup>.

87 Reported fertility and fecundity in men with CAH on routine steroid replacement therapy range from 88 normal to severely impaired. Fertility can be compromised due to primary (hypergonadotropic) hypogonadism or central (hypogonadotropic) hypogonadism <sup>3-11</sup>. In addition, reduced fertility and 89 fecundity rates problems in CAH can be caused by psychosexual factors <sup>4</sup>. One of the commonest 90 91 complications in men with CAH is the presence of Testicular Adrenal Rest Tumor (TART)s, which can 92 cause disturbances of gonadal function, including mechanical obstruction of the seminiferous 93 tubules. The reported prevalence of TART ranges between 12.5% and 94% of the populations 94 studied. Central or secondary hypogonadism is defined as decreased testosterone concentrations in 95 combination with either low or low-normal LH or FSH concentrations. In patients men with CAH, 96 secondary hypogonadism is most likely to be caused by the suppressive effect of elevated adrenal 97 androgens (that are aromatized to estrogens) on the hypothalamic-pituitary-gonadal (HPG)-axis <sup>6</sup>. 98 Differentiation between gonadal and adrenal testosterone is difficult, complicating the diagnosis of hypogonadism in patients men with CAH. One of the commonest complications in men with CAH is 99 100 the presence of Testicular Adrenal Rest Tumor (TART)s, which can cause disturbances of gonadal 101 function, including mechanical obstruction of the seminiferous tubules. The reported prevalence of 102 TARTs ranges between 12.5% and 94% in the populations studied <sup>4-10, 12-22</sup>. Until now, the data on fertility outcome in men with CAH are scarce. Available data are <sup>3-11</sup> and often

Until now, the data on fertility outcome in men with CAH are scarce. Available data are <sup>3-11</sup> and ofte
derived from studies with patients from a single center or country. Our aim was to study gonadal
function in a large European multi-center cohort of male patients with CAH by evaluating hormone
concentrations, semen parameters, and TART frequency.

#### 107 Subjects and Methods

108 Subjects

109 dsd-LIFE is a cross-sectional clinical outcome study of individuals with disorders/differences of sex 110 development (DSD). Fourteen study centers in 6 European countries (France (n=4), Germany (n=4), 111 United Kingdom (n=1), Poland (n=2), Sweden (n=1), and the Netherlands (n=2)) included former and 112 current patients as participants from February 2014 - September 2015. In addition to DSD 113 participants, 121 male participants with CAH (46XY karyotype) aged 16-68 years were recruited as 114 they may face similar clinical challenges as DSD patients, including sex hormone imbalances and 115 fertility problems, although male patients with CAH do not fit into the classification of DSD. Written 116 informed consent was obtained from all participants and/or their parents, with assent of minors. 117 Ethical approvals were obtained as appropriate for each country, e.g. Ethics Commission of the 118 Charité Universitätsmedizin; reference number EA2/069/13. For. The theoretical and methodological framework of the dsd-LIFE study have been published in detail elsewhere see Röhle 2017 et al.<sup>23</sup>. 119 120 Patients were investigated in their local treatment center. Cross-sectional data were obtained for 121 serum hormone concentrations, semen parameters and testicular imaging. The genotype of patients with 210HD was classified into genotype groups null, A, B, and C<sup>24</sup>. Patients were also classified into 122 123 salt-wasting (SW), simple virilizing (SV) or non-classical (NC) based on their main symptoms and time 124 of diagnosis General patient characteristics and clinical parameters included: country of inclusion, 125 age, age at diagnosis, CAH genotype and phenotype, socioeconomic status, and obesity, as well as 126 height, weight, and BMI throughout the years (at diagnosis, 9 months old, 6 years old, Tanner stage 127 2, 16 years old, and current age). Patients' educational levels was established according to the EU 128 classification. We combined the standardized ES-ISCED (international standard classification of 129 education) scale to Low (ES-ISCED I = less than lower secondary and ES-ISCED II = lower secondary); 130 medium (ES-ISCED IIIb = lower tier upper secondary; ES-ISCED IIIA = upper tier upper secondary; ES-131 ISCED IV = advanced vocational, sub-degree) and high (ES-ISCED V1 = lower tertiary education, BA

- level; ES-ISCED V2 = higher tertiary education, >=MA level). Data was collected during medical
   examination at study inclusion (cross-sectional) and retrieved from medical records (retrospective
- 134 <mark>data).</mark>

135 Hormonal analysis

- 136 Blood samples were taken during day time, but mostly in the morning, before intake of the
- 137 glucocorticoid medication <sup>23</sup>. Total testosterone, SHBG, LH, FSH, inhibin B, AMH, androstenedione,
- 138 17-hydroxyprogesterone (17OHP) concentrations, and renin/plasma renin activity were measured in
- the local hospital laboratory and compared to local references. Values are reported in SI or
- 140 international units and reported as "below reference range", "within reference range", "above
- 141 reference range up to twice the upper limit", and "more than twice the upper limit of the reference
- range". To increase the number of patients per category, we combined the latter 2 categories into
- 143 the category "above reference range".
- 144 The serum androstenedione/testosterone ratio (AD/T) was calculated and divided into normal (<0.5;
- 145 interpreted as testosterone mainly of testicular origin), ≥0.5 and <1 (significant fraction of
- 146 testosterone is of adrenal origin), and  $\geq 1$  (testosterone mainly of adrenal origin ) as suggested by
- 147 others <sup>25</sup>.
- 148 Three patients were excluded from part of the analyses as they received testosterone substitution,
- 149 which directly affects testosterone and gonadotropin concentrations. Two of these patients had data
- 150 on TART available; these are described in the results section, but were otherwise excluded from
- 151 further analyses.
- 152 Semen analysis

Semen analysis was performed by the local hospital laboratory and interpreted in accordance with
 the 2010 World Health Organization criteria <sup>26</sup>, including sperm concentration (lower reference limit

(LRL): 15x10<sup>6</sup>/mL), motility (LRL: 40%), morphology (LRL: 4%), vitality (LRL: 58%), and volume (LRL:
1.5 mL).

#### 157 Imaging of testes

- 158 At the study visit, 68 patients (56.2%) underwent testicular ultrasound. The presence of TART at the
- age of 16 years was also reported retrospectively (in 30/68 patients with cross-sectional TART data).
- 160 In addition, retrospective data were available for 12 participants based on ultrasound findings or MRI
- 161 (n=11) and on histological findings (n=1).

#### 162 Paternity

163 Data about paternity and relationships were collected from the dsd-LIFE questionnaires <sup>23</sup>.

#### 164 Medication and estimation of metabolic control in the past

- 165 Patients used different formulations of glucocorticoids, including hydrocortisone, prednisone,
- 166 prednisolone, and dexamethasone. Furthermore, we converted all All glucocorticoid preparations
- 167 were converted to hydrocortisone equivalents for comparison, using the following factors for the
- 168 glucocorticoid equivalent dose: 1 (hydrocortisone), 4 (prednisone or prednisolone), 30
- 169 (dexamethasone), and 15 (fludrocortisone)<sup>27</sup>. We also calculated mineralocorticoid equivalent dose
- using the following factors: 1 (hydrocortisone), 0.8 (prednisone or prednisolone), 0 (dexamethasone),
- and 200 (fludrocortisone)<sup>27</sup>. In addition to the serum 170HP concentrations presented in the section
- 172 hormonal analysis, we also assessed metabolic control by a subjective rating, of metabolic control of
- the local examining physician at 5 different time points: at diagnosis, at the age of 9 months, at
- 174 Tanner stage 2, at age 16 years and at study inclusion, using the following scores: "poor",
- 175 "moderate", "good", "excellent" or "unknown".

#### 176 Statistical Analysis

SPSS Statistics 22 (SPSS Inc., Chicago, IL, USA) was used for all analyses. Descriptive analyses were performed for all variables. Depending on normality, mean and 95% confidence intervals (95%CI) or median and interquartile ranges (IQR) were calculated. We compared patients with values below or above reference range to patients with normal values (within the reference range). Odds ratios (OR) with 95%CI were calculated if at least 3 cases were present in both subgroups. If any cell count in the contingency table was zero, OR and 95%CI were calculated manually by using a continuity correction (+0.5 in each cell).

- 184 Missing data were evaluated for each variable and the total number of participants in a particular
- analysis was reported exactly. Analysis of the variables was <del>only</del> performed <mark>only if</mark> <del>when</del> the number
- 186 of participants was  $\geq$  at least 25% of the total cohort of male patients with CAH.
- 187 Three patients were excluded from part of the analyses as they received testosterone substitution,
- 188 which directly affects testosterone and gonadotropin concentrations. Two of these patients had data
- 189 on TART available; these are described in the results section, but were otherwise excluded from
- 190 further analyses. Furthermore, we excluded 22 patients with missing genotype information and 2
- 191 patients with  $11\beta$ -hydroxylase deficiency from all comparative analyses.

#### 192 Results

#### 193 General characteristics of the male CAH cohort

- 194 A total of 121 male patients were included in the CAH cohort in the dsd-LIFE study. General
- 195 characteristics are shown in Table 1. The median age of the study population was 28 years (IQR: 18.5-
- 196 37.5, range 16-68). Mean height was 170.7 (95%CI: 169.3-172.0) cm and median BMI was 25.6 (IQR:
- 197 22.0-29.2) kg/m<sup>2</sup> (data available for 119 patients). Nearly Almost all patients had 210HD (119/121),
- 198 of which and 97 were confirmed by molecular genetic analysis and 22 were based on phenotype
- 199 alone. The remaining 2 patients had  $11\beta$ -hydroxylase deficiency. Among the 97 patients with
- 200 genetically confirmed 21OHD, Genotype groups null, A, B, and C contained 19.8% 24.7% were
- 201 classified as genotype null, <del>30.6%</del> 38.1% as genotype A, <del>27.3%</del> 34.0% as genotype B, and <del>2.5%</del> 3.1%
- 202 as genotype C. of the 97 patients with genotyping results. The majority of patients (62.0%) were
- 203 classified as having the SW form of CAH, 31.4% had the SV form and 4.1% had the NC form.
- 204 Glucocorticoids were used by 116 (95.9%) patients, most commonly hydrocortisone, followed by
- prednisone or prednisolone, and dexamethasone. Fludrocortisone was used by 86 patients (71.1%).
- 206 The patients' education was intermediate or high in 54.5%, and 22.3% of the participants,
- 207 respectively. Furthermore, 54.6% of the patients were in a relationship at the time of study.
- 208 We analyzed all variables mentioned in the method section, but we only present in detail the data
- 209 that differed between the analyzed groups (no overlap in the confidence intervals). In the following
- 210 sections we will present data regarding hormone concentrations, semen analysis and TART.
- 211 Hormone concentrations
- 212 Univariate descriptive analyses of hormone concentrations were performed. The proportion of
- 213 patients with normal, decreased or increased serum testosterone, LH, FSH, inhibin B, AMH, and SHBG
- 214 concentrations is illustrated in Figure 1A. Hormone concentrations were below the reference range
- in 19/97 (19.6%: testosterone), 8/43 (18.6%: inhibin B), 12/90 (13.3%: LH), 9/90 (10.0%: FSH), and

- 216 1/69 (1.4%: SHBG) of the participants. SHBG concentrations were above the reference range in 14.5%
  217 (10/69).
- 218 Table 2 shows compares testosterone and gonadotropin concentrations in all patients with data on T, 219 LH, and, FSH available. that in p-Seven patients (50%) with decreased testosterone concentrations 220 (19/83), 8 (42.1%) had decreased gonadotropins, while 10-6 (52.6-42.9%) had normal LH and FSH 221 concentrations, and 1 (5.3 7.1%) patient had gonadotropin concentrations above reference range. 222 Normal testosterone concentrations were found in 64/83 55/69 (77.1 79.7%) patients, 50 44 (78.1 80.0%) of whom had normal gonadotropin concentrations, whereas 9 7 (14.1-12.7%) had increased, 223 224 and  $\frac{5}{4}$  (7.38%) had decreased concentrations. Decreased testosterone concentrations were clearly 225 associated with decreased LH and/or decreased FSH concentrations (OR 8-0 12.8, 95%CI: 2.9 - 2-226 <del>29.6</del>57.3). 227 A serum An AD/T ratio was calculated in 49 patients, 22 of whom (44.9%) had an AD/T ratio ≥1. Ten 228 patients (45.5%) with an AD/T ≥1 had decreased gonadotropins, while 11 (50.0%) patients had 229 normal gonadotropins and only 1 (4.5%) patient had increased gonadotropins. Normal AD/T ratios 230 were found in 27/49 (55.1%) patients, 21 of whom had normal gonadotropin concentrations (77.8%), 231 5 had increased concentrations, but none had decreased gonadotropin concentrations. was found in 232 7/8 patients (87.5%) with decreased testosterone and gonadotropins, while 4/5 patients (80.0%) 233 with normal testosterone and decreased gonadotropins had an AD/T ratio ≥1. Moreover, 5/10 234 patients (50.0%) with decreased testosterone and normal gonadotropins had an AD/T ratio ≥1, 235 whereas this was seen in only 11/50 patients (22.0%) with normal testosterone and gonadotropins.
- An AD/T ratio ≥1 was strongly associated with decreased LH and/or decreased FSH concentrations
- 237 (OR <del>16.8</del> <mark>39.3</mark>, 95%CI: <mark>2.1</mark>0 <del>142.5</del> 732.4</mark>).

#### 238 Semen analysis

Semen analysis was performed in approximately one third of the patients (Figure 1B). Normal values
for all known (at least 3 out of 5) semen parameters (normozoospermia) were seen in 11/39 patients

241 in which semen analysis was performed. Sperm concentration, motility, and volume were below the 242 normal ranges in 38.5% (15/39), 35.1% (13/37), and 25.6% (10/39) of the patients, respectively, while 243 morphology and vitality were both impaired in 14.3% (4/28 and 2/14) of the patients. Five of 8 244 patients (62.5%) with decreased testosterone and gonadotropin concentrations underwent semen 245 analysis, with 4 (80.0%) of them showing abnormal semen parameters (Table 3). In only 2/10 246 patients with decreased testosterone, but normal gonadotropin concentrations, semen analysis was 247 performed and both had decreased sperm concentrations (7.0 and  $10.0 \times 10^6$ /mL). No statistically 248 significant associations were found (data not shown).

#### 249 Testicular adrenal rest tumors

250 TARTs were visualized by ultrasound or MRI at cross-sectional investigation in 28/68 patients. For 1 251 patient, the diagnosis was based on retrospective histology data. Furthermore, retrospective imaging 252 data were available for 11 men: TARTs were present in 10 of these individuals. So, in In the total 253 population screened, TARTs were present in 39/80 patients (48.8%) of which 34 were bilateral TARTs 254 (87.2%). Documented retrospective TARTs at age 16 years were reported in 16/30 patients (53.3%), 255 all of which were bilateral. In only 2/16 patients (12.5%) with TART reported to be present at age 16, TART was no longer observed during the cross-sectional investigation: one patient was misdiagnosed 256 257 with TART as it appeared to be a varicocele, and in the other patient TART (size 2 mm) disappeared 258 after treatment with prednisone. This patient was still considered as a TART patient with TART in all 259 analyses.

#### 260 Comparison of patients with and without TART

Table 4 shows associations of TART with various variables in the 78 68 patients with gonadal imaging

262 data (12 patients were excluded due to testosterone substitution, 11β-hydroxylase deficiency or

263 unconfirmed 21-hydroxylase deficiency), comprising <del>37</del> 33 patients with and 41 35 without TARTs.

264 Genotype was <del>clearly</del> associated with the presence of TART: The null genotype group had the highest

prevalence of TART (14/18: 77.8%), while the prevalence was 10/27 (37.0%) for genotype group A,

266 and 7/21 (33.3%) for genotype group B. The odds of having TART in the null genotype group was 6.0 267 [1.5-23.1] and 7.0 [1.7-29.4] times higher compared to the genotype groups A and B, respectively. 268 TARTs were also present in both men in the genotype C group, and also in 1 CYP11B1-deficient 269 patient (the other CYP11B1 patient did not undergo assessment for TART). The OR of having TART 270 when having an a serum androstenedione level concentration above the upper limit of normal at the 271 time of the cross-sectional investigation was 3.6 3 [1.0 - 11.2 12.7]. Similar associations were found 272 for serum 17OHP at the cross-sectional investigation, with an OR of 6.4 28.0 [1.7 3.1 - 24.7 252.5] for 273 having TART when 170HP concentrations were more than twice the upper level of the reference 274 range, and an OR of 4 18.7 [1.3 2.2 - 158.1 16.5] when these concentrations were above the 275 reference range compared to concentrations within the reference range.

#### 276 Paternity

Data on paternity were available for 113 of the 121 patients, 26 (23.0%) of whom (age range 26-68
years) had fathered a total of 43 children. Three couples had used assisted reproductive techniques
(ART) resulting in 4/43 children. One of the men who had used ART had decreased testosterone
concentrations, while another had increased FSH, decreased sperm concentration, and TART. No
information was available about the third patient who had used ART. One man with impaired semen
motility, increased FSH concentrations, and TART had adopted a child.

#### 283 Discussion

This unique and relatively large European multicenter study shows that gonadal dysfunction is a common complication in male patients with CAH. Approximately half of the patients were affected by endocrine disturbances of the HPG axis at an adult age and TARTs were present in approximately half of the patients as well.

The difficulty in diagnosing hypogonadism in men with CAH is related to the fact that testosterone
 measured in serum is a mixture of testosterone of gonadal and adrenal origin <sup>25, 28</sup>. Circulating

290 testosterone in male patients with well-controlled CAH is predominantly derived from testicular 291 production, but when there is poor hormonal control, a relevant contribution arises from adrenal 292 steroidogenesis. Until now, no method is able to discriminate between testosterone derived from 293 the testes or the adrenal gland. Therefore, it has been suggested to use the serum AD/T ratio in male 294 patients with CAH, as this precursor steroid is elevated in serum when serum androgens are predominantly of adrenal origin <sup>25</sup>. Our data point toward an association confirmed a stronger 295 296 association between an AD/T ratio ≥1 (testosterone mainly of adrenal origin) and decreased LH 297 and/or decreased FSH concentrations compared to testosterone concentrations alone, suggesting 298 that adrenal androgens in men with CAH contribute to the suppression of gonadotropins. In 299 approximately half of the patients, either aberrant testosterone or AD/T ratios, or aberrant 300 gonadotropin concentrations, or a combination of both were found. In previous studies, the reported prevalence of endocrine HPG axis disturbances ranged from 20% to 52% <sup>5-7, 9, 10</sup>. However, only 1 301 other report study provided had information on testosterone and gonadotropin concentrations in 302 303 each patient, and also indicated endocrine disturbances hypogonadism in approximately half of the patients <sup>6</sup>. We recommend to including include the evaluation of the AD/T ratio in the regular follow-304 305 <mark>up androstenedione measurements in the gonadal evaluation</mark> of male patients with CAH <del>to calculate</del> 306 the AD/T ratio, and interpret this ratio in combination with gonadotropin concentrations in order to 307 detect a disturbance of the HPG axis. Our study did does not include data information on 11-308 oxygenated androgens, that are generated through conversion of androstenedione, and are reported to be elevated concentrations are found in patients with CAH <sup>29, 30</sup>. Recent studies indicate that 11-309 310 oxygenated androgens are almost entirely derived from the 11beta-hydroxylation of androstenedione in the adrenal, and as they are potent androgens they can contribute to 311 suppression of the HPG axis<sup>31</sup>. However, their exact role in the evaluation of However, their 312 313 associations with hormonal control and gonadal function in men with CAH has to be established in further studies. Serum AMH and inhibin B are also used as markers for male fertility <sup>32</sup>. However, 314 315 literature already showed it has been demonstrated that serum AMH concentrations do not

correlate with sperm concentration and other male fertility parameters <sup>33</sup>. Serum inhibin B, a marker 316 317 of Sertoli cell function, is known to correlate with spermatogenesis in healthy men <sup>34</sup> and was 318 decreased in 18.3% 18.6% of our cohort. Semen quality, assessed in one third of the study cohort, 319 was reduced in 40% of the men. Except for the study of Urban et al.<sup>3</sup>, all other studies on fertility in male patients with CAH showed decreased sperm concentrations ranging from 47.8% to 66% <sup>4-7, 9, 10</sup>. 320 321 More strikingly, in all studies only half of the participants participated in semen analysis. Taken 322 together, these data indicate the need for Therefore, increased awareness on fertility status in 323 patients with CAH, and to start is needed. We recommend performing semen analysis and gonadal 324 <mark>function biomarkers assessment <del>as early as possible in</del> from</mark> adolescence <mark>on, in order</mark> to detect disturbances early and allow semen preservation to be able to preserve semen for later fertility 325 326 purposes. 327 Data from our cohort indicate, in agreement with previous studies The prevalence of TART in the 328 present patient cohort was 48.8%, confirming previous reports that TART is a common complication in male patients with CAH<sup>4-10, 12-22</sup>, that TART is a common complication in males with CAH (with a 329 330 prevalence of 48.8%) and can have onset as early as in adolescence. In fact, Strikingly, 14 patients 331 with TART at the time of the dsd-LIFE study already had TART at the age of 16 years. TARTs 332 disappeared on at 16 years were no longer detectable following treatment with prednisone in only 1 333 patient, thus indicating. This could indicate that complete regression of TART might only be achieved 334 in a small proportion of the patients. Hence, prevention of the development of TART should be 335 pursued, by optimizing treatment strategies already in childhood. Current standard of care does not 336 include imaging of testes, however we recommend incorporating testicular ultrasound in routine 337 clinical practice. 338 In contrast to previous studies, several studies did not find an association between CAH severity and TART<sup>4,9,10</sup>, we observed an association between the CYP21A2 genotype and the presence of TARTs, 339

with the prevalence of this complication being was highest in men with the null *CYP21A2* genotype.

341 This likely confirms supports the current perception that TARTs are more frequently observed in patients with a more severe form of CAH, as these patients are exposed to higher concentrations of 342 ACTH, already in utero, which is thought to be a possible causative factor for TART development <sup>6, 7, 15,</sup> 343 344 <sup>22</sup>. However, a clinically relevant finding in this study is that TARTs occurs even in less severe forms of 345 210HD. In fact, in -In our study, 2 patients in genotype group C with NC CAH (both compound heterozygous for deletion and P30L mutation) had TARTs. In our current dataset, we could not find 346 347 an association between genotype and semen quality or genotype and hypogonadism. Both patients 348 were compound heterozygous (deletion + P30L mutation). Only 1 patient in our cohort had the 349 typical NC mutation, i.e. the V281 mutation (V281/I2Splice). No TARTs were detected in this patient. 350 In our study, we We found an association between increased 170HP concentrations at cross-351 sectional data assessment and the presence of TART. Although a single 170HP measurement may 352 not be representative of overall metabolic control, these results could be interpreted as a possible 353 indicator of the patient's metabolic control in the recent past. Therefore, our results seem to be in 354 accordance with literature reporting higher TART prevalence in patients with poor hormonal control 355 compared to patients with adequate hormonal control <sup>5, 7, 13, 35-38</sup>. The association between increased 356 androstenedione concentrations at cross-sectional data assessment and the presence of TARTs adds 357 evidence to this pathophysiologic concept, even if the AD/T ratios were not clearly associated with 358 TART within this subgroup of patients. Primary gonadal dysfunction may be suggested by raised FSH 359 concentrations. In our dataset, 10 patients (11.1%) had elevated FSH concentrations. Seven of these 360 patients had data on the presence of TART, and 4 had evidence of TART. King et al. found that testicular failure was a consequence of TART in the majority of cases <sup>10</sup>. However, our data are 361 limited and do not allow firm conclusions concerning this issue. We cannot confirm the findings of 362 King as we have only very limited data available. 363

Despite this being the first international multicenter study describing gonadal function in male
 patients with CAH, the study also has some limitations. All centers included in this consortium are

366 tertiary care centers, therefore it is possible that the patient groups were selected and that the 367 patients included were more severely affected. Furthermore, serum hormone concentrations were 368 not measured centrally, but in various centers with a range of different assays. Accounting for this 369 fact, only range variables were used in the data analyses. The median BMI in our patient cohort was 370 25.6 kg/m<sup>2</sup> (range 22.0-29.2), which is slightly overweight. It has been demonstrated that excess of total and abdominal body fat could represent one cause of fertility impairment in men with CAH <sup>25</sup>. 371 372 Serum total testosterone can be decreased in patients with obesity, as a result of the decreased 373 serum concentration of SHBG. In case of increased serum SHBG (induced by hepatitis, 374 hyperthyroidism, or a genetic variant), total testosterone may be increased. Ideally, free testosterone 375 should be measured in these cases, but this requires complex equilibrium dialysis <sup>39</sup>. Free 376 testosterone can also be calculated from the level of total testosterone, SHBG, and albumin 377 concentrations, but it is crucial that the results of such calculations are compared with the normal 378 range of each separate laboratory. Such data were not available. We are aware that assessment of 379 fertility by paternity numbers in our study was is incomplete, as many other factors, of which 380 including female fertility, are important as well. However, these data were not available. 381 Furthermore, participation in the medical examination was not obligatory compulsory for study 382 inclusion. This may have led to even more selection, especially concerning the ultrasound 383 examination and semen analysis. It is likely that only the very motivated patients and the more 384 severely affected patients consented to these additional examinations. Due to the resulting low 385 numbers of available data, multivariable logistic regression analyses were not possible. 386 In summary, impaired gonadal function is common in adult men with CAH. This is indicated by the 387 presence of TART and/or hypogonadotropic or hypergonadotropic hypogonadism. The risk of TART is highest in men with the most severe forms of enzyme deficiencies underlying CAH. Our data suggest 388 389 that an An association with poor previous hormonal control is likely but <del>has to be confirmed</del> requires 390 confirmation by <del>further prospective</del> studies. Determination of the serum AD/T ratio, in addition to 391 serum concentrations of testosterone, androstenedione, LH, and FSH may help to differentiate

- 392 between testicular and adrenal androgens in male patients with CAH and to estimate the degree
- 393 diagnose of gonadal dysfunction. Routinely performed semen analysis, measurement of serum
- inhibin B, and testicular ultrasound investigation already in adolescence are recommended to detect
- 395 upcoming reproductive problems and to allow for fertility preserving measures, such as sperm
- 396 banking.

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#### 404 **Declaration** Conflict of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing theimpartiality of the research reported.

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538

### 540 Figure legends

541	Figure 1: Hormone concentrations (A) and semen quality (B) in male patients with congenital
542	adrenal hyperplasia to assess gonadal function. Stacked bars represent percentage of patients
543	within a category. Numbers in the bars represent the specific number of patients within a category,
544	while the total number of patients included in this analysis is stated underneath the x-axis. A)
545	Hormone concentrations of each patient were measured in the local hospital and compared to the
546	hospitals standard reference ranges. B) Semen analysis was performed and scored according to
547	World Health Organization 2010 criteria <sup>26</sup> : sperm concentration, motility, morphology, and vitality,
548	and semen volume were assessed. Abbreviations: AMH, anti-Müllerian hormone; INHB, inhibin B; N,
549	number of patients; T, testosterone.