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Increased cardiovascular and metabolic morbidity in patients with 21-hydroxylase deficiency : a Swedish population-based national cohort study

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1 Increased cardiovascular and metabolic morbidity in patients with 21hydroxylase deficiency: a Swedish population-based national cohort study 2 3 4 Henrik Falhammar^{1,2}, Louise Frisén^{3,4}, Angelica Linden Hirschberg^{5,6}, Christina Norrby⁷, Catarina 5 Almqvist^{7,8}, Agneta Nordenskjöld^{6,9,10}, and Anna Nordenström^{6,11} 6 7 ¹Department of Endocrinology, Metabolism and Diabetes, Karolinska University Hospital, Stockholm, 8 Sweden; ²Department of Molecular Medicine and Surgery, Karolinska Institutet, Stockholm, Sweden; 9 ³Department of Clinical Neuroscience, Karolinska Institutet, Stockholm, Sweden; ⁴Child and Adolescent Psychiatry Research Center, Stockholm, Sweden; ⁵Department of Obstetrics and 10 11 Gynaecology, Karolinska University Hospital, Stockholm, Sweden; ⁶Department of Women's and 12 Children's Health, Karolinska Institutet, Stockholm, Sweden: ⁷Department of Medical Epidemiology and Biostatistics, Karolinska Institutet, Stockholm, Sweden; ⁸Lung and Allergy Unit, Astrid Lindgren 13 14 Children's Hospital, Karolinska University Hospital, ⁹Center for Molecular Medicine, Karolinska Institutet, Stockholm, Sweden; ¹⁰Paediatric Surgery, Astrid Lindgren Children Hospital, Karolinska 15 University Hospital, Stockholm, Sweden; ¹¹Department of Paediatric Endocrinology, Astrid Lindgren 16 17 Children Hospital, Karolinska University Hospital, Stockholm, Sweden 18 Abbreviated title: Cardiovascular and Metabolic morbidity in CAH 19 Key words: obesity, diabetes, hypertension, venous thromboembolism, glucocorticoid 20 Counts: Abstract word count: 250; Main text word count: 3443; References: 31; Tables: 5; 21 Supplemental Tables: 3. 22 Corresponding author: Associate Professor Henrik Falhammar, MD, PhD, Department of 23 Endocrinology, Metabolism and Diabetes, D02:04, Karolinska University Hospital. SE-171 76 24 Stockholm, Sweden. E-mail: henrik.falhammar@ki.se Phone: +46-851776411, Fax: +46851773096 25 Grants: This project was supported by grants from the Magn. Bergvalls Foundation, Karolinska 26 Institutet, Stockholm Council and the Swedish Research Council through the Swedish

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30 Abstract

Context: Congenital adrenal hyperplasia (CAH) is lethal in its most severe forms if not treated with
 glucocorticoids. However, glucocorticoids may increase the risk of cardiovascular and metabolic
 morbidity.

34 **Objective:** To study cardiovascular and metabolic morbidity in CAH.

35 **Design, Setting and Participants:** Patients with CAH due to 21-hydroxylase deficiency (n=588;

36 >80% with known CYP21A2 mutations) were compared with controls matched for sex, year, and place

37 of birth (n=58,800). Data were obtained by linking national population-based registers. Subgroup

38 analyses were performed regarding sex, clinical severity (salt-wasting, simple virilising, nonclassic),

39 *CYP21A2* genotype (null, I2 splice, I172N, P30L), and stratified by the introduction of neonatal

40 screening, age-groups, and non-obesity.

41 Main Outcome Measures: Cardiovascular and metabolic morbidity.

42 **Results:** In CAH, both any cardiovascular and metabolic disorders (OR 3.9, 95% CI 3.1-5.0), and

43 cardiovascular disease (OR 2.7, 95% CI 1.9-3.9) were increased. Separate analyses of the individual

44 diseases showed higher frequencies in CAH of hypertension, hyperlipidemia, atrial fibrillation, venous

45 thromboembolism, obesity, diabetes (mainly type 2), obstructive sleep disorder, thyrotoxicosis and

46 hypothyroidism. Similar results were seen in the stratified groups. On the subgroup level, females

were generally more affected (especially I172N and the nonclassic group), as were males with the nullgenotype.

49 Conclusions: CAH was associated with excess cardiovascular and metabolic morbidity but the 50 mechanism is not certain as the glucocorticoids were not assessed. Hypothyroidism and obesity may 51 be an effect of close observation. However, more severe conditions were presumably detected equally 52 in patients and controls. Screening for diabetes and other metabolic disorders which increase 53 cardiovascular risk is important.

54 Introduction

55 Congenital adrenal hyperplasia (CAH) is an autosomal recessive disorder. The most common cause, 56 21-hydroxylase deficiency, is characterised by a reduction of cortisol and aldosterone production and 57 concurrently by increased levels of steroid precursors and adrenal androgens (1-3). There is a wide 58 spectrum of severity of the disease. Untreated CAH is fatal in severe cases due to salt-wasting crises. 59 Women with classic CAH, i.e., the salt-wasting (SW) or simple virilizing (SV) phenotype, have 60 varying degrees of virilization of the external genitalia at birth. Individuals with nonclassic (NC) CAH 61 do not present with cortisol deficiency and may never be diagnosed. If NCCAH is diagnosed, it is 62 generally due to symptoms and signs of androgen excess, including infertility; therefore, usually more 63 females rather than males are diagnosed (1, 3). 64 In most cases, glucocorticoid replacement therapy is necessary for survival in classic CAH. 65 However, the physiological circadian rhythm of cortisol cannot be mimicked with oral 66 glucocorticoids, and the doses needed to suppress the androgens are usually higher than normal 67 replacement (3). Hence, the treatment can be expected to be potentially harmful and may result in an 68 increased risk of obesity, type 2 diabetes, hyperlipidemia and hypertension, i.e., the metabolic syndrome, resulting in cardiovascular morbidity and mortality. So far, neither increased cardiovascular 69 70 morbidity nor an increased prevalence of type 2 diabetes has been found in CAH patients in the few 71 studies reporting on cardiovascular morbidity (4, 5). This is to be expected since glucocorticoids were 72 introduced in the 1950s and very few of the studied patients have been over 50 years of age (3). 73 However, risk factors for cardiovascular disease and an increased risk for metabolic disorders have 74 been reported in both children and adults (6-11). We have previously reported an increased risk for 75 gestational diabetes, a strong risk factor for future type 2 diabetes, in women with CAH (4, 12). In 76 contrast, we could not demonstrate a significant increase in cardiovascular mortality in CAH 77 individuals compared to controls (13). Thyroid disease can increase both cardiovascular and metabolic 78 morbidity (14), but it has not been studied in CAH individuals. 79 The aims of this study were to investigate the cardiovascular and metabolic morbidity in a

80 large population-based national cohort of patients with CAH due to 21-hydroxylase deficiency and to

analyze whether the outcomes varied between the different pheno- and genotype groups, or between
the sexes, as well as stratified by the introduction of neonatal screening, age-groups, and non-obesity.

83

84 Methods

85 Subjects

CAH patients with 21-hydroxylase deficiency and a complete personal identity number born between
1910 and 2009 were identified (n=545) using the national CAH register (15). More than 80% had the
diagnosis genetically confirmed by *CYP21A2* mutation analysis. An additional 43 individuals were
included who had received the diagnosis CAH at least three times in the National Patient Register
(NPR) using the International Classification of Diseases, ICD-8 (255.01, 255.08), ICD-9 (2552, 255C)
and ICD-10 (E25.0) and had not subsequently been given other diagnoses, i.e., Addison's disease,

92 Cushing's syndrome, acromegaly, or received glucocorticoid treatment due to malignancies.

93 Most patients with known CYP21A2 mutations, analysed as previously described (15, 16), 94 were allocated to one of the five most prevalent genotype groups (see below). All the different 95 mutations in this cohort have been described in detail elsewhere (15). The mildest mutation defines the 96 genotype group in compound heterozygotes. Generally speaking, null and I2 splice are associated with 97 the SW phenotype, I172N with SV, and V281L with NC. P30L results in a phenotype between SV and 98 NC, but in this study it was defined as SV. Patients with unknown CYP21A2 mutations were given a 99 clinical classification (SW, SV or NC) if clinical data that clearly could be used for classification were 100 accessible. Genetically verified or clinically diagnosed NC disease was combined to the NC group. 101 The data were also stratified after the introduction of neonatal screening for CAH in Sweden (1986), 102 three different age groups (0-19, 20-39, and older than 40 years old), and non-obesity.

103

104 Characteristics of the included patients and controls

All CAH patients (n=588, females n=335) had been diagnosed with 21-hydroxylase deficiency. The
 median age was 26.0 (range 0–92) years. The clinical phenotype could be determined in 482 patients

107 (82.0%). SW, SV and NC phenotypes were diagnosed in 240 (20.7 years, range 0-69), 167 (27.4 years, 108 range 0.4-79), and 75 (22.1 years, range 3.3-92) patients, respectively. In the most common genotype 109 groups, the numbers were: null, n=100 (19.4 years, range 0.1-57); I2 splice, n=122 (20.6 years, range 110 0-69); I172N, n=130 (26.9 years, range 0.7-79); P30L, n=24 (22.4 years, range 0.4–38); and V281L, 111 n=56 (42 females). A similar number of patients were born before and after the introduction of the 112 neonatal screening programme, but those born before it were older (before, n=305, 40.3 [15-95] years, 113 178 females; after, n=283, 14.8 [0-24] years, 157 females). Controls matched for sex, year and place 114 of birth were included from the Total Population Register (n=58,800). The characteristics of this 115 cohort have been reported previously (13, 17, 18).

116

117 Study protocol

118 A matched cohort design was employed, with exposure defined as having the diagnosis CAH in the 119 national CAH Register or in the NPR. Controls matched for birth year, sex and place of birth were 120 identified in the Total Population Register (100 controls per CAH individual). The Migration Register 121 (Statistics Sweden), with all migrations since 1901, was used to control for migration. The Swedish 122 personal identity number enables unambiguous linkage between population-based registers, including 123 the NPR (maintained by the Swedish Board of Health and Welfare). All participants in this study were 124 given an anonymous code number by Statistics Sweden after linkage with the registers. The NPR 125 contains the discharge diagnoses according to the ICD for both in- and outpatient care since 1964 and 126 2001, respectively. The outcomes were a diagnosis of cardiovascular or metabolic disorders. The 127 different ICD codes (Swedish version) used for the separate analyses are shown in Table 1. The study 128 was approved by the Regional Ethical Review Board in Stockholm, Sweden.

129

130 Statistical analysis

131 A conditional logistic regression model was used to estimate the association between CAH and the

132 outcomes in Table 1. The same outcomes were also estimated for the different subgroups. Odds ratios

133 (ORs) were calculated with 95% confidence intervals (CIs). A CI not surpassing 1.0 was considered

134 significant. SAS (version 9.4) was used for all statistical analyses.

135

136 **Results**

137 Cardiovascular and metabolic disorders

The results are shown in detail for the total cohort and for males and females in Table 2, broken down by phenotype group in Table 3 and *CYP21A2* genotype group in Table 4 and the number of patients in each group is shown in Supplemental Table 1.

141 Any cardiovascular and metabolic disorders were increased in CAH patients, with OR

142 3.9(3.1-5.0) for the whole cohort, 4.4(3.2-6.0) for females and 3.3(2.3-4.9) for males. The increase

remained significant on the subgroup level for SW (both genders), SV (females), NC (both genders),

144 null (males), I172N (females) and P30L (males). For cardiovascular disease, the ORs were 2.7(1.9–

145 3.9) for the total CAH cohort and 3.9(2.5–6.1) for females, but were not significant for the males.

146 They remained significant on the subgroup level for SW (males), SV and NC (females), null (males),

147 and I172N (females).

148 Obesity was increased in both genders (OR 10-15[5.5–19.5]) and in all subgroups of patients, 149 except in I2 splice males and P30L females. Obesity was most pronounced in the NC group (both 150 genders) and the P30L males. Obstructive sleep apnoea was increased in the entire group (OR 2.0[1.0-151 4.1)) and in the SV group. In the P30L male group, it almost reached significance. More obese CAH 152 patients had cardiovascular disease than the non-obese CAH patients (16.3% vs 4.0%, OR 4.6 [1.9-153 11.5]). Two thirds of the obese CAH patients had uncomplicated obesity and no other cardiovascular 154 or metabolic disorders. Three CAH patients (one SW, one NC and one with unknown pheno/genotype) 155 had obesity and five other cardiovascular and metabolic disorders, including type 2 diabetes, 156 hypertension, and acute coronary syndrome or stroke. One CAH patient had obesity combined with 157 hypertension and hypothyroidism while five had obesity combined with hypertension, 158 hyperlipidaemia, or venous thromboembolism. The obese patients with CAH are described in detail in

Supplemental Table 2. Moreover, if only the non-obese CAH patients were compared to their non-obese controls the results were similar to the entire cohort (Table 7).

161 Diabetes was more prevalent in the entire cohort and among females (OR 3.0[1.6-5.8] and 162 4.0[1.8–9.1], respectively) compared to controls. The rate in the subgroups was elevated among 163 females in the SV, NC and I172N groups.

Hypertension occurred more often in females (OR 4.1[2.4–7.3]) and remained significant only
for SV and I172N females. The rate of hyperlipidaemia was increased in the total cohort (OR 2.8[1.2–
6.5]) and in the SW and null (males) subgroups. The frequency of stroke was elevated in only the NC
female group.

Venous thromboembolic events were raised in the entire cohort (OR 3.8 [1.6–8.7]) and SW,
NC, null and I2 splice subgroups. Mostly CAH females were affected.

Acute coronary syndrome was only increased in SW (males) and null (males) groups. Cardiac arrest and heart failure were similar to the findings in controls in all groups. Atrial fibrillation was more frequent in the total cohort (OR 2.3[1.0–5.2]), but was not significant at the subgroup level, except in I172N males. The prevalence of heart block and aortic valve disease was similar to that in controls. One case of cardiomyopathy (SW female, null genotype) and one case of pulmonary heart disease (SW female), but no cases of aortic aneurysm and dissection were found in the CAH cohort (all non-significant compared to controls).

Thyroid disease was increased in both males and females. Thyrotoxicosis and hypothyroidism
were increased (OR 4.7[2.4–8.9] and 3.7[2.2–6.3]), being most pronounced in males (OR 15.8[4.7–
53.4] and 12.9[5.8–28.7]). In a subgroup analysis, thyrotoxicosis was more frequent in SW (males),
NC (both genders) and null (males) subgroups. In the subgroup of hypothyroidism only males in SW,
SV, NC, null and I172N groups were affected.

Patients born before the introduction of neonatal screening showed similar results to those in the entire cohort (data not shown), but those born after 1986, showed an increased frequency of any cardiovascular and metabolic disorders, obesity, hypertension and thyrotoxicosis (data not shown). Compared to controls, those born after the introduction of neonatal screening had higher risk (0.34%
vs 0.04%, OR 7.5[1.7–32.9]) of having hypertension compared to those born before (3.6% vs 1.8%,
OR 1.5[1.0–2.4]). Moreover, when the cohort was stratified into different age groups, the CAH
patients in 0-19 year-old group were mainly affected by obesity and thyrotoxicosis, but females had
more hypertension and males hypothyroidism compared to controls. The older groups of CAH patients

190 also had more cardiovascular disease compared to controls (Table 5).

191

192 Discussion

193 This is the first time established cardiovascular disease, and not merely risk factors, have been 194 investigated in a large cohort of CAH patients. We found increased cardiovascular and metabolic 195 morbidity in CAH patients compared to controls with some subgroups being more affected than others 196 (females generally, and specifically 1172N and NC, and males in the null genotype group). Obesity 197 was consistently increased in all subgroups with the NC group and P30L males being most affected.

198 Even though all the separate measured outcomes, except aortic aneurysm, were increased in 199 CAH individuals, some did not reach statistical significance, probably owing to a lack of statistical 200 power. Although this is the largest CAH cohort ever reported, the median age was relatively low and 201 cardiovascular disease occurs more commonly at an older age. Moreover, in the general population, 202 cardiovascular disease is more prevalent in males, which has been attributed to the higher testosterone 203 levels compared to females. CAH results in elevated androgens, and it could be speculated that this is 204 one of the reasons for the increased cardiovascular morbidity demonstrated in this study, illustrated by 205 the NC group being especially affected. Delayed diagnosis, which is frequently seen in the milder 206 pheno- and genotypes and especially before the introduction of neonatal screening, is frequent and 207 results in prolonged hyperandrogenism. However, once treatment with glucocorticoids has been 208 initiated, androgens are usually decreased compared to controls (4, 19).

209 Obesity was markedly increased, which is consistent with many studies reporting an increased 210 body mass index and/or fat mass in CAH children and adults (4, 5, 8, 11, 20-24). NC individuals and

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211 P30L males were most affected by obesity. However, only a minority of the obese CAH patients had 212 been diagnosed with another cardiovascular or metabolic disorder and the non-obese CAH patients 213 were similarly affected as the entire CAH cohort. Obstructive sleep apnoea is prevalent in obesity, but 214 it has only been reported once in a case of CAH (25), although it could be suspected to occur more 215 frequently. However, we did find an increased frequency of obstructive sleep apnoea in this CAH 216 cohort. Similarly, the frequency of diabetes was increased, especially in females with SV (I172N 217 genotype) or NC phenotype. Decreased insulin sensitivity in CAH children and adults has been 218 reported several times previously (4-6, 9, 23, 26); however, this is the first time a raised occurrence of 219 diabetes has been found. Interestingly, Williams et al. reported insulin resistance only in NC but not in 220 classic CAH children (26), and in a Chinese study on newly diagnosed and untreated young adult 221 females with SV, insulin resistance was found. It has also been claimed that androgens in females can 222 result in insulin resistance (27). Thus, all this taken together could suggest that prolonged postnatal 223 hyperandrogenism, and not only supraphysiological glucocorticoid replacement therapy together with 224 obesity, may cause insulin resistance and diabetes in CAH patients. The doses of corticosteroids used 225 are usually similar in the different pheno- and genotypes (5, 19, 28), and it can be speculated that a 226 more unfavourable profile could be explained by a relative overtreatment considering the milder 227 disease. However, doses of corticosteroids are usually those reported only at the time of the study and 228 the cumulative dose during the entire treatment period is generally unknown. As CAH females have 229 more symptoms of hyperandrogenism compared to CAH males, the females may have been exposed 230 to higher doses of corticosteroids, especially during younger ages, which may explain why females 231 were more affected.

An increased rate of hyperlipidaemia was found, especially in our males with null genotype. Hyperlipidaemia in CAH individuals has been reported in some studies (8, 22), yet, most have found similar lipid profiles, compared to controls (4-6, 11, 26, 29). We found an increased frequency of hypertension in CAH individuals, which is similar to other studies (8, 11, 22, 26, 29), but on analysing the different subgroups, only SV (I172N) females and NC females (tendency) had increased blood pressures, while this was rare or non-existent in the more severe pheno- and genotypes. Most previous studies have not compared the results in different pheno-and genotypes; however, a few have indicated 239 a higher blood pressure in patients with milder forms of CAH, i.e., 1172N and NC groups (5, 26). 240 Moreover, classic adult CAH males were recently reported to have lower blood pressure compared to 241 healthy men (24). Mineralocorticoid replacement is generally mandatory in SW and is often 242 recommended in SV cases to minimise the glucocorticoid doses (1-3), and it is sometimes used even 243 in NC patients (4, 5, 8, 26). A more cautious approach to prescribing mineralocorticoids could 244 possibly be employed, but this has to be investigated in studies where the prescribed doses of 245 mineralocorticoids are known and, ultimately, in randomised controlled trials. Obesity was more 246 pronounced among patients with the milder forms and this may contribute to the development of 247 hypertension. One of the main risks of hypertension is stroke, and we did find an increased occurrence 248 of stroke in the NC female group, but not in the other groups. It has been demonstrated that patients 249 with classic CAH and severe mutations have reduced epinephrine production (null and I2splice), 250 whereas those carrying the milder I172N had normal production (1, 5). It could be speculated that not 251 only a later diagnosis of CAH (5, 26), but also differences in epinephrine secretion, could influence 252 the cardiovascular risk profiles (5).

253 Another risk factor for stroke, atrial fibrillation, was increased in CAH individuals and in the 254 I172N group, but this time only in males. Atrial fibrillation has never been studied in CAH before, but 255 heart rates have been reported to be elevated (5, 29). Alcohol can precipitate atrial fibrillation and 256 alcohol misuse was increased in these CAH males, as reported previously by our group (17); however, 257 this was only significant in the I2 splice group. Thyrotoxicosis can also predispose to atrial fibrillation, 258 and this risk was increased in CAH individuals. The frequency of thyrotoxicosis was extremely high 259 compared to controls in null and NC males, while being only moderately raised in NC females. The 260 prevalence of hypothyroidism was also elevated, especially in the male subgroups. However, thyroid 261 disorders are generally more common in females, which may explain the lower OR in CAH females. 262 Thyroid disorders have not been studied before in CAH. The main cause of both hyper- and 263 hypothyroidism is autoimmunity and the question arises of whether there is an increased risk for 264 autoimmune disorders in CAH. The gene responsible for the 21-hydroxylase enzyme, CYP21A2, and 265 its pseudogene, CYP21A1P, is located in the HLA major histocompatibility complex on chromosome 266 6p21.3, about 30 kb apart, next to the C4B and C4A genes, but there are also other genes involved in

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the immune system located in the vicinity (30). A putative link is purely speculative but should bestudied further.

We were able to show for the first time an increased frequency of venous thromboembolic events in CAH individuals. This could be expected, as both Cushing's syndrome and glucocorticoid use have been associated with venous thromboembolism due to a state of hypercoagulability (31). More liberal use of thrombosis prophylaxis may be warranted.

Patients born after the introduction of neonatal CAH screening seemed to be less affected than those born before, which may indicate a benefit of early diagnosis and/or more optimal corticosteroid replacement therapy in recent years. However, there was a difference in mean age of almost 26 years between the two groups. When stratifying the cohort into different age groups, all age groups were equally affected by any cardiovascular and metabolic disorder. However, the older age groups also had an increased risk of cardiovascular disease while the younger mainly were affected by obesity and thyrotoxicosis; females also had more hypertension and males hypothyroidism compared to controls.

280 The major limitations of this study are that all outcome data were derived from national 281 registers, and the ICD coding may have been inadequate. A prerequisite for obtaining approval by the 282 Ethics Committee was that all individuals included were anonymised to protect their privacy. 283 Therefore, it was impossible to analyse the results on an individual level and compare them with 284 medical files. There may be ascertainment bias as the CAH patients were more likely to be under 285 intensive surveillance compared to controls, which may explain some of the differences, e.g. the 286 observation of hypothyroidism and obesity. However, more severe conditions were presumably 287 detected equally in patients and controls. Moreover, the number of patients with cardiovascular 288 disease was low, in spite of the large number of included individuals with CAH due to the low median 289 age of only 26 years, considering that most cardiovascular events occur in the middle and older age 290 groups. Furthermore, the number of patients in the different severity subgroups was low and some of 291 the ICD codes were used only occasionally. Moreover, the many different subgroup analyses 292 performed will, by definition, give rise to some significant results by sheer chance. Hence, the results 293 from the subgroup analysis must be interpreted with caution. However, the present study probably

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- underestimates the cardiovascular morbidity among patients with CAH as we have recently shown
- from the same cohort that this group died 6.5 years earlier compared to controls, mainly due to adrenal
- crisis (13). Thus, the CAH patients have not had the same risk of developing cardiovascular disease. In
- 297 contrast, the strengths of this study are the unique national registry of CAH individuals covering
- almost all CAH patients diagnosed in Sweden, with most registered patients being both pheno- and
- 299 genotyped, and the almost complete coverage of all discharge diagnoses according to ICD of both in-
- 300 and outpatient care by the National Patient Register.
- 301 In conclusion, CAH was associated with excess cardiovascular and metabolic morbidity. Some
- 302 subgroups seemed to be more affected. Regular follow-up is needed with lifestyle intervention to limit
- 303 the onset of weight gain and obesity, screening for diabetes, other metabolic disorders and
- 304 cardiovascular risk factor. Close monitoring of glucocorticoid doses is important.

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Table 1. The discharge diagnoses from the National Patient Registry analysed according to ICDs for

both in- and outpatient care.

	Diagnosis	ICD 8	ICD 9	ICD 10
1	Obesity	277	278	E66
2	Diabetes mellitus	250	250	E10, E11, E13
3	Type 1 and type 2 diabetes			E10 & E11
4	OSA	347.00, 780.60	347, 780F	G47
5	Hyperlipidaemia	272	272	E78
6	Hypertension	401	401, 405	I10, I15
7	Stroke ^{\$}	430-436	430-436	I60-I64, G45
8	ACS	410,411	410,411	I20-I22
9	Cardiac arrest	427.98	427F	I46
10	Heart failure	427.00, 427.10	428	I50
11	Atrial fibrillation*	427.92	427D	I48
12	Heart block	427.20, 427.28	426	I44, I45
13	Aortic valve disease**	395	424B	I35
14	VTE	450, 451	415, 451	126, 180
15	Cardiomyopathy	425	425	I42
16	Aortic aneurysm & dissection	441	441	I71
17	Pulmonary heart diseases	426	416	I27
18	Hypotension	458	458	I95
19	Thyrotoxicosis	242	242	E05
20	Hypothyroidism	244, 245	244, 245	E03, E06

*includes atrial flutter. **non-rheumatic. ^{\$}includes transient cerebral ischaemic attack. ACS, acute

coronary syndrome, i.e. heart attack and unstable angina. VTE, venous thromboembolism. OSA,

obstructive sleep apnoea, but also includes other occasional sleep disorders. Any cardiovascular and/or

metabolic disorder was defined as no. 1–20, and any cardiovascular disease as no. 6–18.

	САН	Controls	Odds ratio	CAH	Controls	Odds ratio	CAH	Controls	Odds ratio
	individuals		(95% CI)	females	females	(95% CI)	males	males	(95% CI)
n	588	58 800		335	33 500		253	25 300	
Any CVD & meta	99(16.8%)	3460(5.9%)	3.9(3.1-5.0)	62(18.5%)	3460(5.9%)	4.4(3.2-6.0)	37(14.6%)	1496(5.9%)	3.3(2.3-4.9)
Any CVD	44(7.5%)	2103(3.6%)	2.7(1.9-3.9)	30(9.0%)	1109(3.3%)	3.9(2.5-6.1)	14(5.5%)	1008(3.9%)	1.6(0.9-2.9)
Obesity	29(4.9%)	281(0.5%)	10.9(7.4-16.2)	18(5.4%)	170(0.5%)	11.3(6.8-18.7)	11(4.3%)	111(0.4%)	10.4(5.5-19.5)
Diabetes	16(2.7%)	741(1.3%)	3.0(1.6-5.8)	10(3.0%)	375(1.1%)	4.0(1.8-9.1)	6(2.4%)	366(1.4%)	2.1(0.7-6.1)
OSA	8(1.4%)	406(0.7%)	2.0(1.0-4.1)	3(0.9%)	154(0.5%)	2.0(0.6-6.2)	5(2.0%)	252(1.0%)	2.0(0.8-5.0)
Hyperlipidaemia	6(1.0%)	230(0.4%)	2.8(1.2-6.5)	3(0.9%)	110(0.3%)	2.9(0.9-9.5)	3(1.2%)	120(0.5%)	2.7(0.8-8.8)
Hypertension	23(3.9%)	1058(1.8%)	2.6(1.6-4.2)	18(5.4%)	582(1.7%)	4.1(2.4-7.3)	5(2.0%)	477(1.9%)	1.1(0.4-2.7)
Stroke ^{\$}	5(0.9%)	423(0.7%)	1.2(0.5-3.1)	4(1.2%)	253(0.8%)	1.7(0.6-4.9)	1(0.4%)	170(0.7%)	0.6(0.1-4.3)
ACS	6(1.0%)	436(0.7%)	1.5(0.6-3.5)	2(0.6%)	198(0.6%)	1.0(0.2-4.4)	4(1.6%)	238(0.9%)	1.9(0.6-5.7)
Cardiac arrest	1(0.2%)	35(0.1%)	2.8(0.4-21.0)	0(0%)	17(0.1%)		1(0.4%)	18(0.1%)	5.6(0.7-42.0)
Heart failure	3(0.5%)	209(0.4%)	1.5(0.4-5.1)	2(0.6%)	109(0.3%)	2.0(0.4-9.3)	1(0.4%)	100(0.4%)	1.0(0.1-7.9)
Atrial fibrillation*	7(1.2%)	331(0.6%)	2.3(1.0-5.2)	3(0.9%)	156(0.5%)	2.1(0.6-7.1)	4(1.6%)	175(0.7%)	2.5(0.9-7.4)
Heart block	3(0.5%)	120(0.2%)	2.5(0.8-8.0)	1(0.3%)	56(0.2%)	1.8(0.2-13.1)	2(0.8%)	64(0.3%)	3.2(0.8-13.0)
Aortic valve dis**	2(0.3%)	101(0.2%)	2.0(0.5-8.3)	1(0.3%)	52(0.2%)	1.0(0.3-14.7)	1(0.4%)	49(0.2%)	2.1(0.3-15.4)
VTE	6(1.0%)	165(0.3%)	3.8(1.6-8.7)	5(1.5%)	106(0.3%)	5.0(2.0-12.7)	1(0.4%)	59(0.2%)	1.7(0.2-12.4)
Thyrotoxicosis	10(1.7%)	223(0.4%)	4.7(2.4-8.9)	7(2.1%)	204(0.6%)	3.5(1.6-7.6)	3(1.2%)	19(0.1%)	15.8(4.7-53.4)
Hypothyroidism	15(2.6%)	418(0.7%)	3.7(2.2-6.3)	8(2.4%)	362(1.1%)	2.3(1.1-4.6)	7(2.8%)	56(0.2%)	12.9(5.8-28.7)

Table 2. Cardiovascular and metabolic disorders in CAH individuals with 21-hydroxylase deficiency, also divided into females and males, compared with age-

and sex-matched controls (100 controls per case).

CI, confidence interval. ^{\$}Includes transient cerebral ischaemic attack. *Includes atrial flutter. **Non-rheumatic. CVD, cardiovascular disease. meta, metabolic

disorder. OSA, obstructive sleep apnoea. ACS, acute coronary syndrome, i.e., heart attack and unstable angina. VTE, venous thromboembolism. Bold, P<0.05.

Italic, P=0.05-0.09. No odds ratio and CI are calculated when no patient had the condition.

Table 3. Cardiovascular and metabolic disorders in CAH individuals with 21-hydroxylase deficiency divided into the three phenotypes, compared with age- and

sex-matched controls (100 controls per case).	
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		SW		SV			NC		
	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio
	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)
	All	Females	Males	All	Females	Males	All	Females	Males
n	240	135	105	167	91	76	75	56	19
Any CVD & meta	3.2(2.1-4.9)	2.4(1.3-4.5)	4.4(2.4-8.0)	2.9(1.8-4.6)	3.9(2.1-7.0)	1.9(0.9-4.0)	5.6(2.9-10.8)	5.9(2.9-12.3)	4.5(1.0-21.0)
Any CVD	2.7(1.4-5.3)	2.1(0.8-6.0)	3.4(1.4-8.2)	1.8(0.9-3.5)	3.3(1.5-7.6)	0.9(0.3-2.5)	2.9(1.0-8.3)	3.7(1.2-11.2)	
Obesity	10.9(5.8-20.7)	8.6(3.4-21.8)	14.1(5.8-34.4)	6.9(3.0-16.0)	5.9(1.8-19.3)	8.3(2.5-27.3)	17.1(5.7-51.8)	15.3(4.3-54.9)	25.0(2.8-224)
Diabetes	1.8(0.2-13.2)		3.2(0.4-24.2)	3.1(1.2-8.4)	6.2(1.8-21.5)	1.6(0.4-7.5)	4.1(1.2-13.7)	5.5(1.6-19.3)	
OSA	0.8(0.1-5.8)	2.0(0.3-14.7)		2.8(1.0-7.6)	2.2(0.3-16.3)	3.0(0.9-9.7)	2.6(0.4-19.2)	3.3(0.4-24.7)	
Hyperlipidaemia	6.2(1.4-27.3)		11.1(2.3-53.2)	1.6(0.4-6.7)	2.2(0.3-16.7)	1.3(0.2-9.8)			
Hypertension	1.2(0.3-4.9)		2.6(0.6-11.2)	1.7(0.7-3.9)	5.0(1.9-13.1)	0.3(0.0-2.2)	2.8(0.8-10.5)	3.2(0.8-12.6)	
Stroke ^{\$}				0.5(0.1-3.8)		0.8(0.1-6.3)	5.8(1.1-30.8)	5.8(1.1-30.8)	
ACS	5.3(1.2-24.3)		9.9(2.0-50.1)	1.4(0.4-4.9)	2.4(0.3-19.7)	1.1(0.2-5.2)			
Heart failure				2.3(0.5-10.8)	6.3(0.8-48.4)	1.3(0.2-10.9)	8.0(0.7-92.8)	9.1(0.7-114)	
Atrial fibrillation*	2.3(0.3-17.4)		3.8(0.5-29.6)	2.3(0.7-8.2)		3.1(0.8-11.6)	3.6(0.4-30.5)	4.3(0.5-38.5)	
Heart block				3.9(0.9-16.4)	4.7(0.6-35.5)	3.4(0.5-25.3)			
VTE	10.5(3.1-35.3)	13.1(2.9-58.9)	7.4(0.9-59.0)				7.0(0.9-55.5)	8.9(1.1-74.7)	
Thyrotoxicosis	4.5(1.4-14.5)	1.6(0.2-11.7)	33.3(6.7-165)				12.3(4.1-36.9)	8.9(2.6-30.8)	326(12.8->1000)
Hypothyroidism	2.2(0.7-7.1)	0.9(0.1-6.3)	8.7(2.1-37.3)	3.4(1.2-9.3)	1.9(0.5-8.0)	11.9(2.6-53.7)	5.7(1.7-19.0)	3.7(0.9-16.5)	50.0(4.5-551)

CI, confidence interval. ^{\$}Includes transient cerebral ischaemic attack. *Includes atrial flutter. CVD, cardiovascular disease. meta, metabolic disorder. OSA,

obstructive sleep apnoea. ACS, acute coronary syndrome, i.e. heart attack and unstable angina. VTE, venous thromboembolism. Bold, P<0.05. Italic, P=0.05-

0.07. No odds ratio and CI were calculated when no patient had the condition.

I2 splice I172N Null Odds ratio (95% CI) All Females Males All Females Males All Females Males 58 100 59 41 122 67 55 130 72 n Any CVD & meta 5.1(2.8-9.5) 2.4(0.8-6.8)10.9(4.8-24.7) 1.9(0.9-3.8) 2.3(0.9-5.8)1.4(0.4-4.7)3.4(2.1-5.6) 4.9(2.6-9.0) 1.9(0.8-4.5)Any CVD 4.9(1.9-13.1) 2.0(0.3-14.6) 8.9(2.7-29.3) 1.3(0.4-4.1)1.6(0.4-6.8)0.8(0.1-6.9)2.2(1.1-4.4)3.8(1.6-8.8) 1.1(0.4-3.4)Obesity 10.1(4.0-26.2) 6.5(1.5-27.9) 16.3(4.6-58.0) 8.2(2.9-23.3) 11.7(3.4-39.8) 4.3(0.6-32.8) 7.0(2.8-17.6) 7.0(2.1-23.1) 7.1(1.7-30.1) Diabetes 1.1(0.1-7.7)2.3(0.3-16.8)2.0(0.4-9.3)3.7(1.4-10.2)7.1(2.0-25.0) OSA 2.8(0.4-21.1) 1.9(0.3-14.3)4.8(0.6-36.2) 2.7(0.8-8.6)2.6(0.6-10.9)Hyperlipidaemia 11.1(1.3-97.5) 18.1(1.7-188) 1.9(0.4-8.0)2.3(0.3-18.2)1.6(0.2-12.1)Hypertension 1.9(0.3-14.7) 3.5(0.4-29.1) 2.0(0.9-4.7)5.5(2.1-14.7) 0.3(0.0-2.7)Stroke^{\$} 0.5(0.1-4.1)0.8(0.1-7.1)ACS 18.1(1.7-188) 34.6(2.1-579) 1.6(0.5-5.8)2.4(0.3-19.7)1.4(0.3-6.5)Heart failure 2.6(0.6-12.3) $6 \cdot 8(0.9 - 52.0)$ 1.5(0.2-12.6)Atrial fibrillation* 3.3(0.4-25.6) 6.1(0.7-50.8) 2.7(0.8-9.5) 3.8(1.0-14.4)

Table 4. Cardiovascular and metabolic disorders in CAH individuals constituting the four most common CYP21A2 genotype groups compared with age- and

sex-matched controls (100 controls per case). Severity of the genotype ranging from left to right.

18.2(2.0-167)

200(18.1->1000)

14.3(1.8-116)

P30L males with odds ratio (95% CI): Any CV & metab 7.1(1.5-34.8); obesity 7.1(1.5-34.8); and OSA 7.6(0.9-63). Women with P30L genotype had none of the

12.5(2.8-56.6)

2.7(0.4-20.5)

21.6(4.5-104)

2.9(0.4-21.8)

studied disorders. CI, confidence interval. ^{\$}Includes transient cerebral ischaemic attack. *Includes atrial flutter. CVD, cardiovascular. meta, metabolic disorder.

OSA, obstructive sleep apnoea. ACS, acute coronary syndrome, i.e., heart attack and unstable angina. VTE, venous thromboembolism. Bold, P<0.05. Italic,

P=0.05-0.07. No odds ratio and CI were calculated when no patient had the condition.

Heart block

Thyrotoxicosis

Hypothyroidism

10.5(1.3-86.6)

8.8(2.0-39.0)

2.0(0.3-14.9)

VTE

4.1(0.5-31.1)

13.6(2.9-62.9)

5.4(0.7-42.2)

2.5(0.6-10.3)

4.7(1.1-19.7)

4.2(1.5-11.8)

Table 5. Cardiovascular and metabolic disorders in CAH individuals with 21-hydroxylase deficiency in different age groups, also divided into females and

		0-19 years old		20-39 years old			40-92 years old		
	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio	Odds ratio
	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)	(95% CI)
	All	Females	Males	All	Females	Males	All	Females	Males
n	228	122	106	239	143	96	121	70	51
Any CVD & meta	4.2(2.4-7.1)	3.7(1.7-8.1)	4.6(2.2-9.6)	3.7(2.5-5.4)	4.2(2.6-6.6)	3.0(1.5-5.8)	3.9(2.6-5.8)	4.9(2.9-8.3)	2.7(1.4-5.2)
Any CVD	2.2(0.5-10.2)	2.9(0.4-21.6)	1.7(0.2-15.9)	3.8(2.1-6.8)	6.0(3.1-11.5)	1.3(0.3-5.3)	2.2(1.4-3.6)	3.0(1.6-5.5)	1.4(0.6-3.1)
Obesity	12.6(6.4-24.8)	11.1(3.8-32.0)	13.9(5.8-33.4)	9.8(4.9-19.7)	9.3(3.9-21.8)	11.0(3.3-36.9)	11.3(5.8-22.3)	14.0(6.5-30.5)	6.5(1.5-27.7)
Diabetes	2.3(0.6-9.2)	2.1(0.3-15.3)	2.5(0.3-18.0)	1.4(0.4-4.4)	2.5(0.8-8.3)		2.9(1.5-5.6)	3.1(1.3-7.5)	2.7(1.0-7.1)
OSA				1.6(0.4-6.4)	1.8(0.3-13.3)		3.5(1.5-8.0)	3.5(0.8-14.7)	3.5(1.2-9.8)
Hyperlipidaemia							2.6(1.0-6.5)	2.1(0.5-8.5)	3.0(0.9-10.2)
Hypertension	6.7(0.9-50.5)	10.0(1.3-78.1)		5.3(2.4-11.8)	8.6(3.5-20.8)		1.9(1.0-3.4)	2.7(1.3-5.4)	1.0(0.3-3.0)
Stroke ^{\$}							1.1(0.4-3.2)	1.0(0.2-4.5)	0.6(0.1-5.0)
ACS							1.5(0.6-3.6)	2.4(0.3-19.7)	1.4(0.3-6.5)
Cardiac arrest							3.7(0.5-27.9)		8.3(1.1-64.9)
Heart failure							1.8(0.6-6.1)	2.3(0.5-11.0)	1.2(0.1-9.8)
Atrial fibrillation*				2.7(0.4-19.6)		4.1(0.6-30.8)	2.1(0.8-5.4)	2.6(0.7-9.3)	3.8(1.0-14.4)
Heart block	3.1(0.3-37.6)		3.7(0.3-49.2)				3.0(0.7-12.6)	3.2(0.4-23.8)	2.9(0.4-21.9)
Aortic valve dis**							2.9(0.7-12.2)	2.7(0.4-21.2)	3.0(0.4-23.1)
VTE				2.4(0.3-17.6)	3.7(0.5-27.2)		4.6(1.8-11.5)	6.0(2.1-17.4)	2.4(0.3-17.7)
Thyrotoxicosis	28.6(5.9-138)	14.3(1.8-116)	201(18.0->1000)	5.1(2.0-12.6)	3.4(1.1-11.0)	16.4(3.7-73.4)	2.8(0.9-9.0)	2.9(0.9-9.5)	
Hypothyroidism	2.1(0.5-9.6)		7.6(1.4-43.1)	3.4(1.5-7.7)	3.1(1.3-7.7)	5.4(0.7-40.7)	4.8(2.2-10.6)	2.2(0.7-7.1)	26.9(8.4-86.1)

males, compared with their age- and sex-matched controls.

CI, confidence interval. ^{\$}Includes transient cerebral ischaemic attack. *Includes atrial flutter. **Non-rheumatic. CVD, cardiovascular disease. meta, metabolic

disorder. OSA, obstructive sleep apnoea. ACS, acute coronary syndrome, i.e., heart attack and unstable angina. VTE, venous thromboembolism. Bold, P<0.05.

Italic, P=0.05-0.07. No odds ratio and CI were calculated when no patient had the condition.