Functional glucocorticoid receptor gene variants do not underlie the high variability of 17-hydroxyprogesterone screening values in healthy newborns

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Abstract

Objective: 17-Hydroxyprogesterone (17-OHP) screening for classical congenital adrenal hyperplasia (CAH) is part of many newborn screening programs worldwide. Cut-off values are relatively high, and screening sensitivity does not reach 100%. Recently, the glucocorticoid receptor (GR) N363S-variant has been linked to relatively low degree of virilization and comparatively lower 17-OHP serum concentrations in clinically diagnosed female CAH patients. We sought to determine whether functional GR gene variants, either increasing (N363S, BclI) or decreasing GR sensitivity (R23K), underlie the variable 17-OHP screening levels in healthy newborns.

Design: GR genotypes were compared with 17-OHP screening values in 1000 random samples from routine screening. 17-OHP was measured by conventional immunoassay (TRFIA) and a liquid chromatography–tandem mass spectrometry method (LC–MS/MS), which has been shown to increase screening specificity by steroid profiling and avoiding cross-reactions of the 17-OHP-antibody.

Results: There was no significant association of 17-OHP with GR genotypes, even after inclusion of gestational and postnatal age as covariates. However, among LC–MS/MS steroid measurements, we observed some unexpected trends, including lower 11-deoxycortisol concentrations in both 363S- and 23K-carriers. For carriers of the frequent BclI variant, linear regression analysis revealed a significant increase of 4-androstenedione levels with every mutant allele inherited.

Conclusions: Functional GR variants do not underlie the variation of 17-OHP values observed in healthy individuals. However, whether and to which extent genetically determined differences in individual GR sensitivity influence 17-OHP screening levels in conditions of a pathological hypothalamus-pituitary-adrenal gland-axis stimulation and thus may explain false-negative screening results in those affected by CAH remains to be investigated.

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Introduction

Classical congenital adrenal hyperplasia (CAH) due to 21-hydroxylase (CYP21) deficiency is an autosomal-recessive inherited disorder and affects ~1 of 14 000 births worldwide. Clinical symptoms are consequences both of reduced corticosteroid synthesis after and of excessive precursor steroid levels before the enzyme block. The more severe salt-wasting CAH newborns are at risk to develop life-threatening salt-loss crises, and female newborns frequently exhibit severely virilized genitalia, which can lead to gender mis-assignment. Excessive secretion of 17-hydroxyprogesterone (17-OHP) enables detection of CAH already in the neonate. 17-OHP screening has therefore become a valuable tool to prevent newborns from life-threatening salt-loss crises as well as gender mis-assignment (1).

We and others have shown that screening sensitivity does not reach 100% (2–4). This can be put down to a high variability of 17-OHP screening levels in healthy newborns and thus the need of relatively high cut-off values. In conventional screening assays, high cut-off values mainly derive from cross-reactions of the 17-OHP antibody. Nonetheless, a considerable variability of 17-OHP levels in healthy individuals is also seen in extracted steroid assays. Moreover, aside from a relatively strong genotype–phenotype correlation concerning the impairment of mineralocorticoid synthesis in classical CAH, there is a high variability in the degree of external genitalia virilization in affected females (5, 6). Background genetic factors such as individual androgen receptor (AR) sensitivity due to nCAG length polymorphisms within the AR gene have been suggested to explain parts of the phenotypical
variability (7). Interestingly, genetically determined increased glucocorticoid receptor (GR) sensitivity in 363S-carriers of the GR N363S variant has recently been found to be associated with less severe virilization and lower 17-OHP levels in females at the time CAH was diagnosed clinically (8). We therefore speculated whether functional GR variants, by modulating the individual setpoint of the neonatal HPA-axis, might explain the wide range of 17-OHP levels observed in unaffected individuals. To test this hypothesis, we compared neonatal steroid hormone levels including 17-OHP with genotypes for three GR gene variants previously shown to either increase (N363S, BclI) or decrease (R23K) GR sensitivity in vivo (9).

**Genotyping**

DNA was extracted from filter paper according to the protocol by Caggana et al. (11), with minor modifications. Briefly, a 2×2 mm scrap was cut from each screening card by scissors and tweezers with cleaning instruments by ethanol dip and flaming until fiery red in order to avoid cross-contamination. After adding 200 μl methanol, samples were air-dried in a fume hood overnight. We added 50 μl water, then samples were boiled for 10 min and centrifuged for 3 min at 1200 g. We used 1.5 μl of the supernatant as PCR template in a total reaction volume of 20 μl. Genotypes for the GR gene variants N363S (rs6195), R23K (rs6190), and BclI (rs41423247) were determined by standard PCR (primers N363S-F: AGTACCTCTGGAGGACAGAT, N363S-R: GTACCATCTTAAGAAACAGA, R23K-F: CCGATCAGGAAGATAATGTGAC, R23K-R: CAGTAGCTCTTCTTCTAGG; BclI-F: GGCCCATCAGTAATCTACTTGAG) and restriction fragment length polymorphism (RFLP), using restriction enzymes Tas1 (N363S), MnlI (R23K), and BclI (BclI). Reaction conditions are available on request. The amplified regions around N363S and R23K harbor more than one enzyme restriction site. Nevertheless, wildtype, heterozygous, and homozygous samples can clearly be distinguished by RFLP (Fig. 1).

From a total of 1002 samples, genotyping was successful in 1000 samples. Two screening cards could not be included due to insufficient DNA amplification. Approximately 5% DNA-preparations and amplifications have been repeated because of lacking or only weak bands in the first PCR-RFLP set. In addition, PCR-RFLP was repeated with all samples homozygous for 363S or 23K on a 3.0% agarose gel (otherwise 2.5%) to confirm homozygosity.

**Statistical analysis**

Statistical analyses were performed using the SPSS software package, version 14 (SPSS Inc., Chicago, IL, USA). Since most of the variables analyzed did not follow a normal distribution, we used the Mann–Whitney U Test and the Kruskal-Wallis non-parametric statistical analysis to compare differences between groups. For some analyses such as ANOVA, we performed log-transformation in order to obtain normal distribution. Statistical significance was assumed for P values <0.05.

Because of the small number of samples homozygous for 363S or 23K, those heterozygous (+/−) and homozygous (+/+) for the variant allele were grouped (+/− and +/+ ) for comparisons of steroid levels between genotype groups. Data in results and tables are expressed as mean ± S.D.
mutants as a possible source of biased analysis, we allowed linear regression analysis in order to test for values differed with a borderline significance between steroid hormone measurements from LC–MS/MS and GR genotypes, although we observed some unexpected trends: whereas mean 11-deoxycortisol values in carriers of the BclI variant (three homozygous samples), and 39 carried the 23K allele (one homozygous). For the more frequent BclI polymorphism, we found 467 heterozygous and 127 homozygous samples. Genotypes were in Hardy-Weinberg equilibrium; allele frequencies (363S 2.6%; 23K 2.0%; BclI 36.1%) were comparable with those reported previously in Western European populations (8, 9, 12, 13).

Table 1 lists gestational age, birth weight, and age of the babies at the time blood samples for screening had been taken, showing no differences between genotype groups. Mean 17-OHP values from both conventional immunoassay and LC–MS/MS did not show any significant association with GR genotypes when analyzed within the total sample of 1000 screening cards. Similarly, there were no significant associations between steroid hormone measurements from LC–MS/MS and GR genotypes, although we observed some unexpected trends: whereas mean 11-deoxycortisol concentrations appeared to be lower in carriers of the sensitizing 363S allele (P = 0.062), a tendency towards the same direction was also seen in carriers of 23K (P = 0.106) which, in contrast, is considered to decrease GR sensitivity. Contrary to the a priori hypothesis (i.e., increased GR sensitivity would be associated with relatively lower steroid hormone concentrations) we also noted a trend towards higher 4-androstenedione levels in carriers of the BclI variant (P = 0.068). When comparing wildtype samples to only those homozygous for this variant, 4-androstenedione values differed with a borderline significance (P = 0.050). The high frequency of the BclI-variant allowed linear regression analysis in order to test for allele-dose effects. However, a significant effect was seen only on 4-androstenedione levels (P = 0.016); TRFIA-17-OHP: P = 0.083; remaining steroids: P > 0.2.

To exclude compound heterozygosity or even double mutants as a possible source of biased analysis, we excluded all samples carrying more than one GR variant and repeated analysis by comparing the pure carriers of each GR variant with real non-carriers, i.e., those who did not carry any of the three variants analyzed (n = 350). Whereas, only one sample was excluded due to heterozygosity for both 363S and 23K alleles, 13 samples carried both BclI and 363S alleles, and another 17 both BclI and 23K alleles. However, apart from further decreasing 11-deoxycortisol values in 23K-carriers now reaching statistical significance, differences between genotype groups remained small and did not reach statistical significance (Table 2).

Considering the high variation of 17-OHP levels, we divided 17-OHP screening values from both TRFIA and LC–MS/MS into tertiles and compared genotype frequencies between tertiles. Consistent with the trends displayed in Table 1, we noted a trend towards an increasing frequency of 363S-carriers (10, 16, and 22 samples) from the lowest to the highest tertile of TRFIA 17-OHP levels (χ² = 4.732; P = 0.094), whereas the opposite trend was observed when using tertiles of LC–MS/MS 17-OHP levels (23, 14, and 11 samples, resp.; χ² = 5.138, P = 0.077). The distribution of 23K-carriers and BclI genotypes did not differ among the 17-OHP tertiles (data not shown).

Gestational age, birth weight, and timing of the screening are known to affect individual 17-OHP levels in routine screening programs. This was also seen within our sample. Spearman’s rank correlation coefficients regarding TRFIA 17-OHP-values were R = −0.22 (P < 0.01) for gestational age, R = −0.12 (P < 0.01) for birth weight, and R = −0.14 (P < 0.01) for postnatal age respectively. Similarly, significant correlations were observed between LC–MS/MS 17-OHP-measurements and gestational age (R = −0.11; P < 0.01) as well as birth weight (R = −0.12; P < 0.01), albeit the relation between 17-OHP and postnatal age at screening did not reach statistical significance in this assay (R = −0.05; P = 0.99). Not unexpectedly, the inter-assay correlation of 17-OHP screening values obtained from TRFIA and LC–MS/MS
was relatively low (RZ 0.28; P! 0.01), which can be explained by the high cross-reactivity of the 17-OHP antibody with steroids other than 17-OHP. Finally, we added gestational age, birth weight, and postnatal age as covariates in a one-way ANOVA, but again, failed to find statistically significant genotype effects on 17-OHP screening values (P values O 0.15).

**Discussion**

We here addressed the question of whether functional GR variants modify neonatal 17-OHP levels, which are routinely used as newborn screening parameter for classical CAH. In a sample of 1000 randomly selected newborn screening cards, we did not find significant associations of these variants with neonatal 17-OHP. Although numerous clinical studies have underlined the functional relevance of human GR gene variants in both physiological and pathological conditions such as metabolism, body composition, risk of cardiovascular disease or even psychiatric conditions (9, 13), little is known of their biological effects in early infancy. Considering elevated 17-OHP levels in newborns affected by CAH, a well established HP A-axis including feedback response via central GRs can be assumed during early stages of life. In our study, the vast majority of the randomly selected individuals are likely to be not affected by classical CAH. 17-OHP screening values revealed the expected broad variability, which could not be explained by the occurrence of three functional GR gene variants. Considering the recent findings of Luczay et al. which strongly indicate that genetically determined differences in glucocorticoid sensitivity can functionally affect the HP A-axis even before birth (8), one could speculate that N363S mediated effects in newborns become apparent only in conditions of a supraphysiologically stimulated HP A-axis such as in CAH, while being undetectable or at least masked by a physiological variation due to other factors in healthy newborns. A similar hypothesis has recently been suggested by Kuningas et al., who strongly indicate that genetically determined differences in glucocorticoid sensitivity can functionally affect the HP A-axis even before birth (8).

We were aware that even within a sample restricted to screening cards taken from babies born at term (37–42 weeks of gestation) with a screening performed between the second and fourth day of life both gestational and postnatal age represent a significant interfering factor concerning steroid hormone measurements (1, 15). To overcome this, we have repeated several statistical analyses within subgroups of equal gestational and postnatal age (e.g., 38, 39 or 40 weeks only; second or third postnatal day only), but still failed to find statistically significant effects on 17-OHP screening values (P values < 0.15).

**Table 1** Birth characteristics, newborn screening measurements (nmol/l), and glucocorticoid receptor (GR) genotypes within the total study sample (n = 1000). Data are given as mean and s.d. (in brackets).

<table>
<thead>
<tr>
<th>N363S</th>
<th>R23K</th>
<th>BclI</th>
</tr>
</thead>
<tbody>
<tr>
<td>(−/− and +/+)</td>
<td>(−/− and +/+)</td>
<td>(−/− and +/+)</td>
</tr>
<tr>
<td>n=952</td>
<td>n=961</td>
<td>n=406</td>
</tr>
<tr>
<td>Gestational age (wk)</td>
<td>39.33 (1.28)</td>
<td>39.33 (1.28)</td>
</tr>
<tr>
<td>Birth weight (g)</td>
<td>3430 (471)</td>
<td>3430 (468)</td>
</tr>
<tr>
<td>Postnatal age (d)</td>
<td>2.35 (0.49)</td>
<td>2.35 (0.49)</td>
</tr>
<tr>
<td>17-OHP (TRFIA)</td>
<td>21.89 (9.35)</td>
<td>21.99 (9.82)</td>
</tr>
<tr>
<td>17-OHP (LC–MS/MS)</td>
<td>4.17 (1.81)</td>
<td>4.16 (1.83)</td>
</tr>
<tr>
<td>F (LC–MS/MS)</td>
<td>77.63 (79.56)</td>
<td>77.05 (79.56)</td>
</tr>
<tr>
<td>4-A (LC–MS/MS)</td>
<td>2.11 (1.84)</td>
<td>2.10 (1.83)</td>
</tr>
<tr>
<td>S (LC–MS/MS)</td>
<td>2.71 (2.34)</td>
<td>2.71 (2.33)</td>
</tr>
<tr>
<td>21-F (LC–MS/MS)</td>
<td>0.42 (0.24)</td>
<td>0.43 (0.24)</td>
</tr>
</tbody>
</table>
| Birth characteristics, newborn screening measurements (nmol/l), and glucocorticoid receptor (GR) genotypes within the total study sample (n = 1000). Data are given as mean and s.d. (in brackets). *P values given for the variable BclI refer to the comparison of non-carriers (−/−) versus carriers (+/− and +/+ of the variant allele.*
to find significant genotype effects (data not shown). In addition, considering that screening samples in our study were taken at a mean age of 2.4 days, placental derived steroids might still circulate in the infant’s blood (16). While the placenta is not under the influence of the HPA-axis feedback regulation, this may add to the fact that no significant GR genotype effects were found. In contrast to lower steroid hormone concentrations (11-21-deoxycortisol were high, both mean concentration differences in the HP A-axis feedback response (11). Although we do not consider the early neonatal life as an entirely non-stressful period, these findings would rather support the assumption of a relatively low stimulated HPA-axis in 2- to 4-day-old neonates than a constellation comparable with that induced by dexamethasone administration.

Moreover, the molecular mechanisms by which 363S, 23K, and the BclI-variant mediate their modulating effects are still poorly understood, and it is not clear whether these variants modulate glucocorticoid sensitivity in a qualitatively comparable manner. N363S and R23K lie within the transactivation domain encoded by exon 2, and in vitro experiments revealed differences in their transactivating capacities as compared with the wildtype protein (19). For N363S, marked changes in gene expression were detected by microarray studies even without ligand incubation (20). Conventional transfection assays are precluded in the case of the BclI-variant due to its intrinsic location. Because BclI, in addition, is not located near a known splicing site, linkage to other functional polymorphisms may underlie its phenotype. Indeed, in the study of Rautanen et al., who investigated the relationship of six GR haplotypes to basal cortisol secretion, the BclI-variant was part of two haplotypes, but only one of these two was associated with higher basal cortisol levels (21). Interestingly, steroid profiling in our study revealed a significant genotype effect for BclI on 4-androstenedione levels, whereas mineralocorticoid and glucocorticoid precursors seemed to be not or only marginally affected. Although the standard deviations for measurements of 4-androstenedione, 11-deoxycortisol, and 21-deoxycortisol were high, both mean concentration in (nmol/l) and the respective s.d. were largely comparable for these three steroids. Thus, this finding may point to an underlying mechanism considerably distinct from those mediated by N363S and R23K.

A limitation of our study is that it does not include a CAH-control group. Only by comparing individual screening values and steroid measurements from the immediate diagnostic work-up within a large cohort of CAH patients, would it be possible to transfer the findings of Luczay et al., whose data derive from patients

### Table 2

Mean steroid hormone concentrations (nmol/l) and s.d. (in brackets) in an adjusted model comparing real non-carriers (3 open circles) versus pure carriers (1 filled circle).

<table>
<thead>
<tr>
<th></th>
<th>Noncarrier</th>
<th>N363S</th>
<th>R23K</th>
<th>BclI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n=350</td>
<td>n=34</td>
<td>n=21</td>
<td>n=564</td>
</tr>
<tr>
<td>17-OHP (TRFIA)</td>
<td>21.20 (9.19)</td>
<td>24.74 (17.76)</td>
<td>0.143</td>
<td>21.32 (8.83)</td>
</tr>
<tr>
<td>17-OHP (LC–MS/MS)</td>
<td>4.15 (1.77)</td>
<td>3.91 (1.90)</td>
<td>0.410</td>
<td>3.98 (1.40)</td>
</tr>
<tr>
<td>21-F (LC–MS/MS)</td>
<td>1.93 (1.67)</td>
<td>1.95 (1.84)</td>
<td>0.841</td>
<td>1.37 (1.43)</td>
</tr>
<tr>
<td>F (LC–MS/MS)</td>
<td>75.21 (72.26)</td>
<td>68.93 (70.79)</td>
<td>0.496</td>
<td>69.33 (66.44)</td>
</tr>
<tr>
<td>21-F (LC–MS/MS)</td>
<td>2.00 (1.76)</td>
<td>1.95 (1.86)</td>
<td>0.726</td>
<td>1.76 (1.57)</td>
</tr>
<tr>
<td>S (LC–MS/MS)</td>
<td>2.63 (2.25)</td>
<td>2.44 (2.09)</td>
<td>0.698</td>
<td>1.53 (1.78)</td>
</tr>
<tr>
<td>21F + 17OHP/F (LC–MS/MS)</td>
<td>0.41 (1.35)</td>
<td>0.43 (0.79)</td>
<td>0.671</td>
<td>0.24 (0.27)</td>
</tr>
</tbody>
</table>
who were diagnosed clinically at a mean age of 1–3 months, from early infancy to the neonatal stage. In addition, disease modifying effects, if any, for R23K and the Bcll-variant in CAH-patients remain to be investigated. Recently, another genomic variant (exon 9 β-polymorphism, rs61998) has been described, which similar to R23K, is related to a relative GR resistance. To date, the molecular mechanisms behind this effect are still incompletely understood. Stabilization of GRβ mRNA resulting in an accumulation of the GRβ protein with a dominant-negative effect on GRα action may play a role (22–24). This polymorphism is linked to R23K but also exists alone. With an allele frequency of up to 15% in European populations, this variant may provide additional information in future studies of individual glucocorticoid sensitivity and its possible relation to the incomplete genotype–phenotype correlation in CAH.

To conclude, our results show that the GR polymorphisms N363S and R23K as well as the more common Bcll-variant do not underlie the wide variation of 17-OHP screening values observed in healthy individuals, which is considered a major cause of the only reduced sensitivity of newborn screening programs for CAH. However, whether and to which extent genetically determined differences in the individual GR sensitivity influence neonatal 17-OHP levels in conditions of a pathological HPA-axis stimulation and thus may explain false-negative screening results in infants affected by CAH remains to be investigated.

Declaration of interest
The authors declare that they have no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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