Molecular genetic testing of congenital adrenal hyperplasia due to 21-hydroxylase deficiency should include CAH-X chimeras

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To the Editor:

We read with great interest the European Molecular Genetics Quality Network practice guidelines for molecular genetic testing of congenital adrenal hyperplasia (CAH) due to CYP21A2 defects [1]. The authors address multiple important issues in the molecular genetic testing of CAH and provide valuable suggested standards based on extensive experience and practices from laboratories worldwide. While in general we agree with the workflow and methods described in the guidelines, we appeal to include evaluation for a contiguous gene deletion syndrome, termed CAH-X, within the scope of the testing. We believe this would be particularly beneficial for individuals carrying a CAH genotype of “30 kb deletion”.

As illustrated in Fig. 2 of the article by Baumgartner-Parzer et al. [1], the terms “30 kb deletion” or “large gene conversion” refer to a CAH genotype of chimeric genes caused by unequal crossover during meiosis due to the structural complexity of the CYP21A2 locus. These deletions account for 20–30% of all CAH alleles and are categorized into two subtypes determined by the location of the junction sites: CAH and CAH-X chimeras [2]. CAH chimeras (CYP21A1P/CYP21A2) are caused by recombination between CYP21A1P and CYP21A2, and impair only CYP21A2; CAH-X chimeras (CYP21A1P-TNXA/TNXB) are caused by recombination between TNXA and TNXB, null the entire CYP21A2 and replace the 3′ of neighboring TNXB with the pseudogene TNXA. This contiguous deletion is hence termed CAH-X, named for both

CAH-X syndrome is composed of two elements: CAH and hypermobility type EDS; the former is autosomal recessive but the latter appears to be autosomal dominant. If carrying a CAH-X allele, phenotypic EDS is expected regardless of the CAH status [2], although in general CAH patients have more severe EDS manifestations than their carrier counterparts [4, 5]. The hypermobility EDS spectrum includes joint hypermobility, arthralgias, frequent joint dislocations, hernias and midline defects including cardiac structural abnormalities. Overall, about 25% of reported CAH-X patients have cardiac abnormalities including congenital heart defects such as structural valve abnormalities, left ventricular diverticulum, and patent foramen ovale [8].

To date, the clinical diagnosis of EDS in CAH-X mostly relies on evaluation of joint hypermobility and subluxations, which are often restricted by factors such as age and time of
onset. Genetic testing for this germline condition in newborn screening is advantageous in terms of offering very early diagnosis to young children before hypermobility evaluation is applicable, and to enable early screening for cardiac defects. In older individuals undergoing CYP21A2 genotyping, identification of CAH-X would provide information regarding the cause of any ongoing connective tissue abnormalities and guidance regarding preventative measures. Treatment for hypermobility type EDS depends on the signs and symptoms present, and a screening echocardiogram should be considered.

To date, most CAH genetic test platforms determine the status of "30 kb deletions". Adding a CAH-X test selectively to "30 kb deletion" positives can be a pinpoint addition. Sharaing a ride with established DNA extraction and even PCR would optimize laboratory efficiency and cost-effectiveness. For instance, laboratories that use a classic CYP779f/tena32F primer pair for PCR can use the same PCR product to conduct Sanger sequencing tests of CAH variants, 30-kb deletions and CAH-X chimeras [9, 10]. A high throughput CAH-X screening has been developed, which has shown excellent accuracy. It is amenable to either qPCR or digital PCR, thus it is fast and easy to operate [6]. If MPLA methodology is used, an up-to-date version now simultaneously checks “30 kb deletion” and a CAH-X chimera of haplinsufficiency [1, 7], therefore labs would only need to test the CAH-X chimeras of misfolding on “30 kb deletion” positives to complete the CAH-X test by either Sanger or qPCR/digital PCR methodologies.

In summary, CAH-X chimeras account for a large portion of “30 kb deletion” CAH alleles and have historically been under-detected. Regardless of their CAH status, individuals who carry a CAH-X allele are expected to suffer from a spectrum of hypermobility EDS manifestations, including cardiac structural defects, joint hypermobility and instability, hernias and gastrointestinal disorders. Including CAH-X determination in CAH molecular genetic testing platforms is convenient and cost-effective, and is particularly beneficial for individuals carrying a “30 kb deletion” allele because many of them are in fact CAH-X. An early diagnosis of CAH-X syndrome is valuable for preventative care and long-term medical management.

Compliance with ethical standards

Conflict of interest DPM received unrelated research funds from Diurnal Limited through the National Institutes of Health Cooperative Research and Development Agreement. Both authors report no potential competing interests in this work.

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