Genetics Corner: Trichothiodystrophy 1 Causes Neutropenia in an Infant with Congenital ichthyosis and Brittle Hair

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Clinical history:
A 10-month old male infant was referred for genetic evaluation for trichothiodystrophy-1.

He was delivered at ~32 weeks' gestation by C-section when his 23-year old G2 P1-->2 mother developed HELLP syndrome. His prenatal course had been complicated by IUGR and decreased fetal movements. Birth weight: 1474 grams (17th %ile). Birth length: 41.9 cm (33rd %ile). Apgar scores were 9 and 9 at 1 and 5 minutes, respectively. He had taut, shiny skin on his upper and lower extremities at birth, resembling collodion, which peeled during resuscitation. He was transferred from the birth hospital to the NICU at a nearby tertiary care center because of congenital ichthyosis, left eyelid fusion, and right cryptorchidism. His CBC and differential were normal.

At 18 days of age, the results of a rapid whole genome sequencing test identifed biallelic variants in ERCC2 (or XPD) c.2164C>T (p.Arg722Trp): pathogenic, paternally inherited c.5+5G>A: a splicing variant of uncertain significance, maternally inherited

The infant was discharged on day 23 of life with a diagnosis of trichothiodystrophy-1.

At ten months of age, he was in generally good health. His mother had rescheduled his genetics clinic visit the previous month, and she had no particular concerns. She was satisfied that his "skin condition" was being managed and was not severe. He had a follow up (video) appointment with a dermatologist at six months of age. An ophthalmology evaluation revealed dry eyes but no cataracts. An otolaryngologist had seen him for recurrent impacted cerumen.

Genetic evaluation:

The infant had been mostly at home without much contact outside his immediate family because of the coronavirus restrictions. He had a history of one prior febrile illness with symptoms of a cold lasting about two weeks; he recovered but still had a persistent cough. He babbled and cooed. He had good eye contact, a social smile, and turned to his name. His mother reported that his developmental progress was "normal," but at ten months of age, he could only maintain a seated position for a few seconds.

The family history was noncontributory. The non-consanguineous parents, mother, age 24, and father, age 25, were of Mexican ancestry. They had one other child, a healthy 4-year old boy.

On physical examination, the growth parameters were poor for a 10-month-old male. Weight <1 %ile (Z = -2.47). Length <1 %ile (Z = -3.45), BMI 16.27 kg/m², 30 %ile (Z = -0.54). The baby was socially aware, interactive, and in no distress. He had deep-set eyes, down-slanting palpebral fissures, and bilateral epicanthal folds. There were sparse eyebrows and eyelashes (Figure 1). The scalp hair was sparse (Figure 2) with thick, waxy flakes and scales on the scalp. He had patchy areas of ichthyosis, mostly on the trunk and antecubital fossa. He had shiny skin on his palms, thin, dystrophic nails (toes>fingers), and shallow, short, concave nails.

The geneticist referred him to Pediatric Hematology and Pediatric Immunology, recommended RSV vaccination, and ordered an absolute neutrophil count (ANC). He had neutropenia: WBC 5.2, hemoglobin 12.8, platelets 312, ANC 0.4 (normal range 2.2-9.2 bil/L), but the ANC had normalized two weeks later at 6.6.

At 11 months of age, he was hospitalized briefly for neutropenia (ANC 0.6), dry cough, and fever to 101.7 degrees without a documented source of infection (cultures were negative). The source of the infection was presumed to be viral. His differential had lymphocyte predominance without excess bands. His growth parameters then were still generally small for his age: Ht 70.5 cm, 2 %ile (Z = -2.10), Wt 7.11 kg, <1 %ile (Z = -2.71), HC 44 cm, 6 %ile (Z = -1.55), BMI 14.31 kg/m² 2 %ile (Z = -2.09). At 13 months of age, he had an in-person visit with the pediatric dermatologist who documented tiger-tail banding on polarized trichoscopy (Figure 3).

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Assessment and counseling:

ERCC2 is a protein that acts as a DNA helicase in transcription-coupled nucleotide excision repair and phosphorylation of nuclear receptors. Cells use nucleotide excision repair to fix damaged DNA, including damage from ultraviolet radiation. Pathogenic variants in ERCC2 (XPD) are associated with a group of autosomal recessive disorders, including trichothiodystrophy (TTD) with sulfur-deficient brittle hair and developmental abnormalities but without skin cancer, xeroderma pigmentosum (XP) with pigmented abnormalities and increased skin cancer, XP/TTD with combined...
Signs and symptoms of TTD vary widely and include ichthyosis, abnormal fingernails and toenails, developmental delays, intellectual disability, dysmyelination, and cerebral atrophy. Some patients with neurological abnormalities are described to have an outgoing, sociable personality, delayed growth, short stature, dry eye, microcornea, nystagmus, congenital cataracts, neutropenia, recurrent, sometimes life-threatening infections, and decreased fertility. Mild cases may involve only brittle, sulfur-deficient hair that displays a diagnostic alternating light and dark banding pattern, called ‘tiger tail banding’ (Figure 3).

More severe presentation of TTD includes delayed development, significant intellectual disability, and recurrent infections. Severely affected individuals may have limited survival. Increased mortality is due to infection. In 2008, Faghri and colleagues published a comprehensive review of 112 patients with TTD, documenting recurrent infections in 35%. They noted early mortality with a 20-fold increased risk of death before the age of 10 years, the majority secondary to infection (pneumonia or sepsis). Randall and colleagues (2018) encourage IVIG or granulocyte colony-stimulating factor therapy for patients with TTD who have severe neutropenia or low levels of IgG. RSV immunization is recommended (personal communication, D.Tamura, NIH) because of the increased mortality associated with this viral infection among infants with TTD. The family was offered enrollment in a clinical trial at the NIH on TTD, xeroderma pigmentosum, and Cockayne syndrome.
Tamura and associates (2011) found a high risk of pregnancy and neonatal complications in their cohort of 27 TTD-associated pregnancies in 23 mothers. There were pregnancy complications in 81%: 56% had a preterm delivery, 30% had preeclampsia, 19% had placental abnormalities, 11% had HELLP syndrome, and 4% had an emergency c-section for fetal distress, while 44% had two or more complications. Only 19% of the pregnancies delivered at term without complications. Eight of the ten pregnancies tested had multiple abnormal screening markers, including elevated human chorionic gonadotrophin levels. Eighty-five percent of the neonates had complications: 70% were low birth weight (<2500 g), 35% had birth weight <10th centile for gestational age, 70% had NICU admission, 67% had a collodion membrane, and 31% of the 16 males had cryptorchidism. Cataracts were present in 54% of the TTD patients examined.

Although the diagnosis of this rare disease, trichothiodystrophy, was made quickly in this baby, he did not fully benefit from having an early diagnosis in the newborn period. He was discharged home from the NICU shortly after the diagnosis was made, and the focus of his subsequent care was on treating his ichthyosis, which was felt to be the major problem of his disorder. The mildness of his skin problems may have falsely reassured his family and his care team. The primary threat to his health was neutropenia and a high risk of infection; these were not appreciated until we alerted his care team. As rapid whole-genome sequencing becomes more available, neonatologists will have to rise to the challenge to realize the benefits and care coordination of early diagnoses for their patients.

**Practical applications:**

1. Understand that the benefits of a rapid genome sequencing test in the neonatal period may not be fully realized without careful phenotyping and coordinated multispecialty follow-up

2. Appreciate that pathogenic variants in one gene can cause several disorders.

3. Realize that trichothiodystrophy poses severe risks of neutropenia, infection, and increased mortality for affected infants. Offer RSV immunization to infants with trichothiodystrophy and follow for neutropenia.

4. Recognize that HELLP syndrome in the mother can be a sign of a genetic disorder, such as trichothiodystrophy, in the fetus.

**References:**


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OPIOIDS and NAS
When reporting on mothers, babies, and substance use
LANGUAGE MATTERS

I am not an addict.
I was exposed to substances in utero.
I am not addicted. Addiction is a set of behaviors associated with having a Substance Use Disorder (SUD).

I was exposed to opioids.
While I was in the womb my mother and I shared a blood supply. I was exposed to the medications and substances she used. I may have become physiologically dependent on some of those substances.

NAS is a temporary and treatable condition.
There are evidence-based pharmacological and non-pharmacological treatments for Neonatal Abstinence Syndrome.

My mother may have a SUD.
She might be receiving Medication-Assisted Treatment (MAT). My NAS may be a side effect of her appropriate medical care. It is not evidence of abuse or mistreatment.

My potential is limitless.
I am so much more than my NAS diagnosis. My drug exposure will not determine my long-term outcomes. But how you treat me will. When you invest in my family’s health and wellbeing by supporting Medicaid and Early Childhood Education you can expect that I will do as well as any of my peers!

Learn more about Neonatal Abstinence Syndrome at www.nationalperinatal.org

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