

RAISE YOUR VOICE FOR RETT SYNDROME

RETT SYNDROME IS A RARE GENETIC NEUROLOGICAL DISORDER

1 IN 10,000 ARE AFFECTED BY RETT

CAUSED BY A MUTATION OF THE MECP2 GENE ON THE X CHROMOSOME

THE HALLMARK OF RETT IS NEAR CONSTANT REPETITIVE HAND MOVEMENTS

THERE IS CURRENTLY NO CURE

TO LEARN MORE VISIT
WWW.RETT.CA



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We are
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to Support
RETT SYNDROME