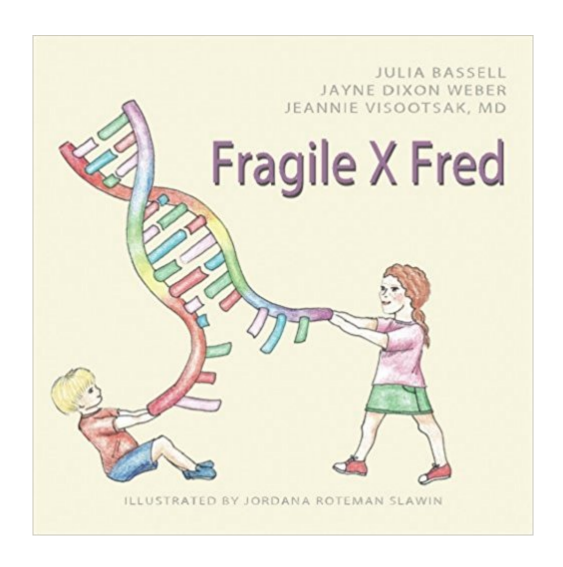


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Fragile X Fred





Synopsis

Look around you. Every person you meet has been molded by an intertwining of genetic information and personal experience. Fragile X syndrome is a genetic disorder that is not widely understood. This book aims to help people of all ages understand that although individuals with fragile X syndrome do have certain challenges, these challenges do not define them. This book is a valuable tool for educators, siblings, parents, and children with fragile X syndrome.

Book Information

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Fitness & Dieting > Diseases & Physical Ailments > Genetic

Customer Reviews

Jayne Dixon Weber, is the Director of Support Services at the National Fragile X Foundation. In addition to assisting with the development of the National Fragile X Foundationââ ¬â,¢s ââ ¬Å*Adolescent and Adult Project,ââ ¬Â*Jayne authored the book Transitioning â⠬ʜSpecialââ ¬â,¢ Children into Elementary School and was editor of the book Children with Fragile X Syndrome: A Parentsââ ¬â,¢ Guide. She is the co-leader of the Colorado Fragile X Community Support Network (parent support) group. Jayne has two childrenâ⠬⠕one, an adult son with fragile X syndrome, and the other, a daughter, who is an occupational therapist. She likes to read, enjoys photography, and goes for a walk every day. Jeannie Visootsak, MD, FAAP, Associate Professor, is a board-certified Developmental-Behavioral Pediatrician at Emory University School of Medicine, Atlanta, GA. She is the Medical Director of the Fragile X Syndrome Clinic and Fragile X Syndrome Clinical Trial Unit at Emory University. The Fragile X Syndrome Clinic was established in 2004 to meet the needs of individuals with fragile X syndrome and their family members. Dr. Visootsak co-founded the Fragile X Clinical and Research Consortium in collaboration with the National Fragile X Foundation. She is also a member of the National Fragile X

Foundationââ ¬â,,¢s Scientific & Clinical Advisory Committee and Executive Committee Council. Additionally, she has received the Jarrett Cole Clinical Award for dedicated service to families in the worldwide Fragile X community. Julia L. Bassell, is a current pre-medical undergraduate student majoring in Neuroscience and Behavioral Biology at Emory University. She has conducted clinical research on fragile X syndrome and Down syndrome, and volunteers with Best Buddies and Running Mates.

This book is a helpful resource for families who are dealing with a fragile X diagnosis. I look forward to reading it to my son's kindergarten class, and all of his classes in the future! A huge thank you to the authors for writing such a delightful, informative, comprehensive book!

Julia is a promising author and on the path to being a brilliant scientist. This is a marvelous book written to help parents, educators and children with fragile x syndrome.

Just what I have been looking for! Ordered 10 more!

Super book to help people understand Fragile X Syndrome.

different but good

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