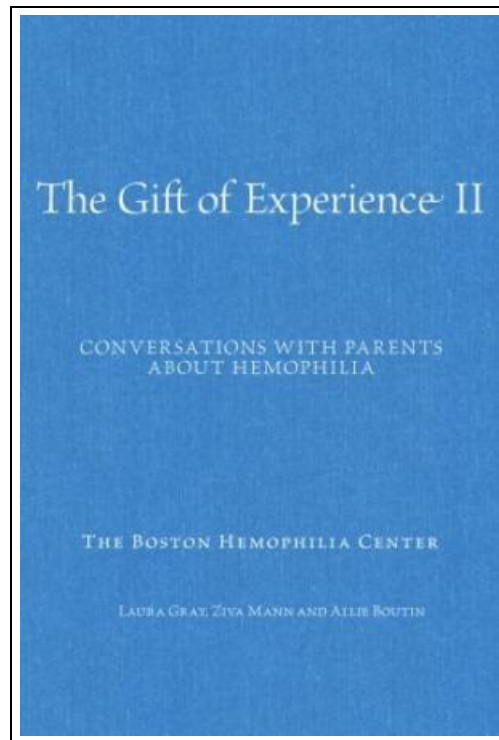


The Gift of Experience II Conversations with Parents about Hemophilia



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Reviews

It is simple in read easier to understand. I am quite late in start reading this one, but better then never. Its been designed in an exceptionally easy way in fact it is just following i finished reading through this publication where basically transformed me, alter the way i really believe.

(Ms. Christy Ondricka DDS)

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CreateSpace Independent Publishing Platform. Paperback. Condition: New. This item is printed on demand. 262 pages. Dimensions: 9.0in. x 6.0in. x 0.6in. The outlook for children with hemophilia keeps getting better. Treatments are safer, easier to administer and new, longer-acting products are becoming readily available. Specialists at comprehensive hemophilia centers all around the country offer innovative treatments while advancing national standards of care. Experts at these centers give patients and their families access to the latest clinical and research developments. By emphasizing the importance of early diagnosis and intervention to prevent complications, these specialists create an enormous impact on patients health and long-term well-being. So its safe to say that there is every reason in the world for parents of children who have been diagnosed with hemophilia to be hopeful about their childs future. Nevertheless, having a child with a chronic illness presents many challenges. There is much to learn, much to understand, a lot to teach and a new reality to adjust to. Concerns about bleeds, joint damage, inhibitors and pain are very real. We interviewed parents whose children are treated at the Boston Hemophilia Center about their experiences. In this book they share what its been like to raise a child with hemophilia from birth to age 6. Through their stories, we appreciate that the journey from receiving an initial diagnosis to feeling capable and in control of the situation can take more than connecting with a team at a treatment center. Home care nurses, other families who have children with hemophilia and myriad of other resources all play important roles in a newly diagnosed familys capacity to cope. These families understand what its like to have a child with hemophilia like no one else can. The guidance, understanding and practical tips they offer from their own experience are gifts...

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