

Heart Warriors: A Family Faces Congenital Heart Disease



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Reviews

*This book is great. I have go through and so i am confident that i will going to read through once again again in the future. I am just easily can get a satisfaction of looking at a written book.
(Miss Vernie Schimmel)*

HEART WARRIORS: A FAMILY FACES CONGENITAL HEART DISEASE



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Behler Publications, United States, 2012. Paperback. Book Condition: New. 213 x 137 mm. Language: English . Brand New Book. Five months pregnant, Amanda Adams and her husband were given two abysmal choices regarding her pregnancy: force her baby to fight for his life through countless invasive and dangerous surgeries, or perform a late term abortion. Despite the fact that Liam was missing half his heart, Amanda chose life. Amanda's emotional plate was full as she found herself redefining the usual expectations a mother has for her child. Instead of wondering where he'd go to college, she wondered if he would survive his first birthday. The eventual acceptance of Amanda's grief helped her accept her new role as a powerful advocate. Over the course of seven years, together, as a family, Amanda and her husband helped Liam endure twelve heart surgeries, each time taking him to the brink of death. Heart Warriors is Amanda's personal and emotional story that initiates a powerful dialogue about infant mortality and hope. Includes a Parent s/Family Field Guide. Amanda Adams is a Heart Mom and a powerful voice within the congenital heart disease (CHD) community by working closely with cardiologists, cardiothoracic surgeons, and nurses from Harvard University, Baylor University, University of Southern California, University of Illinois, and CU Boulder Medical. She founded two CHD organizations, organizes medical conferences for Hypoplastic Right Hearts, attends medical advisory board meetings and national chapter presidents meetings for the Children's Heart Foundation, and chaired the Congenital Heart Walk in Colorado. As a pioneer of CHD advocacy, she is also firmly rooted in the social network of CHD parents and.



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