

**Testimony and Support of HB 2399**  
**House Committee on Health and Human Services - March 5, 2025**

**Karey Padding, LMSW**  
**Parent Advocate and Cystic Fibrosis Mom**  
**Kansas Council on Developmental Disabilities - Chairwoman**  
**Kansas Newborn Screening Advisory Board Member**  
**Associate Executive Director, Cystic Fibrosis Foundation – Heart of America**  
**Chapter - Wichita**

My name is Karey Padding, and I ask you to support HB 2399 because all babies born in the great state of Kansas deserve a healthy start. Newborn screening is truly one of the 20th century's great public health success stories. Newborn screening saves lives and can drastically change the trajectory of life for babies and their families.

An efficient and effective newborn screening program is important to me because I became a mother for the first time on a warm November day in 2010. A quick and simple test approximately 24 hours after Gavin James was born, the newborn screen consisting of a small heel stick and blood spot test, would make a big difference for us in the weeks and months to come. A few days after we arrived home from the hospital, we received a call from our pediatrician's office asking us to bring Gavin back in because one of his tests came back abnormal. We were scared and a bit sleep-deprived, but of course, we complied, and the blood spot test was redone. A few days later, we received a call from our pediatrician, confirming the abnormal screen for cystic fibrosis. We were shocked, devastated, scared, and had so many questions. "What does this mean for Gavin? What do we do next?" We were quickly set up with an appointment at the CF Clinic in Wichita to have Gavin undergo a sweat chloride test, which ultimately confirmed his diagnosis. After the first care center visit, Gavin began a daily regimen of enzymes, inhalers, nebulizers, and CPT (chest physical therapy). A daily regimen he continues to this day. I couldn't believe my three-week-old baby was eating applesauce off of a spoon and taking enzymes with every meal – he wasn't even holding his head up yet!

It took many months to reflect on our birth and diagnosis story to realize how truly lucky we were that Gavin's first few weeks of life set the course for the best possible health outcomes, and it all started with the newborn screen identifying his cystic fibrosis. I have several CF mom friends who have older children who were not screened in Kansas or not screened for CF initially whose babies struggled with their weight, and health and endured numerous medical appointments and hospitalizations before the real cause of their failure to thrive was finally identified as cystic fibrosis. It caused significant financial strain for families as well as taking a toll on their emotional, mental, and physical health.

Gavin's story is a personal success story and a public health success story. Gavin is growing, thriving, and healthy because of an early diagnosis and treatment, and it all began with a few drops of blood and a hearing test to set my baby on a healthy path. In my professional life, I am

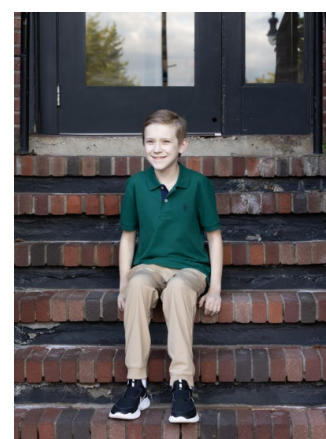
the Associate Executive Director of the Cystic Fibrosis Foundation – Heart of America Chapter in Wichita, KS. I am so thankful for the Newborn Screen and the early detection of Gavin’s cystic fibrosis. I believe he continues to grow today because preventative treatments were started before symptoms.

HB 2399 allows all newborn Kansans to continue having the advantage of early diagnosis of metabolic and genetic diseases which greatly benefit from early treatments and interventions. Each year, approximately 35,000 KS newborns are provided an initial screening, and approximately 2,000 will require repeat screenings to complete the screening process. Please consider increasing the capitation to expand this lifesaving and life-changing program to continue to meet the needs of the many families served through this screening program. I urge you to support House Bill 2399 and I thank you for allowing me to share my story with you today.

Karey Padding, LMSW (CF Mom, Triplet Mom, Concerned Kansan)



Gavin – Age 14 – lives with cystic fibrosis and autism



Padding Triplets – Age 11 and Gavin – Age 14



Padding Family – 2024 Family Photo

