Polycystic kidney disease (PKD) is a chronic, genetic disease causing uncontrolled growth of fluid-filled cysts in the kidneys, often leading to kidney failure. There is no cure. The only life-extending options are dialysis and kidney transplant. PKD affects all racial and ethnic groups equally and affects children as well as adults. PKD in children devastates families and dramatically affects quality of life.

There are two types: autosomal dominant PKD and autosomal recessive PKD.

**ADPKD**  
Autosomal dominant polycystic kidney disease  
Fluid-filled cysts develop and enlarge in kidneys, eventually leading to end-stage renal disease in 50 percent of patients by the age of 50.

**ARPKD**  
Autosomal recessive polycystic kidney disease  
Kidney cysts develop, the liver becomes fibrotic and prenatal lung development can be delayed.

Approximately 600,000 people in America and 12.4 Million people in the world have ADPKD.  
1 in 20,000 children have ARPKD.

Georgia was born with ADPKD.  
Ava was born with ARPKD.
ABOUT ADPKD

Autosomal dominant polycystic kidney disease (ADPKD) can be diagnosed at a very young age or even before birth.

Diagnostic and/or genetic tests performed during pregnancy should be done in conjunction with medical counseling so the test results can be completely understood. For example, knowing your baby may have an ADPKD gene does not determine the course or severity of the disease. There are two different groups of children with ADPKD — those diagnosed before birth or in their first year of life with large cystic kidneys and those diagnosed after their first year.

GENETICS
When a parent has ADPKD, every conception carries a 50 percent chance of the child inheriting the mutated gene and having ADPKD. In ADPKD, only one mutated gene of the pair is required to pass on the disease. Approximately 10 percent of patients with PKD have no family history of the disease, as ADPKD may develop as a spontaneous (new) mutation.

SCREENING AND DIAGNOSIS
Diagnoses are usually made using radiology — ultrasound, CT scan or MRI.

Characters of children diagnosed in first year of life:
> One parent may have severe ADPKD.
> Most are diagnosed in-utero with large kidney cysts. Most develop high blood pressure (hypertension) in childhood, which should be monitored and treated by a doctor or pediatric nephrologist.
> Some patients may develop end-stage renal disease by their teenage years.

Characters of children diagnosed after first year of life:
> Usually, one of the parents is known to be affected with typical ADPKD.
> These cases most likely represent incidental findings due to widespread use of ultrasounds and improved resolution of the scans to detect smaller-sized cysts.
> Almost all children who are diagnosed after the first year of life will have normal kidney function throughout childhood.

SYMPTOMS OF ADPKD
> High blood pressure (hypertension)
> Frequent urinary tract infections
> Blood in urine (hematuria)
> Protein in urine (proteinuria)
> Depression and anxiety (due to stress and emotional impact)
> Mitral valve prolapse
> Hernia
> Back/flank pain
> Kidney stones
> Enlarged kidneys

HEALTH MANAGEMENT
> Have regular appointments with a pediatric nephrologist.
> Maintain a healthy diet: Avoid too much dietary salt and increase water intake.
> Ensure all medications are taken exactly as prescribed.
> Report any and all unusual circumstances to the doctor.

Friends Lena and Brittnee (left) were born with PKD.

Friends Greg and MaryKatherine (left) were born with ADPKD.
ABOUT ARPKD

Autosomal recessive polycystic kidney disease (ARPKD) is a rare form of PKD, affecting approximately 1 in 20,000 children.

ARPKD can cause death in the first month of life. Only 70 percent of ARPKD children survive the newborn period, and approximately one-third of them will need dialysis or transplantation by age 10. Previously thought to be a fatal condition, the prognosis for children with ARPKD has improved dramatically. Twenty years ago, only half of the children born with the disease survived to their 10th birthday, but now that percentage has increased to 85 percent.

GENETICS

Each conception carries a 25 percent chance of the child inheriting both mutated genes and of having ARPKD. In recessive genetic disorders such as ARPKD, affected children must inherit a copy of the disease gene from both parents. Since the parents each have only one copy of the disease gene, they do not have the disease and are referred to as “carriers.” With each conception, there is a 50 percent chance the child will be a carrier and a 25 percent chance both mutated genes are inherited.

SCREENING AND DIAGNOSIS

- MR Cholangiography (x-ray examination of the bile ducts)
- MRI
- CT scan
- Ultrasound

Typically in ARPKD, the kidneys appear to be larger than normal. In some babies, prenatal ultrasound can detect the enlarged kidneys as early as 18 weeks after conception. After birth, the kidneys may be large enough to examine by touch or through an ultrasound. Prenatal genetic testing is possible. These genetic tests can either involve a direct search of the gene for mutations or an indirect association using linkage analysis. For linkage analysis, DNA samples are required from the fetus, a previously affected child, and the parents.

SYMPTOMS OF ARPKD

**Prenatal**

- Diminished amniotic fluid levels during pregnancy.
- Enlarged kidneys on fetal ultrasound.
- Lung immaturity and functioning issues.

**Immediately after birth**

- Enlarged kidneys due to cysts.
- Breathing problems due to lack of space because of enlarged kidneys and decreased urine production. Ventilation is frequently required to sustain life.
- Enlarged spleen with low red blood cell, white blood cell and platelet counts.
- Congenital hepatic fibrosis
- Hypertension
- Growth problems

HEALTH MANAGEMENT

Families should work with specialists in several areas, including liver, kidney and lung specialists, to monitor the child’s health.

Madeline (left) and Cleo (far right) were both born with ARPKD.
The PKD Foundation is the only organization in the United States solely dedicated to finding treatments and a cure for polycystic kidney disease.

Since 1982, we have been dedicated to supporting and improving the lives of patients and families impacted by PKD. These efforts are accomplished through promoting research to find treatments and a cure, as well as providing education, advocacy, and awareness on a national level. We provide direct services to local communities nationwide and are the largest private funder of PKD research.

The Foundation supports PKD-relevant scientific and clinical meetings which bring together PKD scientists from around the world to discuss current findings and encourage research collaboration. Additionally, the Foundation is an online resource featuring numerous webinars and videos for families, along with an extensive website.

PKD PARENTS CHAPTER
The PKD Parents Chapter provides support for parents of children with both ARPKD and ADPKD.

This Chapter has served more than 500 families since 2000. Once a family with a child with PKD reaches out to the Foundation, a volunteer from the PKD Parents Chapter contacts them for support, compassion and guidance. Families can be connected with experts and other local families and are supported in many ways.

CONNECT
Social media

facebook.com/groups/PKDinChildren

Talk with other parents of children with PKD.

Voices of PKD
Find stories about people affected by PKD.

pkdcure.org/learn

The Foundation has invested over $2 million since 2006 in research, support, education and awareness for ARPKD, ADPKD in children and congenital hepatic fibrosis (CHF), a disease closely associated with ARPKD.