Bladder Exstrophy

What is bladder exstrophy?
Bladder exstrophy (x-tro-fee) is a bladder that is not formed right. The bladder and genitals are split in half, turned inside out and sit outside the body. There are many kinds of exstrophy.

How common is bladder exstrophy?
Bladder exstrophy is rare. It happens in about 1 in 10,000 to 1 in 50,000 babies. It is more likely in boys than girls. Cloacal exstrophy (a form of bladder exstrophy) happens in about 1 in 50,000 to 100,000 births.

What causes bladder exstrophy?
We don’t know what causes exstrophy. The problem occurs 4 to 8 weeks after a woman gets pregnant. This is when organs, muscles and tissues begin to form layers that separate, divide and fold. Exstrophy is not caused by something the mother did or did not do while she was pregnant. It does not run in families (it is not hereditary).

What other defects can babies with exstrophy have?
They may have some or all of these defects.

Genital
• Epispadias – In boys, the tube that carries urine from the bladder to the outside of the body (urethra) may be short and split. It opens on the upper
surface of the penis. The split may also involve the two halves of the testicles (scrotum).

- **Epispadias** – In girls, the tube that carries urine from the bladder to the outside of the body (urethra) is located between a split clitoris and labia minora.
- **Chordee** – In boys, the penis may curve up.

**Bladder**
- Your baby may have an incomplete bladder neck and sphincter. The bladder neck is the lower part, or door, of the bladder. The sphincter is a ring of muscles around the bladder neck. These parts control urine flow when they open and close.
- The bladder may hold less urine than normal.

**Vesicoureteral reflux (VUR)**
- Urine travels from the kidneys down tubes called ureters into the bladder. This flow is one way. Reflux is a condition in which urine can flow back up from the bladder to the kidneys.

**Kidneys**
- Some children may have double kidneys or kidneys in a different location.

**Bowel**
- Children with cloacal extrophy may have poorly developed large bowels and no rectum. They often need surgery to make an opening from the intestines to outside the stomach so poop (stool) can come out (colostomy). Regular extrophy does not usually involve the bowels.

**Pelvis**
- The front parts of the pubic bone are wide apart. This is called diastasis.

**Spine**
Some children with cloacal extrophy have a fatty growth (lipoma) on their spinal cord. This may cause problems with their legs, bladder and rectum.
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How will we take care of our baby until they have surgery?

Your baby’s open bladder does not cause them any pain. But it is important to protect it. Until your baby has their surgery to close the bladder, you will put a dressing over their bladder to protect it. You will wash and care for your baby as normal and avoid rubbing their bladder.

Babies with exstrophy are at risk for developing an allergy to latex later in life. Because of this, you should avoid using latex at home when you can, especially when it touches your baby’s mouth, nose, urethra, etc. (mucous membranes). We will also avoid using latex products on your baby when they are in the hospital.

What surgeries will my baby need?

Between the 4 to 6 months of life, your baby may have 1 or more surgeries to:

• Close their bladder and place it in the pelvis. This may include reconnecting the ureters.
• Bring the front parts of the pubic bones back together (called “Reapproximate”) to fix the diastasis.
• Create a bladder neck
• Fix the epispadias
• Close the abdomen
• Create an ostomy (for children with cloacal exstrophy)

Your child’s urologic surgeon and team will help you know what to expect before, during and after each surgery.

What can we expect before surgery?

Your baby will need regular visits at Seattle Children’s to prepare for surgery.

Two months before surgery:

• You will have a clinic visit with your child’s urologic surgeon
• Your baby will need a blood test for anemia by their primary care provider

Two weeks before surgery:

You will need to:

• Have a clinic visit with your child’s urologic surgeon
• Have a clinic visit with your child’s orthopedic surgeon. They will help you order a special car seat for your baby to use after their surgery.
• Visit the Pre-Anesthesia and Surgical Service clinic (PASS Clinic), where the anesthesia team will check that your baby is ready for surgery.
• Your baby may need some labs checked after this clinic
What can we expect after surgery?

Right after surgery, your child will have:

A spica cast

This is a special cast that will be put on by your baby’s orthopedic surgeons. The cast goes from the ankles to the mid-chest, with a hole for the diaper area. This cast will stay on for about 6 weeks. The cast will impact how your baby moves, how you hold them and care for them. Your child’s nurses will teach you how to take care of the cast and give you special tips for diapering. You will get a special type of car seat to use while they have the spica cast.

Catheters

Catheters are tubes to drain urine from your baby. Your baby will have a tube called a supra pubic tube in for about 6 weeks at home. We call this the “SP” tube. Your baby may also have 2 ureteral catheters or stents in place that come out through the urethra. These will fall out on their own in about 10-14 days after surgery.

Intravenous line (IV)

Your baby will have a tube inserted into their vein to give them fluids and antibiotics for several days after surgery.

Cardiac respiratory monitor

You baby may have a cardiac respiratory monitor to check their heart rate, breathing and oxygen levels for several days.
When will we be able to go home from the hospital?

We will teach you all you need to know to care for your baby at home. Your child will stay in the hospital until you feel comfortable taking care of your baby. They also need to meet specific goals for eating, drinking, going to the bathroom and pain control. Before they can leave, they will be fitted for their car seat and your follow up visits will be scheduled.

What can we expect after we leave the hospital?

Your child will have weekly appointments in the Urology clinic for the first 6 weeks. While you are home, we can answer your questions at any time.

Six weeks after surgery:

Your child will return to the hospital for:

- Clinic visit with their orthopedic surgeon to remove their spica cast
- Radiology studies including:
  - X-ray to look at how well their bones are healing
  - Cystogram or dye study to look at how well their bladder is healing and whether your child has vesicoureteral reflux
  - Ultrasound to look at their kidneys
- Clinic visit with their Urologic surgeon to remove the SP tube.

You will also need to see your child’s primary care provider for their on-going well-child checkups.

Will the exstrophy impact my child as they grow up?

Some children who had surgery for exstrophy gain complete control over urine flow, and they have no other issues later that need treatment.

Some children will need other procedures. These may include:

- Children with urine that flows back up from the bladder to the kidneys (vesicoureteral reflux) sometimes need antibiotics. This helps prevent urinary tract infections (UTIs). As they get older, they may need a procedure to reduce reflux.
- If leaking of urine (incontinence) is a problem, your child may need surgery to help them stay dry.
- Some children need surgery to enlarge their bladder (bladder augmentation) or to place a tube (Mitrofanoff channel) so they can empty their bladder by putting a catheter through an opening in their belly.
- Boys may need surgery on their penis if the opening is on the underside, not at the tip (hypospadias).

Surgeries can often be combined.

Your child may get more UTIs than other children. Your child’s primary care provider can help monitor and treat these.
When your child reaches school-age, they should see their pediatric urologist at least once a year. Your child will get a kidney ultra sound to make sure that their kidneys are growing well and to look for any signs of kidney problems.

We partner with adult urologists at University of Washington and Harborview Medical Center to ensure your child gets proper care when they become an adult. Life-long follow-up care is important for good urinary health.

Children and adults with exstrophy can lead normal, healthy, active lives. Our goal is to help your child have healthy kidneys and good bladder health. As an adult, your child should have normal sexual sensation, function and potential for fertility.

Each person with exstrophy is different, and their care and treatment will vary based on their needs. You and your child’s urologist will develop a plan of care just for your child.

**Where can I get more information?**

We can provide you with more information on:

- Bladder exstrophy resources
- Other families of children with exstrophy via the Northwest Exstrophy Support Group
- A national exstrophy organization called Association for the Bladder Exstrophy Community (ABC)
- Latex allergy
- Vesicoureteral reflux
- Spica casts
- Urinary tract infections