EASY READING

If Your Child Has a Wilms Tumor

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What are Wilms tumors?

Cancer starts when cells in the body begin to grow out of control. Cells in nearly any part of the body can become cancer.

Wilms tumors are a type of kidney cancer that is most often found in young children. But rarely these tumors can occur in older children, or even teens or adults.

To understand Wilms tumors, it helps to know a little about the kidneys. The kidneys are 2 bean-shaped organs that are attached to the back wall of the abdomen (see picture). One kidney is just to the left and the other just to the right of the backbone.
The kidneys do a number of things:

- They filter the blood to remove excess water, salt, and waste products, which leave the body as urine.
- They help control blood pressure.
- They help make sure the body has enough red blood cells.

Our kidneys are important, but we need less than one complete kidney to do all of its basic functions.

Most often, Wilms tumors occur in only one kidney. But a small number of children will have tumors in both kidneys.

**Different types of Wilms tumor**

Wilms tumors are grouped into 2 major types\(^1\) based on how they look under a microscope (called their **histology**):

- **Favorable:** The cancer cells in these tumors don’t look quite normal, but they don’t look very abnormal, either. Most Wilms tumors have a favorable histology. The chance of curing children with these tumors is very good.
- **Anaplastic:** In these tumors, the look of the cancer cells varies widely, and parts of the cell tend to be very large and distorted. This is called *anaplasia*. Anaplasia can
be either **focal** (limited to just certain parts of the tumor) or **diffuse** (spread widely through the tumor).

**Questions to ask the doctor**

- How sure are you that my child has a Wilms tumor?
- Is there a chance it’s not a Wilms tumor?
- Would you please write down the kind of tumor you think my child has?
- What will happen next?

**How does the doctor know my child has a Wilms tumor?**

Wilms tumors often grow quite large before causing any symptoms. Often the first sign of a Wilms tumor is **swelling or hardness in the belly**, which might be on one side or both. It’s usually not painful.

Other symptoms might include:

- Fever
- Nausea
- Loss of appetite
- Shortness of breath
- Constipation
- Blood in the urine

Wilms tumors can also sometimes cause **high blood pressure**. This might not cause symptoms on its own, but in rarely the blood pressure might get high enough to cause problems such as headaches, bleeding inside the eye, or even a change in consciousness.

If your child has symptoms that could be from a Wilms tumor, the doctor will want to get a complete medical history to find out more about the symptoms, and will do a physical exam. Tests might be needed as well.

**Tests that may be done**

Here are some of the tests your child may need:
Ultrasound: This is often the first test done if the doctor thinks your child has a tumor in the belly. This test is easy to have, does not use radiation, and it gives the doctor a good view of the kidneys and the other organs in the belly.

CT or CAT scan: This test uses x-rays to make detailed pictures of the inside of the body. This is one of the most useful tests to look for a tumor inside the kidney. It can also show if the cancer has grown into nearby veins or has spread to other organs, such as the lungs.

MRI: This test uses radio waves and strong magnets to make detailed pictures of the inside of the body. This test might be done if the doctor needs to see very detailed pictures of the kidney or nearby areas.

Chest x-ray: This test might be done to look for spread of Wilms tumor to the lungs. A chest x-ray likely won’t be needed if a CT scan of the chest is done.

Lab tests: Blood and urine tests might be done if your child’s doctor suspects a kidney problem. These tests might also be done after a Wilms tumor has been found.

Biopsy: Most of the time, the tests above are enough to tell doctors if a child most likely has a Wilms tumor, and therefore if surgery should be done. But to be sure, a biopsy is needed. This is when a piece of the tumor is removed and checked under a microscope.

The biopsy is most often done as part of the surgery to treat the tumor. But if the doctors aren’t as sure about it being a Wilms tumor, or if they aren’t sure all of the tumor can be removed, a sample of the tumor might be taken during a biopsy done on its own, before surgery.

Questions to ask the doctor

- What tests will my child need?
- Who will do these tests?
- Where will they be done?
- Who can explain them to us?
- How and when will we get the results?
- Who will explain the results to us?
- What do we need to do next?
How serious is my child’s tumor?

If your child has a Wilms tumor, the doctor will want to find out some key pieces of information\(^4\) to help decide how to treat it. The most important of these are:

- The **stage** (extent) of the cancer, which is based on if it is just in one kidney, if it has grown outside the kidney, and if all of it can be removed with surgery
- The **histology** of the tumor (favorable or anaplastic)

Other things that doctors use to decide on treatment and a child’s chances of getting better include:

- The child’s **age**
- If the tumor cells have certain **gene changes**
- The **size** of the tumor

Questions to ask the doctor

- Where exactly is the tumor?
- How big is the tumor?
- Can all of the tumor be removed?
- Has the tumor spread anywhere else?
- What is the stage of the tumor?
- What is the histology of the tumor? What does this mean?
- How do these things affect our treatment options?
- What will happen next?

What kind of treatment will my child need?

The main treatments for Wilms tumors are:

- Surgery
- Chemotherapy (chemo)
- Radiation treatment
Surgery

In the United States, surgery\(^5\) is the first treatment for most children with Wilms tumors. The main goal is to remove the entire Wilms tumor in one piece, if possible. If this can’t be done safely, then other treatments like chemo might be done first to shrink the tumor and make surgery easier.

The main operations to treat Wilms tumors are:

**Radical nephrectomy**: This surgery removes the entire kidney and some nearby structures. This is the most common surgery for a Wilms tumor that’s only in one kidney, as it gives the best chance of making sure all of the tumor is removed.

**Partial nephrectomy (nephron-sparing surgery)**: This surgery removes only part of the kidney(s). It is used most often in children who have Wilms tumors in both kidneys, to try to save some normal kidney tissue.

Ask your doctor what type of surgery your child will need and what to expect.

If chemo is going to be part of treatment, surgery might be done to insert a small tube (called a central venous catheter (CVC)\(^6\) or port) into a large blood vessel – usually under the collar bone. This might be done during the surgery to remove the tumor, or as a separate operation if chemo is going to be given before the surgery.

**Side effects of surgery**

Any type of surgery can have risks and side effects, such as bleeding or infections. Ask your child’s doctor or nurse what to expect. If your child has any problems, let them know. Doctors and nurses who treat children with Wilms tumors should be able to help you with any problems that come up.

Most children do well if only one kidney is removed. But children who have both kidneys removed, or even parts of both kidneys removed, might need regular dialysis treatments to filter their blood. They also might need a kidney transplant at some point.

Chemo

*Chemotherapy*\(^7\) (chemo) is the use of drugs to fight cancer. These drugs are put into the blood and spread through the body.

Most children with Wilms tumors will get chemo at some point during their treatment.
(Some children with very low risk tumors might not need it.)

Most often chemo is given after surgery, but sometimes it’s given before surgery to shrink the tumor and make the surgery easier.

Children with Wilms tumors will get 2 or more chemo drugs as part of their treatment. Chemo is given in cycles or rounds. Each round of treatment is followed by a break. Treatment often lasts for many months.

**Side effects of chemo**

Chemo can make your child feel very tired, sick to their stomach, or cause their hair to fall out. It might also cause other problems. But these tend to go away after treatment ends.

There are ways to treat most chemo side effects. If your child has side effects, talk to the cancer care team so they can help.

**Radiation therapy**

Radiation uses high-energy rays (like x-rays) to kill cancer cells. It may be used:

- After surgery to kill any tumor cells left behind
- Before surgery to try to shrink the tumor to make it easier to remove
- As the part of the main treatment if surgery can’t be done

For Wilms tumors, radiation is aimed at the tumor from a machine outside the body. Most often it’s given 5 days a week for at least a few weeks. Each treatment is a lot like getting an x-ray, and takes about 15 to 30 minutes. The radiation is stronger, but your child will not feel it.

**Side effects of radiation therapy**

Radiation might cause some children to feel tired, or feel sick and throw up. A child might also have loose stools (diarrhea). Radiation can also affect the skin where it enters the body, which can cause changes that are like a sunburn.

Most side effects get better after treatment ends. But some might last longer, or might not show up until years later. Talk to your child’s care team about what to expect during and after treatment. There may be ways to ease some side effects.
Clinical trials

Clinical trials are research studies that test new drugs or other treatments in people. They compare standard treatments with others that may be better.

Clinical trials are one way to get the newest treatments. They are the best way for doctors to find better ways to treat cancer. But they might not be right for everyone. If your child’s doctor talks to you about a clinical trial, it’s up to you whether to take part.

If you would like to learn more about clinical trials that might be right for your child, start by asking the doctor if your child’s clinic or hospital conducts clinical trials. See Clinical Trials to learn more.

What about other treatments I hear about?

When your child has a Wilms tumor, you might hear about other ways to treat it or treat symptoms from it. These may not always be standard medical treatments. These treatments may be vitamins, herbs, diets, and other things. You may wonder about these treatments.

Some of these might help, but many have not been tested. Some have been shown not to help. A few have even been found to be harmful. Talk to your doctor about anything you’re thinking about using, whether it’s a vitamin, a diet, or anything else.

Questions to ask the doctor

• Do we need any other tests before we can decide on treatment?
• What treatment do you think is best for my child?
• What’s the goal of this treatment? How likely is it to cure the tumor?
• Will treatment include surgery? If so, who will do the surgery?
• What will the surgery be like?
• Will my child need other types of treatment, too?
• What will these treatments be like?
• What’s the goal of these treatments?
• What side effects could my child have from these treatments?
• What can we do about side effects that my child might have?
• Is there a clinical trial that might be right for my child?
• What about vitamins or diets that friends tell me about? How will we know if they are safe?
• How soon do we need to start treatment?
What should we do to be ready for treatment?
- Is there anything we can do to help the treatment work better?
- What’s the next step?

What will happen after treatment?

You’ll be glad when treatment is over\(^1\). But it’s hard not to worry about the tumor coming back. Even if it never comes back, you might still worry about it. For years after treatment ends, your child will still need to see the doctor. At first, these visits may be every few months. Then, the longer your child is cancer-free, the less often the visits are needed.

Be sure your child goes to all of these follow-up visits. Your doctors will ask about symptoms, do physical exams, and may do tests to see if the tumor has come back. They may also test to see if the cancer or its treatment has caused any long-term problems. If needed, they will help you and your child learn to deal with the changes.

Hyperlinks

Words to know

Biopsy (BY-op-see): Taking out a piece of an abnormal area to see if there are tumor cells in it.

Central venous catheter (CVC): A small tube that is put into a large blood vessel (usually under the collar bone), with one end staying outside of the body or just under the skin. It can be left in place for months and can be used to give chemo or take blood samples. Also called a venous access device (VAD) or a port.

Chemotherapy (KEY-mo-THAIR-uh-pee): The use of drugs to kill cancer cells. Also called chemo.

Metastasis (muh-TAS-tuh-sis): The spread of cancer cells from where they started to other places in the body.

Nephrectomy (nef-REK-tuh-mee): Surgery to remove a kidney (radical nephrectomy) or part of a kidney (partial nephrectomy).

Radiation (ray-dee-AY-shun) therapy: The use of high-energy rays (like x-rays) to kill cancer cells.

How can I learn more?

We have a lot more information for you. You can find it online at www.cancer.org (http://www.cancer.org)\(^\text{12}\). Or, you can call our toll-free number at 1-800-227-2345 to talk to one of our cancer information specialists.

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Written by

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