Epilepsy and Seizure Disorders

A Wyoming Resource Guide for Parents
Notes and Acknowledgements

Notes to the reader:

► The information contained in this guide is not intended as, and is not a substitute for, professional medical advice. A child’s medical team should be consulted about all decisions regarding clinical care and treatment. No pharmaceutical funding was used in the development of this guide.

► Throughout the text, we use the word “child”. Please understand that we use this term to refer to a person of any age between birth and 21 years.

Photograph acknowledgements:

Many of the photographs in the Guide were taken by University of Wyoming, Public Relations, WIND staff, Project Access partners, parents and appear courtesy of the Wyoming Institute for Disabilities.
Table of Contents

Notes and Acknowledgements (inside front cover)

Introduction 2

Section 1: Understanding Epilepsy/Seizure Disorder 3
- What is epilepsy/seizure disorder? 3
- What are the different kinds of epilepsy/seizure disorder? 3
- How is epilepsy/seizure disorder diagnosed? 5
- What causes epilepsy/seizure disorder? 6
- What might trigger a seizure? 6
- What are the treatment options? 6

Section 2: Health Care for My Child—Access to Care/Access to Services 8
- What do I need to know about taking care of my child? 8
- What organizations can help me find the help we need? 10
- How can I pay for or access the care my child needs? 10
- Who are the members of my child’s health care team? 15
- How can I assure the best health care for my child? 16

Section 3: Advocacy & Support 20
- Where can I find support from other parents like me? 20
- What do I need to know about my child attending school? 23
- How can I educate others about my child’s epilepsy/seizure disorder? 26

References 27

Appendix: Forms 28
Dear Parent or Caregiver,

Children are one of life’s greatest gifts. As parents, we want to do all we can to nurture and take care of our children. As a parent of a child with epilepsy, also referred to as seizure disorders, I know we face the additional challenge of learning as much as we can about our child’s special needs. This guide was written with you in mind to give you some very basic information, including an introduction to epilepsy and seizure disorders. It is not intended to provide medical advice. Any questions related to the medical treatment of your child should always be addressed to your child’s pediatrician, family physician, or neurologist.

While this guide may not provide all of the information you are looking for, it should help you begin your search. The Guide includes information about the additional resources that we hope will be useful as you begin this journey—a journey that is likely to evolve as you become more knowledgeable about your child’s condition. Take information that applies and is valuable to you and your child.

I would like to extend a warm thanks to the following individuals who made this guide possible in Wyoming. The wonderful WIND staff, family, friends, and parents and children with epilepsy who submitted photos to produce our guide: Michelle Ecker, Jessica Rasmussen, and Lori Schaeffer. I would like to recognize the participation of Allison Sedlacek of the Colorado Epilepsy Foundation and Richard Leslie of the Wyoming Epilepsy Association, affiliates in the development and review of the Guide. The Guide would not be possible without the dedicated work of Cary Kreutzer, Project Director and her staff of University of Southern California, and University Center for Excellence in Developmental Disabilities at the Childrens Hospital Los Angeles. A special thanks to Dr. David Wheeler, the Director of the Wyoming Epilepsy Center, Wyoming Neurologic Associates, for his participation in the Wyoming Access Team for his valuable advice and conference presentations. Thanks also goes for the support of families who have a child with epilepsy. Finally, thanks to Athena Lickel and Lilly Zuniga, WIND Project Coordinators, Janet Perkins Corbett, Whitney Buckley-Dutton, and Sara DiRienzo for editing and Elizabeth Ono Rahel for graphic design.

Please use this guide as it helps in your life and the care of your child.

Sincerely,

Sandra Root-Elledge
Associate Director
Understanding Epilepsy/Seizure Disorder

What is epilepsy/seizure disorder?*

The brain contains billions of nerve cells called neurons that communicate electrically and signal to each other. A seizure occurs when there is a sudden and brief excess surge of electrical activity in the brain between nerve cells. This can cause abnormal movements, change in behavior, or loss of consciousness.

Seizures are not a mental health disorder. Instead, epilepsy is a neurological condition that is still not completely understood.

Having a single seizure does not mean that a child has epilepsy. A child has epilepsy when he or she has two or more seizures without a clear cause such as fever, head injury, drug use, or alcohol use. About three million Americans have epilepsy. Of the 200,000 new cases that develop each year, up to 50% are children and adolescents. About 300,000 children under the age of 14 in the United States have this condition. It develops in children of all ages and can affect them in different ways.

What are the different kinds of epilepsy/seizure disorders?

There are many kinds of epilepsy and seizures. They each cause different behaviors and they each need different treatments. Identifying the type (or types) will help your child’s doctor suggest treatment options.

GENERALIZED SEIZURES

This type of seizure involves the whole brain and is the most common type of epilepsy. In a generalized seizure, the abnormal electrical activity affects the entire brain. These seizures produce muscle twitches, convulsions, and loss of consciousness. People with this type of epilepsy do not remember having a seizure.

PARTIAL SEIZURES

This type of seizure involves only part of the brain. Sometimes a partial seizure can spread to involve the whole brain, which is known as a partial seizure that secondarily generalizes.

* Some people use the term “seizure disorder” instead of “epilepsy” to describe this condition. In fact, both terms mean the same thing—an underlying tendency to experience seizures.
The following chart contains more information about the different types of generalized and partial seizures. Also see page 19 for information about first aid and safety specific to seizure type.

<table>
<thead>
<tr>
<th>SEIZURE TYPE</th>
<th>DESCRIPTION</th>
</tr>
</thead>
</table>
| **Tonic-clonic or Grand Mal**
(loss of consciousness) | This is the most common and recognized generalized seizure. The person becomes stiff and falls to the ground. Teeth clench and the arms, and usually the legs, begin to jerk rapidly and rhythmically. The seizure usually lasts no more than a few minutes, after which the jerking slows and stops. |
| **Absence or Petit mal**
(loss of consciousness) | During an absence seizure it might seem like the person is daydreaming. However, in an absence seizure the person cannot be alerted or woken up. They are unconscious for a moment and totally unaware of what is happening around them. These seizures usually last a few seconds. |
| **Myoclonic or Jerks**
(no loss of consciousness) | Myoclonic means a jerking or twitching of a muscle. During this seizure brief shock-like jerks of a muscle or group of muscles occur. These usually involve the neck, shoulders, and upper arms. Myoclonic jerks occur most frequently in the morning and often occur in clusters. Although the seizures are brief, they can be extremely frustrating, resulting in spilt drinks or similar incidents. |
| **Tonic and Atonic**
(loss of consciousness) | Tonic seizures result in all muscles contracting. The body stiffens and the person will fall over if unsupported. Atonic seizures, in a way, are the opposite of tonic seizures. Instead of the body going stiff, all muscle tone is lost and the person simply drops to the ground. Although the person falls heavily, they are usually able to get up again right away. When the body goes limp, it inevitably falls forward causing potential head injuries. |
| **Status Epilepticus**
(loss of consciousness) | This event is characterized by frequent, long-lasting seizures without regaining consciousness between the start and end of seizures. |
| **Simple**
(no loss of consciousness) | In these types of seizures, even though a person’s consciousness is not impaired, it does not mean that the person experiencing this type of seizure is able to stop or control the seizure. Simple partial seizures can be different depending on where in the brain the epileptic activity is occurring. Examples of symptoms are the movement of a limb, tingling, experiencing a smell or taste, and going pale or sweating. |
| **Complex**
(impaired consciousness) | Because a person’s consciousness is impaired in this type of seizure, the person will not remember the seizure or their memory of it will be distorted. Others may believe the person is fully aware of what they are doing, but they are not. Usually the person loses awareness and stares blankly. Most people move their mouth, pick at the air or their clothing, or repeat other purposeless actions. These movements are called “automatisms”. They usually last between 30 seconds and 2 minutes. |
How is epilepsy/seizure disorder diagnosed?

Your doctor will ask a lot of questions when trying to diagnose epilepsy and determine the type(s) of seizures your child is having. The diagnosis is based on several exams and tests in addition to an interview about your child’s condition.

The facts you provide the doctor are very important in diagnosing your child’s epilepsy and deciding on treatment. The following chart outlines some of the tools a doctor uses for diagnosis.

### TOOLS FOR DIAGNOSIS

<table>
<thead>
<tr>
<th>Information about seizure(s)</th>
<th>Questions that may be asked:</th>
</tr>
</thead>
<tbody>
<tr>
<td>The doctor will ask for a complete description of what happened.</td>
<td>- What circumstances surrounded the seizure?</td>
</tr>
<tr>
<td></td>
<td>- What seemed to bring on the seizure?</td>
</tr>
<tr>
<td></td>
<td>- What happened before the seizure?</td>
</tr>
<tr>
<td></td>
<td>- How did your child feel before, during, and after the seizure?</td>
</tr>
<tr>
<td></td>
<td>- Description of seizure behavior.</td>
</tr>
<tr>
<td></td>
<td>- What happened after the seizure?</td>
</tr>
<tr>
<td></td>
<td>- Where was the child when the seizure occurred?</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Medical history</th>
<th>You may be asked about:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>- family history of seizures</td>
</tr>
<tr>
<td></td>
<td>- medical conditions or medications</td>
</tr>
<tr>
<td></td>
<td>- general medical history of your child</td>
</tr>
</tbody>
</table>

An examination of muscle strength, reflexes, eyesight, hearing, and ability to detect various sensations are tested to better understand the cause of the seizures.

<table>
<thead>
<tr>
<th>EEG (Electroencephalogram)</th>
<th>An EEG measures the electrical impulses in the brain. During an EEG, electrodes (small metal disks) are attached to specific locations on the head and monitored. Abnormal electrical spikes indicate seizure activity.</th>
</tr>
</thead>
<tbody>
<tr>
<td>CT or CAT scan (Computerized Axial Tomography)</td>
<td>A CT scan is an X-ray of the brain that creates three-dimensional images of the brain and shows possible abnormalities.</td>
</tr>
<tr>
<td>MRI (Magnetic Resonance Imaging)</td>
<td>An MRI is another imaging method using magnets instead of X-rays. MRI tests provide detailed images of the brain. CT or MRI scans may be used to search for any growths, scars, or other physical conditions in the brain that may be causing the seizures.</td>
</tr>
<tr>
<td>Blood tests</td>
<td>Tests to measure white blood cell count, blood sugar, sodium, calcium and electrolyte levels, and liver and kidney function. Blood tests also help rule out other illnesses.</td>
</tr>
<tr>
<td>Other tests</td>
<td>Other tests may be ordered as needed.</td>
</tr>
</tbody>
</table>
What causes epilepsy/seizure disorder?

Many parents wonder if they have somehow caused their child’s epilepsy. They search for a way to understand why this is happening to their child. But, it is unlikely a parent or anyone did anything to cause the epilepsy.

Finding the cause of epilepsy is difficult. For seven out of ten children with epilepsy, there is no known cause. These children are said to have idiopathic epilepsy. “Idiopathic” means “of unknown cause.”

However, there are many known causes. Understanding and identifying the causes help to diagnose the type(s) of epilepsy. Possible causes include the following:

- problems with brain development before birth
- lack of oxygen during or following birth
- a serious head injury that leaves a scar in the brain
- unusual structures in the brain
- tumors
- a prolonged seizure with fever
- the after-effects of severe brain infections, such as meningitis or encephalitis
- genetic factors

What might trigger a seizure?

Some people report very specific triggers or immediate causes that can bring about a seizure. Children with epilepsy are more likely to have a seizure (have a lower seizure threshold) when they have a cold, the flu, or other common illnesses. Some common seizure triggers include:

- forgetting to take your medicine
- not enough sleep
- food allergies
- stress
- flashing lights
  (e.g., from video games, strobe lights)
- alcohol
- illicit drugs
- dehydration
- poor diet
- inactivity

What are the treatment options?

Childhood epilepsy is usually treated with medications that prevent seizures. If the medications do not work or if the child has too many side effects, there are other treatment options. These include surgery, the ketogenic diet, or vagus nerve stimulation (VNS). Other treatment therapies are being tested, but have not yet been FDA approved.

MEDICATION

Children often take the same antiepileptic medications as adults. Medication may be in the form of tablets, sprinkles, capsules or syrup.
MEDICATION TIPS:

- Make sure you understand the dosages of medicine prescribed by your doctor, such as the number of pills or teaspoons required for each dose, as well as the number of doses and times (e.g., morning, noon, night) to administer each dose. Also, make sure that you understand the best method for administering each medication (e.g., by dropper, mixed with food or liquid, or crushed).

- Ask your doctor or pharmacist about possible side effects associated with each medication your child is taking and what you should do if they occur.

- Keep follow-up appointments. Some medications require periodic blood tests that are important to your child's health. Ask the doctor at the end of each visit when he or she wants to see you both again, and schedule the next appointment.

- Don't change the dose or stop giving seizure medication on your own without first talking with your child's doctor.

- Ask for refills from your pharmacy several days before the medication is due to run out.

- Ask your doctor or pharmacist about over-the-counter medications as they may interfere with the epilepsy medication your child is taking. Also, check with the pharmacist when filling other prescriptions for your child.

- Ask your doctor how to handle fever associated with childhood illness. Ask, too, about the best way to give medication when a child has a stomach virus and cannot keep medication down.

- Find out what you should do if your child misses a dose. Ask whether the dosing schedule is flexible and what to do if your child is scheduled for a dose when he or she is sleeping.

- Do not allow yourself to run out of medicine. It is important that anticonvulsants be given steadily. Ask for a new prescription at each visit and do not order refills of "old" prescriptions.

These medications are designed to prevent seizures. Some are successful with a few seizure types; others have a broader range of action. Whenever possible, doctors try to control seizures with one medication. Some children, however, may need to take more than one.

Children may respond so well to medication that no further seizures occur as long as the medication is taken regularly as directed by the doctor. Not having seizures does not mean that the medication is no longer needed. Always ask the doctor before stopping a seizure medication. Doing so without medical supervision may result in a seizure or another type of reaction.

Recently, there have been concerns regarding differences between brand and generic medications, and different manufacturers of the same generic medicine. For more information about these concerns, medications, and medication side effects, contact your doctor or the Epilepsy Foundation at (800) 332-1000.

KETOCENIC DIET

If medications do not control a child's seizures, a doctor may recommend a special high fat, low carbohydrate diet called a ketogenic diet.

Carbohydrates are strictly limited and parents have to be very careful that the child does not eat foods that are not on the diet or have not been pre-measured and pre-weighed.

The diet requires a team effort—the family, the physician, the dietitian, the nurse, and, if the child is old enough, the child himself or herself all working together to make sure the diet is followed and any side effects are monitored.

WARNING

The ketogenic diet is not a do-it-yourself diet. It must be monitored by a medical team.

EPILEPSY AND SEIZURE DISORDER: A RESOURCE GUIDE FOR PARENTS 7
VAGUS NERVE STIMULATION

Vagus nerve stimulation (VNS) is a type of surgical intervention that may be tried when seizures cannot be controlled by other methods. The treatment works by sending regular small bursts of electrical energy to the vagus nerve, a large nerve in the neck that leads directly into the brain.

The energy comes from a small disk-like generator, about the size of a stop watch, implanted by a surgeon under the skin on the chest. The generator is connected to wires under the skin that are wrapped around the vagus nerve.

SPECIALIZED CARE

Specialized care for children with refractory (difficult to treat) epilepsy is available at special centers around the country. Many offer in-depth evaluations, surgery, the ketogenic diet, and VNS therapy. For information about sources of specialized care near you, call the National Association of Epilepsy Centers or any other organization listed on page 10.

Health Care for My Child

Access to Care/Access to Services

What do I need to know about taking care of my child?

The most important way to help your child and his or her physicians is to be an active member of your child's health team. This means preparing for visits, keeping records, learning to ask questions and advocating for your child. Networking with other parents of children with epilepsy may become a source of information, strength, and inspiration as you learn to live with your child's epilepsy. And most of all, do not forget to take care of yourself—without you, your child's number one advocate is gone.

MAKE OBSERVATIONS

Seeing your child having what appears to be a seizure can be very frightening. However, remaining as calm as possible is important so you can describe exactly what happened to the doctor. The likeliness your doctor will ever see your child have a seizure is small. Your observations and knowledge of your child are vital in helping the doctor reach a diagnosis. Writing down these and all observations about the seizure or subsequent behavior is a good idea so you can tell your doctor exactly what happened.

Two examples of forms to help you track this information are included in the Appendix section of this guide (Seizure Log [F1] and Seizure Description Sheet [F2]). If possible, try to record a video of your child during a seizure so that your child's doctor can observe exactly what happens and what steps you take in response. Make sure that you or someone else is still tending to your child and the seizure during the taping, keeping your child's health and safety a priority at all times.

EDUCATE YOURSELF—ASK QUESTIONS

Feeling overwhelmed and confused is perfectly normal when first learning about epilepsy. But if you ask lots of questions and gather as much information as you can, you are choosing the most effective way to help your child. Learning as much as you can about epilepsy will build
your self-confidence and improve your ability to respond to your child's needs. In this guide you will encounter many Internet resources, as well as physical addresses and phone contact information. If you do not have a computer with Internet access at home or work, you can go to your local library and use one of their public computers to look up the links for free. If you live in a rural area with no library, please contact the Epilepsy Foundation (1 (800) 332-1000) for assistance.

You must be your child’s advocate. Ask for what you want from doctors. Put everything in writing. You may find friends and advocates at the school district, the doctor’s office, or at an Early Intervention program who can help you. If you are not satisfied with the services you receive from your health plan, medical group, or provider, talk to them about your concerns. If you are still not satisfied, document your concerns in writing. Sample letters can be obtained from your family resource center or online (www.hrh.org—click on Consumer Action Guides).

A wealth of information is available about epilepsy in children. Organizations exist on the national, state, and local level that can direct you about whom to call and where to look for the information you need. There are books, brochures, Web sites, training classes, and more to help you find your way and learn about your child and his or her condition.

GET ORGANIZED

One of the most important things you can do for yourself and your child is to organize information you are receiving. Having complete medical records available helps all the doctors who are treating your child.

ATTEND SUPPORT GROUPS AND NETWORKS WITH OTHER PARENTS

Find support groups for yourself, your spouse, and your family. Take care of yourself to avoid being overwhelmed or getting sick.

Your children may sense your stress and it can scare them. You need a place to talk about your fears and concerns. Ask your nearest epilepsy organization about where groups in your area are meeting. Your doctor, nurse, Early Intervention coordinator, or your local hospital may also be able to tell you what associations exist and where support groups may be available. If you go online to a chat room, be sure that there is a medical moderator in the chat room.

TAKE CARE OF YOURSELF

As parents, our needs always seem to come last. But we cannot continue as effective caregivers if we do not take time to care for ourselves. Giving all that we can to our loved ones requires taking care of ourselves first. Here are some basic tips:

- Schedule time for yourself each week—take a walk, read a book, take a yoga class, indulge in a makeover, or even go shopping. Take the time to do something that makes you happy.
- Stay in touch with yourself. Keep a journal in which you can express feelings and thoughts.
- Stay in touch with friends.
- Try to eat healthily and stay physically active to prevent chronic disease.
What organizations can help me find the help we need?

The following organizations can help direct you to information and services in your area.

**Wyoming Epilepsy Association**  
119 W. 17th St., Cheyenne, WY 82001  
(307) 634-5329/(307) 286-8637  
E-mail: admin@wyoeningepilepsy.org  
www.wyomingepilepsy.org

**Epilepsy Foundation of Colorado**  
234 Columbine St. #333, Denver, CO 80206  
(303) 377-9774/1 (888) 378-9779  
Satellite Office: Greeley (970) 352-4646  
E-mail: allison@epilepsycolorado.org  
www.epilepsycolorado.org

**Epilepsy Foundation**  
8301 Professional Place East, Landover MD 20785-2238  
1 (800) 332-1000  
www.epilepsyfoundation.org

**National Association of Epilepsy Centers**  
5775 Wayzata Blvd., Suite 200, Minneapolis, MN 55416  
1 (888) 525-6232  
E-mail: info@naec-epilepsy.org  
www.naec-epilepsy.org/

**Epilepsy Foundation Parent’s Web Site**  
www.epilepsyandmychild.org/

**Family Voices of Wyoming**  
Rick Skagen, (307) 742-4822  
www.familyvoices.org

**UPLIFT**  
4007 Greenway, Suite 201, Cheyenne, WY 82001  
(307) 778-8686/1 (888) UPLIFT3  
www.upliftwy.org/index.htm

**Parent Information Center (PIC)**  
500 W. Lott St., Suite A, Buffalo, WY 82834  
1 (800) 684-5314  
www.wpnc.org

**Parent Education Network (PEN)**  
Wyoming State PIRC  
500 W. Lott St, Suite A, Buffalo, WY 82834  
(307) 684-7441/Toll free 1 (877) 900-9736 (WY)  
www.wpen.net

**Wyoming Epilepsy Center**  
Wyoming Neurologic Associates  
419 S. Washington, Suite 201, Casper, WY 82601  
(307) 265-4343

► Look your best. Although it takes energy, it also boosts your spirit.  
► Accept help—extra hands at doctors’ appointments, grocery shopping, child care, car pool, etc.

Also, parents may benefit from mental health counseling services and are encouraged to bring up their mental health concerns (i.e., depression, anxiety) to their primary health care provider.

**Tips for Caregivers**

The following Web site has some great tips:  
www.americanheart.org/presenter.jhtml?identifier=3039889

How can I pay for or access the care my child needs?

Services are in place to help you find and pay for your child’s health care. Private and public programs provide access to services and payment options for those services. Trying to find your way through all the information can be overwhelming to anyone. Someone from one of the organizations listed in the previous section should be able to help.

Local epilepsy organizations may be able to provide the names of doctors who accept private insurance, Medicaid, Medicare, CSH, Healthy Families, or Healthy Kids, or doctors who are willing to see patients at reduced rates. They may also know of clinics where care is given free of charge. Online resources offer easy ways to search for doctors as well.

Here are some of the important programs and service delivery systems (i.e., Managed Care) you will come across as you go about getting care for your child.
MANAGED CARE

Managed Care is a common way that services are delivered under medical insurance plans or Medicare. Managed Care links coverage with health care providers in its network. If you have health care coverage through this type of plan, you are an enrollee or subscriber. Enrollees receive their health care from a group of approved doctors and hospitals, called a provider network.

Often enrollees in a managed care plan are assigned to a primary care doctor, or gatekeeper. The gatekeeper's job is to see that the enrollee receives routine medical care such as regular check-ups. The gatekeeper is also the one who decides if the enrollee needs to see a specialist—a doctor or other health care provider who has special training in a particular condition.

The two basic types of managed care plans are Health Maintenance Organizations (HMOs) and Preferred Provider Organizations (PPOs). The main difference between these two plans is that HMOs require their enrollees to receive all of their care from within the plan's network, while PPOs give enrollees the option of using providers either within or outside the plan's network of providers.

Another type of managed care called Exclusive Provider Organization (EPO) is very similar to an HMO. In simple terms, an EPO is a much smaller PPO, offering a very limited number of providers, who offer deeper discounts on their rates because they see a higher volume of patients.

CHILDREN’S SPECIAL HEALTHCARE PROGRAM (CSH)

CSH is a program that pays for specialty health care services for eligible children and young adults with serious or chronic medical conditions. There is a CSH program in every county in Wyoming.

The program is open to children and young adults who meet all of the following requirements:

- Wyoming resident under 19 years of age
- Suspected or known to have one of the medically eligible conditions, i.e. chronic illness or disability. Your child may also have insurance, KidCare CHIP, or EqualityCare (Medicaid)
- Meets financial eligibility

MENTAL HEALTH SERVICES FOR CHILDREN

Mental health refers to the successful performance of brain function, resulting in productive activities, fulfilling relationships with other people, and the ability to adapt to change and cope with adversity.

Children may require the services of a mental health provider for a variety of reasons; their needs may be related to stresses in the home, family, or school or they may be related to stress or an emotional disturbance associated with special needs or learning disabilities. Mental health services are available for children through several different programs. Children who have special health care needs may be eligible to receive services through more than one program, therefore coordination of care is important. The Wyoming Department of Health offers programs to support children enrolled in EqualityCare and Kid Care CHIP and adults with mental health conditions. In addition, Children’s Special Healthcare Program, CSH, may provide funding for all mental health issues related to the CSH.
eligible condition. Finally, commercial insurance may also include a mental health benefit. Information about this can be obtained from your member services department (check your health insurance card for the number).

For a listing of Wyoming County Mental Health Departments, go to the following Web site:

Wyoming DMH Listings by County
wdh.state.wy.us/Media.aspx?mediaId=3311.
or
wdh.state.wy.us/mhsa/treatment/MHprovidermap.html

DEVELOPMENTAL DISABILITIES

Developmental Disabilities Division (DDD)

A Division of the Department of Health, DDD’s mission is “to provide funding and guidance responsive to the needs of people with disabilities to live, work, enjoy, and learn in Wyoming communities with their families, friends, and chosen support service and support providers.” Their programs include the Early Intervention and Education Program, Adult Acquired Brain Injury Waiver Program, Children’s Home and Community-Based Waiver, State of Wyoming Respite Program, and Inventory for Client and Agency Planning (ICAP). Each program’s requirements vary.

Wyoming Division of Developmental Disabilities
wdh.state.wy.us/DDD/index.html

The Governor’s Planning Council on Developmental Disabilities

The Council is a governor-appointed body of people with developmental disabilities, family members, and professionals who work together to promote choice, independence, and the full inclusion of all people with developmental disabilities in community life. The council’s mission is to “seek and advocate for activities, opportunities, and systems that promote self-determination, independence, choice, community involvement, and educational opportunities for people with disabilities, their families and the Wyoming community.”

Governor’s Planning Council on Developmental Disabilities
ddcouncil.state.wy.us

EQUALITYCARE (MEDICAID)

KID CARE CHIP AND CHILDREN’S MENTAL HEALTH WAIVER

Wyoming’s Medicaid program (EqualityCare) provides health insurance to low-income families and individuals who lack health insurance. The most commonly used forms of EqualityCare are:

1. Fee-for-services
2. Managed care
3. Share of cost
4. Waiver services

For those families who do not have health insurance and make too much money to qualify for EqualityCare, Kid Care CHIP provides low cost health insurance. Healthy Families offers different plans from which to choose, and includes health, dental, and vision care. Small copayments may be charged for visits. For more information on qualifications:

Department of Family Services
Offices throughout the state:
dfsweb.state.wy.us/dfs/contact.html#localoffices

12     EPILEPSY AND SEIZURE DISORDER: A RESOURCE GUIDE FOR PARENTS
The Children’s Mental Health Waiver provides individualized services and support for children and youth with serious emotional disturbances and their families. Financial eligibility is based on the child’s own resources.

**Wyoming Children’s Mental Health Waiver**
(307) 777-3352
wdh.state.wy.us/mhsa/treatment/waiverindex.html

---

**MEDICATION ASSISTANCE PROGRAMS**

The following organizations offer prescription drug assistance:

**Partnership for Prescription Assistance**
A partnership bringing together America’s pharmaceutical companies, doctors, other health care providers, patient advocacy organizations, and community groups to help qualifying patients who lack prescription coverage get the medications they need through the public or private program that is right for them.
1 (888) 477-2669
www.pparx.org

**Access2wellness**
Access2wellness provides access to one of the broadest selections of assistance programs that offer more than 1,000 prescription medications, for free or at a discount, to those who qualify.
1 (866) 317-2775
www.access2wellness.com

**Rx Assist**
A nonprofit group that provides information to health care providers to help patients get medications.
www.rxassist.org

**Rx Hope**
A patient assistance company partially funded by the pharmaceutical manufacturers. It promises to speed requests for no-cost medications. The Web site includes information about medications covered by assistance programs.
1 (800) 511-2120
www.rxhope.com

**Together RX Access**
A free prescription savings program for eligible individuals and families who lack prescription drug coverage and are not eligible for Medicare.
1 (800) 444-4106
www.togetherrxaccess.com

**Free Medicine Program**
Established by volunteers, the Free Medicine Program helps patients substantially reduce or completely eliminate their prescription drug costs.
1 (800) 921-0072
www.freemedicineprogram.com
What Is HIPAA?

The Health Insurance Portability and Accountability Act (HIPAA) provides rights and protections for participants and beneficiaries in group health plans. HIPAA includes protections for coverage under group health plans that limit exclusions for preexisting conditions, prohibit discrimination against employees and dependents based on their health status, and allow a special opportunity to enroll in a new plan to individuals in certain circumstances. HIPAA may also give you a right to purchase individual coverage if you have no group health plan coverage available and have exhausted COBRA or other continuation coverage.9

The law also gives you rights over your health information. Providers and health insurers who are required to follow this law must comply with your right to the following:

► view or get a copy of your health records
► add corrections to your health information
► receive a notice telling you how your health information may be used and shared
► decide if you want to give permission before your health information can be used or shared for certain purposes.10

If you believe your rights are being denied or your health information is not being protected you can do the following:

► file a complaint with your provider or health insurer
► file a complaint with the US government

Pharmaceutical manufacturers also provide assistance with prescription drugs for low-income families. Try contacting the specific manufacturers for details.

Abbott Laboratories (Depakote)
1 (800) 222-6885
www.helpingpatients.org

Cephalon, Inc (Gabitril)
1 (866) 209-7589

Eisai (Zonegran)
1 (866) 694-2550

GlaxoSmithKline (Lamictal)
1 (888) 825-5249
www.bridgestoaccess.com

Novartis (Tegretol, Trileptal)
1 (800) 277-2254
www.pharma.us.novartis.com/novartis/pap/pap.jsp

Ortho-McNeil (Topamax)
1 (877) 937-9682/1 (800) 652-6227
http://www.access2wellness.com/a2w/

Pfizer (Dilantin, Neurontin, Zarontin)
1 (866) 776-3700
www.pfizerhelpfulanswers.com

UCB Pharma (Keppra)
1 (800) 477-7877

Valeant (Diastat, Mysoline)
1 (800) 511-2120

For a chart of pharmaceutical companies offering assistance programs for epilepsy medications, see Epilepsy: Patient and Family Guide, 2nd edition, pages 135-138, by Orrin Devinsky, MD, F.A. Davis Company, 2002.8
Who are the members of my child’s health care team?
Possible members:

<table>
<thead>
<tr>
<th>TEAM MEMBER</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatrician</td>
<td>This is a doctor who specializes in the care and treatment of children and is usually the first to see your child for evaluation. This person may also be referred to as your child's primary care provider. This doctor may run some tests and if he or she suspects seizures, will probably refer your child to a neurologist.</td>
</tr>
<tr>
<td>Neurologist</td>
<td>This is a doctor with special training in disorders of the brain, including epilepsy, and other parts of the nervous system.</td>
</tr>
<tr>
<td>Pediatric neurologist</td>
<td>This is a neurologist who specializes in children.</td>
</tr>
<tr>
<td>Epileptologist</td>
<td>This is a neurologist with specialty training in epilepsy who spends most of their time treating people with this disorder. For routine treatment of epilepsy, it is usually not necessary to see an epileptologist. However, for a child with a seizure disorder that is difficult to treat (refractory), an epileptologist may provide an added level of comfort. If possible, an initial consultation with an epileptologist is advantageous and encouraged in new onset epilepsy. Also, consultation may be beneficial in other circumstances, such as counseling about pregnancy and childbirth.</td>
</tr>
<tr>
<td>Neuropsychologist</td>
<td>This is a psychologist with training in the neurobiological causes of brain disorders.</td>
</tr>
<tr>
<td>Pediatric nurse</td>
<td>This is a health care provider educated and trained to care for children who are sick or disabled.</td>
</tr>
<tr>
<td>Pediatric epilepsy nurse</td>
<td>This is a nurse with extra training in pediatrics and neurology.</td>
</tr>
<tr>
<td>Nurse practitioner</td>
<td>This is a registered nurse who has completed advanced training in the diagnosis and management of common medical conditions.</td>
</tr>
</tbody>
</table>
The organizations and Web sites listed below offer services to assist you in finding physicians with expertise in epilepsy.

**Epilepsy Foundation**
8301 Professional Place, East Landover, MD 20785
1 (800) 332-1000
www.aesnet.org/go/find-a-dr/epilepsy-foundation
*Use this Web address to go directly to the page that offers an online search for doctors near you.*

**American Epilepsy Society (AES)**
342 N. Main St.
West Hartford, CT 06117-2507
1 (860) 586-7505
www.aesnet.org
*The AES Web site features a searchable member directory where you can find epilepsy specialists near you.*

**National Association of Epilepsy Centers (NAEC)**
5775 Wayzata Blvd., Suite 200
Minneapolis, MN 55416
1 (888) 525-6232
www.naec-epilepsy.org
*The NAEC is a non-profit trade association whose members include more than 60 specialized epilepsy centers.*

**Epilepsy.com**
Epilepsy Therapy Project
P.O. Box 742
Middleburg, VA 20118
(540) 687-8077
Epilepsy.com provides in-depth information and community for people living with epilepsy.

**Wyoming Epilepsy Association**
119 W. 17th St.
Cheyenne, WY 82001
(307) 634-5329/(307) 286-8637
E-mail: admin@wyomingepilepsy.org
www.wyomingepilepsy.org

---

**How can I assure the best health care for my child?**

**COORDINATION OF CARE**

As more health care professionals are included in your child’s health care team, you may feel overwhelmed and confused. Health care professionals have recognized the need to partner with families in caring for children (see Characteristics of a Medical Home on next page), especially those with special health care needs. For children with epilepsy and their families, such care is therefore ideally provided in a way that feels like a “home,” sometimes referred to as a “medical home,” which is not a building, house, or hospital, but a team approach to providing health care. Your child’s primary care doctor or neurologist may assume the role of the “medical home” by taking the lead in coordinating services and helping you to feel connected and supported. The “medical home” is generally the health care provider who best knows your child.

Your role in this partnership is to share your thoughts and concerns, and to be prepared to discuss them, as well as potential solutions, with your child’s primary doctor. You will want to ask about your child’s medical health care plan (health insurance), medications, referrals, tests, and procedures. Other concerns or issues you may want to discuss with your primary care provider may include recreational activities, school concerns, interactions with siblings, and alternative therapies. Your child’s primary care provider should work closely with you and individuals who provide care to your child to assure that all services and supports are coordinated.
A partnership between your family and your child’s primary care clinician

A relationship based on mutual trust and respect

Connections to supports and services to meet your child’s and family’s needs

Respect for your family’s cultural and religious beliefs

After hours and weekend access to medical consultation about your child

Families who feel supported in caring for their child

Your primary care clinician working with your team of other care providers

Knowing What to Do When a Seizure Occurs

An important way to assure your child’s safety is to know what to do when a seizure occurs. Then you can make others aware of the necessary first aid in case your child has a seizure when they are around.

Most seizures are over so quickly that you do not really have much time to do anything. After it is over, you simply make sure that your child was not injured. Tonic-clonic seizures are the most dramatic and frightening of the seizures, and they usually last longer than other seizures.

CPR is rarely used when a child is seizing. Take a look at the table on the next page to review some basic information that should be considered when responding to a child’s seizure.

Please refer to the chart on page 19 for information specific to different seizure types and recommended first aid for each.

How Can I Make Sure Others Know What to Do?

Remember to tell others about the appropriate first aid for your child. Use the Seizure Action Plan (form F3 in the Appendix section of this guide) and customize it to fit your child’s needs or create your own “what to do” form. Step by step directions should be numbered and should include comforting steps for your child as well as who to call after a seizure. If possible, involve your child in writing these instructions. Hand this form out to babysitters, school personnel, and anyone else who may be taking care of your child. It is important that caretakers be informed about how to handle a seizure and what to do afterwards. Be sure to include how to reach you as well as the telephone number of your child’s doctor.

Water Safety

NEVER leave your child alone in the bathtub or in the pool even for a couple of seconds. This is true for all children, but is especially important for children with epilepsy, as seizure occurrences are not predictable.
<table>
<thead>
<tr>
<th>SEIZURE RESPONSE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>WHAT TO DO DURING A SEIZURE</strong></td>
</tr>
<tr>
<td>▶ Stay calm; most seizures only last a few minutes. Try to time the length of the seizure.</td>
</tr>
<tr>
<td>▶ Move things out of the way so your child will not be injured.</td>
</tr>
<tr>
<td>▶ Lay him or her on one side.</td>
</tr>
<tr>
<td>▶ Make your child as comfortable as possible—loosen any tight clothing around the neck and put a pillow or something soft under the head.</td>
</tr>
<tr>
<td>▶ Pay attention to the length of the seizure.</td>
</tr>
<tr>
<td>▶ Keep onlookers away.</td>
</tr>
<tr>
<td>▶ Be sensitive and supportive, and ask others to do the same.</td>
</tr>
</tbody>
</table>

| **WHAT NOT TO DO DURING A SEIZURE** |
| ▶ Do not put anything in the mouth. While your child may bite his or her tongue during a seizure, trying to put something in the mouth probably will not work to prevent this. You may also get bitten, or you may break some of your child’s teeth. |
| ▶ Do not try to hold your child down. People, even children, have remarkable muscular strength during seizures. Trying to pin a child with a seizure to the ground is not easy and restraint can lead to more serious medical complications. |
| ▶ Do not give mouth-to-mouth resuscitation until the seizure is over. After the seizure has ended, give mouth-to-mouth resuscitation only if your child is not breathing. |

| **WHEN TO CALL 911** |
| Most seizures are not medical emergencies, but an ambulance should be called if: |
| ▶ The seizure lasts longer than five minutes or one seizure immediately follows another. |
| ▶ Your child does not resume normal breathing after the seizure ends. |
| ▶ Your child was seriously injured during the seizure. |
| ▶ Your child or adolescent is pregnant or has diabetes. |
| ▶ The seizure happens in water, and your child may have inhaled water. |
| ▶ Do not call an ambulance during a typical seizure. For a lot of people, the first response to seeing a seizure is to call 911. But for the vast majority of seizures, that is not necessary. It is also frightening for a child to spend an afternoon in the hospital unnecessarily. |

| **WHAT TO LOOK FOR AFTER A SEIZURE** |
| The period immediately after a seizure varies depending on the type, duration, and intensity of the seizure as well as other factors. Some children may feel discomfort from muscle soreness, headache, and pain in the tongue or cheek if those areas were bitten. Your child may feel confused or tired, and his or her skin may appear pale or bluish. For some, the symptoms after a seizure may be more noticeable than the seizure itself. Antiepileptic medications may not alter the seizures, but may minimize those symptoms. |
### FIRST AID FOR SEIZURES

<table>
<thead>
<tr>
<th>SEIZURE TYPE</th>
<th>WHAT TO DO</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Tonic-clonic or Grand Mal</strong></td>
<td>- If it is the first time that child has had a seizure, you should get medical help immediately.</td>
</tr>
<tr>
<td>(loss of consciousness)</td>
<td>- Move any hard, sharp, or hot objects away from the child and remove glasses and any tight neckwear. Put something soft under child’s head. Turn child on one side to keep airway clear.</td>
</tr>
<tr>
<td></td>
<td>- Do not put anything in child’s mouth or give liquids or medications during or immediately after the seizure.</td>
</tr>
<tr>
<td></td>
<td>- Do not try to hold the child’s tongue; it cannot be swallowed.</td>
</tr>
<tr>
<td></td>
<td>- Do not restrain movements unless they place the child in danger.</td>
</tr>
<tr>
<td></td>
<td>- Reassure the child when consciousness returns.</td>
</tr>
<tr>
<td></td>
<td>- Usually it is not necessary to call an ambulance if it is known that the child has epilepsy, and the seizure ends after a minute or two.</td>
</tr>
<tr>
<td></td>
<td>- Call for emergency aid if this is the child’s first seizure of unknown cause, if multiple seizures occur, if the seizure lasts longer than five minutes, or if the child seems sick, injured, or unresponsive.</td>
</tr>
<tr>
<td><strong>Absence or Petit Mal</strong></td>
<td>- Observe the child carefully.</td>
</tr>
<tr>
<td>(loss of consciousness)</td>
<td>- Reassure the child if he or she is frightened or confused.</td>
</tr>
<tr>
<td></td>
<td>- Try to count and record episodes.</td>
</tr>
<tr>
<td></td>
<td>- No first aid is necessary, but if this is the first observation of an absence seizure, medical evaluation is recommended.</td>
</tr>
<tr>
<td><strong>Myoclonic or Jerks</strong></td>
<td>- Speak calmly to the child and other children around him or her.</td>
</tr>
<tr>
<td>(no loss of consciousness)</td>
<td>- If the child is walking, guide him or her gently to a safe place.</td>
</tr>
<tr>
<td></td>
<td>- Stay close until the seizure has ended and the child is completely aware of where he or she is and can respond normally when spoken to.</td>
</tr>
<tr>
<td></td>
<td>- First aid is usually not needed, however, a child having a myoclonic seizure for the first time should receive a thorough medical evaluation.</td>
</tr>
<tr>
<td><strong>Tonic and Atonic or Akinetic</strong></td>
<td>- Comfort the child and check to see if he or she is hurt. A medical check-up is recommended.</td>
</tr>
<tr>
<td>(loss of consciousness)</td>
<td>- No first aid is needed (unless there is injury from a fall), but if this is a first tonic or atonic seizure, the child should be given a thorough medical evaluation.</td>
</tr>
<tr>
<td><strong>Status Epilepticus</strong></td>
<td>- This condition requires immediate medical attention. Call for medical help.</td>
</tr>
<tr>
<td>(loss of consciousness)</td>
<td>- Do not attempt to transport an actively seizing child in your car unless an ambulance is not available.</td>
</tr>
<tr>
<td></td>
<td>- If the child has had this type of seizure before, follow advice specifically given by the child’s doctor.</td>
</tr>
<tr>
<td><strong>Simple</strong> (no loss of consciousness)</td>
<td>- No first aid necessary, but if this is the first observation of the seizure, medical evaluation is recommended.</td>
</tr>
<tr>
<td><strong>Complex</strong> (loss of consciousness)</td>
<td>- Speak calmly to the child and other children around him or her.</td>
</tr>
<tr>
<td></td>
<td>- If the child is walking, guide him or her gently to a safe place.</td>
</tr>
<tr>
<td></td>
<td>- Stay close until the seizure has ended and the child is completely aware of where he or she is and can respond normally when spoken to.</td>
</tr>
<tr>
<td></td>
<td>- Pay attention to the duration of the seizure; most partial seizures last a minute or two. However, a child may be confused for up to half an hour afterward. Longer periods of confusion may mean that seizure activity is continuing and the child needs medical attention.</td>
</tr>
</tbody>
</table>
Advocacy and Support

Where can I find support from other parents like me?

Support groups may be in person, chat rooms (an area on the internet where you can chat with other members), electronic mailing lists (users can subscribe to a mailing list by sending an e-mail note to a mailing list they learn about), or message boards (an application that allows users to post and reply to messages from other users and review the flow of a discussion—also known as a discussion group). Groups can be used to share valuable information about doctors, medications, and treatments. The information, experiences, and successes gained through each family’s journey can be shared to help other families in similar situations.

Always remember that support groups and online groups are made up of other parents like you. Do not take medical advice from an untrustworthy source and always check with your doctor before trying something new.

IN PERSON SUPPORT

You can find face-to-face support groups and information about counseling for parents by calling your nearest Epilepsy Foundation affiliate or the Epilepsy Foundation (see page 10 for contact information). You may also want to contact one (or more) of Wyoming’s Centers designed to help parents.

WYOMING CENTERS

Wyoming Institute for Disabilities (WIND)

WIND is the Wyoming University Center for Excellence in Developmental Disabilities (UCEDD). WIND’s mission is “to assist individuals with developmental and other disabilities and their families to achieve their desired quality of life by promoting and supporting full community inclusion, community membership, independence, productivity, and social participation.” WIND works in partnership with many other organizations in the state to achieve their vision of “a Wyoming where all people can participate in everyday community life as they choose.”

UPLIFT

UPLIFT has been providing support, advocacy, information, and referral to Wyoming families since 1990. They have six fully staffed regional offices in an effort to serve families across the state, including the Wind River Indian Reservation. UPLIFT’s focus has been on children and their families with emotional, behavioral, and learning disorders but works with children with all disabilities and special needs throughout Wyoming. UPLIFT works with parents and other community organizations in a culturally and linguistically appropriate manner to build the service and support capacity for families in that region with the ultimate goal of improving educational and independency outcomes. UPLIFT is founded on successful provision of information, referral, support, and advocacy services for families of children with special needs.

Wyoming Epilepsy Association

Wyoming Epilepsy Association works “to improve the quality of life for individuals who are challenged by the struggles of epilepsy or seizure disorders and other health related issues.” They work toward epilepsy advocacy and awareness to empower people and knock down the “invisible barriers” many people with disabilities face. These efforts include fighting discrimination and promoting people-first language.
Parent Information Center (PIC)

PIC is a project of Parents Helping Parents of Wyoming and is part of a national network of Parent Training and Information Centers (PTI's). PIC's main responsibility is informing and educating parents about special education law and the rights and responsibilities they and their child have under the Individuals with Disabilities Education Act (IDEA). PIC services are available to parents of children ages birth through 21 with any disability or chronic health need. PIC's goal is to become as well informed as possible about the law, education, medicine, human services, rehabilitation, and technology so that the parents and professionals to whom they provide information, referral and support have access to the best information available.

Parent Education Network (PEN)

PEN is the Parent Information Resource Center (PIRC) for Wyoming. PEN's purpose is to offer Wyoming parents learning strategies and to provide technical assistance to schools and families to boost parental engagement, and to ultimately increase student success. PEN believes parents have dreams for their children and their children's welfare, have the capacity to support their children's learning, have responsibility to share an equal-partner role with schools, have parental engagement linked to learning, and have responsibility for building partnerships between school and home.

Family Voices

Family Voices believes in keeping families at the center of children's health care for all children or youth with special health care needs or disabilities. Family Voices provides families with tools to make informed decisions, advocates for improved public and private policies, builds partnerships among professionals and families, and serves as a trusted resource on health care. In Wyoming, Family Voices connects families to a support system of other experienced families to assist in navigation. Wyoming also offers support groups, conferences, listserv, newsletters, and a care notebook.

Wyoming Steps, Inc.

Wyoming Steps is a non-profit agency that provides services to children and adults with developmental disabilities and acquired brain injuries. Their mission is “to provide support, training, and educational programs (STEP) to people with disabilities for the purpose of helping to promote a healthier quality of life.” Wyoming Step provides community and home-based waiver services in Albany County and Laramie County.

Wyoming Protection and Advocacy

Wyoming Protection and Advocacy (P&A) strives “to establish, expand, protect and enforce the human and civil rights of persons with disabilities through administrative, legal, and other appropriate remedies.” P&A acts as an advocate for people with disabilities, especially the individual needs of different populations. The Wyoming P&A office employs attorneys and other professional staff who provide a variety of services.

Wyoming Vocational Rehabilitation Services

Wyoming Vocational Rehabilitation (VR) Services is a part of the Department of Workforce Services for Wyoming. VR works yearly with more than 4,000 of Wyoming's citizens with disabilities, and, on average, approximately 700 of those citizens successfully complete a rehabilitation program and enter the workforce. Services provided include personal assistance services, deaf and blind interpretive services, rehabilitation teaching services, occupational licenses, tools and equipment, self-employment opportunities, and much more.
ONLINE SUPPORT

**Epilepsy Foundation eCommunities**

You can interact with individuals affected by epilepsy from around the world through threaded discussions and real-time chats. Select from the following groups:

- **Parents Helping Parents:** A group for parents and guardians of children with epilepsy.
- **Advocacy:** This group discusses various advocacy issues, including local, state, and federal efforts.
- **Friends, Family, and Loved Ones:** Support for those who give support.
- **SUDEP (Sudden Unexplained Death in Epilepsy):** A group for education, support, and information sharing about this serious subject.
- **The Lounge:** An informal place to hang out and get to know other members.
- **Teen Group:** A place just for teens.
- **Entitled2Respect:** Straight talk for teens about acceptance and epilepsy.
- **KidsClub:** The friendly puppy, Scoop, invites kids of all ages to KidsClub. Come out to the playground, visit the funshop, read some stories, and play games with the puppy.

[www.epilepsyfoundation.org/ecommunities](http://www.epilepsyfoundation.org/ecommunities)

**Epilepsytalk**

This electronic mailing list, available globally, is centered on pediatric epilepsy and is designed to benefit families who wish to share comments, concerns, and questions regarding their experiences with pediatric epilepsy and other related issues.

[www.epilepsytalk.org](http://www.epilepsytalk.org)
Epilepsy Support Group on WebMD

This message board includes discussions about the warning signs of epilepsy or a seizure, which medications are effective, living with the aftermath, and reactions from family and friends.

www.webmd.com/community/boards
(click on ALL BOARDS, then EPILEPSY SUPPORT GROUP or go to boards.webmd.com/topic.asp?topic_id=79)

Epilepsy Resource Center on HealingWell.com:

▶ Message Boards & Chat—Features interactive message boards and chat rooms. Find support and share information with others.

www.healingwell.com/epilepsy/

The Vagus Nerve Stimulation Message Board—Share your experience with VNS.

s4.invisionfree.com/VNS_Message_Board/

What do I need to know about my child attending school?

Federal laws exist that protect the rights of students to receive a public education, attending schools in their community. The federal law, the Individuals with Disabilities Education Act (IDEA) 2004, states that all eligible children are entitled to a free and appropriate public education (FAPE) regardless of their disability and should be placed in the least restrictive environment (LRE) with proper supports. Many children with epilepsy/seizure disorder have seizures rarely or not at all and are able to participate in all school activities and may require few supports. The legal document that defines the supports a child requires in order to benefit from his or her education is called the Individual Education Plan or IEP. Some students who need special accommodations as a result of their medical condition may not require an IEP but instead may benefit from having a Federal 504 Plan. This plan defines those special accommodations that are needed in school, for example, a daily check-in with the nurse for medications or additional time to walk from class to class.

Requests for student evaluations should be put in writing and submitted to the school district director of special education. A parent can request an evaluation by the school to determine their child’s eligibility for special education. The school has 15 school calendar days to contact the parent and to get their consent for testing. Once the consent is signed the school has 60 school calendar days to conduct testing, evaluate results, send the results to the parents, and hold the IEP meeting to determine eligibility. If eligible, this meeting would include development of the child’s IEP.

Children who have seizures may be socially isolated at school for various reasons. They may experience self-esteem issues and a lower
level of achievement. Fortunately, many of these problems can be addressed and prevented when the school develops a good prevention and education plan. School staff, particularly the classroom teacher and the school nurse, can be educated and provide support to the child and other students to improve understanding. Students can benefit the most when the family and school work together. For example, school personnel and the family can monitor the effectiveness of medication as well as any side effects.

The IEP or 504 Plan is developed by parents and school personnel working together with the student. The services and supports your child receives are based on his or her individual needs. Schools are required by federal law to provide the services in the IEP or 504 Plan. If the school is not providing all of the services outlined in the plan, you and your child have the right to file a compliance complaint by calling the Procedural Safeguards Referral Service at (800) 926-0648.

For more information on special education, you can get a free copy of “Special Education Rights and Responsibilities” at Wyoming Protection and Advocacy, Inc. (number and Web site in Resources section on the next page).

RESOURCES—IEP AND THE 504 PLAN

Wyoming Department of Education
www.k12.wy.us

Wyoming Parent Training and Information Centers
  Parent Information Center
  wpic.org
  Parent Education Network
  wpen.net
  UPLIFT
  www.upliftwy.org

The Complete IEP Guide: How to Advocate for Your Special Ed, 5th ed.

Families and Advocates Partnership for Education (FAPE), PACER Center
(952) 838-9000
(952) 838-0190 (TTY)
E-mail fape@fape.org
www.fape.org / www.pacer.org
FAPE aims to improve the educational outcomes for children with disabilities. This Web site links families, advocates, and self-advocates to communicate the new focus of the Individuals with Disabilities Education Act (IDEA).

Learning Disabilities Online
www.ldonline.org
This learning disabilities Web site for parents and teachers has great links to information about learning disabilities.

National Dissemination Center for Children with Disabilities (NICHCY)
www.nichcy.org
EDUCATING PEOPLE AT SCHOOL

According to the experts, the best way to prevent misunderstandings about epilepsy at school is to step in early. At the beginning of the year, talk to your child’s teacher and school nurse and explain that your child has epilepsy. You may want to take some brochures about the disorder. Getting the right information to the right people at school early can make a big difference in your child’s school experience. Refer to the next section “How can I educate others about my child’s epilepsy/seizure disorder?”

Products are available to teach other children about epilepsy/seizure disorder so that students and friends understand more about seizures. You may want to explore some of the resources listed below.

Beach Park
www.epilepsy.org.uk/kids/
Web site for kids.

KidsClub
www.epilepsyfoundation.org/
kidsclub/nonflash/home/
Web site for kids.

The Kids on the Block, Inc.
1 (800) 368-KIDS (5437)
www.kotb.com/kob2.htg/epilepsyfeature

Educational programming (puppet show and curriculum). The Kids on the Block Program on epilepsy was developed in conjunction with the Epilepsy Foundation. It addresses a variety of issues surrounding seizure disorders and provides a basic understanding of seizures, demonstrates seizure first aid, emphasizes the importance of supportive and open friendships, and examines issues of discrimination.

Kids on the Block is offered by:
Wyoming Epilepsy Association
1 (866) 634-5329
How can I educate others about my child’s epilepsy/seizure disorder?

Often times you will have to educate others about epilepsy/seizure disorder and your child’s condition. It is important to make sure that everyone is aware of what to do when a seizure occurs. But you also want to make others feel comfortable when they are around your child. The more knowledge a person has about epilepsy, the more comfortable he or she will feel around your child therefore creating a healthier environment for your child.

When you explain your child’s epilepsy, clearly explain the terminology and then explain in simple terms what this means and how seizures might occur. Encourage others to ask questions. Attempt to remove the mystery around your child and his or her seizures. You may want to hand out a brochure or even a video so they can find out a little more on their own and have time to process the information.

The way in which others are informed affects how they treat your child. The same goes for your child as well. If you learn to talk about epilepsy/seizure disorder, your child will too. Children are more likely to feel confident about themselves and accept their epilepsy when they know how to talk about it and are not ashamed.

Children and youth with epilepsy must also deal with the psychological and social aspects of the condition. These include public misperceptions and fear of seizures, uncertain occurrence, loss of self-control during the seizure episode, and compliance with medications.

Openness and honesty are the best ways to combat any stigma surrounding epilepsy/seizure disorder and you will probably find that most people are empathetic and supportive if they understand the condition and know how they can help.

Please make sure your child is given many opportunities to ask questions and discuss his or her concerns or fears. When possible, he or she also needs to be encouraged to take on some aspects of management of the condition.

You may be getting to know a great deal about epilepsy/seizure disorder, but there are still a lot of people out there who do not. As you educate others about your situation and that of your child, you will help spread the word and reduce the stigma surrounding epilepsy. Please do not take others’ lack of knowledge personally.

Materials are available to help make the task of educating others easier. Many find that a good videotape or DVD to hand out to a school teacher or babysitter can help tremendously. For recommendations and to obtain materials, contact your local Epilepsy Foundation affiliate, Parent Training and Information Center (PTIC), or Community Parent Resource Center (CPRC).

The Colorado Epilepsy Foundation affiliate in Colorado (EFCO), and the Wyoming Epilepsy Association (WEA) provide seizure first aid training free of charge. Most of the educational resources are free as well.

A variety of forms are available to help keep track of seizures and other information about your child’s condition. The following pages contain a selection of such forms in reduced size. Choose the pages that work for you and make copies so you can easily track your child’s records. You may print full size copies from PDF format, access the Guide online at the Wyoming Institute for Disabilities (WIND) and The Wyoming Epilepsy Association Web site.
References


Appendix forms

Seizure Records

Seizure Log F1
Seizure Description Sheet F2
Seizure Action Form F3

Going to the Doctor

My Child’s Profile F4
Medication Log F5
Doctor’s Visit Notes and Tips F6

Communications

Encounter Log F7

Please photocopy and use these forms.
# Seizure Log

**Directions:** Use this form to keep track of your child’s seizure activity. Your notes will help your nurse and physician in further treatment of your child.

<table>
<thead>
<tr>
<th>Date</th>
<th>Time Seizure Began</th>
<th>Time Seizure Ended</th>
<th>Current Medication and Dose</th>
<th>Description of Seizure</th>
<th>Any Intervention?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This tool was developed from the federal Maternal and Child Health Bureau, Health Resources and Services Administration under grant #H98MC08579, funded to the USC UCEDD, Children’s Hospital Los Angeles, through its initiative, Project Access: Improving Care for Children with Epilepsy. Support was also provided by two other partners in Project Access: the National Initiative for Children’s Healthcare Quality (NICHQ), under grant #U23MC058893, and the Epilepsy Foundation, under grant #U23MC05909.
Seizure Description Sheet

Directions: Please check (✓) what happens (or happened) during your child’s seizure and bring this sheet to your child’s neurology appointment.

### Description of Spell or Seizure

<table>
<thead>
<tr>
<th>Body</th>
<th>□ white</th>
<th>□ right</th>
<th>□ left</th>
<th>□ can’t tell</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[Picture of a person]</td>
<td>[Picture of a person]</td>
<td>[Picture of a person]</td>
<td>[Picture of a person]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Movement</th>
<th>□ jerking</th>
<th>□ stiffness</th>
<th>□ jerking and stiffness</th>
<th>□ can’t tell</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[Person jerking]</td>
<td>[Person stiff]</td>
<td>[Person jerking and stiff]</td>
<td>[Person can’t tell]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Eyes</th>
<th>□ up ↑</th>
<th>□ closed</th>
<th>□ right →</th>
<th>□ left ←</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[Eye open]</td>
<td>[Eye closed]</td>
<td>[Eye right]</td>
<td>[Eye left]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Eyes</th>
<th>□ stare</th>
<th>□ stare and blink</th>
<th>□ no change</th>
<th>□ can’t tell</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[Eye staring]</td>
<td>[Eye staring and blinking]</td>
<td>[No change]</td>
<td>[Can’t tell]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Skin Color</th>
<th>□ blue</th>
<th>□ no change</th>
<th>□ can’t tell</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[Hand blue]</td>
<td>[Hand no change]</td>
<td>[Hand can’t tell]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Toilet</th>
<th>□ pee-pee</th>
<th>□ poop</th>
<th>□ none</th>
<th>□ can’t tell</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[Urinal]</td>
<td>[Poo]</td>
<td>[No poo]</td>
<td>[Can’t tell]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Mouth</th>
<th>□ dry</th>
<th>□ drool</th>
<th>□ foam</th>
<th>□ bite tongue</th>
<th>□ can’t tell</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[Dry lips]</td>
<td>[Drooling]</td>
<td>[Foam]</td>
<td>[Bite tongue]</td>
<td>[Can’t tell]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>How Often</th>
<th>□ daily</th>
<th>□ weekly</th>
<th>□ monthly</th>
<th>□ other:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[Daily]</td>
<td>[Weekly]</td>
<td>[Monthly]</td>
<td>[Other:]</td>
</tr>
</tbody>
</table>

### After Seizure or Spell

<table>
<thead>
<tr>
<th>□ asleep</th>
<th>□ drowsy</th>
<th>□ alert</th>
<th>□ confused</th>
<th>□ paralyzed</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Asleep]</td>
<td>[Drowsy]</td>
<td>[Alert]</td>
<td>[Confused]</td>
<td>[Paralyzed]</td>
</tr>
</tbody>
</table>
Seizure Action Plan

This form provides information to assist in the safe care of this child during a seizure

Name: ___________________ DOB: ___________ Age: _______ Age during 1st Seizure: _______
Alt. Contact: ___________________ Home: ___________________ Work: ___________ Cell: ___________
Primary Doctor: ___________________ Office: ___________________ Fax: ___________ Alt.: ___________
Neurologist: ___________________ Office: ___________________ Fax: ___________ Alt.: ___________

Seizure Information

<table>
<thead>
<tr>
<th>Seizure Type</th>
<th>Length</th>
<th>Frequency</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Does your child act differently before a seizure? □ Yes □ No Explain: ____________________________________________

How the child acts after a seizure is over: ________________________________________________________________

How other illnesses affect child's seizures: ________________________________________________________________

Seizure First Aid

In addition to Basic Seizure First Aid, the following procedures should be done when the child has a seizure: ________________________________________________________________

Will child need to leave the classroom? □ Yes □ No
Should an extra change of clothes be kept at school? □ Yes □ No
If "yes," where are they kept? ________________________________________________________________

Does child have a VNS? □ Yes □ No
If "yes," when should magnet be used? ________________________________________________________________

Seizure Emergency

A seizure emergency for this child is: ________________________________________________________________

☐ Administer emergency medication listed below
☐ Contact school nurse at: ________________________
☐ If an ambulance is required, transport to: ________________________
☐ Notify parent at: ________________________
☐ Notify emergency contact at: ________________________
☐ Notify doctor: ________________________

Doctor's Name ________________________ Phone ________________________

Basic Seizure First Aid

- Stay calm & track time
- Keep the child safe
- Do not restrain
- Do not put anything in mouth
- Stay with the child until fully conscious
- Record seizure in log
- Turn the child on their side

For tonic-clonic seizures:
- Make sure head is protected
- Keep airway open/watch breathing

A Seizure is generally considered an EMERGENCY when:
- A convulsive (tonic-clonic) seizure lasts longer than 5 minutes
- The child has repeated seizures without regaining consciousness
- The child has a first time seizure
- The child is injured or has diabetes
- The child has breathing difficulties

Emergency Medication Plan

<table>
<thead>
<tr>
<th>Medication</th>
<th>Dosage</th>
<th>Method</th>
<th>When to use</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Special Considerations/Safety Precautions
(with school activities, sports, trips, etc.) ________________________________________________________________

Doctor Name & Signature: ________________________
Parent/Guardian Signature: ________________________
School Nurse Signature: ________________________

This tool was developed from the federal Maternal and Child Health Bureau, Health Resources and Services Administration under grant ‘F98MC8579, funded to the USC UCEDD, Children's Hospital Los Angeles, through its initiative, Project Access: Improving Care for Children with Epilepsy Support was also provided by two other partners in Project Access: the National Initiative for Children's Healthcare Quality (NICHQ), under grant ‘U2MC038893, and the Epilepsy Foundation of America, under grant ‘U2MC039699.'
# My Child’s Profile

<table>
<thead>
<tr>
<th>Name</th>
<th>Eye Color</th>
<th>Blood Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date of Birth</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Height</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Languages Spoken</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Telephone</td>
<td>Home</td>
<td>Work</td>
</tr>
<tr>
<td>Address</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parent/Guardian</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Telephone</td>
<td>Home</td>
<td>Work</td>
</tr>
<tr>
<td>Address</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parent/Guardian</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Telephone</td>
<td>Home</td>
<td>Work</td>
</tr>
<tr>
<td>Address</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Emergency Contact</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Relation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Telephone</td>
<td>Home</td>
<td>Work</td>
</tr>
</tbody>
</table>

## PRIMARY Health Insurance

- Health Insurance Plan
- Plan Number

## SECONDARY Health Insurance

- Health Insurance Plan
- Plan Number

---

This tool was developed from the federal Maternal and Child Health Bureau, Health Resources and Services Administration under grant #H08MC08579, funded to the USC UCEDD, Children’s Hospital Los Angeles, through its initiative, Project Access: Improving Care for Children with Epilepsy. Support was also provided by two other partners in Project Access: the National Initiative for Children’s Healthcare Quality (NICHQ), under grant #U23MC038903, and the Epilepsy Foundation, under grant #U23MC03909.
My Child’s Profile
(Continued)

Primary and Secondary Diagnosis

Medical Alerts

Most Recent Hospitalization History Including Surgeries

<table>
<thead>
<tr>
<th>Date</th>
<th>Reason</th>
<th>Name of Hospital</th>
<th>Attending Doctor</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Primary Care Physician

Office Number

Neurologist

Office Number

Notes
**Medication Log**

**Directions:** Complete the chart below, listing all medications (prescription, over-the-counter, vitamins and supplements) your child takes. If the directions for a medication changes, mark through that row and begin a new row with updated directions. Be sure to bring this sheet with you to all of your child’s medical appointments so you can review the information with your child’s providers.

<table>
<thead>
<tr>
<th>Start Date</th>
<th>Name of Medication</th>
<th>Dosage</th>
<th>Frequency</th>
<th>What is it used for?</th>
<th>End Date</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
# Doctor’s Visit Notes

**Directions:** Use this form to remind yourself about questions you have for your child’s doctor and to record your own notes from the visit.

<table>
<thead>
<tr>
<th>Date of Appointment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Doctor’s Name/Phone Number</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
</tbody>
</table>

## Before the Visit

1. Reason for today’s visit?

2. How long has it been going on?

3. What makes it better/worse?

4. What have you tried so far?

5. What do you hope will happen at this visit?

## Notes from the Visit

1. What is the diagnosis?

2. Does my child need a prescription? If yes, what is the medication and dosage?

3. What should the medication do and when?

4. Is there anything I should watch out for/side effects?

5. If no medication is needed, what should I do for my child to resolve the issue?

6. What should I do if my child gets sick or has more symptoms?

7. Where can I get more information?
Doctor’s Visit Tips

Get Ready

1. Bring any information that you have about your child (My Child’s Profile)
2. Bring pen/pencil and paper to take notes
3. Check your data and be specific about all changes in your child’s health status (Seizure Log, Seizure Description Sheet)
4. Have a list of all medications your child is currently using, including over-the-counter drugs (Medication Log)
5. Have a list of reactions your child has experienced from any medications, prescribed or over-the-counter
6. Write all of your questions down before visiting the doctor (Doctor’s Visit Notes)

Tell the Doctor:

1. How your child has been doing
2. Information about successes and setbacks
3. Detailed information about changes and symptoms that are different from your child’s normal status
4. What you are concerned about
5. When the symptoms started changing
6. How often and when the symptoms occurred
7. What you have tried to relieve the symptoms and your child’s response

Don’t Leave the Doctor’s Office Without:

1. Instructions and name(s) for new and old medication(s)
2. Asking how long the child will be on medication and whether there are refills
3. Asking about possible side effects or cross-reactions of medication(s)
4. Asking what the child can eat with new medication(s)
5. Asking if you need authorization before filling the prescription for your specific insurance provider
6. Understanding all follow-up questions including how to report changes in symptoms
7. Asking the doctor if you need a follow-up appointment
8. Making the next appointment, if needed
# Encounter Log

<table>
<thead>
<tr>
<th>Date</th>
<th>Name or Agency/Phone</th>
<th>Purpose</th>
<th>Type</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Telephone</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Written</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>In person</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Email</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fax</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Telephone</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Written</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>In person</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Email</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fax</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Telephone</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Written</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>In person</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Email</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fax</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Telephone</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Written</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>In person</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Email</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fax</td>
<td></td>
</tr>
</tbody>
</table>

This tool was developed from the federal Maternal and Child Health Bureau, Health Resources and Services Administration under grant #H98MC08579, funded to the USC UCEDI, Childrens Hospital Los Angeles, through its initiative, Project Access: Improving Care for Children with Epilepsy. Support was also provided by two other partners in Project Access: the National Initiative for Children’s Healthcare Quality (NICHQ), under grant #U23MC09893, and the Epilepsy Foundation, under grant #U23MC09899.
University of Southern California
University Center for Excellence in Developmental Disabilities
Childrens Hospital Los Angeles

An electronic version (PDF format) of the Guide can be accessed via the Internet at the WIND Web site:

http://www.uwyo.edu/wind/community/education/ProjectAccess.asp

Project Access: Improving Care for Children with Epilepsy is a grant (#H98MCO8579) from the federal Maternal and Child Health Bureau, Health Resources and Services Administration, awarded to the USC University Center for Excellence in Developmental Disabilities, Childrens Hospital Los Angeles. The grant subcontracts with UCEDDs in Alaska and Wyoming, Family TIES of Nevada, and the Epilepsy Foundation of Northern California. The Epilepsy Foundation Northwest and the Epilepsy Foundation of Colorado also provide support. Two other partners in Project Access also provide support: the National Initiative for Children's Healthcare Quality (NICHQ) and the Epilepsy Foundation.