Partnering with your Doctor

The Medical Home Approach

A guide for families of children with special health care needs

Region 4
Midwest Genetics Collaborative
ACKNOWLEDGEMENTS

Thank you to the families and Region 4 Genetics Collaborative Medical Home Education Workgroup members who contributed their time and effort to produce this helpful guide. The information and resources are intended for educational use only and are provided solely as a service. The information provided should not be used for diagnosing or treating a health problem or disease, and is not a substitute for professional care.

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INTRODUCTION

Partnering with your Doctor: The Medical Home Approach is a guide to help you and your family connect with your doctor, other medical professionals, and caregivers through a medical home.

Some information in the guide might be helpful now, while other sections will be useful in the future.

The Guide is intended to be a user-friendly, hands-on tool to support you and your family to move forward in obtaining and providing a medical home for your child. This guide provides definitions, examples, and tools for you to use when working with your doctor to develop a medical home.

The Guide is designed to:

1. Provide a detailed definition and description of the medical home concept
2. Lay the foundation for understanding and advocating for a medical home
3. Provide tools and examples for parents to use and learn from while creating and maintaining a medical home with their doctor
4. Bring attention to the importance of linking families of children with special health care needs to a medical home
5. Provide advice on how to effectively partner with your child's doctor
6. Define family-centered care and demonstrate how it benefits families
7. Demonstrate how to seek culturally effective, compassionate care
8. Assist families through the transition process by providing resources and tips
9. Provide a list of resources and links to organizations that support families and the medical home concept

To request additional copies of the Guide, please contact Region 4 Staff via email at info@region4genetics.org or phone 517.381.8247. The Guide also may be downloaded as a PDF file free of charge from www.region4genetics.org.

Note: The Region 4 Genetics Collaborative’s mission is focused on children with genetic conditions. Often, children with genetic conditions have special health care needs. This guide was developed with children who have genetic conditions in mind, but would be useful for any children with special health care needs. Hereafter, the broader population of children with special health care needs will be used, rather than children with genetic conditions.
Dear Families:

The birth of my twin girls was both a joyous and overwhelming experience. Dealing with the reality of one child being healthy and the other having birth defects created challenges that we were not prepared for. My daughter was born with a grouping of anomalies: a specific group called V-A-C-T-E-R-L. We didn’t know she had VACTERL until all of the anomalies were identified during her 21-day stay in neonatal intensive care. It was hard to come to terms with the fact that our 3-pound, 9-ounce baby girl could have such a hard start at life from birth. We were determined to give her a full and happy life and began the medical journey to help her.

During the first year of her life, it was a struggle to keep up with all the appointments with specialists for her heart, kidney, lungs, legs, and back. I returned to work shortly after my maternity leave ended, only to resign eight weeks later. The twins needed more one-on-one attention both maternally and medically. There were 12 or more specialists we saw separately. In addition to the specialists, she also was seen by public health representatives, we were involved in some community services available, and she had to see a few therapists for speech and physical therapy.

At each visit/appointment, a history form had to be completed and I had to tell them all about her condition and her current status at the time. After a while it became a very robotic process. Not only was I providing information, but I also was receiving a wealth of information that needed further follow up and created more office visits. Immediate family members also wanted updates. It was all exhausting and the medical journey became unmanageable at times.

My daughter’s primary care doctor was helpful at making sure we had the necessary referrals to the specialists, but I felt that I was left to manage the rest. So, being the organized mother I am, I created a paper file for each representative, service, therapist, specialist, and so on. I also put all appointments on our family calendar. This was my only means to keep it all organized. This worked for many years but something was still needed. My daughter needed more comprehensive primary care. I wanted to be more of a valued partner to access and coordinate specialty care and community services that were important for my daughter’s overall health and well being.

I needed to be part of a medical home! I still do. I am looking forward to using this guide to expand my knowledge of what a medical home is, have access to resources in my state and nationally, and enhance my parent leadership skills to be a participant in a medical home.

Nancy (Mother of Audrey, Lucy and Maureen)
HOW TO USE THIS GUIDE
Each section of this guide describes an important component of the medical home. The sections have been assigned a color to keep the section material organized and easy to find.

- ACCESSIBLE CARE
- CONTINUOUS CARE
- COMPREHENSIVE CARE
- COORDINATED CARE
- COMPASSIONATE CARE
- CULTURALLY EFFECTIVE CARE
- FAMILY-CENTERED CARE
- ADVOCATING FOR YOUR CHILD
- GLOSSARY
- MEDICAL HOME RESOURCES

Blue boxes titled “Parent-to-Parent Tips!” are placed throughout the guide and showcase helpful hints and practical advice from parents who have children with special health care needs.

Below are two icons you will find throughout the guide:

- **Designates a page that includes a tool or list of questions for you and your family to think about, fill out, and possibly share with your doctor and other family members.**

- **Indicates a tip or suggestion for you and your family to consider for a variety of situations, such as working with doctors, planning for life events, and other circumstances.**
THE MEDICAL HOME DEFINED

The American Academy of Pediatrics (AAP) describes the medical home as a model of delivering care that is accessible, continuous, comprehensive, family-centered, coordinated, compassionate, and culturally effective. The AAP has begun many initiatives to promote the adoption and spread of the medical home.

The AAP joined with the American Academy of Family Physicians, American College of Physicians and the American Osteopathic Association to publish a set of principles on the medical home for doctors to focus on: personal physician; physician-directed medical practice; whole person orientation; coordinated care; quality and safety; enhanced access; and appropriate payment. The AAP developed an introduction to the medical home (below) that represents the specific needs of the pediatric population because every child deserves a medical home.

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**American Academy of Pediatrics Introduction to the Medical Home**

*Family-centered partnership:* A medical home provides family-centered care through a trusting, collaborative, working partnership with families, respecting their diversity and recognizing that they are the constant in a child’s life.

*Community-based system:* The medical home is an integral part of the community-based system. A family-centered, coordinated network of community services is designed to promote the healthy development and well being of children and their families. As such, the medical home works with a coordinated team, provides ongoing primary care, and facilitates access to and coordinates with a broad range of specialty, ancillary and related community services.

*Transition:* The goal of transition is to optimize life-long health and well being and potential by providing high-quality, developmentally appropriate health care services that continue uninterrupted as the individual moves along and within systems of services and from adolescence to adulthood.

*Value:* Recognizing the importance of quality health care, appropriate payment for medical home activities is imperative. A high-performance health care system requires appropriate financing to support and sustain medical homes that promote system-wide quality care with optimal health outcomes, family satisfaction, and cost efficiency.²

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[www.pcpcc.net/files/preamble_joint_preamble_0.pdf](http://www.pcpcc.net/files/preamble_joint_preamble_0.pdf)

² What is a Medical Home? National Center for Medical Home Implementation. [www.medicalhomeinfo.org/](http://www.medicalhomeinfo.org/)
HAVING A MEDICAL HOME MEANS YOUR CHILD’S CARE IS: ³

1. **Accessible**
   - Care is provided in your community
   - The doctor’s office accepts your insurance
   - The location meets American Disabilities Act (ADA) requirements
   - Access to health advice or care is available around the clock, 365 days per year
   - If needed, the office is accessible by public transportation

2. **Continuous**
   - The same doctors are available from infancy through adolescence
   - Communication occurs between doctors and specialists to ensure your child’s needs are met at any age
   - Assistance is provided to secure a medical home for adolescents transitioning to adult care

3. **Comprehensive**
   - Preventive and primary care is provided
   - Growth and development assessments are conducted when appropriate
   - Care coordination is emphasized and exercised between all doctors for your child
   - Referrals to specialty-care services are available and offered when needed
   - Child/family counseling and health education are a part of visits or available upon request
   - Your doctor is knowledgeable about community resources and shares information

4. **Coordinated**
   - You and your family are linked to appropriate support, educational, and community services
   - Your doctor’s office staff communicate and work with all agencies to ensure all your child’s needs are met
   - A centralized record containing important information on your child’s care will be maintained by you, your doctor and office staff

5. **Compassionate**
   - The doctor shows concern for the well being of your child and family as a whole
   - The doctor shows understanding for the variety of emotions because of your children with special health care needs
   - Office staff are flexible and respond to requests and needs
   - Office staff offer privacy when requested
   - Appointment times are scheduled with consideration of your and your family’s needs

6. **Culturally Effective**
   - Your family’s cultural background is recognized, valued, and respected
   - Office staff are sensitive to stereotypes and cultural assumptions
   - Multi-language materials and translation services are made available as needed

7. **Family-Centered**
   - Office staff recognize that you and your family are the principle caregivers and the center of strength and support for your child
   - Your doctor and family share the responsibility in decision making

³ Adapted from: What is a medical home? And what does it mean for you and your child? University at Chicago, Division of Specialized Care for Children, publication 40, 16, 2003.
WHAT IS ACCESSIBLE CARE?

Accessible care means having a doctor whose office is available, not only in terms of location, but also meets or exceeds the requirements of disability access. The doctor’s office also accepts your insurance, and you are able to speak with your doctor when you need to. These are all important components to the structure of your medical home.

A medical home is not just an accessible building or hospital, but an approach to providing quality health care services. Your primary care doctor, specialists, and other health care professionals act as partners in a medical home to identify and access all of the medical and non-medical services needed to help your child and family. By working together and identifying barriers, your medical home is creating an accessible space for the care of your child.4

THE ROLES OF FAMILY MEMBERS IN THE MEDICAL HOME

You and your family are recognized as:

- **Constants** in the life of your child
- **Experts** in the individual strengths and needs of your child
- **Supervisors** of those who coordinate your child’s care
- **Visionaries** who see the “big picture” for your child and his future

FAMILY REACTIONS TO THE CONCEPT OF THE MEDICAL HOME

“Sign me up! I’ll take one!”

“Everything is at your fingertips for easy access.”

“Something like this would take the weight off...you could relax a little...”

“It sounds too good to be true!”

“Complete care. Complete care that runs smoothly.”

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4 Adapted from the brochure *Medical Home: Families and Providers Working Together* - developed by the Washington Medical Home Leadership Network. [http://www.medicalhome.org/leadership/brochures.cfm](http://www.medicalhome.org/leadership/brochures.cfm)
Clinical skills & knowledge
Does the doctor have training and/or specialty interests important to your child’s needs?
*Selecting a doctor with an interest in children’s special needs is a benefit.*

Experience
Has the doctor cared for other children who have a diagnosis similar to your child’s or cared for children with developmental delays or disabilities in general?
*Experience can help your doctor to be more aware of resources and services that might benefit your child and your family.*

Mutual respect and sense of connection
Is the doctor approachable? Does the doctor make you feel comfortable?
*Choosing a doctor may mean finding a balance between technical skills, interpersonal skills, and experience. It’s up to you to decide what you and your child need right now and you may find that what you need changes over time.*

Don’t expect perfection
Every relationship has rough times. Be willing to be flexible, if needed.
*Recognize that doctors are human too, and give them the chance to make things better. If you are mostly pleased with your child’s doctor, then it may be worth working through rough times.*

Be a role model
Show by example how you want to be treated and how you want your child to be treated.
*If you want the doctor to listen to you, be a good listener, too. Talk about the good things as well as your concerns. Share pictures and stories so he can appreciate and get to know your child.*

Be understanding
Doctors often must have appointments back-to-back, every 15 minutes or even sooner.
*If it seems that the doctor is in a hurry to move on to the next appointment, you are probably right! If you need more time, let her know and maybe she would be willing to work it out.*

Be a partner
Decide what type of partnership you want with your child’s doctor.
*Whatever type of partnership you have, you should feel comfortable asking questions, sharing your insights, and feeling like you and the doctor are part of a team. In turn, you should be open to the doctor’s questions and insights as part of your child’s team.*

Express gratitude
Say thank you, in person or in writing.
*Let your doctor know what is helpful and when he is doing a good job.*
Call the office you are interested in, explain that you are looking for a new doctor and ask to speak to someone who can answer a few questions for you. Use this sheet as a starting point to interview the members of the office staff (receptionist, nurse and doctor) to get answers that will inform your decision.

Questions to ask the office receptionist

1. Is this practice accepting new patients?

2. Does this office regularly see children with (specify your child’s condition)?

3. If no, ask: Do staff members have experience with children with special health care needs?

4. How long does an average appointment take? How long does an expanded appointment take?

5. Does your office accept (name of your insurance, Medicaid, etc.) for payment of services? Does your office bill my insurance company directly?

6. What is the average cost of an office visit? If the office charges are above what my insurance considers to be “reasonable” are you able to waive the remainder of the balance?

7. Does the office offer same-day sick appointments?

8. What is the average waiting time to have an appointment?

9. What is your telephone call policy? Does the doctor give advice or refill medications over the phone?

10. Does your office charge for advice calls during the day, advice calls after hours, or medication refills?

11. Does your office communicate or respond to questions through email?

12. What are the office hours? Are weekend and/or evening hours available?

13. Will appointments always be at this location? Are there other offices I may have to travel to for appointments?
Don’t forget to include and prioritize your own questions. If you feel comfortable with the responses, ask to speak with one of the medical staff (nurse or doctor) to continue with your “interview”. If the answers you received are not satisfactory, thank them for their time and call another office.

**Questions to ask the nurse**

1. What types of services does your office offer? Which are provided by a physician and which by a physician’s assistant or nurse?

2. Does your office have a care coordinator? Or, is the doctor comfortable being my child’s medical care coordinator?

3. Are tests and lab work done at the office, or do we need to go to another location?

4. Does your office have a specialist referral list available?

5. Does your office provide referrals for other services that might help my child and family? *Example: mental health services, early intervention programs, etc.*

**Questions to ask the doctor**

1. What approach do you take when multiple professionals are involved in the care of my child? *Example: Is there a team approach to care when other doctors, physical therapists, teachers, or home care nurses are involved?*

2. Which hospital(s) do you have admitting privileges to?

3. Who will my child see if you are not available?

4. Are you comfortable discussing alternative treatments? *(Homeopathic remedies/therapies)*

5. Do you allow appointments specifically to ask questions and discuss a plan of care?

6. How would you feel if I ever asked for a second opinion?

7. If I have an after-hours need, will I be able to reach you or another doctor in this office?
DECIDING TO CHANGE DOCTORS?

Parents are often pleased with their child’s doctor, but sometimes things just don’t feel right and your desire to be part of a medical home is not getting through to your doctor. Here are some questions to ask yourself if you are feeling uncertain about the quality of care your child is experiencing.

1. Does the way the office is run work for you?

2. Is the doctor available when needed?

3. Are you often confused about recommendations and why certain treatments are prescribed?

4. Do you feel like your concerns are not taken seriously?

Remember: Trust your own judgment and view yourself as an expert on your child.

If these questions make you doubt your instincts or your doctor’s...trust your gut. If the relationship doesn’t feel right and you haven’t been able to make it better, then it is time to consider changing doctors.

Parent-to-Parent Tip!

Try a positive approach through a phone call or written note. Example: “Thank you for all you have done for my child. We really appreciate the time you have spent with us. But right now this is not the perfect fit for our family – I would like my child to see someone else.”
WHAT IS MEDICAL TRANSITION?

Growing up is not always easy for children and it can be hard on parents, too. When your child has special health care needs, growing up may be more complicated. Still, many parents of children with special health care needs find it helpful to remember the "bottom line," you want your child to be as independent as possible. Independence in adult life (health care, employment, living, and recreation) takes preparation. The process of growing up and becoming independent is sometimes called "transition." The term refers to transition from adolescence to adulthood and is often used when your child is:

- Learning to be more independent
- Learning to manage his own health care
- Changing from pediatric to adult-oriented medical care
- Moving from school to work and other aspects of adult life

The medical home definition of transition is to thoughtfully coordinate what is appropriate for the child to ensure successful transition to:

- Adult health care system
- Work
- Independence
- Inclusion in community life

Transitioning your child to adult-oriented medical care is the doctor’s responsibility. Continuity of care ensures communication across the developmental continuum and among multiple doctors. It guides you through the transition from the child health care system to the adult health care system and provides you and your child with the support needed to make informed decisions about his health care needs. A medical home provides continuity of care and allows your child to successfully transition into the role of his own primary advocate.
MEDICAL TRANSITION, CONTINUED

Youth: managers of their own health care
For teens (who are able to be responsible for their own care) to become independent, they need family members to take less active roles in their medical care. This way they gain independence and skills to plan for and assist in medical transition.

You can assist your child early on by:

- Helping her understand why she is going to the doctor
- Teaching your child basic information about her condition(s)
- Encouraging your child to learn about her medical condition(s)
- Letting your child meet with doctors in private
- Teaching your child the names of medications, the dosage, why the medication is prescribed, and any side effects
- Letting your child ask health-related questions so she may gain important skills in managing her own care

Helping your teen prepare
Teen years are an important time in life. Transition planning is recommended to start by age 14 or younger. Here are tips on medical transition:

1. Help your child create and manage a care plan to keep track of his health care needs
2. Create a calendar for your child’s appointments and give your teen gentle reminders when appointments are coming up
3. Encourage your child to ask health-related questions as well as any other things he may want to know from the doctor
4. Ask your doctors and other adults to talk directly to your teen

See Page 14 for a checklist to help you plan your child’s health care management goals.

“...I have encouraged Cody to understand his condition, and little by little he is taking interest in the management of his own care. He keeps his own medical record at school and in his backpack. Now, I can feel confident giving Cody some independence.” Claire (Cody’s mom)

Genetic counselors and your child
As your teen approaches adulthood and independence, often she becomes interested in meeting with someone to discuss her condition in more detail as she learns about managing her own health care. A genetic counselor can address concerns your child may have as she thinks about starting her own family someday. Ask your primary doctor to refer your child to a genetic counselor or center if you have not been in contact with one.

Parent-to-Parent Tip!
The process of parents gradually “letting go” of the care of their children is critical to the adolescent years. These teens, who will soon be young adults, will take charge of their own lives - including their health. While it can be a hard process, know that you are doing right by your child by transferring responsibilities and management of care to your child.
### HEALTH CARE SKILLS CHECKLIST

<table>
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<th>SKILL/RESPONSIBILITY</th>
<th>PLAN TO START</th>
<th>NEEDS PRACTICE</th>
<th>PERFORMS PARTIALLY</th>
<th>PERFORMS ON OWN</th>
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<tr>
<td>Understands and describes medical condition(s)</td>
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<td>Understands how medical condition affects life</td>
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<td>Accesses medical records &amp; health information</td>
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<td>Prepares and asks questions of doctors</td>
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<td>Knows medications and what they are for, or carries information</td>
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<td>Is responsible for taking own medication</td>
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<td>Is responsible for doing own treatments</td>
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<td>Gets a prescription filled/refilled</td>
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<td>Calls to schedule medical/dental appointments</td>
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<td>Knows how to access transportation for appointment</td>
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<td>Knows height, weight, and birth date</td>
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<td>Knows how to care for self (basic needs: feeding; hygiene; etc.)</td>
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<td>Cleans &amp; maintains own living space</td>
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<td>Makes financial decisions (pays bills, manages checking/savings accounts)</td>
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<td>Knows where income will come from</td>
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<td>Knows where he will live</td>
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<td>Knows what path he is on after high school</td>
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<td>Knows health emergency phone numbers or carries information</td>
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<td>Knows about and understands medical insurance</td>
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<td>Obtains sex education materials/birth control/family planning information as needed</td>
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<td>Makes contact with community resources</td>
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<td>Considered need for health advocate, if necessary</td>
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<td>Has identified physicians for adult care (primary, specialty and gynecological, if applicable)</td>
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<td>Knows how to hire and manage a personal care attendant, if needed</td>
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<td>Has considered guardianship, if needed</td>
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<td>Knows how medical care will be paid for</td>
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5 Adapted from Children’s Hospital in Boston, MA and the PACER Center
There may come a time to talk with your child’s doctor about transitioning to adult-oriented care. The doctor may bring it up during your child’s regular appointments or when he sees your child becoming more independent and taking ownership in his care. You may feel the need to bring up transition as well. Here are some topics to think about and discuss as the time comes to transition your child to adult-oriented care:

1. Is the timing right for transition?

2. What type of doctor should my child see?

3. What can my family and/or my child expect during transition?

4. What will be the same about adult-oriented care?

5. What will be different about adult-oriented care?

6. Will my child meet the new doctor before leaving this practice?

7. What do my child and my family need to know before moving to adult-oriented health care?
Parents: managers of an adult child’s health care
Many children with special health care needs may need assistance with care into adulthood. Below are tips on what you may do to prepare for medical transition:

- Include your child in her care as much as possible
- Have medical information that is readily accessible to your family and to doctors when you are not available
- Be aware of support groups for parents who care for their adult children to provide peer support and resources

Guardianship and alternatives
At age 18 your child becomes her own guardian. At this point some decisions will need to be made, either with your child or on behalf of your child, if she is unable to handle and manage her own care.

If you need to make these decisions, here are some questions to ask yourself:

- What are my main concerns for my child’s future?
- Are there financial, medical, or emotional concerns?
- What decisions will my child be able to make on her own?
- What decisions will my child need assistance in making?

Think about what you would like to secure for your child’s future. Below are some choices your family may consider.

Full guardianship provides full decision-making rights to an appointed guardian.

Partial guardianship provides rights to an appointed guardian in certain areas of decision making.

Durable power of attorney allows health care decisions to be made by appointed guardian. Power of attorney can cover other decisions.

A patient advocate acts for the individual receiving care. Most hospitals and providers have information on this choice. The appointed individual might be a family member or friend.

In a conservatorship, an appointed individual manages the child’s finances.

A representative payee manages the finances of a person with Supplemental Security Income or Social Security Disability Insurance benefits. There may be legal fees and actions that will need to go through the court system to appoint this person.

Health care representative: You may appoint a health care representative to make medical decisions in the event you become incapacitated. You may specify instructions to be followed by the health care representative, who must act in good faith and in your and your child’s best interest, consistent with the terms of the appointment.

You may have more options, depending on which state you live in. For more resources on guardianship please see the National Guardianship Association at www.guardianship.org.
Comprehensive care provides management of all aspects of your child’s care. Ideally, your doctor (or his backup) would be available 24 hours a day, seven days a week, 365 days a year to give your child the care she needs. Your doctor also would be in charge of preventive care, such as immunizations, growth and development assessments, and other appropriate screenings, as well as the monitoring of your child’s complex condition. Often doctors are a secondary advocate for your child. The doctor and office staff are able to provide information about private and public resources, including supplemental funding resources, education programs, waivers and other services for children with special needs. Comprehensive care allows for the doctor to manage your child’s care and incorporates you and your family into the decision-making process.

ONE FAMILY’S EXPERIENCE
We had just been to our child’s doctor the previous week, but the following weekend we noticed our child was running a fever. We were worried, and were not sure anyone from the doctor’s office would be available during the weekend. We called our doctor’s office and to our relief and surprise, we were immediately connected with an on-call doctor from the office. After assessing our child’s symptoms, he suggested we come in first thing the next morning. This brief phone call put us at ease and the next morning, our child was able to get the care he needed! We are so happy that the office has an on-call doctor available after hours so that we can get the information and care we need, when we need it!

“A medical home lets you know that you’re not the only one fighting to get something - that there’s somebody else there doing that with you - being your advocate, it’s very positive.”

“Comprehensive care does not stop when you are comfortable in knowing your child’s care is being managed. It is a process of providing information, referrals, treatments, emotional support and help with decision making.”

Parent-to-Parent Tip!

- Write down your questions ahead of time
- Record questions and answers in one place, such as in your child’s care plan
- Feel okay about needing advice in between appointments
- The best time to talk about comprehensive care is before an emergency so you are prepared with the next steps for your child’s care

Massachusetts Consortium for Children with Special Health Care Needs Resources Center: Medical Home: Compassionate Care www.neserve.org/maconsortium/rc_mh_compassion.htm
BUILDING HEALTH PARTNERSHIPS

It’s okay to ask questions
Family members often leave the doctor’s office irritated with themselves for not asking what they wanted to ask. Sometimes you may forget your question (bring a list!) or on other occasions you may not feel sure if it is okay to ask. Sometimes those questions that you hesitate to ask are the ones you are most concerned about or the ones that could address a need your doctor may not have noticed before. Doctors expect questions and want to work with you to get the answers you need to care for your child.

Below are some examples of questions that were not always asked:

- How much will it cost?
- Is there a less expensive alternative?
- What are the likely benefits or consequences for the proposed treatment?
- Can you repeat/write that down?

Get clear about communication
Ask your doctor how to best communicate about your child’s care. Is there a good time during the day to call? Does she prefer that you first talk to the office nurse if you have questions between appointments? Are there ever situations when the doctor would want you to call her at home? Does your doctor welcome communication via email?

Feel okay about needing advice between appointments
If you have a concern between appointments and feel that you need an answer from your child’s doctor, ask the office staff to have the doctor call you. Explain that you are more worried than usual – for reasons you may not be able to explain just yet – and that you would really appreciate talking directly to the doctor.

Reflecting
It is important that you get the information you need from your child’s appointment. Here are some questions to ask that could help you reflect on your child’s appointment:

- Were you able to discuss your most important questions or issues?
- If not, is this okay with you?
- Is there a plan for how you will get the information you need?
- Did you feel like a partner in your child’s health care team?
- Is there something you might want to do differently next time?

Parent-to-Parent Tip!
Remember that communication is the key to every relationship.
Take good notes, stay organized, and ask questions.
GETTING THE MOST OUT OF AN APPOINTMENT

Your partnership with your child’s doctor is primarily based on visits to the doctor’s office. This page will give you and your family some things to think about when working to get the most out of your child’s doctor appointments.

Scheduling
To cut down on waiting time, schedule your child’s visit with the doctor for the first appointment of the day or right after lunch. If you have questions or concerns that may take more time than usual, ask the office staff to schedule a longer appointment. Your doctor and staff will appreciate the advance notice and you will feel less frustrated about not having enough time.

Preparing
Think about what you would like to get out of the appointment ahead of time:

- Gather questions, research and/or reports that you especially want to discuss. You are the coordinator between your child’s care and the doctor.
- Share questions and concerns. Make a list of questions, concerns and other information you would like to share. Decide what on your list is the most important to you.
- It is okay to bring up things that don’t seem related to health but still matter to you and your child because communication is an important component of care.
- Ask the doctor if he would like a copy of your questions and concerns ahead of time. Emailing, faxing or dropping your list off before the appointment might give more time for the doctor to prepare more complete responses to your questions.

Participating
Share your list of questions and concerns at the start of the appointment. The doctor likely has his own list for what he needs to accomplish during the visit. Together you might need to decide what to discuss during this appointment and what to discuss later. Things can move so quickly during appointments that it can be hard to remember all that is said. Make sure to take notes, record answers to your questions and any other important information your doctor shares with you.

Updating
Tell your doctor about your child’s progress. A fun and memorable way to update your doctor is to send an occasional picture of your child with a note highlighting progress. Here are some examples:

- “Here is my daughter having fun at dinnertime.”
- “Through therapy, I learned how to encourage him to eat more table foods.”
- “She doesn’t gag and cry at the sight of food anymore and I’m not so worried about her growth.”

“Having access to longer appointment times for the complex child is not only beneficial for the family but also for the physician because they can give a quality visit without having to run behind the rest of the day.”  Ashley (Camerynn’s mom)
SECTION 4: COORDINATED CARE

Medical care often is fragmented and doctors do not always have the time or ability to communicate with each other about your child’s care. When you have a child with a complex medical condition, you will see any number of health and other service providers for different reasons. Coordinating all of your child’s multiple doctor visits, tests, procedures, and medical information is a daunting task and one of the most important in keeping your child healthy.  

Coordinated care is a process that links children and their families with the right services and resources in a coordinated effort to achieve good health. Care coordination for children with special health care needs is often complicated because of the child’s complex condition. Your child’s doctor and office team play a vital role in the process of care coordination; there are many effective models for providing it. Some offices have a nurse or other health professional whose job is to coordinate the care of patients’ in other offices, the doctor steps in to provide care coordination. While there are few doctor’s offices that are set up to perform care coordination services that link all of the other activities that impact your child’s health, there are ways that you and your doctor care partner to coordinate the care of your child. With well-coordinated information, everyone involved in your child’s care will be better adapted and more in control of helping you and your child meet the challenges associated with your child’s condition.

WHAT DOES CARE COORDINATION LOOK LIKE?

1. Care coordination is a team process made up of your family, doctors, and other service providers.
2. A plan of care is developed by you, your primary doctor, and any other doctors involved in your child’s care.
3. A central record with pertinent medical information is kept in the primary care office.
4. Your doctor’s office assists when referrals need to be made to other doctors or specialists and helps your family understand the referral recommendations during a follow-up visit.
5. Your doctor’s office coordinates your child’s care plan with other community agencies.

Some doctors employ the expertise of a care coordinator or specify a current employee to promote care coordination. A designated care coordinator manages your child’s information and ensures complete care. If your doctor does not have a care coordinator, perhaps mention the following benefits:
- Care coordinators can provide consistent staff contact for coordinating care
- They promote shared responsibility among the doctor, family and care coordinator
- They allow for direct communication between the doctor and family
- They engage the family in decision making and care planning

Parent-to-Parent Tip!
Stay organized! A really great tool for organizing care is an individual care plan. A care plan lists health problems, tests or procedures, doctors, health care services, equipment, treatment plan and expected outcomes. A flash drive is an inexpensive way to save this information and have it available at all times, without having to carry around heavy binders of paper work.

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7 Adapted from the AAP, Family Voices, MCHB, National Association of Child’s Hospitals and Related Institution for Child: “Comprehensive, coordinated, collaborative care” presentation.
8 Adapted from Donati, B; Passerello, T. & Stille, C. “Coordination of care in the medical home.” October 2003.
SECTION 4: COORDINATED CARE

BASIC STEPS OF COORDINATED CARE

Assessing and identifying needs: Activities performed by your doctor/care coordinator are based upon a complete assessment that includes a discussion with your family about the needs of your child. In your child’s doctor’s office, your care coordinator may use an assessment checklist to gather information to develop a plan of care for your child.

Developing a plan of care: After identifying your family’s needs, a plan of care is developed with your family’s goals for your child in mind. The care coordinator may clarify which action steps will be addressed by your family and which will be addressed by the care coordinator.

Implementation: The plan is implemented and actions are taken to work toward the goals you and your doctor discussed. The care coordinator organizes and assists the family with resources, referrals, coordination of care with specialty physicians, schools, and other agencies.

Evaluation: Your care coordinator will take time to look over the plan of care with you and your family to make changes as new needs are identified.

WHAT IS A CARE PLAN?
A care plan is written information about how to best care for your child’s needs. It is something you and your child’s doctor can develop together. A care plan may include:

- What medicines your child takes and when
- What foods your child should avoid
- What to do for your child in an emergency
- Goals for your child’s health and steps to achieve them

Your child, family, doctor and other care providers will benefit from having a clear, written care plan. The care plan care be written on paper, saved to a data flash drive, or if possible, be web based. There are multiple purposes of the care plans. Care plans are an available source of information for parents to provide to the medical, educational, and other care teams, and a quick reference with child-specific information for a medical emergency. Care plans are an action plan that the entire care team, including the family, develop to use in prioritizing, assigning tasks, and implementing and assessing care. Having a written care plan helps you share important information with others who care for your child. This group may include doctors, nurses, therapists, emergency medics, teachers, child-care providers, friends, and neighbors.

Parent-to-Parent Tip!
Encourage your doctor to have internet access in their office and use it for care coordination. The internet can serve as a tool for ease of access to your child’s care plan, treatment options, and other medical information. Many doctor’s offices have computers, and having information readily accessible is great!
WHAT IS COMPASSIONATE CARE?

Compassionate care is concern for the well being of the child and family expressed in verbal and nonverbal interactions. Efforts are made to understand the feelings and perspectives of the child, as well as the family.

Key components of compassionate care:

1. Your child's doctor treats you, your family, and child with respect and care. You notice other families are treated with the same respect and care.
2. Your child's doctor takes time to listen to you and your family's concerns.
3. The doctor’s office staff takes the time to ask you and your family about any special needs or concerns and helps your family meet those needs.
4. The doctor’s office staff work with community groups to help meet your child’s specific needs.
5. Your child's doctor takes the time to get to know your child's special health care needs.
6. Your child's doctor recognizes the impact your child's chronic health needs have on you and your family.
7. Changes in ways the office staff cares for your child are made in response to your and your family's unique needs and challenges.
8. Office staff help groups in your community learn about the needs and concerns of children with special health care needs and their families.  

Examples of compassionate care: When you come in with your child for an appointment, the doctor is interested in your child, his progress, and any updates. The doctor is attentive to your questions and expresses concern for your child in an understanding manner.

“We have a care plan that is always with us, and the hospital and clinic are aware of the special needs and openly give Miriam that much needed “extra” time and gentleness. All these little changes are making a significant difference not only for Miriam, but for our family.”

Jennifer (Miriam’s mom)

10 From the American Academy of Pediatrics, Family Voices, Maternal and Child Health, Bureau National Association of Children’s Hospitals and Related Institutions, and Shiner's Hospitals for Children: Common Elements Medical Home: www.occrra.org/inf-todd/module/physical_health/Medical_Home_Common_Elements_AdaptedFinal.ppt
WHAT IS CULTURALLY EFFECTIVE CARE?

“Culturally effective care” starts with letting your doctor know that you have certain cultural beliefs or requests you would like him to observe or consider when treating your child. When the doctor is aware of your requests, he works to make you and your family more comfortable with the care of your child.

What is culture?
Culture is a way of life, values, beliefs, behaviors, experiences, tastes, preferences, languages, and customs.

What is diversity?
Diversity is generally defined as differences among people with respect to age, class, ethnicity, gender, physical and mental ability, race, sexual orientation, spiritual practice, and public assistance status. Your doctor demonstrates culturally effective care when he acknowledges, understands, and accepts these differences.

Language and communication
Spoken and written language is our primary form of communication. If you find it difficult to communicate with your doctor, it may be necessary to get the assistance of a trained or certified bilingual translator to help you understand what information your doctor is telling you and your family.

Listed below are some points for your family to keep in mind when seeking to increase cultural awareness in the health care setting.

- Ask questions when you do not understand a word or phrase
- Ask a person to slow down when giving you treatment options
- Bring another person who can help explain things to you
- Ask for visual aids or physical gestures to help you
- Ask the person to rephrase the question or instructions if you do not understand the words he is using

Parent-to-Parent Tip!
It is important for your child’s health care team to be aware of values, beliefs, and practices that are common in your community and important to your family. Although your health care team may be familiar with some of the general health care practices or norms for your culture, it is still very important for you to communicate your personal feelings about how you would like to participate in the health care process.
VALUES AND ATTITUDES

“Family” can be defined in many ways. Share how you define your family so your child’s doctors will understand the role each member may have in your child’s care plan. Let the staff know:

- How important medical decisions are made in your family
- Who needs to hear the information from the doctor
- Who in your family provides you with the most support

“Health and well being” mean different things to different people. Understanding and sharing what this means for you and your family can help a doctor in developing a care plan. Let the doctor know things such as:

- How you handle your emotions in public and in private
- What you consider ‘healthy’ for your child with special health care needs
- What foods you and your child eat to promote good health

Religion and other beliefs influence how families respond to illness, disease, disability, and death. Awareness of your belief system will give your child’s doctors a better understanding of what is important to your family. The doctor and office staff may wish to know:

- How your family typically celebrates the birth of a child
- What activities your community offers for families of children with special health care needs
- What type of response you would get from the community about your child with special health care needs
- What the cultural expectations are for self-help skills such as dressing and feeding

Families bring unique cultural perspectives to service-delivery systems. Each child and family has strength and resilience as well as needs and challenges. It is essential that integrated systems of care within states and communities understand and respond effectively to the extent to which children and families are satisfied with the services and supports they receive. The National Center for Cultural Competence (NCCC) presents a series of articles to increase awareness, knowledge and skills in this important aspect of care.

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12 National Center for Cultural Competence (NCCC) www11.georgetown.edu/research/gucchd/nccc/
INTERPRETATION AND TRANSLATION SERVICES

Most large hospitals and outpatient clinics are able to provide an interpreter for common languages if you request one. Be sure to let the office know an interpreter is needed when scheduling your child’s appointment. Title VI of the Civil Rights Act of 1964 requires health care providers to overcome language barriers to care. This law applies to any provider or hospital receiving federal funds, including providers who treat Medicaid or Medicare patients. The Americans with Disabilities Act (ADA) requires that service providers contract and pay for interpreters to provide medical interpretation for the deaf and hearing impaired. Asking someone to interpret medical information after the patient’s appointment is not acceptable under the ADA.

Many organizations provide language translation services which may include translating written documents, interpreting speech, or both. Some translation agencies charge a fee for their services. Interpreters also may be found through community or government agencies, colleges or universities, or translation-related businesses. Families also may find health care and other assistance offered in their native language through community organizations.

To help you choose your translator, keep these four tips in mind:

**Find a translator who understands the primary language spoken at home:** Medical words and phrases can be confusing. If your translator is new to the language or has just learned how to speak English, she might miss important details. A confused translator can give incorrect information and this can be dangerous.

**Pick a translator who is at least 18 years old:** Some things you will talk about with your doctor can be serious or complicated. Your translator must be mature enough to handle any topics that come up. Your translator also must understand everything and be able to translate the doctor’s advice correctly so you can understand it.

**Choose someone you trust:** Office visits are often routine, however, sometimes you may have to answer tough questions or talk about personal topics. Make sure you are comfortable with including your translator in these private conversations.

**Think carefully before bringing a family member:** Many people bring family members as translators. If your family member is an adult, this may be an easy decision. But if you have a younger family member, remember that some topics you talk about with your doctor may be sensitive.

If the translator you bring doesn’t fully understand the topic or information given, ask your doctor to write down the important information and have it translated later. Use these tips to make a list of the people you know who may be right for the job. If you don’t have a friend or family member who meets your needs, check with your doctor’s office, and they may be able to assist you.

**Parent-to-Parent Tip!**

Good communication is an important key to better health outcomes. Sometimes your child’s doctor may come from a different ethnic or cultural background than your family. A language difference may make it difficult for you to understand one another, but remember that what you have in common is a desire to help your child.
WHAT IS FAMILY-CENTERED CARE?

Family-centered care assures the health and well-being of children and their families through a respectful partnership between the family and doctor. It honors the strengths, cultures, traditions and expertise that everyone brings to the relationship. Family-centered care is the standard of practice that results in high-quality services.¹³

Key components of family-centered care

- Your family is essential to your child’s care and is a constant in your child’s life
- Your doctor acknowledges key family members
- Your doctor asks your family about values
- Decision making is shared

Example of family-centered care: Upon bringing your child into the doctor’s office for a concern about your child’s health, the doctor asks you what you have observed, what you think could be the cause of your concern and what you think could be a possible course of action to take. By communicating with you, the doctor is utilizing your knowledge and is able to provide quality treatment and care for your child.

How to make family-centered care work:

**Keep the lines of communication open and honest**
- *Share your thoughts and ideas about decision making for your child*

**Be aware of office patient procedures**
- *Request to schedule a longer appointment to go over your list of concerns, updates, and questions*

**Participate on the clinic’s advisory board**
- *This is a great chance to be involved in discussions, get to know other families with similar situations, and learn from them!*

**Complete patient- and family-satisfaction surveys**
- *Giving and receiving feedback is important to your doctor and office staff so they may continue to provide you and your child with quality, family-centered care!*

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¹³ Adapted from American Academy of Pediatrics, Medical Home Initiatives Every Child Deserves a Medical Home Training Manual, Appendix D: Family Centered Care Fact Sheet

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**A Definition of Family**

“Families are big, small, extended, nuclear, and multi-generational with one parent, two parents, and grandparents. We live under one roof or many. A family can be as temporary as a few weeks or as permanent as forever. We become part of a family by birth, adoption, marriage or from a desire of mutual support. A family is culture unto itself, with different values and unique ways of realizing its dreams together, our families become the sources of our rich cultural heritage and spiritual diversity...Our families create neighborhoods, communities, states, and a nation”.

Polly Arango, Family Voices
CORE PRINCIPLES OF FAMILY-CENTERED CARE

This set of values describes the family-centered care that a medical home practice provides. Use this list as a conversation starter with your doctor.

1. **Respecting** each child and his family.
2. **Honoring** racial, ethnic, and cultural diversity and its affect on the family’s experiences and views on care.
3. **Recognizing** and building on the strengths of each child and family member, even in difficult and challenging situations.
4. **Supporting** the choices of the family about approaches to care and decision making.
5. **Ensuring** flexibility in office policies, procedures, doctor practices, and services tailored to the needs, beliefs, and cultural values of each child and family.
6. **Sharing** honest and unbiased information with families on an ongoing basis and in ways they find useful and assuring.
7. **Providing** and/or ensuring formal (example: doctor-to-patient relationship and referrals to other services) and informal support (example: family-to-family support) for the child and family during pregnancy, childbirth, infancy, childhood, adolescence, and young adulthood.
8. **Collaborating** with families at all levels of the medical care of the individual child and in professional education, policy making, and program development.
9. **Empowering** each child and family to discover their own strengths, build confidence, and make choices and decisions about their health.
10. **Participating** in decision making and celebrating achievements.

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**Parent-to-Parent Tip!**

Family-centered care starts with the family and doctor communicating and building a successful partnership. As a parent, you will prepare, organize, schedule, and advocate for your child. Never forget that you are an expert on your child and your opinion is important to the health and happiness of your child and family.

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14 Core principles of family-centered care. American Academy of Pediatrics. [http://aappolicy.aappublications.org/cgi/content/full/pediatrics;112/3/691#SEC3](http://aappolicy.aappublications.org/cgi/content/full/pediatrics;112/3/691#SEC3)
Advocacy means educating others about a need, small or large, in an effort to help meet that need. It can refer to community issues, like making a plea for a stop sign at a dangerous intersection or improving the accessibility of a playground so that all children can enjoy the space. Advocacy also can be as simple as asking for written instructions to compliment what you are being told, whether you are at a pharmacy or at the car repair shop. For families of children with special health care needs, advocacy becomes an everyday effort to improve the quality of life for their children and others like them.

Advocacy happens at many levels, from a conversation in the doctor's office, up to a visit with your legislators, but any effort you make is a step in the right direction. You can write letters, send emails, make phone calls, or attend support group meetings. The more you do to voice your needs, the more attention your issue will receive.

LEVELS OF ADVOCACY

The most basic level of advocacy affects your own child. You may need to advocate when:

- Your child needs to see a doctor after hours or when no appointments are available
- You would like your insurance to make an exception on a denied payment
- You wish to see your doctor’s office take on the concept of the medical home

When you are advocating for your child’s medical home, take Angela’s letter (Page 30) with you to break the ice. It will give your doctor a full picture of what you would like to see happen for your child’s care.

The next level of advocacy could also affect other children and their families. You are advocating at the policy level when:

- You request your doctor’s office to hire a care coordinator or specify a current employee to promote care coordination. Share how this person’s time will assist families in managing the complex care of their children.

- You ask your insurance company to update their policy to cover the purchase of sterile water for use in a ventilator, which is critical for a child needing humidification. Again, the policy change you are asking for will help others in a similar situation.

The next level of advocacy involves asking for a system change. A change in a system might be involved when:

- You go to state agencies and government entities to bring attention to the financial burden of medical care for families who have children with special health care needs. You explain how care coordination in your doctor’s office could save costs for families and the health care system as a whole.

- You explain your needs (which also represent the needs of many others) to a state agency that influences how funding is dispersed. Reimbursement from insurance providers on medical foods and formulas or care coordination are issues that families face every time they visit the doctor’s office.
SECTION 8: ADVOCATING FOR YOUR CHILD

BASIC GUIDELINES FOR ADVOCACY

1. **Choose and learn about your issue(s)**
   Identify the issue you most need to focus on and learn as much as you can about it. Talk to other parents, find information through your local or hospital library, the internet, health newsletters, and local parent support groups. Locate existing advocacy groups and find out what has already been done on your issue.

2. **Identify decision makers**
   Find out who has authority to make decisions, whether it's a supervisor, a program director, a chief administrator, or your local legislator. These are the people to whom you must appeal for changes in the system. Your local parent support groups can help you identify the appropriate decision makers.

3. **Learn how to navigate the system**
   Navigating the system can mean anything from learning the right vocabulary when calling about an insurance statement, to asking for the same office administrator when you need help at your doctor's office. It also can mean learning how the legislative process works so you can influence potential legislation.

4. **Communicate your views**
   When making your request, be brief, to the point, and polite. Identify yourself, your issue, and give one or more reasons for your position. Sometimes it helps to write this down first, to make sure you can summarize what you want and why. And remember, change takes time, so you may need to be patiently persistent.

5. **Thank those who have helped**
   Always remember to thank those who have taken the time to listen to your needs. A simple written note of thanks can make a difference when the next person comes to ask for help.
Dear Parents,

I was sent into a tail spin when my third child was diagnosed with congenital adrenal hyperplasia. My other two children were born healthy and I had no experience with any medical disorder. I didn’t even know which pediatrician to contact as a primary care doctor as my other children were under the care of a family doctor.

Luckily, I found a fantastic pediatrician who practiced using the medical home model. When my daughter was born, I was less focused on the medical home model and more focused on her medical condition. It didn’t take long before I realized what a wonderful concept the medical home model is.

Through our medical home, my daughter was assigned a care coordinator. This nurse is our lifeline to medical care. She met with us during the first appointment to explain how the office did things differently. First, she showed us the private waiting room that we could utilize so that our daughter didn’t have to be exposed to germs. Next, she told us how to directly contact her and what to do in her absence. She went on to discuss what appointments would entail and how to manage the paper work involved. Prior to each visit she contacts me to find out if my daughter has seen a specialist, had an emergency room visit, or if I have any special concerns for the doctor. This is all coordinated prior to the appointment so that we can have all the facts in front of us when we see the doctor. This pre-visit coordination helps tremendously with information flow. The patient care coordinator also contacts specialists to follow up on their reports regarding my daughter’s care. We don’t have a medical folder; instead, we have a medical binder and any medical paperwork regarding my daughter’s care from inside or outside of the pediatrician’s office is in that binder. Paperwork from hospital visits and other specialist visits are all included so that the pediatrician has a total picture of my daughter’s care.

Our primary doctor is fantastic. Upon the first visit he introduced us to a secondary doctor that he wanted us to see in his absence. Although the office has seven doctors, he wanted to have her care managed by only two doctors who would understand everything about her condition. Together, both doctors spent ample time with my daughter on her initial visit. They also worked with me to discuss a medical plan for her. We discussed when to call them, what to do after hours, medicine dosages, and how to handle emergency room visits. I left the office feeling as if both doctors truly had a vested interest in seeing my daughter thrive.

Under the medical home model the office also holds quarterly parent advisory group meetings. This is a time for staff and families of special needs children to come together to educate and share information with each other. The staff contacts outside organizations to come speak to parents on a variety of topics. We have had many guest speakers, ranging from camps that provide for children with special needs to government agencies. Occasionally, parents will request certain topics to be addressed and the staff will coordinate a workshop. Recently, I asked for a refresher training on performing injections. The staff not only provided this to my entire extended family, but they also invited other families who could benefit from this training. These meetings also give parents the time to collaborate, share, and support each other on their journey to providing the best possible care for their child.

While my daughter’s pediatrician may be the exception, I would encourage you to find an office that is practicing the medical home concept. This office should be the central hub of your child’s care. They should collaborate with your family and your child’s specialists. They should support you in finding resources, second opinions, and advanced specialists in your area. And most importantly, your pediatrician’s office should be connected with and supportive of your medical treatment endeavors.

Best wishes,
Angela (Mother of a daughter with congenital adrenal hyperplasia)
<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Accessible Care</td>
<td>Accessible care encourages your doctor and his or her office to be available in terms of transportation, insurance acceptance, and disability access.</td>
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<tr>
<td>Care Coordinator</td>
<td>A person who assists in linking children and their families with appropriate services and resources in a coordinated effort to achieve good health.</td>
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<tr>
<td>Comprehensive Care</td>
<td>Comprehensive care ensures your doctor will manage all aspects of your child’s care and be available when needed.</td>
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<tr>
<td>Coordinated Care</td>
<td>Coordinated care involves the family, doctors, teachers, social service professionals and other caregivers to ensure access to appropriate services and planning.</td>
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<tr>
<td>Continuous Care</td>
<td>Continuous care encourages doctors to know about their patients’ other medical visits, procedures, and medications so they can provide the best care possible.</td>
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<tr>
<td>Culturally Effective Care</td>
<td>Culturally effective care encourages communication between the doctor and patient to share certain cultural beliefs and ensure they are observed or considered during treatment.</td>
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<tr>
<td>Genetic Condition</td>
<td>An illness caused by abnormalities in genes or chromosomes. Most disorders are quite rare and affect one person in every several thousands or millions.</td>
</tr>
<tr>
<td>Health care Professionals</td>
<td>Workers who have special health care skills, such as doctors and nurses.</td>
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<tr>
<td>Legislator</td>
<td>A person who writes and passes laws, especially someone who is a member of a legislature. Legislators are usually politicians and are often elected by a population.</td>
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<tr>
<td>Local Health Department</td>
<td>This office is a local resource to connect you to health-related services available to you in your community.</td>
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<tr>
<td>Occupational Therapist</td>
<td>Occupational therapy is a health profession with the goal of helping people achieve independence, meaning, and satisfaction in all aspects of their lives. The occupational therapist provides the patient with skills for the job of living - those necessary to function in the community or in the client's environment.</td>
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<tr>
<td>Pediatric</td>
<td>The branch of medicine dealing with the care of children.</td>
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<tr>
<td>Primary Care</td>
<td>General or basic health care. Traditionally provided by a pediatrician, internist, or family practitioner.</td>
</tr>
<tr>
<td>Primary Care Provider</td>
<td>General or basic health care professional who is involved with overall care, as well as coordination of care when specialists are involved.</td>
</tr>
<tr>
<td>Specialty Care</td>
<td>A medical practitioner whose practice is limited to a particular class of patients or of diseases or of technique</td>
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NATIONAL RESOURCES

National Center of Medical Home Initiatives for Children with Special Needs
www.medicalhomeinfo.org/
The National Center works to ensure that children and youth with special health care needs have access to a medical home. Through the National Center, physicians, parents, administrators, and other healthcare professionals have access to educational, resource, and advocacy materials, guidelines for care, evaluation tools, and technical assistance.

Center for Medical Home Improvement (CMHI)
www.medicalhomeimprovement.org/
CMHI is committed to supporting practices, patients, and families to develop strong and sustainable medical homes. CMHI aims to link you to the most helpful resources for medical home development. CMHI has developed a Parent Partners Guide to help parents (and physicians) better understand their critical role as parent partners.

Family Voices
www.familyvoices.org
Family Voices aims to achieve family-centered care for all children and youth with special health care needs and/or disabilities.

Institute for Family-Centered Care
www.familycenteredcare.org/
The Institute for Family-Centered Care provides leadership to advance the understanding and practice of patient- and family-centered care in hospitals and other health settings.

Medical Home Portal
www.medicalhomeportal.org/
The Medical Home Portal aims to provide ready access to reliable and useful information for professionals and families to help them care and advocate for children and youth with special health care needs, as partners in the medical home model.

National Center for Family/Professional Partnerships
www.fv-ncfpp.org
The National Center for Family/Professional Partnerships (NCFPP) promotes families as partners in the decision-making of healthcare for children and youth with special health care needs (CYSHCN) at all levels of care.

National Initiative for Children’s Healthcare Quality (NICHQ)
www.nichq.org/
Provides information about the medical home as it pertains to children and youth with special health care needs.
OTHER RESOURCES & TOOLKITS

A new way… A better way: The Medical Home Partnership
http://fha.dhmh.maryland.gov/genetics/docs/NESSERVE_Maryland_Families.pdf?Mobile=1
This brochure describes what a medical home is and provides information about what you should expect from a medical home.

Building Early Intervention Partnerships with your child’s Doctor: Tips from and for Parents
This booklet gives suggestions and ideas for getting started with a new doctor and for improving a partnership that you already have.

Does your child have a medical home?
www.medicalhome.org/4Download/medicalhomebro.pdf
Provides information about what a medical home is, benefits from having a medical home, and a checklist to determine whether or not your child has a medical home.

Extraordinary care: Improving your Medical Home
The learning guide is designed to assist families and others in their education about the basics of “medical home.” It also suggests specific activities that help to strengthen their medical home or advocate for stronger primary care services.

Families partnering with providers
https://org2.democracyinaction.org/o/6739/images/PartneringWithProviders.pdf
This brochure provides tips to help families build effective partnerships with their child’s doctors.

Family-Centered Care Self-Assessment Tool - Family Tool
https://org2.democracyinaction.org/o/6739/images/fcca_FamilyTool.pdf
Families, both individually and within family support and advocacy organizations, can use this tool to increase awareness and knowledge of the specifics of family-centered care.

Family-Centered Care Self-Assessment Tool - User’s Guide
The user’s guide provides a detailed set of steps to use the family and provider tools for a full assessment of a practice setting or family. In addition, it may be used as an educational tool to build awareness and knowledge of family-centered care for families, youth, doctors, health plans, and policy-makers.

What is a medical home? And what does it mean for you and your child?
http://internet.dscc.uic.edu/forms/medicalhome/4016.pdf
This brochure provides information about the components of a medical home.