FAMILY-CENTERED CARE SELF-ASSESSMENT TOOL

Developed by
FAMILY VOICES®
...keeping families at the center of children’s health care

with funding from Maternal and Child Health Bureau (MCHB), Health Resources and Services Administration, U.S. Department of Health and Human Services under Cooperative Agreement #U40MC00149-09-00

OCTOBER 2008
INTRODUCTION

Health care visits for children, youth and their families can be more than getting shots, having ears examined or treating the physical symptoms of an illness. Each visit is an opportunity for families, youth and health care providers to partner to assure quality health care for the child and to support the family’s needs in raising their child. This enhanced aspect of the family and health care provider relationship is called family-centered care. The foundation of family-centered care is the partnership between families and professionals. Key to this partnership are the following:

• Families and professionals work together in the best interest of the child and the family.
• As the child grows, s/he assumes a partnership role.
• There is mutual respect for the skills and expertise each partner brings to the relationship.
• Trust is fundamental.
• Communication and information sharing are open and objective.
• Participants make decisions together.
• There is a willingness to negotiate.

Within that framework, ten components of family-centered care have been identified. (National Center for Family-Centered Care (1989); Bishop, Woll and Arango (1993)) Family-centered care accomplishes the following:

1. Acknowledges the family as the constant in a child’s life.
2. Builds on family strengths.
3. Supports the child in learning about and participating in his/her care and decision-making.
4. Honors cultural diversity and family traditions.
5. Recognizes the importance of community-based services.
6. Promotes an individual and developmental approach.
7. Encourages family-to-family and peer support.
8. Supports youth as they transition to adulthood.
9. Develops policies, practices, and systems that are family-friendly and family-centered in all settings.
10. Celebrates successes.

PURPOSE OF THE FAMILY-CENTERED CARE SELF-ASSESSMENT TOOL

Family-centered care is a key aspect of quality in health care for children, youth and their families. This tool is designed to:

1. Increase outpatient health care settings’ and families’ awareness about the implementation of family-centered care and,
2. Provide an organized way for health care settings to assess current areas of strength and identify areas for growth, plan future efforts and to track progress.

This tool is not designed to provide a score but is meant as an opportunity for reflection and quality improvement activities related to family-centered care within outpatient health care practices. It can also be used by families to assess their own skills and strengths, the care their children and youth receive, and to engage in discussions within health care settings and with policy makers in organizations, health plans and community and state agencies about ways to improve health care services and supports. The tool is intended to assess care for all children and youth and also has some questions that are specific to the needs of children and youth with special health care needs and their families. Questions on the tool address the ten components of family-centered care and the key aspects of family/youth/provider partnerships.
DESCRIPTION OF THE FAMILY-CENTERED CARE SELF-ASSESSMENT TOOL

The tool is divided into three major sections: 1) Family/Provider Partnership, 2) Care Setting Practices and Policies and 3) Community Systems of Services and Supports. Within each major section, there are several subtopics that address family-centered care.

Sections and Subtopics of the Family-Centered Care Self-Assessment Tool

Family/Provider Partnership
- The decision-making team
- Supporting the family as the constant in the child’s life
- Family-to-family and peer support
- Supporting transition to adulthood
- Sharing successes

Care Setting Practices and Policies
- Giving a diagnosis
- Ongoing care and support
- Addressing child/youth development
- Access to records
- Appointment schedules
- Feedback on care setting practices
- Care setting policies to support family-centered care
- Addressing culture and language in care

Community Systems of Services and Supports
- Information and referral and community-based services
- Community systems integration and care coordination

Each section contains a series of questions that ask about concrete actions that reflect family-centered care. Each section is coded with numbers that indicate which of the ten elements of family-centered care it addresses.

DEFINITIONS OF TERMS

Alternative Healing is the use of herbs, aromatherapy, acupuncture, massage and other remedies that are not considered part of conventional (Western) healthcare treatments.
http://www.aap.org/healthtopics/complementarymedicine.cfm

Care Setting is the physical location where a family and child/youth receives outpatient health services. These services can be provided by a pediatrician, other physician, physician’s assistant, nurse, social worker, care coordinator, or any other staff person at this setting.

Children and Youth with Special Health Care Needs have or are at increased risk for chronic physical, developmental, behavioral, or emotional conditions that require health and related services of a type or amount beyond that required by children generally.

Community-based Services are local, non-medical services that help children and families accomplish daily activities. These services can range from school, childcare, after-school activities, family and peer support and advocacy groups, early intervention or Head Start (a program that helps families learn about and promote the development of their young children).

Consent is approval for healthcare decisions. Individual must have legal authority to provide consent. Generally, the parents or legal guardians of minor children (younger than 18 years old) provide consent. For individuals 18 and older who cannot make their own decisions, a legal guardian or proxy can provide consent.

Cultural Competence requires organizations to have a defined set of values and principles and demonstrate behaviors, attitudes, policies and structures that enable them to work effectively cross-culturally. They should:
- Value diversity,
- Conduct self-assessment,
- Manage the dynamics of difference,
- Acquire and institutionalize cultural knowledge and
- Adapt to diversity and the cultural contexts of the communities they serve.
Organizations should incorporate the above in all aspects of policymaking, administration, practice, service delivery and involve systematically consumers, key stakeholders and communities. Cultural competence is a developmental process that evolves over an extended period. Both individuals and organizations are at various levels of awareness, knowledge and skills along the cultural competence continuum.

(Adapted from Cross et al., 1989) http://www11.georgetown.edu/research/gucchd/nccc/foundations/frameworks.html

Family—Families are big, small, extended, nuclear, 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, as permanent as forever. We become part of a family by birth, adoption, marriage, or from a desire for mutual support. As family members, we nurture, protect, and influence each other. Families are dynamic and are cultures unto themselves, with different values and unique ways of realizing dreams. Together, our families become the source of our rich cultural heritage and spiritual diversity. Each family has strengths and qualities that flow from individual members and from the family as a unit. Our families create neighborhoods, communities, states, and nations.

Developed and adopted by the New Mexico Legislative Young Children’s Continuum and New Mexico Coalition for Children, June 1990.

Family Supports are services that strengthen and support the family’s role as caregiver and decision-maker on behalf of their children.

Family-to-Family and Peer Supports are ways to bring together families, youth and others who share similar life situations so they can share their knowledge, concerns, and experiences with each other.

HIPAA (The Health Insurance Portability and Accountability Act). This is a federal law that protects the privacy of personal health information. You must give permission before personal health information can be shared with:

- Health providers,
- Hospitals,
- Insurance companies, state and federal agencies,
- Schools,
- Employers, or
- Anyone else.

You also have the right to read your medical record and make corrections. For more information visit: http://www.hhs.gov/ocr/hipaa

Linguistic Competence is the capacity of an organization and its personnel to communicate effectively, and convey information in a manner that is easily understood by diverse audiences including persons of limited English proficiency, those who have low literacy skills or are not literate, and individuals with disabilities. Linguistic competency requires organizational and provider capacity to respond effectively to the health literacy needs of populations served. The organization must have policy, structures, practices, procedures and dedicated resources to support this capacity.


Provider—in the context of this tool is any person that provides services within the care setting.

Transition to Adulthood—The period of time during which families, youth and providers plan for and develop the process to assure that youth will:

1. Be able to successfully manage all aspects of their healthcare and be prepared to take on adult responsibilities, and ultimately, live as independently as possible and
2. Continue to receive high quality healthcare services.
**Directions:** Please answer each question by choosing only one answer. It may be difficult to rate some items; just answer each question based on your knowledge, experiences and opinions. There are no right or wrong answers.

**Note:** In each question the term “provider” refers to the health care professionals and other staff within your health care setting.

## THE FAMILY/PROVIDER PARTNERSHIP

### The Decision-Making Team

<table>
<thead>
<tr>
<th></th>
<th>FCC Components: 1, 2, 4, 9</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Do you and your staff:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>A. Partner with families to help define their role in their child’s care?</td>
<td>Never</td>
</tr>
<tr>
<td>B. Honor families’ requests for others (extended family, community elders, faith leaders or traditional healers that are designated by the family) to participate in the process that leads to decisions about care?</td>
<td>Never</td>
</tr>
<tr>
<td>C. Help families advocate for services and work to improve systems of care, if they so choose?</td>
<td>Never</td>
</tr>
<tr>
<td>D. Act to support each family’s chosen role?</td>
<td>Never</td>
</tr>
<tr>
<td>2. Do you and your staff:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>A. Work in partnership with families/youth to make health care decisions?</td>
<td>Never</td>
</tr>
<tr>
<td>B. Talk about the range of treatment and care choices for the child/youth?</td>
<td>Never</td>
</tr>
<tr>
<td>C. Discuss which treatment and care choices would be best for the family and child/youth?</td>
<td>Never</td>
</tr>
<tr>
<td>D. Make sure the family and child/youth understand the range of treatment and care choices?</td>
<td>Never</td>
</tr>
<tr>
<td>E. When deciding on treatment options, do you and your staff work with the family and child/youth to decide what the desired outcomes are (e.g., improved health status, better school attendance, less pain, or better involvement with social or sports activities)?</td>
<td>Never</td>
</tr>
<tr>
<td>3. Do choices of diagnostic and treatment approaches take into account:</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>A. Family/child youth preferences for site of care, type of provider (gender, language spoken, etc.)?</td>
<td>Never</td>
</tr>
<tr>
<td>B. Child/youth’s ability to tolerate the procedure?</td>
<td>Never</td>
</tr>
<tr>
<td>C. Any follow up medical treatment the child/youth will need?</td>
<td>Never</td>
</tr>
<tr>
<td>D. Family insurance status and economic situation?</td>
<td>Never</td>
</tr>
<tr>
<td>E. Family and child/youth work and school schedules?</td>
<td>Never</td>
</tr>
</tbody>
</table>

| 4. Do you and your staff make sure all the family’s and child/youth’s questions have been answered before they leave the office? | Never | Some of the Time | Most of the Time | Always |
The Decision-Making Team Continued

5. Do you and your staff make sure the family and child/youth feel comfortable enough to let you know if they disagree with medical advice and recommendations for treatment and care?  

   ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

6. Is there a respectful negotiation process to resolve any disagreements about a child/youth’s treatment and care?  

   ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

7. Does your partnership with families change over time as their experiences, knowledge and skills change?  

   ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

Supporting the Family as the Constant in the Child’s Life  

FCC Components: 1, 2, 4, 6

1. Do you and your staff ask about:

   A. The family’s well-being (adults and other children) and their needs for support?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

   B. The family’s support network and the role of faith/religion or other cultural supports?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

   C. The family’s concerns and any stresses or successes they may experience as a caregiver?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

   D. Depression, domestic violence, substance abuse, housing or food insecurity?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

2. Do you and your staff offer families:

   A. Information about health and wellness appropriate to child/youth’s developmental stage? (This includes information about child development, mental health, healthy weight and nutrition, physical activity, sexual development and sexuality, safety/injury prevention, and oral health.)  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

   B. Opportunities to become more knowledgeable about promoting their child’s healthy development, such as written information, classes, or connect them with other opportunities in the community?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

Family-to-Family and Peer Support  

FCC Components: 2, 4, 7, 8

1. Do you and your staff:

   A. Have a process to identify the strengths of families within the practice that they can share with other families?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

   B. Actively assist in linking families to other families who share similar life situations, for example, new mothers, children with special health care needs and disabilities, single parent families, grandparents raising grandchildren, etc?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

   C. Link families in a way that that reflects the cultural and language preferences of each family?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

   D. Help families/youth make the first connection to support?  

      ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always
E. Offer peer mentoring/support opportunities for families/caregivers to be supported in their changing roles from decision-makers to supporting their children/youth in making decisions?

F. Offer peer mentoring/support opportunities to help children/youth learn how to be active in decision-making about their own care?

### Supporting Transition to Adulthood

**FCC Components: 1, 3, 4, 6, 7, 8**

#### 1. Do you and your staff:

<table>
<thead>
<tr>
<th>A. Ask families and children (from an early age)/youth about their vision for the future? (For example, take time to discuss the family’s and child’s hopes and dreams about education, social relationships, development of independent living skills, meaningful work, and health care.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>B. Connect families and youth with information and resources to support achieving their vision in ways that respect their cultural beliefs about roles and expectations in adulthood?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>C. Offer opportunities for families and youth to meet with older youth and young adults as role models for achieving future goals?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
</tbody>
</table>

#### 2. In preparation for transition, do you and your staff:

<table>
<thead>
<tr>
<th>A. Help youth learn about: managing their health, dealing with the logistics of care settings, insurance and decision-making?</th>
</tr>
</thead>
<tbody>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>B. Work with the family to assure that the youth knows about any diagnoses, current treatments and can discuss those with the care provider?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>C. Work with the family and youth to develop a formal healthcare transition plan by age 14 that documents provider, family and youth roles?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>D. Have a formal mechanism to identify adult health care providers for youth in transition?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>E. Have a process to share information with adult care providers including: current care plans, transition plans, medical records, key health issues and current family and youth roles in managing care?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>F. Help youth develop a portable medical record to support their interactions with new providers?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>G. Work with adult providers so they are involved in a youth’s care during transition?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
<tr>
<td>H. Help families plan for needed legal solutions and supports if their child/youth will be unable to independently manage his/her care?</td>
</tr>
<tr>
<td>○ Never ○ Some of the Time ○ Most of the Time ○ Always</td>
</tr>
</tbody>
</table>
### Sharing Successes of the Family/Provider Partnership

**FCC Component: 10**

1. **Do you and your staff take time to document and share successes with families?**
   - Never  Some of the Time  Most of the Time  Always

2. **Do you and your staff take time to document and share successes with each other?**
   - Never  Some of the Time  Most of the Time  Always

### CARE SETTING PRACTICES AND POLICIES

#### Giving a Diagnosis

**FCC Components: 1, 2, 6**

1. **Do you and your staff:**
   - A. Fully inform families about all test results, positive and negative?  
     - Never  Some of the Time  Most of the Time  Always
   - B. Fully inform families about any diagnosis in a way they can understand?  
     - Never  Some of the Time  Most of the Time  Always
   - C. Help families name and explain their child’s diagnosis to others?  
     - Never  Some of the Time  Most of the Time  Always
   - D. Work together with families so they can explain how their child’s diagnosis might affect how s/he will be able to participate in school, community and faith-based activities?  
     - Never  Some of the Time  Most of the Time  Always

#### Ongoing Care and Support

**FCC Components: 1, 2, 3, 4, 6**

1. **Do you and your staff:**
   - A. Help families identify their strengths, skills and knowledge related to their child’s health care?  
     - Never  Some of the Time  Most of the Time  Always
   - B. Ask families what is working well?  
     - Never  Some of the Time  Most of the Time  Always
   - C. Help families identify areas where they may need additional support?  
     - Never  Some of the Time  Most of the Time  Always
   - D. Ask families to provide input about their child’s care based on their knowledge about their child’s temperament, behavior and reactions, and other current personal and family needs and priorities?  
     - Never  Some of the Time  Most of the Time  Always
   - E. Ask families, children and youth to share information, such as a change in daily routine or new stress that may provide insight into the interpretation of test results or diagnostic procedures?  
     - Never  Some of the Time  Most of the Time  Always
   - F. Encourage families to initiate consultation appointments or other meetings to discuss changes in their child’s care, for example, changes in medications, or other daily procedures?  
     - Never  Some of the Time  Most of the Time  Always
   - G. Fully inform families about diagnostic and treatment options in a way they can understand?  
     - Never  Some of the Time  Most of the Time  Always
   - H. Encourage questions about treatment options and the need for any procedures?  
     - Never  Some of the Time  Most of the Time  Always
   - I. Offer developmentally appropriate information (for example: stories, workbooks, videos, web-resources, tours) to prepare the family/child/youth for medical testing and procedures?  
     - Never  Some of the Time  Most of the Time  Always
2. Do you and your staff offer house calls, home care, or other ways that care/treatment can be provided in natural settings?  

3. If a family has difficulty traveling to specialty care settings outside of your community, can you and your staff receive partner with specialists to enhance the amount of care you can provide within your care setting?

4. If a family uses alternative and traditional medicine, do you and your staff work with the family to integrate traditional and alternative healing treatments into the child/youth’s overall care?

### Addressing Child/Youth Development

FCC Components: 3, 4, 6, 8

1. Do you and your staff:

   A. Assess and document the developmental status of each child/youth?  

   B. Reassess care approaches at key developmental milestones and transitions? (For example, when the child begins to walk, talk, begins school, enters puberty, and begins middle or high school.)

   C. Offer children/youth educational opportunities to support self-care? (For example, know how to make appointments, learn about insurance, medications, diet, and other aspects of health care.)

   D. Honor the family’s request about how much they want their child involved in decision-making?

   E. Include child in making decisions in keeping with age and development?

   F. Explain changes in who can give consent for care as the family’s child gets older?
### Access to Records

**FCC Component: 9**

**1. Do you and your staff:**

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Explain family’s rights and responsibilities under HIPAA in a way they can understand?</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
<tr>
<td>B. Have a policy to assure that financial issues, such as copying costs, do not prevent families from receiving copies of records?</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
<tr>
<td>C. Have a process to assist families in understanding and interpreting their child’s medical record? (For example, provide explanations of medical terms and answer questions about content.)</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
<tr>
<td>D. Let families or youth add information to their medical records?</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
<tr>
<td>E. Assure that all staff are aware of policies and procedures related to access to medical records?</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
</tbody>
</table>

### Appointment Schedules

**FCC Components: 6, 9**

**1. Do you and your staff:**

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Recognize families’ schedules and cultural events as important factors related to scheduling appointments?</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
<tr>
<td>B. Respond to families’ concerns about proposed scheduling of appointments and procedures?</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
<tr>
<td>C. Honor families’ requests for longer appointments to discuss complex issues?</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
<tr>
<td>D. Schedule appointments tailored to the needs of the child/youth and family? (For example, have “no wait” appointments for patients who may have immune system problems or developmental/behavioral issues that make it difficult to sit in the waiting room.)</td>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
</tr>
</tbody>
</table>

### Feedback on Care Setting Practices

**FCC Components: 4, 9**

**1. Are there opportunities for families/youth to provide feedback about care? (Examples include surveys, focus groups, suggestion boxes.)**

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
<td>Always</td>
</tr>
</tbody>
</table>

**2. Does your practice have a formal advisory committee for families and youth to provide input on policies and practices?**

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
<td>Always</td>
</tr>
</tbody>
</table>

**3. Are families/youth offered mentoring, or other supports and training to effectively participate on the advisory committee?**

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
<td>Always</td>
</tr>
</tbody>
</table>

**4. Do you and your staff make accommodations for families and youth who need supports to participate in advisory activities? (For example, English or sign language translation, alternate ways to participate, like via conference call or surveys.)**

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Never</td>
<td>Some of the Time</td>
<td>Most of the Time</td>
<td>Always</td>
</tr>
</tbody>
</table>
# Care Setting Policies to Support Family-Centered Care

### FCC Components: 4, 9

## 1. Do you and your staff:

| A. Make sure families are knowledgeable about care setting policies and practices? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| B. Have a policy about delivering family-centered care? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| C. Share information with families about elements of family-centered and culturally and linguistically competent care? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| D. Create job descriptions that include roles and responsibilities for all providers and staff related to family-centered and culturally and linguistically competent care? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| E. Provide orientation, training and support to staff and others related to family-centered and culturally and linguistically competent care? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| F. Evaluate staff performance and effectiveness in providing family-centered care and culturally and linguistically competent care? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| G. Invite families or youth to give presentations for staff to learn about the family perspective? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| H. Invite families or youth to do presentations for staff to learn how their cultures and values influence decision-making around their health care needs? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| I. Provide stipends to family and youth who share their knowledge and expertise with your staff? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |

## Addressing Culture and Language in Care

### FCC Components: 4

## 1. Do you and your staff:

| A. Ask families about any cultural values, beliefs or practices that might relate to their child/youth’s care? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| B. Honor families beliefs and practices when developing diagnostic and treatment plans? (This includes respect for diet; customs regarding eating, bathing, dressing; religious rituals related to health; modesty concerns such as gender of the provider, expected roles related to gender or generation.) | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| C. Offer trained interpretation (foreign language or sign)? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| D. Document the need for interpretation services when making referrals? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| E. Help families find information and educational materials they can easily read or that don’t need reading, like videos to watch? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |
| F. Employ staff that reflect the diversity of your community? | ○ Never ○ Some of the Time ○ Most of the Time ○ Always |

## 3. Do you have policies to institutionalize these care setting practices?  ○ Yes ○ No

## 4. Are all staff informed about Care Setting Policies and Practices?  ○ Yes ○ No

## 5. Are all families informed about Care Setting Policies and Practices?  ○ Yes ○ No
COMMUNITY SYSTEMS OF SERVICES AND SUPPORTS

**Information and Referral and Community-Based Services**  
**FCC Component: 5**

1. **Do you and your staff:**
   
   - A. Work with families to identify needed community-based services?  
     - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always
   
   - B. Help families make the first contact with community-based services?  
     - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always
   
   - C. Follow up to see if the family and child/youth has successfully connected with the service?  
     - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always
   
   - D. Follow up to see if the service was easy to access by the family?  
     - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always
   
   - E. Follow up to see if the service was useful to the family?  
     - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always
   
   - F. Follow up to see if the service was respectful of the family’s culture and values?  
     - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

2. **As family circumstances change (changing diagnoses, functional level, child transitions, change in family make up, etc.), do you and your staff work with the family to review current services and help identify new community-based services they may need?**  
   - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

**Community Systems Integration and Care Coordination**  
**FCC Component: 5**

1. **Do you and your staff:**
   
   - A. Document the community-based services families use?  
     (Examples include early intervention, Head Start, childcare, community recreation programs, vocational rehab, faith-based activities.)  
     - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always
   
   - B. Have a staff person or care coordinator that helps families coordinate care?  
     - ○ Never  ○ Some of the Time  ○ Most of the Time  ○ Always

Thank you for taking the time to complete this self-assessment tool.  
We welcome your feedback. Visit www.familyvoices.org to share your ideas and suggestions for how we might improve future versions.