Involving Patients and Families in Improving Care

...1...

Open Door Community Health Centers
Patient Voice
Jay Molofsky
Determinants of Health and Their Contribution to Premature Death

- Genetic: 30%
- Medical: 15%
- Environmental: 5%
- Behavior: 10%
- Social: 30%

Schroeder, NEJM 357; 12
Something to Pray For...

Patient Centered Care

Patients actively involved in managing their conditions on a day-to-day basis, developing the confidence to sustain healthy behaviors for a lifetime.
Humboldt Open Door Clinic Goal

We will develop collaborative relationships with patients and families to empower them to excel in the day-to-day management of their own conditions, and we will partner with patients and family advisors in quality improvement to create truly patient-centered care delivery.
What is Self Management Support?

Self Management support is a way of providing health care that “aids and inspires patient to become informed about their conditions and take an active role in their treatment.”
Example of Self Management Support

**MY ACTION PLAN**

| I ___________________________ and ________________________ |
| (name) | (name of clinician) |

I have agreed that to improve my health I will:

1. **Choose one of the activities below:**
   - Work on something that’s bothering me:
   - ___________________________
   - Stay more physically active!
   - Take my medications.
   - Improve my food choices.
   - Reduce my stress.
   - Cut down on smoking.

2. **Choose your confidence level:**
   - This is how sure I am that I will be able to do my action plan:
   - 10 VERY SURE
   - 5 SOMETHAT SURE
   - 0 NOT SURE AT ALL

3. **Complete this box for the chosen activity:**
   - What: ___________________________
   - ___________________________
   - How much: ___________________________
   - When: ___________________________
   - ___________________________
   - How often: ___________________________
   - ___________________________

(Signature)

(Signature of clinician)
Examples of On-going Support

Outside the visit through web-based resources that patients design, review and update, peer programs, telephone follow-up by the care team, and other community programs.
Humboldt Open Door Clinic

Patient Advisory Group
Established June 5, 2009
Recruitment

- Identify a Patient and Family Liaison as the point person to work with advisors.
- Ask providers and staff to refer patients to the group.
- Create a sign or poster to let patients know about the group and who to contact if they are interested.
Patient and Family Liaison

- Enables patients to have direct input and influence on the policies, programs, and practices that affect the care and services they receive.
- Organizes patient advisory group (attendees, agendas, meeting space, etc.)
- Point person for patient advisors to contact.
You are invited to join the Humboldt Open Door Clinic Patient and Family Care Advisory Board.

- Would you like to help guide the programs and services of Humboldt Open Door Clinic?
- Do you have ideas about ways the clinic can help you reach your health care goals?
- HODC has an opportunity for you to volunteer your time and energy as a Patient and Family Care Advisor
HODC is looking for patients who want to:

- Share both good and bad experiences in a constructive way.
- Work with others whose backgrounds, experiences, and styles may be very different from your own.
- Help us improve medical provider and staff communication skills with patients.
- Help us develop computer and community resources that patients trust.
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- Help us develop computer and community resources that patients trust.

For more information call Jacqui John, Patient and Family Liaison 826-0610 x1111 or email: patientboard@opendoorhealth.com

Current Projects

- Helping Patients...
- Better manage their disease.
- Prepare for their visit
- Explore Online Resources
- Get the most out of their visit
Humboldt Open Door Clinic
Patient Advisory Group Goals

• Help patients develop skills in managing their chronic diseases.
• Build relationships in which patients, their family members, Humboldt Open Door Clinic staff, and the community work together to provide the best services for patients and their family.
• Improve clinic services for patients and their families.
• Help develop clinic, community, and online resources patients need and trust.
Ideas in Action

As a result of the patient advisors’ clinic walk through, let’s look at some of the changes happening at Humboldt Open Door Clinic…
Advisor Feedback:
Make the clipboards more accessible

• Completed!
Advisor Feedback:
Lower the Hand Sanitizers and Masks

• Completed!
Advisor Feedback:
Place hooks on the doors in the restrooms for coats etc.

• Completed!
Advisor Feedback:
More and varied reading materials in waiting rooms
• Completed!
Idea:

“Things I Want to Remember” form for patients to organize their thoughts.
Things I want to remember for my appointment

What is the main reason for your visit today?

________

________

________

________

________

________

Other things you would like your provider to know about

________

________

________

________

________

Patient use only

Provider use only

This form was useful: YES NO

This form was useful: YES NO

Things to consider

Test Results Counseling Symptoms Side Effects

Forms Concerns Referrals Insurance

Prescriptions Family Needs My Care Plan Nutritionist
Idea: “Get The Most Out of Your Visit” posters to help patients prepare for their visit and make the most of the time they have with their provider.
Make the Most Out of Your Appointment

**Plan Ahead**
- Check in is 15 minutes before the appointment.
- Be on time to avoid late fees.
- Be on time to avoid late fees.
- Review your medical history.
- Be clear on what will be happening during your visit.

**Bring**
- A friend or family member can assist you.
- Bring your current medications or a current list of your medications.
- Know what you are taking and why.
- List allergies (how much you take and how often).
- Include medications you are no longer taking.
- List allergies (how much you take and how often).

**Write down symptoms/break a chart or diagram**
- Sometimes it is difficult to remember everything during the visit.
- Keep a journal of your symptoms and what you are doing to treat them.
- Please describe what you think is happening.

**Know your insurance plan**
- Know your insurance plan.
- Bring your insurance card.
- Have your co-payment ready.

**During Your Visit**
- Focus on the medical main points.
- Typical visits are 15 minutes.
- Discuss the most important issues first.
- (If you see the doctor first, you may see the doctor later).
- Limit the conversation to things the doctor can address.

**After the Visit**
- Make your next appointment.
- If you need more time, ask for an extended appointment.

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**Compassion**

**Access to Care**

**Respect**

**Excellence**

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Aproveche Su Consulta al Máximo

Prepárese con Tiempo

Llegue con Anticipo
- El registro se hace 15 minutos antes de la cita.
- Organice sus ideas mientras espera.
- Traiga algo para entretenimiento a sus hijos.
- Traiga a un amigo.
- Un amigo o paciente pueden ayudarle.

Traiga los frascos o una lista de las medicinas que está tomando ahora.
- Sepa lo que está tomando, y para qué.
- Escriba la dosis de cada uno, y cada vez que las toma.
- Diga cada medicación, ya sea tomada.
- Diga si tiene alergia a alguna medicina.

Escriba su síntoma/s: Lleve un diario o dibuje/llée estadísticas.
- A veces es difícil recordar toda una síntoma.
- Mantenga un diario de sus síntomas y lo que está haciendo para tratarlos.
- Describa lo que usted piensa que está sucediendo.

Entrese de cuanto dinero cubre su seguro.
- Traiga su tarjeta de seguro.
- Traiga el dinero para pagar su co-pago.

**Durante su Visita**

Concentrése en el principal motivo médico de su visita.
- Las visitas duran 15 minutos generalmente.
- Diga el motivo principal primero; después, puedo no tener tiempo.
- Limita su conversación a los temas relacionados con la visita.

**Después de la Visita**

Solicite su próxima cita.
- Si necesita más tiempo, solicite una cita más larga.
Waiting Room Poster
Make the most from your visit today

Plan Ahead

Arrive early
• Check in is usually 15 minutes before the appointment
• Use the time to gather your thoughts
• Focus on what you need from this visit

Bring a friend
• A friend or family member can assist you
Plan Ahead

Bring your current medications or a current list of your medications.

• Know what you are taking and why

• List the dosages (how much you take and how often)

• Include medications you are no longer taking

• Know allergic reactions to medications
Plan Ahead

Write down symptoms/bring a journal/make a chart or diagram

• Sometimes it is difficult to remember everything during the visit

• Keep a journal of your symptoms and what you are doing to treat them

• Describe what you think is happening
Plan Ahead

Know your insurance plan coverage

- Bring your insurance card

- Have your copayment ready
During Your Visit

• Typical visits are 15 minutes

• Discuss the most important issue first (If you save this one for last you may run out of time)

• Limit the conversation to things the provider can address
• Make your next appointment

• If you need more time, ask for an extended appointment

• Ask for an *After Visit Summary Report*
We are partners.

We care.

Compassion
Access to Care
Respect
Excellence

Open Door puts the patient and family at the center of their health care team.

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Open Door Community Health Centers
Current Projects

• Online resource guide regularly reviewed and revised by patients
• Wallet sized clinic information card for patients
• Posters for waiting rooms to inform patients of our website
• Patient led art therapy group at local Wellness Center
• Patient advisors invited to corporate retreat (2/11) to help set 2011 Open Door dashboard.
HODC Patient Advisory Group
Some of our members
PARTNERING WITH PATIENT ADVISORS IN QUALITY IMPROVEMENT

Ellen H. Chen, MD
UCSF Department of Family and Community Medicine
On San Francisco General Hospital campus

- 10,000+ patients served
- 40,000 visits per year
- Full scope family practice
- Teaching clinic: 41 family practice residents and many medical and nursing students
- Diverse patient population
  - 41% Latino, 27% Asian, 15% Caucasian, 12% African American
  - 71% public insurance, 29% uninsured
  - 26 different languages spoken
    - 47% English, 30% Spanish, 9% Cantonese/Mandarin
Multidisciplinary team as the referral base for recruiting patient and family advisors.

30 patient and family advisors

Informational PAB meeting

7-10 Spanish speaking patients active since February
Engaging patient advisors

- The individual relationship
- Build motivation-
  - Framing participation as an opportunity for skills and support
- Facilitating logistics
SEGUNDO ENTREGABLE:
PLANIFICACIÓN Y CONDUCCIÓN DE REUNIONES EFECTIVAS

Alumno: Carlos Augusto Lone Sáenz
Fecha de Entrega: 06 / 09 / 2009
Mesa Directiva de Pacientes

Gracias por venir a nuestra reunión de la Mesa Directiva de Pacientes. Trabajar junto con los pacientes nos ayuda a mejorar su satisfacción y a alcanzar nuestra meta de brindar excelencia en el cuidado médico y servicio. Para ayudarlo/a a prepararse, la parte superior de este formulario es completada por uno de los miembros de la Mesa Directiva. La parte inferior, con la lista de cosas a chequear, es para que la use usted de la forma que prefiera.

Nuestra Mesa Directiva es:


Los principales objetivos en los que nos enfocamos son:


Algunos de los temas pasados que hemos tratado son:


------------------------------------------------------------------------------------------------------------------

Su información será aún más útil para nosotros si usted:

- [ ] Nos comenta brevemente en qué está trabajando
- [ ] Viene preparado con ideas sobre la manera en que los pacientes y/o sus familias y amigos pueden ayudar a mejorar la clínica y sus servicios
- [ ] Tiene definidas las preguntas que quiere que le contesten los miembros de la Mesa Directiva
- [ ] Escucha las preguntas y las inquietudes de los miembros de la Mesa Directiva
- [ ] Se enfoca en temas que se están tratando en el momento
- [ ] Realiza comentarios acerca de lo hablado en el final de la reunión. Provee una dirección de correo electrónico (e-mail) u otra información de contacto para que se puedan comunicar con usted los miembros que no hayan podido estar presentes en la reunión (hasta 2 semanas después de la misma).
- [ ] Está preparado para compartir ideas acerca de cómo realizar el seguimiento (por ejemplo, a través de nuevas visitas o de notas de progreso)

¡Muchas gracias por compartir sus experiencias y conocimientos con nosotros!
Quality improvement

- Engaging frontline clinic staff
  - Staff communication training
  - Agenda-setting form
- Resident education
- Building partnership with CBO to enhance computer literacy
- Future input: staff recognition program; referrals to community resources for self-management
Partnering with CBO

SFGH Family Health Center
Computer Skills Class to Support Information Seeking and Access to Community Resources
This path is not linear!
SFGH Family Health Center Walking Group
“Running” with an Idea from the Patient and Family Advisory Board
Acknowledgements

This work would not be possible without the contributions of Jorge Palacios, Irma Largaespada, Hali Hammer, and an entire multidisciplinary team at the FHC.

Thank you to the Institute for Patient and Family Centered Care and the California Healthcare Foundation.
ADVANCING THE PRACTICE OF PATIENT- AND FAMILY-CENTERED CARE IN PRIMARY CARE AND OTHER AMBULATORY SETTINGS

How to Get Started…

Institute for Patient- and Family-Centered Care
6917 Arlington Road
Bethesda, MD 20814
(301) 652-0281
www.ipfcc.org

For further information about patient- and family-centered care in primary care and other ambulatory settings: http://www.ipfcc.org/advance/topics/primary-care.html
The Institute for Patient- and Family-Centered Care provides essential leadership for advancing the practice of patient- and family-centered care. Through the development and dissemination of materials, policy and research initiatives, training, technical assistance, and on-site consultation, the Institute serves as a central resource for increasing the understanding and skills necessary to build effective partnerships with patients and families.

For well over a decade, there has been growing recognition of the enormous benefits patient- and family-centered care offers to health care providers, patients, and families in all areas of health care. As hospitals, primary care practices, other outpatient settings, and health systems struggle with issues related to quality, safety, HIPAA compliance, workforce capacity, the use of technology, the need to renovate or build new facilities, and cost control, they are recognizing that patient- and family-centered approaches and the perspectives of patients and families are essential to their efforts.

Visit our website at www.ipfcc.org for additional resources, tools, schedule of events, profiles of patient and family advisors and leaders, and profiles of organizational change.
What is patient- and family-centered care? Why does it matter? How does it fit with our overall mission? And finally, what can our practice or organization do to advance the practice of patient- and family-centered care? Where do we start?

Today, health care leaders and patients and families nationwide are asking these questions. The purpose of this document is to provide some answers.

Part I provides a rationale for a patient- and family-centered approach to care and defines its core concepts.

Part II outlines steps an organization can take to begin to create partnerships with patients and families and offers practical suggestions for getting started.

Part III, “Where Do We Stand?,” provides an assessment tool that primary and other ambulatory care centers can use to determine the degree to which patient- and family-centered approaches are embedded in their current organizational culture.

Part IV lists selected print and web resources.

Part V, “Selecting, Preparing, and Supporting Patient and Family Advisors,” offers practical guidance for beginning the process of identifying, recruiting, and sustaining the involvement of advisors.

Part VI, “A Checklist for Attitudes About Partnering with Patients and Families,” provides a tool for gathering information about the perceptions and attitudes of staff and administrative leaders.

PART I: WHAT IS PATIENT- AND FAMILY-CENTERED CARE?

Rationale

In their efforts to improve health care quality and safety, health care leaders today increasingly realize the importance of including a perspective too long missing from the health care equation: the perspective of patients and families. The experience of care, as perceived by the patient and family, is a key factor in health care quality and safety.

Bringing the perspectives of patients and families directly into the planning, delivery, and evaluation of health care, and thereby improving its quality and safety, is what patient- and family-centered care is all about. Studies increasingly show that when health care administrators, providers, and patients and families work in partnership, the quality and safety of health care rise, costs decrease, and provider and patient satisfaction increase.
Core Concepts

▼ **Dignity and Respect.** Health care practitioners listen to and honor patient and family perspectives and choices. Patient and family knowledge, values, beliefs, and cultural backgrounds are incorporated into the planning and delivery of care.

▼ **Information Sharing.** Health care practitioners communicate and share complete and unbiased information with patients and families in ways that are affirming and useful. Patients and families receive timely, complete, and accurate information in order to effectively participate in care and decision-making.

▼ **Participation.** Patients and families are encouraged and supported in participating in care and decision-making at the level they choose.

▼ **Collaboration.** Patients, families, health care practitioners, and health care leaders collaborate in policy and program development, implementation, and evaluation; in facility design; and in professional education, as well as in the delivery of care.

PART II: MOVING FORWARD WITH PATIENT- AND FAMILY-CENTERED CARE: ONE STEP AT A TIME

Establishing patient- and family-centered care requires a long-term commitment. It entails transforming the organizational culture. This approach to care is a journey, not a destination—one that requires continual exploration and evaluation of new ways to collaborate with patients and families.

The following steps can help set a clinic or health system on its journey toward patient- and family-centered care.

1. Implement a process for all senior leaders to learn about patient- and family-centered care. Include patients, families, physicians, and staff from all disciplines in this process.

2. Appoint a patient- and family-centered steering committee comprised of patients and families and formal and informal leaders of the organization.

3. Assess the extent to which the concepts and principles of patient- and family-centered care are currently implemented within your clinic or health system. (A brief initial assessment tool appears in the next section of this document.)

4. On the basis of the assessment, set priorities and develop an action plan for establishing patient- and family-centered care at your organization.

5. Using the action plan as a guide, begin to incorporate patient- and family-centered concepts and strategies into the organization’s strategic priorities. Make sure that these concepts are integrated into your organization’s mission, philosophy of care, and definition of quality.

6. Invite patients and families to serve as advisors in a variety of ways. Appoint some of these individuals to key committees and task forces.
7. Provide education and support to patients, families, and staff on patient- and family-centered care and on how to collaborate effectively in quality improvement and health care redesign. For example, provide opportunities for administrators and clinical staff to hear patients and family members share stories of their health care experiences during orientation and continuing education programs.

8. Monitor changes made, evaluate processes, measure the impact, continue to advance practice, and celebrate and recognize success.

A template and example for *Creating an Action Plan* can be downloaded from www.ipfcc.org/tools/downloads.htm.

**PART III: WHERE DO WE STAND? AN ASSESSMENT TOOL FOR GOVERNING BOARDS, ADMINISTRATORS, PROVIDERS, STAFF, AND PATIENT AND FAMILY LEADERS**

An effective action plan for moving forward with patient- and family-centered care is based on a thoughtful assessment of the degree to which an organization has already incorporated key principles of this approach to care and of the areas in which progress remains to be made.

Here are some questions that can serve as a springboard for such an assessment. Ideally, the assessment should be completed individually by administrators, managers, physicians, staff, and patient and family advisors. Representatives of each of these groups should then convene to discuss the responses and, together, develop an action plan.

**INITIAL ASSESSMENT FOR PRIMARY CARE AND OTHER AMBULATORY CARE SETTINGS**

**Leadership in the Organization**

☐ Do the practice’s/clinic’s vision, mission, and philosophy of care statements reflect the principles of patient- and family-centered care and promote partnerships with patients and families?

☐ Has the practice/clinic defined quality health care?

☐ Does this definition include how patients and families will experience care?

☐ Has the definition of quality and philosophy of care been communicated clearly throughout the practice/clinic to staff and clinicians, patients and families, and others in the community?

☐ Do the practice’s/clinic’s leaders invest time and financial and personnel resources in patient- and family-centered initiatives?

☐ Do the practice’s/clinic’s leaders, through their words and actions, hold staff and clinicians accountable for collaborating with patients and families?
Patients and Families as Advisors

☐ Is there an organizational Patient and Family Advisory Council?
   △ If there is a Patient and Family Advisory Council, is patient safety a regular item?

☐ Do patients and families serve on committees and work groups involved in:
   △ Patient/family education?
   △ Care of chronic conditions (e.g., self-management support)?
   △ Peer-led education and support?
   △ Planning group visits?
   △ Patient safety?
   △ Transition planning?
   △ Information technology?
   △ Oversight of culturally and linguistically appropriate services?
   △ Community services and programs?
   △ Staff orientation and education?
   △ Policy and procedure development?
   △ Primary or ambulatory care redesign?
   △ Quality improvement?
   △ Facility design?

Architecture and Design

☐ Does the practice's/clinic's architecture and design:
   △ Create positive and welcoming impressions throughout for patients and families from diverse cultural and linguistic backgrounds?
   △ Display messages that communicate to patients and families that they are essential members of the health care team?
   △ Provide signage in the language(s) of the communities served?
   △ Provide for the privacy and comfort of patients and families?
   △ Support the presence and participation of families?
   △ Enhance patient and family access to useful information?
   △ Promote efficient workflow?
   △ Support collaboration among clinicians and staff across disciplines?
Patient and Family Participation in Care and Decision-Making

☐ Do staff and clinicians view patients and families as essential members of the health care team?

☐ Are patients and their families, according to patient preference, encouraged and supported to participate in care planning and decision-making?

☐ Are the cultural and spiritual practices of patients and families respected and incorporated into care planning and decision-making?

☐ Do providers encourage and support patients and their families, according to patient preference, to set goals and create action plans for self-management of chronic conditions?

☐ Are policies, programs, and staff practices consistent with the view that patients and families are allies for patient health, safety, and well-being?

Patient and Family Access to Information, Education, and Support

☐ Are there systems in place to ensure that patients and their families have access to complete, unbiased, and useful information?

☐ Are a range of informational and educational programs and materials consistently available to patients and families?

☐ Are written, audiovisual, and web-based information and educational resources provided in primary languages and at appropriate literacy levels for patients and families served by the practice/clinic?

☐ Do the written, audiovisual, and web-based information and educational resources include examples and images that reflect the diversity of patients and families served by the practice/clinic?

☐ Are trained interpreters available?

☐ Are peer-led educational programs available and accessible to patients and families?

☐ Are web-based opportunities for information-sharing and networking offered to patients and families?

Documentation Systems and Charting

☐ Does the registration/sign in process for practice/clinic visits capture the patient’s priorities, goals, and concerns for the visit?

☐ Do charting and documentation systems and the use of information technology support the recording of patients’ and families’ observations, goals, and priorities for care?
Do patients and families, according to patient preference, have easy and continuing access to their medical records?

Are patients and families offered a way to collect and organize personal health information (e.g., patient portals to the electronic medical record, electronic personal health record)?

Human Resources

Does the practice’s/clinic’s human resources system support and encourage collaboration with patients and families?

Are patient and family advisors involved in the hiring process for:

- staff leaders?
- clinical leaders?

Do position descriptions and performance appraisal processes clearly articulate the necessity of collaborating with patients and families of diverse cultural and linguistic backgrounds in care and decision-making?

Do position descriptions and performance appraisal processes clearly articulate the necessity of collaborating with patients and families in program development, professional education, and quality improvement?

Do position descriptions and performance appraisal processes clearly articulate the necessity of collaborating with staff across disciplines and departments?

Do position descriptions and performance appraisal processes clearly articulate the necessity of collaborating with community agencies?

Do staff reflect the diversity of patients and families served by the practice/clinic?

Does the practice/clinic employ doulas, promotores, and/or other lay health workers?

Are there rewards and recognition for partnering with patients and families?

Are there rewards and recognition for supporting self-management of chronic conditions?
Education of Staff, Clinicians, Students and Trainees

☐ Do orientation and education programs prepare the following people for collaboration with patients and families in care and decision-making:
  ▼ Staff?
  ▼ Clinicians?
  ▼ Students and Trainees?

☐ Do orientation and education programs prepare the following people for culturally responsive practice:
  ▼ Staff?
  ▼ Clinicians?
  ▼ Students and Trainees?

☐ Do orientation and education programs prepare the following people to collaborate with patients and families to develop goals and action plans for self-management of chronic conditions:
  ▼ Staff?
  ▼ Clinicians?
  ▼ Students and Trainees?

☐ Are patients and families involved as faculty in orientation and educational programs?

Research and Evaluation

☐ In research and evaluation, do patients and families participate in:
  ▼ Shaping the agenda?
  ▼ Conducting the research or evaluation?
  ▼ Analyzing the data?
  ▼ Disseminating the results?

Adapted from Patient- and Family-Centered Ambulatory Care: A Self-Assessment Inventory. Available from the Institute for Patient- and Family-Centered Care at www.ipfcc.org/tools/downloads.htm.
PART IV: SELECTED RESOURCES AND WEBSITES

Resources

For the most recent references on this topic, please see the Institute’s Primary Care Bibliography at http://www.ipfcc.org/advance/supporting.html.


Minnesota Rules Chapter 4764, Minnesota Department of Health (MDH), 34 SR 951 (January 11, 2010).


Websites

Agency for Healthcare Research and Quality (AHRQ)
www.ahrq.gov
AHRQ funds, conducts, and disseminates research to improve the quality, safety, efficiency, and effectiveness of health care. The information gathered from this work and made available on the website assists all key stakeholders — patients, families, clinicians, leaders, purchasers, and policymakers — make informed decisions about health care.

American Academy of Pediatrics National Center of Medical Home Initiatives for Children with Special Needs — Tools for Youth
www.medicalhomeinfo.org/tools/youthindex.html
This special section of the Medical Home website is devoted to providing youth with special health care needs and their families with information and tools to help as they transition into adult health care.

California HealthCare Foundation (CHCF)
www.chcf.org
The California HealthCare Foundation is an independent philanthropic organization that supports improvements in health care for all Californians. They offer resources on their site that will be helpful to anyone interested in chronic care, care for the underserved, and health information technology.

Center for Medical Home Improvement (CMHI)
www.medicalhomeimprovement.org
The “medical home” began as a model for providing comprehensive primary care to children and youth with special health care needs. This site has practical assessments and resources for providers in community practices and families serving on improvement teams. The complete Medical Home Improvement Kit including measurements, strategies, and A Guide for Parent and Practice “Partners” Working to Build Medical Homes for CSHCN can be downloaded from the site.

The Commonwealth Fund — Patient-Centered Coordinated Care Program
www.commonwealthfund.org/Content/Program-Areas/Delivery-System-Innovation-and-Improvement/Patient-Centered-Coordinated-Care.aspx
The Commonwealth Fund supports activities focused on improving the quality of primary health care in the United States, including efforts to make care more centered around the needs and preferences of patients and families. This section of the website contains information about current projects and grant opportunities.

Consumer Assessment of Healthcare Providers and Systems (CAHPS)
www.cahps.ahrq.gov/
The Consumer Assessment of Healthcare Providers and Systems program develops and supports the use of a comprehensive and evolving family of standardized surveys that ask consumers and patients to report on their experiences with health care.
Family Voices

www.familyvoices.org

Family Voices offers a variety of information and resources for families and providers caring for children with special health care needs. Family Voices activities embody the principles of patient- and family-centered care.

Foundation for Informed Medical Decision Making (FIMDM)

www.fimdm.org

The mission of FIMDM is to strengthen the ability of patients in making decisions about their health care. The site offers evidence-based information about treatment options, interviews with patients, decision-aids to guide decision-making, and other tools on a variety of conditions.

Health Care Transitions Initiative

http://hctransitions.ichp.ufl.edu

Established at the University of Florida, this program’s mission is to increase awareness of, gain knowledge about, and promote cooperative efforts to improve the transition from pediatric to adult health care. It has audiovisual and print materials for adolescents, young adults, parents, and providers.

Improving Chronic Illness Care

http://www.improvingchroniccare.org

As a national program of the Robert Wood Johnson Foundation, Improving Chronic Illness Care (ICIC) is dedicated to improving the experience of chronic illness care for patients and their families. Based at the MacColl Institute for Healthcare Improvement in Seattle, ICIC promotes the use of evidence-based change concepts to enhance care. This site provides comprehensive resources about the Chronic Care Model, which views the patient as a partner with providers in decision-making, participation in care, and quality improvement. Useful tools and strategies for change and evaluation are shared.

Institute for Patient- and Family-Centered Care

http://www.ipfcc.org

The Institute extends its efforts to provide leadership to advance the understanding and practice of patient- and family-centered care through its website. The site includes a wealth of practical resources, effective strategies, and profiles from families, providers, and organizations. Information and resources on Primary Care can be found in the Special Topics section under Advancing the Practice.
Institute for Healthcare Improvement (IHI)
http://www.ihi.org
Founded in 1991, IHI has been a leader in advancing the improvement of health care. IHI’s ever-expanding website has a wealth of information on patient and family involvement in quality improvement and research. This includes strategies to capture the patient and family experience of care as well as to involve patients and families on research and evaluation teams. Particularly related to primary care are the topic sections on chronic conditions, office practices, patient-centered care, and self-management support.

The John D. Stoeckle Center for Primary Care Innovation
http://www.mgh.harvard.edu/stoecklecenter
The Stoeckle Center, based at Massachusetts General Hospital (MGH) works with physicians, patients and primary care practices at MGH and elsewhere and partners with other organizations to explore approaches—to revitalize primary care and develop a model of primary care for the future.

National Center for Cultural Competence
http://www11.georgetown.edu/research/gucchd/nccc
This center’s work is focused on increasing the capacity of health and mental health programs to design, implement, and evaluate culturally and linguistically competent service delivery systems. They offer numerous and valuable online resources for improving primary health care, including self-assessment tools and publications and on-site training and education.

National Initiative for Children’s Healthcare Quality (NICHQ)
www.nichq.org
NICHQ has the mission to eliminate the gap between what is and what can be in health care for all children. Many of their initiatives support family involvement in quality improvement.

National Patient Safety Foundation
http://www.npsf.org
The Foundation’s mission is to improve the safety and welfare of patients in the health care system. Its website provides, among other resources, an area devoted solely to resources for patients and families who wish to get involved in patient safety initiatives.

National Working Group on Evidence-Based Health Care
www.evidencebasedhealthcare.org
This site provides a clearinghouse for consumers, voluntary health organizations, providers, and researchers to obtain information and resources about evidence-based health care. Useful resources include the guidance, *The Role of the Patient/Consumer in Establishing a Dynamic Clinical Research Continuum: Models of Patient/Consumer Inclusion.*
New Health Partnerships
http://www.newhealthpartnerships.org
New Health Partnerships offers an online community for patients, families, and health care providers dedicated to improving the health care and lives of people with chronic conditions. Profiles of individuals and organizations, information, tools, and other resources promoting collaborative self-management support and strategies for involving patients and families in improvement efforts are offered.

Patient-Centered Medical Home Resource Center
http://www.pcmh.ahrq.gov/portal/server.pt/community/pcmh__home/1483
This section of the AHRQ website provides access to evidence-based resources about the medical home and its potential to transform primary care and improve the quality, safety, efficiency, and effectiveness of U.S. health care.

Patient-Centered Primary Care Collaborative
http://www.pcpcc.net/
This Collaborative works to promote the development of “patient-centered medical homes,” primary care settings in which partnerships between providers, patients, and families are central. Information to facilitate patient and family understanding about the medical home model and to encourage their involvement in transformation include a short video, What is a Patient-Centered Medical Home, and a written guide, A Collaborative Partnership: Resources to Help Consumers Thrive in the Medical Home. (See http://www.pcpcc.net/consumers-and-patients)

Patient Decision Aids
http://decisionaid.ohri.ca
This site is part of the Ottawa Health Research Institute and was created to assist clinicians and patients make difficult health care decisions. The program is research-based and the site offers online tools, clinician training programs, and other resources.

Portland State Research and Training Center
www.rtc.pdx.edu/index.php
The Center promotes effective community-based, culturally competent, family-centered services for families and their children who are, or may be affected by mental, emotional or behavioral disorders. This goal is accomplished through collaborative research partnerships with family members, service providers, policy makers, and other concerned persons.

Professionals with Personal Experience in Chronic Care (PPECC)
www.ppecc.org
This group of health care professionals established PPECC to advocate for improved systems of care after personal and family experiences with chronic illness and long-term care. Health care professionals are encouraged to share their personal experiences with the health care system in order to promote greatly needed change.
Robert Wood Johnson Foundation (RWJF)

www.rwjf.org

RWJF has been working for over 30 years to improve U.S. health care through supporting research, advocacy, and policy work. RWJF has funded innovative programs that integrate patient- and family-centered care into their initiatives. A good example can be found in a current project, Designing the 21st Century Hospital: Serving Patients and Staff and is a great resource for leaders and managers undergoing a design project, http://www.rwjf.org/files/publications/other/Design21CenturyHospital.pdf.

Safety Net Medical Home Initiative

www.qhmedicalhome.org/safety-net/index.cfm

In 2008, The Commonwealth Fund, Qualis Health and the MacColl Institute for Healthcare Innovation at the Group Health Research Institute launched an initiative to support primary care safety net clinics in becoming high-performing patient-centered medical homes. The goal is to develop a replicable and sustainable implementation model for medical home transformation. Practical tools and resources are available on the website.

PART V: SELECTING, PREPARING, AND SUPPORTING PATIENT AND FAMILY ADVISORS IN PRIMARY CARE AND OTHER AMBULATORY SETTINGS

Hospitals, clinics, and other community-based ambulatory care practices are increasing efforts to partner with patients and families in policy and program development, patient safety, quality improvement, health care redesign, professional education, facility design planning, and research and evaluation. They are asking patients and families to serve on patient and family advisory councils, committees, task forces, and project teams. Appropriate selection, preparation, and support of patient and family advisors are key to effective partnerships.

Selecting Patient and Family Advisors

A patient or family advisor is an individual or family member who has experienced care in the clinic or other ambulatory care setting. In identifying patient and family advisors, look for individuals who have demonstrated an interest in partnering with providers in their care or the care of a family member. Consider those who have offered constructive ideas for change and who have a special ability to help staff and physicians better understand the patient or family perspective.

Seek individuals who are able to:

▼ Share insights and information about their experiences in ways that others can learn from them.
▼ See beyond their personal experiences.
▼ Show concern for more than one issue.
▼ Listen well.
▼ Respect the perspectives of others.
▼ Interact well with many different kinds of people.
▼ Speak comfortably in a group with candor.
▼ Show a positive outlook on life and a sense of humor.
▼ Work in partnership with others.

To find individuals with these qualities and skills, ask physicians and other clinicians for suggestions. Reviewing the patient registry or their panel of patients may help clinicians identify potential advisors. Community outreach workers, promotores and other lay health workers, and current patient and family advisors may also be able to identify potential advisors. Contacting community support groups and health and wellness and chronic care education programs is another way to find individuals who might be interested in serving as advisors.

For additional ideas for recruiting patients and families, see Tips for Recruiting Patients and Families to Serve in Advisory Roles, available at www.ipfcc.org/tools/downloads.html.

**Informing Potential Patient and Family Advisors About Roles and Responsibilities**

Before individuals can make decisions about whether they wish to participate on a patient safety committee, a quality improvement team, or in other redesign projects, they should be informed of the responsibilities and privileges associated with the role. A fact sheet containing the following information, as relevant, can be prepared and offered to individuals who are being asked to participate:

▼ Mission and goals of the group or project.
▼ Expectations for their participation.
▼ Meeting times, frequency, and duration.
▼ Travel dates.
▼ Expectations for communication among team members between meetings (including means of communication—email, mail, phone, etc.).
▼ Time commitment beyond meeting times.
▼ Compensation offered.
Benefits of participation (i.e., what are the expected outcomes of their involvement).

Training and support to be provided.

Compensation

At a minimum, the program should reimburse patients and families for expenses incurred in association with their work with the team (e.g., parking, transportation, child care). Many programs also offer stipends or honoraria for participation in meetings. These payments typically range from $12 – $25 per meeting. Consider the needs of the patient or family advisor and ask about their preferences. If they have no means to cash a check, stipends will have to be offered in an alternative way (e.g., store voucher, cash, etc.).

Preparing and Supporting Patient And Family Advisors

In order for patients and families to participate effectively as advisors, appropriate orientation, training, preparation, and support should be offered. They should have a chance to discuss their questions or thoughts about the work with an identified staff member who is willing to serve as a liaison to the advisors and has time dedicated to coordinate activities with advisors.

The orientation for patient and family advisors should include information on the following as relevant:

- The mission, goals, and priorities of the clinic, community-based ambulatory practice, or health system.
- Patient- and family-centered care.
- Overview of patient safety issues and strategies.
- Models of care for patients with chronic conditions or special health care needs (e.g., collaborative self-management support, medical home).
- HIPAA and the importance of privacy and confidentiality.
- Specific skills and knowledge needed to be an effective team member (e.g., quality improvement methodology for those who will be participating on a quality improvement team).

If the organization has a volunteer program, its orientation and training may be very useful for patient and family advisors. Other training topics may include:
Speaking the organization's language, “Jargon 101.” While it is best to reduce the amount of jargon used in meetings, sometimes it is impossible to completely eliminate jargon. If there are terms that will be used frequently, make sure that patient and family advisors understand them. Encourage them to ask for an explanation of anything they don't understand.

Who’s who in the organization or on the project team and how to contact other team members.

How to prepare for a meeting: what to wear, what to do ahead of time, and what to bring.

How meetings are conducted: format, agenda, minutes, roles (e.g., facilitator, secretary, timekeeper).

Technologies that will be used (e.g., conference calls, web-based tools).

Effective communication skills:

- Expressing your perspective so others will listen.
- How to ask tough questions.
- What to do when you don’t agree.
- Listening, to and learning from, the perspectives of others.
- Thinking beyond your own experience.

How to prepare for any conferences or other events — making travel arrangements, all logistical information (e.g., hotel, transportation from airport to hotel), expenses that are covered, reimbursement procedures, what to wear, and what to bring. Some patients and families may not have credit cards and, therefore, may have difficulty arranging travel and will need assistance in planning travel and checking in to a hotel.

It is extremely helpful for new patient and family advisors to have a “coach” or mentor who can provide informal, ongoing support to them. A member of the staff or project team who has experience working on collaborative initiatives and is willing to serve as a liaison can be assigned to this role. An experienced patient or family advisor can also fulfill this role. This person can insure that patient and family advisors are prepared for each meeting. During meetings, this person can actively encourage participation of the advisor. Also they can debrief after each meeting to determine what additional information or resources patient and family advisors need. Most importantly, they can support patient and family advisors in participating fully on the team by providing feedback and encouragement.

The guidance tools, *Staff Liaison to Patient and Family Advisory Councils and Other Collaborative Endeavors* and *Tips for Group Leaders and Facilitators on Involving Patients*
Believe Patient and Family Participation Is Essential

The single most important guideline for involving patients and families in advisory roles and as members of improvement and redesign teams is to believe that their participation is essential to the design and delivery of optimal care and services. Without sustained patient and family participation in all aspects of policy and program development and evaluation, the health care system will fail to respond to the real needs and concerns of those it is intended to serve. Effective patient/family and provider partnerships will help to redesign health care and safety and quality. It will lead to better outcomes and enhance efficiency and cost-effectiveness. Providers will also discover a more gratifying, creative, and inspiring way to practice.

Involving patients and families as partners and advisors will…

▼ Bring important perspectives about the experience of care.
▼ Teach how systems really work.
▼ Inspire and energize staff.
▼ Keep staff grounded in reality.
▼ Provide timely feedback and ideas.
▼ Lessen the burden on staff to fix the problems… staff don’t have to have all the answers.
▼ Bring connections with the community.
▼ Offer an opportunity for patients and families to “give back.”

The tool, A Checklist for Attitudes About Partnering with Patients and Families in Primary Care and Other Ambulatory Settings, can be used to help physicians and staff explore their attitudes and beliefs about partnering with patients and families (see page 16, Part VI).

This material has been adapted from two resources: Developing and Sustaining a Patient and Family Advisory Council and Essential Allies: Families as Advisors published by the Institute for Family-Centered Care.
PART VI: A CHECKLIST FOR ATTITUDES ABOUT PARTNERING WITH PATIENTS AND FAMILIES IN PRIMARY CARE AND OTHER AMBULATORY SETTINGS

Use this tool to explore attitudes about patient and family involvement in their own health care and as advisors and/or members of committees and improvement teams. It can be used for self-reflection and as a way to spark discussion among staff and physicians before beginning to work with patients and families as members of quality improvement, policy and program development, and health care redesign teams.

Answer and discuss the following questions:

At each clinic visit:

☐ Do I believe that patients and family members bring unique perspectives and expertise to the clinical relationship?

☐ Do I encourage patients and families to speak freely?

☐ Do I listen respectfully to the opinions of patients and family members?

☐ Do I encourage patients and family members to participate in decision-making about their care?

At the organizational level:

☐ Do I consistently let colleagues know that I value the insights of patients and families?

☐ Do I believe in the importance of patient and family participation in planning and decision-making at the program and policy level?

☐ Do I believe that patients and families bring a perspective to a project that no one else can provide?

☐ Do I believe that patients and family members can look beyond their own experiences and issues?

☐ Do I believe that the perspectives and opinions of patients, families, and providers are equally valid in planning and decision-making at the program and policy level?

If you have experience working with patients and families as advisors on committees and teams, answer and discuss these additional questions:

☐ Do I understand what is required and expected of patients and families who serve as advisors and/or members on committees and teams?

☐ Do I help patients and families set clear goals for their roles in these efforts?
☐ Do I feel comfortable delegating responsibility to patient and family advisors?

☐ Do I understand that an illness or other family demands may require patients and family members to take time off from their responsibilities on committees and teams?

Adapted from Jeppson, E., & Thomas, J. (1994). Essential Allies: Families as Advisors. Available from Institute for Patient- and Family-Centered Care, Bethesda, MD.