The formulation of quality-of-care goals in medicine has expanded to include values such as patient centeredness (1). National data suggest that care for most medical conditions meets accepted standards for technical quality for only about half of patients (2). The prevailing approach to this problem is quality improvement programs, such as the collaborative approach to chronic disease management (3,4). Quality improvement programs have been evaluated for many conditions, including depression (5–12). In primary care, depression serves as an excellent tracer for studying quality problems, because although it can be treated effectively and involves high social costs, it is often undetected and untreated in primary care settings (13–17). Quality improvement programs for depression can improve quality of care and health outcomes for two years and beyond (5–12). However, few studies have evaluated the effects of quality improvement on ethics goals (18–20).

In this article we address this gap by illustrating how four ethics goals can be incorporated as outcomes in demonstrations of quality improvement, using depression in primary care as an example (21–27). Two ethics goals—beneficence and avoiding harm—are fundamental to quality improvement and might not seem to require separate attention. However, aspects of these goals fall outside the utilitarian framework that guides most quality improvement efforts and

The Partners in Care Approach to Ethics Outcomes in Quality Improvement Programs for Depression

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Objective: Patient centeredness and equity are major quality goals, but little is known about how these goals are affected by efforts to improve the quality of care. The authors describe an approach to addressing these goals in a randomized trial of quality improvement for depressed primary care patients. Methods: For four ethics goals (autonomy, distributive justice, beneficence, and avoiding harm), the authors identify intervention features, study measures, and hypotheses implemented in Partners in Care, a randomized trial of two quality improvement interventions, relative to usual care and summarize published findings pertinent to these outcomes. Results: To implement an ethics framework, modifications were required in study design and in measures and analysis plans, particularly to address the autonomy and justice goals. Extra resources were needed for sample recruitment, for intervention and survey materials, and to fund an ethics coinvestigator. The interventions were associated with improvements in all four ethics areas. Patients who received the interventions were significantly more likely to receive the treatment they had indicated at baseline as their preferred treatment (autonomy goal). Intervention-associated benefits occurred more rapidly among sicker patients and extended to patients from ethnic minority groups, resulting in a reduction in ethnic-group disparities in health outcomes relative to usual care (distributive justice goal). The interventions were associated with improved quality of care and health outcomes (beneficence goal) and with reduced use of long-term minor tranquilizers (goal of avoiding harm). Conclusions: It is feasible to explicitly address ethics outcomes in quality improvement programs for depression, but substantial marginal resources may be required. Nevertheless, interventions so modified can increase a practice’s ability to realize ethics goals. (Psychiatric Services 55:532–539, 2004)
merit separate attention under a broader perspective. Patient autonomy and distributive justice are our major focus and have rarely been included as goals in quality improvement evaluations. Yet leaders in health policy and public interest groups have emphasized the importance of addressing client centeredness (one aspect of autonomy) and equity (one aspect of distributive justice) as quality goals (1,17,28,29).

The Institute of Medicine and others have noted that promoting autonomy can compete with promoting equity, so special care is needed when addressing both goals in quality improvement (1,30). Here we address such issues by reviewing how Partners in Care (PIC) (31), a Patient Outcomes Research Team II (PORT-II) project of the Agency for Healthcare Research and Quality, adopted an ethics framework. We review the PIC study design and, for each of four ethics goals, discuss the framework that informed the study and how we modified the design and measures, and we summarize our hypotheses and relevant published findings.

Methods

Design of Partners in Care

PIC is a group-level randomized trial of the effects of quality improvement programs for depressed patients in primary care on cost, health outcomes, and quality of care (31–33). Seven managed primary care group practice organizations were recruited, selected to produce a practice sample that was diverse in organization and location and a patient sample that was overrepresentative of Latino persons, particularly Mexican Americans, while including African Americans at each site.

Across organizations, 46 matched primary care clinics were randomly assigned to provide either care as usual or one of two quality improvement interventions. A total of 181 primary care providers were enrolled. In each clinic, a consecutive patient sample was screened for depression with use of a self-report depressive symptom measure followed by a standard diagnostic interview. A total of 1,356 ongoing care patients with depression who had eligible insurance (that is, insurance or a public program that would potentially reimburse for services from intervention providers) were enrolled between June 1996 and April 1997. Participating patients were asked to complete baseline and semiannual follow-up surveys and interviews over two years and at 37 months.

Two quality improvement interventions were developed—one involving medications and the other involving therapy. The interventions provided information and resources for obtaining appropriate care for depression—either antidepressant medication or psychotherapy—without assigning or requiring treatment. Each intervention included four components: commitment of in-kind resources from the practices to support quality improvement; local clinician teams (consisting of a mental health specialist, a primary care clinician, a nurse, and in some practices an administrator) trained by study investigators to implement quality improvement strategies and to adapt them to local practice priorities and resources; a toolkit of clinician and patient education materials; and trained depression nurse specialists to educate, assess, and activate patients and facilitate initial treatment—that is, medication management or referral to therapy if indicated. In addition, each intervention had supplemental resources.

The resources for the medication intervention consisted of an extension of the nurse specialist role to provide follow-up on compliance with psychotropic medications for six or 12 months. The resources for the therapy intervention consisted of a reduction in copayment for up to 12 sessions of therapy from a practice therapist who was trained by the study in individual and group cognitive-behavioral therapy and was also trained to communicate regularly with the primary care provider (31–33).

Ethics consultation

The study funded an ethics coinvestigator (the first author, at 10 to 20 percent for three years) to educate investigators, consult about conflicts inherent in resource allocation, and observe project planning meetings to identify and help resolve ethical concerns. The coinvestigator identified opportunities for measuring ethics outcomes, participated in pretesting activities to observe consumers’ reactions to study materials, and presented the overall ethics perspective and goals to practice representatives at the first advisory board before the study was implemented. The consultation benefited from existing literature on ethical issues in the development of practice guidelines (23–25). The ethicist on this study was a psychiatrist and ethicist who had also trained in health services research. The study was approved by the institutional review boards of RAND and the participating practices and institutions.

Results

The four ethics goals, the intervention and study design modifications, and key hypotheses are outlined in Table 1. Ethics outcome measures and summaries of published findings are shown in Table 2. Below we briefly describe the study framework and highlight key points from the two tables.
Autonomy and respect for persons

The concept of autonomy, or self-rule, is often applied narrowly in medicine to a single encounter in which a patient gives adequately informed consent—or refusal—free from paternalistic interference. This emphasis on noninterference ignores the advocacy efforts by providers that may be needed to ensure exercise of autonomy by populations whose self-advocacy is limited—for example, because of cognitive deficits during depression. Furthermore, a broader conception of respect for autonomy is needed to respond to patients’ needs, not only the need to give informed consent at one moment in time but also the need to receive types of treatment that promote their own life goals over time (34,35).

We approached autonomy from a systems perspective. Patients’ options depend on physicians’ knowledge and attitudes, and findings from focus groups suggested that some managed care physicians tended not to offer psychotherapy, because they perceived it as less effective and less accessible than medication. We therefore thought that optimal practice culture should educate both physicians and patients to support alternative, effective treatments (different evidence-based medications and psychotherapies) and ongoing access to information about their costs and benefits.

Interventions. The interventions for the autonomy goal were designed to facilitate informed choice and genuine treatment options over time for patients and clinicians: patients and clinicians had full choice over treatment (including the option of no treatment), and practices could modify the interventions to fit their priorities and resources. The nurse spe-

| Table 1 |

<table>
<thead>
<tr>
<th>Ethics goal</th>
<th>Definition</th>
<th>Intervention and study design features</th>
<th>PIC hypothesis</th>
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<tbody>
<tr>
<td>Autonomy</td>
<td>Patients and providers have the opportunity to make free, informed decisions about treatment</td>
<td>Emphasis on patient and provider education. Emphasis on patient activation and preference by a nurse specialist. Patients and providers are allowed to choose among treatments or choose no treatment in all intervention conditions. Consumer and provider feedback on intervention materials. Clinics randomized to usual care or interventions that enhance information and resources for treatment while preserving patients’ and providers’ choice of treatments or no treatment.</td>
<td>Patients in intervention clinics will have better knowledge about depression treatments than those in usual care clinics. Providers in intervention clinics will have better knowledge of depression treatments than those in usual care clinics. Patients in intervention clinics will be more likely to receive the treatment they most preferred at baseline than those in usual care clinics.</td>
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<tr>
<td>Distributive justice</td>
<td>Improving access to treatment and outcomes for those with the greatest need; minimizing disparities in access to treatment; reducing disparities in health outcomes</td>
<td>Screening for depressive symptoms and disorder and identifying cases to participating practices. Minority investigators. Educational materials and therapy manuals available in English and Spanish. Cognitive-behavioral therapy developed for low-income patients from ethnic minorities. Practices asked to ensure access to Spanish-speaking nurse specialists and therapists. Patient videotape developed by African-American and Latino providers. Selection of sites with a high proportion of Latino patients and some representation of African Americans (across sites); selection of public- and private-sector systems of care.</td>
<td>The effect of the interventions relative to usual care on use of guideline-concordant care will be greater among patients who had depressive disorder at baseline than among those who had depressive symptoms only. The interventions will improve rates of appropriate treatment for minority patients as well as for whites. The interventions will reduce preexisting disparities in health outcomes for minority patients compared with whites.</td>
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<td>Beneficence</td>
<td>Maximizing opportunities for appropriate treatment</td>
<td>Use of treatment guidelines. Provider training. Depression nurse specialists.</td>
<td>Patients in intervention clinics will be more likely to receive guideline-concordant treatment than those in usual care clinics.</td>
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<tr>
<td>Avoiding harm</td>
<td>Decreasing the likelihood of using treatments that are known to worsen patients’ outcomes</td>
<td>Treatment guidelines discouraged long-term use of minor tranquilizers in the absence of a co-occurring anxiety disorder.</td>
<td>Patients in intervention clinics will be less likely to receive long-term minor tranquilizers (benzodiazepines) than those in usual care clinics.</td>
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<tr>
<td>Ethics outcome (reference)</td>
<td>PIC measures</td>
<td>Findings</td>
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<tr>
<td>Autonomy</td>
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<td>Patient knowledge gain</td>
<td>Change in patients’ knowledge about depression and its treatment (change in knowledge score from baseline to six months)</td>
<td>Controlling for baseline patient characteristics and study site, the quality improvement intervention involving therapy (QI-therapy) was associated with improved patient knowledge (t=2, df=998, p=.05); findings were similar for the pooled interventions relative to usual care (t=1.97, df=998 p=.06).</td>
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<td>Clinician knowledge gain</td>
<td>Change in provider knowledge score from baseline to 18 months</td>
<td>The combined QI interventions, relative to usual care, were associated with greater knowledge about psychotherapy (p=.006) and a tendency toward increased overall knowledge (p=.06).</td>
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<td>Use of preferred treatments</td>
<td>Likelihood that at six-month follow-up patients will be receiving the treatment they most preferred at baseline</td>
<td>At six-month follow-up, each QI intervention was associated with an increase in the percentage of patients who received the treatment they most preferred at baseline (50.7 percent to 54.2 percent of intervention patients compared with 40.5 percent of usual care patients, p&lt;.05 for each).</td>
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<tr>
<td>Beneficence: use of appropriate treatments; health outcomes</td>
<td>Likelihood of receiving guideline-concordant treatments at six and 12 months and over two years</td>
<td>At six- and 12-month follow-up, the intervention was associated with an increased rate of appropriate care and improved health outcomes for the sample as a whole, relative to usual care—for example, at six months, 50.9 percent of patients in the combined QI interventions but 39.7 percent of those receiving usual care had appropriate treatment (p&lt;.001); 55.4 percent of patients in the combined QI interventions but 64.4 percent of those with usual care exceeded the cutoff for clinical depression as measured by the Centers for Epidemiological Studies Depression scale (CES-D) (p=.005). Improvements in mental-health–related quality of life and employment status also improved with the QI interventions.</td>
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<td>Avoiding harm: use of long-term minor tranquilizers</td>
<td>Likelihood of use of minor tranquilizers for more than three months during the previous six months, measured at each follow-up for three years (controlling for co-occurring anxiety disorder)</td>
<td>Patients in the QI intervention involving medications (QI-meds) showed decreased use of long-term minor tranquilizers over time (from 4.6 percent at baseline to 2.5 percent at two years), whereas patients who received usual care or QI-therapy showed little change (4 percent to 7 percent). At two years, statistical differences were noted between QI-meds and usual care (p=.06) and between QI-meds and QI-therapy (p=.04).</td>
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<td>Distributive justice</td>
<td>Relative effects of the interventions on receiving appropriate care and having probable depressive disorder at six and 12 months among patients with disorder rather than symptoms only at baseline</td>
<td>Sicker patients—that is, those with depressive disorder at 12 months rather than symptoms only—tended to show greater improvements in appropriateness of care and outcomes under QI in the first six months relative to those with symptoms only; but improvements associated with interventions were comparable at 12-month follow-up between those who had disorder at baseline and those who had symptoms only.</td>
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<td>Effects of the intervention on underserved minorities</td>
<td>Effects of the intervention on use of appropriate treatments and having probable depressive disorder at six and 12 months among Latinos and African Americans</td>
<td>At six and 12 months, no significant differences were noted in the effects of the combined QI interventions on appropriateness of care for whites and for Latinos and African Americans combined, but the role of QI in reducing the percentage of patients with probable disorder at six and 12 months was greater among Latinos and African Americans combined compared with whites (p&lt;.05 for the interaction term). The effects of the intervention in reducing unemployment were significant only for white patients (p&lt;.05), although a similar trend was observed for Latinos and African Americans. Thus the intervention was effective among these minority groups and was associated with reduced disparity in their clinical—but not employment—outcomes relative to whites.</td>
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<tr>
<td>Effects of the intervention on outcomes for underserved minorities compared with whites</td>
<td>Comparison of the effect of the intervention on outcomes (such as having probable depressive disorder) at six and 12 months for Latinos and African Americans relative to whites</td>
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**Table 2**

Partners in Care (PIC) measures and findings for ethics outcomes
cials, through their communication activities and education toolkits for patients, enabled informed choice over time. Clinician training materials also encouraged attention to patients’ preferences (Table 1).

**Study design and measures.** The interventions randomly assigned clinics to information and resources that encouraged appropriate care rather than to treatments. To measure patients’ treatment preferences and patients’ and providers’ knowledge, survey items were added at baseline and at six-month patient follow-up and 18-month provider follow-up.

**Hypotheses and findings.** We hypothesized that the interventions would improve patients’ and providers’ knowledge about treatment and increase the proportion of patients receiving the treatment that they initially preferred (Table 1). The intervention involving therapy was associated with a significant increase in patients’ knowledge (t=2, df=998, p=.05; a similar trend was observed for the pooled interventions (p=.056). (These are previously unreported results.) In addition, the results showed that the combined quality improvement interventions improved clinicians’ knowledge about counseling and tended to improve overall knowledge (36). Furthermore, the interventions increased the proportion of patients at six and 12 months who received the treatment that they had preferred at baseline, relative to the patients who received usual care (37).

**Distributive justice**

Our working assumption was that explicit attention to justice is needed in designing quality improvement interventions, because individuals from underserved ethnic minority groups are likely to face ongoing societal barriers when attempting to obtain better care. Furthermore, interventions based on practice guidelines that have been formulated for the average patient may lead to insufficient attention being given to the particular needs of the sickest or more complex patients, such as those with suicidal ideation.

PIC’s approach to the distributive justice goal was influenced by the work of Rawls (38). From this perspective, it is unfair to allow people to be saddled with disadvantages that are not the result of their own choices and that prevent them from meeting their basic needs when such an outcome is preventable. Thus far, thinkers such as Daniels (39) have applied Rawlsian ideas to health care by using a definition of need based on severity of illness and functional limitations. We expand this conception to include a focus on disparities that reflect historical and current disadvantage (17,40).

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**Explicit attention to justice is needed in designing quality improvement interventions, because individuals from underserved ethnic minority groups are likely to face ongoing societal barriers when attempting to obtain better care.**

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**Intervention.** Practice administrators were encouraged to develop methods to facilitate access, particularly to psychotherapy, among patients from ethnic minority groups. When available, quality improvement components were selected that had been evaluated among low-income minority patients—for example, cognitive-behavioral therapy (41,42). Examples of modifications made by the practices include increasing the number of bilingual nurses or therapists, having therapy sessions in primary care settings, or conducting therapy by telephone. The expert leader, intervention staff, and clinician training materials included suggestions for addressing needs of sicker and more complex patients. The therapy intervention included a form of cognitive-behavioral therapy (four sessions) that was tailored to patients with minor depression. Practice therapists preferred having something to offer such patients. This modification was based on evidence that cognitive-behavioral therapy in minor depression may reduce symptoms of depression (43) and that individual components of cognitive-behavioral therapy may be as effective as full cognitive-behavioral therapy (44).

**Study design and measures.** PIC included a large number of Latino persons (primarily Mexican Americans, who constituted about a third of the sample, and included one public-sector organization with uninsured patients. The costs of modifying study materials for Spanish-speaking patients were equivalent to about half the (marginal) implementation costs of adding one site to the study, or $100,000. All surveys were field tested in Spanish and English, type fonts were enlarged for elderly patients, telephone assistance was available, and patients with hearing or other impairments could participate through in-person interviews in a location of their choice. The study included patients who at baseline had had current depressive symptoms and depressive disorder within the previous 12 months as well as those who had had current depressive symptoms without disorder within the previous 12 months. These patients were included to determine how practices prioritized the care of sicker patients and because the interventions were designed to facilitate management of a group of patients who were at high risk of developing depressive disorder, which included tracking patients who had symptoms only to watch for early evidence of disorder and to initiate treatment. No special measures were necessary to examine these outcomes.
Hypotheses and findings. We formulated three hypotheses about equity goals, which are listed in Table 1. First, we hypothesized that the benefits of quality improvement, in terms of improved quality of care and outcomes, would extend to and include Latinos and African Americans. We found that quality of care improved in all ethnic groups, with a greater gain in clinical outcomes in the first follow-up year under quality improvement for these minority groups relative to white persons. Benefits in the area of employment were significant among whites; a similar trend was found among the minority groups, but the statistical power to detect this effect was low, and the finding was not significant (45).

The second hypothesis was that the interventions would reduce disparities in health outcomes between white patients and patients from ethnic minority groups (17). We found that under usual care, African-American and Latino persons had worse outcomes than white persons in the first year but that this disparity was reduced for participants in clinics with the quality improvement initiatives (45).

Our third hypothesis was that the quality improvement interventions would be associated with greater improvements in quality of care among sicker patients who initially had depressive disorder than among patients who had only depressive symptoms. We found that in the first six months, the effects of the intervention on quality of care and health outcomes were stronger among patients who had had depressive disorder for the previous year at baseline than among those who had depressive symptoms without disorder. However, by the 12-month follow-up, the effects associated with the quality improvement interventions were comparable. Thus sicker patients initially were given higher priority for treatments. After the first six months, more equal outcome effects by baseline disorder status could reflect greater initial improvement among persons with a disorder and progression to a disorder among some of the patients who initially had only symptoms.

Beneficence
Beneficence refers to the positive duty of fiduciaries to benefit each person whom they are entrusted to serve and to the utilitarian goal of optimizing outcomes for a population (46). The former use of beneficence reflects an individual client perspective that may resonate with practice goals of individual clinicians but that is easy to overlook when such programs are implemented in the context of practice strategies to contain rising costs. In this respect, the latter use of beneficence in reference populations is likely to be the perspective that guides outcome goals related to quality improvement at the practice administrator level. The study focused on improving average outcomes while expanding benefits to specific vulnerable populations but respected the right of practices to serve patients according to their cultural norms and resources. From an ethics perspective, this approach represents a compromise. However, it facilitated a separate scientific study goal of evaluating the impact of quality improvement interventions as naturalistically implemented by practices.

Interventions. For the beneficence goal, the interventions provided information and resources to facilitate the recognition and assessment of depression and to increase appropriate treatment (31,32).

Hypotheses and findings. As we hypothesized, the interventions were associated with improvements in quality of care and mental-health–related outcomes and with an increase in the proportion of patients who were employed (33). Improved health outcomes persisted into the second follow-up year for patients in the therapy intervention, and benefits in the area of employment continued into the second year for both interventions. Higher use of antidepressant medication continued into the second follow-up year for patients who received the medication intervention (47–50).

Avoiding harm
Nonmaleficence is neither absolute nor independent of beneficence. Many medical procedures involve minor or harm in producing larger benefits. Furthermore, interventions that aim to maximize cost-effectiveness may conflict with interventions that seek to protect each individual patient from harm. We did not face this conflict in PIC, because our interventions sought to reduce harm by reducing the use of minor tranquilizers in the absence of comorbid anxiety, and this was also expected to improve cost-effectiveness (5,15).

Intervention features. Our quality improvement manual for clinics specifically recommended avoidance of long-term minor tranquilizers in the absence of comorbid anxiety disorder. In the medication intervention, the nurse specialist reinforced this recommendation through follow-up (Table 2).

Study design and measures. No special modifications were necessary other than collecting information about the long-term use of minor tranquilizers in each follow-up survey and assessing comorbid anxiety at baseline.

Hypothesis and findings. As we hypothesized, the medication intervention tended to be associated with lower use of long-term minor tranquilizers, especially at two years, compared with the therapy intervention and usual care (49).

Discussion
We found that it was feasible to adopt a standard ethics framework for developing practice guidelines for application in a major demonstration of quality improvement for depression in primary care. Adopting this framework required modification of the interventions, study design, and measures. We formulated new hypotheses and analysis plans to address the less commonly studied domains of autonomy and distributive justice. Our model for these modifications may serve as a template for other evaluations of quality improvement programs. Our empirical findings suggest that these modifications permit practices to more fully realize these ethics goals. For example, we found that this type of quality improvement intervention was associated with an increase in the percentage of patients who received their most preferred...
treatment among reasonable alternatives and with reductions in preexisting disparities in health outcomes by benefiting Latino and African-American persons in particular. Documenting such achievements could have other benefits for practices, such as increasing enrollment.

Addressing these ethics goals—particularly autonomy and distributive justice—and integrating them fully with the scientific goals of the project required substantial modifications to the study and substantial resources to be set aside. One key innovation was random assignment of opportunities for care rather than treatment per se. This approach enabled us to evaluate the impact of the interventions on realization of patients’ preferences. The sample size that was required to enable observation of the intervention effect—given that “encouragement” rather than treatment was randomized—was large (N=1,356). These design modifications were also necessary to address other study goal of estimating, through observational analyses, the effectiveness of treatments for depression under naturalistic care conditions (51).

The realization that this design afforded a unique opportunity to examine implications of quality improvement for patients who were being offered genuine options and then receiving preferred care—aspects of autonomy—is one example of the benefits of proactive ethics collaboration. Furthermore, the study identified patients through routine screening of visitors to the practices rather than the more common approach of clinician referral. This approach was selected to increase the generalizability of findings to consecutive patients visiting these practices and to reduce unobserved sources of heterogeneity due to provider referral criteria. However, it also represented an opportunity to optimize beneficence among patients whose depression was not already detected by their provider.

The inclusion of a large sample of patients from ethnic minority groups and modifications for Spanish-speaking patients derived more exclusively from an ethics (justice) objective, and study modification costs were substantial (we think at least $100,000). Implementing the more conventional ethics goals of beneficence and avoiding harm only trivially increased study costs. Formulating and implementing the full framework required three years of funding for an ethics coinvestigator, not the usual few days of consultation. In addition, it was necessary to work collaboratively with the practices to discuss the value of addressing these goals in the study design phase.

The PIC ethical framework, as implemented, had some important limitations. The intervention methods did not include an explicit method, such as a computerized decision analysis tool, to inform and address diverse values of patients or clinicians. Although others have been developing computerized decision aids on the basis of guidelines for depression care (52)—and these decision support tools can incorporate information about patients’ values—few actually do so. Such developments seem to represent an important and feasible direction for future quality improvement programs. In addition, our measures of ethics outcomes were self-reported by patients or providers.

Furthermore, as noted above, the study’s approach to beneficence represented a compromise from an ethics standpoint in that it promoted quality while respecting the practices’ organizational values and resource constraints. Such a position means that conflicts between the value of an individual patient’s needs and a plan’s resources can remain unresolved. However, it also represents a form of “neutrality” that permits the real stakeholders to reveal their priorities through the study data and findings while facilitating the study goal of observing practice-directed implementation of quality improvement. Thus the patients, providers, and practices themselves achieved the ethics outcomes reported by the study.

Conclusions

In conclusion, it is feasible to explicitly incorporate ethics goals into quality improvement demonstrations fielded in community practices. Furthermore, such programs can contribute to substantial, measurable improvements in ethics goals in the care of depressed patients. Although incorporating an ethics perspective entails real costs, the interventions achieved cost-effectiveness ratios in the range of widely used medical therapies (53). Thus it is possible to approach clinical, economic, and ethics goals of health care through programs that encourage use of guideline-concordant care by providing education and resources to enable flexible choice of treatment while encouraging inclusion of diverse populations. However, the impetus for substantial commitment to autonomy, justice, and other ethics domains as outcomes of quality improvement programs ultimately depends on how important such goals are to patients, providers, and policy makers. By sharing our approach, we hope to increase debate on the value of this commitment.

Acknowledgments

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References


