The Patient’s Voice: Legal Implications of Patient-Reported Outcome Measures

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Abstract:
In recent years, the medical community has paid increasing attention to patients’ own assessments of their health status. Even regulatory agencies, such as the Food and Drug Administration and the Centers for Medicare and Medicaid Services, are now interested in patient self-reports. The legal implications of this shift, however, have received little attention. This article begins to fill that gap. It introduces to the legal literature a discussion that has been ongoing in the health care field.

Patient-reported outcome measures (PROMs) are reports of patients’ symptoms, treatment outcomes, and health status that are documented directly by patients, typically through electronic questionnaires. In this era of growing efforts to control health care costs, improve care delivery, and combat physician burnout, patients’ own input can be invaluable for clinicians as well as researchers, regulators, and insurers. At the same time, however, PROMs have several pitfalls, and the implementation of PROM programs is challenging and complex.

The article argues that health care providers should be keenly aware of potential medical malpractice risks associated with PROMs. In addition, because PROMs collect a plethora of sensitive information about pain, sexual function, anxiety, and other matters, the HIPAA Privacy Rule should be revised to address PROMs specifically. The Article further posits that it would be premature for regulatory agencies or private insurers to require PROM submission at this time. It also details strategies, such as use of artificial intelligence, to strengthen PROMs and facilitate their integration into clinical practice and other arenas.

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INTRODUCTION

Anyone who reads the news or follows policy debates is aware of grave concerns about the U.S. healthcare system. A typical article from *Harvard Health Publishing* begins as follows: “Here’s a question that’s been on my mind and perhaps yours: Is the US healthcare system expensive, complicated, dysfunctional, or broken? The simple answer is yes to all.” In an effort to address some of the system’s grave shortcomings, health care and policy experts have developed concepts such as value-based care and comparative effectiveness research. They are also harnessing big data and artificial intelligence to benefit patients. Improving the system using any of these strategies, however, will depend on validly and reliably measuring health care outcomes.

This Article focuses on a particular means of assessing health care outcomes, called patient-reported outcome measures (PROMs). Little has been written thus far about the legal implications of PROM use. This Article begins to fill that


3 INSTITUTE OF MED., INITIAL NATIONAL PRIORITIES FOR COMPARATIVE EFFECTIVENESS RESEARCH 13 (2009) (“Comparative effectiveness research (CER) is the generation and synthesis of evidence that compares the benefits and harms of alternative methods to prevent, diagnose, treat, and monitor a clinical condition or to improve the delivery of care.”); Amit Dang & Kirandeep Kaur, *Comparative Effectiveness Research and its Utility in In-Clinic Practice*, 7 PERSPECT. CLINICAL RSCH. 9, 9-10 (2016).


5 42 U.S.C. § 1320e(a)(2)(A) (explaining that comparative effectiveness research involves “evaluating and comparing health outcomes and the clinical effectiveness, risks, and benefits of 2 or more medical treatments, services, and items . . . ”); Thomas Davenport & Ravi Kalakota, *The Potential for Artificial Intelligence in Healthcare*, 6 FUTURE HEALTHCARE J. 94, 94 (2019) (explaining that machine learning applications (a common form of AI) most often need to be trained on datasets with known outcome variables); Pantaleon, supra note 2, at 356 (“In value-based care, the only true measures of quality are the outcomes that matter to patients.”).


7 NAT. QUALITY F., PATIENT-REPORTED OUTCOMES: BEST PRACTICES ON SELECTION AND DATA
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gap, providing an overview of legal and technical PROM-related concerns. It introduces to the legal literature a discussion that has been ongoing in the medical community. Such analysis is particularly timely because the U.S. Food and Drug Administration (FDA) and the Centers for Medicare and Medicaid Services (CMS) have begun using PROMS for regulatory purposes. For example, approximately twenty-six percent of new drugs approved between 2016 and 2020 included patient reported outcome-related statements in labeling. Furthermore, physicians, who are increasingly pressed for time, may soon come to rely on PROMS as a partial replacement for extensive face-to-face conversations with patients.

Traditionally, individual and population health care outcomes have been assessed based on clinical measures such as mortality, number of hospital-acquired infections, number of avoidable hospital readmissions, blood pressure changes, and blood sugar levels. But what about patients’ own voices? Aren’t patients’ opinions about whether medical interventions improved or diminished their quality of life equally significant? And what about important conditions that cannot be clinically measured, such as pain, anxiety, or impaired sexual functioning?

In some instances, patients receive medications for their ailments (e.g., a rash or joint pain) but are not asked to return for follow-up visits. In those instances, physicians may obtain no information about treatment outcomes at all. In the absence of follow-up assessments, it may be difficult to determine which therapies work best for patients. This is a problem not only for individual patients and physicians but also for medical science in general.

See infra Parts III.B and IV.


 footnote texts
PROMs can fill these data vacuums. PROMs can be defined as reports of the "status of a patient’s health condition that come [] directly from the patient without interpretation of the patient’s response by a clinician or anyone else." Promoting an additional type of input is the patient-reported experience measure (PREM), which refers to patients’ perceptions of their interactions with the health care system or clinicians. This Article focuses on PROMs, which measure patients’ symptoms, functionality, and quality of life.

PROMs typically take the form of questionnaires that patients are asked to complete. They can be used for a variety of purposes. First and foremost, they are used in clinical care to inform physicians about patients’ conditions and assist them in making diagnostic and treatment decisions. In addition, PROMs are employed for purposes of 1) clinical research, including comparative effectiveness studies, 2) quality improvement initiatives, 3) FDA oversight and labeling, and 4) performance measurement and other assessments by insurers.

PROMs have many potential benefits, especially when employed in conjunction with clinician-reported outcomes and administrative data. In addition to the benefits discussed above, they can promote more informed clinical decision making, improve physician-patient communications, and foster patient empowerment. As a potent example of PROM benefits, one study found that monitoring PROMs increased the survival of metastatic cancer patients by

17 See infra notes 36-40.
18 Ian Porter et al., Integrating Patient Reported Outcome Measures (PROMs) into Routine Nurse-Led Primary Care for Patients with Multimorbidity: A Feasibility and Acceptability Study, 19 HEALTH QUALITY LIFE OUTCOMES 133, 134 (2021).
19 Neubert et al., supra note 17, at 1; Lee Squitieri et al., The Role of Patient-Reported Outcome Measures in Value-Based Payment Reform, 20 VALUE HEALTH 834, 834 (2017); Rahma Warsame & Anita D’Souza, Patient Reported Outcomes Have Arrived: A Practical Overview for Clinicians in Using Patient Reported Outcomes in Oncology, 94 MAYO CLINICAL PROC. 2291, 2292-98 (2019); MASS. MED. SOC., PATIENT-REPORTED OUTCOME MEASURES: CURRENT STATE AND MMS PRINCIPLES (2018), https://www.massmed.org/proms/.
20 Fatima Al Sayah et al., A Multi-Level Approach for the Use of Routinely Collected Patient-reported Outcome Measures (PROMs) Data in Healthcare Systems, 5 J. PATIENT-REPORTED OUTCOMES 98, 98 (Supp. 2 2021).
21 See supra notes 21-25 and accompanying text.
22 See infra Part I.B.
5.2 months. But PROMs come with a number of pitfalls and shortcomings.

One of us has personal experience with PROMs. Professor Podgurski has Parkinson’s disease. One neurologist’s office routinely gave him a tablet computer and asked him to complete long questionnaires prior to each of his appointments. He did this with difficulty because of his limited dexterity and because he felt pressured to complete the survey quickly, before being called in to see the doctor. Yet, the doctor never mentioned the PROMs and seemed unaware of the information Professor Podgurski provided. When doctors disregard PROMs that patients have worked hard to complete, patients may feel frustrated and resentful.

More serious shortcomings exist as well. For example, PROM questionnaires may not be validated and reliable and thus be of poor quality. Patients may not fully answer all questions, thus providing incomplete data. Patients’ responses may be biased by a desire to please the physician or by personality traits that influence their tolerance for discomfort. An additional problem for research initiatives is that the group of patients who are able and willing to complete PROMs may not be representative of the patient population as a whole, thereby yielding biased research results. Some patients do not have access to the technology needed to complete PROMs or have disabilities or language barriers that prevent them from doing so.

Health care providers may have their own difficulties with PROMs. Physicians may not know how to interpret PROM scores or determine if score

23 Ethan Basch & Allison M. Deal, Overall Survival Results of a Trial Assessing Patient-Reported Outcomes for Symptom Monitoring During Routine Cancer Treatment, 318 JAMA 197, 198 (2017) (“Median overall survival was 31.2 months (95% CI, 24.5-39.6) in the PRO group and 26.0 months (95% CI, 22.1-30.9) in the usual care group.”). See also infra note 111 and accompanying text.

24 Al Sayah et al., supra note 20, at 4-5.

25 Olalekan Lee Aiyegbusi et al., Patient and Clinician Opinions of Patient Reported Outcome Measures (PROMs) in the Management of Patients with Rare Diseases: A Qualitative Study, 18 HEALTH & QUALITY LIFE OUTCOMES 177, 185 (2020) (“The time constraints during clinics could prevent clinicians from acting on ePROM results and this could become a barrier to the use of ePROMs.”); Sara Heath, Only 1% of Docs Use Patient-Reported Outcomes Measures (PROMs), PATIENT ENGAGEMENT HIT (Nov. 2, 2022), https://patientengagementhit.com/news/only-1-of-docs-use-patient-reported-outcomes-measures-proms#:~:text=Only%201%25%20Docs%20Use,%20Outcomes%20Measures%20(PROMs). See also infra note 207 and accompanying text.

26 See infra Part I.C.1.a.

27 See infra Part I.C.1.d (discussing missing data).


29 See infra notes 166-186 and accompanying text.

30 See infra notes 179-182 and accompanying text.
changes are clinically meaningful. Clinicians may also feel that they are already overwhelmed and burnt out and that adding PROM use to their workload stretches them further towards the breaking point.

These challenges and others generate several legal concerns. Because PROMs may solicit sensitive information about patients’ quality of life, they raise questions about the adequacy of medical privacy protections. In addition, clinicians may rightly worry about medical malpractice liability associated with PROMs. The appropriateness of using PROMs for regulatory or reimbursement purposes is also open to debate.

The remainder of the paper proceeds as follows. Part I discusses the attributes, benefits, and risks of PROM use. Part II focuses on the clinical use of PROMs and analyzes privacy and medical malpractice concerns. It examines relevant HIPAA Privacy Rule requirements and exemptions that could threaten PROM confidentiality. In addition, this section posits that PROM use can generate malpractice risks for clinicians and health care entities under a variety of circumstances. These include health care providers ignoring data that are disclosed in PROMs because of time and workload constraints, relying on PROMs excessively when other diagnostic tools should have been used, or failing to implement PROMs when doing so has become the standard of care.

Part III assesses PROM use in research and FDA regulation. It highlights critiques of current PROM utilization in clinical studies. It also discusses the FDA’s acceptance of PROMs for medical device assessment and labeling purposes. Part IV focuses on PROM use for performance measurement and insurance coverage.

Part V formulates recommendations to address PROM-related legal concerns. It develops technical and administrative recommendations for PROM selection and implementation that would reduce the likelihood of malpractice claims and enhance PROM integrity. These include automation of PROM review using artificial intelligence, psychometric evaluations, pilot programs, stakeholder input, and more. Part V also recommends enhanced vigilance regarding data security, a modification to the HIPAA Privacy Rule, the development of clinical practice guidelines regarding PROM use, and patient education and notice concerning PROMs. Additionally, it outlines how PROMs might be used to support either plaintiffs or defendants in malpractice litigation. Part V further argues that it is premature for the FDA and CMS to mandate PROM use because of this tool’s potential weaknesses. At the same time, financial incentive programs for voluntary PROM adopters are desirable. Part VI concludes.

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31 See infra Part I.C.1.e (discussing PROM interpretability).
32 See infra Part I.C.2.b.
I. PROMS Attributes, Benefits, and Risks

PROMs can offer important insights into patient welfare, but they must be expertly selected and implemented so that they reflect human-centered design.33 This Part discusses the nature of PROMs along with their benefits and pitfalls.

A. What Are PROMs?

PROMs are usually standardized questionnaires that solicit patients’ input about their general health status and specific medical conditions.34 They focus on patients’ perceptions of their symptoms, ability to function, health behaviors, health care experience, and health-related quality of life.35 PROM scores can be compared over time to determine the efficacy of medical interventions.36 Patients can be asked to answer questionnaires online before or after their visits or can be given tablet computers to use at the clinician’s office.37 Administrators can also use paper forms, though many find electronic PROMs preferable.38

One example is the following short form sleep survey.39

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33 Lauren Landry, What Is Human-Centered Design?, HARV. BUS. SCH. ONLINE (Dec. 15, 2020), https://online.hbs.edu/blog/post/what-is-human-centered-design (“Human-centered design is a problem-solving technique that puts real people at the center of the development process, enabling you to create products and services that resonate and are tailored to your audience’s needs.”).
38 Jennifer Y. Yu et al., Electronic Forms for Patient Reported Outcome Measures (PROMs) are an Effective, Time-Efficient, and Cost Minimizing Alternative to Paper Forms, 19 PEDIATRIC RHEUMATOLOGY 67, 67 (2021).
A second example is the Oxford hip score, which uses twelve questions to evaluate hip pain and function in patients that may need hip replacements. Patients are asked to rate different types of hip pain (e.g., nighttime pain, shooting pain) and how it affects various functions, such as walking, climbing stairs, bathing, and shopping and are given five choices for each answer to indicate range of discomfort severity. Patients’ ratings in response to the individual questions are combined to generate an overall score. Thus, in the hip survey, scores in the range of 40-48 indicated that treatment is most likely not needed, and, at the other end of the spectrum, scores in the range of 0-19 indicate the presence of severe arthritis and a likely need for surgery.

PROMs can systematically collect information that would otherwise be difficult to obtain. For example, PROMs are particularly useful for those treating pain because pain cannot be objectively measured. Information about patients’ symptoms, functionality, and quality of life can also be invaluable in the specialties of oncology, cardiology, neurology, rheumatology, and more.

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41 Id.
42 Dawson et al., supra note 36, at 464.
43 OXFORD HIP SCORE, supra note 40.
44 Michelle M. Holmes et al., The Impact of Patient-Reported Outcome Measures in Clinical Practice for Pain: A Systematic Review, 26 QUALITY LIFE Rsch. 245, 249 (2017).
45 Roxanne E. Jensen et al., Review of Electronic Patient-Reported Outcomes Systems Used in Cancer Clinical Care, 10 J. ONCOLOGY PRAC. e215, e215 (2014); Warsame & D’Souza, supra note
In recent draft guidance, the FDA stated that PROMs are the best means of assessing the following:

- A feeling or experience known only to the patient, such as pain, itch, shortness of breath as no one else has direct access to feelings except for the patient
- Any type of functioning or activity that is part of the patients’ day-to-day life
- The patients’ satisfaction or dissatisfaction with their treatment and/or functioning
- Degree of impact on day-to-day life associated with one or more symptoms.49

PROMs are not a novel concept, and they have been embraced internationally. As early as 1975, Sweden incorporated PROMs into clinical databases that were disease specific.50 By 2000, PROMs were used by some U.S. practices, and since 2009, the United Kingdom has required that PROMs be collected for patients that undergo certain elective surgeries.51 The International Consortium for Health Outcomes Measurement (ICHOM), founded in 2012, states that its mission is to “unlock the potential of value-based health care by defining global Patient-Centered Outcome Measures . . . that really matter to patients for the most relevant medical conditions and by driving adoption and reporting of these measures worldwide.”52 To that end, ICHOM focuses on PROMs – outcomes that are reported directly by patients without being interpreted by clinicians.53
In 2004, the National Institutes of Health (NIH) launched the Patient Reported Outcomes Measurement Information System (PROMIS).\textsuperscript{54} Researchers used advanced psychometric\textsuperscript{55} techniques to validate existing survey instruments and to create better tools.\textsuperscript{56} As of this writing, the PROMIS website features 559 English-language surveys relating to anxiety, depression, fatigue, pain, sleep disturbance, physical functioning, satisfaction with participation in social roles, and much more.\textsuperscript{57} These are available free of charge to anyone who wishes to access them.\textsuperscript{58} Many experts consider PROMIS to be the gold standard for patient-generated assessments.\textsuperscript{59} PROMIS aims to standardize PROMs just as blood chemistry outcomes are standardized.\textsuperscript{60} PROMIS measures produce T-scores, which can be defined as “standard scores with a mean of 50 and standard deviation of 10 in a reference population (usually U.S. general population).”\textsuperscript{61} This enables comparison of an individual’s health status to that of the general population, or in some cases, a sub-population of interest (e.g., cancer patients).\textsuperscript{62}

Other PROM tools exist as well. One is the Medicare Health Outcomes Survey (HOS).\textsuperscript{63} The HOS is used in Medicare Advantage plans in order to gather health status data for purposes of quality improvement, monitoring and

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\textsuperscript{54} Patient Reported Outcomes Measurement Information System: Program Snapshot, NAT. INSTITUTES HEALTH, https://commonfund.nih.gov/promis/index (last visited Jan. 29, 2019);

\textsuperscript{55} Psychometrics is “the branch of psychology concerned with the quantification and measurement of mental attributes, behavior, performance, and the like, as well as with the design, analysis, and improvement of the tests, questionnaires, and other instruments used in such measurement.” Psychometrics, AM. PSYCH. ASS’N, https://dictionary.apa.org/psychometrics (last visited Dec. 11, 2022).

\textsuperscript{56} Lawson, supra note 54, at 16.

\textsuperscript{57} Id. at 17; Intro to PROMIS, HEALTH MEASURES, https://www.healthmeasures.net/explore-measurement-systems/promis/intro-to-promis (last visited Dec. 11, 2022); Search and View Measures, HEALTH MEASURES, https://www.healthmeasures.net/search-view-measures (last visited Apr. 22, 2023) (<under “search parameters,” select “English language,” and enter the following terms separately: anxiety, depression, fatigue, pain, sleep disturbance, physical functioning, satisfaction with participation in social roles.>) [hereinafter View Measures].

\textsuperscript{58} Lawson, supra note 54, at 16.


\textsuperscript{60} Id. at 346.

\textsuperscript{61} Nan E. Rothrock et al., Development and Validation of an Interpretive Guide for PROMIS Scores, 4 J. PATIENT-REPORTED OUTCOMES 1, 2 (2020).

\textsuperscript{62} Thi Xuan Mai Tran et al., Utility of the Patient-Reported Outcomes Measurement Information System (PROMIS) to Measure Primary Health Outcomes in Cancer Patients: A Systematic Review, 29 SUPPORTIVE CARE CANCER 1723, 1723 (2021).

\textsuperscript{63} Welcome to the Medicare Health Outcomes Survey (HOS) Website, CENTERS FOR MEDICARE & MEDICAID SERVS., https://www.hosonline.org/ (last modified Oct. 20, 2022).
rewarding plan performance, and helping participants make informed decisions. Each year a random sample of participants is surveyed, and the respondents are surveyed again after two years. Respondents are asked questions about their quality of life and daily functioning including matters such as mental health, incontinence, exercise, fall risks and more. After the second survey, change scores are calculated and each participant’s physical and mental health status is rated as “better than expected,” “as expected,” or “worse than expected.” CMS calculates summary HOS results for each Medicare Advantage Organization based on its members’ aggregated outcomes. CMS includes HOS measures in the Medicare Star Ratings program. The program scores Medicare Advantage plans using a range of one to five stars, and consumers can consult these ratings for purposes of plan selection. An additional tool is Focus on Therapeutic Outcomes (FOTO), which collects self-reported data from patients who underwent outpatient rehabilitation. FOTO assesses functional status changes in patients by comparing PROMs collected before, during, and after rehabilitation.

The extent of PROM use in the United States is unclear. According to one source, in 2016 only one-fifth of hospitals routinely used PROMs. A 2020 study noted that PROM adoption has been “limited” and that there is a “paucity of information on large-scale systemwide implementations that include diverse specialties and clinical settings.” The slow rate of PROM adoption is likely

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64 Id.
65 Id.
67 Reamer, supra note 66.
68 Id.
69 Id.
74 Rachel C. Sisodia et al., Factors Associated with Increased Collection of Patient-Reported Outcomes Within a Large Health Care System, 3 JAMA NETWORK OPEN e202764 (2020); see also
attributable to a variety of barriers that are discussed in Part I.C below.\textsuperscript{75}

To ease the burden of PROM completion and minimize the number of questions presented to patients, PROMs can leverage computer adaptive technology (CAT).\textsuperscript{76} Sometimes trained through machine learning (a type of artificial intelligence), CAT adapts the questions asked of each patient to the individual’s prior responses.\textsuperscript{77} Tailoring questionnaires to the responder’s symptoms and circumstances and eliminating irrelevant standardized queries can cut completion time by as much as fifty percent.\textsuperscript{78}

PROMs should be integrated into patients’ electronic health records (EHR) so that clinicians can easily review and maintain documentation concerning patient-reported information.\textsuperscript{79} Institutions can design their own integration mechanisms, can opt for EHR systems that embed PROMs, or can purchase independent commercial products to deploy PROMs.\textsuperscript{80} For example, experts at the University of Minnesota and other colleagues developed the Patient Reporting and Insight System from Minnesota (PRISM).\textsuperscript{81} PRISM enables patients to use a mobile app to fill out questionnaires and then integrates the responses into patients’ EHRs.\textsuperscript{82} Integrating PROMs into EHRs, however, can be challenging because of cost, logistics, and technological complexities.\textsuperscript{83}

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\textsuperscript{75} See infra Part I.C; Heath, supra note 25.

\textsuperscript{76} Liam T. Kane et al., \textit{Use of Computerized Adaptive Testing to Develop More Concise Patient-Reported Outcome Measures}, 5 \textit{JBJS Open Access} 2020, at 1.

\textsuperscript{77} Id. at 3; see also Conrad Harrison et al., \textit{Maximizing the Potential of Patient-Reported Assessments by Using the Open-Source Concerto Platform with Computerized Adaptive Testing and Machine Learning}, 22 \textit{J. Med. Internet Res.} 2020, at 2.

\textsuperscript{78} Id.

\textsuperscript{79} Marzyeh Amini et al., \textit{Facilitators and Barriers for Implementing Patient-Reported Outcome Measures in Clinical Care: An Academic Center’s Initial Experience}, 125 \textit{Health Policy} 1247, 1254 (2021); Heather Tafret Gold et al., \textit{Implementation and Early Adaptation of Patient-Reported Outcome Measures into an Electronic Health Record: A Technical Report}, 26 \textit{J. Health Informatics} 129, 130 (2020); National Quality Forum, supra note 7, at 21-22; Josef Stehlik et al., \textit{Implementation of Real-Time Assessment of Patient-Reported Outcomes in a Heart Failure Clinic: A Feasibility Study}, 23 J. Cardiac Failure 813, 815 (2017).


\textsuperscript{81} PRISM, \textit{Univ. Minn. Institute Health Informatics}, https://healthinformatics.umn.edu/research/research-projects/prism (last visited Dec. 11, 2022).

\textsuperscript{82} Id.

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B. PROM Benefits

PROMs can assist physicians in making medical decisions. Based on patients’ ratings of their discomfort and other quality of life indicators, doctors may change their course of treatment. Some outcomes, such as mortality, infections, and disease recurrence can be measured objectively. But outcomes such as pain levels and psychological wellbeing cannot be objectively assessed, and thus PROMs can complement objective measures and provide valuable insights about patients.

Ideally, physicians should be able to gather comprehensive information about patients’ perceptions of their health status by questioning them extensively during office visits, but sadly, that is often not possible in practice. Contemporary physicians are generally pressed for time and are often pressured by employers to limit the duration of visits to increase patient volume and profits. The average primary care visit, for instance, lasts only fifteen to twenty minutes. Therefore, PROMs may be the only way for clinicians to collect in-depth information about patients’ quality of life.

PROMs enable physicians to focus on symptoms, side effects, and outcomes that matter most to patients. To illustrate, a prostate cancer patient may care deeply not only about survival, but also about impotence and incontinence after

Outcomes 1, 12 (2022) (citing Daniell C. Lavallee et al., Incorporating Patient-Reported Outcomes into Health Care to Engage Patients and Enhance Care, 35 Health Aff. 575, 578-80 (2016)).

84 Holmes et al., supra note 44 at 252.


89 Id. at 88.
90 Youssef Ben Bouazza et al., Patient-Reported Outcome Measures (PROMs) in the Management of Lung Cancer: A Systematic Review, 113 Lung Cancer 140, 146 (2017) (discussing the benefits of PROMs).
treatment.\textsuperscript{91} If doctors collect PROMs about these complications, they will be better equipped to discuss them with patients and to tailor treatment recommendations to patients’ concerns.

In some cases, PROMs may save costs.\textsuperscript{92} One study found that careful surveillance of lung cancer patients using PROMs reduced the need for follow-up clinical visits and imaging.\textsuperscript{93} Although patients in the experimental arm of the study had a higher number of visits, their costs were lower because their symptoms were better controlled.\textsuperscript{94} In other cases, patients with knee, hip, or back pain whose PROMs reveal that they are high functioning and that their pain is tolerable could be spared expensive, unnecessary, and sometimes risky surgeries.\textsuperscript{95} A group of researchers focusing on spine surgery noted that PROMs have recently become an important tool in assessing the cost-effectiveness of procedures such as cervical and lumbar fusions.\textsuperscript{96} Information about quality of life outcomes can thus inform decisions about treatment options.\textsuperscript{97}

Since PROMs come directly from patients, they are free of any bias that might be introduced by clinicians interpreting what patients tell them.\textsuperscript{98} At least in some instances, therefore, they can provide better data than physicians’ descriptions of symptoms.\textsuperscript{99} More accurate information can better enable clinicians to make sound treatment decisions.\textsuperscript{100}

PROMs can potentially improve the physician-patient relationship by enhancing communication and patient engagement.\textsuperscript{101} PROM questionnaires can help patients remember their symptoms and drug side effects.\textsuperscript{102} They can induce
patients to think more deeply about their health status and to deepen their understanding of their medical conditions.\textsuperscript{103} PROMs can also make patients feel empowered to discuss concerns with their physicians because clinicians have solicited their views through the questionnaires.\textsuperscript{104} PROMs can help patients articulate their concerns and raise problems they may have otherwise been reluctant to report.\textsuperscript{105} They can therefore facilitate conversations with clinicians, enhance shared decision making, and increase patients’ satisfaction with their care.\textsuperscript{106}

One study focused on PROM use for rheumatology patients at the Cleveland Clinic.\textsuperscript{107} It revealed that seventy-eight percent agreed or strongly agreed that answering PROM queries improved communication with their physicians, and seventy percent agreed or strongly agreed that doing so made them feel that they had more control over their own care.\textsuperscript{108}

According to some estimates, oncologists miss symptoms, impaired functioning, and adverse effects of treatment fifty to seventy-four percent of the time.\textsuperscript{109} Physician awareness and response to these matters can generate dramatic benefits for patients. In one study, monitoring patient-reported outcomes increased the survival of individuals with metastatic cancer by 5.2 months.\textsuperscript{110} Participants in this study were randomly assigned either to receive usual care or to answer questions concerning twelve common symptoms via a web-based platform at and between office visits.\textsuperscript{111} Reports of severe or worsening symptoms would trigger emails to clinical nurses, and oncologists received summaries of patients’ symptom histories at each appointment.\textsuperscript{112}

PROMs can also provide invaluable information concerning emerging diseases, such as COVID-19. A 2020 study, for example, showed that seventy-six percent of patients who had been hospitalized for COVID-19 continued to
have abnormal PROMs three months after the onset of their initial symptoms.  
A third of these individuals reported “at least moderate impairment in major dimensions of quality of life.” Clinicians could learn a great deal about long COVID from such responses and use them as a guide for treating patients and alleviating their symptoms.

Public access to anonymized or summarized PROMs could enable patients to make more educated choices with respect to clinicians, medical facilities, and therapeutic options and to have realistic expectations about treatments and recovery. Individuals could select providers based on patient accounts of their post-treatment quality of life, such as whether they suffered incontinence or impotence after prostate surgery. Patients could also gain insight concerning others’ experiences during treatment and recovery, so that they know what to anticipate and can perhaps be less anxious or concerned.

Insurers may use PROMs to determine which health care providers and services to include in their networks. Insurers may also use PROMs to create profiles of high-risk patients that will incur high costs and to develop programs and interventions that might improve their health. While proactive interventions could help patients, one might worry that insurers will at the same time use PROM-based high-risk patient profiles as a justification for raising group premium rates.

Quality improvement initiatives can benefit from PROMs as well. Patients’ own perceptions regarding treatment outcomes and the care they receive are an important component of assessing the performance of health care providers.

114 Id.
115 Phillip Berry, Use Patient Reported Outcome Measures (PROMs) in Treatment of Long Covid, 373 BMJ n1260 (2021) (“If there was ever a condition where the use of PROMs should be prioritised, and traditional economic models challenged, it is post-covid-19.”).
117 Health Catalyst Editors, supra note 91.
118 Id.
119 Neubert et al., supra note 17, at 7.
120 Id. at 7-8.
121 How Insurance Rates Are Determined, OHIO DEP’T INS., https://insurance.ohio.gov/consumers/resources/how-insurance-rates-are-determined (last visited Apr. 20, 2023) (“All insurance companies use data and statistics to predict levels of risk for various individuals or groups. This risk calculation information is also used to develop rating plans.”).
providers and identifying areas for improvement.123

C. PROM Shortcomings and Concerns

Despite their many potential benefits, PROMs face strong critics who voice significant concerns about the tools and their implementation.124 PROM data can be particularly challenging because they consist of patients’ subjective assessments rather than objective medical test or examination results. This Part analyzes data quality and administrative challenges that constitute barriers to PROM implementation in both clinical and other contexts.

1. Data Quality

A large number of shortcomings can taint data quality and undermine their usefulness in clinical and other settings. This section analyzes the primary sources of data quality problems.

a. Reliability, Responsiveness, and Validity

To be useful, PROMs must be reliable, responsive, and valid.125 Not all PROMs are of equal quality.126 Reliability means the degree to which a measure is internally consistent and reproducible.127 Internal consistency refers to “correlation between different items in the measure.”128 If a survey is internally consistent, responders will answer items that test the same value similarly.129 For example, if the survey tests optimism, optimistic respondents will give high ratings to optimism indicators and low ratings to pessimism indicators throughout.130

123 Id. at 1027; Patient-reported Outcome Measures (PROMs), CANADIAN INSTITUTE HEALTH INFO., https://www.cihi.ca/en/patient-reported-outcome-measures-proms (last visited Dec. 11, 2022).
124 See Al Sayah et al., supra note 20, at 4-5.
126 Laith Alrubaiy et al., Assessing Patient Reported Outcome Measures: A Practical Guide for Gastroenterologists, 2 UNITED EUR. GASTROENTEROLOGY J. 463, 463 (2014) (“Not all PROM instruments currently used in research and clinical practice in gastroenterology have gone through a rigorous development methodology.”).
127 Ju & Tong, supra note 125, at 1883.
128 Id.
129 Id.
130 Fiona Middleton, The 4 Types of Reliability, Definitions, Examples, Methods, SCRIBBR
Reproducibility refers to a tool’s ability to generate the same result when it is used multiple times in similar circumstances. Thus, if a person takes a survey repeatedly without any change in health status, the individual’s responses should be very similar.

Responsiveness is a measure’s ability to discern outcome changes over time. This includes both changes in health status and changes in response to medical interventions. Responsiveness may be limited by a variety of factors, such as questions that offer too few answer choices and do not enable patients to indicate subtle alterations in their condition. Similarly, questionnaires that are administered too frequently may not give patients time to note meaningful differences in how they feel.

Validity is the extent to which a measure actually assesses what it claims to evaluate. This attribute can further be broken down into several categories. Criterion validity is the degree to which a measure relates to a gold standard, if one exists. Content validity refers to a measure’s ability to cover all dimensions that are important to the condition in question. Construct validity is the degree to which the measure evaluates the intended outcome (e.g., fatigue). External validity has to do with whether identified causal relationships can be generalized to other patients and circumstances. Internal validity is the extent to which observed results truly represent a causal relationship. Other forms of validity have also been recognized.

Experts use special techniques to validate survey instruments. For
instance, validity can sometimes be measured by comparing PROM scores to other related variables, such as clinical outcomes noted in EHRs. To illustrate, one study focused on sleep and compared self-reports to objective measures of sleep. It found that on average, participants slept six hours but reported sleeping 0.8 hours longer than they did. Analysts who are aware of such discrepancies might determine that a sleep PROM is not valid or adjust for the discrepancies when analyzing data.

Not all PROMs are validated with equal rigor. Furthermore, if a PROM is used for different purposes (e.g., clinical care, research, performance measures) or multiple populations (e.g., older patients, people with different underlying diseases), it may require different validations.

b. Response Shift and Response Bias

A phenomenon known as response shift can impact PROMs’ integrity as well. Response shift occurs because of a change in a responder’s perspective, for example, because of an alteration in the individual’s internal measurement standards or values. Therefore, response variations over time may reflect differences in a patient’s attitude rather than health status.

Response bias is yet another challenge. At times, individuals’ responses aim to reflect what they think the questioner wants to hear or what will impress the questioner rather than to be completely truthful. This bias may also be called “social desirability bias.” In the voting arena, for example, researchers have found that individuals untruthfully claim to have voted when they have not gone to the polls because they believe that is the correct and admirable answer to

143 Ju & Tong, supra note 125, at 1884.
145 Kate Churruca et al., Patient-Reported Outcome Measures (PROMs): A Review of Generic and Condition-Specific Measures and a Discussion of Trends and Issues, 24 HEALTH EXPECTATIONS 1015, 1021 (2021); Ju & Tong, supra note 125, at 1882.
146 Churruca et al., supra note 145, at 1021; John T. Farrar, Advances in Clinical Research Methodology for Pain Clinical Trials, 6 NATURE MED. 1284, 1289 (2010) (“[C]areful consideration should be given to each particular use, as subtle changes in the questions used or the population of interest can affect the results.”).
147 Cella et al., supra note 37, at 33.
148 Id.
150 Holbrook & Krosnick, supra note 149, at 37.
provide.\textsuperscript{151} Similarly, some physicians feel that patients’ answers are influenced by a desire to please the physician or gain some benefit by overstating or understating their symptoms.\textsuperscript{152}

c. PROM Selection

Determining which PROMs will best fit patients’ and clinicians’ needs is a challenging task.\textsuperscript{153} Given the breadth of choices, it is difficult to identify PROMs that are the most appropriate, valid, and illuminating for each condition, treatment, and practice.\textsuperscript{154} One group of experts suggests a general approach to PROM selection including:

1. Establish PROMs selection committee;
2. Identify the focus, scope, and type of PROM measurement;
3. Identify potential PROM(s);
4. Review practical considerations for each of the identified PROMs;
5. Review measurement properties of shortlisted PROMs;
6. Review patient acceptance of shortlisted PROMs;
7. Recommend a PROM(s); and
8. Pilot the selected PROM(s).\textsuperscript{155}

Other experts emphasize that selected PROMs must be reliable, responsive, and valid and must minimize the burdens of administering, answering (including for those with cultural and language barriers), reviewing, and incorporating PROM questionnaires into EHRs.\textsuperscript{156}

Beyond such general recommendations, however, there is no consensus as to

\textsuperscript{151} Id.
\textsuperscript{152} Turner et al., supra note 149, at 7.
\textsuperscript{154} Fatima Al Sayah et al., Selection of Patient-Reported Outcome Measures (PROMs) for Use in Health Systems, 5 J. Patient Rep. Outcomes 99, 99 (Supp. 2 2021); Ju & Tong, supra note 125, at 1882 (“[S]electing a robust and validated PROM from the plethora of available measures is challenging”); Tran et al., supra note 62, at 1724 (“The selection of a meaningful PRO instrument that provides accurate assessment and, at the same time, maximizes feasibility for clinical use is, thus, a challenge.”).
\textsuperscript{155} Al Sayah et al., supra note 154, at 99.
\textsuperscript{156} CELLA ET AL., supra note 36 at 38 (Table 4: Primary Criteria for Evaluating and Selecting Patient-Reported Outcome Measures (PROMS) for Use in Performance Measurement); see also supra Parts I.C.1.a (discussing reliability, responsiveness, and validity), I.C.1.e (discussing interpretability), and I.C.2 (discussing administrative challenges for respondents and health care providers).
PROM choices for particular conditions and no standardized PROM sets that are endorsed by professional organizations.\textsuperscript{157} Thus, researchers continue to explore and compare PROMs. The NIH states that its PROMIS project has generated over four-hundred publications.\textsuperscript{158} For example, one study compared PROMIS general health questionnaires for individuals who underwent carpal tunnel hand surgery with “the performance of region- and condition-specific PROMs such as the Michigan Hand Questionnaire (MHQ) and the Boston Carpal Tunnel Questionnaire (BCTQ).”\textsuperscript{159} It found that the PROMIS physical function PROMs were not useful for evaluating these surgical patients but the upper extremity and pain interference domains were.\textsuperscript{160}

PROMs can address generic health status or specific symptoms and conditions.\textsuperscript{161} Generic health status measures are broad and relevant to a variety of conditions, assessing degree of impairment and quality of life.\textsuperscript{162} Some experts recommend use of a combination of generic and condition-specific PROMs to obtain the most meaningful data.\textsuperscript{163}

d. Missing Data and PROM Timing

Some health care providers resist PROMs adoption because of concern about the accuracy and comprehensiveness of the data.\textsuperscript{164} While patients can be asked to complete PROMs, they are not forced to do so or to answer every query in the questionnaire.

Several studies highlight the problem of missing data.\textsuperscript{165} Some respondents

\begin{itemize}
\item \textsuperscript{157} Massachusetts Medical Society, \textit{supra} note 19, at 9.
\item \textsuperscript{159} David N. Bernstein et al., \textit{Responsiveness of the PROMIS and its Concurrent Validity with Other Region- and Condition-specific PROMs in Patients Undergoing Carpal Tunnel Release}, \textsc{477 Clinical Orthopedic Related Res.}, 2544, 2544 (2019).
\item \textsuperscript{160} Id. at 2545.
\item \textsuperscript{161} Ju & Tong, \textit{supra} note 125, at 1882.
\item \textsuperscript{162} Id. (providing the examples of “the 36-Item Short Form Health Survey (SF-36) and the Sickness Impact Profile).
\item \textsuperscript{163} Cella et al., \textit{supra} note 37, at 48.
\item \textsuperscript{164} Ryan P. Jacobson et al., \textit{Can Patient-Reported Outcomes Measurement Information System\textsuperscript{®} (PROMIS) Measures Accurately Enhance Understanding of Acceptable Symptoms and Functioning in Primary Care?}, \textsc{4 J Patient-Reported Outcomes} 1, 2 (2020).
\item \textsuperscript{165} See Fatima Al Sayah et al., \textit{supra} note 20 at 5; Olawale F. Ayilara, et. al, \textit{Impact of Missing Data on Bias and Precision when Estimating Change in Patient-Reported Outcomes from a Clinical Registry}, \textsc{17 Health & Quality Life Outcomes} 106, 107 (2019); Ethan Basch et al., \textit{Methods for Developing Patient-Reported Outcome-Based Performance Measures (PRO-PMs)}, \textsc{18 Value Health} 493, 501 (2015).
\end{itemize}
may skip questions or stop answering questionnaires prematurely because they are fatigued, confused, bored with the activity, or are called into their appointment and thus run out of time. In addition, some patients may choose not to respond to questionnaires or be unable to do so because of disabilities, language barriers, or lack of access to technology. One determinant of response rates may be the degree to which health care providers encourage patients to answer PROMs.

Response rate discrepancies can skew results in research studies or oversight initiatives that compare health care providers. Treatment outcomes of those who diligently employ PROMs, including with very sick patients, may look worse than outcomes from entities that are more lax about urging patients to fill out PROMs. At the same time, resource-poor organizations may not have the funds to implement PROMs and may not be included in clinical trials that solicit PROMs. If that is the case, little to no data would be gathered from important segments of the population that suffer socioeconomic disadvantages. The results of such research would be of questionable external validity and likely would not be generalizable to excluded populations.

Furthermore, vital details may be missing from PROM questionnaires. To illustrate, hip replacement surgery may not be as helpful for individuals who have other conditions that affect mobility, but questionnaires may not ask patients about these comorbidities. Cultural background may also influence how people answer PROMs, causing some people to interpret questions differently from others or to be reluctant to respond negatively about their health or treatment.

A further complication is that multiple choice questions, which are the format for many PROMs, may not capture all necessary information. A study relating to pain concluded that narrative descriptions of pain provided the best

166 See infra notes 199-202 and accompanying text (discussing survey fatigue).
167 See Basch et al., supra note 165, at 503; infra notes 193-198 and accompanying text (discussing various access barriers).
168 Basch et al., supra note 165, at 503.
169 Id. at 501.
170 See infra notes 211-12 and accompanying text (discussing implementation costs).
171 Rivera et al., supra note 10, at 1911 (“PRO research may not reflect the perspectives of underserved groups such as older individuals, socioeconomically disadvantaged populations, and racial and ethnic minority groups which could threaten the scientific validity of results.”).
172 Id.; see also Sharona Hoffman & Andy Podgurski, The Use and Misuse of Biomedical Data: Is Bigger Really Better?, 39 AM. J. L. MED 497, 521-23 (2013) (discussing selection bias, which occurs “when the subset of individuals studied is not representative of the patient population of interest.”). See supra note 121 and accompanying text for discussion of external validity.
173 Dawson et al., supra note 36, at 466.
174 Cella et al., supra note 37, at 39.
insight into patients’ experiences. If analysts do not collect appropriate auxiliary data about responders, they may not be able to contextualize and interpret PROM results correctly.

Using PROMs for purposes other than clinical care (such as research or FDA oversight) can be problematic for additional reasons as well. Survey responders may be a self-selected group that differs from non-responders in important ways, including health status, socioeconomic status, or other attributes. Individuals with low literacy or with language barriers are unlikely to complete PROMs. Individuals with cognitive decline or other intellectual or physical disabilities may also be unable to complete PROMs. If many potential participants face these barriers, PROM responders would not be representative of the relevant patient population at large (e.g., all patients with heart failure), and there will be significant gaps in the data collected.

Comparison and assessment of treatment outcomes may also be hindered by the timing of PROM collection. If different patients submit PROMs at different intervals following a medical intervention, they will not provide information that is easy to synthesize. Determining the appropriate point at which to solicit PROMs is itself complicated. Collecting PROMs too soon after an intervention may not provide complete data as to its impact, but collecting them after significant time has passed makes it difficult to attribute all reported phenomena to the intervention at issue rather than to other intervening factors.

176 Dawson et al., supra note 36, at 466.
177 Id. at 466.
178 CELLA ET AL., supra note 37, at 28-31.
179 Id. at 31; Jessica M. Kramer & Ariel Schwartz, Reducing Barriers to Patient-Reported Outcome Measures for People with Cognitive Impairments, 98 ARCHIVES PHYSICAL MED. & REHAB. 1705, 1705 (2017); Hahn Nguyen et al., A Review of the Barriers to Using Patient-Reported Outcomes (PROs) and Patient-Reported Outcome Measures (PROMs) in Routine Cancer Care, 68 J. MED. RADIATION SCI. 186, 188 (2021).
180 Id.; Walter F. Stewart et al., supra note 87, at 140 (discussing significant differences between respondents and non-respondents that resulted in differences between respondents and the total source population).
181 Al Sayah et al., supra note 154, at 5 (referring to “varying time points of PROM(s) measurement”); Dawson et al., supra note 36, at 466 (“Follow-up times should be the same for all patients in relation to the intervention or other key event.”).
182 Dawson et al., supra note 36, at 466.
184 Id.
e. Interpretability

In order to be useful, PROMs data must be available in formats that are accessible and easy to interpret. In many cases, clinicians do not know how to interpret PROMs and integrate them into patient care. Clinicians must easily be able to determine what changes in PROM scores mean and whether they indicate significant improvement or deterioration in a patient’s condition. As the National Quality Forum noted, “PROM scores and results must be integrated and viewed as actionable values upon a quick glance to successfully be incorporated into the clinical treatment plan.” Ideally, patients should also be able to view and understand their PROMs. Raw scores alone, without explanation and contextualization, might be of little value to providers and the patients they serve.

2. Administrative Challenges

Implementing a PROMs program can be challenging, even with high-quality PROMs. PROMs might face resistance from both patients and providers, as detailed below. Health care organizations should be keenly aware of these challenges and proceed with caution.

a. Patient Concerns

A variety of obstacles may hinder PROM completion. Patients may find that PROMs are collected through a platform that is inaccessible or difficult to use or that questions are hard to understand.

If patients are not given tablet computers at the clinician’s office or are not able to seek assistance while using them, they may ignore requests for PROMs. In one instance, Mass General Brigham found that when it discontinued tablet use because of COVID-19, significant racial disparities in the rate of PROM

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185 See Basch et al., supra note 165, at 503; NATIONAL QUALITY FORUM, supra note 7, at 20.
186 Nguyen et al., supra note 179, at 191.
187 NATIONAL QUALITY FORUM, supra note 7, at 20.
188 Id.; see Stehlik et al., supra note 79, at 815 (“It will also be important to determine the best approaches with which to share the results with the patients so that the understand the meaning of the scores and remain engaged in the process of serial PRO assessment.”).
189 NATIONAL QUALITY FORUM, supra note 7, at 20.
190 Id. (“Real-time information and interpretation must be available to accompany PROM scores.”).
191 Stine Thstrup Hansen et al., User Experiences on Implementation of Patient Reported Outcome Measures (PROMs) in a Haematological Outpatient Clinic, 4 J. PATIENT-REPORTED OUTCOMES 87, 96 (2020); MASS. MED. SOC., supra note 19, at 6-7.
192 MASS. MED. SOC., supra note 19, at 6-7.
completion developed.\textsuperscript{193} Patients identifying as Black provided PROMs at half the rate of White patients, and self-identifying Hispanics essentially did not fill them out at all, perhaps because of problems accessing computers and the Internet at home.\textsuperscript{194} On the other hand, patients with certain disabilities such as Parkinson’s disease may not have the dexterity to work with tablet computers in the clinic and might prefer to use their home computers.\textsuperscript{195} Others with learning disabilities, cognitive decline, or mental health conditions may not be able to complete PROMs on their own at all.\textsuperscript{196}

Survey fatigue is an additional concern.\textsuperscript{197} If patients are inundated with requests for PROMs, they may fill out questionnaires as quickly as possible without adequate thought, respond to only some of the queries, or ignore questionnaires altogether.\textsuperscript{198} According to one source, respondents generally stop answering questions after thirty queries.\textsuperscript{199} Thus, survey fatigue could contribute to low response rates, missing data, and poor data quality in PROMs.\textsuperscript{200} Note that in the research context, however, participants will have different expectations and may be willing to fill out longer PROMs.\textsuperscript{201}

b. Health Care Provider Concerns

Although PROMS can provide valuable information to health care providers,\textsuperscript{202} clinicians and staff members may find PROMs to be burdensome and unwelcome additions to their workloads.\textsuperscript{203} Burnout among physicians and other health care providers has received increasing attention in recent years.\textsuperscript{204} Already over-stretched providers might feel that the added tasks of processing and reviewing PROMs and responding to patient-reported concerns will be unmanageable for them.\textsuperscript{205} For example, in one study of an orthopedic medical

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\item \textsuperscript{193} Sisodia et al., supra note 74, at 2266.
\item \textsuperscript{194} Id.
\item \textsuperscript{195} \textit{Cella et al.}, supra note 37, at 31 (discussing functional abilities and PROMs completion).
\item \textsuperscript{196} Nguyen et al., supra note 179, at 188.
\item \textsuperscript{197} \textit{Mass. Med. Soc.}, supra note 19, at 10.
\item \textsuperscript{198} Vikas N. O’Reilly-Shah, \textit{Factors Influencing Healthcare Provider Respondent Fatigue Answering a Globally Administered In-App Survey}, 5 PEERJ 2017, at 2 (“Respondent fatigue, also known as survey fatigue, is a common problem in the collection of survey data.”).
\item \textsuperscript{199} Health Catalyst Editors, \textit{supra} note 91.
\item \textsuperscript{200} Rosaline de Koning et al., \textit{Survey Fatigue During the COVID-19 Pandemic: An Analysis of Neurosurgery Survey Response Rates}, 8 FRONTIERS SURGERY 1, 2 (2021).
\item \textsuperscript{201} \textit{Cella et al.}, supra note 37, at 42.
\item \textsuperscript{202} See \textit{supra} notes 92-102 and accompanying text.
\item \textsuperscript{203} \textit{Mass. Med. Soc.}, supra note 19, at 7.
\item \textsuperscript{204} Hoffman, \textit{supra} note 88, at 56.
\item \textsuperscript{205} Hansen et al., \textit{supra} note 191, at 96 (“[N]urses in this study did not use the PROM data
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center in Minnesota, researchers found that despite a high patient response rate (68-55%, depending on questionnaire timing), only one percent of physicians used PROM responses in their clinical work. Another study, which involved fourteen U.S. primary care clinics, found that while PROMs readily captured patients’ reports of their fall risks and urinary incontinence, this information was coded in EHRs only between three and fourteen percent of the time.

The cost of implementation is another concern. Institutions that adopt PROMs need information technology experts, personnel to maintain the program, and equipment such as tablet computers.

While PROMs can improve the physician-patient relationship by focusing doctors’ attention on patient concerns, there is also a risk that they will further diminish the time physicians spend face-to-face with patients. If patients are asked to complete PROMs during their appointments, they may have less time to speak with clinicians than they would otherwise. Patients may find this to be disappointing and frustrating because many prefer as much in-person communication with their providers as possible. Diminished opportunities for such communication can adversely affect the physician-patient relationship. It can also affect treatment outcomes if doctors have less time to examine the patient, speak with the patient, and provide explanations and advice.

Furthermore, health careemployers might require doctors to review PROMs online to obtain data about patients’ progress and complaints and then reduce the length of already rushed office visits. Many health care organizations pressure

and explained that lack of time required a focus on mandatory tasks related to treatment, control and documentation.

206 Heath, supra note 25.
207 Paul J. Barr et al., No Date for the PROM: the Association between Patient-Reported Health Events and Clinical Coding in Primary Care, 4 J. PATIENT-REPORTED OUTCOMES 17, 17 (2020).
208 See generally, Baumhauer et al., supra note 80.
209 MASS. MED. SOC., supra note 19, at 8 (discussing implementation costs and barriers to PROM adoption).
210 Id. at 9.
211 Evelyn Sharples et al., A Qualitative Exploration of Attitudes Towards the Use of Outcome Measures in Child and Adolescent Mental Health Services, 22 CLINICAL CHILD PSYCH. & PSYCHIATRY 219, 222 (2017) (noting that PROMs could take time away from psychotherapy session discussions).
214 Id.
215 Sharples et al., supra note 211, at 222; Hoffman, supra note 88, at 88 (noting that the
physicians to see more patients and generate more income, and they may consider PROMs a means to further those ends.

II. CLINICAL USE OF PROMS: PRIVACY AND MALPRACTICE IMPLICATIONS

Health care providers should recognize both the benefits and the shortcomings of PROMs when considering their implementation. In addition, clinical use of PROMs raises important legal questions. This Part provides an overview of two vital issues: privacy and malpractice concerns.

A. Privacy

Patients who complete PROMs may be concerned about the privacy of the information they provide. PROM questionnaires often ask patients to disclose information about their pain, ability to function, depression, anxiety, sexual satisfaction, and other sensitive matters. Once PROMs are completed, they are available electronically to multiple clinicians. If appropriate security measures are not implemented, they could also be inadvertently or intentionally disclosed to third parties or compromised through hacking. All identifiable medical information creates privacy concerns. But PROMS may intensify contemporary worries about privacy because of the volume and sensitivity of the collected data.

PROMs are covered by the HIPAA Privacy and Security Rules, whether or not they are integrated into patients’ EHRs. Both the Privacy and Security rules apply to health plans, health care clearinghouses, health care providers who transmit health information electronically for purposes of HIPAA-relevant transactions, and their business associates. Business associates would include

average primary care visit lasts only 15-20 minutes); Nguyen et al., supra note 179, at 188 (noting that a frequent complaint is “the time for patients to complete PROMs”).
216 Hoffman, supra note 88, at 90-91.
217 See supra Parts I.B and I.C.
218 Nguyen et al., supra note 179, at 191.
223 See supra notes 79-83 and accompanying text.
all entities that work with health care providers to collect, process, and store PROMs.\textsuperscript{225}

The Privacy Rule establishes that, in general, covered entities must obtain patients’ permission before disclosing their medical data to others.\textsuperscript{226} The HIPAA Security Rule requires administrative, physical, and technical safeguards to protect the confidentiality and integrity of electronic health information.\textsuperscript{227} Consequently, PROMs should not be disclosed to most third parties, such as employers or marketers, without patient consent and should be stored securely.

However, patients should be aware of significant exceptions to the HIPAA regulations. First, covered entities are permitted to divulge patients’ medical information without consent for purposes of treatment, payment, and health care operations.\textsuperscript{228} Thus, physicians can consult colleagues about patients, and facilities can send treatment information to insurers or use data for quality improvement activities without patients’ knowledge. In addition, the Privacy Rule lists a variety of other requests to which covered entities can respond without patient authorization, such as those made for purposes of public health activities, judicial and administrative proceedings, or law enforcement.\textsuperscript{229} There is no limit to the number of individuals who can view medical data for these permitted purposes.\textsuperscript{230} By some estimates, between 150 and 400 individuals view each patient’s EHR.\textsuperscript{231}

At the same time, the HIPAA Privacy Rule’s “minimum necessary” standard attempts to limit the extent of lawful disclosures. It provides that entities that disclose protected health information pursuant to a legitimate request “must make reasonable efforts to limit protected health information to the minimum necessary to accomplish the intended purpose.”\textsuperscript{232} There are certain exceptions to the

\begin{itemize}
  \item \textsuperscript{225} 42 U.S.C. §17934(a) (2018); 45 C.F.R. 160.103 (2022). Note that that the privacy of PROMs collected for non-clinical purposes (e.g., research) is also protected. The Privacy Act of 1974 prohibits federal agencies from disclosing individuals’ data without their consent unless particular exceptions apply. 5 U.S.C. § 552a(b) (2018). This safeguard would protect PROMs that are handled by the FDA and by federal programs such as Medicare or Medicaid. In addition, the federal research regulations, also known as the Common Rule, require that study participants provide informed consent for the use of any identifiable private information, which would include PROMs. 45 C.F.R. §46.116 (2022).
  \item \textsuperscript{226} 45 C.F.R. §§ 164.508–510 (2022).
  \item \textsuperscript{227} 45 C.F.R. §§ 164.302–318 (2022).
  \item \textsuperscript{229} 45 C.F.R. § 164.512 (2022).
  \item \textsuperscript{230} Id.; 45 C.F.R. § 164.506 (2022).
  \item \textsuperscript{231} Merida L. Johns, Privacy and Security of Health Information, in JEROME H. CARTER, ELECTRONIC HEALTH RECORDS: A GUIDE FOR CLINICIANS AND ADMINISTRATORS 298 (2008).
  \item \textsuperscript{232} 45 C.F.R. § 164.502(b) (2022).
\end{itemize}
minimum necessary requirement, such as disclosures to clinicians for treatment purposes and disclosures required by law.233

De-identified data constitute another major carve-out and are entirely exempt from HIPAA coverage.234 Therefore, they can be disclosed without patient authorization and stored in ways that do not comply with HIPAA Security Rule standards. It is thus possible that healthcare providers will disclose de-identified PROMs to third parties for research, marketing, or other purposes.

In theory, de-identification in compliance with HIPAA instructions thoroughly protects health information. However, there can never be a one-hundred percent guarantee that data will not be re-identified.235 In some cases, skilled attackers may be able to re-identify data by matching them to publicly available information, such as voter registration records or news stories about individuals with illnesses or injuries.236

Sadly, there is also no guarantee that HIPAA-covered data will not be compromised by hacking or other unlawful disclosures due to security lapses. According to one source, “[i]n 2022, an average of 1.94 healthcare data breaches of 500 or more records were reported each day.”237 But data breach risks are not unique to PROMs and are the cost of having so many data-rich medical resources.238

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234 45 C.F.R. § 160.103 (2022) (defining protected health information as “individually identifiable health information” that is electronically or otherwise transmitted or maintained). The HIPAA Privacy Rule provides detailed guidance regarding de-identification. It states that health information is de-identified if: (1) a qualified expert determines that there is only a “very small” risk that the data can be re-identified, and (2) the expert documents the analysis used to make this determination. 45 C.F.R. § 164.514(b)(1) (2022). As an alternative de-identification method, the HIPAA Privacy Rule lists eighteen items that should be removed to render data anonymized. These include names, geographic information, phone numbers, email addresses, Social Security Numbers, medical record numbers, and more. 45 C.F.R. § 164.514(b)(2)(i) (2022).
B. Medical Malpractice

For health care providers, a primary concern is medical malpractice. Both clinicians and health care entities can be held liable for malpractice. PROMs could potentially constitute a liability minefield for the medical community. Claims might arise because clinicians ignore PROMs that could influence important medical decisions, rely on them excessively, or fail to adopt them. This Section considers the malpractice implications of PROMs use.

1. Clinician Liability

Providers that ask patients to complete PROMs but do not review and react appropriately to them could potentially be vulnerable to liability if patients experience adverse events after reporting that their symptoms are not improving or are worsening. 239 For example, clinicians could potentially be sued if patients report suicidal ideation in PROM questionnaires and then, in the absence of intervention, commit suicide. 240 At the same time, liability could arise from inappropriate reliance on PROMs. To illustrate, psychiatrists may improperly fail to provide aggressive treatment for clinical depression if patients inaccurately score their depression as being low-grade in PROMs. Arguably, had the doctors had thorough face-to-face conversations with such patients, they may have discerned that their problems were more serious than the scores indicated. Similarly, surgeons may decide against needed surgery because patients do not report a high enough level of discomfort in PROMs. 241 In both cases, PROMs should be used as a tool, but fact finders may determine that clinicians should have also conducted other testing or had face-to-face conversations with patients. 242 Recall that patients sometimes experience survey fatigue and fail to answer questions carefully and thoughtfully. 243

A third possibility is that plaintiffs will bring claims against clinicians who failed to adopt PROMs that would have been helpful to their treatment. For example, PROMs concerning pain or mental health could be critical to medical decision making because these conditions are difficult to assess without patients’ subjective input. 244 Patients who feel they were injured because their doctor

239 Rivera et al., supra note 10, at 1922 (“If concerning data are not managed appropriately, those data could lead to suboptimal . . . care”).
240 See National Quality Forum, supra note 7, at 23.
241 See Safran & Higgins, supra note 74 (noting that PROMs can inform clinical decisions).
242 See Black, supra note 183, at 4 (“While some patients will not benefit from surgery, unfortunately they cannot necessarily be identified preoperatively using PROMs.”).
243 See supra notes 197-200 (discussing survey fatigue).
244 See supra notes 46-50, 87 and accompanying text.
failed to solicit their thorough input might sue for negligence. Medical malpractice plaintiffs suing health care professionals must establish the four elements of a negligence case. These are:

1) The defendant owes a duty of care to the plaintiff;
2) The defendant breached that duty through conduct that fails to meet the applicable standard of care;
3) The plaintiff suffered harm or injury; and
4) There is a causal link between the injury and the breach of duty.

Courts will need to grapple with the novel and complicated question of what the standard of care with respect to PROM use will be. The standard of care in each case is determined through an assessment of whether the defendant exercised “that reasonable degree of skill, knowledge and care ordinarily possessed and exercised by members of their profession under similar circumstances.” This assessment generally requires expert testimony. Fact-finders, therefore, should not judge clinicians based on whether they provided optimal care, but rather, on whether they provided reasonably competent care in light of the particulars of the specific case. The standard of care is to be “objectively determined by reference to the availability of medical and practical knowledge which would be brought to bear in the treatment of like or similar patients under like or similar circumstances by . . . physicians in the same field, given the facilities, resources and options available.”

Because PROMs are not yet a routine part of patient care, there is no clear standard of care concerning their use. Whenever emerging technologies begin to

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246 Id. at 1534; McDowell v. Brown, 392 F.3d 1283, 1295 (11th Cir. 2004); Hanson v. Grode, 90 Cal. Rptr. 2d 396, 400 (Cal. Ct. App. 1999).
248 Scott, 180 Cal. Rptr. 3d 479 at 498-99.
249 See supra note 242.
250 Hall v. Hilbun, 466 So.2d 856, 872 (Miss. 1985).
251 See supra notes 73-74 and accompanying text (discussing the limited extent to which PROMs have been adopted in the United States).
With respect to claims that clinicians ignored information in PROMs, clinicians will likely argue that it is impossible to review and respond to all PROMs and that doing so should not be considered the standard of care. Arguably, instead of assuming that providers are scrutinizing all PROMs, patients who require attention should call the office. According to one study, family physicians have a mean of approximately 2300 patients each, and consequently, reviewing PROMs could be an overwhelming and unmanageable task unless it is largely automated, as suggested later in this Article.

In contrast, patients will posit that there is no point in taking the time to complete PROMs if clinicians simply ignore them. Arguably, requests for PROMs imply that clinicians will read and respond to them.

While there is currently no precedent involving PROMs, a few cases concerning physicians’ communication with patients suggest that an argument for PROM-related liability may be viable. In Gaffney v. Giles, a Louisiana court of appeals upheld a lower court’s determination that a physician’s failure to return a patient’s phone calls constituted a breach of the standard of care. The patient was awarded damages because his condition deteriorated as he tried and failed to reach his doctor. In an older case, St. Charles v. Kender, the court held that an HMO patient who suffered a miscarriage could assert a breach of contract claim against a doctor who ignored her phone calls. By extension, if patients are led to believe that health care providers will review their PROMs, plaintiffs might successfully bring medical malpractice claims based on clinicians’ failure to respond to alarming PROM information.

Claims that clinicians did the opposite and relied excessively on PROMs in making diagnostic or treatment decisions and neglected to investigate other indicators would be treated like all claims relating to erroneous medical decision making.
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making. Courts would need to assess the degree to which reliance on the tool of PROMs to the exclusion of other tools complies with the standard of care.259

Interesting questions could also arise with respect to patients who fail to fill out PROM questionnaires or do not answer all questions after being informed that clinicians rely on PROMs for decision making purposes. Would sending reminders to patients or incentivizing survey completion become part of the standard of care? Would courts apply the doctrine of contributory negligence or comparative fault to patients who do not complete PROMs after being told of their importance?260

Claims that plaintiffs were injured because physicians failed to implement PROMs and thereby to gather vital information would be assessed in the same manner as claims regarding other new medical technologies. For example, in Washington v. Washington Hospital Center, the court ruled that reasonable jurors could find that the standard of care in 1987 required hospitals to use end-tidal carbon dioxide monitors for anesthetized patients during surgery.261 It thus upheld a jury verdict for a patient who suffered permanent brain injuries because of oxygen deprivation.262

It is possible that malpractice concerns will accelerate widespread adoption of PROMs.263 If courts come to expect that health care providers collect PROMs and integrate them into clinical decision making, providers will be more likely to adopt PROMs quickly to avoid deviating from the standard of care.

Ultimately, the courts will have to determine what the standard of care is in the context of PROMs.264 If litigation is brought by plaintiffs who feel they were injured and the harm is linked to PROMs, case law will help establish the legal standards for managing this data tool.

259. See George Maliha et al., Artificial Intelligence and Liability in Medicine: Balancing Safety and Innovation, 99 MILBANK Q. 629, 632 (2021) (discussing the use of artificial intelligence and machine learning and noting that a “physician who in good faith relies on an AI/ML system to provide recommendations may still face liability if the actions the physician takes fall below the standard of care and other elements of medical malpractice are met.”).


262. Id. at 177.

263. Ryan Abbott, The Reasonable Computer: Disrupting the Paradigm of Tort Liability, 86 GEO. WASH. L. REV. 1, 12 (2018) (“In its quest to reduce accidents, tort law can either accelerate the introduction of new technologies, as was the case with the use of glaucoma testing and pulse oximeters, or it can discourage the use of new technologies, as is usually the case where the standard of care is based on custom.”).

264. See Sokol & Molzen, supra note 252, at 469 (“The reality that the health care industry has not uniformly embraced information technology will cause courts to reexamine the standard of care and how to shape it.”)
2. Liability of Health Care Entities

Aggrieved plaintiffs may wish to assert medical malpractice claims not only against clinicians, but also against health care entities. First, plaintiffs can sue health care organizations such as hospitals and clinics for the negligence of their employees, and, under agency principles, employers can be held vicariously liable for their employees’ acts.\textsuperscript{265} Thus, if courts determine that clinicians can be liable for failing to react appropriately to information captured in PROMs, failing to adopt PROMs, or over-relying on PROMs, patients could use vicarious liability theories to sue health care entities. When clinicians are employees of the entity, plaintiffs can allege actual agency,\textsuperscript{266} and if clinicians are independent contractors, claimants may attempt to prove apparent agency.\textsuperscript{267}

Alternatively, plaintiffs may wish to sue health care facilities directly if they believe entities have mishandled PROMs, have faulty PROM policies, or do not enforce policies appropriately. The corporate negligence doctrine, which is recognized by most states,\textsuperscript{268} establishes that health care entities are liable for failing to provide treatment that meets the standard of care.\textsuperscript{269} Hospitals (and other medical entities) have the following four duties:

- (1) a duty to use reasonable care in the maintenance of safe and adequate facilities and equipment;
- (2) a duty to select and retain only competent physicians;
- (3) a duty to oversee all persons who practice medicine within its walls as to patient care; and
- (4) a duty to formulate, adopt and enforce adequate rules and policies to ensure quality care for the patients.\textsuperscript{270}

To establish a prima facie case of corporate negligence, plaintiffs must show (1) that the hospital deviated from the standard of care; (2) that the hospital has actual or constructive knowledge of the flaws or procedures that caused the injury; and (3) that a causal link exists between the conduct in question and the

\textsuperscript{265} CLARK, supra note 260, at 231-32.
\textsuperscript{266} Scott v. SSM Healthcare St. Louis, 70 S.W. 3d 560, 566-67 (Mo. Ct. App. 2002).
\textsuperscript{267} See Burless v. West Virginia U. Hosp., Inc., 601 S.E.2d 85, 92-96 (W. Va. 2004). To prevail on a theory of apparent agency, a plaintiff must establish two elements: (1) The hospital either committed an act that would cause a reasonable person to believe that the physician in question was an agent of the hospital, or, by failing to take an action, created a circumstance that would allow a reasonable person to hold such a belief, and (2) the plaintiff relied on the apparent agency relationship.
\textsuperscript{270} Id.
harm. Plaintiffs could sue health care organizations for mishandling or neglecting PROMs if they feel that fault lies with the entity itself. Claims relating to failure to review and respond to PROMs or excessive reliance on PROMs could arguably fall under the duty to oversee personnel properly or to have suitable rules and policies. Failure to implement a PROMs program in the first place (if doing so has become the standard of care) could potentially be considered a breach of the latter duty as well as the duty to maintain adequate equipment.

III. PROM USE IN RESEARCH AND FDA OVERSIGHT

PROMs can serve many purposes outside the clinical setting. They are frequently employed in research studies to obtain quality of life data directly from patients. The FDA has also begun to accept PROMs for certain oversight functions. This section critiques PROM use in research and FDA oversight.

A. Incorporating PROMs into Research

Many researchers are enthusiastic about incorporating PROMs into research. They note that patients have much to contribute in assessing their own symptoms and adverse events and that PROMs are an important adjunct to clinician-reported outcomes. To that end, the National Cancer Institute developed the Patient-Reported Outcomes version of the Common Terminology Criteria for Adverse Events (PRO-CTCAE). The PRO-CTCAE contains “124 items representing 78 symptomatic toxicities” and is designed to be a companion to the physician-reported CTCAE.

There is also a pediatric module for self-reporting by minors who are seven to seventeen years old ((Ped-PRO-CTCAE®) and a module for caregivers of minors who cannot self-report (Ped-PRO-CTCAE®[Caregiver]), and all versions are publicly available.

PROMs may be particularly useful for comparative effectiveness research in which different

272 Thompson, 591 A.2d at 707.
273 Id.
274 Kluetz et al., supra note 87, at 743.
275 Id. at 743.
278 NCI, supra note 276.
medical interventions are directly compared to determine which are of greatest benefits or harm to particular patients.279

Nevertheless, some experts are highly critical of the way PROMs are currently used in research.280 According to one article, thousands of new PROM questionnaires are produced, many of which are used for only one study, and they have little impact on medical research.281 The authors note that while PROMs are very widely employed in studies, their results are rarely reported in publications, and when they are discussed, there is often no comparison of score changes between study arms.282 This article is not alone in noting that PROM data are often neglected in research publications.283

Others express additional concerns. One international consortium developed recommendations for identifying suitable statistical methods for PROM analysis, managing missing data, and other challenges.284 However, it noted that there is “no consensus on standards and unclear guidelines on how to analyse and interpret PRO data” collected in cancer clinical trials.285 It concluded that it is critical that robust findings “be derived consistently across studies to yield meaningful results” and that a great deal of work has yet to be done to finetune PROM standards for cancer studies.286

B. PROM Use in FDA Drug and Device Assessment and Labeling

At their best, patients’ own voices, expressed through PROMs, can play a vital role in research and regulatory oversight. PROMs are increasingly used for FDA regulatory purposes.287 The 21st Century Cures Act established a program

279 Hostetter & Klein, supra note 14; Albert W. Wu et al., Adding the Patient Perspective to Comparative Effectiveness Research, 29 HEALTH AFFS. 1863, 1863 (2010).
281 Id.
282 Id.
283 Rivera et al., supra note 10, at 1911 (“A 2019 evaluation of 160 cancer trials showed nearly 50,000 participants were included in studies that failed to publish their PRO data”); Thi Xuan Mai et al., Utility of the Patient-Reported Outcomes Measurement Information System (PROMIS) to Measure Primary Health Outcomes in Cancer Patients: A Systematic Review, 29 SUPPORTIVE CARE CANCER 1723, 1736 (2021) (“Non-reporting of PRO results is prevalent, and this devalues the considerable contribution of participants who spend time and effort to provide their PRO information.”).
284 Carneel Coens et al., International Standards for the Analysis of Quality-of-Life and Patient-Reported Outcome Endpoints in Cancer Randomised Controlled Trials: Recommendations of the SISAQOL Consortium, 21 LANCET e83, e83 (2020).
285 Id.
286 Id. at e94.
under which the FDA is to evaluate the use of real world evidence to support new uses of approved drugs and to help conduct post approval studies. The Act defines “real world evidence” as “data regarding the usage, or the potential benefits or risks, of a drug derived from sources other than traditional clinical trials.” This data includes information that is generated by patients themselves.

In 2022 guidance regarding medical devices, the FDA stated that use of patient-reported outcomes (PROs) is voluntary, and thus they are not currently required for any FDA purpose. However, the FDA supports and recommends PROMs in many circumstances.

Under the Medical Device Development Tools program, PROMs qualify for use in the development and assessment of medical devices. PROM-based research can be valuable for purposes of designing and developing devices that will best serve patient needs. In addition, PROMs can significantly contribute to post market surveillance, providing data about treatment success or failure after products are deployed in clinical care.

If developers wish to use PROMs to meet regulatory requirements such as medical device evaluation, the FDA will determine what validity evidence is needed to render them “fit-for-purpose.” In addition, the FDA runs the Clinical

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290 Real-World Evidence, U.S. FOOD & DRUG ADMIN., https://www.fda.gov/science-research/science-and-research-special-topics/real-world-evidence (last viewed Oct. 19, 2022) (noting that “real-world data” can include data gathered from digital health technologies, which could include PROMs data).
292 Id.
294 FDA 2022, supra note 291, at 3-4.
295 Id. The FDA acknowledges that not all side effects of drugs and devices can be discerned “based on preapproval studies involving only several hundred to several thousand patients.” Consequently, it has post marketing surveillance and risk assessment programs designed to identify adverse events that did not manifest before a drug or device was approved. Postmarket Surveillance Programs, U.S. FOOD & DRUG ADMIN., https://www.fda.gov/drugs/surveillance/postmarketing-surveillance-programs (last visited Apr. 20, 2020).
296 FDA 2022, supra note 291, at 4-5 (“By assessing the similarities and differences between the population in the clinical study and in the development of the PRO instrument, FDA can determine whether the PRO instrument is fit-for-purpose.”). “Fit-for-Purpose” is defined as a “conclusion that the level of validation associated with a medical product development tool is sufficient to support its context of use.” Id. at 12.
Outcome Assessment (COA) Qualification Program.\textsuperscript{297} The FDA explains that “COA qualification represents a conclusion that within the stated context of use, results of assessment can be relied upon to measure a specific concept and have a specific interpretation and application in drug development and regulatory decision-making.”\textsuperscript{298}

There is no consensus as to which PROMs should be used for FDA approval.\textsuperscript{299} The FDA offers several key principles that should guide incorporation of PROMs into device evaluation. They are:

1. Establish and define the concept of interest (COI) the PRO instrument is intended to capture;
2. Clearly identify the role of the PRO (e.g., primary, secondary, ancillary, effectiveness, safety) in the clinical study protocol and statistical analysis plan;
3. Provide evidence showing that the PRO instrument reliably assesses the COI; and
4. Effectively and appropriately communicate the PRO-related results in the [product] labeling to inform healthcare provider and patient decision making.\textsuperscript{300}

Drug and device “labeling” includes not only labels pasted on containers, but also other written, printed, or graphic material on items, their containers, wrappers, or other matter that accompany them.\textsuperscript{301} In 2009 the FDA issued guidance that describes how the FDA reviews and assesses PROM instruments that are used to develop evidence for claims in medical product labeling.\textsuperscript{302} According to one source, approximately twenty-six percent of new drugs approved from 2016 to 2020 included PRO-related statements in labeling.\textsuperscript{303}

The FDA is developing further guidance regarding PROM use. These include draft guidance on “Core Patient-Reported Outcomes in Cancer Clinical Trials”\textsuperscript{304} and a “Patient-Focused Drug Development Guidance Series for

\textsuperscript{298} Id.
\textsuperscript{299} Warsame & D’Souza, supra note 19, at 2291.
\textsuperscript{300} FDA 2022, supra note 291, at 4.
\textsuperscript{302} U.S. FOOD & DRUG ADMIN., GUIDANCE FOR INDUSTRY PATIENT-REPORTED OUTCOME MEASURES: USE IN MEDICAL PRODUCT DEVELOPMENT TO SUPPORT LABELING CLAIMS (2009), https://www.fda.gov/media/77832/download.
\textsuperscript{303} Gnanasakthy et al., supra note 12, at 650.
\textsuperscript{304} U.S. FOOD & DRUG ADMIN., CORE PATIENT-REPORTED OUTCOMES IN CANCER CLINICAL TRIALS: GUIDANCE FOR INDUSTRY, DRAFT GUIDANCE (June 2021), https://www.fda.gov/media/149994/download.
Enhancing the Incorporation of the Patient’s Voice in Medical Product Development and Regulatory Decision Making.” Consequently, it is not inconceivable that the FDA will ultimately require PROM use for some regulatory purposes once it refines its approach to this tool.

IV. PROM USE FOR PERFORMANCE MEASUREMENT AND INSURANCE COVERAGE

Policy makers in the U.S. have long expressed a commitment to achieving value-based care that rewards health care providers for high-quality services and outcome improvements. Such a system requires the ability to measure quality of care and health outcomes accurately, and, according to some advocates, PROMs are a critical component of these measurements. Thus, the concept of patient-reported outcome performance measures (PRO-PM) has emerged. A PRO-PM is a “performance measure that is based on patient-reported outcomes assessed through data, often collected through a PROM and then aggregated for . . . [a] healthcare entity.” CMS endorses the use of PRO-PMs for performance improvement and accountability purposes.

Under the CMS Quality Payment Program (QPP), created by the Medicare Access and CHIP Reauthorization Act of 2015, CMS rewards clinicians for high performance levels and reduces payments for sub-standard performance. Clinicians have two QPP options: 1) the Merit-based Incentive Payment System (MIPS) or 2) Advanced Alternative Payment Models. PROMs are a priority measurement category for MIPS. Furthermore, CMS is incorporating PRO-
PMs into its Meaningful Measures 2.0 initiative, which aims to streamline quality measures and “promote innovation and modernization of all aspects of quality.”  

Additionally, CMS and the National Quality Forum have undertaken an initiative called “Building a Roadmap from Patient-Reported Outcome Measures to Patient-Reported Outcome Performance Measures.” The project aims to provide guidance regarding PRO-PMs that will be used in CMS accountability programs such as alternative payment models and was scheduled for completion in late 2022. To date, however, PRO-PMs have constituted only five percent of the measures that were used by federal programs and endorsed by the National Quality Forum.

Private insurers have used PROMs as well. In 2013, Blue Cross Blue Shield of Massachusetts (BCBSMA) and providers participating in its Alternative Quality Contract (AQC) program collaboratively selected conditions for initial PROM implementation. The chosen conditions were depression and knee/hip pain, which had well-recognized, validated PROMs. BCBSMA paid providers to participate in the PROM program, and, during 2013-2015 participation was voluntary. In 2016, BCBSMA transitioned to requiring participation from AQC providers, expanded the number of conditions for PROM adoption, and continued to pay providers for participation. It did not

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make any payment adjustments based on performance as reflected in PROM scores so that clinicians would not be concerned that participation could lead to financial penalties. BCBSMA plans to roll out its PROM program in three phases: 1) paying providers for PROM adoption, data sharing, and learning; 2) using collected data to inform clinical decision making; and 3) eventually, using collected data to adjust payment for performance outcomes and promote accountability. Some insurers may also use PROMs to determine which physicians should be included in their networks.

As noted earlier, advocates argue that validated PROMs that are implemented correctly can have a positive impact on clinical decision making and cost savings, which would also benefit health care payers. For example, BCBSMA found that patients whose PROMs indicated that they were high functioning at baseline (approximately eight percent of its cohort) did not benefit from hip and knee replacement surgery and could feel worse because of the procedure. Thus, PROM assessment could spare some patients from undergoing a painful and expensive surgery and recovery period at the same time that it spares insurers from paying for unnecessary procedures.

Other commentators caution against use of PROMs for insurance purposes at this time. The American College of Physicians (ACP) asserts that more data are needed to establish that PRO-PMs in truth enhance quality of care and can be used to compare clinician performance accurately. The ACP notes that outcomes can be affected by factors that are out of the physicians’ control, such as patient compliance with treatment protocols or access to family and other support systems. Moreover, some physicians could wrongly be penalized because they treat very sick patients or members of vulnerable communities whose outcomes are likely to be suboptimal even if they receive excellent care. The Special Needs Plan Alliance studied use of the Medicare Health Outcome Survey and found that it was problematic for special needs plans because “they serve diverse, low-income, disabled, and chronic care, complex, or advanced-

324 Id.
325 MASS. MED. SOC., supra note 19, at 6.
326 Neubert et al., supra note 17, at 7.
327 Id. at 5 (noting that preliminary European studies show that “PROMs do support more evidence-based decision-making and value-based care delivery”); see supra Part I.B.
328 Safran & Higgins, supra note 74.
329 Id.
330 Qaseem et al., supra note 317, at 1161.
331 Id. at 1161; see also Holmes et al., supra note 44, at 254 (“There is no definitive evidence as to whether PROMs have an impact on health status, with only some studies showing significant differences.”).
332 Qaseem et al., supra note 317, at 1161.
333 Id.
illness populations.” Skilled analysts would need to adjust for such factors.

The Massachusetts Medical Society warns against unintended consequences of using PROMs for performance measurement purposes. If reimbursement were to depend on PROM scores, some medical decisions might be driven by health care providers’ desire to maximize their earnings, and such decisions may not always be in patients’ best interest. Thus, clinicians may opt for the least uncomfortable diagnostic tests so that patients do not report increased anxiety or pain, even if more uncomfortable tests may have been better diagnostic tools. This is not merely a hypothetical concern, as clinicians frequently respond to incentives despite adverse effects on patient care. For example, a United Kingdom initiative that linked financial rewards to swift access to care may have eroded continuity of care, which is important for many patients with complex needs. Health care organizations were incentivized to furnish access to any provider as quickly as possible, so patients were given appointments with clinicians who knew nothing about them.

PROMs require extensive validation, and their use requires sound risk adjustment strategies. PROM programs that are poorly implemented by insurers could penalize clinicians that are providing the best care possible under the circumstances. They could also deprive patients of needed treatments because of erroneous PROM-based assumptions about their functionality or discomfort. Both the ACP and the Massachusetts Medical Society caution that it is premature to rely on PROMs for insurance purposes.

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335 MASS. MED. SOC., supra note 19, at 9.

336 Id.


338 MASS. MED. SOC., supra note 19, at 8.

339 Id.

340 Qaseem et al., supra note 317, at 1161-62. See also supra notes 342-344 and accompanying text (discussing validation). Risk adjustment can be defined as “A statistical process that takes into account the underlying health status and health spending of the enrollees in an insurance plan when looking at their health care outcomes or health care costs.” Risk Adjustment, HEALTHCARE.GOV, https://www.healthcare.gov/glossary/risk-adjustment/ (last visited Dec. 11, 2022).

341 See Black et al., supra note 183, at 3 (cautioning against using PROMs to crudely ration care and relating that UK PROM data was “misinterpreted as showing that 20,000 hernia and varicose vein operations and 16,000 hip and knee replacements each year should not take place.”).

342 MASS. MED. SOC., supra note 19, at 10 (“[S]ince PROMs implementation remains in its infancy . . . PROMs results should not be used to compare providers or outcomes for payment”); Qaseem et al., supra note 317, at 1162 (advising caution “until PRO-PMs are developed in a rigorous manner and physicians can seamlessly integrate patient-reported data collection into
V. RECOMMENDATIONS

In an ideal world, physicians or other skilled clinicians would have ample
time to speak with patients about their symptoms, complaints, and medical
progress. But medicine is all too often a profit-driven industry, pressuring
providers to limit the duration of patient encounters and pack their schedules. In light of these realities, PROMs can potentially fill important data gaps. But much work remains to be done to address considerable PROM deficiencies and concerns that can lead to liability. Whether these shortcomings can be consistently overcome is still in question. This part formulates recommendations for technical and administrative improvements as well as legal and policy interventions. As PROM programs are increasingly adopted by health care providers and regulators, it is vital to ensure that they are appropriately implemented and do not have unintended adverse consequences for patients and clinicians.

A. Technical and Administrative Recommendations

Many experts have offered recommendations to assist health care providers and researchers in establishing PROM programs. Thoughtful selection and implementation of PROMs by qualified experts should provide a degree of protection against liability risks and render PROMs better fit for research, use by the FDA and CMS, and other purposes.

1. PROM Selection

Selecting appropriate PROMs for inclusion in questionnaires can be very challenging and is vital to the effectiveness of any PROM initiative. Hundreds of potentially relevant PROMs are often available, and their quality may be difficult to discern. Those tasked with PROM selection (called “implementers” below) must carefully contemplate what they hope to achieve, including what specific information they wish to gather and how it will be used. PROM selection practice.

343 Hoffman, supra note 88, at 87-92.
344 See supra Part I.B (discussing PROM benefits).
345 Danny Mou et al., Impetus of US Hospital Leaders to Invest in Patient-Reported Outcome Measures (PROMs): A Qualitative Study, 12 BMJ OPEN 1 (2022) (“[H]ospital leaders feel a strong moral imperative to collect PROMs [which] can be used to demonstrate the value of their services to payors and patients); supra Parts III and IV.
346 See, e.g., Al Sayah et al., supra note 20, at 3-4; MASS. MED. SOC., supra note 19; NATIONAL QUALITY FORUM, supra note 7, at 5-23; Rivera et al., supra note 10.
347 Churruca et al., supra note 145, at 1021.
348 Churruca et al., supra note 145, at 1021.
requires a literature review and thorough research.\textsuperscript{349} Below are several key components of a successful selection process.

a. Obtain Stakeholder Input

PROMs selection requires input from diverse stakeholders.\textsuperscript{350} These can include clinicians, patients, computer system administrators, technical experts, family members, caregivers, and others.\textsuperscript{351} It may be prudent to establish a formal selection committee to ensure that such input is obtained.\textsuperscript{352} It is particularly important to engage with patients to determine whether they will view PROMs favorably.\textsuperscript{353} Patients should be asked whether they find proposed PROMs to be accessible, understandable, or offensive in any way.\textsuperscript{354} Additionally, academic and industry researchers should continue to examine how PROMs can best respond to patient needs, abilities, and preferences.\textsuperscript{355}

b. Select PROMs that Align with Goals

Implementers should identify the “focus, scope, and type” of PROMs that will support both treatment of individual patients and institutional goals.\textsuperscript{356} For example, a key decision is whether to use generic PROMs, condition-specific PROMs, or a combination of both.\textsuperscript{357} PROM selection should be informed by a clear understanding of what outcomes clinicians or researchers wish to measure.\textsuperscript{358} Institutional goals might include performance evaluation, health care

\begin{thebibliography}{99}
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\bibitem{footnote} \textsuperscript{349} \textit{National Quality Forum}, \textit{supra} note 7, at 7.
\bibitem{footnote} \textsuperscript{350} Sivan et al., \textit{supra} note 35, at 1.
\bibitem{footnote} \textsuperscript{351} Al Sayah et al., \textit{supra} note 20, at 3; \textit{National Quality Forum}, \textit{supra} note 7, at 9.
\bibitem{footnote} \textsuperscript{352} Al Sayah et al., \textit{supra} note 20, at 3.
\bibitem{footnote} \textsuperscript{353} Id. at 102; CMS 2022, \textit{supra} note 6, at 7; Rivera et al., \textit{supra} note 10, at 1915 (discussing the need for patient input regarding PROMs that will be used in research).
\bibitem{footnote} \textsuperscript{354} Al Sayah et al., \textit{supra} note 20, at 4.
\bibitem{footnote} \textsuperscript{355} San Keller et al., \textit{Selecting Patient-Reported Outcome Measures to Contribute to Primary Care Performance Measurement: A Mixed Methods Approach}, 35 J Gen. Internal Med. 2687, 2694 (2020) (discussing the need for future research); Brocha Z. Stern, \textit{Clinical Potential of Patient-Reported Outcome Measures in Occupational Therapy}, 76 Am. J. Occupational Therapy 1 (2022) ("Looking forward, clinicians should collaborate with multiple stakeholders, from patients to health system leaders, to meaningfully and equitably integrate PROMs into routine clinical care. Researchers should evaluate best practices for selecting, interpreting, implementing, and applying PROMs to maximize both individual-level and aggregate-level value.").
\bibitem{footnote} \textsuperscript{356} Id. at 3.
\bibitem{footnote} \textsuperscript{357} Churruca et al., \textit{supra} note 145, at 1021. See also \textit{supra} notes 161-163 and accompanying text (discussing generic and condition-specific PROMs).
\bibitem{footnote} \textsuperscript{358} \textit{National Quality Forum}, \textit{supra} note 7, at 10; Rivera et al., \textit{supra} note 10, at 1913 (discussing the importance of clear research questions, rationales for PROM assessment, objectives, and hypotheses).
\end{thebibliography}
delivery improvements, and treatment cost analyses.

c. Select PROMs that Meet Practical Needs

Practical considerations are of vital importance.359 PROM questions should be written in clear, accessible language, and for some patient populations, multiple languages will be needed.360 Some practices or research projects include many patients with cognitive decline and, to the extent possible, their PROM queries should be appropriate for such patients.361

In addition, patients may have limited attention spans and tolerance for answering queries or may have impairments that hinder their response abilities, so PROM questionnaires must not be excessively lengthy.362 Computer adaptive technology can be helpful in limiting patient burden because it tailors questionnaires to particular patients based on their responses.363 For example, to avoid survey fatigue, PROMIS often limits the number of queries to four to six when computer adaptive technology is used.364 However, implementers must also ensure that thoroughness is not sacrificed for the sake of brevity.

Another practical consideration is cost. Implementers must determine whether PROMs will strain their budget and may opt for PROMs that are publicly available rather than those that require licensing fees.365

d. Evaluate PROM Attributes Prior to Selection

Implementers must examine the psychometric properties of proposed PROMs.366 Implementers should look for evidence of reliability, validity, responsiveness, interpretability, and appropriateness for particular patient

populations and diseases.\textsuperscript{367} To that end, PROMs endorsed by PROMIS are often a good choice.\textsuperscript{368} In addition, implementers should verify that selected PROMs have been used successfully by other entities in similar circumstances.\textsuperscript{369} Further guidance for PROM review is found in a variety of resources, two of which are the “COSMIN Guideline for Systematic Reviews of Patient-Reported Outcome Measures”\textsuperscript{370} and the Terwee criteria for measurement properties of health status questionnaires.\textsuperscript{371}

e. Conduct a Pilot Program

Prior to full-scale launch of PROMs, implementers should conduct a pilot program to identify any pitfalls that were missed during the selection process.\textsuperscript{372} The pilot program should evaluate how easily PROMs can be integrated into clinical workflow and how well they serve their intended purposes.\textsuperscript{373}

2. \textit{PROM Implementation}

Implementing PROMs can be no less challenging than selecting them. The following are several essential components of the implementation process.

a. Cultivate Stakeholder Buy-In

Implementers should build enthusiasm for PROMs among all stakeholders, including providers, staff, patients, and technical experts.\textsuperscript{374} It is particularly important to have one or more clinician champions to promote appreciation of

\begin{itemize}
  \item \textsuperscript{367} Al Sayah et al., \textit{supra} note 20, at 4; Basch et al., \textit{supra} note 165, at 500-01. See \textit{supra} Parts I.C.1.a and I.C.1.e for a discussion of reliability, validity, responsiveness, and interpretability.
  \item \textsuperscript{368} Evans et al., \textit{supra} note 59, at 350 (noting that PROMIS is the gold-standard for PROMs); \textit{MAGS. MED. SOC.}, \textit{supra} note 19, at 6; Wong & Meeker, \textit{supra} note 83, at 1 (finding that PROMIS physical health computerized adaptive test domains “are reliable, responsive, and interpretable in most contexts of patient care throughout all orthopaedic surgery subspecialties.”).
  \item \textsuperscript{369} Basch et al., \textit{supra} note 165, at 500; \textit{NATIONAL QUALITY FORUM}, \textit{supra} note 7, at 9.
  \item \textsuperscript{370} C. A. C. Prinsen et al., \textit{COSMIN Guideline for Systematic Reviews of Patient-Reported Outcome Measures}, 27 QUALITY LIFE R SCH. 1147, 1148-56 (2018).
  \item \textsuperscript{372} Al Sayah et al., \textit{supra} note 20, at 4; CMS 2022, \textit{supra} note 6, at 6.
  \item \textsuperscript{373} Al Sayah et al., \textit{supra} note 20, at 4 (“It is important to test these tools with the population on which the measure focuses.”).
  \item \textsuperscript{374} \textit{NATIONAL QUALITY FORUM}, \textit{supra} note 7, at 14.
\end{itemize}
PROMs’ benefits and acceptance of the program.\textsuperscript{375}

b. Minimize Burdens Associated with PROMs

PROM completion should be minimally burdensome for patients.\textsuperscript{376} To that end, implementers might provide patients with options, such as using either a tablet computer or a patient portal and completing PROMs either at the clinical visit or at home.\textsuperscript{377} Implementers should also be mindful of the frequency of PROM administration to avoid redundant and unnecessary data collection.\textsuperscript{378} Thus, administration frequency should be included in PROM specifications. The value of PROMs should be explained to patients, and clinicians should demonstrate their usefulness by referring to patients’ PROM scores during visits.\textsuperscript{379}

Health care organizations should also ensure that PROMs are not excessively cumbersome for clinicians.\textsuperscript{380} Staff members should be tasked with the work of educating patients about PROMs, asking them to complete PROMs, and sending reminders if necessary.\textsuperscript{381}

Initial PROM review could be assigned to someone other than the physician. Trusted nurses or other clinicians could read completed PROM questionnaires and create short summaries for physicians. They would then alert doctors to any responses that require special attention.

c. Harness Artificial Intelligence

Potentially, an even better approach is to automate PROM review using artificial intelligence (AI). AI algorithms can analyze vast amounts of information and make decisions based on the data.\textsuperscript{382} AI could assess each patient’s PROMs, provide physicians with very brief summaries, and alert clinicians to any alarming data that should not be ignored. The alert could appear prominently on the opening screen of the patient’s EHR.

\textsuperscript{375} Id. at 14-15; MASS. MED. SOC., supra note 19, at 7.
\textsuperscript{376} MASS. MED. SOC., supra note 19, at 7; NATIONAL QUALITY FORUM, supra note 7, at 16.
\textsuperscript{377} NATIONAL QUALITY FORUM, supra note 7, at 16.
\textsuperscript{378} Id. at 10.
\textsuperscript{379} Id. at 17; MASS. MED. SOC., supra note 19, at 7.
\textsuperscript{380} NATIONAL QUALITY FORUM, supra note 7, at 17.
\textsuperscript{381} Id.
Furthermore, AI could discern patterns. It could highlight responses or trends in responses that indicate the failure of treatment or worsening of the patient’s condition. It could also identify patterns of responses that are characteristic of particular conditions that the patient might have.

d. Adopt Strategies for PROM Interpretation, Risk Adjustment, and Missing Data

In the clinical setting, physicians must be able to understand PROM scores and know how to respond to them. They must be able to determine whether score changes over time are clinically meaningful and actionable. Implementers should ensure that educational materials are available to train clinicians with respect to PROM interpretation.

If PROMs will be used for nonclinical purposes, such as performance measurement, research, or quality improvement, a proper analysis plan must be in place. This includes statistical adjustment for problems such as response bias and nonresponders as well as mechanisms to address missing data. For example, to compensate for missing data, analysts may collect auxiliary information that is associated with the patient-reported outcome in question (e.g., diagnostic test results) or use statistical machine learning techniques to make adjustments. The process of estimating missing data based on known data points is called imputation.

e. Incorporate PROMs data into EHR Systems

To be optimally useful in clinical practice, PROMs data should be incorporated into EHR systems. Such integration helps physicians use PROMs

383 Id.
384 NATIONAL QUALITY FORUM, supra note 7, at 20.
385 Id.
386 Id.
387 Basch et al., supra note 165, at 500.
388 Id.; Rivera et al., supra note 10, at 1916 (discussing methods to minimize missing data in research studies, such as reminders and notifications to participants). See supra note 342 and accompanying text for discussion of risk adjustment. See supra notes 150-152 for discussion of response bias and notes 142-153 and accompanying text for discussion of missing data and nonresponders.
389 Ayilara et al., supra note 165, at 107; see also CELLA ET AL., supra note 37, at 35-36 (discussing “statistical methods of adjustment”).
391 CELLA ET AL., supra note 37, at 52-54; NATIONAL QUALITY FORUM, supra note 7, at 21-22; see also supra notes 79-83 (discussing integration of PROMs into EHRs).
because they can view them when checking other information in patients’ records. It also facilitates PROM use in research and quality improvement initiatives that will utilize EHRs. PROM developers should adopt user-centered design approaches\textsuperscript{392} so that PROM scores are easy to access, read, and understand.\textsuperscript{393}

\textbf{B. Legal and Policy Interventions}

The legal and policy communities can employ several strategies to facilitate PROM implementation and address its legal implications. Key areas of focus are enhancing privacy protections, addressing medical malpractice concerns, and considering financial incentives for PROM adoption.

\textit{1. Privacy}

PROMs can include a plethora of data about deeply private matters.\textsuperscript{394} A search of the PROMIS database reveals that a very large number of the featured PROMs relate to depression, anxiety, ability to participate in activities, alcohol use, irritability, relationships, positive affect, stress, self-efficacy, sexual functioning and satisfaction, and other sensitive attributes.\textsuperscript{395} Routine inclusion of such patient-provided information in EHRs raises acute privacy concerns.

In response, the HIPAA Privacy Rule’s minimum necessary provision should be modified.\textsuperscript{396} Entities that request patient records and are entitled to receive them because of patient consent or a HIPAA exception should not automatically receive PROMs. Instead, PROMs should be disclosed to requesters only if they have asked for them specifically and explained why they need them. Covered entities should be empowered to assess justifications for PROM requests to approve or deny them just as they already are tasked with determining what constitutes the minimum necessary response for all requests.\textsuperscript{397} These

\begin{itemize}
  \item \textsuperscript{392}User-Centered Design Basics, USABILITY, https://www.usability.gov/what-and-why/user-centered-design.html (last visited Apr. 22, 2023) ("[User-centered design] is based upon an explicit understanding of users, tasks, and environments; is driven and refined by user-centered evaluation; and addresses the whole user experience.").
  \item \textsuperscript{393}National Quality Forum, supra note 7, at 22.
  \item \textsuperscript{394}See, e.g., Rasa Ruseckaitë et al., Evaluation of the Acceptability of Patient-Reported Outcome Measures in Women Following Pelvic Floor Procedures, 31 Quality Life Rsch. 2213, 2214, 2217 (2022).
  \item \textsuperscript{395}View Measures, supra note 57.
  \item \textsuperscript{396}45 C.F.R. § 164.502(b) (2022); see supra notes 396-397 and accompanying text.
  \item \textsuperscript{397}Minimum Necessary Requirement, U.S. Dep’t Health & Hum. Servs. https://www.hhs.gov/hipaa/forprofessionals/privacy/guidance/minimumnecessaryrequirement/index.html (last viewed Jul. 26, 2013) ("For non-routine disclosures and requests, covered entities must develop reasonable criteria for determining and limiting the disclosure or request to only the

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assessments should be carefully conducted by experts such as privacy officers so that disclosures are not simply rubberstamped. PROMs should be stored in EHRs in ways that make them easy to identify and withhold when other data are disclosed.

Establishing a default withholding rule for PROMs has several benefits. First, it would encourage patients to answer questionnaires candidly. This approach would not be unprecedented, as psychotherapy notes already receive a higher degree of privacy protection than less sensitive information. 398

Second, a default withholding rule may often spare health care providers from the work of reviewing all a patient’s PROM responses to determine which should be disclosed under the minimum necessary standard. 399 Patient medical records might include a multitude of PROMs that could make the review task very burdensome. Likewise, a default exclusion rule would save requesters from having to process voluminous unwanted information upon receipt of disclosures. Employers, for example, would likely need only objective clinical data that reveals whether their applicants are qualified for physically demanding jobs. They are unlikely to be able to interpret PROMs accurately and to find them helpful in making employment decisions.

The proliferation of sensitive data provided through PROMs could further intensify data security concerns. To promote compliance with the HIPAA Security Rule, the U.S. Department of Health and Human Services offers numerous data security resources on its website. 401 It should continue to update these resources as technology changes and experts develop new recommendations.

Health care providers must also be vigilant about data security and ensure that skilled professionals are tasked with its maintenance. Some commentators have decried health care providers’ lack of preparedness for cybersecurity attacks. 402 According to one report, seventy-nine percent of data breaches in 2020

minimum amount of protected health information necessary to accomplish the purpose of a non-routine disclosure or request.

398 45 C.F.R. § 164.508(a)(2) (2022) (significantly restricting covered entities’ ability to use psychotherapy notes for treatment, payment, or healthcare operations without patient consent even though the HIPAA Privacy Rule allows for such uses in the case of most protected health information).

399 45 C.F.R. § 164.502(b) (2022) (describing minimum necessary standard).

400 See supra Part I.C (discussing PROM shortcomings and concerns).


402 Devin Partida, 5 Biggest Challenges of Health Care Data Security in 2022, HEALTH IT ANSWERS (Feb. 23, 2022), https://www.healthitanswers.net/5-biggest-challenges-of-health-care-data-security-in-2022/ (“Medical organizations’ vast amounts of sensitive patient data make them prime targets, and many lack the expertise and tools necessary to protect themselves”); Emily
involved healthcare organizations. As others have noted, “Just as hand washing is a foundational element of modern medicine, cyber hygiene must be regarded as a basic and essential component of a functioning medical system.”

2. Medical Malpractice Liability

Clinicians and health care entities should be aware of the potential for malpractice liability associated with PROMs. Liability could arise from failure to review and address data provided in PROMs, excessive reliance on PROMs, or failure to adopt PROMs that have become the standard of care. Medical malpractice attorneys should learn to investigate PROM use when representing both plaintiffs and defendants. For its part, the medical community should undertake efforts to minimize the risk of PROM-related litigation, including formulating clinical practice guidelines for health care providers about PROM implementation and educating patients about PROM use.

a. The Role of PROMs in Litigation

In preparing for litigation, both plaintiffs’ attorneys and defense attorneys should investigate whether PROMs were used during treatment. Plaintiffs’ attorneys should ask clients whether they completed PROMs, what information they provided, whether physicians discussed PROMs with them, and whether they believe their doctors ignored PROM data. Defense attorneys should likewise ask clients whether they used PROMs and how they handled data provided through PROMs. Discovery should routinely include queries about PROMs, such as whether they were utilized, reviewed, or served as the basis for any decision.

b. Clinical Practice Guidelines

Health care providers should proceed with caution when implementing

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404 Skahill & West, supra note 402.

405 See supra Part II.B.

PROM programs and selecting PROMs. Ideally, trustworthy professional organizations and government entities will develop clinical practice guidelines (CPG) that providers can follow in implementing PROM programs. Providers would benefit from guidance regarding the incorporation of PROMs into clinical practice. CPGs could include the technical guidance regarding PROM selection and administration provided above. They could also address how to induce as many patients as possible to complete PROMs, how to review PROMs efficiently, how to determine whether PROM scores require any response, the extent to which PROMs should be discussed during office visits, and more.

It is unclear whether following CPGs could support a defense in a medical malpractice lawsuit. Some experts argue that CPGs should never be admissible in court as evidence of the standard of care because they constitute recommendations rather than proof of actual customary medical practice. Nevertheless, several courts have permitted litigants to use CPGs as evidence regarding the standard of care. Regardless of CPGs’ admissibility, carefully formulated and widely disseminated guidance would be valuable for health care providers as they transition to implementing PROMs. It could prevent them from making obvious mistakes that could lead to malpractice litigation and help them operate in ways that promote patients’ trust and cooperation.

c. Patient Education and Notice

Providers would be wise to communicate clearly with patients regarding how PROMs will be used and what expectations patients should have with respect to them. Patients who are asked to complete PROMs should be given

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411 Hoffman & Podgurski, supra note 245, at 1570-72; McMenamin et al., supra note 410, at 23-24; Mello, supra note 406, at 663-67 (discussing the role of CPGs in litigation).
412 See supra Part II.B.1 and accompanying text (discussing liability concerns relating to physicians’ management of PROMs).
verbal and written explanations of whether doctors will review PROMs in a timely fashion and contact patients about them when appropriate. If PROMs will not be routinely reviewed, patients should be told why they are being asked to complete PROMs (e.g., for quality improvement purposes) and instructed that they should not assume their physicians are aware of all the data they have provided in PROM questionnaires.

On the other hand, if doctors plan to rely on PROMs in making medical decisions because they do not have adequate time for lengthy discussions during patient encounters, it is particularly important that patients be clearly informed that it is vital that they complete their PROM questionnaires. Patients must be warned that their care might be compromised if they ignore requests for PROMs or answer questionnaires only partially, thereby withholding important information from clinicians.

Such notice would be consistent with other notice practices in the medical arena. The HIPAA Privacy Rule requires health care providers to give patients notice of their privacy practices.\(^{413}\) The American Medical Association’s Code of Medical Ethics Opinion 2.3.1 addresses electronic communication with patients.\(^{414}\) It advises physicians to “[n]otify the patient of the inherent limitations of electronic communication, including possible breach of privacy or confidentiality . . . and possible delays in response.”\(^{415}\) A similar notice regarding PROMs would help patients understand their function and limitations and potentially prevent litigation. Written notices should preferably be accompanied by verbal explanations and perhaps training videos to reinforce patient understanding and learning.\(^{416}\) Documentation showing that patients received this guidance could also serve as compelling evidence in clinicians’ defense.

3. **PROM Use by Regulatory Agencies**

The FDA and CMS do not presently require PROMs for any oversight purpose, though regulated entities have the option of submitting them to meet certain requirements.\(^{417}\) Given the current shortcomings and pitfalls of PROMs, it is premature for the FDA and CMS to make them mandatory. The agencies

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413 45 C.F.R. § 164.520(a) (2022).
415 Id.
416 Anne Johnson et al., Written and Verbal Information Versus Verbal Information Only for Patients Being Discharged from Acute Hospital Settings to Home, COCHRANE DATABASE SYSTEMATIC REV., 2003, at 2 (recommending that patients be given both written and verbal instructions).
417 See supra notes 289-300 and 313-319 and accompanying text.
should continue to work with experts to produce PROM guidance for regulated entities so that PROMs that are used voluntarily provide sound data.  

In addition, the FDA should continue to scrutinize any PROMs that are used to meet regulatory requirements and to provide assessments as to whether they are “fit-for-purpose.” CMS would be wise to undertake a similar review and approval process for any PROMs it accepts for payment programs. Note that a determination that a PROM is fit for purposes of FDA or CMS determinations will not necessarily mean that it is also an appropriate choice for clinical care.

4. Financial Incentives

Both the federal government and private insurers can institute financial incentive programs to promote PROM adoption. This section posits that a government program akin to the one established for EHRs is unlikely. Private insurers, however, may well opt to pay providers bonuses for PROM use, though they should not penalize providers for deficient PROM scores at this time.

a. Government Incentives

To accelerate the adoption of PROMs, Congress could pass legislation that establishes a federal incentives program and regulations for PROM adoption. This approach would follow the precedent set by the Health Information Technology for Economic and Clinical Health (HITECH) Act in 2009. The statute dedicated $27 billion to the promotion of health information technology. The funding was used to award generous incentive payments to providers who adopted certified EHR systems and met regulatory specifications for their use. In conjunction with the HITECH Act, CMS enacted the Meaningful Use regulations that detailed objectives that clinicians had to meet with respect to EHR system operation to receive payments. In addition, it established a process for the certification of EHR systems.

Congress could adopt the same approach with respect to PROMs. It could

418 See supra notes 293, 304-307, and 317 and accompanying text (listing several existing and developing guidance documents).
419 See supra notes 296-297 and accompanying text.
421 SHARONA HOFFMAN, ELECTRONIC HEALTH RECORDS AND MEDICAL BIG DATA 2 (2016).
422 Id. at 42-46; 42 C.F.R. §§ 495.2-495.370 (2022).
424 NATIONAL QUALITY FORUM, supra note 7, at 23; Wu et al., supra note 279, at 1869.
enact legislation that empowered CMS to establish a financial incentive program along with regulations for PROM implementation. PROMs would be certified if they met particular requirements such as those outlined above. Providers who work with Medicare and Medicaid patients could receive payments to offset PROM-related investments of time and money. CMS regulations would strive to ensure that providers not only collect suitable PROM data but also employ them to improve patient care.

A PROMs incentive program, however, is improbable. First, such an initiative would require an investment of billions of dollars, and PROMs implementation is unlikely to be a high priority for Congress in this divisive and crisis-prone era. Second, it is doubtful that clinicians who are already overburdened will be receptive to additional regulatory requirements, even if they are accompanied by incentive payments. The meaningful use regulations were widely criticized and resented. PROMs regulations are likely to receive a similar reception. Health care providers would be even more resentful of regulatory mandates that are not accompanied by financial payments to compensate for PROM implementation costs. At this time, CPGs and government agency guidelines may remain the better option.

b. Private Payer Incentives

Alternatively, private payers could offer health care providers financial incentives to implement PROMs. This could be an attractive option for payers that believe PROMs can improve health outcomes and save costs. As discussed above, BCBSMA already piloted such an incentive program.

BCBSMA paid providers for participating in the PROMs initiative but did not adjust insurance coverage based on PROM data. This policy encouraged PROM adoption because it did not create any risk of penalty for providers, even if their patients’ PROM scores appeared unfavorable. Given the many existing challenges of PROM implementation, this is a prudent approach.

It is important to understand that financial incentives for PROM adoption

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425 See supra note 421 and accompanying text.
426 HOFFMAN, supra note 421, at 49-50 (noting that some clinicians called the regulations the “meaningless abuse” regulations); Srinivas Emani et al., Physician Beliefs about the Meaningful Use of the Electronic Health Record: A Follow-Up Study, 8 APPLIED CLINICAL INFORMATICS 1044, 1050 (2017) (“Only a fifth of the physicians responding to our survey agreed or strongly agreed that the meaningful use of the EHR would improve patient-centered care and the quality of care.”).
427 See supra notes 84-87, 92-97 and 327-329 and accompanying text (discussing medical benefits and cost savings associated with PROMs).
428 See supra notes 319-328 and accompanying text.
429 See supra notes 323-324 and accompanying text.
alone do not guarantee that PROMs will be collected consistently or used effectively to promote health care improvements. A 2020 study found that incentives increased PROM collection but did not necessarily lead to successful PROM programs.430 Successful clinics were defined as those with a “mean collection rate in the 6 months prior to January 2019 [that] was 50% or greater.”431 According to the study, health care organizations are most likely to be successful if they engage physicians in building enthusiasm for the benefits of PROMs and provide training regarding PROM use.432 Physician enthusiasm will likely depend on how cumbersome PROM review is and on the availability of tools such as AI that facilitate PROM use. Nevertheless, if employed in conjunction with some of the strategies described above, monetary inducements can play a useful role in encouraging clinicians to embrace PROMs and build a productive PROMs program.

CONCLUSION

PROMs hold promise as an emerging clinical tool that can also contribute to research, health care administration, and regulation. As other scholars have noted, PROMs “directly support the primary goal of much of health care: to improve health-related quality of life,” because “[n]o one can judge this better than the patient.”433 The emergence of PROMS is particularly timely because physicians have ever-shrinking amounts of time to collect data from patients in face-to-face visits.

But PROMs currently have significant pitfalls, and their implementation is complex. This Article has argued that providers should be keenly aware of medical malpractice risks associated with PROMs and that the HIPAA Privacy Rule’s minimum necessary provision should be revised to address PROMs specifically. It further posits that it would be premature for the FDA, CMS, or private insurers to require PROM submission at this time.

Many strategies can be employed to strengthen PROMs and facilitate their integration into clinical practice and other arenas. These include clinical practice guidelines, patient education, financial incentives, PROM analysis by AI, stakeholder input, pilot programs, psychometric evaluations, and a variety of other safeguards relating to PROM selection and implementation. It remains to be seen whether PROMs can become a consistently reliable tool for clinicians, researchers, and others. But with careful planning and execution by qualified professionals.

430 Sisodia et al., supra note 74, at 1.
431 Id. at 3.
432 Id. at 6.
433 Wu et al., supra note 279, at 1864.
experts, PROMs may be able to fulfill their promise of serving as an important instrument to promote health care delivery improvements and bolster efforts to control medical costs.