

Enhancing Oncology Model (EOM) Implementing ePROs

This factsheet provides an overview of electronic patient reported outcomes (ePROs) in EOM, as well as an example timeline for the implementation of ePROs within the model.

What are Patient Reported Outcomes?



Patient Reported Outcomes (PROs) are measurements based on a report that comes directly from the patient, without amendment or interpretation of the patient's response.¹

ePROs are the *electronic capture* of this data.

Why are ePROs important to EOM?



ePROs have been studied in a variety of practice settings -- including community-based practices -- and found to have a number of benefits.²

Using ePRO tools in oncology settings can lead to better identification of patients' needs, improving patient-provider communication, care management, and patient satisfaction, as well as advancing other positive cancer outcomes, such as decreased emergency department visits and improved survival, sometimes exceeding the benefits of oncology drugs.^{3,4}

Immediate benefits of ePROs include, but are not limited to:



Prompting discussions with a clinician



Streamlining consultations



Increasing awareness and triaging of symptoms



Facilitating interprofessional communication



Clinicians in community settings report that utilization of ePROs has been shown to be **helpful for clinical documentation.** Studies also show high levels of **patient engagement**, for patients who are 65 years and older.⁵



Patients report that utilization of ePROs improved discussions with providers and made them feel more in control of their care.

Which ePROs tool should EOM participants use?

CMS is **not** requiring the use of a specific ePROs tool, however participants must use tools that capture, where applicable, outcomes for each of the following domains:

Symptoms and/or Toxicity

Examples: frequency, severity, activity interference, presence/absence of symptoms

Functioning

Examples: physical functioning, role functioning*

*refers to an individual's ability to work or pursue social and/or personal functions

Health-Related Social Needs (HRSN)

Examples: financial toxicity, transportation, food insecurity

Behavioral Health

Examples: psychosocial functioning, anxiety, depression, other behavioral health conditions

Examples of validated and publicly available ePROs tools include the Patient-Reported Outcomes version of the Common Terminology Criteria for Adverse Events (PRO-CTCAE®) and the Patient-Reported Outcomes Measurement Information System (PROMIS®).

Note: The ePROs tools listed here are for example only and do not constitute an endorsement by CMS or CMS affiliates.



Model participants must integrate ePROs data into electronic health records (EHRs). However, EOM participants **do not need** to report the data to CMS at this time.

¹ BEST (Biomarkers, Endpoints, and other Tools) Resource. (2016). Glossary. Retrieved from https://www.ncbi.nlm.nih.gov/books/NBK338448/#IX-P
Basch E, Deal AM, Dueck AC, et al. (2017) Overall Survivial Results of a Trial Assessing Patient-Reported Outcomes for Symptom Monitoring During Routine Cancer Treatment. JAMA. 318(2):197–198. Retrieved from: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5817466/

² Patt D, Wilfong L, Hudson KE, et al: Implementation of electronic patient-reported outcomes for symptom monitoring in a large multisite community oncology practice: Dancing the Texas two-step through a pandemic. JCO Clin Cancer Inform 5:615-621, 2021.

³ Ladanie, A., Schmitt, A.M., Speich, B. (2020). Clinical trial evidence supporting US Food and Drug Administration approval of novel cancer therapies between 2000 and 2016. JAMA. Retrieved from: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7656288/



Enhancing Oncology Model (EOM) ePROs implementation

The sections below provide options for collecting ePROs and an example implementation timeline for EOM participants to integrate ePROs into practice decision-making.

How can participants collect ePROs from patients?

CMS requires that ePROs be administered in an electronic format, including but not limited to the following:



Screen-Based
Reporting Devices
(e.g., via the patient portal on a smart phone or computer)



Interactive Voice Response Systems (e.g., calls to a patient who responds to phone prompts)



SMS Text Systems (e.g., patient provides information via text on mobile device)



ePRO Collection In the Waiting Room (e.g., patient provides data via a tablet while waiting for office visit)

Example ePROs Implementation Timeline

CMS requires that ePROs be collected using a gradual implementation approach. Below is an **example** implementation timeline for ePROs collection:

Performance Period (PP)	Model Year (MY)	ePROs Data Collection Requirement	
1	Year 1	Optional pre-implementation years	EOM participants will collect data using a gradual implementation approach, including an optional pre-implementation period. During the pre-implementation period, if EOM participants are not administering ePROs to their EOM beneficiaries, then they should be building the capabilities to do so beginning in PP5. EOM participants are not required to report data to CMS at this time.
2			
3	Year 2		
4			
5	Year 3	35%	Beginning in PP5 (MY3), participants will be required to implement ePROs prior to each visit where one or more qualifying E&M services are furnished to the EOM beneficiary.
6		attributed EOM beneficiaries*	
7	Year 4	50%	Similar to the pre-implementation period, EOM
8		attributed EOM beneficiaries*	participants are not required to report data to CMS at this time.
9	Year 5	75% attributed EOM beneficiaries*	Note: This does not include the beneficiary's first visit with the EOM participant, however it does include subsequent visits.
10			
* Note: This timeline includes example percentages of ePROs data collection beginning in PP5.			