

QOPI® REPORTING REGISTRY (QCDR) 2018

MEASURE SPECIFICATIONS

| Measure # | Measure Title | Measure Description | Denominator Exclusion/Exceptions | Denominator | Numerator |
|-----------|---|---|---|---|--|
| QOPI5 | Chemotherapy administered to patients with metastatic solid tumor with performance status of 3, 4, or undocumented (Lower Score - Better) | Percentage of adult patients with metastatic solid tumors and performance status of 3, 4, or undocumented who receive chemotherapy (Lower score - Better) | Diagnosis of malignant neoplasm of placenta/trophoblastic neoplasm, testicular carcinoma, leukemia, Hodgkin and non-Hodgkin's lymphoma | Solid tumor AND ((Intent not documented AND Stage IV at initial diagnosis or development of distant metastases = Yes) OR Intent = non-curative) AND Patient received chemotherapy for stage IV or distant metastatic disease | (Performance status documented within 2 weeks of most recent chemotherapy administration for distant metastatic disease = 3 or 4 OR Not documented) AND (Patient received chemotherapy for metastatic disease as part of IRB approved protocol = No OR Patient received chemotherapy for metastatic disease as part of IRB approved protocol = Unknown) |
| QOPI11 | Combination chemotherapy received within 4 months of diagnosis by women under 70 with AJCC stage IA (T1c) to III ER/PR negative breast cancer | Percentage of adult women under 70 with a diagnosis of AJCC stage IA (T1c) to III ER/PR negative breast cancer, who receive combination chemotherapy within 4 months of diagnosis | Transfer-in Status ≠ Reporting practice has/had primary responsibility for the initial course of the patient's medical oncology care OR M-Stage at breast cancer diagnosis = M1 OR Diagnosis of malignant phyllodes, cystosarcoma phyllodes, tubular carcinoma, mucinous carcinoma OR ((Multi-agent Breast Chemotherapy administered = Chemotherapy NOT administered AND | Patients 18-69 at diagnosis AND Breast cancer diagnosis AND ((AJCC stage at breast cancer diagnosis = IIA - IIIC) OR (AJCC stage at breast cancer diagnosis = (IA and T-Stage at breast cancer diagnosis=T1c) or IB) OR (T-Stage at breast cancer diagnosis = T1c, T2-T4d and N-Stage at breast cancer diagnosis = N0) OR (N-Stage at breast cancer diagnosis = N1-N3c) OR | Chemotherapy administered during initial treatment course (Breast cancer) = Multi-agent chemotherapy administered AND Date the chemotherapy was initiated (multi-agent) - Date of Diagnosis ≤ 124 days OR Alternative treatment according to clinical trial protocol |

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| | | | (Abstraction date – diagnosis date < 124 days OR Deceased date – diagnosis date < 124 days OR Date of first visit – diagnosis date > 124 days) OR (Reason Multi-Agent Chemotherapy NOT Administered = Patient declined or Patient died or transferred or Contraindication or other clinical exclusion or Null)) | (T1c and N1mi)) AND (ER status = ER negative and PR status = PR negative) | |
| QOPI15 | GCSF administered to patients who received chemotherapy for metastatic cancer (Lower Score - Better) | Percentage of adult patients with metastatic cancer who are administered chemotherapy and who receive a colony stimulating factor (Lower score - Better) | Diagnosis of malignant neoplasm of placenta/trophoblastic neoplasm, testicular carcinoma, leukemia, Hodgkin and non-Hodgkin’s lymphoma | Patients ≥ 18 at cancer diagnosis AND ((Metastatic/advanced disease AND Chemotherapy administered) OR Palliative intent chemotherapy administered) | GCSF received = Yes |

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| QOPI16 | Oncology: Treatment Summary Communication – Radiation Oncology | Percentage of patients, regardless of age, with a diagnosis of cancer who have undergone brachytherapy or external beam radiation therapy who have a treatment summary report in the chart that was communicated to the physician(s) providing continuing care and to the patient within one month of completing treatment | Documentation of a patient reason(s) for not communicating the treatment summary report to the physician(s) providing continuing care (e.g., patient requests that report not be sent) and to the patient within one month of completing treatment OR Documentation of a system reason(s) for not communicating the treatment summary report to the physician(s) providing continuing care (e.g., patient does not have any physician responsible for providing continuing care) and to the patient within one month of completing treatment | All patients, regardless of age, with a diagnosis of cancer who have undergone brachytherapy or external beam radiation therapy | Patients who have a treatment summary report in the chart that was communicated to the physician(s) providing continuing care and to the patient within one month of completing treatment |
| QOPI17 | External Beam Radiotherapy for Bone Metastases | Percentage of patients, regardless of age, with a diagnosis of painful bone metastases and no history of previous radiation who receive external beam radiation therapy (EBRT) with an acceptable fractionation scheme as defined by the guideline. | The medical reasons for denominator exclusions are: 1) Previous radiation treatment to the same anatomic site; 2) Patients with femoral axis cortical involvement greater than 3 cm in length; 3) Patients who have undergone a surgical stabilization procedure; and 4) Patients with spinal cord compression, cauda equina compression or radicular pain | All patients with painful bone metastases and no previous radiation to the same anatomic site who receive EBRT | All patients, regardless of age, with painful bone metastases, and no previous radiation to the same anatomic site who receive EBRT with any of the following recommended fractionation schemes: 30Gy/10fxns, 24Gy/6fxns, 20Gy/5fxns, 8Gy/1fxn. |

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| QPP 47 | Care Plan | Percentage of patients aged 65 years and older who have an advance care plan or surrogate decision maker documented in the medical record or documentation in the medical record that an advance care plan was discussed but the patient did not wish or was not able to name a surrogate decision maker or provide an advance care plan | Hospice services received by patient any time during the measurement period: G9692 | <p>Patients aged ≥ 65 years on date of encounter</p> <p>AND</p> <p>Patient encounter during the performance period (CPT or HCPCS): 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215, 99218, 99219, 99220, 99221, 99222, 99223, 99231, 99232, 99233, 99234, 99235, 99236, 99291, 99304, 99305, 99306, 99307, 99308, 99309, 99310, 99324, 99325, 99326, 99327, 99328, 99334, 99335, 99336, 99337, 99341, 99342, 99343, 99344, 99345, 99347, 99348, 99349, 99350, G0402, G0438, G0439</p> | <p>Advance Care Planning discussed and documented; advance care plan or surrogate decision maker documented in the medical record (1123F)</p> <p>OR</p> <p>Advance Care Planning discussed and documented in the medical record; patient did not wish or was not able to name a surrogate decision maker or provide an advance care plan (1124F)</p> |
| QPP 67 | Hematology: Myelodysplastic Syndrome (MDS) and Acute Leukemias: Baseline Cytogenetic Testing Performed on Bone Marrow | Percentage of patients aged 18 years and older with a diagnosis of myelodysplastic syndrome (MDS) or an acute leukemia who had baseline cytogenetic testing performed on bone marrow | <p><i>Denominator Exceptions:</i></p> <p>Documentation of medical reason(s) for not performing baseline cytogenetic testing on bone marrow (e.g., no liquid bone marrow or fibrotic marrow) (3155F with 1P)</p> <p>OR</p> <p>Documentation of patient reason(s) for not performing baseline cytogenetic testing on bone marrow (e.g., at time of diagnosis receiving palliative care or not receiving treatment as defined above) (3155F with 2P)</p> <p>OR</p> <p>Documentation of system reason(s) for not performing</p> | <p>Patients aged ≥ 18 years on date of encounter</p> <p>AND</p> <p>Diagnosis for MDS or acute leukemia – not in remission (ICD-10-CM)</p> <p>AND</p> <p>Patient encounter during the performance period (CPT)</p> | <p>Cytogenetic testing performed on bone marrow at time of diagnosis or prior to initiating treatment (3155F)</p> |

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| | | | baseline cytogenetic testing on bone marrow (e.g., patient previously treated by another physician at the time cytogenetic testing performed) (3155F with 3P) | | |
| QPP 69 | Hematology: Multiple Myeloma: Treatment with Bisphosphonates | Percentage of patients aged 18 years and older with a diagnosis of multiple myeloma, not in remission, who were prescribed or received intravenous bisphosphonate therapy within the 12 month reporting period. | <i>Denominator Exception:</i> Documentation of medical reason(s) for not prescribing bisphosphonates (e.g., patients who do not have bone disease, patients with dental disease, patients with renal insufficiency) (4100F with 1P) OR Documentation of patient reason(s) for not prescribing bisphosphonates (4100F with 2P) | Patients aged ≥ 18 years on date of encounter AND Diagnosis for multiple myeloma – not in remission (ICD-10-CM): C90.00, C90.02 AND Patient encounter during the performance period (CPT): 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215 | Bisphosphonate therapy, intravenous, ordered or received (4100F) |
| QPP 70 | Hematology: Chronic Lymphocytic Leukemia (CLL): Baseline Flow Cytometry | Percentage of patients aged 18 years and older, seen within a 12 month reporting period, with a diagnosis of chronic lymphocytic leukemia (CLL) made at any time during or prior to the reporting period who had baseline flow cytometry studies performed and documented in the chart | <i>Denominator Exceptions:</i> Documentation of medical reason(s) for not performing baseline flow cytometry studies (3170F with 1P) OR Documentation of patient reason(s) for not performing baseline flow cytometry studies (e.g., receiving palliative care or not receiving treatment as defined above) (3170F with 2P) OR Documentation of system reason(s) for not performing | Patients aged ≥ 18 years on date of encounter AND Diagnosis for CLL – not in remission (ICD-10-CM): C91.10, C91.12 AND Patient encounter during the performance period (CPT): 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215 | Flow cytometry studies performed at time of diagnosis or prior to initiating treatment (3170F) |

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| | | | baseline flow cytometry studies (e.g., patient previously treated by another physician at the time baseline flow cytometry studies were performed) (3170F with 3P) | | |
| QPP 102 | Prostate Cancer: Avoidance of Overuse of Bone Scan for Staging Low Risk Prostate Cancer Patients | Percentage of patients, regardless of age, with a diagnosis of prostate cancer at low (or very low) risk of recurrence receiving interstitial prostate brachytherapy, OR external beam radiotherapy to the prostate, OR radical prostatectomy, OR cryotherapy who did not have a bone scan performed at any time since diagnosis of prostate cancer | <i>Denominator Exceptions:</i> Documentation of medical reason(s) for performing a bone scan (including documented pain, salvage therapy, other medical reasons) (3269F with 1P) OR Documentation of system reason(s) for performing a bone scan (including bone scan ordered by someone other than the reporting physician) (3269F with 3P) | Diagnosis for prostate cancer (ICD-10-CM): C61 AND Patient encounter during the performance period AND Low (very low) risk of recurrence = Yes AND Receiving interstitial prostate brachytherapy = Yes OR Receiving external beam radiotherapy to the prostate = Yes OR Receiving radical prostatectomy = Yes OR Receiving cryotherapy = Yes | Bone scan not performed prior to initiation of treatment nor at any time since diagnosis of prostate cancer |

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|----------------|--|--|---|---|---|
| QPP 104 | Prostate Cancer: Adjuvant Hormonal Therapy for High Risk or Very High Risk Prostate Cancer | Percentage of patients, regardless of age, with a diagnosis of prostate cancer at high or very high risk of recurrence receiving external beam radiotherapy to the prostate who were prescribed adjuvant hormonal therapy (GnRH [gonadotropin-releasing hormone] agonist or antagonist) | Diagnosis for metastatic cancer <i>Denominator Exception:</i> Documentation of medical reason(s) for not prescribing/administering adjuvant hormonal therapy (e.g., salvage therapy) OR Documentation of patient reason(s) for not prescribing/administering adjuvant hormonal therapy | Diagnosis for prostate cancer (ICD-10-CM): C61 AND Patient encounter during the performance period (CPT): 77427, 77435 AND High or very high risk of recurrence = Yes AND Receiving external beam radiotherapy to the prostate = Yes | Adjuvant (i.e., in combination with external beam radiotherapy to the prostate for prostate cancer) hormonal therapy (gonadotropin-releasing hormone[GnRH] agonist or antagonist) prescribed/administered (4164F) |
| QPP 130 | Documentation of Current Medications in the Medical Record | Percentage of visits for patients aged 18 years and older for which the eligible professional attests to documenting a list of current medications using all immediate resources available on the date of the encounter. This list must include ALL known prescriptions, over-the-counters, herbals, and vitamin/mineral/dietary (nutritional) supplements AND must contain the medications' name, dosage, frequency and route of administration | <i>Denominator Exception:</i> Patient Not Eligible: Eligible clinician attests to documenting in the medical record the patient is not eligible for a current list of medications being obtained, updated, or reviewed by the eligible clinician. | Diagnosis of cancer AND Patients aged ≥ 18 years on date of encounter AND Patient encounter during the performance period | Eligible clinician attests to documenting in the medical record they obtained, updated, or reviewed the patient's current medications (G8427) |

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| QPP 134 | Preventive Care and Screening: Screening for Clinical Depression and Follow-Up Plan | Percentage of patients aged 12 years and older screened for depression on the date of the encounter using an age appropriate standardized depression screening tool AND if positive, a follow-up plan is documented on the date of the positive screen | Documentation stating the patient has an active diagnosis of depression or has a diagnosed bipolar disorder, therefore screening or follow-up not required: G9717 <i>Denominator Exception:</i> Screening for depression not completed, documented reason (G8433) | Patients aged ≥ 12 years on date of encounter AND Patient encounter during the performance period (CPT or HCPCS): 90791, 90792, 90832, 90834, 90837, 92625, 96116, 96118, 96150, 96151, 97165, 97166, 97167, 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215, G0101, G0402, G0438, G0439, G0444 | Screening for depression is documented as being positive AND a follow-up plan is documented (G8431) OR Screening for depression is documented as negative, a follow-up plan is not required (G8510) |
| QPP 143 | Oncology: Medical and Radiation - Pain Intensity Quantified | Percentage of patient visits, regardless of patient age, with a diagnosis of cancer currently receiving chemotherapy or radiation therapy in which pain intensity is quantified | None | Diagnosis of cancer AND Each office visit during the performance period AND (Receiving chemotherapy = Yes OR Receiving radiation therapy = Yes) | Pain severity quantified; pain present (1125F) OR Pain severity quantified; no pain present (1126F) |
| QPP 226 | Preventive Care and Screening: Tobacco Use: Screening and Cessation Intervention | Percentage of patients aged 18 years and older who were screened for tobacco use one or more times within 24 months AND who received cessation counseling intervention if identified as a tobacco user | None <i>Denominator Exception:</i> Documentation of medical reason(s) for not screening for tobacco use (e.g., limited life expectancy, other medical reason) | Patients ≥ 18 on date of encounter AND (Patient encounters during performance period ≥ 2 OR Preventive encounter during performance period ≥ 1) | Patient screened for tobacco use AND received tobacco cessation intervention (counseling, pharmacotherapy, or both), if identified as a tobacco user (4004F) OR Current tobacco non-user (1036F) |

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| QPP 317 | Preventive Care and Screening: Screening for High Blood Pressure and Follow-Up Documented | Percentage of patients aged 18 years and older seen during the reporting period who were screened for high blood pressure AND a recommended follow-up plan is documented based on the current blood pressure (BP) reading as indicated | <p>Active diagnosis of hypertension</p> <p><i>Denominator Exception:</i> Documented reason for not screening or recommending a follow-up for high blood pressure.</p> <ul style="list-style-type: none"> • Patient refuses to participate (either BP measurement or follow-up) • Patient is in an urgent or emergent situation where time is of the essence and to delay treatment would jeopardize the patient's health status. This may include but is not limited to severely elevated BP when immediate medical treatment is indicated | <p>Patients ≥ 18</p> <p>AND</p> <p>Patient encounter during the performance period</p> | <p>Normal blood pressure reading documented, follow-up not required (G8783)</p> <p>OR</p> <p>Pre-Hypertensive or Hypertensive blood pressure reading documented, AND the indicated follow-up is documented (G8950)</p> |

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| QPP 408 | Opioid Therapy Follow-up Evaluation | All patients 18 and older prescribed opiates for longer than six weeks duration who had a follow-up evaluation conducted at least every three months during Opioid Therapy documented in the medical record | None | <p>Patients aged ≥ 18 years on date of encounter</p> <p>AND</p> <p>Patient encounter during the performance period (CPT): 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215, 99304, 99305, 99306, 99307, 99308, 99309, 99310, 99324, 99325, 99326, 99327, 99328, 99334, 99335, 99336, 99337, 99341, 99342, 99343, 99344, 99345, 99347, 99348, 99349, 99350</p> <p>AND</p> <p>Patients prescribed opiates for longer than six weeks: G9561</p> | Patients who had a follow-up evaluation conducted at least every three months during opioid therapy (G9562) |
| QPP 449 | HER2 Negative or Undocumented Breast Cancer Patients Spared Treatment with HER2-Targeted Therapies | Proportion of female patients (aged 18 years and older) with breast cancer who are human epidermal growth factor receptor 2 (HER2)/neu negative who are not administered HER2-targeted therapies | Patient transferred to practice after initiation of chemotherapy | <p>Female Patients ≥ 18 years on date of encounter</p> <p>AND</p> <p>Diagnosis of Breast Cancer</p> <p>AND</p> <p>Two or more encounters at the reporting site</p> <p>AND</p> <p>HER-2/neu = Negative</p> <p>OR</p> <p>HER-2/neu = Undocumented</p> <p>OR</p> <p>HER-2/neu =Unknown</p> | HER2-targeted therapies not administered during the initial course of treatment (G9827) |

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| QPP 450 | Trastuzumab Received By Patients With AJCC Stage I (T1c) - III And HER2 Positive Breast Cancer Receiving Adjuvant Chemotherapy | Proportion of female patients (aged 18 years and older) with AJCC stage I (T1c) – III, human epidermal growth factor receptor 2 (HER2) positive breast cancer receiving adjuvant chemotherapy who are also receiving trastuzumab | Reason for not administering Trastuzumab documented (e.g. patient declined, patient died, patient transferred, contraindication or other clinical exclusion, neoadjuvant chemotherapy or radiation NOT complete) (G9836) <i>Denominator Exception:</i> Reason for not administering Trastuzumab documented (e.g. patient declined, patient died, patient transferred, contraindication or other clinical exclusion, neoadjuvant chemotherapy or radiation NOT complete) (G9836) | Female Patients aged ≥ 18 years on date of encounter AND Diagnosis of breast cancer AND Patient encounter during performance period (CPT): 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215 AND Two or more encounters at the reporting site AND Breast Adjuvant Chemotherapy administered= Yes AND HER-2/neu = Positive AND AJCC stage at breast cancer diagnosis = II or III OR AJCC stage at breast cancer diagnosis = I (IA or IB) and T-Stage at breast cancer diagnosis does NOT equal = T1, T1a, T1b | Trastuzumab administered within 12 months of diagnosis (G9835) |

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| QPP 451 | KRAS Gene Mutation Testing Performed for Patients with Metastatic Colorectal Cancer who receive Anti-epidermal Growth Factor Receptor (EGFR) Monoclonal Antibody Therapy | Percentage of adult patients (aged 18 or over) with metastatic colorectal cancer who receive anti-epidermal growth factor receptor monoclonal antibody therapy for whom KRAS gene mutation testing was performed | None | Patients aged ≥ 18 years on date of encounter AND Diagnosis of Initial colon or rectal cancer diagnosis AND Two or more encounters at the reporting site AND Patient has metastatic disease at diagnosis AND Anti-EGFR monoclonal antibody therapy | KRAS gene mutation testing performed before initiation of anti-EGFR MoAb (G9840) |
| QPP 452 | Patients with Metastatic Colorectal Cancer and KRAS Gene Mutation Spared Treatment with Anti-epidermal Growth Factor Receptor (EGFR) Monoclonal Antibodies | Percentage of adult patients (aged 18 or over) with metastatic colorectal cancer and KRAS gene mutation spared treatment with anti-EGFR monoclonal antibodies | None | Patients aged ≥ 18 years on date of encounter AND Diagnosis of colon or rectal cancer (ICD-10 CM): C18.0, C18.2, C18.3, C18.4, C18.5, C18.6, C18.7, C18.8, C18.9, C19, C20 AND Patient encounter during the performance period: 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215 AND Two or more encounters at the reporting site AND Patient has metastatic disease at diagnosis: G9842 AND KRAS gene mutation: G9843 | Patient did not receive anti-EGFR monoclonal antibody therapy ((Anti-EGFR monoclonal antibody-cetuximab or panitumumab) G9844) |

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| QPP 453 | Proportion Receiving Chemotherapy in the Last 14 Days of Life (Lower score - Better) | Proportion of patients who died from cancer receiving chemotherapy in the last 14 days of life (Lower score - Better) | None | Diagnosis of cancer AND Two or more encounters at the reporting site AND Patients who died from cancer: G9846 | Patient received chemotherapy in the last 14 days of life (G9847) |
| QPP 456 | Proportion Not Admitted To Hospice | Proportion of patients who died from cancer not admitted to hospice | None | Diagnosis of cancer AND Patient encounter during the performance period (CPT): 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215 AND Two or more encounters at the reporting site AND Patients who died from cancer: G9855 | Patient was not admitted to hospice (G9856) |
| QPP 457 | Proportion Admitted to Hospice for less than 3 days (Lower score - Better) | Proportion of patients who died from cancer, and admitted to hospice and spent less than 3 days there (Lower score - Better) | None | Diagnosis of cancer AND Two or more encounters at the reporting site AND Patient enrolled in hospice: G9858 AND Patients who died from cancer: G9859 | Patient spent less than three days in hospice care (G9860) |

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| QPP 462 | Bone Density Evaluation for Patients with Prostate Cancer and Receiving Androgen Therapy | Patients determined as having prostate cancer who are currently starting or undergoing androgen deprivation therapy (ADT), for an anticipated period of 12 months or greater and who receive an annual bone density evaluation. The bone density evaluation must be prior to the start of ADT or within 3 months of the start of ADT. | None | Patients determined as having prostate cancer who are currently starting or undergoing androgen deprivation therapy (ADT), for an anticipated period of 12 months or greater | Patients with a bone density evaluation within the two years prior to the start of or less than three months after the start of ADT Treatment |