## The American College of Radiology

## Lung Cancer Screening Registry (LCSR)

## **Measures**

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American College of Radiology 1891 Preston White Drive Reston, VA 20191-4397

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#### AMERICAN COLLEGE OF RADIOLOGY

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## 1. Appropriateness of screening by USPSTF 2013 criteria

Data Elements	Clinical Performance Measure
<ul> <li>Patient's date of birth</li> <li>Number of pack-years of smoking</li> <li>Number of years since quit</li> <li>Smoking status</li> </ul>	Measure Description: Percent of screening exams done on adults who are aged 55 to 80 years, have a 30 packyear smoking history, and currently smoke or have quit within the past 15 years.  Numerator: Number of exams done on patients aged 55 to 80 years who have a 30 pack-year smoking history and currently smoke or have quit within the past 15 years.  Denominator: Number of screening exams  Numerator and Denominator Exclusion: None

## 2. Appropriateness of screening by USPSTF 2021 criteria

Data Elements	Clinical Performance Measure
<ul> <li>Patient's date of birth</li> <li>Number of pack-years of smoking</li> <li>Number of years since quit</li> <li>Smoking status</li> </ul>	Measure Description: Percent of screening exams done on adults who are aged 50 to 80 years, have a 20 packyear smoking history, and currently smoke or have quit within the past 15 years.  Numerator: Number of exams done on patients aged 55 to 80 years who have a 30 pack-year smoking history and currently smoke or have quit within the past 15 years.  Denominator: Number of screening exams  Numerator and Denominator Exclusion: None

## 3. Smoking cessation offered

Data Elements	Clinical Performance Measure
Did physician provide smoking cessation guidance to patient?	Measure description: Percent of exams where patients are offered smoking cessation guidance.  Numerator: Number of exams where smoking cessation guidance were offered
	<b>Denominator</b> : Number of screening exams
	Numerator and Denominator Exclusion: None

## 4. Smoking cessation offered among current smokers

Data Elements	Clinical Performance Measure
<ul> <li>Did physician provide smoking cessation guidance to patient?</li> <li>Smoking status</li> </ul>	Measure description: Percent of screening exams done on current smokers who are offered smoking cessation guidance.  Numerator: Number of screening exams where smoking cessation guidance were offered  Denominator: Number of screening exams where the patient currently smokes  Numerator and Denominator Exclusion: None

#### 5. Non-smoking rate

Data Elements	Clinical Performance Measure
Smoking status	<b>Measure description</b> : Percent of patients reporting as Former Smoker out of all patients reporting as Current Smoker, Former Smoker, or Smoker, Current Status Unknown
	<b>Numerator:</b> Number of screening exams among former smokers
	<b>Denominator</b> : Number of screening exams among patients reporting as Current Smoker, Former Smoker, or Smoker, Current Status Unknown
	Numerator and Denominator Exclusion: None

## 6. Adherence to annual screening

Data Elements	Clinical Performance Measure
<ul><li>Examination date</li><li>CT exam results by Lung-RADS category</li></ul>	<b>Measure description</b> : Percent of patients with follow- up exam 11-15 months after previous screening (where previous screening met appropriateness criteria and had Lung RADS 1 or 2)
	<b>Numerator:</b> Number of patients with follow-up exam 11-15 months after previous screening smokers
	<b>Denominator</b> : Number of patients where previous screening met appropriateness criteria and had Lung RADS 1 or 2
	Numerator and Denominator Exclusion: None

## 7. Radiation exposure 1: Mean CTDIvol overall

Data Elements	Clinical Performance Measure
• CTDIvol (mGy)	Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening exams performed
	<b>Numerator:</b> Total CTDIvol for all screening exams performed
	<b>Denominator</b> : Number of screening exams
	<b>Numerator and Denominator Exclusion:</b> None

#### 8. Radiation exposure: Mean CTDIvol underweight (BMI<18.5)

Data Elements	Clinical Performance Measure
<ul><li>CTDIvol (mGy)</li><li>Patient height (inches)</li><li>Patient weight (lbs)</li></ul>	Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening exams performed on underweight patients (BMI<18.5)
	<b>Numerator:</b> Total CTDIvol across all screening exams performed on underweight patients
	<b>Denominator</b> : Number of screening exams where patients were underweight (BMI<18.5)
	Numerator and Denominator Exclusion: None

## 9. Radiation exposure: Mean CTDIvol healthy weight (BMI 18.5-24.9)

Data Elements	Clinical Performance Measure
<ul><li>CTDIvol (mGy)</li><li>Patient height (inches)</li><li>Patient weight (lbs)</li></ul>	Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening exams performed on normal weight patients (BMI 18.5-24.9)
	<b>Numerator:</b> Total CTDIvol across all screening exams performed on normal-weight patients
	<b>Denominator</b> : Number of screening exams where patients were normal-weight (BMI 18.5-24.9)
	Numerator and Denominator Exclusion: None

#### 10. Radiation exposure: Mean CTDIvol overweight (BMI 25-29.9)

Clinical Performance Measure
Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening exams performed on overweight patients (BMI 25-29.9)  Numerator: Total CTDIvol across all screening exams performed on overweight patients  Denominator: Number of screening exams where
patients were overweight (BMI 25-29.9)  Numerator and Denominator Exclusion: None

## 11. Radiation exposure: Mean CTDIvol obese (BMI of 30 or greater)

Data Elements	Clinical Performance Measure
<ul> <li>CTDIvol (mGy)</li> <li>Patient height (inches)</li> <li>Patient weight (lbs)</li> </ul>	Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening exams performed on obese patients (BMI of 30 or greater)  Numerator: Total CTDIvol across all screening exams performed on obese patients  Denominator: Number of screening exams where patients were obese (BMI of 30 or greater)
	Numerator and Denominator Exclusion: None

#### 12. Radiation exposure: Mean CTDIvol obese class I (BMI of 30-34.9)

Data Elements	Clinical Performance Measure
<ul><li>CTDIvol (mGy)</li><li>Patient height (inches)</li><li>Patient weight (lbs)</li></ul>	<b>Measure description:</b> Mean Volumetric CT Dose Index (CTDIvol) across all screening exams performed on obese patients class I (BMI of 30-34.9)
	<b>Numerator:</b> Total CTDIvol across all screening exams performed on obese patients
	<b>Denominator</b> : Number of screening exams where patients were obese class I (BMI of 30-34.9)
	Numerator and Denominator Exclusion: None

#### 13. Radiation exposure: Mean CTDIvol obese class II (BMI of 35-39.9)

Data Elements	Clinical Performance Measure
<ul> <li>CTDIvol (mGy)</li> <li>Patient height (inches)</li> <li>Patient weight (lbs)</li> </ul>	Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening exams performed on obese patients class II (BMI of 35-39.9)  Numerator: Total CTDIvol across all screening exams performed on obese patients
	<b>Denominator</b> : Number of screening exams where patients were obese class II (BMI of 35-39.9) <b>Numerator and Denominator Exclusion</b> : None

#### 14. Radiation exposure: Mean CTDIvol obese class III (BMI of 40 or greater)

Data Elements	Clinical Performance Measure
<ul><li>CTDIvol (mGy)</li><li>Patient height (inches)</li><li>Patient weight (lbs)</li></ul>	Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening exams performed on obese patients class III (BMI of 40 or greater)  Numerator: Total CTDIvol across all screening exams performed on obese patients
	<b>Denominator</b> : Number of screening exams where patients were obese class III (BMI of 40 or greater) <b>Numerator and Denominator Exclusion</b> : None

## 15. Radiation exposure: Mean CTDIvol low dose chest exams

Data Elements	Clinical Performance Measure
<ul><li>CTDIvol (mGy)</li><li>Modality</li></ul>	Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening other than routine chest exams performed
	<b>Numerator:</b> Total CTDIvol for all screening other than routine chest exams performed
	<b>Denominator</b> : Number of screening other than routine chest exams
	Numerator and Denominator Exclusion: None

#### 16. Radiation exposure: Mean CTDIvol routine (not low dose) chest exams

Data Elements	Clinical Performance Measure
<ul><li>CTDIvol (mGy)</li><li>Modality</li></ul>	Measure description: Mean Volumetric CT Dose Index (CTDIvol) across all screening routine chest exams performed  Numerator: Total CTDIvol for all screening routine chest exams performed
	<b>Denominator</b> : Number of screening routine chest exams <b>Numerator and Denominator Exclusion</b> : None

#### 17. Radiation exposure: Mean DLP overall

Data Elements	Clinical Performance Measure
• DLP (mGy*cm)	Measure description: Mean Dose Length Product (DLP) across all screening exams performed
	Numerator: Total DLP for all screening exams performed
	Denominator: Number of screening exams
	Numerator and Denominator Exclusion: None

## 18. Radiation exposure: Mean DLP underweight (BMI<18.5)

Data Elements	Clinical Performance Measure
<ul><li>DLP (mGy*cm)</li><li>Patient height (inches)</li><li>Patient weight (lbs)</li></ul>	Measure description: Mean Dose Length Product (DLP) across all screening exams performed on underweight patients (BMI<18.5)
	<b>Numerator:</b> Total DLP across all screening exams performed on underweight patients
	<b>Denominator</b> : Number of screening exams where patients were underweight (BMI<18.5)
	Numerator and Denominator Exclusion: None

## 19. Radiation exposure: Mean DLP healthy weight (BMI 18.5-24.9)

Data Elements	Clinical Performance Measure
<ul> <li>DLP (mGy*cm)</li> <li>Patient height (inches)</li> <li>Patient weight (lbs)</li> </ul>	Measure description: Mean Dose Length Product (DLP) across all screening exams performed on normal weight patients (BMI 18.5-24.9)  Numerator: Total DLP across all screening exams performed on normal-weight patients  Denominator: Number of screening exams where patients were normal-weight (BMI 18.5-24.9)  Numerator and Denominator Exclusion: None
	Numerator and Denominator Exclusion: None

## 20. Radiation exposure: Mean DLP overweight (BMI 25-29.9)

Data Elements	Clinical Performance Measure
<ul> <li>DLP (mGy*cm)</li> <li>Patient height (inches)</li> <li>Patient weight (lbs)</li> </ul>	Measure description: Mean Dose Length Product (DLP) across all screening exams performed on overweight patients (BMI 25-29.9)  Numerator: Total DLP across all screening exams performed on overweight patients  Denominator: Number of screening exams where patients were overweight (BMI 25-29.9)  Numerator and Denominator Exclusion: None

## 21. Radiation exposure: Mean DLP obese (BMI of 30 or greater)

Data Elements	Clinical Performance Measure
<ul><li>DLP (mGy*cm)</li><li>Patient height (inches)</li><li>Patient weight (lbs)</li></ul>	<b>Measure description:</b> Mean Dose Length Product (DLP) across all screening exams performed on obese patients (BMI 30 or greater)
	<b>Numerator:</b> Total DLP across all screening exams performed on obese patients
	<b>Denominator</b> : Number of screening exams where patients were obese (BMI of 30 or greater)
	Numerator and Denominator Exclusion: None

## 22. Radiation exposure: Mean DLP obese class I (BMI of 30-34.9)

Data Elements	Clinical Performance Measure
<ul><li>DLP (mGy*cm)</li><li>Patient height (inches)</li><li>Patient weight (lbs)</li></ul>	<b>Measure description:</b> Mean Dose Length Product (DLP) across all screening exams performed on obese class I patients (BMI 30-34.9)
	<b>Numerator:</b> Total DLP across all screening exams performed on obese patients
	<b>Denominator</b> : Number of screening exams where patients were obese (BMI of 30-34.9)
	Numerator and Denominator Exclusion: None

## 23. Radiation exposure: Mean DLP obese class II (BMI of 35-39.9)

Data Elements	Clinical Performance Measure
<ul> <li>DLP (mGy*cm)</li> <li>Patient height (inches)</li> <li>Patient weight (lbs)</li> </ul>	Measure description: Mean Dose Length Product (DLP) across all screening exams performed on obese class II patients (BMI 35-39.9)  Numerator: Total DLP across all screening exams performed on obese patients
	<b>Denominator</b> : Number of screening exams where patients were obese (BMI of 35-39.9)
	Numerator and Denominator Exclusion: None

#### 24. Radiation exposure: Mean DLP obese class III (BMI of 40 or greater)

Data Elements	Clinical Performance Measure
<ul><li>DLP (mGy*cm)</li><li>Patient height (inches)</li><li>Patient weight (lbs)</li></ul>	Measure description: Mean Dose Length Product (DLP) across all screening exams performed on obese class III patients (BMI 40 or greater)  Numerator: Total DLP across all screening exams
	performed on obese patients
	<b>Denominator</b> : Number of screening exams where patients were obese (BMI of 40 or greater)
	Numerator and Denominator Exclusion: None

## 25. Radiation exposure: Mean DLP routine chest exams

Data Elements	Clinical Performance Measure
<ul><li>DLP (mGy*cm)</li><li>Modality</li></ul>	Measure description: Mean Dose Length Product (DLP) across all screening routine chest exams performed
	Numerator: Total DLP for all routine chest screening exams performed
	Denominator: Number of screening routine chest exams
	Numerator and Denominator Exclusion: None

#### 26. Radiation exposure: Mean DLP other than routine chest exams

Data Elements	Clinical Performance Measure
<ul><li>DLP (mGy*cm)</li><li>Modality</li></ul>	Measure description: Mean Dose Length Product (DLP) across all screening other than routine chest exams performed
	Numerator: Total DLP for all other than routine chest
	screening exams performed
	Denominator: Number of screening other than routine chest exams
	Numerator and Denominator Exclusion: None

#### 27. Lung Cancer Screening Abnormal Interpretation Rate

Data Elements	Clinical Performance Measure
CT exam result by Lung-RADS category	<b>Measure description:</b> Percent of screening exams that have a Lung-RADS assessment of 3, 4a, 4b, or 4x (assessment categories that may lead to additional imaging or biopsy).
	<b>Numerator</b> : Number of exams that have a Lung-RADS assessment of 3, 4a, 4b, or 4x (assessment categories that may lead to additional imaging or biopsy).
	<b>Denominator</b> : Number of screening exams
	Numerator and Denominator Exclusion: None

#### 28. Lung Cancer Screening Abnormal Interpretation Rate during baseline exam

Data Elements	Clinical Performance Measure

CT exam result by Lung-RADS category
 Measure description: Percent of baseline screening exams that have a Lung-RADS assessment of 3, 4a, 4b, or 4x (assessment categories that may lead to additional imaging or biopsy).
 Numerator: Number of baseline exams that have a Lung-RADS assessment of 3, 4a, 4b, or 4x (assessment categories that may lead to additional imaging or biopsy).
 Denominator: Number of baseline screening exams
 Numerator and Denominator Exclusion: None

#### 29. Lung Cancer Screening Abnormal Interpretation Rate during annual exam

Data Elements	Clinical Performance Measure
CT exam result by Lung-RADS category	Measure description: Percent of annual screening exams that have a Lung-RADS assessment of 3, 4a, 4b, or 4x (assessment categories that may lead to additional imaging or biopsy).  Numerator: Number of annual exams that have a Lung-RADS assessment of 3, 4a, 4b, or 4x (assessment categories that may lead to additional imaging or biopsy).  Denominator: Number of annual screening exams  Numerator and Denominator Exclusion: None

## 30. Cancer Detection Rate (CDR) per 1,000 screening exams

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer within 365 days of screening exam, per 1,000 lung cancer screening exams.  Numerator: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer within 365 days of screening exam.  Denominator: Number of screening exams  Numerator and Denominator Exclusion: None

# 31. CDR for prevalent cancers, detected at baseline exam per 1,000 baseline screening exams

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Indication for exam</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Number of baseline screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer within 365 days of baseline screening exam, per 1,000 baseline screening exams.  Numerator: Number of baseline screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer within 365 days of baseline screening exam.  Denominator: Number of baseline screening exams  Numerator and Denominator Exclusion: None

# 32. CDR for incident cancers, detected at annual exam per 1,000 annual screening exams

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Indication for exam</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Number of annual screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer within 365 days of annual screening exam, per 1,000 annual screening exams.  Numerator: Number of annual screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer within 365 days of annual screening exam.  Denominator: Number of annual screening exams  Numerator and Denominator Exclusion: None

#### 33.CDR stage 0

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> <li>Overall stage</li> </ul>	Measure Description: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 0 within 365 days of screening exam, per 1,000 lung cancer screening exams.  Numerator: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 0 within 365 days of screening exam.
	<b>Denominator</b> : Number of screening exams
	Numerator and Denominator Exclusion: None

## 34. CDR stage IA, IA1, IA2, IA3, IB

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> <li>Overall stage</li> </ul>	Measure Description: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 1 within 365 days of screening exam, per 1,000 lung cancer screening exams.  Numerator: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 1 within 365 days of screening exam.
	<b>Denominator</b> : Number of screening exams
	Numerator and Denominator Exclusion: None

## 35. CDR stage IIA, IIB

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> <li>Overall stage</li> </ul>	Measure Description: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 2 within 365 days of screening exam, per 1,000 lung cancer screening exams.  Numerator: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 2 within 365 days of screening exam.
	<b>Denominator</b> : Number of screening exams
	Numerator and Denominator Exclusion: None

## 36. CDR stage IIIA, IIIB, IIIC

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> <li>Overall stage</li> </ul>	Measure Description: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 3 within 365 days of screening exam, per 1,000 lung cancer screening exams.  Numerator: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 3 within 365 days of screening exam.
	<b>Denominator</b> : Number of screening exams
	Numerator and Denominator Exclusion: None

## 37. CDR stage IV, IVA, IVB

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> <li>Overall stage</li> </ul>	Measure Description: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 4 within 365 days of screening exam, per 1,000 lung cancer screening exams.  Numerator: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of cancer stage 4 within 365 days of screening exam.  Denominator: Number of screening exams  Numerator and Denominator Exclusion: None

## 38.CDR for unknown stage

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> <li>Overall stage</li> </ul>	Measure Description: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of unknown cancer stage within 365 days of screening exam, per 1,000 lung cancer screening exams.  Numerator: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had a tissue diagnosis of unknown cancer stage within 365 days of screening exam.
	<b>Denominator</b> : Number of screening exams
	Numerator and Denominator Exclusion: None

## 39. CDR for presumptive case

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had other than a tissue diagnosis of cancer within 365 days of screening exam, per 1,000 lung cancer screening exams.  Numerator: Number of screening exams with a Lung-RADS assessment category of 3 or 4 that had other than a tissue diagnosis of cancer within 365 days of screening exam.
	<b>Denominator</b> : Number of screening exams
	Numerator and Denominator Exclusion: None

## 40. Positive Predictive Value (PPV1)

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Percent of patients recommended for biopsy who are subsequently diagnosed with lung cancer within 365 days of screening exam  Numerator: Number of screening exams that had a Lung-RADS assessment category of 3 or 4 and a tissue diagnosis of cancer within 365 days of screening exam  Denominator: Number of screening exams that had a Lung-RADS assessment category of 3 or 4  Numerator and Denominator Exclusion: None

## 41. PPV1 for lung cancers detected surgically

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Tissue diagnosis method</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Percent of patients recommended for biopsy who are subsequently diagnosed with lung cancer by surgical tissue diagnosis method within 365 days of screening exam  Numerator: Number of screening exams that had a Lung-RADS assessment category of 3 or 4 and a tissue diagnosis of cancer within 365 days of screening exam where tissue diagnosis method was surgical
	<b>Denominator:</b> Number of screening exams that had a Lung-RADS assessment category of 3 or 4
	Numerator and Denominator Exclusion: None

## 42. PPV1 for lung cancers detected on percutaneous biopsies

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Tissue diagnosis method</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Percent of patients recommended for biopsy who are subsequently diagnosed with lung cancer by percutaneous biopsies within 365 days of screening exam  Numerator: Number of screening exams that had a Lung-RADS assessment category of 3 or 4 and a tissue diagnosis of cancer with in 365 days of screening exam where tissue diagnosis method was percutaneous
	<b>Denominator</b> : Number of screening exams that had a Lung-RADS assessment category of 3 or 4
	Numerator and Denominator Exclusion: None

## 43. PPV1 for lung cancers detected on bronchoscopies

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Tissue diagnosis method</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Percent of patients recommended for biopsy who are subsequently diagnosed with lung cancer by bronchoscopy within 365 days of screening exam  Numerator: Number of screening exams that had Lung-RADS assessment category of 3 or 4 and a tissue diagnosis of cancer within 365 days of screening exam where tissue diagnosis method was bronchoscopy
	<b>Denominator</b> : Number of screening exams that had a Lung-RADS assessment category of 3 or 4 <b>Numerator and Denominator Exclusion</b> : None

## 44. Positive Predictive Value 2a (PPV2a)

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Percent of patients that had a Lung-RADS assessment category of 3 or 4a (typically associated with additional CT recommendation or PET/CT recommendation) and a tissue diagnosis of cancer within 365 days of screening exam  Numerator: Number of screening exams that had a Lung-RADS assessment category of 3 or 4a and a tissue diagnosis of cancer within 365 days of screening exam  Denominator: Number of screening exams that had a Lung-RADS assessment category of 3, 4a  Numerator and Denominator Exclusion: None

## 45. Positive Predictive Value 2b (PPV2b)

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Percent of patients that had a Lung-RADS assessment category of 4B or 4X (typically associated with biopsy recommendation) and a tissue diagnosis of cancer within 365 days of screening exam  Numerator: Number of screening exams that had a Lung-RADS assessment category of 4B or 4X and a tissue diagnosis of cancer within 365 days of screening exam  Denominator: Number of screening exams that had a Lung-RADS assessment category of 4B or 4X  Numerator and Denominator Exclusion: None

## 46. Positive Predictive Value 3 (PPV3)

Data Elements	Clinical Performance Measure
<ul> <li>CT exam result by Lung-RADS category</li> <li>Tissue diagnosis</li> <li>Follow-up diagnostic</li> <li>Exam date/Date of screening</li> <li>Date of follow-up</li> </ul>	Measure Description: Percent of patients that had a Lung-RADS assessment category of 4B or 4X (typically associated with biopsy recommendation) and with record of biopsy performed that had a tissue diagnosis of cancer within 365 days of screening exam  Numerator: Number of screening exams that had a Lung-RADS assessment category of 4B or 4X and with a record of biopsy performed that had a tissue diagnosis of cancer within 365 days of screening exam  Denominator: Number of screening exams that had a Lung-RADS assessment category of 4B or 4X and with record of biopsy performed  Numerator and Denominator Exclusion: None

#### **GLOSSARY OF TERMS**

1. **Abnormal Interpretation Rate**: The percentage of exams interpreted as positive. For lung cancer screening CT, positive exams include Lung-RADS® Categories 3 and 4 assessments.

Abnormal Interpretation Rate = (positive exams) / (all exams)

- 2. **Cancer:** Tissue diagnosis of non-small cell or small cell lung cancer.
- 3. **Cancer Detection Rate:** The number of cancers correctly detected at screening CT per 1,000 patients examined at screening CT. This may be calculated separately for PREVALENT cancers (those found at a baseline or first time lung cancer screening CT exam) and for INCIDENT cancers (those found at subsequent screening exams performed at or close to the recommended screening interval).
- 4. **False Positive (FP):** This includes each of the following three definitions.
  - FP<sub>1</sub>: No known or presumed tissue diagnosis of lung cancer within 1 year of a positive screening exam (Lung-RADS® Category 3 or 4).
  - FP<sub>2</sub>: No known tissue diagnosis of cancer within 1 year after recommendation for biopsy or surgical consultation on the basis of a positive examination (LungRADS® Category 4B or 4X)
  - FP<sub>3</sub>: Benign tissue diagnosis within 1 year after recommendation for biopsy on the basis of a positive examination (LungRADS® Category 4B or 4X).
- 5. **Lung cancer screening CT exam:** is one performed on an asymptomatic individual to detect early, clinically unsuspected lung cancer.
- 6. **Lung RADS category:** This is a quality assurance tool designed to standardize lung cancer screening CT reporting and management recommendations, reduce confusion in lung cancer screening CT interpretations and facilitate outcome monitoring
  - 0: recalls (incomplete screen)
  - 1: normal, continue annual screening
  - 2: benign appearance or behavior, continue annual screening
  - 3: 6 month CT recommended
  - 4A: 3 month CT recommended; may consider PET/CT
  - 4B: Additional diagnostics and/or tissue sampling recommended
  - 4X: Additional diagnostics and/or tissue sampling recommended

\*It is our recommendation that the CT report contain the Lung-RADS category. If the category is not specifically stated in the report, then assign a category to the LCSR registry case record (this is sufficient documentation from the registry perspective)

#### https://www.acr.org/Clinical-Resources/Reporting-and-Data-Systems/Lung-Rads

Lung-RADS™ Version 1.	O Accocoment Categorie	e Dolonco dato: Anvil 20	2014
Lung-KADS version 1.	u Assessment Categorie	s kelease date: Abril 28.	2014

Category	Category Descriptor	Category	Findings	Management	Probability of Malignancy	Estimated Population Prevalence
Incomplete -		0	prior chest CT examination(s) being located for comparison	Additional lung cancer screening CT images and/or	n/a	1%
incomplete		v	part or all of lungs cannot be evaluated	comparison to prior chest CT examinations is needed	n/a	170
Negative	No nodules and definitely benign nodules	1	no lung nodules nodule(s) with specific calcifications: complete, central, popcorn, concentric rings and fat containing nodules		,	
Benign Appearance or Behavior	Nodules with a very low likelihood of becoming a clinically active cancer due to size or lack of growth	2	solid nodule(s):  < 6 mm  new < 4 mm  part solid nodule(s):  < 6 mm total diameter on baseline screening  non solid nodule(s) (GGN):  < 20 mm OR  ≥ 20 mm and unchanged or slowly growing  category 3 or 4 nodules unchanged for ≥ 3 months	Continue annual screening with LDCT in 12 months	<1%	90%
Probably Benign	Probably benign finding(s) - short term follow up suggested; includes nodules with a low likelihood of becoming a clinically active cancer	3	solid nodule(s):  ≥ 6 to < 8 mm at baseline OR new 4 mm to < 6 mm  part solid nodule(s)  ≥ 6 mm total diameter with solid component < 6 mm OR new < 6 mm total diameter non solid nodule(s) (GGN) ≥ 20 mm on baseline CT or new	6 month LDCT	1-2%	5%
additional diagno Suspicious testing and/or tis sampling is	Findings for which additional diagnostic	4A	solid nodule(s):  ≥ 8 to < 15 mm at baseline OR growing < 8 mm OR new 6 to < 8 mm part solid nodule(s:  ≥ 6 mm with solid component ≥ 6 mm to < 8 mm OR with a new or growing < 4 mm solid component endobronchial nodule	3 month LDCT; PET/CT may be used when there is a ≥ 8 mm solid component	5-15%	2%
		4B	solid nodule(s)  ≥ 15 mm OR  new or growing, and ≥ 8 mm  part solid nodule(s) with:  a solid component ≥ 8 mm OR  a new or growing ≥ 4 mm solid component  Category 3 or 4 nodules with additional features or imaging findings that	chest CT with or without contrast, PET/CT and/or tissue sampling depending on the *probability of malignancy and comorbidities. PET/CT may be used when there is a ≥ 8 mm solid component.	> 15%	2%
Other	Clinically Significant or Potentially Clinically Significant Findings (non lung cancer)	5	increases the suspicion of malignancy modifier - may add on to category 0-4 coding	As appropriate to the specific finding	n/a	10%
Prior Lung Cancer	Modifier for patients with a prior diagnosis of lung cancer who return to screening	С	modifier - may add on to category 0-4 coding	٠	120	1211

- 7. **Negative lung cancer screening CT exam** is one that is negative or has benign findings (Lung-RADS® Categories 1 and 2).
- 8. **Per 1,000 exams:** Measures defined *per 1,000 exams* are calculated as below: If there are 12 screening exams with a Lung-RADS category of 3 or 4 that had a tissue diagnosis of cancer within 365 days of screening (numerator) exam out of 2500 screening exams (denominator), the CDR per 1000 exams =12/2500\*1000= 4.8
- 9. **Positive Predictive Value (PPV):** This includes the following three definitions:
  - a.  $PPV_1$  (abnormal findings at screening): The percentage of all positive screening exams (LungRADS® Categories 3 and 4) that result in a tissue diagnosis of cancer within 1 year.

An initial screening assessment of Category 3 or 4 is expected to occur approximately 10% of the time, and is more likely to occur on a baseline lung cancer screening CT exam for which the stability of baseline findings has not yet been established.

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PPV1 = TP / (number of positive screening exams)

OR

PPV1 = TP / (TP + FP1)
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• PPV<sub>2</sub> (additional CT imaging recommended): The percentage of all lung cancer screening CT examinations recommended for 6 month LDCT, or 3 month LDCT with or without PET/CT, (LungRADS® Categories 3 and 4A) that resulted in a tissue diagnosis of cancer within one year.

PPV2 = TP / (number of screening examinations recommended for additional CT imaging)

OR

$$PPV2 = TP/(TP + FP2)$$

PPV<sub>3</sub>: The percentage of all known biopsies done as a result of positive screening with or without subsequent PET/CT or diagnostic chest CT examinations (LungRADS® Categories 4B and 4X) that resulted in a tissue diagnosis of cancer within 1 year. PPV3 is also known as the Biopsy Yield of Malignancy or the Positive Biopsy Rate (PBR).

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PPV3 = TP / (number of biopsies)

OR

PPV3 = TP / (TP + FP3)
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#### 10. Positive screening exam is

- one that requires a 6 month interval low dose chest CT (Lung-RADS® Category 3),
- one that requires a 3 month interval low dose chest CT (Lung-RADS® Category 4A), or
- one that requires chest CT with or without intravenous contrast, PET/CT and/or tissue sampling (Lung-RADS® Category 4B)

- 11. **Tissue diagnosis**: A pathologic diagnosis rendered after any type of interventional procedure, such as CT or fluoroscopy guided biopsy, bronchoscope biopsy, lobar Broncho alveolar lavage or excisional biopsy (VATS or open resection)
- 12. **True Positive (TP):** Tissue diagnosis of cancer within 1 year after a positive exam (Lung-RADS® Category 3 or 4).