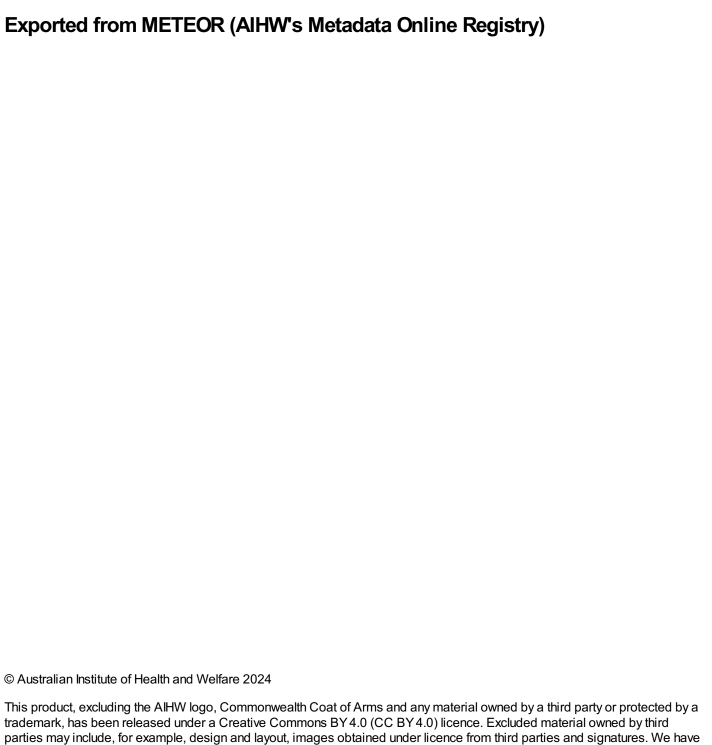
# National Healthcare Agreement: P04-Incidence of selected cancers, 2010



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# National Healthcare Agreement: P04-Incidence of selected cancers, 2010

### Identifying and definitional attributes

Metadata item type: Indicator Indicator type: Indicator

**Short name:** Incidence of selected cancers, 2010

METEOR identifier: 393532

Registration status: Health, Superseded 08/06/2011

**Description:** Incidence of selected cancers of public health importance.

Indicator set: National Healthcare Agreement (2010)

Health, Superseded 08/06/2011

Outcome area: Prevention

Health, Standard 07/07/2010

Data quality statement: National Healthcare Agreement: P04-Incidence of selected cancers, 2010 QS

Health, Superseded 08/06/2011

## Collection and usage attributes

**Computation description:** Selected cancers of public health importance are: melanoma of the skin, bowel

cancer, lung cancer, cervical cancer and breast cancer diagnosed in females.

For melanoma, bowel cancer and lung cancer, the numerator is the number of new primary cases diagnosed in the Australian population in the reported year. The

denominator is the total Australian population for the same year.

For cervical and breast cancer in females, the numerator is the number of new primary cases diagnosed in the Australian female population in the reported year. The denominator is the total Australian female population for the same year.

Calculation is 100,000 × (Numerator ÷ Denominator), calculated separately for each type of cancer, presented as a rate per 100,000 and age-standardised to the Australian population as at 30 June 2001, using 5-year age groups from 0-4 years to 85+ years. Indigenous disaggregations were standardised using 0-4 years to 75+ years due to small populations at older ages in some jurisdictions.

Rates are directly age-standardised to the Australian population as at 30 June

2001.

**Computation:** 100,000 × (Numerator ÷ Denominator)

Calculated separately for each type of cancer.

Presented per 100,000 population.

**Numerator:** For melanoma, bowel and lung cancer: Number of new cases diagnosed in the

reported year

For cervical and breast cancer in females: Number of new cases diagnosed in

females in the reported year

#### Numerator data elements:

#### Data Element / Data Set-

Patient—diagnosis date (cancer), DDMMYYYY

**Data Source** 

Australian Cancer Database

Guide for use

Data source: Administrative by-product data

#### Data Element / Data Set-

Person—person identifier, XXXXXX[X(14)]

**Data Source** 

Australian Cancer Database

Guide for use

Data source: Administrative by-product data

#### Data Element / Data Set-

Episode of care—additional diagnosis, code (ICD-10-AM 7th edn) ANN{.N[N]}

**Data Source** 

Australian Cancer Database

Guide for use

Data source: Administrative by-product data

#### Data Element / Data Set-

Episode of care—principal diagnosis, code (ICD-10-AM 7th edn) ANN{.N[N]}

**Data Source** 

Australian Cancer Database

Guide for use

Data source: Administrative by-product data

**Denominator:** 

For melanoma, bowel and lung cancer: Total population

For cervical and breast cancer in females: Total female population

# Denominator data elements:

#### Data Element / Data Set

#### **Data Element**

Person—estimated resident population of Australia

#### **Data Source**

ABS Estimated resident population (total population)

#### Guide for use

Data source type: Census-based plus administrative by-product data

#### Data Element / Data Set-

#### **Data Element**

Person—estimated resident population of Australia

#### **Data Source**

ABS Indigenous experimental estimates and projections (2001 Censusbased)

#### Guide for use

Data source type: Census-based plus administrative by-product data

#### Data Element / Data Set-

Person-sex, code N

#### **Data Source**

ABS Estimated resident population (total population)

#### Guide for use

Data source type: Census-based plus administrative by-product data

#### Data Element / Data Set-

Person—sex, code N

#### **Data Source**

ABS Indigenous experimental estimates and projections (2001 Censusbased)

#### Guide for use

Data source type: Census-based plus administrative by-product data

# Disaggregation data elements:

#### Data Element / Data Set

Person (address)—Australian postcode, code (Postcode datafile) {NNNN}

**Data Source** 

Australian Cancer Database

Guide for use

Data source: Administrative by-product data

Used for disaggregation by state/territroy, remoteness area and SEIFA of

residence

#### Data Element / Data Set

Person-Indigenous status, code N

**Data Source** 

Australian Cancer Database

Guide for use

Data source: Administrative by-product data

Comments:

Specified disaggregation: Nationally and by state/territory: sex, Indigenous status,

remoteness area and SEIFA of residence.

Available disaggregation: Nationally and by state/territory: Indigenous status,

remoteness area and SEIFA of residence.

Data for 2007 will be available in 2011.

Most recent data available for 2010 CRC baseline report: 2006.

# Representational attributes

Representation class: Incidence

Data type: Real

Unit of measure: Disease type

Format: N[NN].N

## Indicator conceptual framework

Framework and

**Health conditions** 

dimensions:

#### **Data source attributes**

Data sources:

**Data Source** 

ABS Estimated resident population (total population)

Frequency

Quarterly

Data quality statement

ABS Estimated resident population (total population), QS

Data custodian

Australian Bureau of Statistics

#### **Data Source**

Australian Cancer Database

Frequency

Annual

Data custodian

Australian Institute of Health and Welfare

#### **Data Source**

ABS Indigenous experimental estimates and projections (2001 Censusbased)

Frequency

Periodic

Data quality statement

ABS Indigenous experimental estimates and projections, QS

Data custodian

Australian Bureau of Statistics

# **Accountability attributes**

Reporting requirements: National Healthcare Agreement

Organisation responsible

for providing data:

Australian Institute of Health and Welfare

Further data development / collection required:

Specification: Long-term

The quality of Indigenous identification varies by jurisdiction and will require

improvement to enable robust reporting.

Other issues caveats: The quality of Indigenous identification varies between jurisdictions. National

disaggregation by Indigenous status is based on jurisdictions with adequate data

quality.

Remoteness area and SEIFA of residence are based on postcode of residential

address at the time of diagnosis.

Disaggregation by Indigenous status, remoteness area and SEIFA of residence within individual states/territories is subject to data quality

considerations.

Some disaggregations could result in numbers too small for publication.

# Relational attributes

Related metadata references:

Has been superseded by National Healthcare Agreement: PI 04-Incidence of selected cancers, 2011

Health, Superseded 30/10/2011

See also National Healthcare Agreement: P10-Breast cancer screening rates.

2010

Health, Superseded 08/06/2011

See also National Healthcare Agreement: P11-Cervical screening rates, 2010

Health, Superseded 08/06/2011

See also National Healthcare Agreement: P12-Bowel cancer screening rates,

**2010** 

Health, Superseded 08/06/2011