

Person with cancer—personal genetic syndrome indicator, yes/no/unknown/not stated/inadequately described code N

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Person with cancer—personal genetic syndrome indicator, yes/no/unknown/not stated/inadequately described code N

Identifying and definitional attributes

Metadata item type:	Data Element
Short name:	Personal genetic syndrome indicator
METEOR identifier:	457065
Registration status:	Health , Standard 04/02/2015
Definition:	An indicator of whether the person with cancer has been diagnosed with a genetic syndrome , as represented by a code.
Data Element Concept:	Person with cancer—personal genetic syndrome indicator
Value Domain:	Yes/no/unknown/not stated/inadequately described code N

Value domain attributes

Representational attributes

Representation class:	Code	
Data type:	Number	
Format:	N	
Maximum character length:	1	
	Value	Meaning
Permissible values:	1	Yes
	2	No
Supplementary values:	8	Unknown
	9	Not stated/inadequately described

Source and reference attributes

Submitting organisation:	Australian Institute of Health and Welfare
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Data element attributes

Collection and usage attributes

Guide for use:	Record yes when a person has been diagnosed with any genetic syndrome. This includes, but is not exclusive to: • Peutz-Jehgers syndrome • B-Raf mutation • Undescended testes • Von Recklinghausen syndrome • Xeroderma pigmentosa • Bloom syndrome • Ataxia telangiectasia • Beckwith-Wiedemann syndrome • Down syndrome • Klinefelter syndrome • Neurofibromatosis type I (NF-1) • Von Hippel–Lindau disease (VHL) • RET proto-oncogene • Adenomatous polyposis coli (APC) • Hereditary nonpolyposis colorectal cancer (HNPCC or Lynch syndrome) • Breast cancer gene 1 (BRCA1) • Breast cancer gene 2 (BRCA2) • Neurofibromatosis • Li-Fraumeni syndrome • Fanconi anaemia If Yes is selected for personal genetic syndrome indicator, record the type of specific syndrome/s at Person with cancer—hereditary genetic events type, code N[N] or Person with cancer—hereditary genetic events type, text X[X(39)].
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Collection methods: Collect from patient medical records.

Source and reference attributes

Submitting organisation: Cancer Australia

Relational attributes

Related metadata references:	See also Person with cancer—personal genetic syndrome type, genetic event type code N[N] Health, Standard 04/02/2015 See also Person with cancer—personal genetic syndrome type, text X[X(39)] Health, Standard 04/02/2015
Implementation in Data Set Specifications:	Adolescent and young adult cancer (clinical) DSS Health, Superseded 14/05/2015 Adolescent and young adult cancer (clinical) NBPDS Health, Standard 14/05/2015