Person with cancer—hereditary genetic events type, code N[N]

Exported from METEOR (AIHW's Metadata Online Registry)

© Australian Institute of Health and Welfare 2024

This product, excluding the AIHW logo, Commonwealth Coat of Arms and any material owned by a third party or protected by a trademark, has been released under a Creative Commons BY4.0 (CC BY4.0) licence. Excluded material owned by third parties may include, for example, design and layout, images obtained under licence from third parties and signatures. We have made all reasonable efforts to identify and label material owned by third parties.

You may distribute, remix and build on this website's material but must attribute the AIHW as the copyright holder, in line with our attribution policy. The full terms and conditions of this licence are available at https://creativecommons.org/licenses/by/4.0/.

Enquiries relating to copyright should be addressed to info@aihw.gov.au.

Enquiries or comments on the METEOR metadata or download should be directed to the METEOR team at meteor@aihw.gov.au.

Person with cancer—hereditary genetic events type, code N[N]

Identifying and definitional attributes

Metadata item type:	Data Element
Short name:	Hereditary genetic events type code
METEOR identifier:	467855
Registration status:	Health, Standard 04/02/2015
Definition:	The type of hereditary genetic event present in a <u>first degree relative</u> or <u>second</u> <u>degree relative</u> of a person with cancer, as represented by a code.
Data Element Concept:	Person with cancer—hereditary genetic events type
Value Domain:	Genetic event type code N[N]

Value domain attributes

Representational attributes

Representation class:	Code	
Data type:	Number	
Format:	N[N]	
Maximum character length:	2	
	Value	Meaning
Permissible values:	1	Peutz-Jehgers syndrome
	2	B-Raf mutation
	3	Undescended testes
	4	Von Recklinghausen syndrome
	5	Xeroderma pigmentosa
	6	Bloom syndrome
	7	Ataxia telangiectasia
	8	Beckwith-Wiedemann syndrome
	9	Down syndrome
	10	Klinefelter syndrome
	11	Neurofibromatosis type I (NF-1)
	12	Von Hippel-Lindau disease (VHL)
	13	RET proto-oncogene
	14	Adenomatous polyposis coli (APC)
	15	Hereditary nonpolyposis colorectal cancer (HNPCC or Lynch syndrome)
	16	Breast cancer gene 1 (BRCA1)
	17	Breast cancer gene 2 (BRCA2)
	18	Neurofibromatosis
	19	Li-Fraumeni syndrome
	20	Fanconi anaemia
	88	Other genetic events or syndromes
data 1679EE	Dee	

Supplementary values:	97	No genetic events or syndromes
	98	Unknown
	99	Genetic event or syndrome recorded but type inadequately described

Source and reference attributes

Submitting organisation: Cancer Australia

Data element attributes

Collection and usage attributes

 Guide for use:
 Record the type of hereditary genetic event or syndrome that is present in any first degree or second degree relative.

 Collection methods:
 Collect from patient medical records.

Source and reference attributes

Submitting organisation: Cancer Australia

Relational attributes

Related metadata references:	See also <u>Person with cancer</u> <u>family history of hereditary genetic events indicator</u> , <u>yes/no/unknown/not stated/inadequately described code N</u> <u>Health</u> , Standard 04/02/2015
	See also <u>Person with cancer—hereditary genetic events type, text X[X(39)]</u> <u>Health</u> , Standard 04/02/2015
Implementation in Data Set Specifications:	Adolescent and young adult cancer (clinical) DSS Health, Superseded 14/05/2015 Conditional obligation: Complete if Person with cancer—family history of hereditary genetic events indicator, yes/no/unknown/not stated/inadequately described code N equals 'yes'.
	Adolescent and young adult cancer (clinical) NBPDS Health, Standard 14/05/2015 Conditional obligation:
	Complete if <u>Person with cancer—family history of hereditary genetic events</u> indicator, yes/no/unknown/not stated/inadequately described code N equals 'yes'.