- Cancer Registry of the Manche (qualified registry)

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General	
Identification	
Detailed name	Cancer Registry of the Manche (qualified registry)
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	332500
General Aspects	
Medical area	Cancer research

Scientific	investigator(s)
(Contact)	

Others (details)

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Cancer

Collaborations		
Fundina		

Governance of the database	
Details	InVS, l'INCa et l'ARKM
Funding status	Public

Sponsor(s) or organisation(s) responsible	Centre Hospitalier Public du Cotentin
Organisation status	Public

Additional contact	
Main features	
Type of database	
Type of database	Morbidity registers
Database recruitment is carried out by an intermediary	A selection of health institutions and services
Database recruitment is carried out as part of an interventional study	No
Additional information regarding sample selection.	Selection of subjects having the required inclusion criteria. Several sources are used to identify cases: - Anatomopathological laboratories - Information Departments (DIM: PMSI data) - Radiotherapy centers - Permanent cancer survey (EPC) of the CLCC (cancer research centers) - Healthcare institutions (ES) - Specialists (private-practicing and hospital-practicing) and general practitioners - Specialized registries - Summaries of the Pluridisciplinary Concertation Meetings (RCP)
Database objective	
Main objective	The Registry of Cancers for la Manche was created

in 1994. It stems from a local medical need, in order to

supplement the incidence rates measured in the neighboring département of Calvados and to answer the public health questions raised by the industrial activity in the département (the nuclear industry predominantly and the use of asbestos in civil and military ship building).

The registry performs a permanent and complete collection of all of the new diagnoses of cancer in the general population residing in the département of la Manche. It has a dual objective of describing and monitoring the risk of cancer and of conducting research using the analysis of the data collected or one-off surveys.

Objectives of the Registry in terms of public health:

- Participate in the epidemiological monitoring of cancers on a local and national level within the framework of the Francim network, through

	publishing incidence indicators by cancer location, sex, age and year of diagnosis. - Contribute to assessing primary and secondary prevention initiatives (organized cancer screenings), caring for patients and the care needs for the general population.
Inclusion criteria	All malignant tumors diagnosed in patients residing in the département of la Manche at the time of the diagnosis. These are invasive malignant tumors, insitu, benign tumors and intermediate malignancy of the bladder and of the central nervous system. Skin tumors of the basal cell type are excluded.

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Population type	
Age	Newborns (birth to 28 days) Infant (28 days to 2 years) Early childhood (2 to 5 years) Childhood (6 to 13 years) Adolescence (13 to 18 years) Adulthood (19 to 24 years) Adulthood (25 to 44 years) Adulthood (45 to 64 years) Elderly (65 to 79 years) Great age (80 years and more)
Population covered	Sick population
Gender	Male Woman
Geography area	Departmental
French regions covered by the database	Normandie
Detail of the geography area	Department of La Manche
Data collection	
Dates	
Date of first collection (YYYY or MM/YYYY)	1994
Date of last collection (YYYY or MM/YYYY)	2008
Size of the database	

individuals)

Size of the database (number of Greater than 20 000 individuals

Details of the number of 1994-2008: 45 078 individuals Data Data collection completed Database activity Type of data collected Clinical data Administrative data Clinical data (detail) Direct physical measures Administrative data (detail) Identification data Presence of a biobank No Health parameters studied Health event/morbidity Health event/mortality **Procedures** Data collection method Active or passive collection (CNIL Authorization): -Reception of the anatomocytopathological reports, lists of stays (PMSI). Lists via encrypted computer transmission. - Active search for additional regular information in the medical records. Participant monitoring Yes - Vital status, date of latest news - An active follow-Details on monitoring of up is performed on a sample (High Resolution participants Francim surveys). - A passive follow-up is conducted using information received at the Registry. Links to administrative sources Yes Promotion and access Promotion

Access

Terms of data access (charter for data provision, format of data, availability delay) The incidence data is available in an aggregate form, by cancer location, year, sex and age. These are published on a regular basis on a local and national level (cf. Find out more). Regional documents can be downloaded from the registry website. The data recorded in the Registry database can be used for specific studies. The methods for collaboration have to be defined.

Access to aggregated data	Access on specific project only
Access to individual data	Access on specific project only