- CALVADOS GENERAL CANCER REGISTRY (CERTIFIED REGISTRY 2013-2016)

Head :Guizard Anne-Valérie, Registre Général des Tumeurs du Calvados et Enquête Permanente Cancer

Governance of the database

Last update : 04/12/2012 Version : 3 ID : 1035		
General		
Identification		
Detailed name	CALVADOS GENERAL CANCER REGISTRY (CERTIFIED REGISTRY 2013-2016)	
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	115.299	
General Aspects		
Medical area	Cancer research	
Scientific investigator(s) (Contact)		
Name of the director	Guizard	
Surname	Anne-Valérie	
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Unit	Registre Général des Tumeurs du Calvados et Enquête Permanente Cancer	
Organization	Centre de Lutte Contre le Cancer François	
Collaborations		
Funding		
Funding status	Public	
Details	InVS, INCA and INSERM.	

Sponsor(s) or organisation(s) responsible

Centre de Lutte Contre le Cancer François Baclesse

Organisation status

Private

Additional contact

Main features

Type of database

Type of database

Morbidity registers

Additional information regarding sample selection.

Selection of subjects meeting the required inclusion criteria. Several sources used to identify cases:Anatomopathology laboratories - Private clinics (medical information department/DIM)- General hospitals (DIM)- Cancer Prevention Centres- Private radiotherapy centres- University Hospital Centres (DIM)- Paediatric oncology department- Health Insurance Funds- Minutes from the Multidisciplinary Consultation Meeting.

Database objective

Main objective

The main purpose of the registry is to produce, through active and exhaustive collection of cases, data enabling indicators to be calculated for monitoring population health (incidence, survival and prevalence). In this regard, the Registry takes part in the national cancer surveillance programme from the French Institute for Public Health Surveillance (InVS).

The Registry's data is included in the database of the French cancer registry network (FRANCIM), which is managed by the Biostatistics Laboratory of the Lyon-Sud Faculty of Medicine in connection with the InVS. It is used to estimate cancer incidence across France as a whole and per region, and to study incidence trends over the last 25 years.

It is also used to estimate prevalence and analyse survival on a regular basis by the FRANCIM network for France and by the EUROCARE group for Europe.

The registry data is also sent at regular intervals to the International Agency for Research on Cancer (Lyon) for publication in Cancer Incidence in Five Continents from the first available five-year period.

The Registry takes part in cooperative studies conducted as part of the FRANCIM and EUROCARE networks aimed at evaluating medical practices in

the population, providing recommendations at Consensus conferences and measuring their impact on public health. These studies focussed on the following sites: thyroid, breast, lung, prostate, testicle and skin melanoma.

Objectives of the registry in terms of research:

- Social inequalities and access to healthcare;
- Reintegration of patients in long-term remission or who have recovered from cancer;
- Pesticide exposure and cancer risk in the agricultural environment (EPI95 cohort).

Inclusion criteria

New cases of invasive and in situ cancer as well as all brain tumours, regardless of behaviour (except digestive cancer).

Basocellular skin carcinoma have not been recorded since 1990:

Haematological diseases have not been recorded since 2001 (Lower Normandy Regional Registry for Haematological Diseases creation date).

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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

Calvados

Data collection

Dates

Date of first collection (YYYY or MM/YYYY)

1978

Size of the database	
Size of the database (number of individuals)	Greater than 20 000 individuals
Details of the number of individuals	61,910 incident cases since beginning of registration.
Data	
Database activity	Current data collection
Type of data collected	Clinical data Administrative data
Clinical data (detail)	Direct physical measures
Administrative data (detail)	- Last name, first name - maiden name - date of birth - town (country) of birth- address- sex - profession at the time of diagnosis.
Presence of a biobank	No
Health parameters studied	Health event/morbidity Health event/mortality
Procedures	
Data collection method	- Computer file - paper files- database consultation.
Participant monitoring	Yes
Details on monitoring of participants	Vital status.
Links to administrative sources	No
Promotion and access	
Promotion	
Link to the document	http://tinyurl.com/PUBMED-CGCR
Description	Liste des publications dans Pubmed
Access	
Terms of data access (charter for data provision, format of data, availability delay)	Incidence data is available in aggregate form according to cancer location, year, sex and age. It is published at regular intervals at local and national level, French Institute for Public Health Surveillance/InVS and IARC.

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