

- Finistère Registry for Digestive Tumours (Certified Registry 2013-2016)

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General

Identification

Detailed name	Finistère Registry for Digestive Tumours (Certified Registry 2013-2016)
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	908270

General Aspects

Medical area	Cancer research Gastroenterology et hepatology
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Scientific investigator(s) (Contact)

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Collaborations

Funding

Funding status	Public
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Details	French Institute for Public Health Surveillance - INVS National Cancer Institute - InCA League
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Governance of the database

Sponsor(s) or organisation(s) responsible	CHU Brest
Organisation status	Public

Additional contact

Main features

Type of database

Type of database	Morbidity registers
Additional information regarding sample selection.	Selection of subjects meeting the inclusion criteria. Several sources are used to identify cases:- Clinical departments - University Hospital Centres (UHC)- Other public clinical departments- Private clinical departments- Private gastroenterologist practices ?

General practitioner surgeries - Medical Information Department (DIM) of the UHC - DIM of other institutions - Anatomopathology laboratory - Regional oncology networks ? Civil registry - National PMSI database - Local medical departments of various health insurance funds (CPAM in Finistère, MSA, RSI, etc.).

Database objective

Main objective

Objectives of the registry in terms of public health (surveillance, evaluation):
Thanks to the multi-source collection of digestive cancer cases, the registry's aim is to produce quality data for calculating health indicators: it allows for the exact measurement of incidence trends for digestive cancers depending on the tumour site or anatomopathological type as well as patient survival. By recording incident cancer cases, in the years to come, we will be able to evaluate the efficacy of HémoCCult screening for colorectal cancer, conducted in the département since June 2004 (Cancer Screening Association ADEC 29).

- Objectives of the registry in terms of research:
The digestive tumour registry has already been used for medical theses and publications. Locally, incidence and survival studies are undertaken on a regular basis by the registry's team. We also have projects with local partners: the Brest University Hospital Centre team, University of West Brittany (prognostic factor of statin prescription on the occurrence of cancers).

The registry's data covers the entire validated period (1984-2006) and is sent annually to the national cancer incidence base managed by the FRANCIM network, the French Institute for Public Health Surveillance and Lyon Civil Hospices. As such, we are participating in the national cancer surveillance programme and contributing to national research.

Inclusion criteria

The registry identifies all patients affected with a primitive malignant tumour, including in situ cancer of the digestive tract, liver, pancreas, biliary tracts, peritoneum and retroperitoneal tissue.

The date of incidence must fall after 01 January 1984.

Patients must live in Finistère and consent to the registration of their cases.

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)
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Population covered	Sick population
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Gender	Male Woman
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Geography area	Departmental
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French regions covered by the database	Bretagne
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Detail of the geography area	Finistère and bordering départements.
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Data collection

Dates

Date of first collection (YYYY or MM/YYYY)	1984
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Size of the database

Size of the database (number of individuals)	Greater than 20 000 individuals
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Details of the number of individuals	1984-2010: 27371 reported cases.
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Data

Database activity	Current data collection
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Type of data collected	Clinical data Paraclinical data Biological data Administrative data
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Clinical data (detail)	Direct physical measures
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Paraclinical data (detail)	Medical imaging.
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Biological data (detail)	Tumour markers.
Administrative data (detail)	Identification data.
Presence of a biobank	No
Health parameters studied	Health event/morbidity Health event/mortality

Procedures

Data collection method	Active, the registry shall research information from different sources.
Classifications used	ICD-O'3
Participant monitoring	Yes
Details on monitoring of participants	- Systematic for invasive malignant tumour and in situ carcinoma cases - vital status.
Links to administrative sources	Yes
Linked administrative sources (detail)	RNIPP.

Promotion and access

Promotion

Link to the document	http://tinyurl.com/PUBMED-RFTD
Description	Liste des publications dans Pubmed

Access

Terms of data access (charter for data provision, format of data, availability delay)	Data promotion and distribution through scientific publications and oral communication.
Access to aggregated data	Access on specific project only
Access to individual data	Access on specific project only