- Hérault Cancer Registry (Certified Registry 2010-2013)

Head :Trétarre Brigitte

Sponsor(s) or organisation(s)

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General		
Identification		
Detailed name	Hérault Cancer Registry (Certified Registry 2010- 2013)	
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	997109	
General Aspects		
Medical area	Cancer research	
Health determinants	Geography	
Keywords	population registry, survival, incidence, mortality	
Scientific investigator(s) (Contact)		
Name of the director	Trétarre	
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Collaborations		
Funding		
Funding status	Public	
Details	InVS and INCA.	
Governance of the database		

Centre de Lutte Contre le Cancer Val d'Aurelle

responsible

Organisation status

Both

Additional contact

Main features

Type of database

Type of database

Additional information regarding sample selection.

Morbidity registers

Selection of subjects fulfilling the required inclusion criteria. Several sources are used to identify cases:-anatomopathology laboratories- private radiotherapy practices- haematology laboratory - regional oncology network (ONCO-LR)- organised mass screening facilities for breast cancer and colon cancer (34 Screening and IMIM)- European Randomised Study of Screening for Prostate Cancer (ERSPC) - PMSI data - health insurance funds - clinical departments (3 private and 3 public) - national brain tumour registration - multidisciplinary urology meetings - Toulouse Cancer Prevention Centre (CLCC) - Castres Hospital- Nîmes University Hospital Centre (UHC) - Gustave Roussy Institute.

Database objective

Main objective

Objectives of the registry in terms of public health (surveillance, assessment):

The tumour registry is the only facility that can assess all of the actions taken over the last 20 years within the département in the field of prevention and screening (breast cancer, prostate cancer, colon-rectum cancer).

The registry monitors incidence and survival for all cancers over time according to all prognostic factors, and provides information on all interval cancers occurring after a negative screening examination.

- Objectives of the registry in terms of research: The registry is conducting a number of studies (alone or together with other French or European registry networks) to address the need for epidemiological knowledge:
- Descriptive epidemiology: all data concerning cancer in the Hérault département is published and distributed at regular intervals to the whole medical community every two years i.e., incidence and mortality data.
- Analytical epidemiology: relative survival,

prevalence, cohort and case-control studies, interval cancers.

- Prognostic factor studies: medico-economic studies, studies on therapies and healthcare channels.

Data from the registry is used by the FRANCIM network in France and by the EUROCARE group at European level to estimate national and regional cancer incidence and prevalence in France, as well as survival analysis.

Inclusion criteria

All new invasive tumours, as well as in situ breast, cervix, colorectal and bladder tumours, skin melanomas, bladder pTa and benign tumours of the central nervous system that are diagnosed in individuals living in Hérault, regardless of where they are treated. Only basocellular skin carcinoma are excluded from the registry. Contralateral breast tumours in the same patient are recorded but not counted in incidents, and are not sent to the FRANCIM common database, in accordance with the rules defined by the European Network Cancer Registry (ENCR) and FRANCIM (French cancer registry network certified by the National Registries Committee).

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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 Languedoc-Roussillon Midi-Pyrénées database

Detail of the geography area Hérault

Data collection

Dates	
Date of first collection (YYYY or MM/YYYY)	1985
Size of the database	
Size of the database (number of individuals)	Greater than 20 000 individuals
Details of the number of individuals	1987-2008: 96,535 patients, 104,252 cases (95,136 invasive).
Data	
Database activity	Current data collection
Type of data collected	Clinical data Paraclinical data Biological data Administrative data
Clinical data (detail)	Direct physical measures
Paraclinical data (detail)	Paraclinical examinations for determining tumour stage.
Biological data (detail)	Tumour histology, grade, 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 or passive collection: - computerised files - computerised listing - paper factsheets - paper listing.
Participant monitoring	Yes
Details on monitoring of participants	Survival.
Links to administrative sources	Yes
Linked administrative sources (detail)	Cépi DC, RNIPP.

Promotion and access	
Promotion	
Link to the document	http://www.registre-tumeurs-herault.fr/
Link to the document	http://tinyurl.com/PUBMED-RTH
Description	Liste des publications dans Pubmed
Link to the document	http://tinyurl.com/HAL-RTH
Description	Liste des publications dans HAL
Access	
Terms of data access (charter for data provision, format of data, availability delay)	Website: downloadable data.
Access to aggregated data	Free access
Access to individual data	Access on specific project only