## - Limousin Region General Cancer Registry (CERTIFIED REGISTRY 2015-2020)

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General	
Identification	
Detailed name	Limousin Region General Cancer Registry (CERTIFIED REGISTRY 2015-2020)
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	999305
General Aspects	
Medical area	Cancer research
Health determinants	Medicine
Keywords	oncogeriatrics
Scientific investigator(s) (Contact)	
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Unit	Pôle Biologie Cancer
Organization	CHU de Limoges
Collaborations	
Funding	
Funding status	Public

Details	French Institute for Public Health Surveillance (InVS), French National Cancer Institute (INCa) and the Limousin Regional Health Agency (ARS).
Governance of the database	
Sponsor(s) or organisation(s) responsible	CHU, Hôpital Le Cluzeau, Limoges
Organisation status	Public
Additional contact	
Main features	
Type of database	
Type of database	Morbidity registers
Additional information regarding sample selection.	Selection of subjects fulfilling required criteria. Several sources are used to identify cases: - Anatomic cytopathology laboratories - molecular, haematology and cytogenetic laboratories - medical information departments (PMSI) - health insurance services (ALD) - summaries from multidisciplinary meetings (RCP) - clinical services - treating physicians - private radiotherapy centres - childhood cancer registries.
Database objective	
Main objective	Registry objectives regarding public health (surveillance, assessment): - To participate in epidemiological cancer surveillance locally (Haute-Vienne and Limousin region), nationally (French network of cancer registries, FRANCIM) and internationally (European Network of Cancer Registries (ENCR)) and the International Agency for Research on Cancer (IARC) by publishing data on incidence, prevalence and survival according to location, sex, age and year of diagnosis. The registry may also identify possible cancer clusters and contribute to studies on measuring population exposure to probable or possible carcinogens, as well as the existence and nature of the causal relationship. Both Limousin and Haute-Vienne are concerned by the presence of natural radioactivity (radon) and former uranium mining sites.

	<ul> <li>second oldest European region (after Liguria),</li> <li>Limousin is an indication of the future of geriatric oncology in France.</li> <li>Registry research objectives: <ul> <li>To participate in the FRANCIM network in national survival studies and "high resolution" studies on cancer treatment.</li> <li>To identify specific demographic traits in the territory covered by the registry that may further expertise in geriatric oncology.</li> </ul> </li> </ul>
Inclusion criteria	All cases of primary invasive tumours (haematological malignancies and solid tumours, except basocellular skin carcinoma), in situ tumours (breast, cervix, colorectal, bladder and urinary tract, melanoma), benign tumours or unpredictable developments in the central nervous system and the bladder, as well as borderline tumours in the ovaries occurring in patients living in Haute-Vienne at the time of diagnosis, regardless of treatment location.
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	Aquitaine Limousin Poitou-Charentes
Detail of the geography area	Haute-Vienne, 375,914 inhabitants as of 01 January 2009, source: INSEE.
Data collection	
Dates	
Date of first collection (XXXX or	2003

MM/YYYY)	
Date of last collection (YYYY or MM/YYYY)	2008
Size of the database	
Size of the database (number of individuals)	[10 000-20 000[ individuals
Details of the number of individuals	2003-2008 : 14,381
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)	All further examinations that provide information on tumour coding.
Biological data (detail)	All further examinations that provide information on tumour coding.
Administrative data (detail)	Identification data (surname, first name, date of birth, place of birth, address).
Presence of a biobank	No
Health parameters studied	Health event/morbidity Health event/mortality
Procedures	
Data collection method	Active or passive collection (CNIL approval): - receipt of anatomic cytopathological and cytohaematological reports, lists of admissions (PMSI) and long-term diseases. Data transferred by encrypted digital files active search for additional information in medical files.
Classifications used	CIM-O, TNM.
Participant monitoring	Yes
Monitoring procedures	Monitoring by contact with the referring doctor

	Monitoring by crossing with a medical- administrative database
Details on monitoring of participants	Vital status, death, date of latest developments. Active follow-up is carried out on the sample (FRANCIM "High Resolution" surveys). Passive follow-up is carried out on information received by the registry.
Links to administrative sources	Yes
Linked administrative sources (detail)	PMSI, ALD, RNIPP.
Promotion and access	
Promotion	
Link to the document	http://www.ncbi.nlm.nih.gov/pubmed/22123138
Link to the document	http://www.ncbi.nlm.nih.gov/pubmed/18449095
Link to the document	http://www.ncbi.nlm.nih.gov/pubmed/12108336
Link to the document	http://www.ncbi.nlm.nih.gov/pubmed?term
Link to the document	http://tinyurl.com/PUBMED-RGCRL
Link to the document	<u>Survie-des-personnes-atteintes-de-cancer-France-</u> <u>metropolitaine-1989-2013-tumeurs-</u> <u>solides_2016.pdf</u>
Access	
Terms of data access (charter for data provision, format of data, availability delay)	Incidence data are available in aggregate format on the registry website, according to year, location, sex and age. Haute-Vienne data is included in the common French network cancer registry database: data may be accessed through FRANCIM. Data are published locally (in collaboration with the Limousin Health Monitoring Agency (ORS)), internationally (FRANCIM publications) and internationally (ENCR).
Access to aggregated data	Free access
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Access to individual data

Access on specific project only