## **RCT - Tarn Cancer Registry (Certified Registry 2020-2025)**

Head :GROSCLAUDE Pascale, U1027 EPIDEMIOLOGIE ET ANALYSES EN SANTE PUBLIQUE : RISQUES, MALADIES CHRONIQUES ET HANDICAPS

Daubisse-Marliac Laetitia, U1027 EPIDEMIOLOGIE ET ANALYSES EN SANTE PUBLIQUE : RISQUES, MALADIES CHRONIQUES ET HANDICAPS

Last update : 10/29/2020 | Version : 2 | ID : 172

Last update . 10/29/2020   Version . 2   ID	. 1/2
General	
Identification	
Detailed name	Tarn Cancer Registry (Certified Registry 2020-2025)
Sign or acronym	RCT
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	99 80 15 (12/1998), 99 80 15 Version 2 (10/2003)
General Aspects	
Medical area	Cancer research
Keywords	Francim, cancer epidemiology
Scientific investigator(s) (Contact)	
Name of the director	GROSCLAUDE
Surname	Pascale
Phone	+33 (0)5 63 47 59 51
Email	pascale.grosclaude@inserm.fr; registre81@orange.fr
Unit	U1027 EPIDEMIOLOGIE ET ANALYSES EN SANTE PUBLIQUE : RISQUES, MALADIES CHRONIQUES ET HANDICAPS
Organization	INSERM - Institut National de la Santé et de la Recherche Médicale
Name of the director	Daubisse-Marliac
Surname	Laetitia

Address	11 rue Lavazière 81012 Albi cedex
Phone	05 63 47 59 51
Email	laetitia.daubisse-marliac@inserm.fr
Unit	U1027 EPIDEMIOLOGIE ET ANALYSES EN SANTE PUBLIQUE : RISQUES, MALADIES CHRONIQUES ET HANDICAPS
Organization	INSERM - Institut National de la Santé et de la Recherche Médicale
Collaborations	
Funding	
Funding status	Public
Details	Inca, SPF, Tarn general council, French League against Cancer.
Governance of the database	
Sponsor(s) or organisation(s) responsible	Institut Claudius Regaud (Centre Régional de lutte contre le cancer)
Organisation status	Public
Sponsor(s) or organisation(s) responsible	Association recherche en épidémiologie et prévention
Organisation status	Private
Additional contact	
Main features	
Type of database	
Type of database	Morbidity registers
Database objective	
Main objective	In the area of public health: (1) To produce quality data to calculate health indicators for incidence, survival and prevalence. (2) To participate in the French network of cancer registries (FRANCIM) and to collaborate with SPF and INCa as part of the national cancer surveillance programme. (3) To work with local stakeholders at a regional level

(ARS, Regional cancer network, screening sites) and to concentrate on the local use of the generated knowledge (SROS cancer research, assessment of requirements, initial and continuous training).

(4) To assess the impact of the disease on the population scale: prevalence, survival with or without the disease.

In the area of research:

(1) To learn about the distribution of therapeutic innovations throughout the population and its impact from a health and economic standpoint (2) To assess the heterogeneity factors in the distribution of cancer treatment developments and their impact in terms of social health inequality in primary and secondary healthcare usage.
(3) To participate in studies initiated by others by providing them with local knowledge from the registry team and data collected in Tarn.

Inclusion criteria

Data collection

All cases of invasive malignant tumors (except basocellular skin carcinoma) + in situ breast, colorectal, cervical, urinary tract and borderline ovarian tumours.

	ovarian cumours.
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
Pathology	C00-C97 - Malignant neoplasms
Gender	Male Woman
Geography area	Departmental
French regions covered by the database	Languedoc-Roussillon Midi-Pyrénées
Detail of the geography area	Tarn

Dates	
Date of first collection (YYYY or MM/YYYY)	1982
Size of the database	
Size of the database (number of individuals)	Greater than 20 000 individuals
Details of the number of individuals	60,000
Data	
Database activity	Current data collection
Type of data collected	Clinical data Paraclinical data Biological data Administrative data
Clinical data (detail)	Direct physical measures Medical registration
Paraclinical data (detail)	Diagnostic assessment, anatomopathological report.
Biological data (detail)	Markers.
Administrative data (detail)	Surname, first name, date and place of birth, address.
Presence of a biobank	No
Health parameters studied	Health event/morbidity Health care consumption and services
Care consumption (detail)	Hospitalization
Procedures	
Data collection method	Systematic collection of information to identify cases is carried out using: (1) ALD information sources (2) PMSI information sources (3) the regional cancer treatment network (4) anatomic pathology laboratories (6) clinical and hospital departments (7) radiotherapy centres (8) specialist practices and (9) electoral registers.
Classifications used	CIM-O; TNM

Quality procedure(s) used	FRANCIM procedure and ENCR rules
Participant monitoring	Yes
Details on monitoring of participants	Vital status.
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
Access	
Access  Dedicated website	https://www.onco-occitanie.fr/pro/page/registredes-cancers-du-tarn
Dedicated website  Terms of data access (charter for data provision, format of	des-cancers-du-tarn