## - General Cancer Registry of French Polynesia

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General	

#### Identification

Detailed name General Cancer Registry of French Polynesia

CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation

**CNIL** 

## **General Aspects**

Medical area Cancer research

Health determinants Geography

Keywords Polynesia, incidence, diagnosis, screening

### Scientific investigator(s) (Contact)

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Institut National de la Santé et de la Recherche Organization

#### Collaborations

Funding

Funding status	Public
Details	Health Directorate of French Polynesia
Governance of the database	
Sponsor(s) or organisation(s) responsible	Direction de la Santé de la Polynésie française
Organisation status	Public
Sponsor(s) or organisation(s) responsible	Office de Protection contre les Rayonnements Ionisants
Organisation status	Public
Sponsor(s) or organisation(s) responsible	Institut National de la Santé et de la Recherche Médicale
Organisation status	Public
Additional contact	
Main features	
Type of database	
Type of database	Morbidity registers
Additional information regarding sample selection.	Cancer cases are identified based on reports from private and public physicians, examination reports from anatomopathology laboratories throughout the territory, PMSI data from the Mammao Regional Hospital Centre (Centre Hospitalier Territorial de Mammao), medical evacuations and death certificates.
Database objective	
Main objective	To identify all new cancer cases among individuals living in French Polynesia. To conduct descriptive, analytic and evaluative epidemiological work with this data. Assessment of healthcare and diagnostic practices. Assessment of screening practices. Participation in etiological and prognostic research.
Inclusion criteria	<ul><li>male and female</li><li>living in French Polynesia at time of diagnosis</li><li>subject with cancer</li></ul>

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
Detail of the geography area	French Polynesia
Data collection	
Dates	
Date of first collection (YYYY or MM/YYYY)	1985
Size of the database	
Size of the database (number of individuals)	[10 000-20 000[ individuals
Details of the number of individuals	475/an/year
Data	
Database activity	Current data collection
Type of data collected	Clinical data
Clinical data (detail)	Direct physical measures
Details of collected clinical data	Patient sociodemographic characteristics (surname, maiden name, first name, age, sex, place of residence, date and place of birth, vital status)
	Cancer morphology and topography. Patient's vital status.
Presence of a biobank	Cancer morphology and topography. Patient's vital

# Health event/mortality

Procedures	
Data collection method	Data are gathered through a report form, including patient identification, topographical characteristics and tumour histology, treatment, monitoring and report source.
Quality procedure(s) used	- Internal quality control for data validation by establishing quality indicators once a year Using quality criteria from the FRANCIM network and French National Committee of Registries Data validation protocol: comprehensive consultation of data sources from laboratories and DIMs (Medical Information Departments); missing and duplicated data control, inconsistency control performed using IARC Tools and corrections carried out.
Participant monitoring	Yes
Links to administrative sources	No
Promotion and access	
Promotion and access Promotion	
	http://onlinelibrary.wiley.com/doi/10.1046/j.1365- 3156.2000.00624.x/full
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