

France Cancer Barometer - Cross-Sectional Study on Patients Treated in French Healthcare Institutions

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General

Identification

Detailed name	Cross-Sectional Study on Patients Treated in French Healthcare Institutions
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Sign or acronym	France Cancer Barometer
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CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	1493177
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General Aspects

Medical area	Cancer research Hematology
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Pathology (details)	By body: epidemiology, current treatments, stage at diagnosis, current stage, patient characteristics (age, sex, ECOG), clinical profile of disease (histology, changes, metastases, etc.)
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Health determinants	Iatrogenic Medicine
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Keywords	clinical profile, agents (chemotherapy/targeted therapy), haematology, oncology, therapeutic strategies
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Scientific investigator(s) (Contact)

Name of the director	Bonnelye
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Surname	Geneviève
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Organization	Kantar Health
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Organization	Kantar Health
Collaborations	
Funding	
Funding status	Private
Details	Kantar Health
Governance of the database	
Sponsor(s) or organisation(s) responsible	Kantar Health France
Organisation status	Private
Additional contact	
Main features	
Type of database	
Type of database	Study databases
Study databases (details)	Repeated cross-sectional studies (except case control studies)
Database recruitment is carried out by an intermediary	A selection of health institutions and services
Database recruitment is carried out as part of an interventional study	No
Additional information regarding sample selection.	quotas
Database objective	

Main objective	To determine the epidemiology, clinical profile and therapeutic strategies across 18 procedures (oncology and haematology).
Inclusion criteria	Patients currently treated within the active cancer population.
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	National
Detail of the geography area	France
Data collection	
Dates	
Date of first collection (YYYY or MM/YYYY)	1993
Date of last collection (YYYY or MM/YYYY)	2013
Size of the database	
Size of the database (number of individuals)	[1000-10 000[individuals
Details of the number of individuals	4,430
Data	
Database activity	Data collection completed
Type of data collected	Clinical data Declarative data

Biological data

Clinical data (detail)	Direct physical measures
Details of collected clinical data	metastases, stage, treatment
Declarative data (detail)	Paper self-questionnaire Face to face interview
Biological data (detail)	changes, histology
Presence of a biobank	No
Health parameters studied	Health event/morbidity Quality of life/health perception Others
Quality of life/perceived health (detail)	administration criteria, support treatment, healthcare locations, associated anti-tumour treatment, clinical profile (histology, changes, metastases, procedures, stage at diagnosis and current stage).)
Other (detail)	administration criteria, support treatment, healthcare locations, associated anti-tumour treatment, clinical profile (histology, changes, metastases, procedures, stage at diagnosis and current stage).

Procedures

Data collection method	Face-to-face/self-administered questionnaire/registry/patient data sheets Data collected every 2 years
Quality procedure(s) used	ISO 20-232
Participant monitoring	No
Links to administrative sources	Yes
Linked administrative sources (detail)	DREES

Promotion and access

Promotion

Access

Terms of data access (charter for data provision, format of	Contact the scientist in charge at Kantar Health France.
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data, availability delay)

Access to aggregated data

Access on specific project only

Access to individual data

Access on specific project only