

SEINON - Registry on Breast Cancer Patients from The Antoine Lacassagne Centre

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General

Identification

Detailed name Registry on Breast Cancer Patients from The Antoine Lacassagne Centre

Sign or acronym SEINON

CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation CIL declaration (Mr. Olivier Prou)

General Aspects

Medical area Cancer research

Health determinants Genetic
Iatrogenic
Medicine

Keywords breast cancer, survival, metastasis, recurrence, characteristics, treatment

Scientific investigator(s) (Contact)

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Unit Epidemiology and Biostatistics Unit (UEB)

Organization Antoine Lacassagne Centre

Collaborations

Funding

Funding status	Public
Details	Antoine Lacassagne Centre
Governance of the database	
Sponsor(s) or organisation(s) responsible	Centre Antoine Lacassagne
Organisation status	Public
Additional contact	
Main features	
Type of database	
Type of database	Morbidity registers
Database objective	
Main objective	The general objective is to provide data on women with breast cancer. These data will be very useful for developing epidemiological studies.
Inclusion criteria	All patients affected with a first breast cancer and operated on at the Antoine Lacassagne Centre (ALC) were entered in the database. The database has since expanded and includes all patients at the ALC for breast cancer, regardless of whether it is primary or recurring.
Population type	
Age	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	Woman
Geography area	Local
French regions covered by the database	Provence - Alpes - Côte d'Azur
Detail of the geography area	Antoine Lacassagne Centre (Nice)
Data collection	

Dates

Date of first collection (YYYY or MM/YYYY) 1975

Date of last collection (YYYY or MM/YYYY) 2015

Size of the database

Size of the database (number of individuals) [10 000-20 000[individuals

Details of the number of individuals 12,739

Data

Database activity Current data collection

Type of data collected
Clinical data
Paraclinical data
Biological data
Administrative data

Clinical data (detail) Medical registration

Details of collected clinical data Pathology; surgery; treatment; biopsies; latest updates

Paraclinical data (detail) Radiological examination

Biological data (detail) HER2; mutations

Administrative data (detail) Only last name; first name and date of birth

Presence of a biobank Yes

Contents of biobank Tissues

Health parameters studied
Health event/mortality
Health care consumption and services

Care consumption (detail)
Hospitalization
Medicines consumption

Procedures

Data collection method Initial creation and collection via Access® and then by Clinsight® software.

Participant monitoring Yes

Details on monitoring of participants	Updates to patient records carried out as part of therapeutic treatment at ALC.
Links to administrative sources	Yes
Promotion and access	
Promotion	
Link to the document	http://www.ncbi.nlm.nih.gov/entrez/eutils/erss.cgi?rss_guid
Link to the document	http://www.ncbi.nlm.nih.gov/entrez/eutils/erss.cgi?rss_guid
Access	
Terms of data access (charter for data provision, format of data, availability delay)	Data cannot be accessed externally. Requests must be sent to Mr. Chamorey if researchers wish to conduct ancillary research. Data export may only be carried out upon signing an agreement (e.g. ALC internal agreement) and obtaining approval on Chapters IX or X from the CNIL. Charges may be applied.
Access to aggregated data	Access on specific project only
Access to individual data	Access on specific project only