## **PROGENE - National Study on Prostate Cancer**

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Last update : 08/12/2014   Version : 1   ID :	: 60134
General	
Identification	
Detailed name	National Study on Prostate Cancer
Sign or acronym	PROGENE
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	CNIL: 27/06/2001
General Aspects	
Medical area	Cancer research
Health determinants	Genetic Social and psychosocial factors
Others (details)	Prostate cancer
Keywords	anatomopathological, data, weight, height, Health episodes, clinical, family history
Scientific investigator(s) (Contact)	
Name of the director	Cussenot
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Unit	CEREPP
Organization	APHP

## Collaborations

Funding

Funding status	Mixed
Details	AP-HP CEREPP
Governance of the database	
Sponsor(s) or organisation(s) responsible	CEREPP
Organisation status	Private
Additional contact	
Main features	
Type of database	
Type of database	Study databases
Study databases (details)	Cohort study
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.	Inclusion method: Prospective Other bodies active in creating this cohort: CHU, CHG, INDEPENDENT PHYSICIANS, CNAM
Database objective	
Main objective	General objective: To initiate a collection of informative families with at least one member with prostate cancer and apparently health subjects. Secondary objectives: - To identify genes for predisposition and susceptibility to prostate cancer - To identify genetic markers associated with recurrence after treatment - To develop a genetic screening test for these pathologies
Inclusion criteria	Men with histologically verified prostate cancer Healthy subjects with PSA levels below 4 ng/ml
Population type	
Age	Adulthood (19 to 24 years) Adulthood (25 to 44 years) Adulthood (45 to 64 years)

Population covered	Sick population
Gender	Male
Geography area	National
Detail of the geography area	France
Data collection	
Dates	
Date of first collection (YYYY or MM/YYYY)	06/2001
Size of the database	
Size of the database (number of individuals)	[1000-10 000[ individuals
Details of the number of individuals	5965
Data	
Database activity	Data collection completed
Type of data collected	Clinical data Declarative data Biological data
Clinical data (detail)	Direct physical measures Medical registration
Declarative data (detail)	Paper self-questionnaire
Biological data (detail)	Type of samples taken: Blood and/or saliva and/or urine
Presence of a biobank	Yes
Contents of biobank	Serum DNA
Details of biobank content	Serum bank, DNA bank
Health parameters studied	Health event/morbidity Health event/mortality
Procedures	
Data collection method	Self-administered questionnaire: Entry from a paper

	questionnaire (Manual input) Clinical examination: Handwritten (Manual input) Biological Analysis: Handwritten (Manual input)
Participant monitoring	Yes
Details on monitoring of participants	(Indefinite duration)
Links to administrative sources	No
Promotion and access	
Promotion	
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Link to the document	http://www.ncbi.nlm.nih.gov/pubmed/?term
	http://www.ncbi.nlm.nih.gov/pubmed/?term
Link to the document	http://www.ncbi.nlm.nih.gov/pubmed/?term  Data may be used by academic teams. Access based on contract collaboration. Data may not be used by industrial teams
Link to the document  Access  Terms of data access (charter for data provision, format of	Data may be used by academic teams. Access based on contract collaboration. Data may not be