

CEPACS - Cohort of children with chromosomal structure abnormality

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General	
Identification	
Detailed name	Cohort of children with chromosomal structure abnormality
Sign or acronym	CEPACS
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	Date de réception de l'avis favorable de la CNIL : 23/05/2007
General Aspects	
Medical area	Immunology
Health determinants	Genetic
Keywords	Health events, cytogenetic reorganization, motor cognitive and physical development, psychometric assessments, education, institutional caring
Scientific investigator(s) (Contact)	
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Unit	INSERM U676 (PHYSIOPATHOLOGIE, CONSEQUENCES FONCTIONNELLES ET NEUROPROTECTION DES ATTEINTES DU CERVEAU EN DEVELOPPEMENT) DEPARTEMENT DE GENETIQUE

Organization	INSERM
Collaborations	
Participation in projects, networks and consortia	Yes
Others	Participation in a cohort network: CEMARA - platform shared by 32 reference centers for rare diseases
Funding	
Funding status	Public
Details	----
Governance of the database	
Sponsor(s) or organisation(s) responsible	ASSISTANCE PUBLIQUE HOPITAUX DE PARIS
Organisation status	Public
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.	Prospective
Database objective	
Main objective	The main objective of this cohort is clinical 1) Acquire information about motor, cognitive and physical development of patients carriers of micro-reorganizations depending on the type of anomaly declared 2) Define, through a longitudinal follow-up, the morbid complications (neurological progress or

degradation, epilepsy, incidence of common pathologies) and mortality rate of these patients 3) Specify psychological, medical and social caring parameters, education and socialization of these patients, medico-economic consequences. Secondary objective : Introduce, at the level of French population, the overall impact, the distribution by type of abnormality and by chromosomal region, the effects of environment variables such as parents age,...

Inclusion criteria

Chromosomal structure micro-reorganizations detected through molecular cytogenetic (Fish or CGH Array)

Population type

Age

Newborns (birth to 28 days)
Childhood (6 to 13 years)
Adolescence (13 to 18 years)

Population covered

Sick population

Gender

Male
Woman

Geography area

National

Detail of the geography area

French multi-center cohort (34 centers)

Data collection

Dates

Date of first collection (YYYY or MM/YYYY)

05/2007

Size of the database

Size of the database (number of individuals)

[1000-10 000[individuals

Details of the number of individuals

1000

Data

Database activity

Data collection completed

Type of data collected

Clinical data
Declarative data
Paraclinical data
Biological data

Clinical data (detail)	Direct physical measures Medical registration
Details of collected clinical data	Clinical examination at inclusion and during the follow-up. Information collected during the clinical examination : indirect collection through parents, most of the time (mentally disabled patients)
Declarative data (detail)	Face to face interview
Details of collected declarative data	Clinical examination at inclusion and during the follow-up. Information collected during the clinical examination : indirect collection through parents, most of the time (mentally disabled patients)
Paraclinical data (detail)	Imaging: different examinations can be collected depending on clinical constraints. No examination is collected without being justified by medical follow-up good practice.
Biological data (detail)	Samples: karyotype and DNA collected with diagnostic purposes, in the context of an etiologic checkup. The results of this examination are a prerequisite for the inclusion in the cohort
Presence of a biobank	Yes
Contents of biobank	DNA
Details of biobank content	DNA bank
Health parameters studied	Health event/morbidity Health event/mortality
Procedures	
Data collection method	Interviews: manually entered paper questionnaire Clinical examinations: hand-written step Biological examinations: hand-written step
Participant monitoring	Yes
Details on monitoring of participants	Undetermined period
Links to administrative sources	Yes
Linked administrative sources (detail)	CépiDC
Promotion and access	

Promotion

Access

Dedicated website

<https://cemara.org>

Terms of data access (charter for data provision, format of data, availability delay)

Possible data utilization by academic teams? Yes.
Contractual access conditions.
Data utilization available for industry sectors? Non

Access to aggregated data

Access on specific project only

Access to individual data

Access on specific project only