

# - Registry of Congenital Malformations in Brittany (qualified registry 2013-2015)

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## General

### Identification

Detailed name	Registry of Congenital Malformations in Brittany (qualified registry 2013-2015)
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	CNIL n°910138 du 11/8 /2010 ; CCTIRS n° 10.145 du 11/3/2010

### General Aspects

Medical area	Disability/handicap
Health determinants	Geography Lifestyle and behavior
Keywords	heart defects, diaphragmatic hernia, hypospadia, Spina bifida

### Scientific investigator(s) (Contact)

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Organization	CHU de

### Collaborations

### Funding

Funding status	Public
Details	- Institut de veille sanitaire - InVS- Agence régionale de santé - ARS- Conseil général
<b>Governance of the database</b>	
Sponsor(s) or organisation(s) responsible	CHU de Rennes
Organisation status	Public
<b>Additional contact</b>	
<b>Main features</b>	
<b>Type of database</b>	
Type of database	Morbidity registers
Additional information regarding sample selection.	Cases referred by practitioners and researcher completeness checks.
<b>Database objective</b>	
Main objective	To carry out a health watch, monitor prevalency according to time and space and identify tetrogenic factors. To assess medical treatment requirements. To study environmental factors and begin a geospatial analyses for certain malformations in connection with surrounding activities.
Inclusion criteria	- Live births or foetal deaths with gestational age at $\geq$ 22 weeks and pregnancy terminations for medical reasons - born on or after 1st September 2010 - with congenital anomaly (EUROCAT criteria) diagnosed during the first year of life - mother living in Brittany.
<b>Population type</b>	
Age	Newborns (birth to 28 days) Infant (28 days to 2 years)
Population covered	Sick population
Gender	Male Woman
Geography area	Regional
French regions covered by the	Bretagne

## database

Detail of the geography area	Brittany: Ille et Vilaine and Côtes d'Armor (106 registered centres)
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## Data collection

### Dates

Date of first collection (YYYY or MM/YYYY)	09/2010
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Date of last collection (YYYY or MM/YYYY)	10/2014
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### Size of the database

Size of the database (number of individuals)	[500-1000[ individuals
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Details of the number of individuals	700 /an /year
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### Data

Database activity	Current data collection
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Type of data collected	Clinical data Declarative data
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Clinical data (detail)	Medical registration
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Declarative data (detail)	Phone interview
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Presence of a biobank	No
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Health parameters studied	Health event/morbidity
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### Procedures

Data collection method	---
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Participant monitoring	Yes
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Links to administrative sources	No
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## Promotion and access

### Promotion

Link to the document	<a href="http://www.cclinouest.com/JOSE/Page_2012/Rouget_registre_phrc_JOSE3.pdf">http://www.cclinouest.com/JOSE/Page_2012/Rouget_registre_phrc_JOSE3.pdf</a>
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Link to the document <http://tinyurl.com/HAL-RMCB>

Description Liste des publications dans HAL

Link to the document <http://tinyurl.com/PUBMED-RMCB>

Description Liste des publications dans Pubmed

## Access

Terms of data access (charter for data provision, format of data, availability delay) Contact the scientist in charge

Access to aggregated data Access on specific project only

Access to individual data Access on specific project only