

PREG-Med - Case-Control Study on the Impact of Pregnancy on Prescribing Psychotropic Medication

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General

Identification

Detailed name Case-Control Study on the Impact of Pregnancy on Prescribing Psychotropic Medication

Sign or acronym PREG-Med

CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation

CNIL

General Aspects

Medical area Geriatrics
Psychology and psychiatry

Health determinants Intoxication
Lifestyle and behavior
Medicine

Keywords psychotropic medication prescription; post-partum; antenatal; pregnancy

Scientific investigator(s) (Contact)

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Organization Erasme

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Organization	Institute of Psychology

Collaborations

Funding

Funding status	Public
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Details	Erasme, Institute of Psychology
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Governance of the database

Sponsor(s) or organisation(s) responsible	Erasme
Organisation status	Public

Additional contact

Main features

Type of database

Type of database	Study databases
Study databases (details)	Case control study

Database recruitment is carried out by an intermediary	An administrative base or a register
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Base or register (detail)	Member of a social security scheme
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Database recruitment is carried out as part of an interventional study	No
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Database objective

Main objective	To determine if and how the prescription of psychotropic medication changes during pregnancy.
Inclusion criteria	Women covered by a social security scheme.
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	General population
Gender	Woman
Geography area	National
Detail of the geography area	France
Data collection	
Dates	
Size of the database	
Size of the database (number of individuals)	Greater than 20 000 individuals
Details of the number of individuals	87,213 (pregnant women); 87,213 controls
Data	
Database activity	Data collection completed
Type of data collected	Administrative data
Administrative data (detail)	Prescription of psychotropic medication
Presence of a biobank	No
Health parameters studied	Health event/morbidity
Procedures	
Participant monitoring	Yes
Details on monitoring of participants	4 months before pregnancy; 9 months during pregnancy and 4 months after pregnancy.

Links to administrative sources	Yes
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Linked administrative sources (detail)	CRAMIF database
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Promotion and access

Promotion

Link to the document	http://www.journal-therapie.org/articles/therapie/abs/first/therapie140012/therapie140012.html
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Access

Terms of data access (charter for data provision, format of data, availability delay)	Contact the scientist in charge (risk that data may be removed after two years due to lack of archiving space in a rolling system).
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Access to aggregated data	Access on specific project only
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Access to individual data	Access on specific project only
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