COroFET - Clinical and laboratory epidemiological monitoring of pregnant women with evaluation of the obstetric, foetal and neonatal risk associated with SARS-CoV-2 during the COVID-19 pandemic- COroFET

Head :DUBUCS Charlotte, IUCT-Oncopole Pathology Department

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General	
Identification	
Detailed name	Clinical and laboratory epidemiological monitoring of pregnant women with evaluation of the obstetric, foetal and neonatal risk associated with SARS-CoV-2 during the COVID-19 pandemic- COroFET
Sign or acronym	COroFET
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	CPP2020-04-039 / 2020-A00870-39 / 20.04.07.83722
General Aspects	
Medical area	Anatomy - Cytology Gynecology/ obstetrics Infectious diseases
Study in connection with Covid- 19	Yes
Scientific investigator(s) (Contact)	
Name of the director	DUBUCS
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Unit	IUCT-Oncopole Pathology Department
Organization	Toulouse University Hospital

Collaborations	
Funding	
Governance of the database	
Sponsor(s) or organisation(s) responsible	Toulouse University Hospital
Organisation status	Public
Additional contact	
Main features	
Type of database	
Type of database	Others
Specify	Clinical and laboratory database (containing clinical and paraclinical data associated with a biological and tissue collection on pregnancy outcomes)
Database recruitment is carried out as part of an interventional study	No
Database objective	
Main objective	In this situation in which data are crucial to understanding the effects of SARS-CoV-2 in the pregnant female population, our primary objective is to collect clinical and paraclinical data from a large sample of women recruited from our level 3 maternity unit, and to create biological and tissue collections with a view to responding to a series of questions, partly explained in the research programme, but which may also evolve as knowledge progresses.
Inclusion criteria	Adult females aged 18 years or over at the date of inclusion Pregnant women giving birth at the Paule de Viguier maternity unit, Toulouse University Hospital, in the study, between April 2020 and April 2021, regardless of pregnancy outcome (live births, intrauterine foetal death, termination of pregnancy, i.e. miscarriages, medical termination of pregnancy) and term Women having given their consent to take part in the study Women registered with a social security scheme (including the state welfare scheme)

Population type	
Age	Adulthood (19 to 24 years) Adulthood (25 to 44 years)
Population covered	General population
Pathology	
Gender	Woman
Geography area	Regional
French regions covered by the database	Languedoc-Roussillon Midi-Pyrénées
Detail of the geography area	Toulouse
Data collection	
Dates	
Size of the database	
Size of the database (number of individuals)	[1000-10 000[individuals
Details of the number of individuals	target 3,600 by end April 2021
Data	
Database activity	Current data collection
Type of data collected	Clinical data Paraclinical data Biological data
Clinical data (detail)	Direct physical measures
Presence of a biobank	Yes
Contents of biobank	Serum Cord blood Fluids (saliva, urine, amniotic fluid, ?) Tissues
Procedures	
Participant monitoring	No

Links to administrative sources No

Promotion and access

Promotion

Access