

DIVAT - Données Informatisées et VALidées en Transplantation

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General

Identification

Detailed name	Données Informatisées et VALidées en Transplantation
Sign or acronym	DIVAT
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	CNIL (17/09/2004) n°891735, Network DIVAT : 10.16.618

General Aspects

Medical area	Urology, andrology and nephrology
Keywords	Transplantation

Scientific investigator(s) (Contact)

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Unit	Institut de Transplantation Urologie Néphrologie, RTRS ?Centaure?, UMR Inserm 1064Centre d'Investigation Clinique et de Biothérapie
Organization	Centre Hospitalier Universitaire Hôtel Dieu

Collaborations

Funding

Funding status	Mixed
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Details	Each center supports its own costs of creation, follow-up and development. We occasionally receive a financial support from private pharmaceutical lab.
Governance of the database	
Sponsor(s) or organisation(s) responsible	Institut de Transplantation Urologie Néphrologie, RTRS ?Centaure?, UMR Inserm 1064
Organisation status	Public
Additional contact	
Main features	
Type of database	
Type of database	Morbidity registers
Additional information regarding sample selection.	All recipients of a kidney and/or a pancreas transplant are included in the cohort.
Database objective	
Main objective	The eight centers established between them collaboration in order to be able to collect and exchange information, to conduct clinical and epidemiological studies all together, to communicate and exchange data and results with other academic researchers, to construct partnership with pharmaceutical industries and to create a biocollection database linked to clinical data (for 3 of the 8 centers).
Inclusion criteria	Kidney and/or pancreas transplant recipients
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	Sick population
Gender	Male Woman
Geography area	National
Detail of the geography area	Nantes, Nancy, Montpellier, Toulouse, Necker

(Paris), Lyon, Saint-Louis (Paris) et Nice

Data collection

Dates

Date of first collection (YYYY or MM/YYYY) 1994

Size of the database

Size of the database (number of individuals) [10 000-20 000[individuals

Details of the number of individuals 11000

Data

Database activity Current data collection

Type of data collected
Clinical data
Paraclinical data
Biological data
Administrative data

Clinical data (detail)
Direct physical measures
Medical registration

Paraclinical data (detail)
Biopsies of the transplant at pre- or post-transplantation. Identification of bacterial, viral, parasitic or mycotic infectious agents.

Biological data (detail)
At pre-transplantation : viral status, immunization and only for pancreas transplantations (C-peptide, HbA1C, serum creatinin, serum proteins, serum albumin, LDL, HDL, cholesterol, triglycerides). At post-transplantation : serum creatinin, proteinuria, hemoglobin, LDL, triglycerides, immunization and only for pancreas transplantations (glycemia, C-peptide, HbA1C, insuline, serum amylase, serum lipase, cholesterol).

Administrative data (detail)
File number, Cristal number (registration on waiting list), patient contact details and name of referring doctor.

Presence of a biobank Yes

Contents of biobank
Whole blood
Serum
Plasma
Blood cells isolated

Fluids (saliva, urine, amniotic fluid, ?)

Details of biobank content

DMSO CELLS, SERUM, WHOLE BLOOD, TRIZOL CELLS, URINE, PAXGENE, PLASMA

Health parameters studied

Health event/morbidity
Health event/mortality

Procedures

Data collection method

Data are collected at pre-transplantation and prospectively at each follow-up visit.

Participant monitoring

Yes

Details on monitoring of participants

Follow-up parameters are collected at 3 months, 6 months, 1 year and then every year.

Links to administrative sources

No

Promotion and access

Promotion

Link to the document

<http://tinyurl.com/PUBMED-DIVAT>

Description

Liste des publications dans Pubmed

Link to the document

<http://tinyurl.com/HAL-DIVAT>

Description

Liste des publications dans HAL

Access

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

The Divat network encourage collaborations and enables interested researchers to submit a research project. Guidelines for requesting data analyses from the registry will be soon available on the DIVAT website.

Access to aggregated data

Access on specific project only

Access to individual data

Access on specific project only