- Hereditary Ichthyosis Cohort

Head :Mazereeuw-Hautier Juliette, Unité « Différenciation Epidermique et Autoimmunité Rhumatoïde » (UDEAR) UMR 5165 CNRS ? 1056 INSERM - Université Toulouse III, Equipe « Kératinocyte granuleux et barrière épidermique » (N Jonca) Jonca Nathalie

Last update : 08/08/2014 Version : 1 ID : 5215		
General		
Identification		
Detailed name	Hereditary Ichthyosis Cohort	
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	en cours	
General Aspects		
Health determinants	Genetic	
Others (details)	Ichthyosis	
Keywords	Hereditary, progression, complications	
Scientific investigator(s) (Contact)		
Name of the director	Mazereeuw-Hautier	
Surname	Juliette	
Address	Service de Dermatologie, Centre de Référence des Maladies Rares de la Peau 24, chemin de Pouvourville TSA 30300, Toulouse Cedex 09	
Phone	+ 33 (0)5 67 77 81 41/21	
Email	Mazereeuw-hautier.j@chu-toulouse.fr	
Unit	Unité « Différenciation Epidermique et Autoimmunité Rhumatoïde » (UDEAR)	

Toulouse III,

épidermique » (N Jonca)

UMR 5165 CNRS ? 1056 INSERM - Université

Equipe « Kératinocyte granuleux et barrière

Organization	CHU
Name of the director	Jonca
Surname	Nathalie
Collaborations	
Funding	
Funding status	Mixed
Details	Financement ministériel du centre de référence maladies rares pour du personnel.
Governance of the database	
Sponsor(s) or organisation(s) responsible	CHU Toulouse
Organisation status	Public
Additional contact	
Main features	
Type of database	
Type of database	Study databases
Study databases (details)	Longitudinal study (except cohorts)
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.	All patients with this disease are included
Database objective	
Main objective	To form a usable database for hereditary ichthyosis on which to base future research projects, such as epidemiological, clinical (disease progression follow-up and onset of complications) physiopathological or therapeutic projects.
Inclusion criteria	All hereditary ichthyosis patients, regardless of clinical type, severity, age, gender, disease

progression or treatment (to ensure a b	road	
representative cohort). Patients attendin	g	
consultation or hospitalised during their	first visit or	
medical follow-up in the Referral Centre of		
centres throughout France. Signed consent to		
participate in the study.		

_				
DO	nııl.	stia	n t	ype
FU	DUIG	コロロ	11 6	vue
-				, , , ,

Age Newborns (birth to 28 days)

Infant (28 days to 2 years)
Early childhood (2 to 5 years)
Childhood (6 to 13 years)
Adolescence (13 to 18 years)
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 France

Data collection

Dates

Date of first collection (YYYY or

MM/YYYY)

09/2009

Size of the database

Size of the database (number of

individuals)

< 500 individuals

Details of the number of

individuals

100

Data

Database activity Current data collection

Type of data collected Clinical data

Declarative data
Paraclinical data
Biological data
Administrative data

Clinical data (detail)	Direct physical measures Medical registration
Declarative data (detail)	Face to face interview
Paraclinical data (detail)	Data collected from regular patient follow-up
Biological data (detail)	Regular biological monitoring
Administrative data (detail)	Socio-demographic data
Presence of a biobank	Yes
Contents of biobank	Whole blood Tissues DNA
Details of biobank content	Blood and skin samples
Health parameters studied	Health event/morbidity Health event/mortality Health care consumption and services Quality of life/health perception
Care consumption (detail)	Hospitalization Medical/paramedical consultation Medicines consumption
Procedures	
Data collection method	By physician during annual visit. Data are recorded in real-time in an Excel database for clinical and biological characteristics.
Participant monitoring	Yes
Details on monitoring of participants	Re-assessment of patient's clinical and biological characteristics and quality of life during regular follow-up visit (normally an annual visit). Follow-up over 10 years.
Links to administrative sources	No
Promotion and access	

Promotion

Access

Terms of data access (charter for data provision, format of

Publications: Data available between CRMRP and INSERM teams. The creation of a database allows

data, availability delay)	ichthyosis studies to be carried out jointly by these units. The transfer of data to other parties for research purposes is not planned unless part of a collaborative project.
Access to aggregated data	Access on specific project only
Access to individual data	Access on specific project only