

FRESH QOLA - Cross-cultural Validation of Systemic Lupus Activity (SLAQ) and Lupus Quality of Life (Lupus QOL) Questionnaires in French

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General	
Identification	
Detailed name	Cross-cultural Validation of Systemic Lupus Activity (SLAQ) and Lupus Quality of Life (Lupus QOL) Questionnaires in French
Sign or acronym	FRESH QOLA
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	CNIL : 28/01/2009
General Aspects	
Medical area	Immunology
Health determinants	Occupation Social and psychosocial factors
Keywords	Advancement, translation, French, quality of life, questionnaire
Scientific investigator(s) (Contact)	
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Collaborations	
Funding	
Funding status	Private
Details	LUPUS FRANCE
Governance of the database	
Sponsor(s) or organisation(s) responsible	CHU Dijon
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.	Prospective

Database objective

Main objective	General objective: To translate and validate the Systemic Lupus Activity questionnaire and the Lupus Quality of Life into French
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Inclusion criteria	- Patients fulfilling ACR 1997 criteria for systemic lupus erythematosus classification; - patients that understand written and spoken French; - patients between 18 and 75 years of age; - patients who received oral and written information regarding research.
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Population type

Age	Adulthood (19 to 24 years) Adulthood (25 to 44 years) Adulthood (45 to 64 years) Elderly (65 to 79 years)
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Population covered	Sick population
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Gender	Male Woman
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Geography area	Regional
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French regions covered by the database	Alsace Champagne-Ardenne Lorraine Bourgogne Franche-Comté
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Detail of the geography area	Multicentric cohort throughout the Eastern Region of France (6 centres)
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Data collection

Dates

Date of first collection (YYYY or MM/YYYY)	02/2009
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Date of last collection (YYYY or MM/YYYY)	01/2010
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Size of the database

Size of the database (number of	< 500 individuals
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individuals)

Details of the number of individuals	13
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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)	Direct physical measures Medical registration
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Declarative data (detail)	Paper self-questionnaire Face to face interview
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Presence of a biobank	No
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Health parameters studied	Quality of life/health perception
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Procedures

Data collection method	Self-administered questionnaire: from a paper questionnaire (manual input) with double data entry Interview: from a paper questionnaire (manual input) with double data entry Clinical Examinations: handwritten (manual input) with double data entry
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Participant monitoring	Yes
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Details on monitoring of participants	Follow-up duration: 6 months
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Links to administrative sources	No
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Promotion and access

Promotion

Link to the document	http://tinyurl.com/Hal-FRESH-QOLA
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Description	List of publications in HAL
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Link to the document	http://tinyurl.com/Pubmed-FRESH-QOLA
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Description	List of publications in Pubmed
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Access

Terms of data access (charter	To be decided if data may be used by academic
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for data provision, format of data, availability delay)

teams Data may not be used by industrial teams

Access to aggregated data

Access on specific project only

Access to individual data

Access on specific project only