

# Stratified medicine in primary Sjogren's syndrome

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		<input type="checkbox"/> Protocol
<b>Registration date</b> 08/08/2023	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
<b>Last Edited</b> 05/09/2023	<b>Condition category</b> Musculoskeletal Diseases	<input type="checkbox"/> Individual participant data
		<input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Primary Sjögren's syndrome (PSS) is a chronic, complex immune-mediated disease with no effective treatment available. The effects vary greatly among PSS sufferers with some reporting disabling fatigue, some unbearable pain and dryness, while others report fewer symptoms. Some PSS patients develop lymphoma or major extra-glandular organ damage while some have disease confined to the exocrine glands. Although many biological abnormalities have been described in PSS, their relationships with specific clinical manifestations remain unclear. This creates challenges in finding appropriate ways of measuring the effectiveness of therapies and the burden of disease. Using detailed clinical data from 600 PSS patients from the UK, researchers have identified four clinical PSS subtypes which differ in characteristics. The data suggest that outcome and response to therapies may differ depending on the patient's subtype. The aim of this study is to develop a PSS-specific patient-reported questionnaire that captures the disease burden/impact in each of the PSS subtypes.

### Who can participate?

Patients aged 18 years and over with PSS and clinicians across Sweden, Norway and France

### What does the study involve?

Completing questionnaires about quality of life and disease impact at baseline and follow-up.

### What are the possible benefits and risks of participating?

There are no direct benefits and no risks of taking part.

### Where is the study run from?

Newcastle upon Tyne Hospitals NHS Foundation Trust (UK)

### When is the study starting and how long is it expected to run?

January 2019 to October 2024

### Who is funding the study?

European Commission

Who is the main contact?  
Professor Wan-Fai Ng, wan-fai.ng@newcastle.ac.uk

**Study website**  
Nil known

## Contact information

**Type(s)**  
Principal Investigator

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## Additional identifiers

**EudraCT/CTIS number**  
Nil known

**IRAS number**  
264215

**ClinicalTrials.gov number**  
Nil known

## Secondary identifying numbers

IRAS 264215, CPMS 48622

# Study information

## Scientific Title

Stratified medicine in primary Sjogren's syndrome: a mixed methods quantitative/qualitative study to develop a patient-reported questionnaire looking at disease burden within the Primary Sjogren's Syndrome subtypes

## Acronym

FOREUM

## Study objectives

This study aims to further characterise the clinical significance and the underpinning pathotypes of the 4 primary Sjogren's syndrome (PSS) subtypes. The researchers will develop a patient questionnaire that enables them to quantitatively measure the impact and disease burden of PSS within the different subtypes.

## Ethics approval required

Ethics approval required

## Ethics approval(s)

Approved 09/04/2021, East of England - Cambridge East Research Ethics Committee (The Old Chapel, Royal Standard Place, Nottingham, NG1 6FS, United Kingdom; +44 (0)2071048096; cambridgeeast.rec@hra.nhs.uk), ref: 21/EE/0069

## Study design

Observational multicentre mixed methods study

## Primary study design

Observational

## Secondary study design

Longitudinal study

## Study setting(s)

Home, Hospital

## Study type(s)

Other

## Participant information sheet

Not available in web format, please use contact details to request a participant information sheet.

## Health condition(s) or problem(s) studied

Primary Sjogren's Syndrome

## Interventions

The aim of the study is to collect information on the disease impact and burden of PSS on patients. This information will be collected using questionnaires due to the large cohort (600 participants) across numerous countries. Patient public involvement is crucial to the design of the study. Patient representatives from each country are involved in the steering committee with the clinicians and will work together to develop the questionnaire.

The main categories of the questionnaire were decided at an initial face-to-face meeting with clinicians, academics, statisticians and patient representatives. These categories form the phases of the questionnaire development. The four phases will be discussed in more detail to establish questions suitable to collect the data for the categories. These discussions will take place by teleconference between the patient representatives, chief investigator, health economist and project manager.

Opinions on the draft questionnaire will first be sought from PSS patient support groups in Norway, Sweden, France and the UK. The completed questionnaire will then be submitted to REC for approval and piloted first in the UK. This will then be enrolled out to the other EU sites (Norway, Sweden and France) to complete the subtype data collection (600 participants, 150 in each subgroup).

Baseline data will be collected and a follow-up questionnaire administered. Questionnaires will be posted or given in clinic to complete at home. The timing of the follow-up will depend on the questions of interest identified for the categories during the development process. Questionnaire development and recruitment will take place over a period of two years to allow one year for data analysis.

Participants will be identified from established PSS registries across the centres. The participants of these registries have already consented to receive future invitations for PSS research. A letter of invitation, patient information sheet and written consent form will be posted to the patient or given during clinic by the clinical research team. If the participant consents then the questionnaire will be posted to them.

## **Intervention Type**

Other

## **Primary outcome measure**

The following data is collected through a one off questionnaire by post.

1. Quality of life measured using EQ5D
2. Wellbeing measured using ICECAPA
3. Anxiety and depression measured using HADs
4. Sjogren's symptoms measured using ESSPRI
5. Fatigue is measured using PROFAD
6. Patient developed measure to include to include; Clinical Health Services (primary and secondary), extra services not covered by the health service – such as cost of prescriptions, dental, opticians, and alternative therapies, employment, family life and activities – such as holidays.

## **Secondary outcome measures**

There are no secondary outcome measures

## **Overall study start date**

10/01/2019

**Completion date**

31/10/2024

## Eligibility

**Key inclusion criteria**

1. Age  $\geq 18$  years
2. Confirmed diagnosis of primary Sjögren's syndrome
3. Willing and able to provide informed written consent
4. Registered within a PSS registry

**Participant type(s)**

Patient

**Age group**

Adult

**Lower age limit**

18 Years

**Sex**

Both

**Target number of participants**

600

**Key exclusion criteria**

1. Unable to provide written consent

**Date of first enrolment**

01/03/2021

**Date of final enrolment**

31/10/2023

## Locations

**Countries of recruitment**

England

France

Norway

Sweden

United Kingdom

**Study participating centre**

The Newcastle upon Tyne Hospitals NHS Foundation Trust

Freeman Hospital

Freeman Road

High Heaton

Newcastle upon Tyne

United Kingdom

NE7 7DN

## Sponsor information

**Organisation**

Newcastle upon Tyne Hospitals NHS Foundation Trust

**Sponsor details**

Joint Research Office

Regent Point

Regent Farm Road

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NE3 3HD

+44 (0)1912824926

nuth.genericqueries@nhs.net

**Sponsor type**

Hospital/treatment centre

**Website**

<http://www.newcastle-hospitals.org.uk/>

**ROR**

<https://ror.org/05p40t847>

## Funder(s)

**Funder type**

Other

**Funder Name**

European Commission

**Alternative Name(s)**

European Union, Comisión Europea, Europäische Kommission, EU-Kommissionen, Euroopa Komisjoni, Ευρωπαϊκή Επιτροπή, Европейская комиссия, Evropské komise, Commission européenne, Choimisiúin Eorpaigh, Europskoj komisiji, Commissione europea, La Commissione europea, Eiropas Komisiju, Europos Komisijos, Európai Bizottságrol, Europese Commissie, Komisja Europejska, Comissão Europeia, Comisia Europeană, Európskej komisii, Evropski komisiji, Euroopan komission, Europeiska kommissionen, EC, EU

**Funding Body Type**

Government organisation

**Funding Body Subtype**

National government

**Location**

## Results and Publications

**Publication and dissemination plan**

Planned publication in peer-reviewed scientific journals.

**Intention to publish date**

31/10/2024

**Individual participant data (IPD) sharing plan**

The datasets generated and/or analysed during the current study will be available upon request from Victoria Macrae ([victoria.macrae@newcastle.ac.uk](mailto:victoria.macrae@newcastle.ac.uk))

**IPD sharing plan summary**

Available on request