# Systematic genetic testing for personalised ovarian cancer therapy (SIGNPOsT)

Submission date	Recruitment status No longer recruiting	<ul><li>Prospectively registered</li></ul>		
07/06/2017		Protocol		
Registration date	Overall study status Ongoing  Condition category Cancer	Statistical analysis plan		
29/06/2017		Results		
Last Edited		Individual participant data		
27/09/2023		[] Record updated in last year		

#### Plain English summary of protocol

https://www.cancerresearchuk.org/about-cancer/find-a-clinical-trial/a-study-looking-at-genetic-testing-for-ovarian-cancer-signpost

# Contact information

#### Type(s)

Scientific

#### Contact name

Dr Ranjit Manchanda

#### **ORCID ID**

http://orcid.org/0000-0003-3381-5057

#### Contact details

Queen Mary University of London Room 4 Basement Old Anatomy Building Charterhouse square London United Kingdom EC1M 6BQ +44 20 7882 5555 r.manchanda@qmul.ac.uk

# Additional identifiers

EudraCT/CTIS number

IRAS number

#### ClinicalTrials.gov number

#### Secondary identifying numbers

ReDA11776

# Study information

#### Scientific Title

Systematic GeNetic testing for Personalised Ovarian cancer Therapy: A prospective cohort study investigating the impact of systematic germline panel and concomitant somatic testing in epithelial ovarian cancer on psychological health and quality of life

#### Acronym

**SIGNPOsT** 

#### Study objectives

Null hypothesis:

There is no difference in psychological health between mutation carriers detected on panel-based genetic testing and concomitant somatic testing (of high grade non-mucinous epithelial ovarian cancer) compared to non-carriers.

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

National Research Ethics Committee - London Riverside, 29/03/2017, ref: 17/LO/0405

#### Study design

Prospective cohort study

# Primary study design

Observational

# Secondary study design

Cohort study

# Study setting(s)

Hospital

# Study type(s)

Other

# Participant information sheet

Not available in web format, please use the contact details below to request a patient information sheet

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

Epithelial ovarian cancer

#### **Interventions**

Participants attend a baseline visit where they undergo systematic genetic germline panel testing for BRCA1, BRCA2, RADL51C, RAD51D, BRIP1 gene mutations and concomitant somatic genetic testing for BRCA1 and BRCA2. They also have demographic data, family history and clinical outcomes collected. Participants are asked to fill out surveys about their anxiety, quality of life, and counselling satisfaction scale. Participants who decline the genetic test complete a survey called "reasons for declining genetic test". This visit takes around 55-60 minutes.

After the results of the genetic testing is complete, participants then repeat the anxiety and quality of life surveys as well as have a psychological health test using the Multi-dimensional impact of Cancer Risks Assessment (MICRA) scale. This visit takes around 30 minutes.

Six months after the test, participants again repeat the anxiety and quality of life surveys as well as have a psychological health test using the Multi-dimensional impact of Cancer Risks Assessment (MICRA) scale. This visits takes around 30 minutes.

Participants then are asked to attend annual visits where they are surveyed for their patient satisfaction and if they regret their decisions. Participants again repeat the anxiety and quality of life tests. Only at the first annual visit since the genetic testings, participants again complete the MICRA test to have their psychological health assessed.

The total follow up period is five years.

#### **Intervention Type**

Behavioural

#### Primary outcome measure

Psychological health following test result is assessed by MICRA Multi-dimensional impact of Cancer Risk Assessment (MICRA) scale at 7-days post-result, 6-months and 12 months.

#### Secondary outcome measures

- 1. Patient reported quality of life is assessed by EORTC QLQ C-30, OV28, EN-24, and EQ5D-5L questionnaires at baseline, post-result, 6-months, 12 months and annually
- 2. Anxiety and depression are assessed using the Hospital anxiety, depression scale (HADS) at baseline, post-result, 6-months, 12 months and annually
- 3. Cost effectiveness is reported using the Incremental cost effectiveness ratio per quality adjusted life years (ICER/QALY) at the end of the study
- 4. Germline and somatic BRCA1, BRCA2 detection rate is measured as the number of carriers /number of cases at the end of the study
- 5. Uptake is measured as the proportion of eligible EOC patients who accepted the offer of genetic testing at baseline
- 6. Patient satisfaction is assessed by the adapted genetic counselling satisfaction scale—measured post recruitment. A decision regret scale is used at 12 months.
- 7. PARP Inhibitor uptake is assessed by proportion of participants commenced on PARP inhibitors and reported at the end of the study

Overall study start date 05/05/2017

Completion date 04/05/2026

# **Eligibility**

#### Key inclusion criteria

- 1. Aged over 18 years
- 2. Histological diagnosis of high-grade non-mucinous epithelial ovarian cancer/fallopian tube cancer/primary peritoneal cancer

#### Participant type(s)

**Patient** 

#### Age group

Adult

#### Lower age limit

18 Years

#### Sex

Female

#### Target number of participants

280

#### Key exclusion criteria

- 1. Women with non-epithelial ovarian cancer
- 2. Women with low grade or mucinous epithelial ovarian cancer
- 3. Unable to consent

#### Date of first enrolment

08/05/2017

#### Date of final enrolment

04/05/2021

# Locations

#### Countries of recruitment

England

United Kingdom

#### Study participating centre St Bartholomew's Hospital

West Smithfield London United Kingdom EC1a 7BE

# Study participating centre Royal London Hospital

Whitechapel Road Whitechapel London United Kingdom E1 1BB

# Study participating centre Queens Hospital

Romvalley Way Romford United Kingdom RM70AG

# Sponsor information

#### Organisation

Queen Mary University of London

#### Sponsor details

Director of Research Services & Business Development Joint Research Management Office (JRMO) QM Innovation Building, 5 Walden Street London England United Kingdom E1 2EF

# Sponsor type

University/education

#### **ROR**

https://ror.org/026zzn846

# Funder(s)

# Funder type

Charity

#### **Funder Name**

# **Results and Publications**

#### Publication and dissemination plan

Results of the study will be presented at scientific meetings and conferences and published in scientific journals. They will also be distributed through supporting charities, patient groups and the Barts Cancer Institute, QMUL web site.

#### Intention to publish date

04/05/2026

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

The datasets generated during and/or analysed during the current study are/will be available upon request from r.manchanda@qmul.ac.uk after publication of all results

#### IPD sharing plan summary

Available on request

# **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient- facing?
HRA research summary			28/06 /2023	No	No
Other publications	Cohort study set within the recruitment to SIGNPOST	26/09 /2023	27/09 /2023	Yes	No