

Checklist for enrolling a participant in the BSRBR-RA

Please include this form with the documents for each participant you wish to register with the BSRBR-RA

I have given my patient the Participant Information Sheet and answered any questions

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The participant has consented to be involved in the study

Please note: only participants who agree to all sections of the consent form can be enrolled in the study. Each box on the consent form should be initialled to indicate consent.

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I have enclosed the following completed documents:

1. Signed Patient Consent Form (version 9)

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2. Completed* Clinical Baseline Form (Clinical Baseline Form V11.2 for patients new to the study or Short Baseline Form V3.1 for re-registering patients to a new cohort)

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3. HAQ

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4. EQ5D

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*Please ensure that the baseline form is completed with as much information as possible. If there are important data missing such as biologic therapy, disease severity measures or existing co-morbidities, it may be necessary to exclude these important patients in future BSRBR-RA analyses, thus losing all their information collected to date. Where essential data is missing, the registration documents will be returned to you for completion.

Please send the registration documentation to the BSRBR-RA study offices:

BSRBR-RA
Unit 4 Rutherford House
40 Pencroft Way
Manchester Science Park
Manchester
M15 6SZ

Fax: 0161 2751640



The participant will be registered and an ID number assigned. You will be sent follow up forms to complete on a six monthly basis for 3 years and then annually thereafter.

If you have any questions about eligibility, recruitment, follow up or anything else please do not hesitate to contact the project assistant team who will be happy to help

Tel: +44 (0)161 275 1652/7390

Email: biologics.register@manchester.ac.uk

Please ensure that a copy of the original MREC approval and subsequent amendment approval letters have been sent to your Trust R & D department.