

Evaluation of HIS-UK Study
Self-Referral Case Report Form (SR-CRF)

Site ID -

Study Participant ID

1. Pre-consultation data transfer checklist

- Chlamydia screening:** Single Triple
- Trial group:** Usual care (control) proHIS eHIS
- Digital recording:** Required Not required

2. Consultation bookings

Date	Method (online / in-clinic)	Consultation Outcome (completed / partial completion / not completed)

3. Consent

Date: ____/____/____

Name of person re-confirming consent: _____

- Study documents were thoroughly reviewed by the subject** Yes No
- Subject understands the nature of the study and what is involved** Yes No
- Subject is able to give informed consent** Yes No

Comments:

4. Patient registration

Patient Record ID

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6. Samples collected

Urine:

- Yes Subject refused Not required Postal-kit requested Postal-kit provided

Anal swab:

- Yes Subject refused Not required Postal-kit requested Postal-kit provided

Oral swab:

- Yes Subject refused Not required Postal-kit requested Postal-kit provided

6. Intervention delivery

Trial group:

- Usual care (control) proHIS eHIS

Digital recording:

- Yes No Not required

7. Protocol Deviation

Comments /reasons:

Site PI Signature: _____ Date: _____

This form should be completed for each participant screened

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Study Participant ID

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Chlamydia screening

Date of screening / kit sent	Method (Postal/in-clinic)	Test Type (Single/Triple)	Date of results transfer to study team
<i>14/05/19</i>	<i>Postal</i>	<i>Single</i>	<i>28/05/19</i>
<i>20/05/19</i>	<i>In-clinic</i>	<i>Triple</i>	<i>31/05/19</i>

Comments:

Site PI Signature: _____ **Date:** _____