

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

Site ID

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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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**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:** \_\_\_\_\_