

Haematuria Biomarker Study (HaBio)

Submission date 16/11/2012	Recruitment status No longer recruiting	<input type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
Registration date 03/01/2013	Overall study status Completed	<input type="checkbox"/> Statistical analysis plan <input checked="" type="checkbox"/> Results
Last Edited 31/10/2022	Condition category Signs and Symptoms	<input type="checkbox"/> Individual participant data

Plain English summary of protocol

<http://www.cancerresearchuk.org/cancer-help/trials/a-study-help-develop-test-find-causes-blood-in-urine-habio>

Contact information

Type(s)

Scientific

Contact name

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Additional identifiers

Study information

Scientific Title

A study investigating the use of protein measurements as diagnostic tests for the causes of blood in the urine

Acronym

HaBio

Study objectives

Collectives of biomarkers aligned with demographics and/or clinical variables can predict bladder cancer in haematuria patients with 90% accuracy.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Office of Research Ethics Committee Northern Ireland, 23/12/2012, ref: 11/NI/0164

Study design

Case control study

Primary study design

Observational

Study type(s)

Screening

Health condition(s) or problem(s) studied

Haematuria

Interventions

At the clinic the Research Nurse will record height, weight and blood pressure, medical history and ask questions about lifestyle and occupations before collecting one urine and one blood sample. The first 20 patients in this part of the study will be asked to provide approximately 25ml urine sample (approximately 5 teaspoons), and a 45ml blood sample (9 teaspoons). All patients after this will be asked to provide approximately 25ml sample of urine (5 teaspoons) and approximately 35ml sample of blood (7 teaspoons). If diagnosis is confirmed using bladder tissue removed during the surgical procedure the researchers will review the tissue and use small samples to identify protein and other constituents including DNA and RNA within the tissue structures. The patients notes will be reviewed by members of the HaBio clinical team. The HaBio clinical team will inform the patients consultant of their review findings should this be appropriate. This will be beneficial for patients. Members of the HaBio clinical team will review the patients notes for a second time 3 years after recruitment to obtain updated information about the patients health.

Intervention Type

Other

Phase

Not Applicable

Primary outcome(s)

Single biomarkers and/or multivariate algorithms will significantly improve on the predictive power of an algorithm based on demographics for prediction of causes of haematuria in patients presenting with haematuria.

Key secondary outcome(s)

1. Patient age, gender, current medications, date of cystoscopy and date of initial diagnosis of bladder cancer (if relevant) are recorded from the patient notes by a Clinical Research Nurse

(CRN) or the HaBio clinician at the time of recruitment

2. Blood pressure and body mass index (BMI) measurements, together with dip-stick analyses are undertaken by the CRN or the HaBio clinician at the time of recruitment
3. A detailed history of smoking, cancers, hypertension, renal stones, pelvic irradiation, and urinary tract infections; weekly alcohol consumption; present and past occupations; lower urinary symptoms; and whether the patient presented with visible or non-visible haematuria are recorded by the CRN or the HaBio clinician following discussion with each patient at their time of recruitment
4. Urine and blood samples are obtained from each patient by the CRN or the HaBio clinician at the time of recruitment
5. Scientists complete analyses of ~ 60 biomarker levels for each patient at Randox Laboratories Ltd (Country Antrim, Northern Ireland) as soon as possible after recruitment
6. Investigation results, details of positive micro-organism findings, causes of infection, causes of benign disease and levels of clinical biomarkers are recorded following review of the patient notes by a consultant urologist at least six weeks after recruitment
7. One or more causes of haematuria are recorded for each patient following review of the patient notes by a consultant urologist at least six weeks after recruitment
8. Where applicable, cause of death, details of chronic kidney disease, non-urological cancers, biomarkers measured in NHS, bladder cancer treatments, prostate and kidney pathology are recorded following review of the patient notes by the HaBio Clinician at the time of follow up
9. Where applicable, pathological review to obtain dates of recurrences and progression and pathology reports for tissue removed at the time of recruitment are recorded for all patients following review of the patient notes by the HaBio Clinician at the time of follow up
10. A detailed pathological review including details of pathological variants is completed by a consultant pathologist following completion of follow up
11. Patterns of expression and cell type location of key biomarkers using appropriate immunohistochemistry (IHC) on sections from diagnostic bladder tissue samples from patients is completed for each patient after completion of the diagnostic review by a Consultant Pathologist

Completion date

21/02/2020

Eligibility

Key inclusion criteria

Bladder cancer patients

Patients with pathologically proven bladder cancer, newly diagnosed or recurrent, will be recruited prior to transurethral resection of the bladder tumour at pre-assessment clinics, as in-patients on urology wards or at planned cystoscopy sessions.

1. Written informed consent to participate in the study
2. Aged between 40 and 80 years
3. Current haematuria or a history of haematuria
4. Cystoscopy within the last 6 months or planned cystoscopy
5. No chemo- or radio- therapy in the three weeks prior to recruitment
6. No previous history of cancers other than bladder cancer
7. Suspicion of bladder cancer or proven bladder cancer

Control patients

These patients will be recruited from haematuria clinics following negative cystoscopy and negative findings for other bladder cancer investigations.

1. Written informed consent to participate in the study
2. No previous history of cancer
3. Of the same gender, approximate age and smoking status (where possible) to a bladder cancer patient already recruited to HaBio
4. Current haematuria or a history of haematuria
5. Negative cystoscopy within the last 3 months, but at least 48h after the procedure
6. No chemo- or radio- therapy in the three weeks prior to recruitment

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Sex

All

Total final enrolment

677

Key exclusion criteria

Bladder cancer patients:

1. No written informed consent to participate in the study
2. Aged <40 or >85 years
3. No history of haematuria
4. No recent or planned cystoscopy
5. Chemo- or radio- therapy in the three weeks prior to recruitment
6. Previous history of cancer(s), other than bladder cancer
7. No suspicion of bladder cancer or proven bladder cancer

Control patients:

1. No written informed consent to participate in the study
2. Previous history of any cancer
3. Not of the same gender, approximate age and smoking status of a patients already recruited as a bladder cancer patient
4. No history of haematuria
5. No recent or planned cystoscopy
6. Chemo- or radio- therapy in the three weeks prior to recruitment

Date of first enrolment

27/11/2012

Date of final enrolment

28/06/2017

Locations

Countries of recruitment

United Kingdom

England

Northern Ireland

Study participating centre

Belfast City Hospital

Department of Urology

Lisburn Road

Belfast

United Kingdom

BT9 7AB

Study participating centre

Ulster Hospital Dundonald

Upper Newtownards Road

Dundonald

United Kingdom

B16 1RH

Study participating centre

Craigavon Hospital

68 Lurgan Road

Portadown

United Kingdom

B63 5QQ

Sponsor information

Organisation

Queens University Belfast (UK)

ROR

<https://ror.org/00hswnk62>

Funder(s)

Funder type

Industry

Funder Name

Randox Laboratories Ltd (UK)

Results and Publications

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study may be available for research collaborations (k.williamson@qub.ac.uk)

IPD sharing plan summary

Available on request

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article		01/09/2022	31/10/2022	Yes	No