



# DIRECT

DIABETES RESEARCH ON PATIENT STRATIFICATION

## CONSENT FORM (version 3.0)

**NB. This form must be completed and signed by the research participant in the presence of someone with knowledge of the research designated by the Principal Investigator. This may be a doctor, nurse, clinical research assistant or other member of the research team who must countersign the form as witness to the participant's signature**

Patient Identification Number for this study: \_\_\_\_\_

**Please initial box**

I confirm that I have read and understood the information sheet dated 16<sup>th</sup> May 2012 (version 3.0) for the above study. I have had the opportunity to consider the information, ask questions and have had these answered to my satisfaction

I confirm that I am aware that I will need to attend the hospital or clinical research facility up to 4 occasions.

I confirm that I am aware that I will receive an MRI scan (body scan)

I understand that my participation is voluntary and that I am free to withdraw my consent at any time without giving any reason, without any medical care or legal rights being affected.

**I understand that:**

- My GP will be informed that I am taking part in this research study
- Information about my diabetes can be accessed from my health records by the research team during the study period.
- Apart from the routine clinical blood tests which will be made available to my health care team, my blood and urine will be anonymised and stored for the duration of the DIRECT project.
- Information regarding any abnormal measurements (e.g. Blood pressure) and important findings on the MRI scan will be communicated to my GP and/or hospital physician.

DIRECT: T2DMProg

- The information I provide and data about me will be stored for the duration of the DIRECT project on secure electronic databases, which can be accessed for this study by researchers who are members of the DIRECT consortium, which includes pharmaceutical companies. This information will *not* include my name, NHS number or address or date of birth.
- My blood and urine samples and anonymised data may be used in future projects by other researchers collaborating with the DIRECT consortium, including commercial companies, into diabetes and related conditions including genetic studies on DNA as described in the Information Sheet.

I understand that the Newcastle NHS Hospitals Foundation Trust, as sponsor to this research, may require access to my medical records for study monitoring purposes only thereby ensuring that the research is properly conducted and the interests of those taking part are adequately protected.

I understand that I am allowing transfer and storage of a specimen of my samples including the hereditary information DNA and RNA for analysis in medical research on the assumption that it is free of any legal claim on my part and without expectation of personal financial gain.

**I agree to take part in the above study.**

\_\_\_\_\_  
Participant Name (PRINT)

\_\_\_\_\_  
Date

Signature\_\_\_\_\_

**THANK YOU for agreeing to take part in this research**

I confirm that the nature of the research, the required additional procedures and the voluntary nature of the study have been explained in terms understandable to the patient.

\_\_\_\_\_  
Name of person taking consent (PRINT)

\_\_\_\_\_  
Date

Signature\_\_\_\_\_