Access to health data for research and innovation – frequently asked questions

January 2020

Introduction

Health Data Research UK is the national institute for health data science in the UK. Its mission is to unite the UK’s health data to enable discoveries that improve people’s lives. By working in partnership with the NHS, industry, academia and patients, and providing safe and secure access to rich health data, it aims to:

- Enable better care
- Improve public health
- Understand the causes of disease
- Support better, faster, more efficient clinical trials

Health Data Research UK is leading a £37.5 million government-funded programme to build a world-leading, robust, trusted health data research infrastructure for the UK to enable discoveries that improve people’s lives. This work is funded by UK Research and Innovation’s Industrial Strategy Challenge Fund (ISCF).

What this means for me

1. I live in England/Scotland/Wales/Northern Ireland. Is my health data available to researchers?

Most people in the UK who use the NHS – whether for an ongoing condition, maternity services, an operation, or an emergency – will have some data recorded about their care which is already available to researchers on an anonymised basis.

This could include, for instance: how long it took an ambulance to get you to hospital, how long you waited for admission to a bed or for your operation or mental health intervention, what sort of treatment you had, how long you stayed in hospital (if you did), whether you were re-admitted soon afterwards and what ongoing treatment you have for health conditions such as diabetes (if any).

Controlled access to data held in patient records improves an individual’s care and helps the health and care system to plan local services. Information can also be provided to researchers to enable them to find out more about diseases, their causes and risks, and develop new treatments.
For example, some GP practices across the UK take part in a 30-year long, nationwide project called the Clinical Research Practice Datalink to support innovative research. This uses de-identified information about patients from those practices.

You may also have agreed to be part of a clinical trial, in which case you will already know what you agreed to, and what its purpose is.

The NHS and other organisations which use data to enable health research – such as Clinical Research Practice Datalink, UK Biobank, and Cystic Fibrosis Trust – already make their de-identified data available to researchers.

2. **Why is health data needed for research and innovation?**

If data from many patients are linked up and pooled, researchers and doctors can look for patterns in the data, helping them develop new ways of predicting or diagnosing illness, and identify ways to improve clinical care. The information from health data is really valuable to help understand more about disease, to develop new treatments, to monitor safety, to plan services and to evaluate NHS policy.

Before a researcher is granted access, their study is assessed by an independent review committee or other decision-making group, who check that the reason for using the data is appropriate. Wherever possible, identifying information will be removed from that data, and researchers should only be given the minimum amount necessary to answer a question. Data must be stored securely, and a legal contract must be signed before data can be transferred or accessed.

Often a study will need to use data about an individual that is held in more than one dataset. When this happens, a trusted third party, usually NHS Digital, links the data using a unique identifier (such as NHS number which is then removed) to make sure the researcher cannot re-identify individuals.

Health Data Research UK is leading work to improve access to data by researchers to improve prevention, care and treatments, helping people lead longer, healthier lives.

3. **What is Health Data Research UK doing that is different?**

Health Data Research UK is working with:

- the organisations which hold health data
- healthcare professionals, academics, charities and industry which want to carry out research to improve prevention, care and treatments
- patients and the public to shape how access to data should work.

Its aim is to enable medical discoveries that improve people’s lives by making it possible for researchers to:

- develop new ways of managing and extracting knowledge and insights from complex and diverse health data
use data from big datasets to address important health research questions that cannot be tackled by individual research organisations.

It will do this by:

- bringing together information about existing health datasets so that researchers know what’s available and how they can access it
- improving the quality of UK health data, and developing its use, by working with the organisations that hold UK health data (known as ‘data custodians’) and helping them work towards common standards for allowing access to the data
- enabling researchers to use the health data safely and securely
- ensuring we have the skills and ways of working to work with data in the most efficient and responsible way to solve health challenges
- developing tools – like Artificial Intelligence – that can help draw new insights

4. **What’s in it for me?**

If researchers have access to larger, more diverse datasets, the more likely it is that they will find something that can help you, your family, your community or someone else with your condition. If research identifies something which could improve care for individual patients taking part in research, wherever possible this will be fed back to the team treating them, so they can benefit directly from it.

The fuller the data that the NHS has about its services and patients, the better it can plan for the future. Evidence shows that people are generally comfortable with anonymised data from medical records being used for improving health, care and services, for example for research, provided there is a public benefit.1

5. **What’s in it for us – the people of the UK?**

The NHS has some of the richest data records in the world, but much of it is very hard to access and it’s not joined up. Health data is held in thousands of different places across the UK.

Part of the aim of Health Data Research UK is to help provide links across these datasets, make it more straightforward for researchers to access and help increase the speed and scale at which research and innovation can take place. The end result will be that people benefit sooner from improved treatments, devices, interventions and health system management.

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How my data is protected

6. Who is in charge of keeping my data safe?

The organisation which currently has your data (known as the ‘data controller’ or ‘controller’) – your GP practice, community health services, mental health or hospital trust, medical research charity, UK Biobank or disease registry – is responsible for keeping your data safe.

Every member of staff who works for these organisations has a legal obligation to keep information about you confidential. In the NHS, organisations maintain a duty of confidentiality by conducting annual training and awareness, ensuring access to personal data is limited to the appropriate staff and information is only shared with organisations and individuals that have a legitimate and legal basis for access.

Where systems have been set up to collect data from GP practices, NHS trusts, Health Boards and so on – for instance by NHS Digital in England, the Information Services Division of NHS Scotland, NHS Wales, and Health and Social Care Northern Ireland – the organisations collecting the data have responsibility for it. It is collected and, in most cases, de-identified before being released for research.

These organisations and others are part of the UK Health Data Research Alliance which brings together the organisations that hold the UK’s health data. They are working together and with the public to develop standardised tools, techniques, conventions and technologies for research and innovation.

These are the principles guiding their work, called ‘Principles for Participation’, which all organisations working with Health Data Research UK have to sign up to.

7. How do I find out what data is held about me?

Under General Data Protection Regulations you have the right to ask an organisation whether or not they are using or storing your personal information. You can also ask them for copies of your personal information, verbally or in writing. This is called the right of access and is commonly known as making a subject access request or SAR.

This guide from the Information Commissioner explains how to make a subject access request.

8. How will data be accessed?

Decisions about how health data is used will always put patient privacy, safety and benefit first and we have multiple safeguards in place to ensure this happens.

Health Data Research UK is enabling researchers to discover and request access to data via the Health Data Research Innovation Gateway. This holds information or descriptions of the different types of datasets in the UK – which will include those held by the NHS, charities, and disease registries, for example – to enable researchers to see what’s available and how they can access it. The Gateway goes not hold or store any patient or health data.
If a researcher wants to access a dataset, they can send a request via the Innovation Gateway and this will be considered by the organisation that looks after that dataset.

Research on the data is carried out in what’s known as a Trusted Research Environment or Safe Haven. These are highly secure places – either physical servers often in a locked room or on a Safe Cloud – that can only be used by researchers who have been permitted entry. Any technology companies involved in providing or supporting the Safe Havens will not be able to see or access the data.

The aim is to enable maximum security, through multiple layers, and to minimise the risk of anyone’s data being misused.

9. What happens if there is a security breach?

The organisations responsible for managing the data (the ‘data custodians’) already have tried and tested data protection plans in place to respond to threats to data security, including significant data breaches or near misses. These plans are regularly reviewed by the NHS and other organisations involved as part of their commitment to cybersecurity, and their legal duty under the Data Protection Act and General Data Protection Regulations to protect people’s data.

If ever there is a security breach, or a risk of a security breach, these plans will come into place and will likely involve reporting the incident to the Information Commissioners Office.

Public involvement in the programme

10. How has the public been involved in developing these plans?

Public engagement and involvement are at the heart of the work of Health Data Research UK, which has these values:

- we are transparent
- we are optimistic
- we are respectful
- we act with courage
- we act with humility.

We want people to understand what we are doing and why so that:

- patients and the public are our partners in research and innovation
- the needs, values and interests of patients and the public are understood and embedded in our work
- people have trust and confidence in the use of health data for research and innovation
- people are excited about their data being used in research and innovation.

Health Data Research UK’s Public Advisory Board, which was recruited from patients and members of the public from across the country, has shaped the work of the programme so far. Patient/public events were
held to gain input and feedback from people about how they would want their data to be accessed and
used for research. Each of the Health Data Research Hubs is expected to involve patients and the public in
making decisions on how their data is used, and they will be reporting on this on a quarterly basis.

11. Why haven’t I heard about these plans?

We are planning a wide ranging campaign in 2020 with our partners, patients and the public to open up the
discussion.

12. How can I get involved?

If you would like to get involved in Health Data Research UK’s work, please email enquiries@hdruk.ac.uk

Concerns about data

13. Could my health data be linked up with other sources of data to identify me?

The General Data Protection Regulation (GDPR) specifically makes it a criminal offence to deliberately
attempt to, or to re-identify someone (conditions apply).

There are multiple safeguards in place – both by Health Data Research UK and the organisations that
manage the data – to reduce the risk of data being reidentified. The Innovation Gateway will not hold or
store any patient or health data. If access is provided by the data custodian, research on the data will be
carried out in what’s known as a Trusted Research Environment or Safe Haven. These are highly secure
places – either physical servers often in a locked room or on a Safe Cloud – that can only be used by
researchers who have been permitted entry. Any technology companies involved in providing or
supporting the Safe Havens will not be able to see or access the data.

This minimises the risk that any data can be downloaded or transferred and then used to identify any
individual patient.

For a helpful description of what is meant by identifiable data, please read Understanding Patient Data’s
factsheet.

14. Is this going to affect my employability/life insurance/pension?

The safeguards we are putting in place to avoid identification of any individual patient and minimise the risk
of any data being misused make it extremely unlikely that there would be any adverse consequences for
people.
15. How can I be sure?

You can make a request to your healthcare provider (your GP or hospital) and ask who has had access to your identifiable NHS data.

16. What if I don’t want my health data to be used?

If you live in England, you can opt out of sharing the data held in personal records about you and you can do this online via the NHS Digital website https://digital.nhs.uk/services/national-data-opt-out

Although Scotland doesn’t have an opt out mechanism, you can contact your GP or hospital to discuss your wishes or visit https://spire.scot/my-choices/ to opt out of sharing GP records only. For further information about use of other data in Scotland you can read the NSS data protection notice and the ISD/HPS Data Protection Notice

Likewise, if you live in Wales you can opt out of your summary GP record being shared by visiting http://www.gprecord.wales.nhs.uk/home

For information about the lawful bases for processing information please visit this guidance on the Information Commissioner’s website.

Industry partners

17. Who are your industry partners?

Health Data Research UK has many industry partners to boost capabilities in a variety of ways – from data analytics tools to research skills and resources. Many UK-based small and medium-size enterprises are involved, as well as large companies with an international presence. Forty per cent of the companies involved in the Health Data Research Hubs are UK-founded, and all must have a UK base, thereby helping to ensuring that this programme enhances the UK’s life sciences sector.

18. What is in it for the big tech companies?

Health Data Research UK is working with some big technology companies, such as Google, Microsoft and IBM to develop services and tools for health data research in the UK. Like the other companies we work with, they benefit from working in partnership with UK organisations by gaining insights into the NHS and how to develop innovative products for patients.
19. Why do we need them? Why can't the NHS and universities do the work without them?

Industry is critical in rapidly translating pioneering research into mainstream care. It is the biomedical and pharmaceutical industries that have taken pioneering research and turned it into many of the medical innovations – such as the X-ray machine, MRI scanner and penicillin – that we take for granted today.

Without the involvement of industry, the NHS would not be able to get the new medicines, treatments and diagnostics for patients. Medicines and diagnostics are always developed outside the NHS and government by the private sector.

The involvement of industry partners in Health Data Research UK’s work does not mean that they will automatically be able to access data. They are providing a specific service, and – like all the partners and companies involved – won’t have any privileged access to information. The same principles and safeguards will apply to anyone inside or outside the partnership. Any decisions about how the data will be used will always put patient privacy, safety and benefit first and we have multiple safeguards in place to ensure this happens.

20. Are you selling data?

The data will not be available for free and there will be some form of charge by the organisation that manages the data (the 'Data Controller') for access or services or both. This is the case currently with most data controllers in the UK.

Health Data Research UK is working with the data controllers and the Hubs to look at an approach to making its work sustainable – so that it does not have to rely on the UK taxpayer for its funding over the long-term – and to achieve this there will be a commercial model for access to data. Anyone requesting access to data must demonstrate the benefits that their research will bring to patients and the NHS. Health Data Research UK is also working closely with the Government to supports its policy development to realise the benefits and value of UK health data.

21. Are private companies going to profit from using my data?

Any researchers working for private companies wishing to access the data will need to demonstrate how their research will benefit patients and the NHS. While companies are likely to profit (for example, by developing new drugs), they will also be delivering vital benefits to patients.

Contact details

Email: enquiries@hdruk.ac.uk

https://twitter.com/hdr_uk

https://www.linkedin.com/company/healthdataresearchuk/