General Data Protection Regulation (GDPR) and research

This note seeks to help prepare researchers to ensure regulatory compliance following the introduction of the General Data Protection Regulation (GDPR / “the Regulation”); and to meet best practice in research ethics and governance.

This note is not intended as the final authority on data protection regulations. Researchers should always consult with the Regulation, Information Commissioner’s Office guidance, guidance from the regulatory authorities such as the Health Research Authority, and colleagues in Legal and Compliance for definitive guidance on the Regulation. This note should be read in conjunction with the resources set out in Appendix 1.

Background

The EU General Data Protection Regulation replaced the Data Protection Directive 95/46/EC and is designed to harmonise data privacy laws across Europe, to protect and empower all EU citizens’ data privacy, and to reshape the way organisations across the region approach data privacy.¹

The introduction of the Regulation requires the University to review the robustness of our personal data processing practices. ‘Personal data’ includes pseudonymised data, and is defined as:

“… any information relating to an identified or identifiable natural person (‘data subject’); an identifiable natural person is one who can be identified, directly or indirectly, in particular by reference to an identifier such as a name, an identification number, location data, an online identifier or to one or more factors specific to the physical, physiological, genetic, mental, economic, cultural or social identity of that natural person.”²

¹ https://www.eugdpr.org/
² General Data Protection Regulation, Article 4 (a definition of pseudonymised is provided in Article 4)
Urgent action needed to prepare for the Regulation

The General Data Protection Regulation, which entered into force on 25th May 2018, introduces and reinforces statutory requirements for the processing of personal data. While most of the data protection requirements are unchanged from previous legislation, there are additional compliance requirements that will require urgent action following the introduction of the Regulation.

These include the requirements to:

- update your participant consent forms and participant information sheets
- safeguard any transfer of data outside the EU, and ensure you have informed consent from participants for any transfer of data outside the EU
- review your data processing arrangements

Further information on each of these requirements is set out below.
**General actions related to the processing of personal data**

The General Data Protection Regulation offers the opportunity to ensure you are regularly reviewing the steps you are taking to protect personal data. Your consent process and your mechanisms for data storage and data use should be subject to ongoing review to ensure that good practice standards are met and that participant’s rights and wishes are respected.

**Providing information on the lawful basis for processing personal data to research participants**

The General Data Protection Regulation requires each activity of processing data to have a lawful basis.

For studies falling under the Department of Health framework, the Health Research Authority have produced [guidance](#) on the requirements for providing information on the lawful basis to research participants, and this guidance must be followed for studies requiring review by the Health Research Authority.

The University processes personal data as part of its research and teaching activities in accordance with the lawful basis of ‘public task’, and in accordance with the University’s [Supplemental Charter](#) which states that the purpose of the University “shall be to advance education, learning and research for the public benefit”.

Further information on fulfilling the requirements for using public task as your lawful basis for processing personal data can be found in [Appendix 2](#).

In order to inform research participants about the lawful basis on which you are processing their personal data, you will need to update your participant consent forms and participant information sheets to ensure that they meet the new University (or HRA) template forms.

When recruiting new participants, you should ensure that you use the new template consent and participant information sheets. For ongoing studies where participants are still in the study (and there is active data collection), you should update your participant information sheet in line with the new template, and provide a copy of the new sheet to participants.
It should be noted that if you have an ongoing study, there is no need to re-consent participants, unless you are making changes to your study processes or arrangements as a result of the Regulation which were not part of the informed consent process (for example: changing what data you collect or how you will hold or use it). However, consent originally obtained should be reviewed, especially where tiered-consent approaches have been adopted (for example, where the consent process allows participants to opt-out of specific elements of the study) to ensure that the data are being managed in a way which is consistent with the terms of the consent.

For University approved studies, the use of new information sheet and consent forms for ongoing studies, and any changes to the research proposal, would need to be submitted to the research ethics committee which originally reviewed the study and any other committees which approved the research.

For studies falling under the Health Research Authority guidance, the HRA have advised that where their GDPR guidance has been followed, the R&D office of participating NHS/HSC organisations does not need to be notified as these will be classified as non-notifiable, non-substantial amendments in the vast majority of cases. However, if you are unsure, please contact the HRA for further guidance.
Participant information sheets and consent forms

The introduction of the General Data Protection Regulation provides the opportunity to review your participant information sheets and participant consent forms and ensure that you are using the latest templates provided – either by the University for studies approved by a University research ethics committee; or by the Health Regulatory Authority for studies approved by a NHS research ethics committee.

All information provided to participants must be concise, transparent, in easily accessible form and made using clear and plain language to meet the needs of the audience. Please see the information below on informed consent for further guidance.

For new studies or ongoing studies where participants are still in the study (which have received approval from a University research ethics committee) you must use the University’s template participant information sheets and consent forms, available at:

https://www.liverpool.ac.uk/intranet/research-support-office/research-ethics/ethics-application-submission/.

The only exception to the requirement to use the University templates would be where these documents need to be tailored to meet the needs of the research population; for example, child-friendly forms which may include pictures and diagrams.

In order to use any new participant information sheets and participant consent forms, you will need to submit an amendment to the relevant research ethics committee to cover the use of the new sheets.

When submitting this amendment in the University online system for research ethics applications, you should indicate within section 2.4 that “this amendment will not significantly affect any aspects of the study”.

International transfer of data

The General Data Protection Regulation applies whenever personal data is transferred in or out of the EU, and the Regulation imposes specific restrictions on the transfer of personal data outside the European Union to third countries or international organisations. Personal data may only be transferred outside of the EU in compliance with the conditions for transfer set out in Chapter V of the General Data Protection Regulation.³

Transferring personal data outside the EU is only permitted where the transfer is:

- made with the individual’s informed consent
- necessary for important reasons of public interest⁴

When transferring personal data outside the EU, you must ensure that you have informed consent from research participants to cover the transfer. You must also ensure that the arrangements for protecting the confidentiality of the data meet the highest levels of confidentiality and security.

The Computer Services Department can provide advice on the mechanisms that can be used to protect personal data when transferring data outside of the EU.

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³ General Data Protection Regulation, Chapter V
⁴ Information Commissioner’s Office: International transfers
General good practice in data collection and management

The General Data Protection Regulation offers an opportunity to review and refresh existing practices to ensure that they meet recommended best practice standards and regulatory requirements in the collection and management of personal data collected during research. Please visit the University’s research data management webpages for additional guidance.

Informed consent

The definition of consent outlined when using consent as the lawful basis for processing personal data has been refined in the Regulation as: “any freely given, specific, informed and unambiguous indication of the data subject’s wishes by which he or she, by a statement or by a clear affirmative action, signifies agreement to the processing of personal data relating to him or her”.5 Whilst this represents best practice in obtaining consent, is recognised that not all research studies can be designed to meet these requirements, which is why the University uses ‘public task’ as it’s lawful basis for processing personal data. However, it should be emphasised that this does not affect the ethical importance of consent or the common law requirements for consent.

Given the additional ethical imperative to obtain consent whenever possible, researchers should take this as an opportunity to review and improve their consent procedures in line with recommended best practice. Consent means offering individuals real choice and control. Genuine consent should put individuals in charge, build trust and engagement, and enable participants to decide whether or not to take part in the study.

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5 General Data Protection Regulation, Article 4
The following extract of the guidance outlines some of the good practice considerations for obtaining informed consent:

- Consent should be a positive opt-in

- Explicit consent requires a very clear and specific statement of consent in words, rather than by any other positive action

- Keep your consent requests separate from other terms and conditions
  - Avoid making consent to processing a precondition of any service you are offering

- Keep evidence of consent – who, when, how, and what you told people.
  - Participant consent forms should be stored securely and confidentially

- The participant information sheet and consent forms should:
  - Outline the lawful basis, ‘public task’, on which the University processes personal data (and the condition for processing if sensitive data is collected – see page 12)
  - Be specific and granular where possible to get separate consent for separate things
  - Explain why you want the data (purpose), and you will do with it (intended use), and how long the data will be stored
  - Explain who, if anyone, the data will be shared with; and in what format the data will be shared
  - Highlight what you are doing to ensure the security of personal information
  - Be clear, concise, user friendly
  - Make it easy for people to withdraw consent to participate in research and tell them how they can withdraw their participation (explaining any limitations to withdrawing or deleting their data)
  - Explain that the participant has the right to complain to the University and the Information Commissioner’s Office if they are unhappy with the data management
  - Contain the contact details of the Principal Investigator and the University of Liverpool Data Protection Officer

- Keep consent under review, and refresh it if anything changes.
You must keep clear records to demonstrate consent; and these must be stored securely and confidentially.

It should be noted that it may not always be possible to achieve the gold standard criteria for consent as outlined above. Explicit and granular consent is not always compatible with recommended best practices in certain types of research. For example, consent obtained for research using human material samples often lacks ‘explicit consent’ as to do so could lead to the unnecessary destruction of unique resource. In such cases, a broader consent is obtained to allow the future use and sharing of personal data under certain conditions which have been reviewed and approved by a research ethics committee.
Research data management

Under the General Data Protection Regulation, there is a greater emphasis on implementing safeguards for personal data. This means that you should give consideration to the arrangements for the security and storage of data; ensure that data are pseudonymised or anonymised wherever possible; and that personal data are only collected when needed (known as ‘data minimisation’). If you can undertake some or all of your research activities without using identifiable personal data, you should make arrangements to do so.6

Primary responsibility for the management of data produced during research activities lies with the Principal Investigator (or Supervisor). Where research is conducted with other institutions and independent researchers, Liverpool researchers are responsible for the management of research data held by UoL that is under their own control.7

The following extract outlines some of the good practice considerations for research data management:

- Wherever possible, data should be anonymised or pseudonymised. Personal data can only be disclosed when explicit and documented permission to disclose is part of the consent procedure.8

- Store data on a secure and regularly backed up site - this should be on server systems operated by the University’s Computing Services Department (University network drive)9
  - Storage of data on locations other than the University networked drives should be approved by Information Security colleagues in the Computing Services Department
  - Further information on the correct processes for storing your research data can be found on the Research Data Management webpages

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6 Health Research Authority: Guidance for Researchers
7 University Research Data Management Policy
8 University Policy on Research Ethics
9 University Information Security Policy
- apply technical controls to limit access to the data
  - University network drives and Microsoft SharePoint contain features which enable users to limit access to the data

- use encryption to digitally secure the data
  - Further information on encryption can be found on the Computing Services webpages

- ensure that hard copies of any data are held in a physically secure location
  - For student projects, hard copies of any personal data should be kept in a locked filing cabinet in the Supervisor’s office

- provide secure deletion and destruction facilities
  - Colleagues in records management can advise on retention and disposal of research data

- Sharing personal data should only be done with the consent of research participants
  - A research ethics committee (or in the case of NHS research, the Confidential Advisory Group) should review any proposal to share data without participant consent

The confidentiality of the information supplied by research participants and the anonymity of respondents must be respected.
Sensitive personal data

The Regulation refers to sensitive personal data as “special categories of personal data” in Article 9 of the regulation. Sensitive personal data includes information revealing an individual’s:

- racial or ethnic origin;
- political opinions;
- religious or philosophical beliefs;
- trade union membership;

... or involves the processing of:

- genetic data;
- biometric data for the purpose of uniquely identifying a natural person;
- data concerning health
- data concerning a natural person’s sex life
- data concerning a natural person’s sexual orientation.10

Special category data is personal data which the Regulation says is more sensitive, and so needs more protection as this type of data could create more significant risks to a person’s fundamental rights and freedoms.

A data protection impact assessment is required for processing that is likely to result in a high risk to individuals – for example, where any special category data is processed. The data security measures should be as rigorous as possible when processing sensitive personal data.

If you are processing special category data, you will need to outline both the lawful basis for processing (‘public task’) and the separate condition for processing this data. The condition on which the University processes special category data is that the “processing is necessary for archiving purposes in the public interest”.

10 General Data Protection Regulation, Article 9
Please refer to the Information Commissioner’s Office guidance on special category data for further information on the conditions.

When processing special category or criminal offence data, it must be recognised that the risk to the rights and freedoms of persons are heightened from processing this data; as processing may give rise to discrimination, financial loss, damage to the reputation, loss of confidentiality of personal data, and any other significant economic or social disadvantage. Therefore the likelihood and severity of the risk to the rights and freedoms of the data subject should be carefully considered alongside the nature, scope, context and purposes of the processing to determine whether processing is necessary and whether the safeguards mitigate the risk.

Criminal offence data

Article 10 applies to personal data relating to criminal convictions and offences, or related security measures. Criminal offence data includes the type of data about criminal allegations, proceedings or convictions that would have been sensitive personal data under the 1998 Act; including personal data linked to related security measures.

Processing of personal data relating to criminal convictions and offences can be carried out only under the control of official authority or when the processing is authorised by Union or Member State law providing for appropriate safeguards for the rights and freedoms of data subjects.11

If you are processing criminal offence data, you will need both a lawful basis for processing (‘public task’) and a separate condition for processing this data under Article 10. Please refer to the Information Commissioner’s Office guidance on criminal offence data for further information.

11 General Data Protection Regulation, Article 10
Human Material, consent and GDPR

The consent provisions for the collection and storage of human material are unchanged by the implementation of the General Data Protection Regulations (GDPR). Consent remains a requirement of Common Law and the common law duty of confidentiality is not affected by the implementation of GDPR. The Human Tissue Authority (HTA) have therefore not provided any changes to the current advice or guidance on this matter (HTA COP A: Guiding Principles and the Fundamental Principle of Consent).

The University’s guidance on best practice for consent involving human material can be found by referring to -The University of Liverpool Human Material Code of Practice-HTA003 and The University of Liverpool supporting document-Consenting for research SDS001.

As outlined in earlier sections, the Implementation of GDPR does change the requirements for organisations to hold and process personal data and special categories of personal data.

Consent to participation in research is not the same as consent as the legal basis for processing under data protection legislation. Consent is obtained for participation in research, but the lawful basis which the data collected will be processed under is defined in the study transparency statement.

Guidance produced by the Medical Research Council (MRC) and the Health Research Authority (HRA) state that for public authorities such as Universities, NHS organisations, Research Council institutes or other public authority the lawful basis under which they hold and use personal data is most likely to be GDPR Article 6(1) (e) a ‘task in the public interest’.

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12 https://www.health-ni.gov.uk/articles/common-law-duty-confidentiality
14 https://www.liverpool.ac.uk/intranet/media/intranet/humanmaterialgovernance/HumanMaterialCodeofPractice.pdf
**GDPR Article 6(1) (e)** “Processing is necessary for the performance of a task carried out in the public interest or in the exercise of official authority vested the controller;”

You should note that if it would be possible to undertake your research without processing personal data then your intended legal basis will not be valid.

The MRC\(^5\) have also provided guidance on which of the separate conditions from Article 9 would most likely be used by public authorities to hold and use special categories of personal data

**GDPR Article 9(2) (j)**

“Processing is necessary for archiving purposes in the public interest, scientific or historical research purposes or statistical purposes in accordance with Article 89(1) based on Union or Member State law which shall be proportionate to the aim pursued, respect the essence of the right to data protection and provide for suitable and specific measures to safeguard the fundamental rights and the interests of the data subject”\(^{18}\).

**Further processing of data**

When personal data has been collected from a data subject but the controller (either the University or the Sponsor) intends to further process the data for a different purpose, the controller must also give the data subject information about that further purpose before the data is processed. An example is that researchers may wish to use personal data originally collected for clinical or local audit for research.

However, if the information about further processing is in fact the same as the information for the original processing, the data controller does not need to give the data subject that information again\(^{19}\).


Reporting requirements

The Regulation introduces a duty on all organisations to report personal data breaches to the relevant supervisory authority. You must do this within 72 hours of becoming aware of the breach, where feasible.

As soon as you become aware of a personal data breach, you must report this to the Director of Legal and Compliance, Mr Kevan Ryan (kevan.ryan@liverpool.ac.uk) and the University Data Protection Officer, Mrs Victoria Heath (V.Heath@liverpool.ac.uk) who will then advise on the next steps for handling the breach.

Additional notes

Although the theme of the General Data Protection Regulation is around empowering individuals’ data rights and reshaping the way organisations approach personal data processing, there are a number of areas where the Regulation does not provide specific and conclusive authority with regard to research.

For example, there are no specific provisions to cover the collection of data obtained from behavioural observation studies, the use of data which is available in the public domain, etc. In such areas where there is no specific legislative provision, the spirit of the Regulation, existing common law, and best practice guidance should be considered when reviewing the processing of the personal data. Relevant considerations should be reflected upon in your research ethics applications.
Appendix 1: Resources

- Consumer Data Research Centre: The General Data Protection Regulation & Social Science Research

- General Data Protection Regulation: The full text

- Information Commissioner’s Office: Guide to the General Data Protection Regulation

- Information Commissioner’s Office: 12 steps to take now

- Information Commissioner’s Office: Lawful basis interactive guidance tool

- Health Research Authority: Guidance for Researchers

- University of Liverpool data protection webpages

- University of Liverpool Research Ethics Policy

- University of Liverpool Research Data Management Policy

- University of Liverpool Information Security Policy

- University of Liverpool research data management webpages

- University of Liverpool: Getting ready for GDPR training (obligatory training module)
Appendix 2: Determining your lawful basis for processing personal data

The General Data Protection Regulation requires each activity of processing data to have a lawful basis. There are around six lawful basis for processing personal data. The Information Commissioner’s Office have produced a ‘Lawful basis guidance tool’ to help determine the most appropriate lawful basis for your processing.

For studies falling under the Department of Health framework, please see the Health Research Authority guidance on the lawful basis for processing.

The University processes personal data as part of its research and teaching activities in accordance with the lawful basis of ‘public task’, and in accordance with the University’s Supplemental Charter which states that the purpose of the University “shall be to advance education, learning and research for the public benefit”. Further information on fulfilling the requirements for using public task as your lawful basis for processing personal data can be found below.

Public interest

The Health Research Authority note “For health and social care research, the legal basis is determined by the type of organisation: for universities, NHS organisations or Research Council institutes the processing of personal data for research will be a ‘task in the public interest’”.20

When relying on ‘public interest’ as the lawful basis for processing, the following points need to be considered:

- Are you processing the data to carry out your official tasks or functions, or other specific tasks in the public interest?
  - The University considers the collection of personal data for the purposes of advancing education, learning and research to be a public task
- Can you point to a clear basis in law for your task or function?

20 Health Research Authority: GDPR Operational Guidance
The University considers the advancement of education, learning and research to be the basis in law for the collection of personal data in research.

- Is there another reasonable way to perform your tasks or functions without processing the data?
  - The processing must be necessary. If you could reasonably perform your tasks or exercise your powers in a less intrusive way, this lawful basis does not apply.

You need to be sure that you can demonstrate why processing is necessary to perform a task in the public interest and that there is no other reasonable way to perform the task without processing personal data. Remember to include information about your purposes and lawful basis in your participant information sheets.

Refer to the ICO’s guidance for further information on the use of ‘public interest’ as the lawful basis for processing.

It is important to note that although ‘public interest’ – and not ‘consent’ is likely to be the lawful basis under which personal data is held and processed, the ethical and common law requirements of consent are not reduced.