OPINION ARTICLE

Pathology and regulation for research in the UK: an overview
[version 1; peer review: 2 approved]

Owen John Driskell 1-3, Jessica Lee 4, Karin Oien 5, Andy Hall 6, Clare Verrill 7, CM-Path QA Panel

1Department of Clinical Biochemistry, University Hospitals of North Midlands, Stoke-on-Trent, Staffordshire, UK
2Institute for Applied Clinical Sciences, University of Keele, Stoke-on-Trent, Staffordshire, UK
3National Institute for Health Research Clinical Research Network West Midlands, NIHR CRN, Albrighton, Shropshire, UK
4Strategy and Initiatives, National Cancer Research Institute, London, UK
5Institute of Cancer Sciences, University of Glasgow, Glasgow, UK
6Newcastle University, Newcastle, UK
7Nuffield Department of Surgical Sciences, University of Oxford, Oxford, UK

Abstract

The input of pathologists is essential for the conduct of many forms of research, including clinical trials. As the custodians of patient samples, pathology departments have a duty to ensure compliance with the relevant regulations, standards and guidelines to ensure the ethical and effective use for their intended investigational analysis. This includes where patients are participating in a research study. The results of research studies have impacts beyond the research study itself as they may inform changes in policy and practice or support the licensing of medicines and devices. Compliance with regulations and standards provides public assurance that the rights, safety and wellbeing of research participants are protected, that the data have been collected and processed to ensure their integrity and that the research will achieve its purpose. The requirements of the regulatory environment should not be seen as a barrier to research and should not significantly impact on the work of the laboratory once established and integrated into practice. This paper highlights important regulations, policy, standards and available guidance documents that apply to research involving NHS pathology departments and academic laboratories that are contributing to research involving human subjects.

Keywords

Research regulation, Research governance, MHRA, GCP, HTA, Quality Management, Clinical Trial, CTIMP

This article is included in the Royal College of Pathologists gateway.
Corresponding author: Owen John Driskell (owen.driskell@nihr.ac.uk)

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Competing interests: No competing interests were disclosed.

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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Introduction
Evidence from research is needed to justify changes in practice and is a driving force behind improvements in patient care. Pathology can be involved throughout the research pathway - from developing research ideas, designing, setting up and running research studies, to analysing samples, interpreting the data and publishing the results (Figure 1).

Research may be initiated by pathologists, but a large amount of research is led by researchers from outside pathology. This research often requires services delivered by pathology departments. For some pathology staff this service delivery, for example the release of material from hospital archives, may be their only experience of research. Data generated from pathology departments supports research in a wide variety of ways. This could be informing study eligibility assessments, companion diagnostics, pharmacokinetics, surrogate or other endpoints or the provision of data, results or tissue for further assessment. Results may already be in the health record as part of routine care or they may require further analysis or the release of tissue to central laboratories for a standardised assessment.

As well as the increase in the overall workload of pathology departments, there have been increases in clinical trials requiring pathology input. As research needs to be carried out in accordance with the appropriate regulatory requirements and standards, pathology departments need to have risk-based and pragmatic procedures in place to manage these requests. Regulations change and so it is important that pathology departments maintain an understanding of the current regulatory environment around research.

UK Policy Framework for Health and Social Care Research
In the NHS research should be carried out in accordance with the UK Policy Framework for Health and Social Care Research, which replaced the Department of Health Research Governance Framework in October 2017. The Framework sets out the principles of best practice for the management and conduct of all health and social care research in the UK, to help organisations meet their legal requirements. It covers research in the NHS and other health and social care environments across all four nations. The Framework includes fifteen principles that apply to all research and a further four that apply only to interventional research. It provides a clear definition and outline of the roles and responsibilities for individuals and organisations involved in health and social care research including the sponsor, the funder, the research team, research sites, clinical research organisations and regulators. It states there should be clear designation of responsibility and accountability with clear lines of communication between all those involved in research.

The Framework is not a regulation - it is statutory guidance published in order to help organisations meet their legal obligations. It is a common set of principles that apply to any study. Though they aren’t identical, these will be recognisable to those already familiar with the principles of Good Clinical Practice. The framework is an important document for all those involved in NHS research as it provides a common language for research and enhances the possibilities for a unified and streamlined approach to research delivery.

Good Clinical Practice, the UK Clinical Trials Regulations and the MHRA
The International Council for Harmonisation (ICH) of Technical Requirements for Pharmaceuticals for Human Use Good Clinical Practice (GCP) is an international standard for the design, conduct, performance, monitoring, auditing, recording, analysis, and reporting of clinical trials. These internationally recognised standards followed on from the Nuremberg Code and originate from the World Medical Association.
Declaration of Helsinki 1964 and were an attempt to overcome international inconsistencies in GCP.

The Guideline for Good Clinical Practice E6 from the European Medicines Agency includes the 13 Principles of ICH GCP. These guidelines refer to ‘clinical trials’ but may also be applied to other clinical investigations that may have an impact on the safety and well-being of human participants.

ICH GCP was incorporated into EU Directives 2001/20/EC and 2005/28/EC aimed at harmonising the regulation of Clinical Trials in the EU. These were transposed into UK Law via The Medicines for Human Use (Clinical Trials) Regulations in 2004 and have since been further amended. The UK has adopted the principles of GCP, but an additional principle relating to sponsor indemnity was added. This did not define ICH GCP as the legal standard unlike other European countries. As with the UK Policy Framework for Health and Social Care Research these regulations also include definitions and roles and responsibilities for research.

The Clinical Trial regulations define a clinical trial of an investigational medicinal product (CTIMP) as any investigation in human subjects, other than a non-interventional trial, intended (a) to discover or verify the clinical, pharmacological or other pharmacodynamic effects of one or more medicinal products, (b) to identify any adverse reactions to one or more such products, or (c) to study absorption, distribution, metabolism and excretion of one or more such products, with the object of ascertaining the safety or efficacy of those products. The Medicines and Healthcare products Regulatory Agency (MHRA) provide a tool for whether research is a CTIMP and can provide further assistance if required.

The regulations have had subsequent amendments in response to developments in healthcare, for example research interventions in pandemic disease and advanced therapies such as gene therapy and tissue engineering. Europe will be implementing a new Clinical Trial Regulation (536/2014) during 2020 and this may be implemented in the UK dependent on Brexit negotiations and future agreements. If the new Regulation does not come into force during the implementation period, the Government has confirmed that UK law will remain aligned with parts of the EU’s CTR legislation that are within the UK’s control, in order that researchers conducting clinical trials can plan with greater certainty.

The Medicines for Human Use Regulations permit inspection of clinical trials and their conduct, including work undertaken in a laboratory, by the MHRA GCP Inspectorate. It is therefore prudent for a laboratory to know which trials they support are CTIMPs and so are subject to these regulations and potential inspection.

The primary reference for describing the requirements of GCP in the context of laboratories is the EMA Reflection Paper for Laboratories that Perform the Analysis or Evaluation of Clinical Trial Samples published in 2012. This was developed based on the MHRA Guidance on the Maintenance of Regulatory Compliance in Laboratories.

The primary focus of inspection by the MHRA GCP Inspectorate is the analysis of samples that support primary and secondary endpoints and objectives of the trial. Routine sample analyses for safety testing within clinical chemistry and haematology laboratories are not the central focus of this inspection programme. The legislation does not differentiate between the purpose of different types of sample analysis; this is a pragmatic approach by the MHRA Inspectorate and potentially any aspect may be reviewed during an inspection. It is expected that laboratories consider and implement the guidance to support all forms of sample analysis linked to clinical trials whilst ensuring that this is appropriate and proportionate for the work being undertaken.

Further information in relation to the conduct of clinical trials can be found in the Good Clinical Practice Guide (otherwise known as the “Grey Guide”) produced by the MHRA. This covers all aspects of GCP and is a recommended reference for laboratories involved in clinical research (including those not involved in primary and secondary endpoint analysis). The MHRA also continues to publish blogs on a variety of topics including those directly relevant to laboratory involvement in CTIMPs.

Human Tissue Act 2004 and the Human Tissue Authority

The Human Tissue Act (2004) relates to the removal, storage and use of ‘relevant material’ for Scheduled Purposes. The Act makes consent the fundamental principle to the use of human tissue. Although there are caveats, relevant material generally refers to samples, other than gametes, which include intact human cells. This includes most types of samples held in histology departments. Samples rendered acellular are not considered to be relevant material. There are eleven Scheduled Purposes for which consent is required, including ‘Research in connection with disorders, or the functioning, of the human body’. Samples collected as part of routine clinical care are not covered by the Human Tissue Act though they could subsequently be used for a Scheduled Purpose. The Act only applies in England, Wales and Northern Ireland as Scotland has separate legislation which only applies to samples collected from the deceased.

The Human Tissue Authority (HTA) was created by the Human Tissue Act to regulate the removal, storage and use of human tissue. They grant licences to establishments for a range of specific activities involving bodies and relevant material, including undertaking post-mortem examinations and storage of relevant material for Scheduled Purposes. HTA-licensed establishments include hospitals (both NHS and private), tissue banks, NHS blood and transfusion premises, private pathology services, clinical research facilities, commercial life sciences laboratories and universities.
Institutions handling human samples must ensure that they have any necessary licences for the activities carried out. The HTA does not license the ‘use’ of tissue for research or approve individual research projects or clinical trials. Neither do they have a role in the ethical approval of research. A research sector licence covers the storage of the relevant material. Storage of relevant material for research may be exempt from the need for a HTA licence if the sample is being stored according to the protocol of a research study approved by an NHS Research Ethics Committee. The HTA standards and guidance for research are contained within Code A: Guiding Principles and the fundamental principle of consent and Code E: Research. There is a useful decision chart for whether you need a research licence on the HTA website (Code E Annex C).

**Other regulation**

The regulations and guidelines that relate to research do not replace the existing regulations and guidelines that govern laboratory practice. All the existing professional standards and regulations still apply and those that relate specifically to research must be applied in addition to these. Examples include ISO-15189, the Data Protection Act implementing the General Data Protection Directive (GDPR) (as well as being custodian of patient samples Pathology is also custodian of their data) and the Transport of Dangerous Goods regulations. Researchers must be aware of, and adhere to, local NHS Trust policies and relevant professional standards such as those set by the General Medical Council (GMC), Royal College of Pathologists, the Institute of Biomedical Science (IBMS) and the Association of Clinical Scientists (ACS). These professional standards include the need to be appropriately qualified for their job roles and the need to maintain continued professional development mirroring requirements in research regulation. Further, international trials may be subject to the scrutiny of other agencies such as the US Food and Drug Administration (FDA) so it is important to consider differing regulatory requirements depending on the intended use of the data.

**Accreditation and research**

NHS Pathology laboratories and those laboratories who are providing clinical services are used to what was Clinical Pathology Accreditation and is now Accreditation to ISO 15189 (Medical Laboratories - requirements for quality and competence in medical laboratories) through the United Kingdom Accreditation Service (UKAS). These standards are about Quality Management supporting a culture and systems that can also be used to support the needs of research regulations and standards.

There are no laboratory accreditation schemes for the analysis of clinical trial samples run by the MHRA. For CTIMPs, laboratories are required to be compliant with all relevant legislation which includes working to the principles of GCP. Where analysis is undertaken for the purposes of patient safety during a CTIMP and is part of a routine repertoire of practice, UKAS accreditation is taken into account. The regulations permit this work to be inspected but often there are other activities which are prioritised for inspection due to the risk-based approach taken.

Study sponsors frequently ask for the UKAS accreditation status of a clinical laboratory as part of their study set up process. However, ISO 15189 itself makes no specific mention of research. While the standards are complementary, laboratory practice also needs to quality manage areas of research practice not covered by ISO 15189. Examples include provision for the management of result reporting in CTIMP studies where blinding may be compromised, data integrity considerations and the strict processing of trial samples according to the trial protocol.

Laboratories can be accredited to Good Clinical Laboratory Practice (GCLP) by third party providers. GCLP is a term for various guidance documents with their origins in a set of standards produced by the World Health Organisation (WHO) for the purposes of supporting safety and efficacy in international studies in developing countries and subsequently reproduced by a number of non-government organisations. Good Laboratory Practice (GLP) applies to all non-clinical safety studies which are designed to determine the effects of a chemical on human health, animal health and the environment. GLP specifically excludes clinical experiments. Both GLP and GCP are statutory requirements. In contrast, adherence to this GCLP is not a statutory requirement and is not assessed during regulatory GCP laboratory inspections performed by organisations like the MHRA.

**Conclusion**

There has been an increased emphasis on the importance of research to the NHS. Pathology’s part in healthcare and research is set to become even more important with diagnostics being central to the rise in genomic, personalised and stratified medicine. It is important that laboratories can demonstrate they have oversight of the research activity undertaken in their department. Quality management, with clear and recognisable processes, record keeping (and therefore traceability) and good communication all support compliance with regulation and guidance. The regulation and guidance around research continues to evolve, reflecting developments in healthcare. It is important that pathologists remain informed about the regulatory requirements, standards and guidance and maintain their professionalism towards the research activities they are involved with. This paper is an overview to highlight some of the regulations and policies around research rather than a comprehensive review. Digital and molecular pathology are areas of research shaping future clinical practice. Through such continued leadership in research, pathology is and will remain a central scientific discipline in clinical practice.

**Data availability**

No data are associated with this article.

**Grant information**

The CM-Path initiative and QA Panel is supported by the National Cancer Research Institute (NCRI).
OJD is funded by the National Institute for Health Research (NIHR) Clinical Research Network West Midlands and is a member of the CM-Path Clinical Trials Working Group supported by the NCRI. CV is funded by the Oxford NIHR Biomedical Research Centre.

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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We would like to thank Jason Wakelin-Smith from the GCP & GLP Inspectorate, at the Medicines and Healthcare products Regulatory Agency and Christopher Birkett from the Human Tissue Authority for their helpful comments and suggestions in the preparation of this paper.

A list of CM-Path QA Panel members is available here.

References

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Bridget Wilkins
St Thomas' Hospital London and Royal Hampshire Country Hospital, Winchester, UK

This is a valuable brief summary of the regulatory landscape in the UK within which research use of pathology-related biosamples sits. It highlights the core contributions of the national bodies that regulate clinical trials, and the requirements of laboratory accreditation in the UK that underpin quality assurance of pathology samples. The cited sources will provide a useful and regularly updated source of information regarding MHRA, HRA, HTA and UKAS/ISO 15189.

I think that it would be helpful to extend the focus more explicitly beyond the field of clinical trials, to provide confidence for pathologists working in diagnostic settings who wish to undertake research work unrelated to clinical trials. To this end, a little more discussion about how NHS pathology laboratories can achieve HTA licensing for research storage of 'surplus' tissue, the topic of generic/specific consent, and HRA processes for achieving ethical review of research (particularly the processes for research deemed to raise minimal ethical concerns) would all be valuable.

Legitimate access to a useful dataset of clinical information to accompany tissue samples is often a perceived barrier to pathology-based research, which could usefully be 'myth busted' in an article such as this (HRA and GDPR considerations, in particular).

In my opinion, the topics I've touched on above are all inextricably linked to the regulation of tissue-based research and to the encouragement of pathologists to engage in well-regulated research - the 'holy grail' of the NCRI's CM-Path initiative.

Is the topic of the opinion article discussed accurately in the context of the current literature?  
Yes

Are all factual statements correct and adequately supported by citations?  
Yes

Are arguments sufficiently supported by evidence from the published literature?  
Yes
Are the conclusions drawn balanced and justified on the basis of the presented arguments?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Governance of biosample-based research within/outwith CTIMPs, patient involvement with consent for tissue-based research, biobanking quality standards, research in the field of haematopathology (specifically, myeloproliferative neoplasms).

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Author Response 12 Aug 2019

**Owen Driskell, NIHR CRN, Albrighton, UK**

We thank the reviewer and agree with their comments.

The constraints in the scope for this individual article did not permit an exhaustive exploration of the wider details specifically facing NHS laboratories hosting diagnostic archives. We agree wholeheartedly they represent a great resource for potential research that would benefit NHS services and ultimately patients and the care they receive. Further clarity in education and ‘myth-busting’ will benefit the cause of breaking down the perceived barriers to pathology-based research.

**Competing Interests:** No competing interests

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**Mark Sobel**
American Society for Investigative Pathology (ASIP), Rockville, MD, USA

The Opinion Article by Owen Driskell and colleagues cites and briefly explains the relevant regulations in the UK concerning research conducted in pathology departments, with an emphasis on clinical trials. Editorially, and consistent with the principles of the UK policy framework for health and social care research (reference 1 - https://www.hra.nhs.uk/planning-and-improving-research/policies-standards-legislation/uk-policy-framework/) the authors state that the “requirements of the regulatory environment should not be seen as a barrier to research and should not significantly impact on the work of the laboratory once established and integrated into practice.”
The article is organized, generally well written and succinct, and provides links (in the references) to the appropriate statutes and regulations.

This reviewer recommends (in the section “Human Tissue Act 2004 and the Human Tissue Authority”) that the authors clarify whether the Human Tissue Act covers analytes stored in pathology departments such as DNA, RNA, and protein that have been isolated from human tissues. The Human Tissue Act does not consider acellular samples to be “relevant material;” however the Act clearly covers the use of DNA in research.

Editing suggestions for consideration by the authors:

- Abstract (sentences 2-3): “As the custodians of patient samples, pathology departments have a duty to ensure compliance with the relevant regulations, standards and guidelines to ensure the ethical and effective use for their intended investigational analysis, including when patients are participating in a research study.”

- Paragraph 2 (sentences 1-2): “In addition to research initiated by pathologists, a large amount of research is led by researchers from outside pathology requiring services delivered by pathology departments.”

- Paragraph 2 (sentence 4): “Data generated from pathology departments support research in a wide variety of ways.” (“Data” is plural.)

- Paragraph 13 (beginning “The primary focus of inspection by the MHRA…”) the second and third sentences could be misinterpreted. I suggest rewording as follows: “Although routine sample analyses for safety testing within clinical chemistry and haematology laboratories are not the central focus of this inspection programme, the legislation does not differentiate between the purpose of different types of sample analysis; this is a pragmatic approach by the MHRA Inspectorate and potentially any aspects may be reviewed during an inspection.”

- Paragraph 17 (sentence 2): All the existing professional standards and regulations still apply and those that relate specifically to research must also be applied.”

Is the topic of the opinion article discussed accurately in the context of the current literature?  
Yes

Are all factual statements correct and adequately supported by citations?  
Yes

Are arguments sufficiently supported by evidence from the published literature?  
Yes

Are the conclusions drawn balanced and justified on the basis of the presented arguments?  
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Biomedical research ethics, biobanking, molecular pathology, neoplasia
I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 12 Aug 2019

Owen Driskell, NIHR CRN, Albrighton, UK

Thank you for the reviewers helpful comments.

In response to the question of DNA.

DNA itself - as opposed to the cellular material from which it originates - is not considered to be relevant material under the HT Act. Its storage is therefore not subject to HTA licensing.

However, the reviewer is correct. The Act does state to anyone holding cellular material without the qualifying consent of the person/s concerned, intending to analyse the DNA and use the results, may be breaking the law. It is an offence to analyse DNA without qualifying consent unless it is for an excepted purpose.

Having human tissue, including hair, nail, and gametes (i.e. cells connected with sexual reproduction), with the intention of its DNA being analysed without the consent of the person from whom the tissue came or of those close to them if they have died is DNA Theft (Medical diagnosis and treatment, criminal investigations, etc., are excluded).

Appropriate informed consent is at the heart of Human Tissue Regulation and at the heart of Research Regulation. Pathology professionals must understand their role in research, and the systems in place by which informed consent is achieved, to give confidence to their compliance with this regulation.

RNA is not covered by the Human Tissue Act 2004 but the guidance produced by HTA also applies to RNA analysis where it is to be used to provide information about DNA.

Competing Interests: No competing interests.

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