
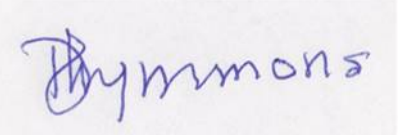


Standard Operating Procedure

Number:	UM/UoM RegisteringnonUoMCTIMPs/SOP04/5.0		
Title:	Registration of a non-University sponsored CTIMP		
Version:	5.0 (March 2018)	Effective Date	March 2018
Author:	Dr Mohammed Zubair / Victoria Sheard	Review Date	March 2020
Reviewed by :	Dr Mohammed Zubair		
Approved By:	Prof Deborah Symmons		
Position:	Research Governance, Ethics and Integrity Manager	Position:	Chair Clinical Trials Management Group
Signature:			
			

Version	Date	Reason for change
2.0	January 2013	Update of web links and office details
2.1	May 2014	Addition of version control statement for SOP
3.0	October 2015	Update of web links and office details
4.0	October 2016	Update of web links and office details
5.0	March 2018	Review and update of content

When using this document please ensure that the version you are using is the most up to date either by checking on the [Research Governance, Ethics and Integrity](http://www.staffnet.manchester.ac.uk/services/rbess/governance/) Team website (<http://www.staffnet.manchester.ac.uk/services/rbess/governance/>) for any new versions or contacting the author to confirm the current version.

UM/UoMRegistrationnonSponsoredCTIMPs/SOP04/5.0

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1.0 Background

The EU Good Clinical Practice (GCP) Directive 2001/20/EC was introduced to establish standardisation of research activity in Clinical Trials throughout the European Union. It was transposed into UK law as the Medicines for Human Use (Clinical Trials) Regulations 2004 (SI 2004/1031) which came into force on 1st May 2004. The Medicines for Human Use (Clinical Trials) Regulations together with subsequent amendments will be referred to as the Regulations in the rest of this document.

Regulation 3, as amended by Statutory Instrument 2006/1928, requires that research which falls under the regulation has a “Sponsor”. As stated in the Regulations, the Sponsor “in relation to a clinical trial, [is] the person who takes responsibility for the initiation, management and financing (or arranging the financing) of that trial”. Therefore all proposed research which falls within the scope of the Regulations must have a confirmed Sponsor.

University of Manchester employees are required to register proposed research projects which are Clinical Trials of Investigational Medical Products (CTIMPs) or Devices with the Research Governance, Ethics and Integrity Team, whether or not the University of Manchester is being requested to act as Sponsor. If the University is being requested or may be requested to act as Sponsor or co-sponsor for a CTIMP please follow the University Sponsorship of a Clinical trial SOP (Processing requests for confirmation of sponsorship for funding applications for Clinical Trials of an Investigational Medicinal Product or Device).

The Medicines and Healthcare products Regulatory Authority (MHRA) expects the University to be aware of all CTIMPs which are being led by its employees whether or not the University is acting as Sponsor of the study. This is particularly important if any of the funding for the trial is coming via the University.

2.0 Purpose

This SOP describes the process for registration of involvement or proposed involvement in a CTIMP for which the University of Manchester (UoM) is not and will not be asked to take on the role of Sponsor. This registration should be completed if all of the following apply:

- The Chief Investigator or other lead investigators (e.g. Laboratory Lead Investigators) of the CTIMP is an employee of the UoM.
- Full sponsorship of the trial is being undertaken by an organisation(s) other than the University
- The study falls under the UK Policy Framework for Health and Social Care Research and The Medicines for Human Use (Clinical Trials) Regulations 2004.

UoM employees whose only involvement in a CTIMP is to recruit, treat or follow patients according to a standard protocol (for example a commercial trial or an investigator led trial) need not register that study.

Each CTIMP need only be registered once by the most senior University of Manchester investigator.

This process of registration will allow the University of Manchester to:

- Retain a record of clinical trials undertaken by its staff
- Where appropriate, discuss UoM processes with the UoM lead investigator
- Where appropriate, discuss research governance issues with the UoM lead investigator

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- Allow the UoM, as the employer of the investigator, to have oversight of the research activities
- Ensure that the investigator(s) have suitable insurance in place before the project starts
- Have oversight of samples being stored and/or analysed on University premises, where such samples form primary or secondary endpoints of a CTIMP.

3.0 Procedure

This procedure only concerns studies that fall into (or potentially fall into) the category of clinical trial of an investigational medicinal product or a medical device, as defined by the Clinical Trials Regulations.

3.1 The University of Manchester Investigator should register their research with the Research Governance, Ethics and Integrity team by completing the Study Registration Form (Appendix 1) and send through the study protocol. If in doubt as to whether or not to register a study please contact the [Research Governance, Ethics and Integrity team](#).

3.2 Registration should take place at the time of application for funding if the submission for funding is being processed via the UoM. Registration should take place after funding has been awarded if another institution is taking the lead with regards to securing funding.

3.3 The [Research Governance, Ethics and Integrity Manager](#) may require evidence of external sponsorship and details of the funding of the study (e.g. a completed faculty research approvals form such as the Pan Man form).

4.0 Appendix

Appendix I: Registration form for a CTIMP in which the University of Manchester is not acting as Sponsor

5.0 References:

- UK Policy Framework for Health and Social Care Research
- Clinical Trials Regulations

Appendix I. Registration form for a CTIMP in which the University of Manchester is not acting as Sponsor

Please answer as many questions as possible and forward this form to [Research Governance, Ethics and Integrity Manager \(Mohammed.Zubair@Manchester.ac.uk\)](mailto:Research.Governance.Ethics.and.Integrity.Manager@Manchester.ac.uk) together with the study protocol and evidence of external sponsorship and Faculty/School approval.

Study Details	
Full Project Title:	
EudraCT Number	
REC Ref:	
Chief Investigator:	
Institution of the chief investigator (if not the University of Manchester)	
Senior University of Manchester investigator (if not the CI)	
Role of the senior University of Manchester investigator in the trial (if not the CI)	
Funding body/bodies	
Sponsor(s):	

		Yes	No
1	Is the attached protocol the final version?	<input type="checkbox"/>	<input type="checkbox"/>
2	Is the research a clinical trial of an investigational medical product (CTIMP)?	<input type="checkbox"/>	<input type="checkbox"/>
3	Is the research a Medical device trial?	<input type="checkbox"/>	<input type="checkbox"/>
4	Does the research involve non University of Manchester researchers?	<input type="checkbox"/>	<input type="checkbox"/>
5	Has funding been awarded for this trial?	<input type="checkbox"/>	<input type="checkbox"/>
6	Is any/all of the confirmed/potential funding commercial?	<input type="checkbox"/>	<input type="checkbox"/>
7	Is any/all of the confirmed/potential funding non-commercial?	<input type="checkbox"/>	<input type="checkbox"/>

8	Does any of the confirmed/potential funding for the trial come via the University of Manchester?	<input type="checkbox"/>	<input type="checkbox"/>
9	Is the University of Manchester the lead institution with respect to the confirmed/potential funding?	<input type="checkbox"/>	<input type="checkbox"/>
10	If funding is via the University is there evidence of School level approval (e.g. a PanMan form)?	<input type="checkbox"/>	<input type="checkbox"/>
11	Are any samples stored and/or analysed on University premises that form primary or secondary endpoints associated with a CTIMP?	<input type="checkbox"/>	<input type="checkbox"/>

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