standard pack and should also be given verbally by the HCP. This means that the additional information provided to the patient about the *research* component (randomisation, data collection and use, additional risks etc.) can be relatively brief.

This also applies to pragmatic trials involving ‘unlicensed medicines’¹ i.e. medicines that are used outside the terms of their UK licence or which have no licence for use in the UK but are commonly used in some areas of medicine, particularly paediatrics, psychiatry and palliative care due to the absence of suitable licensed treatments.

Where the outcome data can be extracted anonymously via electronic records or via the patient’s HCP, the consent process can be focussed on the research intervention itself. However, in other studies, where it is not possible to extract outcome data in an anonymised way, informed consent will also need to be sought for accessing and sharing the patient’s identifiable data and/or samples in addition to the intervention.

The following is an example of a short Participant Information Sheet that may be adapted and used in a pragmatic trial conducted to compare two medicines or other treatments that are routinely prescribed within the NHS.

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**We are inviting you to take part in a research project called [Trial name].**

**You do not have to take part if you do not want to.**

**Please read this information which will help you decide.**

**Research Title:** [e.g. A research study to find out if [X] is better than [Y] for treating people with [medical condition]].

**IRAS Reference Number:**
EudraCT No./EU trial number/Other registry No. [As applicable]

**Why am I being asked to take part in this research?**

You and your doctor have agreed that you would benefit from treatment for [patient’s medical condition]. [X] and [Y] are [two] licensed/commonly used treatments routinely used to treat [patient’s medical condition] and they are believed to be equally good. However, we do not know which is best.

In order to find out whether [X] or [Y] is better we are inviting patients like you to take part in a research project in which some patients will be given [X] and some patients [Y] and the two groups of patients compared.

Although you would not receive any extra benefit from taking part, research like this helps to continually improve the treatments and care provided to all patients now and in the future.

**Do I have to take part?**

No.

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¹ See the General Medical Council’s ‘Good practice in prescribing and managing medicines and devices (2013)’ - Prescribing unlicensed medicines: [http://www.gmc-uk.org/guidance/ethical_guidance/14327.asp](http://www.gmc-uk.org/guidance/ethical_guidance/14327.asp) for further information

² Required by forthcoming **EU Clinical Trials Regulation**

Please check HRA PIS and consent guidance for updates (this version was released on 10th February 2017)
It is entirely up to you to decide. If you do not want to take part that’s OK. Your decision will not affect the quality of care you receive.

If you decide NOT to take part you and your [GP/Doctor/healthcare professional] will agree on which treatment you will receive. This may be the same as the treatment you would have received by taking part in this research project.

If you do decide to take part you are free to withdraw at any time, without giving a reason, by contacting your [GP practice/Doctor/healthcare professional].

What will I need to do if I take part?

If you agree to take part in this research you will be given either [X] or [Y] both of which are used to treat [patient’s medical condition].

[Or if cluster design33: If you agree to take part in this research you will be given [X]/[Y] which is routinely used to treat [patient’s medical condition] in the NHS but may not be the treatment usually prescribed by [your GP/GP practice/Doctor/this hospital etc.].

Everybody taking part in this study, in this [describe cluster unit: ward/hospital/GP practice etc.] will be treated with [X].]

You do not need to do anything more. All the information needed for the research (but not anything that could identify you) will be collected from your medical records and shared with the researchers.

[Describe any additional samples/tests etc. beyond normal care]

If you choose to take part in this study, it will last for [duration of individual participant’s involvement]. The entire research will last for [duration of study]. You will not have to make any extra visits to your doctor over and above those needed for your normal care.

At the end of the research, or earlier if you experience any unpleasant side effects, your [GP/Doctor/healthcare professional] will discuss with you whether you should continue with the treatment you are taking or switch to another treatment.

What are the disadvantages/risks?

[There are no extra risks involved in taking part in this research.]

[There are only minimal risks involved in this research. These are (provide detail of any potential risks due to additional research procedures)]

The possible side-effects of the medicine you are given will be explained by your [GP/Doctor/healthcare professional] and are also provided in the information leaflet that comes with that medicine.

If we do find that one treatment is better than the other for you the trial will be stopped [and you will be switched to the better treatment]

A summary of the results of this research will be made available to all those taking part who would like to receive this34. [Provide details of how the results will be made available]

What will happen to information collected about me during the study?

33 A type of research design that randomises the drugs or treatments being investigated to different groups or clusters of individuals (such as households, primary care practices, hospital wards, classrooms, neighbourhoods or communities), rather than individuals.


Please check HRA PIS and consent guidance for updates (this version was released on 10th February 2017)
Your medical information will be kept strictly confidential by your doctor. The researchers will only be given as much information from your medical records as is needed for this research and that information will be anonymised. They will not be given your name, where you live or anything that could identify you.

Who is organising and funding the research?

This study is being carried out by [details of researcher(s), Sponsor and institution(s)].

[If applicable: The researchers will pay your GP/GP practice/Hospital etc. £[amount] for including you in this study.]

The research is funded by [name of funder (if different from Sponsor)].

Thank you for reading this information and for considering taking part in this research

Further Information: You can ask your [GP/Doctor/healthcare professional or other nominated person] any questions you may have about the study.

You may also obtain more detailed information about this research, including how your medical information will be used, your privacy protected, and the compensation arrangements in the unlikely event that anything goes wrong from [this website: [URL] and/or your GP/Doctor/healthcare professional etc.]

Contact Details:

Your [GP/Doctor/healthcare professional]:

Chief Investigator:

PIS Version No. …………… Date………………

2.6. Consent in postal/self-completion surveys

For postal/online surveys or self-administered questionnaire-based research, it is not necessary to include a separate Participant Information Sheet or consent form. Participants should still be provided with sufficient information to enable them to reach an informed decision whether to complete and return the survey/questionnaire or not but this may be included as a short introductory paragraph as part of the survey/questionnaire itself or included in a short covering letter.

Provided that the information adequately describes in broad terms the nature and purpose of the research, why they are being invited to take part, how the information collected will be used and stored, and how the findings might be made available to them, then completion and return of the survey/questionnaire will indicate consent on behalf of the participant.

Where the research involves sensitive questions and/or potentially greater threats to participant confidentiality then this should be clearly spelt out in the covering letter to participants; there is still no need to seek separate written consent to take part in a postal survey prior to the participant completing the questionnaire.
2.7. **Good Clinical Practice (GCP) training for those seeking consent: A proportionate approach**

Good Clinical Practice (GCP) is an international ethical and scientific quality standard for designing, conducting, recording and reporting clinical trials that involve the participation of human subjects.\(^{35}\)

The International Conference on Harmonisation GCP Guideline (ICH GCP) (as adopted by the Committee for Medicinal Products for Human Use (CHMP)) is part of European guidance, as an element of EudraLex Volume 10, and as such should be taken into consideration, where appropriate, as an established standard for GCP. In particular, if a study is to be included as part of a marketing authorisation application then it is an expectation that ICH GCP should be complied with, and this is referred to in the annexes to the Notice to Applicants (Volume 2B) for the Common Technical Document.

Both the HRA and the MHRA advocate a proportionate approach to the application of GCP to the conduct of clinical trials and the appropriate training of staff involved, including those seeking consent from potential participants.

Sponsors of CTIMPs which are **not** to be included as part of a marketing authorisation application can choose to comply with ICH GCP as a standard in its entirety or they can take a more proportionate approach depending on the nature of the trial. Further information about this can be found in the MHRA guidance on risk adapted approaches in the management of CTIMPs.\(^{36}\)

However, it is important to emphasise that for all CTIMPs it is the **“conditions and principles”** of GCP set out in the The UK Clinical Trials Regulations (SI 2004/1031, as amended) (see Annex A) that **must** be complied with. The **principles** of GCP are high level and may be interpreted in relation to the individual trial and **in proportion to the risks posed to the participants and to the integrity of the results**.

The UK Clinical Trials Regulations stipulate that:

> “each individual involved in conducting a trial shall be qualified by education, training and experience to perform his tasks” (Schedule 1, Part 2, 2).

Staff involved in the conduct of clinical trials need to be appropriately trained so that all investigators know what is expected of them in relation to trial procedures, and in order to ensure that the conditions and principles of GCP can be applied to any trial in a proportionate manner.

The training required does not need to follow a generic syllabus, format or prescribed timing but should be appropriate and proportionate to the activities undertaken by staff involved in the clinical trial. It should be tailored to the specific roles and responsibilities being undertaken by an individual. For example, it may be appropriate that some staff only receive an overview of the clinical trial, which could be in the form of a written summary; or they could simply be made aware of the local trial team contacts and have an awareness of, rather than a detailed knowledge of, ICH GCP requirements.

In the case of pragmatic trials, involving only minimal risk related to the research, it may be appropriate for the HCP to simply have an awareness of GCP requirements (which could be

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\(^{35}\) ICH Harmonised Tripartite Guideline - Guideline For Good Clinical Practice E6(R1) Current Step 4 Version Dated 10 June 1996

\(^{36}\) MHRA- Risk-adapted Approaches to the Management of Clinical Trials of Investigational Medicinal Products, Oct 2011.

*Please check HRA PIS and consent guidance for updates (this version was released on 10th February 2017)*