

Approvals from other authorities

All changes to a non-first-in-human trial have been reviewed and approved as part of a substantial amendment or modification in the EU, EEA, or USA, provided that the UK modification includes the same version of the documents and does not include any UK-specific aspects

Changes to the protocol

A change to the primary objective of the clinical trial

Use of new measurements for the primary endpoint

Changes to the trial design which have a significant impact on statistical consideration

Changes in the definition of the end of the trial (except a change to the planned end date, which is minor)

In trials involving an IMP authorised for use in the UK and used according to that authorisation, a change to the number of planned interactions with the participants to assess their ongoing safety in the trial, unless the change is in response to a new safety concern (which is a Route A substantial modification)

Changes to the list of concomitant medications that the participant can or cannot take

Inclusion of a UK-specific addendum or protocol into a global protocol version, unless the addendum or protocol results in a change to approved safety reporting

Changes to the Investigator's Brochure (IB) or Summary of Product Characteristics (SmPC)

Addition of new toxicological or pharmacological data relating to the IMP (including new interpretations of data) of relevance for the investigator, unless changes to the protocol are required from a safety perspective (which is a Route A substantial modification)

Changes to the reference safety information (RSI) involving an increase in frequencies with no new expected adverse reactions

Changes to the IB that do not change the initial risk and benefit assessment of the study or the safety profile of the IMP that was approved by the authorities

Updates to section 4.8 in the SmPC when this is used as the supporting document in place of an IB for an IMP (i.e. when section 4.8 is considered the RSI)