

Medicines & Healthcare products Regulatory Agency

Additional criteria

Any of the IMPs used in a trial that is eligible for automatic approval under Condition A are placebos. This is a potential safety concern because use of a placebo in these trials (but not trials eligible for automatic authorisation under Condition B or C) can change the risk-benefit profile, as there is no benefit for patients on a placebo

For a trial eligible for automatic approval under Condition C, the application uses a different version of one or more of the:

- protocol
- Investigator's Brochure (IB)
- IMP dossier (for EU or EEA approvals)
- IMP manufacturing process and controls (for USA approvals)

compared to the trial that was already approved in the USA, EU or EEA state. This is a potential safety concern because in this scenario, the document(s) being used for the trial will not have been approved by any authority

The applicant considers that the application may require the licensing authority or ethics committee to consult with a relevant committee or specialist group before issuing a decision

The application is for a trial that combines both an IMP and an investigational medical device or investigational in-vitro diagnostic device

The sponsor is requesting to disapply or vary the standard labelling requirements for at least one IMP involved in the trial that is not exclusively administered in a hospital or healthcare setting

The sponsor is intending to deviate from standard reporting of suspected unexpected serious adverse reactions