



TERMINOLOGIES USED IN CLINICAL TRIAL

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TOPICS COVERED

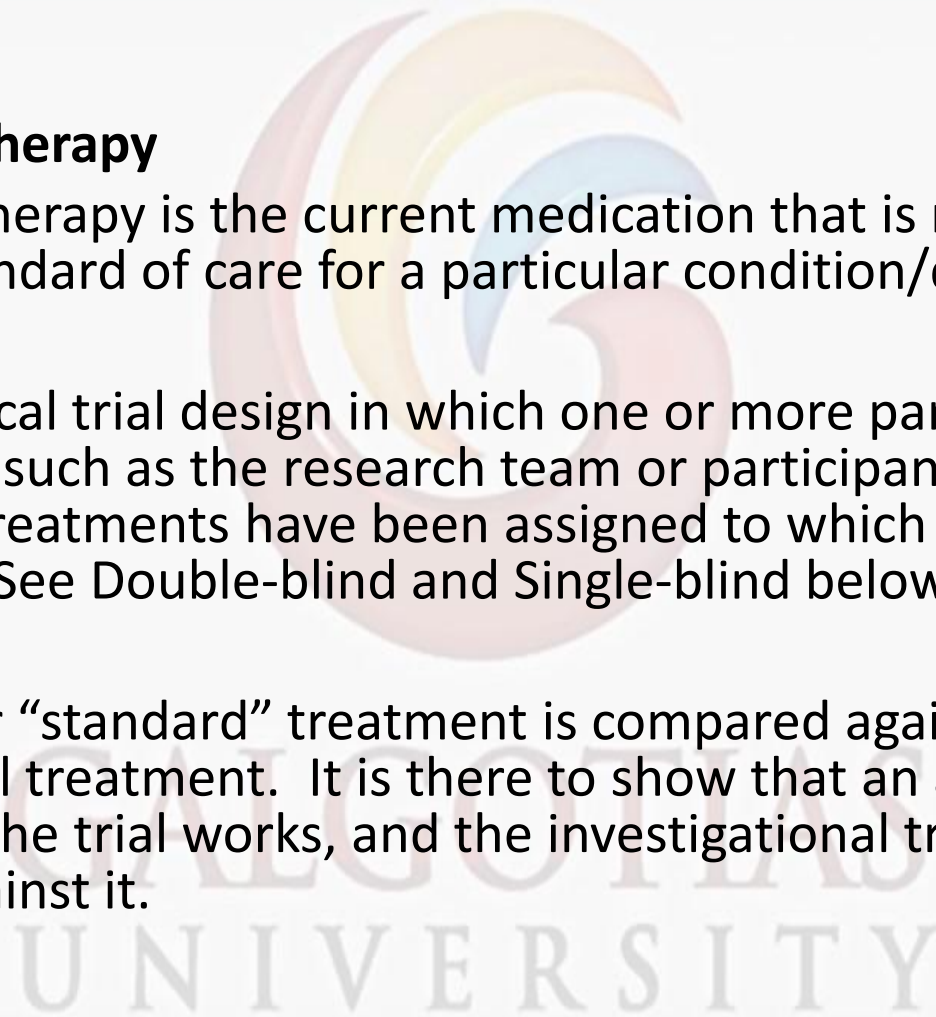
- DIFFERENT TERMS AND DEFINITION
COMMONLY USED IN A CLINICAL TRIAL

The logo of Galgotias University is a stylized, circular emblem. It features a central blue swirl that transitions into a yellow and orange swirl, all enclosed within a larger, light-colored circular frame. The overall design is modern and abstract.

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- **Adverse Event**
- A negative change or medical occurrence that happens during a clinical trial or within a certain time period after the trial has ended. An adverse event may or may not be caused by the treatment being studied.
- **Arm assignment**
- The assignment of a group or subgroup of participants in a clinical trial to receive interventions, or no interventions, as specified in the study protocol.
- **Assessment**
- A procedure (e.g. a blood test, scan, etc.) used to generate data required by the trial.

- **Background therapy**
- Background therapy is the current medication that is routinely taken as a standard of care for a particular condition/disease.
- **Blinding**
- A type of clinical trial design in which one or more parties involved with the trial, such as the research team or participant, do not know which treatments have been assigned to which participants. See Double-blind and Single-blind below.
- **Control**
- The control or “standard” treatment is compared against the investigational treatment. It is there to show that an approved treatment in the trial works, and the investigational treatment is compared against it.



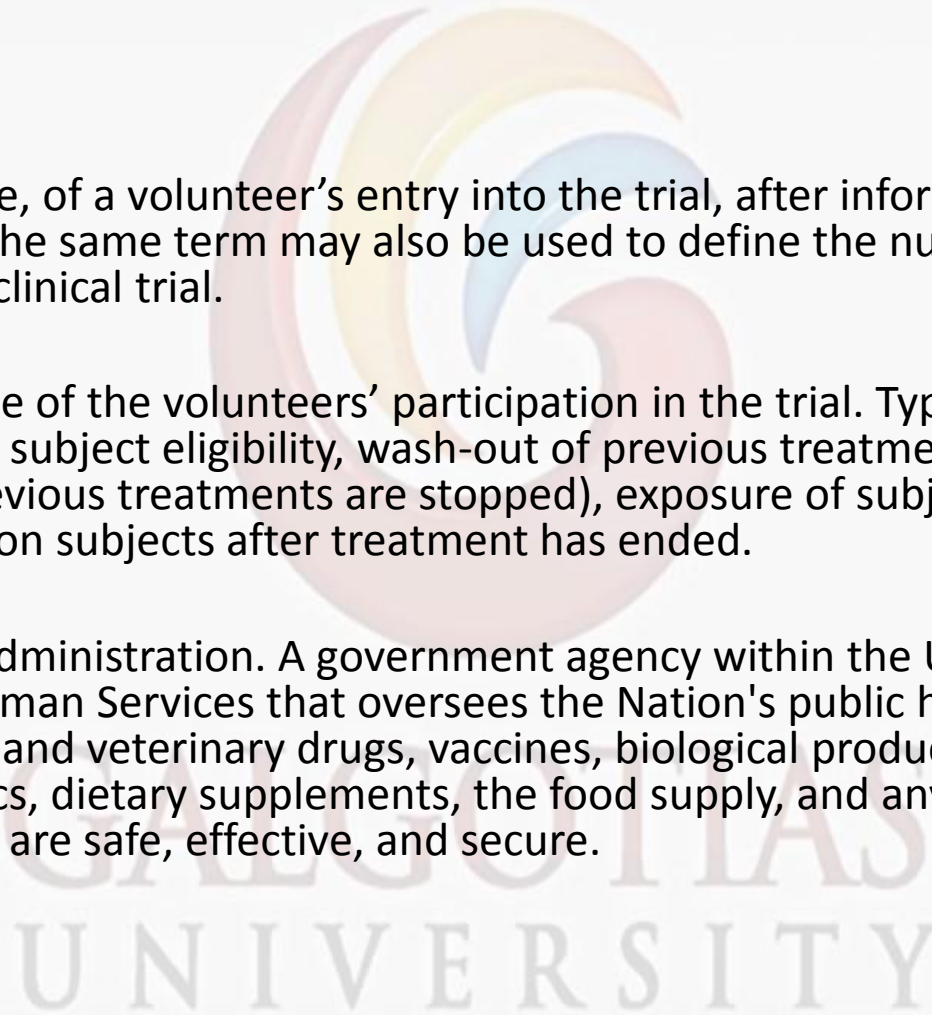
- **Clinical study**
- A research study conducted in human volunteers to answer specific health questions. Interventional studies determine whether experimental treatments or new ways of using known therapies are safe and effective under controlled environments.
- **Cross-over trial**
- A clinical trial where groups of volunteers are administered two or more interventions in a specific order. For example, a “two-by-two” cross-over trial design is where one group receives drug A at the beginning of the trial and then receives drug B for the rest of the trial. In the second group, participants receive drug B first and then drug A. Thus, the term “cross-over” is used to describe the order in which they are assigned; for example drug A and then drug B, or drug B and then drug A. All participants receive both drugs during the study.

- **Dosing discontinuation**
- Point/time when a patient volunteer permanently stops taking study drug for any reason. This may be at the end of the study or before the end if the patient wants to stop taking the medicine for some reason.
- **Double-blind**
- In a double-blind trial, only the study pharmacist knows what study medication a participant is receiving; the participants, doctors, nurses, and other clinical trial staff are not informed.

- **Early patient withdrawal (premature withdrawal)**
- Point/time when a patient exits from a trial prior to the planned completion of all investigational/trial drug administration and all assessments (including follow-up).
- **Eligibility Criteria**
- The requirements that people who want to participate in a clinical study must meet. Eligibility Criteria include both inclusion criteria and exclusion criteria and are defined in the protocol.
- **EMA**
- European Medicines Agency. An agency of the European Union that oversees the use of medicinal products.

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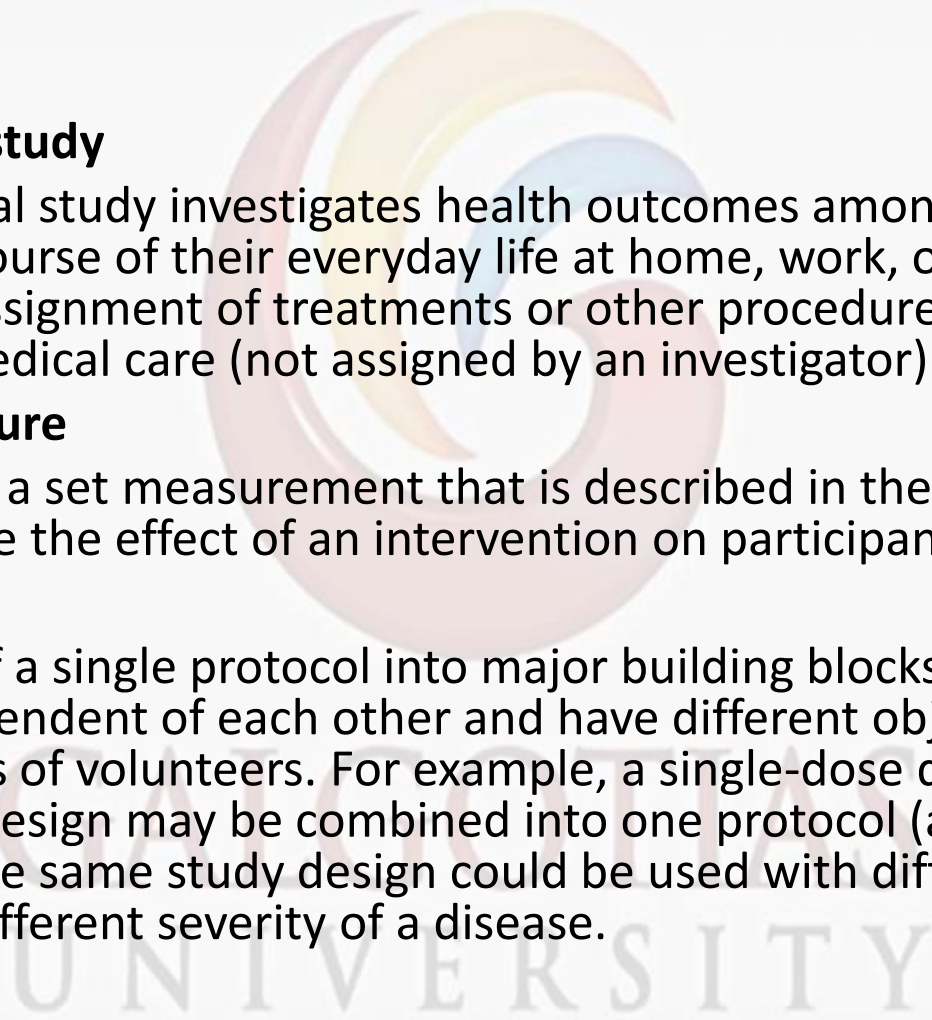
- **Enrollment**
- The point, or time, of a volunteer's entry into the trial, after informed consent has been obtained. The same term may also be used to define the number of participants in a clinical trial.
- **Epoch**
- The planned stage of the volunteers' participation in the trial. Typical epochs are: determination of subject eligibility, wash-out of previous treatments (i.e., a period of time when previous treatments are stopped), exposure of subject to treatment, or the follow-up on subjects after treatment has ended.
- **FDA**
- Food and Drug Administration. A government agency within the U.S. Department of Health and Human Services that oversees the Nation's public health by making sure that human and veterinary drugs, vaccines, biological products, medical devices, cosmetics, dietary supplements, the food supply, and any products that give off radiation are safe, effective, and secure.



- **Health Authority**
- A national or international health agency that has authority over and regulates a clinical study.
- **Indication**
- A disease, symptom, or particular set of circumstances that make a particular test, medication, procedure, or surgery advisable. For a treatment, an indication refers to the use of that treatment in treating a particular disease.
- **Informed consent**
- Informed consent is used by researchers to explain the clinical trial to potential volunteers. Its purpose is to protect the participant. It is used when somebody who is interested in participating first asks about the study and it continues throughout the study, until the study ends. The research team will review the details of the trial with the potential participant and will answer any questions. This information is also written in a document, known as the informed consent form, which is designed to be clear and easy to understand. If a person decides to enroll in a clinical trial, they will sign the informed consent form to acknowledge that they understand the details of the trial and consent to participating. The informed consent form is not a contract and the participant can withdraw from the trial at any time, and for any reason.
- **Institutional Review Board (IRB)**
- An IRB (also known as an independent ethics committee (IEC), ethical review board (ERB) or research ethics board (REB)) is a group of doctors, scientists, advocates, researchers, and members of the community that has been formally designated to review and monitor all research involving humans. IRBs are in place to provide ethical oversight and to minimize risk to participants.

- **Interventional study**
- Also known as a clinical trial, a type of clinical study in which participants receive one or more interventions, according to the protocol and group that they are assigned to, so that researchers can evaluate the effects of the intervention on a health condition.
- **Investigational drug**
- The drug being evaluated in the trial; this definition is synonymous with “investigational new drug” or “investigational medicinal product.”
- **Medication number**
- A unique number on the label of each investigational drug package that is used in a trial to dispense and track medication. The number is used to make sure the drug is supplied in the right quantities to different research centers.

- **Observational study**
- An observational study investigates health outcomes amongst groups of people in the course of their everyday life at home, work, or the doctor's office, where assignment of treatments or other procedures is as part of their regular medical care (not assigned by an investigator).
- **Outcome measure**
- In clinical trials, a set measurement that is described in the protocol and is used to evaluate the effect of an intervention on participants.
- **Part**
- A subdivision of a single protocol into major building blocks. These parts often are independent of each other and have different objectives or different groups of volunteers. For example, a single-dose design and a multiple-dose design may be combined into one protocol (a protocol with two parts) or the same study design could be used with different groups of patients with different severity of a disease.



REFERENCES

- Guide to Clinical Trials (Volume-I &II), DCGI
- Modules of Clinical trial methodology and management, RHE Life Science (CRO)

A large, faint watermark logo of Galgotias University is centered on the slide. It features a stylized circular emblem with three curved, overlapping bands in shades of yellow, blue, and red, resembling a 'G' or a spiral. Below the emblem, the words 'GALGOTIAS' and 'UNIVERSITY' are written in a large, light grey, serif font, stacked vertically.

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