# New Drugs: Their Development & Evaluation



#### New Drug Development

- Idea or hypothesis
- Design & synthesis of substances
- Studies on tissues & animal (preclinical studies)
- Studies on man (clinical studies)
- Official license (registration & market authorization)
- Post-marketing studies

#### Aims of Therapeutic Evaluation

- To assess efficacy, safety & quality of new drugs
- To expand indications for the use of current drugs
- To protect public health

#### Drug Development

- Drugs are chemical substances useful in prevention & diagnosis & treatment of diseases
- The process of drug development may be abandoned at any stage including after marketing (safety, inadequate efficacy)

#### Drug Development

- New drug development is <u>enormously expensive</u>
- Cost of development of a new chemical entity from synthesis to market <u>US \$ 500 million</u>
- The process may take 10-15 years

### Origin of Drugs

- Natural sources:
  - Plant origin like morphine, digoxin, atropine
  - Micoorganisms as fungi & bacteria synthesizing antibiotics
  - Animal origin like hormones (insulin), heparin
  - Mineral origin like iron, calcium

#### Origin of Drugs

- Synthetic when synthesized chemically in laboratories
  - These represent majority of drugs, as they are easily manufactured & cheaper like aspirin, paracetamol & propranolol

#### Medicines

- Medicines are drugs formulated in a suitable way for administration & use by patients
- Medicines consist of the active drug combined with excipients that give it shape, size, stability & other criteria as starch, Arabic gum & many other substances

### Therapeutic Investigation

- There are three questions to be answered during drug development:
  - 1. Does the drug work?
  - 2. Is it safe?
  - 3. What is the dose?

### Phases of Drug Development

- 1. Pre-clinical studies in animals
- 2. Clinical studies in human

#### 1. Pre-clinical studies in animals including:

#### A. General pharmacology studies:

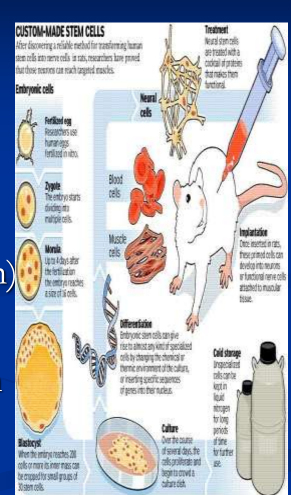
- Pharmacokinetic studies
- Pharmacodynamic studies
- Dose, preparation & routes of administration



#### 1. Pre-clinical Studies in Animals including:

#### B. Toxicological studies

- Acute toxicity
  - Special toxicity studies:
    - Reproductive system
    - Mutagenesis (mutation production)
    - Oncogenesis (malignancy)
    - **Teratogenicity** (harmful effects on foetus)



#### 2. Clinical Studies in Human

- These are carried out in humans in clinical trials centers & in hospitals under supervision of qualified investigators
- □ These include:



#### 2. Clinical studies in human

- Phase 1 studies
- Phase 2 studies
- Phase 3 studies
- Phase 4 studies

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## Phase 1 Studies (Human pharmacology)

These are performed on <u>a limited number of</u> <u>healthy volunteers</u> (20-50 subjects)

The aims of these trials are:

- Study of the general pharmacology of drug
- Pharmacokinetics (ADME)
- Pharmacodynamics (biological effect)
- Tolerability, efficacy & safety (associated adverse effects)

## Phase 2 Studies (Therapeutic exploration)

- These are carried out on a limited number of patients (50-300) to:
- General pharmacology of drug in patients
- Pharmacokinetics
- Pharmacodynamics
- Establish safety of drugs
- Assess potential therapeutic effects, comparison with placebo

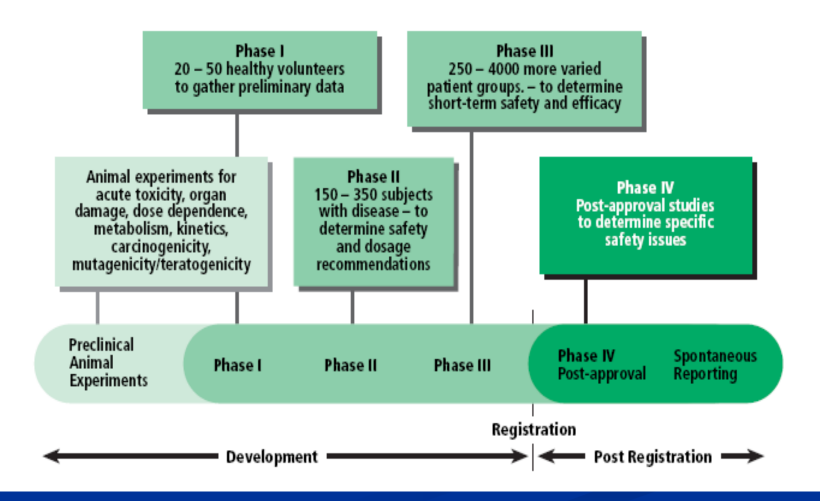
## Phase 3 Studies (Therapeutic confirmation)

- Randomized control trials
- These include multi-centre comparative studies on a large number of patients (250-1000) to establish therapeutic efficacy & safety, comparison with existing drugs
- Short term efficacy & safety

### Phase 4 studies (Therapeutic use)

- These include <u>post-marketing surveillance</u> (post-authorization studies) (2000- 10,000) to look for possible long term effects of drugs
- Long term efficacy & safety

#### Figure 1 Clinical development of medicines



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#### Clinical Trials

Clinical trials are carefully and ethically designed <u>controlled experiments</u> performed on <u>human beings</u> to evaluate certain aspects of drug studies

#### Aims of clinical trials

- Whether treatment is of value
- Magnitude of that value compared with other remedies
- Type of patients in whom it is of value
- Best method of applying treatment (how often, dosage of drug)
- Disadvantages & dangers of treatment

#### Fundamental to any clinical trial are:

- An hypothesis
- Definition of primary endpoints
- Method of analysis
- A protocol

#### Other factors when designing a trial:

- Characteristics of patients
- Size of trial
- Duration
- Method of monitoring
- Use of interim analyses

#### Subjects included in the studies are either:

- Healthy normal volunteers or
- Patients

#### Patients excluded from clinical trials include:

- Children
- Pregnant women
- Mentally ill patients

#### Techniques to avoid bias

- **■** Randomization:
- Introducing element of <u>chance</u> into <u>selection & allocation of subjects to treatments</u>
- Blinding

#### Criteria of clinical trials (CCT)

- Objective: should be clear & limited to one aim
- Careful design: A protocol should be prepared that shows design of the CCT prepared by clinical pharmacologist, physician & statistician
- Crossover design: when each subject is randomized to a sequence of two or more treatment, and he acts as his own control for treatment comparisons

#### Criteria of clinical trials

- Clinical trials may be of non-crossover design recruiting different subjects as a control group
- Balanced regarding sex, age, weight & disease state
- Double-blind technique when neither investigator nor subject knows about treatments they are receiving. This technique is important to:
  - Eliminate investigator bias
  - Eliminate patients or subject bias
  - Allow the use of placebo

- Single-blind technique is described when investigator knows but patient does not know treatment given to him
- Control group is used who will receive either placebo or a standard therapy
- Statistical analysis should be planned initially including the proper tests used

#### The use of placebo



It is a pharmacologically inert substance identical in all aspects to the active treatment indistinguishable from it

#### It is intended to:

- Eliminate observer or investigator bias
- Detect non-pharmacological effects of drugs (placebo effects)
- A control for statistical comparison

### Conditions that do not require use of placebo

- Therapeutic studies as it is unethical to deprive patients of treatments. A standard therapy is chosen instead of placebo
- When the active compound can be identified e.g. a vasodilator, alkaptonuria (nitisinone)
- Dose-finding studies
- Pharmacokinetic studies

#### **Ethical Considerations in Clinical Trials**

- Declaration of Helsinki
- The declaration of Helsinki (1964, 1975) sought to clarify the ethical principles governing clinical research involving human subjects emphasizing informed consent & proper scientific research design. It is the mission of doctor to safeguard health of people. The doctor's knowledge & conscience are dedicated to the fulfillment of this mission

### Recommendations are essential as a guide to doctors in clinical research:

- Risks & benefits must be carefully assessed
- Nature, purpose & possible hazards must be explained to subjects by doctor

### Recommendations are essential as a guide to doctors in clinical research:

- Informed written consent must be obtained
- Subjects must be free to withdraw from clinical trial anytime
- The investigators should discontinue research, if in their judgment it may if continued be harmful to subjects





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<u>Home Search Study Topics Glossary</u>

Search

ClinicalTrials.gov is a registry and <u>results database</u> of federally and privately supported clinical trials conducted in the United States and around the world. ClinicalTrials.gov gives you information about a trial's purpose, who may participate, locations, and phone numbers for more details. This information should be used in conjunction with advice from health care professionals. Read more...

Search for Clinical Trials

Find trials for a specific medical condition or other criteria in the ClinicalTrials.gov registry. ClinicalTrials.gov currently has 114,546 trials with locations in 177 countries.

Investigator Instructions

Get instructions for clinical trial investigators/sponsors about how to register trials in ClinicalTrials.gov. Learn about mandatory registration and results reporting requirements and US Public Law 110-85 (FDAAA).

Background Information

#### Resources:

**Understanding Clinical Trials** 

What's New

Glossary

#### Study Topics:

List studies by Condition

List studies by Drug Intervention

List studies by Sponsor

List studies by Location