Making a medicine. Step 3 and 4: Choosing a molecule or lead

Introduction

It takes over 12 years and on average costs over €1 billion to do all the research and development necessary before a new medicine is available for patients to use.

Medicines development is a high-risk venture. The majority of substances (around 98%) being developed do not make it to the market as new medicines. This is mostly because when you look at the benefits and risks (negative side effects) found during development do not compare well with medicines that are already available to patients.

The development of a new medicine can be divided into 10 different steps. The following article covers Steps 3 & 4: Lead generation and optimisation.



It takes well over 10 years of careful planning and research for a medicine to go from molecule to a marketable treatment.

Step 3: Lead generation

This step consists of finding a molecule that will interact with the target. The molecule might come from a natural source, for example a plant, or might have been made by chemists. These are called 'small molecules'. Hundreds of thousands of molecules will be tested to find 'leads' – molecules that interact with the target. Testing for leads is called a screening process. Modern robotic technology allows for 'high throughput' screening. This means that millions of molecules can be tested quickly. Once the leads have been generated or found, the process can move to the next step.

It is also possible to find large molecules (proteins) that interact with the target. These large molecules are much

more complicated to make. They are produced by biotechnology processes and are therefore described as 'biologics'. In biotechnology, the large molecules are produced by host cells of a living organism — such as bacteria, yeasts, or animal cells — in large fermentation vats. The proteins are then separated and purified. The purified proteins can then be used to explore interactions with the target.



Erythropoietin is a naturally occurring protein, but it has been used as a biologic therapy in patients whose bodies cannot produce sufficient amounts of it. The cells in the fermentation vats have been genetically engineered so that they produce large amounts of erythropoietin.

Traditionally, the pharmaceutical industry produced 'small molecules' which were then developed as new medicines. These days, large molecules (proteins) or 'biologics' are also produced. Both small molecules and biologics are important in medicines development.

Step 4: Lead optimisation

During lead optimisation, modifications are made to molecules to improve their effects. The screening process identifies 'leads' - molecules which interact with the target. However, these molecules often have only a weak effect and would not be suitable for further development. Chemists therefore alter the selected 'lead' molecule by adding or deleting elements. This creates a range of slightly different molecules. The molecule of an existing medicine could also be modified to improve or change its effect. Computer technology can help to design these molecules.

These modified molecules are then tested to determine which structure has the best efficacy and is better tolerated (safety). These studies help scientists gain understanding of the pharmacology of the molecule, meaning the way the molecule works in the body. The molecules with better efficacy and safety can then proceed for further testing as a 'candidate drug'. Around this time, the scientific and technical information of the candidate compound, for instance its molecular structure and effects, are usually registered, or patented, in order protect it as intellectual property.



Indomethacin

Optimisation of indomethacin to a potent CRTH2

antagonist. The original molecule on the left (called indomethacin) has been chemically altered (changes shown in red) to turn it into a candidate drug for a development project.

Remember, at each stage in the development process the results from a series of experiments are reviewed and a decision has to be made whether or not to proceed. A 'go' decision means that further investments will be made to fund the next set of experiments. If the information gathered during the experiments does not support further work, a 'no-go' decision is made and the project will be stopped.

Summary: Steps 1-4

A candidate medicine emerges only once the right target has been selected and the best lead compound has been identified. By this stage, the medicines discovery process will have:

- taken an average of 4.5 years
- involved testing a large number of molecules (5,000 to 10,000 – or more for small molecules), and
- cost an average of €500 million.

The candidate compound may be either a small molecule or a biologic.

References

1. Edwards, L., Fox, A., & Stonier, P. (Eds.). (2010).
Principles and practice of pharmaceutical medicine (3rd
ed.). Oxford: Wiley-Blackwell.

Attachments

Fact Sheet: Medicine discovery

Size: 905,797 bytes, Format: .docx

Medicine discovery. This fact sheet covers the steps of the medicines discovery and development process that occur before a compound can be tested in humans – from pre-discovery (gathering information on a disease) to non-clinical safety testing in animals.

Presentation: The basic principles of medicine discovery and development
 Size: 918,164 bytes, Format: .pptx
 The basic principles of medicine discovery and development. It takes over 12 years and over €1 billion to do all the research and development necessary before a new medicine is available for patients to use. This presentation details the process from discovery to release of a new medicine onto the market and beyond.

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