Animal testing is still the best way to find new treatments for patients

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**ABSTRACT**

Experimental research proceeds by hypotheses formulated on the basis of previous or new knowledge and then tested. If they are accepted, they serve as the basis for further hypotheses, and if they are rejected new hypotheses can be developed. In other words, when we are at the frontiers of knowledge the path is forged by “trial and error”. When a trial shows a hypothesis is wrong, this is a step toward making fewer errors.

This process also applies to drug development. There is no magic formula at present to predict - at the pre-clinical level - the therapeutic value of a drug for people with a disease. However, pre-clinical studies are needed in order to formulate hypotheses that justify clinical trials. Without these preliminary studies in selected animal species it would be unethical to test still unproven chemicals in humans.

**Keywords:**

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**1. Introduction**

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This process also applies to drug development. There is no magic formula at present to predict - at the pre-clinical level - the therapeutic value of a drug for people with a disease. However, pre-clinical studies are needed in order to formulate hypotheses that justify clinical trials. Without these preliminary studies in vitro and in vivo in selected animal species it would be unethical to test still unproven chemicals in humans.

There has recently been a shift in drug development. Historically, drugs were discovered by identifying the active ingredient from traditional remedies or serendipitously. Later, series of chemicals were screened on intact cells, isolated organs and whole organisms (functional screening) to identify substances with a desirable therapeutic effect. The sequencing of the human genome has permitted rapid cloning and purification of large quantities of proteins, and it has become common to use high-throughput screening of large chemical libraries against isolated biological targets which hypothetically are disease-modifying, in a process known as reverse pharmacology (targeted screening). Hits from these screenings are then tested in cells and animals for efficacy. In recent years scientists have been able to see the three-dimensional structure of target molecules and use that knowledge to design drug candidates.

Independently from the procedure followed, the discovery of new drugs has always been based on a series of variable interactions among data collected in patients, tissues, organs or cell culture and different animal species. However, in a large majority of cases clinical studies have been preceded by studies in mice, rats and other animal species which have led to suggestions for drugs to be tested in patients. Not always have the animal results been translated into effective drugs but the failures themselves have helped to reformulate the model or the experimental conditions or the type of chemical. The problem is selecting, for a given human target or function, the animal species that most closely resembles man, which should in principle be possible, drawing on the diversity of the animal kingdom.

The use of animals has always aroused controversy on ethical or technical grounds. Since the ethical issue is not the subject of this article, we shall analyze how animal experiments could be improved in order to increase their probability of predicting useful clinical results.

Are in vitro experiments alternative or complementary to animal tests?

Modern technology enables us to cultivate in vitro almost every kind of cell from all animal species including man. These cells can provide very useful preliminary information or help us understand how chemicals interact on the cell metabolism or functions, such as secretion of proteins, motility or enzyme activity. A few comments are necessary about whether in vitro tests offer an alternative and can therefore replace in vivo experiments.

First, drugs are easily available to cells in vitro while in vivo this is not always the case. For example, isolated cancer cells are more sensitive to an anticancer agent than in vivo because the complexity of a solid tumor, with the presence of inflammatory cells, inadequate vascularization,
fibrosis and other factors, limits the drug penetration into the cancer cells. Old studies already demonstrated that several anticancer agents are not distributed evenly in a growing tumor, reaching higher concentrations on the surface than in the center. More recent studies have shown that paclitaxel distributes unevenly in ex vivo slices of tumor, indicating that not all the targets may be available to the drug [1]; this situation is different from in vitro conditions where the drug is distributed more uniformly. The difference is extremely important when deciding about the probable efficacy of an anticancer agent and therefore its potential as a candidate for clinical trials.

Second, when a drug is given in vivo it encounters a number of barriers that are hard to reproduce in vitro. The blood-brain barrier is an example - it is supposed to protect the brain from exposure to exogenous chemicals. The barriers can sometimes be overcome because there are transport mechanisms. Another example is the intestinal barrier for drugs that are taken orally. In this case too it is difficult to mimic intestinal absorption in vitro. The drug may interact with the microbiome, affect intestinal motility, or be absorbed by fibers in food, metabolized by cytochrome P450 (Cyp) in the intestine, transported or rejected by the multidrug resistance (MDR) complex.

Third, when a drug is absorbed by the intestine it may bind to circulating proteins and distribute to various organs. The first pass is in the liver, where drugs can be profoundly metabolized to form several metabolites, depending on the Cyp system. These metabolites may have similar or different activities and in some cases they may be toxic or even counteract the action of the parent drug. Therefore in vivo, in contrast to the in vitro condition, the action of a drug may be related not just to a single chemical species but to a mixture of effects depending on other chemical species formed (the metabolites) and their interactions. To summarize, in vitro drugs are faced with a static system, but in vivo they are subject to a very dynamic condition where absorption, distribution, metabolism and excretion change with time.

Fourth, cells or tissue cultures cannot mimic the complexity of a living organism where cells are assembled in organs, under the influence of nerve, hormonal, immunological and circulatory systems. In particular, interactions between drugs and functional activities such as blood pressure, sleep or cognitive activities cannot at present be studied in vitro.

Fifth, the animal species employed for preclinical tests have many features similar to man. They have similar organs: brain, lung, heart, liver, etc., similar functions such as circulation, hormonal set-up, peripheral nervous system, immunological functions. The genomic organization too is common, although with different degrees of complexity; animal proteins are in most cases homologous to man; metabolic processes are similar.

All the reasons that distinguish the complexity of living animals from in vitro conditions also apply to the various animal species and strains, which can differ in their absorption, transport, distribution, metabolism and excretion as well as in the way in which they respond to the same drug. All these considerations imply that while cells or tissue culture are useful for studying drug activity they are complementary, not alternative, to in vivo studies. At present, animals are still the best model - however imperfect - to predict activity in man.

Why does translation from animals to man sometimes fail?

Hackman and Redelmeier [2] analyzed this question in a quantitative manner. Out of 76 animal studies retrieved from top journals, 37% were replicated in clinical randomized trials, 18% were contradicted and 45% remained untested. It is logical to assume that for therapies against bacterial, fungal or viral infections the translation from animals to man is more likely to be effective. Vaccines against poliomyelitis, meningitis and rotavirus are outstanding examples, as are a number of antibiotics and the recent agents against HIV and hepatitis C viruses. They illustrate the striking concordance between animal results and human benefits. Translation has also been fairly good with agonists or antagonists of chemical mediators. Beta-adrenergic blockers for the treatment of tachycardia, alpha-adrenergic blockers for hypertension, and beta-adrenergic agonists for asthma are also good examples of concordant results between animals and man. Similarly, serotonin antagonists have antiemetic activity and serotonin-uptake inhibitors act on some symptoms of human depression. Antihistamines are useful for the treatment of allergic reactions. Drugs acting on metabolism, such as insulin, oral anti-diabetic agents and cholesterol-lowering drugs have been successfully translated from animals to man. The history of today’s therapeutic armamentarium has always involved animal testing.

At the other extreme, however, we cannot overlook the poor correlations between results in animals and man in several diseases such as stroke, amyotrophic lateral sclerosis (ALS) and Alzheimer disease. A recent study found that the success rate of new drugs entering phase 1 clinical trials for diseases of the central nervous system is only 8% [2,3].

Several analyses have set out to understand why in many cases the results in animals and man differ. One obvious reason is the difference not so much in organ composition and functions but the greater complexity of man compared to all the animal species that could be used. For logistic and economic reasons mice and rats are the most widely used laboratory animals partly because their genomic, proteomic and metabolomic profiles as well as their organic functions and behavior are better known than for other species. The significant number of rodent strains extends the range of experimental testing. In addition, mice can be genetically mutated so we can investigate the functional role of single and combined genes. Mutated human genes responsible for or involved in human diseases can be transferred into mice, and recently “humanized” mice have been developed, which are useful models to reproduce some aspects of neonatal sepsis [4,5]. All these and future improvements may enable researchers to reproduce at least some features of most human diseases awaiting better treatments.

It must be admitted that an important area of discrepancy is the poor quality of some animal investigations [6]. For instance, amlodipine was tested in 22 trials of cerebral hemorrhage in man, with negative results - in apparent contrast with animal findings. However, when the animal results were systematically reviewed, it became clear that there was no benefit, indicating that animal findings did in fact overlap those in man. Similarly, out of nine drugs found effective in an animal model of ALS only one, riluzole, actually appeared to prolong patients’ survival – to a limited extent. But then, when these nine drugs were tested in mice by the ALS Therapy Development Institute, all of them, with the partial exception of riluzole, were inactive. Therefore, the discrepancy was due to the fact that the borderline results had been interpreted optimistically. In general, there is a regrettable tendency to over-rate the value of products that in fact show only marginal efficacy for a pathology that has no treatment.

It must be stressed that the progress made in controlling bias in clinical trials has not been translated to animal trials. Although animal publications far exceed the number of clinical trials, systematic reviews and meta-analyses are ten times less frequent for animals than for clinical research. Bias related to randomization, double blinding, surrogate end-points, calculation of sample size, statistical analysis, and non-publication of negative results still greatly limits the extrapolation of animal findings to man. The over-enthusiastic attitude of scientists, together with economic interests, have in several cases led to premature clinical tests.

In some cases the treatment schedule is inadequate, or blood and tissue concentrations are too low to affect a target or too high to be tolerated in man. In addition, the target in man may not follow the same pathway of events, with a different functional effect from that seen in pre-clinical studies. Frequently, for instance in experimental cancer research, treatments are preventive or are tried at too early a stage of tumor development – differently from the clinical condition. Therefore the inadequacy of the model or the time and doses of the treatment – as well as a critical evaluation of results – may explain the discrepancy rather than the translation itself.

How could we improve in vivo studies? There may be at least four essential approaches.
First, we need to intensify studies and develop techniques to improve and reduce the use of animals, following the 3R rule. For instance, identification of the insulin structure abolished the need to measure insulin’s potency on blood glucose in rabbits (R = replacement). Powerful analytical techniques such as the various types of mass spectrometry now permit drug measurements in very small amounts of blood while before rats and mice had to be killed to obtain large enough samples. Nuclear magnetic resonance and other non-invasive diagnostic tools in small animals have enabled us to follow the progression of a disease in the same group of animals (R = reduction).

Human organoids can now be developed using different kinds of stem cells and may serve to study normal and diseased organs in order to verify the therapeutic and toxic effects of drugs as an intermediate step between in vitro and in vivo tests [7]. In general, any technological development may open the way to new methods to improve the value of in vitro studies and/or reduce, refine or replace the need to use animals.

Second, animal testing needs to be improved by incorporating all the rules developed to improve clinical trials in order to minimize bias. As already indicated, systematic reviews and meta-analyses of preclinical studies show up publication bias in laboratory animals. An analysis of 76 animal studies published in top journals between 1980 and 2000 and relevant to translation from animals to the clinic rated only 48% as having good methodological quality, and regrettably the quality did not improve with time [2]. In another analysis, random allocation was used only in 12% of the tests and blinded outcome assessment only in 14% [8].

It is important to bear in mind that the objective of experimental research is to establish therapeutic value. For instance, in a meta-analysis on the tests to study statins there was a redundancy of publications on the lowering of cholesterol and other lipids but only very few studies on therapeutic targets such as myocardial infarction or stroke [9].

More comparative studies are needed on the metabolic, inflammatory and hormonal pathways in order to select the animal species closest to man for each pathway. Multicentric studies are needed, similar to clinical trials, to establish the reproducibility of results. An example is available concerning a new potential drug for stroke [10]. For more realistic pre-clinical studies it may also be possible to organize trials that test new drugs in pet animals with chronic diseases. Another way to reduce the number of animals is to establish bio-banks to store blood and organs from a given experiment: should it be necessary to do further biochemical analysis later, there will be no need to repeat the same experiment. It is also worth recalling that new drugs are often tested in animals against placebo, while comparative trials should really be done with already known drugs in order to establish what kind of evidence suggests the translation from animals to man.

Third, further research is needed to improve the translation of animal research to patients. The NIH recently launched a program to train preclinical scientists to plan their experimental trials better by applying the same rules as for clinical trials. The NIH has also invited applications to prolong life. Public opinion must be made aware that hypotheses have affected the epidemiology of human pathology, contributing to their “ivory tower” to explain the complexity of translating research results from animals to man. At the same time it cannot be stressed enough that animal studies have led to the production of drugs that have affected the epidemiology of human pathology, contributing to prolonging life. Public opinion must be made aware that hypotheses in the biomedical field are just as likely to fail as in any other field of research. Only continuing trial- and-error to understand errors will show the best way to reach a goal. Limitations to the use of animals, particularly other than rodents, are an obstacle to obtaining a wider spectrum of activity across species which may help in deciding when a treatment is suitable for patients. Nevertheless, there is certainly ample room for substantial improvement in the protocols of animal tests to boost their credibility and reproducibility. The validity of animal testing is not limited to translation to man, but also to their value for the treatment of animal pathology: a large number of drugs employed in animals are the same as those used in man [16].

Given the limited usefulness of computer and in vitro models, for the time being animal models remain the best alternative and their use must continue, considering that patients cannot just wait for better tests to cure their suffering.
Conflict of interests

I have no conflict of interest as regards the manuscript submitted.

References


