Clinical effectiveness of Traditional Medicines

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There is a general conception that, herbs are natural therefore, they are completely safe. In fact, herbs or herbal preparations also can cause toxic adverse effects, serious allergic reactions and adverse drug/diet interactions.

The efficacy of a drug substance is its capacity to produce a desired therapeutic effect, or the relative ability of a drug receptor complex to produce maximum functional response. Nowadays, in health care systems the term 'clinical effectiveness' or 'clinical governance' is preferred instead of efficacy, though both are having similar meanings. The term clinical effectiveness includes the sum total of the pharmacological and the non medical effects of bioactive compounds that may act synergistically, otherwise termed as placebo effect, or sometimes antagonistically or the nocebo effect.

The meaning response

The non therapeutic function of medicines plays a major role in executing the 'meaning response'. The 'meaning response' mainly arise from the physiological or psychological perception of mind in the treatment of illness. This includes mainly the brain mechanisms like- expectation, anxiety, and reward, in addition to other learning phenomena.

In regards to traditional herbal medicines, the socio cultural aspects are likely to elicit a physiological response in addition to the intrinsic pharmacological activity of the herb, since they often exist within religious and mythical traditions creating a collaborative 'meaning response'. Meaning response plays a very important role in the clinical effectiveness of traditional herbal medicines, for e.g. traditional healer may make magical claims, gestures, chanting or prayers while giving a crude herbal extract to the needed person, finally result in the so called meaning response.

Even the packaging, shape, size, color, odor, of a herbal dosage form may affect the clinical effectiveness of the drug, a theme which can entertain many levels of pharmacy practice research.
Evidence

In an era of evidence based health care, systematic reviews of randomized control trials (RCT) are becoming increasingly important as a source of evidence for decision making. The evidence of clinical effectiveness mainly comes from the use of high quality herbal ingredients in the formulations, systematic reviews and Meta analysis of randomized controlled clinical trials.

To establish the true effectiveness of herbs as medicines, they must be tested in prospective, double blind, randomized, placebo-controlled clinical trials (RCTs).

The main reliable reviews about the effectiveness of herbal drugs may be obtained from the well known Cochrane reviews and Medline reviews considered as the bench mark of high quality informations on health care systems. Cochrane reviews are systematic reviews of primary research in human health care and health policy and are internationally recognized as the highest standard in evidence based health care.

The major setback for the clinical effectiveness of herbal medicines is the lack of quality controlled trials or more precisely the lack of phase 3 and phase 4 trials. The official documents and quality reviews related to the randomized trials and post marketing surveillance studies may contribute improved clinical effectiveness indications about traditional herbal remedies.

Cochrane reviews indicated the effectiveness of devil's claw (Harpagophytum procumbens DC), white yellow bark (Salix alba), and Capsicum frutescense for reducing pain, compared to placebo, in short term trials. The review on St.John's wort (Hypericum perforatum) indicated its effectiveness in treating patients with depression which is found equivalent to selective serotonin uptake inhibitors. Feverfew (Tanacetum parthenium) indicated in migraine, found better than placebo. Reviews on Saw Palmetto (Serenoa repens) indicated its effectiveness in Prostatic hyperplasia.

The evidence of effectiveness was strong in the reviews for Harpagophytum procumbens, (grapple plant, wood spider or devil's claw) in the treatment of painful osteoarthritis and chronic low back pain. Harpagophytum preparations containing > 50 mg harpagoside in the daily dosage proved effective in reducing pain associated with osteoarthritis.

Evidences again include references from approved Pharmacopoeias, approved monographs, and independent written histories of use in the classical or traditional medical literature. Mainly three different categories of clinical trials are performed to identify the effectiveness of therapeutic claims. They are Interventional studies, Observational studies, and Research synthesis studies.

Interventional studies

These studies are considered as the gold standard of therapeutic efficacy which involves the randomized controlled clinical trial (RCT). In a randomized controlled trial, subjects similar to each other are randomly assigned either to receive the intervention or not to receive the participation.
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**Observational studies**

*In observational studies* of prospective nature, the investigators recruit subjects and observe them prior to the occurrence of the effect or outcome. In retrospective observational studies, investigators review the records of subjects and interview subjects after the effect has occurred. Various types of observational studies are available which include *Cohort studies, Case control studies, Cross sectional studies, Time series studies, In ecological studies, Descriptive epidemiology, Case reports* etc.

**Research synthesis studies**

Research synthesis studies including meta analysis may be useful as supporting evidence for a health claim. A meta-analysis may be conducted on several clinical trials of a medical treatment, in an effort to obtain a better understanding of how well the treatment works.

**Major setbacks of traditional medicines**

The absence of, or very few clinical evidences for herbal medicines is not considered as a serious problem by practioners or consumers in the complementary and alternative medicine segment, since it is believed that traditional use over many generations without known side effects or health threats, reflects that they are safe. The side effects of some plant medicines are not immediate, for example the production of nephrites by aristolochic acid, a phytochemical present in many plant species of Aristolochia genus.

Another segment lacking informations in the case of herbal medicines is the unavailability of the pharmacokinetic and pharmacodynamic data. Only for a few phytoconstituents or herbal products, the mechanism of action is well documented. But for the rest of herbal products the mechanism of action or the pharmacodynamics is not established clearly due to the complexity of multiple compounds present in the extracts.

In both Ayurvedic and Chinese medicines the complexity is multifarious by the fact that each preparation usually contains multiple herbal /other natural ingredients with each ingredient containing a number of active constituents. This increased level of complexity creates difficulties in establishing valid standards, and also due to the unknown toxicity of constituents, safety may be compromised, which directly reflects on the unavailability of registered Ayurvedic and Chinese herbal medicines in western market.

Treatment in Ayurvedic system is holistic, involving natural medicines, massage, diet and the regulation of life style. It sees each individual as having a unique mind body constitution and set of life conditions.

Other hindering factors for the non acceptance of these treatments by main stream western health care providers include the unstable nature of therapeutic efficacy of different preparations of Ayurvedic medicines and TCM and the lack of standardized clinical trials. The chemical constituents of plants vary depending on the species, variety and part of the plant, with conditions of growth (soil, water and temperature), and with the age of the plant. These complexities and variations of chemical content make standardization essential.
Newer scientific trends about herbal medicines

Application of novel trial designs such as Point Of Care clinical trial (POC –CT ), a newly derived clinical trial approach developed by Dr.Philip Lavori of Stanford University, compares treatments that doctors are already using, and collects data on which treatments work best within the context of real-world, everyday practice. Enrollment and randomization of study volunteers occurs during regular care - within the framework of a patient’s visit to their usual health care provider.

It is assumed that herbal medicines interact with bacteria present in the gut flora, leading to a kind of bioprocessing inside, may result in the conversion of inactive to active medicine compounds, or sometimes improve the population of beneficial bacteria which in turn improve the immune system performance. Phytopharmaceuticals proved useful in several cases as these compounds act synergistically with the synthetic drugs and may reduce the adverse reactions to some extent.

Reports again indicated that the phytoconstituent, piperine present in Piper longum and Piper nigrum, increases the bioavailability, $C_{\text{max}}$ and $\text{AUC}$ of several synthetic drugs amoxicillin, phenytoin, propranol and theophylline. Licorice has been shown to increase the plasma concentration of prednisolone.

Recent research on several Ayurvedic medicines indicated that, the total synergistic effect of the constituents lead to the inhibition of Stress activated Protein Kinase there by blocking the NFκB (Nuclear Factor kappa B) signal transduction pathway, finally results in apoptosis, and anti inflammatory responses.

The emergence of modern standardization tools like proteomics, bioinformatics and various chemical docking studies may improve the identification of the evidence based herbal constituents and products in the future.