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NEEM

The Divine Tree
Azadirachta indica

H.S.Puri
Herba Indica
India
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FOREWORD

For medicinal and aromatic plants the clock has slowly, but surely, turned a full circle. The use of plants for health care started, as recorded in Indian and Chinese treatises available of that time, about 2 to 3 millennia before Christ and reached its zenith in the first millennium AD. These mention the use of plants and plant-based preparations in human and animal health care and occasionally for the preparation of house-hold products. Thereafter for several centuries the plant materials occupied a pre-eminent position in the trousseau of a traditional practicing healer. It was only at the beginning of the second millennium that alchemic and mineral-based products also started appearing. The plants and herbals nonetheless continued to be widely used until the industrial revolution in Europe brought in synthetic products for various kinds of usages. These products, because of their ease of preparation and administration, led to a slide in the popularity of plant and herbal-based products. Among other reasons responsible for this was a belief (which still persists to a large extent) that most of the plant-based products/herbals are non-standard preparations and hence lack quality and efficacy. Noticeable batch-to-batch variations for the same products and lack of therapeutic consistency further eroded to some extent the credibility of these products.

At the present time, however, the increasing environmental degradation due to a burgeoning synthetic products industry has rung alarm bells the world over. Several scientists in various countries are now engaged in discovering or rediscovering the usefulness of plants and herbals for value-added products. Their quest for rediscovering the usefulness of plant materials has its basis in those leads or references which are mentioned in folklore or traditional systems as indigenous cures for several ailments. This resurgence of interest has enormous economic and commercial implications as well. However, at the same time the public at large and also scientists are conscious of the fact that if indiscriminately commercially exploited, this plant wealth may not last long. This has given rise to a paradoxical situation where, on the one hand, the public wants ‘green’ products, be it for medicinal use, personal hygiene or for its palate, but on the other, dwindling resources make us wary of environmental denudation. A balance has to be struck between demand and supply and in my view it can be best taken care of by sustained and structured ‘social forestry’ programs with an emphasis on planting those species which are proven sources of herbal drugs or phytomedicines.

If we dwell on this further, we find that this attitudinal change in the learned and lay public towards products originating from plant sources is basically because of a belief that ‘green’ products have distinct therapeutic advantages over allopathy in treating ailments like hepatitis, asthma, diabetes, arthritis, immune disorders, some tumours, etc. Likewise, cosmetics and biocides from plant sources are popular because they are soft for human use and ecofriendly. Commercial estimates indicate that 70 to 80 percent of the population in developing countries, accounting for over 50 percent of world population, depends partly or entirely on herbal remedies. According to Indian Medicinal Plants: A Sectoral Study, a report recently brought out by India’s Exim Bank, the global trade in medicinal plants is estimated to be approximately
US$60 billion, of which India’s share is about US$700 million. The world demand for herbal products has been quoted to be growing at a rate of 7 percent per annum. This growth in demand, especially in developing countries, is partly due to the ready availability of herbal remedies, a shortage of practitioners of modern medicines in many of these countries, and the socio-cultural background of the users. Farnsworth et al. (in bulletin of the World Health Organisation 63, 965–85, 1985) have mentioned that even in developed countries, plant drugs are proving to be of great importance. In the USA, for example, 25 percent of all prescriptions contained plant extracts or active principles derived from higher plants. At this juncture, the importance of a plant like Neems come to the fore.

Neem, a large evergreen tree, commonly found throughout the Indo-Malaysia region, has been the subject matter of numerous scientific studies. Scientists the world over have carried out extensive work on its botanical, medicinal, industrial and agricultural usages. Practitioners of the Indian ayurvedic system advise the use of Panchang (five parts) of neem, i.e. leaves, bark, fruit, flower and root, for various applications. The seed is another extremely useful part, especially for its oil. Extracts of various parts of neem have proven medicinal properties—anthelmintic, antifungal, antidiabetic, antibacterial, antiviral, antifertility, etc. It is for these properties only that the practitioners of ayurvedic, siddha, unani tibb and homeopathic systems of medicine make extensive use of its parts. It is my firm belief that it is a only matter of time before even the allopathic system starts to make use of its medicinal properties in a regular way. Neem’s use as an insecticide and pesticide is also well documented. There is no gainsaying the fact that its economic and commercial value lies in every single part having a proven utility.

I find this comprehensive treatise on Neem (Azadirachta indica) to be an excellent collation of the recorded observations, research efforts and accomplishments of scores of individual botanists, taxonomists, traditional medicine men, and medicinal chemists of past and present. It involves a massive effort on the part of the author wherein he has mapped the Neem in its entirety. Be it the botanical description, chemical constituents and products derived from its various parts, usage in human and animal healthcare, personal hygiene, household remedies or other commercially valuable products—the coverage has been extensive. As is observed in any monograph of this kind, it offers us a well-documented retrospect of the work carried out. The chapters are full of ideas which provide further leads to a researcher, good questions to an inquisitive mind, alternative ways to combat various ills to a physician, etc. I feel this monograph has come at the most appropriate time when everyone—scientists, the political system and citizens as part of non-governmental organisations—is concerned about dwindling natural resources and would like to have a direction on the sustainable use of what Mother Earth has provided.

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PREFACE TO THE SERIES

There is increasing interest in industry, academia and the health sciences in medicinal and aromatic plants. In passing from plant production to the eventual product used by the public, many sciences are involved. This series brings together information which is currently scattered through an ever increasing number of journals. Each volume gives an in-depth look at one plant genus, about which an area specialist has assembled information ranging from the production of the plant to market trends and quality control.

Many industries are involved such as forestry, agriculture, chemical, food, flavour, beverage, pharmaceutical, cosmetic and fragrance. The plant raw materials are roots, rhizomes, bulbs, leaves, stems, barks, wood, flowers, fruits and seeds. These yield gums, resins, essential (volatile) oils, fixed oils, waxes, juices, extracts and spices for medicinal and aromatic purposes. All these commodities are traded world-wide. A dealer’s market report for an item may say “Drought in the country of origin has forced up prices”.

Natural products do not mean safe products and account of this has to be taken by the above industries, which are subject to regulation. For example, a number of plants which are approved for use in medicine must not be used in cosmetic products.

The assessment of safe to use starts with the harvested plant material which has to comply with an official monograph. This may require absence of, or prescribed limits of, radioactive materials, heavy metals, aflatoxins, pesticide residue, as well as the required level of active principle. This analytical control is costly and tends to exclude small batches of plant material. Large scale contracted mechanised cultivation with designated seed or plantlets is now preferable.

Today, plant selection is not only for the yield of active principle, but for the plant’s ability to overcome disease, climatic stress and the hazards caused by mankind. Such methods as in vitro fertilisation, meristem cultures and somatic embryogenesis are used. The transfer of sections of DNA is giving rise to controversy in the case of some end-uses of the plant material.

Some suppliers of plant raw material are now able to certify that they are supplying organically-farmed medicinal plants, herbs and spices. The Economic Union directive (CVO/EU No 2092/91) details the specifications for the obligatory quality controls to be carried out at all stages of production and processing of organic products.

Fascinating plant folklore and ethnopharmacology leads to medicinal potential. Examples are the muscle relaxants based on the arrow poison, curare, from species of Chondrodendron, and the antimalarials derived from species of Cinchona and Artemisia. The methods of detection of pharmacological activity have become increasingly reliable and specific, frequently involving enzymes in bioassays and avoiding the use of laboratory animals. By using bioassay linked fractionation of crude plant juices or extracts, compounds can be specifically targeted which, for example, inhibit blood platelet aggregation, or have antitumour, or antiviral, or any other required activity. With the assistance of robotic devices, all the members of a genus may be readily screened. However, the plant material must be fully authenticated by a specialist.

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The medicinal traditions of ancient civilisations such as those of China and India have a large armamentaria of plants in their pharmacopoeias which are used throughout South East Asia. A similar situation exists in Africa and South America. Thus, a very high percentage of the World’s population relies on medicinal and aromatic plants for their medicine. Western medicine is also responding. Already in Germany all medical practitioners have to pass an examination in phytotherapy before being allowed to practise. It is noticeable that throughout Europe and the USA, medical, pharmacy and health related schools are increasingly offering training in phytotherapy.

Multinational pharmaceutical companies have become less enamoured of the single compound magic bullet cure. The high costs of such ventures and the endless competition from me too compounds from rival companies often discourage the attempt. Independent phytomedicine companies have been very strong in Germany. However, by the end of 1995, eleven (almost all) had been acquired by the multinational pharmaceutical firms, acknowledging the lay public’s growing demand for phytomedicines in the Western World.

The business of dietary supplements in the Western World has expanded from the Health Store to the pharmacy. Alternative medicine includes plant based products. Appropriate measures to ensure the quality, safety and efficacy of these either already exist or are being answered by greater legislative control by such bodies as the Food and Drug Administration of the USA and the recently created European Agency for the Evaluation of Medicinal Products, based in London.

In the USA, the Dietary Supplement and Health Education Act of 1994 recognised the class of phytotherapeutic agents derived from medicinal and aromatic plants. Furthermore, under public pressure, the US Congress set up an Office of Alternative Medicine and this office in 1994 assisted the filing of several Investigational New Drug (IND) applications, required for clinical trials of some Chinese herbal preparations. The significance of these applications was that each Chinese preparation involved several plants and yet was handled as a single IND. A demonstration of the contribution to efficacy, of each ingredient of each plant, was not required. This was a major step forward towards more sensible regulations in regard to phytomedicines.

My thanks are due to the staff of Harwood Academic Publishers who have made this series possible and especially to the volume editors and their chapter contributors for the authoritative information.

Roland Hardman
PREFACE

The study of herbs and alternative systems of medicine has experienced radical changes during the past decade. The developments in organic chemistry during the second world war and afterwards changed the lifestyle of most of the people of the world, and a feeling developed that every ill has a pill. This was not the end; projects for synthetic foods were on the way. These developments not only made the study of herbs and the other natural products outdated in the developed countries, but had a serious effect in the developing world as well, where centuries-old medicinal plant gardens were destroyed and agricultural practices of growing herbs ignored.

With the revival of interest in natural products and the ‘back to nature’ call, researchers started looking into the herbal literature of oriental civilisations, particularly those of India and China, in addition to European and other sources. It was gratifying to find that Harwood Academic Publishers also decided to publish books on the industrial profiles of well-known medicinal and aromatic plants, including those of the Ayurveda.

When Dr R. Hardman, the book series editor and my former guide at the University of Bath (UK), contacted me about writing a book on important Ayurvedic plants, I suggested Neem to him, to which he readily agreed. I was aware of the old medicinal virtues of this tree and the recently discovered insecticidal activity. Quite a number of books and other publications are available on the chemistry and agricultural uses of azadirachtin, the major constituent of neem. But the Western world knew little about the medicinal uses and the recent pharmacological and therapeutic researches into this important tree. I planned the book for these and other, not so well-known aspects. In view of the enormous amount of literature available, I decided to cover all the topics briefly, with a complete bibliography so that the interested reader may find the details from the original source.

I must thank Dr Naresh of the Institute of Microbial Technology (IMTECH), Chandigarh (India) and his staff for library facilities and technical help in the preparation of the manuscript. He also agreed to write the foreword to the book and critically corrected first proof. Ms Vibha provided information on the extraction process of azadirachtin. Dr Pushpinder S. Puri, Corporate Research Director, Air Products Inc. (USA) was helpful in linguistic revision of some parts of the book. The Research Foundation for Science Technology and Natural Resource Policy, India, is thanked for permission to print the poster regarding patent rights on Neem. The computer typing of the whole text, and the layout of the book was only possible with the assistance of my daughter Navneet and some help from my son Avneet. My wife Harminder made important contributions in some places.

H.S. Puri
1. INTRODUCTION

Azadirachta indica, called Neem or Nim in most parts of the world, is one of the very few trees known in the Indian subcontinent since antiquity. During excavation of the site at Mohandjodaro, now in Pakistan, which is as old as 2000 BC (Puri, 1969), in the era of proto-Australoid and proto-Dravidian culture, neem leaves were found. As is evident from Hindu mythology, on their arrival in the Indo-Gangetic plains, the Aryans also attached great importance to neem. They considered this tree to be of divine origin. It was said that Amrita (ambrosia, the elixir of immortality) was being carried by Garuda (demi-god of Hindu mythology which is part human, part bird) to heaven and a few drops of this Amrita fell on the neem tree. In another story, Amrita was sprinkled by Indira (the celestial king) on the earth, which gave rise to the neem tree, yet in another instance neem tree is related to Dhanvantri (the Aryan god of medicine). It was also mentioned that the sun god took refuge in the neem tree to escape from the awesome power of demons (Vijayalakshmi et al., 1995).

The neem tree was considered a gift of God and Sreva roga nivarni (the panacea for all diseases). In an old proverb it was said,

The land where the neem tree abound,
Can death, disease there in be found?

Neem has also been called “Heal all”, “Divine Tree”, “Village Pharmacy” and even “Nature’s drugstore” (Rawat, 1995).

The ancient Indian found many therapeutic uses for the tree and also observed that the tree could survive in very dry and arid conditions. In due course of time, the name and fame of neem spread, not only in the remote areas of the Indian subcontinent but also in the adjoining countries in Asia, now known as Sri Lanka, Malaysia, Indonesia and Thailand. Since ancient times, India has had cultural and commercial relations with the people of these countries.

Whereas in folklore mainly the leaves and to some extent the oil was used in Ayurveda (the Indian system of medicine), Siddha (the system of medicine practiced in some parts of south India) and Unani Tibb (the Greco-Persian system of medicine), polyherbal preparations containing one, two or all five parts of the plant, i.e. leaves, bark, flower, fruit and root, called panchang in Ayurveda, were used. In the traditional systems of medicine, some of the preparations were for internal administration, while others such as nasal drops, medicated oils or fats were for external application.

European colonizers, on their arrival in India in the sixteenth century, also noticed this important tree and they called it Margosa. This term has been widely used in the subsequent literature and until recently neem was called Margosa indica and neem oil was known as margosa oil. European physicians in India, as well as Indian physicians trained in the orthodox system of medicine (allopathy) and in homoeopathy, saw great virtues in the nineteenth century in the bark of neem both from the stem and the root, but mainly stem bark was used, because of its easy accessibility. The bark was considered a substitute for cinchona, widely prescribed for malaria and other fevers at that time. Neem bark was included in the Indian Pharmacopoeia, the
Indian Homoeopathic Pharmacopoeia and even in the British Pharmaceutical Codex. At one time it was in the US National Formulary, but it is doubtful if the source of this drug was neem or the closely allied *Melia azedarach*, also called China berry, with which neem has very often been confused.

Keeping in view the importance of neem in Indian culture, some studies were carried out in the earlier part of the twentieth century to establish the therapeutic efficacy of the various claims made about it in the traditional systems of medicine. The researches showed that neem lacked profound pharmacological activity, which was considered important at that time for a herb to be a source of a drug. Neem was also not found effective against any disease, as compared to the other drugs available at that time. The oil with its foetid odor was not acceptable in any form, even for external application.

During the second world war, because of the scarcity of various raw materials and war needs, research work on the industrial utilization of neem oil started again (Siddiqui and Mitra, 1945a, 1945b, 1945c). These workers filed patents for the pharmaceutical use of neem bitters and for refining the oil. Mitra (1963) published a book on neem, under the auspices of the Oil Technological Institute in India, covering all aspects of information available at that time, which included history, post-harvest technology and processing of the seed to obtain a good quality oil, detailed chemistry and technology of the oil. The main aim of this book was to provide information to the general public on the proper use of neem products, particularly the use of oil for making soap and other industrial products.

Ketkar (1976) in the organisation Neem Mission, tried to popularize neem products, keeping in view the large number of trees growing in India, and the amount of oil and seed cake they can yield. The Neem Mission propagated the idea of making neem soap from the oil at the village level as a small-scale cottage industry, and the utilization of seed cake, left after the extraction of oil, as a manure and as a denitrifying agent for nitrogenous fertilizers. Due to this effort and that of other agencies, neem soap for toilet purposes became a household name in India.

It was well known to Indian farmers that during invasion avoid the neem tree and that it has an antifeedant property. Pradhan and Jotwani (1962, 1968) brought this fact to the notice of the scientific world. Radwaski (1977, 1977a) gave a detailed account of the tree. Thakur *et al.* (1981) reviewed the literature. Slangen and Kerkhoff (1984) found the nitrification inhibitory activity in neem cake. Parmar (1987) gave an overview of neem research and use in India.

The research on neem got a new stimulus, when out of 2000 plants investigated for their action against insects, only neem gave promising results. It was found that it was not only effective against insects but also quite safe for human beings and other warmblooded animals. The active compound was later isolated and identified as azadirachtin.

Azadirachtin attracted the attention of workers all over the world, and various studies were published on it; the most important of these are Warthen (1979, 1989), Marzu (1989), Kraus (1983), Ascher (1987, 1992), Morgan and Mandava (1987), Schmutterer (1987), Schmutterer and Ascher (1987), Siddiqui *et al.* (1988), Jacobson (1989), Ascher and Misner (1989) and Arnason and Philogene (1991). The commercialization of azadirachtin under the trade name Margosan-O and its clearance...
by the Environmental Protection Agency (EPA) of the USA (Larson, 1987) started a new era of non-hazardous insect controlling agents from plants.

In due course of time, neem was found to be a multi-purpose tree (Ahmed and Grainge, 1986), which could be used for the day-to-day activity of human beings (Fig. 1). It was observed that neem could adapt itself to a dry, harsh and hostile climate and degraded soil, particularly in the dry arid regions of the world, where availability of water is quite poor. It could also be planted for soil reclamation (Sastry and Kanathekar, 1990). The tree could provide much-wanted shade to cattle and man in scorching heat and support undergrowth vegetation. The leaves could be used as fodder for ruminants, particularly at times of scarcity. During a recent drought in Gujarat, a west Indian state, a large number of cattle were saved by feeding them neem leaves. It acts as a wind-breaker, an avenue tree, and the dry leaves that fall on the ground provide organic matter for the soil to support vegetation. The wood can be used as fuel, so scarce in arid regions, and also as timber for household furniture, and for agricultural implements. The seed (Axtell and Fairman, 1992) can provide oil for use in household lamps for illumination, as a lubricant for agricultural machinery, against various pests and diseases and for soap. The oil when applied to leather goods prolongs their life and is also useful as a first-aid medicine for healing wounds and skin diseases of man and domestic animals. The seed cake, after washing, can be used in small amounts in poultry and cattle feed. It may be used as such as an organic manure. It not only provides nutrition to plants, but helps in the conservation of nitrogenous fertilizers and the elimination of nematodes.

![Figure 1 Possible uses of neem](image)
International conferences on neem were held in Germany in 1980 and 1983 and in Kenya in 1986. In the proceedings of IUFRO (Salazar, 1990), cultivation of neem as a multi-purpose tree for arid zones was recommended. Koul et al. (1990) and Nat et al. (1991) in their review article dealt with various aspects of the chemical constituents of neem, followed by their chemistry and pharmacological activity. Hedin (1991) edited a book, based on two American Chemical Society symposia on naturally occurring pest regulators, which included neem. Isman et al. (1991) brought to light the various steps taken for the development of neem-based insecticide for Canada.

The book *Neem—a tree for solving global problems* was published by the National Research Council of the USA (Vietmeyer, 1992). Tewari (1992) in his book on neem, in addition to other details, gave an exhaustive account of botanical and forestry aspects, which were not available earlier. Under the auspices of the Indian Society of Tobacco Science, a symposium on the problems and prospects of botanical pesticides in integrated pest management was organized in 1990, in which the role of neem in agriculture was highlighted. In another seminar at Bangalore (India) in 1993, the theme was “Neem and Environment”. Parmar and Singh (1993) also gave an account of the importance of neem in Indian agriculture, while Mordue and Blackwell (1993) presented an update on azadirachtin. Schmutterer and Doll (1993) described an allied species, *Azadirachta excelsa*, as a new source of insecticides. Read (1993) gave an account of the genetic improvement of neem at an international conference.

Randhawa and Parmar (1993) edited the book *Neem Research and Development* for the Society of Pesticide Science, India. Thomsen and Souvannavong (1994) described the activities of the international network on neem. Mariappan (1995) gave an account of neem in the management of crop diseases, while the journal *Indian forester* came out with a special issue, “Neem—gift of the gods” (Khullar, 1995). Schmutterer (1995), who has done extensive research work on the pesticide activities of this tree, edited a book on neem. This book is a compilation of the research carried out by various workers all over the world, on all aspects of this wonderful plant. The book is a very authoritative source of information, and details have been given crop wise, as well as pest wise. An account of research carried out in applied entomology in the tropics is also available (Schmutterer, 1995a). Vijayalakshmi et al. (1995) published a user’s manual for neem for household purposes and for small farmers.

The proceedings of the seminar held in Bangalore (India) in 1993 have been edited by Singh et al. (1995), while Kleeberg (1996) edited the proceedings of a workshop held in Germany. The book has chapters on the control of plant pests, including those found in greenhouses and others such as lice in children, fungicidal activity and the control of nematodes.

The University Grants Commission of India has awarded a book-writing project entitled “Neem—the wonder tree” to Dr M.D.Kharya. The book intends to cover those aspects not covered earlier (personal communication).

The publications mentioned above indicate the importance of this tree in the modern world. Exhaustive information is available now, particularly on the chemistry, pesticide activity and the role of neem in agriculture. Some topics such as history, detailed uses in Indian systems of medicine, botanical studies, mass propagation techniques, use for the preservation of food, pollution prevention, poultry and cattle feed, fertilizer,
soil conservation, pharmacology, etc. have not been touched in most of the earlier publications; the details of these are given here. The well-known aspects like chemistry and use as a pesticide in agriculture have been touched on briefly.

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2. PLANT SOURCES

BOTANICAL NAMES

*Azadirachta indica* A.Juss

*Melia indica* Brandis, *Forest Flora*, 67

CLASSIFICATION

Family: Meliaceae, sub-family: Melioideae, Tribe: Melieae
Genus: *Azadirachta* A.Juss in Mem. Mus. Par. XIX (1830) 220 (Mel.68) 1830-Melia
Species: *indica* A.Juss lice 221, 69-Melia azadirachta

ORIGIN

According to Gamble (1902), the center of origin of *A. indica* is in the forests of Karnataka (south India) or the dried inland forests of Burma (Myanmar). Other authors were of the opinion that this tree originated in the forests of the Shivalik hills (foothills of the western Himalayas) or on east coast of south India. The great variety in the shape of the leaves and other morphological features support the theory of the origin of *A. indica* in upper Myanmar (Schmutterer, 1995); later it became naturalized in the forests of central and western India.

ETHNOBOTANICAL STUDIES

A detailed account of ethnobotanical studies is available in Watt (1889), Dymock *et al.* (1890), Dey (1896), and in many other old publications. In some of the recent botanical surveys in India, some more information has been collected by Billore and Audichya (1978), Tewari and Chaturvedi (1981) and Venkatraghavan and Sundersan (1981). Deka *et al.* (1983) reported it as an Ayurvedic plant from Assam, Sudhakar and Rao (1985) from the upper east Godavari district in Andhra Pradesh, Aminudin *et al.* (1993) from eight districts of Orissa and Singh (1994) documented its use by the tribal Kols of Uttar Pradesh.

In some surveys, stress was laid on finding the cure for certain diseases. Oommachan *et al.* (1990) reported the use of neem for jaundice and Aminudin *et al.* (1993) for malaria.

The use of neem in leather technology at the village level for preservation of leather goods is well known. Pal (1994) noted that it was also used for curing snake skin.
ETYMOLOGY

The present popular name “neem”, also spelled earlier as “nim”, has been derived from the Sanskrit word “nimba” which means sprinkler, which is the short term for “sprinkler of nectar (ambrosia)”. The other Sanskrit synonyms for the tree, as given in the chapter on Ayurveda, refer to its habitat and the use of it in ancient India.

The meaning of the generic name *Azadirachta* does not appear to be interpreted properly in most of the literature. It is often said that it is from the Persian words *azad*—free, and *drakhat*—tree, i.e. free tree, and when the specific name *indica* is added to it, the meaning of the botanical name becomes the free tree from India, which does not convey any specific significance of the name.

The views expressed by Watt (1889) appear to be more convincing, according to which the Persians were well conversant with the allied tree *Melia azedarach* also commonly known as the China berry, but in Persian as “Azadarakhdt” (the corrupted form of it in most of the north Indian languages is Dharek). As discussed with a Persian scholar in Panjab University, India, *aza* means bitter in Persian and *drakhat* means tree, so the name of the China berry in Persian stood for “bitter tree”. When the neem was introduced into Iran, to distinguish it from the China berry, which it resembled to a major extent, neem was called *Aza-drakhat Hindi*, i.e. the bitter tree from India, which led to the present botanical name *Azadirachta indica*.

NAMES IN OTHER LANGUAGES

In the Indian subcontinent, it is largely understood by the name neem or nimb, but in some areas, particularly in the south and east of India, there are regional names for the tree. These as well as the names from other parts of the world are given in Table 1.

RECENT BOTANICAL STUDIES

After the morphological and taxonomic description of neem, under *Melia azadirachta* by Hooker (1872) in *Flora of British India*, no serious efforts were made to investigate its taxonomy. Pennington and Styles (1975), and Pennington (1981) at the New York Botanical Garden, gave the generic monographs of the various members of Meliaceae.


HABITAT

Neem is a large-sized evergreen tree (*Fig. 2*), but younger trees in dry localities may become leafless for a short period, and new leaves may appear in March-April,
which are pinkish green in color. The tree may grow up to a height of 20m and a girth of 2.5m.

(a) Stem

The color of the bark varies according to the part of the plant, its age and locality. The younger branches have a lighter color bark but in a mature trunk it may be grey to greyish black, rough, feebly fissured, and exfoliating. The inner surface of the bark is fibrous and pinkish brown. Small deposits of gum may be present on the stem in some places, but occasionally in some trees, which are quite old and in a humid climate, a fetid sap may be exuded from the trunk.

Bisht et al. (1993) described the branching pattern of a six-year-old tree.

(b) Root

It is normally dicotyledonous in nature, but in more than half of the population, vesicular—arbuscular mycorrhizal (VAM) infection is present due to *Glomus* and *Cigaspora* at 250 cm length. The intensity of infection varies with the availability of water. Neem appears to be a highly mycorrhizal-dependent species. Bala et al. (1989) concluded that a deep-rooted growth habit along with VAM infection may be a survival mechanism when competing for nutrients and water with shallow-rooted and fast-growing plant species. Mohan et al. (1995) carried out a survey of root and rhizosphere soil samples from nursery and plantation. It appears that VAM not only increases the nutrient uptake of the plant but also makes the tree tolerant to root diseases, transplant shock, toxicity of heavy metals and seasonal extremities like

<table>
<thead>
<tr>
<th>Name of the language</th>
<th>Geographical location</th>
<th>Name for neem</th>
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<tbody>
<tr>
<td>Arabic</td>
<td>Middle east</td>
<td><em>Aça darakhul hind</em></td>
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<tr>
<td>Assamese</td>
<td>India (east)</td>
<td><em>Neem gach</em></td>
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<td>Bali</td>
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<td>Bengali</td>
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<td><em>Neem gach</em></td>
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<td>Burmese</td>
<td>Myanmar</td>
<td><em>Thinbau, Tamahin, Kamakha</em></td>
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<tr>
<td>English</td>
<td>Europe</td>
<td><em>Margosa, Indian Lilac Tree</em></td>
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<td>French</td>
<td>Europe</td>
<td><em>Azadran d'Inde, Margosier</em></td>
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<td><em>Indischer Zedrich</em></td>
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<td>Kanad</td>
<td>India (south)</td>
<td><em>Vêpa, Bery, Hebhuw, Kai beru</em></td>
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<td>Konkan</td>
<td>India (south)</td>
<td><em>Bena rooku</em></td>
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<td>Marathi</td>
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<td>India (east)</td>
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<td>Persian</td>
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<td>Portuguese</td>
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<td><em>Amargoesira, Margosa</em></td>
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<td>Sinhali</td>
<td>Sri Lanka</td>
<td><em>Kohunma</em></td>
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<td>Tamil</td>
<td>India (south)</td>
<td><em>Vênhu, Vêppam, Nimhamu</em></td>
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<tr>
<td>Telugu</td>
<td>India (south)</td>
<td><em>Vêpa, Œppa, Nimanuv</em></td>
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drought, etc. It has also been said that *Phytophthora tinnamoni* which is destroying neem worldwide, may be attacking those trees that lack mycorrhiza, as VAM forms a cover on the root and thus protects the plant from pathogens.

(c) Leaves

These are alternate, exstipulate, on a long slender petiole (Fig. 3), dorsal side darker green, ventral light in color, leaves 20–40 cm long, dense at the end of branches, alternate, leaflets 7–15, sometimes up to 17, variable in shape, particularly with respect to the central axis (Fig. 4). The leaves appear smooth but closer examination of young leaves near the shoot apex reveals the presence of resin-secreting glands (Figs. 5A and B). The lower portion of the leaf stalk is covered over with extra floral nectaries.

Leaves are bitter to the taste but Jacob (1941) observed a tree without these principles. Soni et al. (1981) observed the flattening of twigs and crowding of leaves
with prominent ridges and furrows with a disturbed phyllotaxy. The leaflets are 2–7 cm long and 1–4 cm broad, imparipinnate, lanceolate, upper side bigger than the lower but it may vary within a population, often alternate, obliquely falcate, coarsely and bluntly serrate. The breadth of lamina and the degree of dentation on the margin of leaflets vary from locality to locality. In general, leaflets from dry arid areas have narrow lamina and sharp teeth along the sides.

A study of the herbarium at New York Botanical Gardens and Panjab University revealed that the trees from arid areas at the foothills of the Himalayas in general have narrow leaflets with sharp dentation in their serrate margin, as compared to the

Figure 3 *Azadirachta indica*, a small portion of branch
trees from a humid climate which have comparatively broader leaflets with a less sharp serrate margin.

Sarma et al. (1992) studied the leaf architecture in relation to taxonomy. The major venation pattern was pinnate eucamptodemous. The authors have given a key for identification and differentiation of closely allied species on the basis of aréoles, the presence and absence of bundle sheath and the leaf margin.
According to Schmutterer (1995), natural hybrids between *A. indica* and *A. siamensis* are found in upper Myanmar where both species grow together. The shape and consistency of leaflets in *A. indica* in this area is of intermediate type.

**d) Flowering**

Trees growing in areas with a warm winter bloom first, followed by areas where the winter is comparatively cooler. Depending on the locality, flowering may range from January to May (Gupta *et al.*, 1995). Sporadic flowering in September-October has been observed quite often, in addition to that in February-March. Shanthi *et al.* (1996) reported abnormal seedlings in December from these trees and suggested that these trees may be used as germ plasm. In the Murshidabad area of West Bengal (India), the tree flowered throughout the year (Guhabakshi, 1984).

Generally the tree starts flowering at three to five years of age and becomes fully productive at the age of about ten years.

**e) The Flower**

The buds are small, hermaphrodite, numerous, stalked, arranged in long, slender, very lax elongated axillary panicles, shorter than the leaves, bracts minute, deciduous. Flower buds open in the evening and are more scented at night. These buds give rise to 4–5 mm long whitish pink flowers.

*Calyx:* sepals 3 to 5 wide, imbricate, rounded, blunt, ciliate, sepals smooth and thin.

*Corolla:* petals 5, imbricate and oblong, oval in the bud, spreading, spathulate, somewhat twisted with a conduplicate claw, smooth outside, finely pubescent within. According to Gill *et al.* (1993) the number of petals may be 4–8, which is not genetically based.

*Androecium:* stamens 10, situated at the base of a hypogynous disk, the staminal filament combined into a long, cylindrical, erect tube, somewhat dilated below as well as at the top, furrowed and smooth externally, hairy within, terminating above in 10 blunt, thick, recurved trifid lobes, anther smooth erect, closely placed, introse, oblong, two celled. According to Pennington and Style (1975), the pollen grains are 3–4 corporate, prolate-spheroidal or sub-prolate, apocolpium medium, exine smooth, slightly thickened at the aperture.

*Gynoecium:* carpels, 3–5, syncarpous, superior, as many locule with 2 ovules in each loculus, style about the length of the staminal tube, stigma 5 lobed, placentation parietal (Nair, 1956). Garudamma (1956) and Rangaswamy and Promila (1972) have studied the embryology in detail. The ovary is trilocular at the base, becoming unilocular at the ovule-bearing region. The pollen tube is monosiphonous, and enters the ovule through a micropyle. One of the synergids is destroyed. Syngamy precedes triple fusion, resulting in an enlarged zygote. Twin embryos occur commonly (Nair and Kanta, 1961); the number of seed per embryo may be 1–3 (Gill *et al.*, 1993; Vijayan and Rehill, 1987). The gametophyte develops in the usual way; the embryo sac is of Polygonum type. Study of premature fallen fruits indicated that embryo abortion is common (Gill *et al.*, 1993).
Pollination
The flowers are cross-pollinated in general, in spite of bisexual flowers and the absence of self-incompatibility. Pollination is occasionally entomophilous but usually anemophilous.

Seed Development
A study of seed development has shown a steady increase in fruit/seed length, breadth, fresh and dry weight up to 12 weeks. The moisture content after this period started declining, with an increase in seed oil, protein and carbohydrates (Sivasamy and Karivaratharyu, 1993).

Fruit
It is an ovoid drupe, bluntly pointed, 1–2cm long, when young and unripe smooth and green with white milky juice, yellow to brown when ripe, epicarp thin, mesocarp with scanty mucilaginous sweetish pulp, endocarp hard enclosing the seed. The fruit gets darker in color and wrinkled on maturity. Variability in seed size among different provenance was studied by Sindhu Veerendra (1995). The seed length varied between 11 and 18mm, width 4.5–8.5mm, and weight 100–530 mg. Among a three-year-old population, Gupta et al. (1995) observed that 92.3 percent of the trees had 1–100 fruits/tree, 4.36 percent 101–200 fruits/tree and 0.49 percent (one only) had more than 400 seed/tree.

Seed Dispersal
Most of the seed fall on the ground under the tree, where at that time the soil is water logged or there may be rain streams. The fruit may remain in moist conditions under the tree or occasionally may travel some distance with rain water. Since there is no dormancy, most of the seed may germinate immediately, but perish because of lack of conditions for further seedling growth. Occasionally, some fruits are swallowed by birds for their sweet pulp and the seeds are passed out of the body, undigested, because of the hard endocarp. The seeds so dropped are far away from the trees; if they germinate, the seedlings have much better chances of surviving and producing plants, as compared to undispersed seed. Gupta et al. (1995) studied the time of flowering and fruiting in a provenance trial of a neem population of three-year-old trees. In this population, 25.22 percent of trees exhibited fruits and 3.7 percent flowering, but in 0.25 percent, both flowering and the maturity of the fruits occurred at the same time.

Seedling Morphology
Germination is epigeal (Troup, 1921). Deb and Paria (1986) gave an account of seedling morphology. The fresh mature seed, if in humid conditions, may start germinating within a day or so, but fully dry seed may germinate from the 7th day of sowing and complete it in 25 days. Cotyledons are plano-convex, sub-opposite, lowest one sessile, blade obovate-oblong, stem erect. A few seedlings have an alternate cotyledon; in these, the lower cotyledon has a unilacunar two-trace node and the other a trilacunar
three-trace node. In those cases where cotyledons are opposite, both cotyledons have a unilacunar two-trace or trilacunar three-trace condition (Bansal and Pillai, 1986). During the further course of development, the cotyledon along with the endocarp is pushed above the soil because of the elongation of the hypocotyl in the lower region. The plumule emerging from the cotyledon dislodges the endocarp. The top of the seedling at this stage is green, glabrous or with minute odorless glands. These glands are common in younger leaflets but become fewer in number as the leaf matures. The phenomenon of twin seedlings has also been observed, which may be as high as 11.27 percent (Vijayan and Rehill, 1987; Gurudev Singh et al., 1995). This may be due to the development of one or more than one ovules, out of five of the ovaries (pentacarpellary), giving rise to more than one seed under the same endocarp, but according to Pushpakar and Babekey (1995) it is due to polyembryony, with a frequency of 1 out of 800.

Three mutants with white hypocotyles and leaves in a progeny of 1220 from a single tree have been observed by Kulkarni and Srimathi (1986, 1987). These seedlings survived for one month.

CYTOLOGY

The genetic chromosome number is n-14 and somatic 2n-28, 2n-30 have also been reported from the root tip mitosis (Pathak and Singh, 1949; Mukherji, 1952; Stylos and Vosa, 1971; Mehra et al., 1972). According to Gill et al. (1993), this species has small-sized chromosomes but a high chromosome number. The loss in combination due to a low chiasmata index is compensated by an increase in the number of linkage groups and the allogamous nature. The open system of genetic recombination is also operating.

OTHER SPECIES

Two more species of Azadirachta, i.e. A., excelsa and A., siamensis have been recognized (Schmutterer, 1995).

(a) A. excelsa (Jack) Jacobs (A. integrifoliola Merr.), sometimes erroneously spelled as A. integrifolia is also called marrango or the Philippine neem tree. It was described as Melia excelsa Merill, and later as A. integrifoliola. A. excelsa is distinguished from A. indica by its entire leaflets, its panicle much longer than the leaves, and by its long flowers. Whereas A. indica thrives in hot dry regions, A. excelsa is a plant of lowland monsoon forests, and tolerates greater rainfall (Schmutterer and Doll, 1993). It is an endangered species because of over-exploitation in the Philippines.

(b) A. siamensis Val. is also called the Thai neem tree. It was identified earlier as A. indica var. siamensis, but Schmutterer (1995) recognized it as a distinct species, on the basis of a number of morphological, anatomical, histological, phytochemical and host infestation characters. In some areas, hybridization has occurred between A. indica and A. siamensis. The black neem reported from Thailand is probably a mutant of the Thai neem or a cross between the Thai and Indian neem (Schmutterer, 1995).
ANATOMICAL STUDIES

Both leaves and bark are medicinal, so have been subjected to detailed pharmacognostic studies, which mainly dealt with anatomical features. Wood as a source of timber has also been studied in detail by Brandis (1906), and recently by Tewari (1992). Some information about it has been provided in Chapter 7 on neem as a source of timber and fuel, in this book.

Naryana Aiyer et al. (1957), while dealing with the pharmacognostic studies of Ayurvedic medicinal plants, described both stem and root bark. Pandey (1969) presented the cuticular character of the leaf surface, while Farooqui (1981) gave an account of the epidermal structure. Purushothman et al. (1988) touched on both pharmacognostic and chemotaxonomic aspects of the bark, and compared it with the closely allied species Melia azedarach. Sarkar and Datta (1989) gave a morphological and anatomical account. Patel (1977) gave the diagnostic key for identification of bark powder, while Bagchi et al. (1992) studied the morphology of calcium oxalate crystal in neem and the other barks to distinguish them from each other.

Secretory Cells

Neem terpenes are present in all parts of living tissues but are most abundant in the seed kernel. The site of synthesis and accumulation of these has been identified as secretory cells. The study of secretory cells is thus of significance, because these accumulate triterpenoids. A new isoprenylated flavone was isolated from the resin gland and its structure was characterized by Balasubramanian (1993a).

Shah (1983), Inamdar et al. (1986), Arumugasamy et al. (1993) and Balasubramanian et al. (1993) gave an account of gum and resin secretions. Dayanandan et al. (1993), on the basis of microscopic and histochemical studies, established the following secretory cells:

1. Secretory cells that accumulate triterpenoids
2. Glandular trichomes, which secrete resin and protect young leaves
3. Gum ducts
4. Lacticifers which develop only in pericarp. These are articulate, and contain milky white latex.
5. Extrafloral nectaries

The secretory cells are found in the parenchymatous cells of the cortex and the pith of the stem, the petiole and cortex of the root, between the palisade and spongy tissue in leaves and all over the cotyledons. These cells are oval in shape and may contain pale orange brown vesicles, which are age dependent as they become larger in the older cotyledons.

STUDIES ON GUM FORMATION

The neem tree yields a meager amount of gum, only in humid areas. Gum resin in many other trees is induced by a primitive method of acid treatment, which in many cases kills
the plant by dehydration. Nair et al. (1980), Nair and Shah (1983) and Nair et al. (1985) tried ethephon and paraquat for this purpose. Ethephon administered to bark increased gum production by 100 percent. Histological and histochemical studies showed the disappearance of starch grains in the ethephon-treated plant, accompanied by the breakdown of cells and mass dissolution. The degradation products of all these tissues lead to gum formation. In heartwood treated with paraquat, gum ducts were observed, along with desiccation, the disappearance of starch grains and the accumulation of lipid insoluble polysaccharides and phenolics. When gum cavities were induced in sapwood by drilling holes, and treated with ethephon or paraquat, abundant high protein gum was produced after seven days in schizo-lysigenous cavities.

Nair (1988) studied wood anatomy and heartwood formation.

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3. CHEMICAL CONSTITUENTS

INTRODUCTION

The earlier Indian chemists concentrated on the bitter principles of neem oil. Sen and Banerjee (1931) gave an account of these. Siddiqui (1942) isolated the crystalline bitter compound nimbin, and later developed a method for separation of these bitters (Siddiqui, 1945a,b,c). Mitra (1963) described all these activities in his book on neem. According to Govindachari (1992), after the work on nimbin, interest in neem constituents became dormant except for the isolation of two new compounds, vilasinin and vepenin. With the discovery of the activity of the neem compound that suppressed feeding in locusts (Butterworth and Morgan, 1968), the detailed chemistry of these received a new stimulus. Very exhaustive studies were undertaken, particularly on the known active compound azadirachtin (Figure 6). Some of the important publications in this direction were by Lavie et al. (1971), Connolly (1983), Kraus (1983), Kraus et al. (1985, 1987), Morgan and Mandava (1987), Siddiqui et al. (1988) and Balandrin et al. (1988). Rastogi and Mehotra (1990, 1991, 1995) in the Compendium of Indian Medicinal Plants in three volumes gave various stages of this study. In their monographic account of neem, Koul et al. (1990), Nat et al. (1991), Tewari (1992) and Govindachari (1992) gave a brief summary of the salient points of research on the chemistry of neem. Schmutterer (1995) has also given the relevant account of compounds which gave promising pesticidal results. Broadly speaking, the following types of compounds have been reported.

MAJOR CONSTITUENTS

Terpenoid constituents

(A) Protolimonoids
(B) Limonoids
(C) Pentatriterpenoids
(D) Hexatriterpenoids

Non terpenoid constituents

(A) Hydrocarbons
(B) Fatty acid
(C) Steroids
(D) Phenols
(E) Flavonoids
(F) Other compounds

Keeping in view the exhaustive literature on the chemistry of neem, for the present purposes only a brief summary of the research on chemistry has been given. The interested reader may get detailed information from the references given above.
SEED OIL

Dasa Rao and Seshadri (1942) and Child and Nathanael (1944) gave the fatty oil composition of oil. Skellon et al. (1962) and Mitra (1963) described the other compounds isolated from neem, most of which are now of historical importance. The important ones of these were nimbin mp 192° amorphous (1.2–1.6 percent) and nimbidin mp 90°. Hegnauer (1969) described sulphurous compounds. The most important compound isolated later on was azadirachtin.

AZADIRACHTINS

Butterworth and Morgan (1968) isolated azadirachtin (Figure 6), the active constituent, in crystalline form. The process of isolation included chromatographic fractionation of neem kernel extract, monitored by antifeedant assay with the desert locust, *Schistocerca gregaria*. Butterworth et al. (1972) established the molecular formula for azadirachtin which was further modified by Kraus et al. (1985).

Remboldt et al. (1987b) determined the structure of azadirachtin B. The relationship between the structure and biological activity of azadirachtin A and B was established (Remboldt et al., 1987a; Remboldt, 1989). Yamasaki and Klocke (1987) prepared eight derivatives of azadirachtin and bio-assayed them for their growth-inhibiting activity. The results showed that a hydroxyl group in azadirachtin was essential for the activity but for maximum action a lipophilic region is required. Broughton et al. (1986) also described the chemical structure. Barnby et al. (1989) and Remboldt (1989) studied the mode of action of the isomers of azadirachtin by exposing them to various wavelengths of ultra-violet radiation. These isomers were then analyzed for structural degradation and loss of biological activity. Further details on the pesticidal constituents can be had from Mordue and Blakewell (1993), Remboldt et al. (1993), Kobeleswaran et al. (1994) and Ley (1994).

Estimation of azadirachtin

Warthen et al. (1984) described a method of estimating azadirachtin in neem extract and formulations. Yamasaki et al. (1986) developed a rapid and inexpensive method

![Figure 6 Structure of azadirachtin](image.png)
for the isolation and purification of azadirachtin by using Flash Chromatography and High Performance Liquid Chromatography (HPLC). Schroeder and Nakanishi (1987) and Schneider and Ermel (1987) simplified the method of quantitative determination of azadirachtin. Ermel et al. (1987) determined this compound from various geographical regions of the world, and found a large variation in different countries. The highest concentration was from Nicaragua and Indonesia (4.7 percent) and the lowest from Sudan and Nigeria (1.5–1.9 per cent). Govindachari et al. (1990) developed a method of direct preparative HPLC without resorting to column Chromatography. Govindachari et al. (1991, 1992, 1994) improved this method further to isolate azadirachtin A, B, D, H and I. The last two isomers were isolated for the first time and their structure was determined. Azadirachtin can also be estimated in other natural products (Sundaram and Curry, 1993) and in commercial preparation, where it is emulsified with surfactants, by a method of Azam et al. (1995).

Isolation of azadirachtins

On a small scale, azadirachtin may be isolated from the seed by the method given in Figure 7. The seed, after extraction with hexane, yields oil and a paste. This paste is first extracted with ethanol and then with chloroform to get crude azadirachtin, which may be about 20 gm from one kg of seed in some samples from India, but the yield varies from sample to sample.

Variation in azadirachtin in different samples

Twenty-one samples of neem seed, representing eight agro-ecological zones of India, were tested for azadirachtin and biological activity (Ketkar and Ketkar, 1993). In another study, azadirachtin varied from 0.05 to 4.24 percent in seed (Rengasamy et al., 1993). Azadirachtin was also studied at different stages of flowering and fruiting by Rangaswamy and Parmar (1994); it was not detectable until forty days after anthesis but was present in green and yellow fruit. Yakkundi et al. (1995) detected azadirachtin in nine-week-old developing fruit; it started increasing with age and was at its maximum after the 19th week, when fruit started changing colour from green to yellow. Sidu and Behl (1996) studied seasonal variations in azadirachtins in seed in phenotypes that produced seed at the normal time in July-August (monsoon rains) and again in November-December. Monsoon seed yield 1.53 percent of azadirachtin-rich fraction as compared to winter seed (1.26 percent). Azadirachtin A was the major metabolite in the rainy season seed. Azadirachtin A and B were in equal proportion in winter seed. Concentration of azadirachtin F increased more than twofold in winter. Winter stress appeared to favor synthesis of azadirachtin B and F.

OTHER COMPOUNDS

Another important compound, salannin, has been studied by Henderson et al. (1968) and nimbin by Harris et al. (1968), who also described their stereochemistry. Gedunin, which has been found effective against the malarial parasite, was isolated by Lavie et al.
Yamasaki et al. (1988) described a method of isolation and purification of salannin which is 0.95 percent in seed oil.

CONSTITUENTS IN DIFFERENT ORGANS

Oil and seed cake

Skellon et al. (1962) described fatty acid of neem oil. Thakur and Godrej (1972) patented a method of purifying the oil. Keeping in view the importance of oil and oil cake, the Indian Standards Institute gave the specifications for both of these

Leaves

The smell of the leaves is due to essential oil (0.13 percent), as reported by Dakshinamurty (1954). The leaves have been investigated for other phytoconstituents also, the details of which can be had from Basak and Chakraborty (1968), Awasthi and Mitra (1971), Tirimanne (1984), Pant et al. (1988), Vashi and Patel (1988), and Katani and Padhya (1988).

Flower

The essential oil (0.025 percent) distilled from air-dried blossoms was reported by Subramanian and Rangaswamy (1947) to contain, besides tetrasulphides, kaempferol, thioamyl alcohol, benzyl alcohol, benzyl acetate and an unidentified alcohol.

Bark

Bitter principles, essential oils, and other constituents have been reported from bark, but the important compounds are polysaccharides, which are water soluble. These anti-inflammatory polysaccharides consist of glucose, arabinose and fructose in a molar ratio of 1:1:1 (Fujiwara, 1982, 1984). Subramanian and Lakshmanan (1993) reported the other constituents.

Heartwood

The wood is composed of cellulose, β-cellulose, hemicellulose A, B and lignin (Wealth of India, 1948). Bhola Nath (1955) chemically examined the heartwood.

Gum

It is amber-coloured, non-bitter and gets blackened with age; it is water soluble and resembles gum acacia in some physical properties. The gum was investigated in detail by Pattabiraman et al. (1968). The gum is rich in protein. Bajpai et al. (1970) isolated pure aldobiuronic acid and aldotriuronic acids. The other studies were carried out by Anderson and Hendric (1971) and Andersen et al. (1972, 1986), who gave the amino acid and amino sugar composition.

Sap (stem exudate)

The sap has a strong smell of fermented liquor. It is slightly sweet and contains many amino acids (Wealth of India, 1948). Ali and Qadry (1988) investigated it in detail, and identified sugars (fructose, mannose and xylose), acids (citric, malonic, succinic, fumaric, and acetic), steroids (β-sitosterol and methylenecycloartenol), limonoids (nimbin, azadirone, and gedunin), free amino acid (aminobutyric acid, glycine and a minor amount of argenine, glutamine, lysine and threonine) and crude proteins 3.57g/100ml.
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4. CULTIVATION

INTRODUCTION

Soil in many parts of the world has been degraded because of over-exploitation by man and deforestation. Neem has been found to be a tree of choice for afforestation. It has been found suitable for arid regions and for large-scale cultivation (Radwansky and Wickens, 1981; Chaturvedi, 1985) and the agroforestry of neem has been given a new term, “Margoculture”. Cultivation of neem has been recommended for the following reasons:

1. Multifarious uses: leaves for fodder (Nehra et al., 1987), wood for timber and fuel, and seed for oil, manure and pesticide.
2. Tolerance of tree to high temperatures, aridity and high concentration of salts in the soil.
3. Avenue tree, which also acts as a wind-breaker (Oboho and Nwoboshi, 1991), shelter belt, canal-side plantation and sand dune stabilizer.
4. Leaf litter enriches the soil by providing organic matter without any harm to friendly insects, earthworms etc.
5. It adapts itself very well to degraded and acid soil and can be used as a tree for reclamation of wasteland (Sastry and Kanathekar, 1990).

Because of the above, margoculture has been included in agroforestry projects in most of the developing countries of the world with dry and arid zones. In developed countries, wherever possible, it is being cultivated as a raw material for harmless pesticides which can be obtained from the seed. Scientific studies are being conducted for methods of mass propagation, ideal conditions for growth and development, and also for its impact on the eco-balance, environment and agricultural crops. Werner and Muller (1990) described it as a fast-growing tree. Ketkar (1982) gave the preparation and uses of its products and by-products. Randhawa and Parmar (1993) gave an account of research on neem.

METHODS OF CULTIVATION

From Seed

Neem seed, in general, loses its germination capacity after a couple of weeks of maturity. Efforts have been made to find out the reasons for this small period of viability and how the germination period of seed can be extended by modifying storage conditions (Jha and Chaudhuri, 1995).
Reasons for Loss of Germination Capacity

Various views have been expressed; some of the important ones are:

1. Fermentation

It has been postulated that in the kernel enclosed in the endocarp, a breakdown of sulfur-containing compounds takes place, causing damage to seed. A viable seed has a green cotyledon, but it turns brown after a couple of days in hot and humid conditions (Smith, 1939). The dry viable seed, when immersed in water, emits a garlic-like smell, which is not present in non-viable seed, indicating loss of some sulfur-containing compounds.

2. High Temperature

Seed stored at low temperature remain viable for a longer period, indicating thermal sensitivity of the embryo. It has been postulated that high temperature increases the metabolic rate of the seed embryo, causing its death.

   Seed of the same batch, stored in India at an average room temperature of 30°C with low humidity and in Florida (USA) at an average room temperature of 20°C with high humidity, when studied after couple of months were found to have a green cotyledon in most of the seed in Florida, but the seed cotyledons had turned brown in India, with most of the seed full of fungal spores.

3. Dm to Pulp

Seed without pulp give a much better germination percentage, as compared to those with pulp (Prasad, 1941). The fruit swallowed by birds, for its sweetish mucilage, gets de-pulped in the alimentary canal of the birds and, when dropped, these seed have a higher germination rate. It has been postulated that the germination percentage in seed with pulp decreases because of the meager amount of moisture present. This amount of moisture is sufficient to start the early stages of embryo development but cannot sustain it until completion of germination, thus causing death of the embryo in the early stages, resulting in loss of germination capacity.

4. Endocarp

It has been known to form a water- and air-tight compartment. If the seed are dried at high temperature, as has been observed by Puri and Hardman (1980) in *Trigonella foenum graecum*, they become dormant and are said to be “hard”. These seed fail to germinate until the endocarp is punctured by some physical or chemical means. A similar phenomenon may be operating in this case also. It has been seen that in neem, in those cases where the endocarp was removed, the seed showed a better germination.

5. Inhibitory Compounds

These may be present in the seed coat or in pulp or in the endocarp and may not allow the seed to germinate (Ponnuswamy *et al.*, 1993).
6. Biochemical and Physiological Carriers

There may be some peculiar mechanism operating in the embryo which does not allow further development after a fixed period of time, causing loss of germination capacity.

Storage and Other Conditions for Germination

Chaisurisri et al. (1986) dried seed at 25°C in an air-conditioned room and in a glasshouse under other conditions. The most effective treatment was drying the fresh seed in sunlight for 3 days at 46.16 percent moisture before storing them in cotton bags at 15°C. The seed stored in this way retained their viability for more than four months, giving 62 percent germination. Nagaveni et al. (1987) collected greenish yellow seed, when they were not fully ripe, de-pulped them and dried them in shade for 2 days. A germination rate of up to 80 percent could be achieved from these, even after four months. After six to seven months germination decreased to 50 percent. Maithani et al. (1989) studied the effect of fruit maturity, temperature and the container on germination. The maximum germination was found from seed 10–12 weeks old after flowering, when fruit turned yellow. Seed stored at 15°C or 50°C in sealed or perforated plastic bags, cardboard boxes, etc., deteriorated, while those stored in aerated containers or at 15°C, showed about 15 percent germination after six months. Venkatesh et al. (1990) also obtained good germination by storing de-pulped seed in cotton bags for up to four months, but after that there was a decline.

Chaney and Knudson (1988) and Radhamani et al. (1990) found that removal of the endocarp improves germination considerably. The authors postulated that the endocarp develops a physical barrier for water, gases, enzymes and inhibitors and for metabolism of fats.

Ponnuswamy et al. (1991) graded the de-pulped seed on the basis of their behavior when immersed in water. Those remaining on the surface were labeled “floater” and the others “sinker”. The floaters were 18 percent of the total and had low viability as compared to sinkers which had 90 percent germination. After 3 months of storage, the germination rate dropped to 15 percent. Seed stored in earthen pots and buried in moist sand recorded 62 percent germination even after 3 months of storage.

Aflatoxin may be one of the causes for lowering the quality of seed when stored in an enclosed atmosphere at high temperature and humidity. Chourasia and Roy (1991) studied the effect of temperature, light and humidity on the storage of neem seed and the production of aflatoxin Bl. The highest production of this compound was recorded at 30°C and relative humidity of 96 percent.

Nursery Techniques

Seed after harvesting should be immediately de-pulped, washed with water and dried in shade with a free flow of water, to such an extent that the outer surface of the seed coat should not be wet to the touch. These seed can be sown in nursery beds, about 2.5 cm deep and at a distance of 2.5 cm each. The soil should be loose and sprinkled with water, but there should be no water logging. Seed can be sown in small containers (flower pots) or in thick plastic bags filled with soil, at the rate of 2–3 seed per
container. In the majority of cases, the seed germinate within couple of days, but some seed may take a longer time. When the seedlings are about 2cm high, they can be transplanted from the nursery beds to bigger beds at a distance of 15 cm×15 cm. In cases where more than one seed is sown in pots/bags, the most robust seedling should be retained, the rest pulled out and discarded.

According to Vijayalakshmi et al. (1995), raised beds of 10×1 m in area and 15cm in height should be prepared. Farmyard manure, sand and local soil should be mixed in the ratio of 1:1:3. This mixture should be put on top of soil to a height of 2.5–5 cm. For transplanting, the bag should be filled with soil containing silt, sand, clay and farmyard manure in the ratio of 1:1:1:1. The plastic bags should be of 150–200 gauge thick so that they have enough mechanical strength and they do not burst during shipping.

In areas where there is a possibility of attack by insects on young seedlings or by ants and termites and the environmental conditions are not ideal, the nursery seedlings can be raised in trays, placed on a platform in a greenhouse (Figure 8). A greenhouse can be economically erected by making a frame of water pipes covered over with thick transparent plastic sheet. On one side exhaust fans or a big blower should be installed so that there is free movement of air. Water can be sprinkled with sprinkler-attached pipes inside the greenhouse or by hand sprinkler. This type of improvised

![Figure 8](image_url)

**Figure 8** A simple green house for young seedlings Abbreviations: BL=blower, DR=door, FP=flower pot, PI=pipes for supporting the structure, SL=transparent plastic sheet to cover the greenhouse, SP=sprinkler, ST=seed tray, RP=concrete stand
greenhouse is very useful in areas with high humidity, heavy rainfall and frost. Water mist has to be created inside, otherwise the seedlings get dehydrated by the greenhouse effect, which is evident by the development of pink pigment in the leaves and hypocotyl, and the stunted growth of the seedlings.

Further details about nursery techniques can be obtained from Maithani et al. (1988), who have given the method of sowing neem in moist tropical climatic conditions. Joshi and Prakash (1992) studied extracts of some trees on germination and seedling growth. The seedlings are ready for transplanting in pits when 6 months old and 15–25 cm in height. The pits of 30x30x30 cm should be dug at intervals of 3x3 m, preferably before rain, and watered frequently.

Direct Sowing

For degraded areas, direct sowing has been recommended by Tewari (1992), provided there is protection and enough moisture during the initial stages. This may be done by dibbling in bushes, by making small holes in a bush like Euphorbia, and then sowing 2–3 seed and sealing the hole. Broadcast sowing can be done on ploughed or unploughed soil, while sowing in lines can be done along the ploughed fields of crops. In heavy soils, sowing on mounds or ridges was practiced, whereas in dry sites trenches were preferred for retaining the moisture. Contour ditches have also been tried in some areas (Nagaraju Kumar et al., 1992).

Vegetative Propagation

Neem seed has a very small period of viability and various methods have been developed to prolong it, which have been discussed earlier. Bhardwaj and Gurdev Chand (1995) suggested cryogenic cold storage. Even if seed germination is achieved, a plantation sown from seed may have a lot of variability. Vegetative propagation may be helpful because of early maturity, true-to-type stock and uniformity of the growth characteristics of the plantation. Good mother trees can be selected on the basis of the desired characteristics such as good vigor, disease resistance, age, better crown shape, well distributed branches, and big, abundant seed rich in azadirachtin. Jha and Chaudhari (1990) carried out trials on stump planting and Swaminathan and Surendram (1989) carried out studies on rooting response to growth regulators. For rapid multiplication of the elite tree, Mohinderpal (1995) prescribed the following methods:

1. **Grafting and budding**: Early summer is the best season. Wedge grafting and patch budding techniques gave good results. Pong-Anant et al. (1989) gave further details.

2. **Air layering**: One cm wide bark was cut by Shanmungavelu (1967) and the exposed tissues were treated with indole butyric acid (IBA) or 1 naphthalene acetic acid (NAA) and covered over by moist moss or coconut fibers. Only those air layers that were treated with 0.1 percent NAA and IBA produced roots.

3. **Rooting branch cuttings**: Hardwood, semi-hardwood, softwood and juvenile cuttings are used:
   (a) juvenile root suckers were planted in a moist layer of sandy soil (Rao, 1958; Pal et al., 1994).
(b) **Hardwood:** Cuttings were treated with IBA and planted in a high humidity area. Sivaganam *et al.*, (1989) observed that these cuttings rooted under mist in 135 days with 1000 ppm of IAA and IBA. The treatment, given by a drip method, stimulated rootings.

(c) **Semi-hardwood:** Cuttings treated as above but for a short duration.

(d) **Softwood cuttings:** In this case, long binodal leafy cuttings are taken. Pal *et al.* (1992) stimulated the root response by treatment with phenolic compounds like naphthol and salicyclic acid, under mist. Verma *et al.* (1996) experimented with the effect of auxin on rootings of cuttings in the spring season by applying IBA and NAA. The 100 ppm of both the hormones significantly increased rooting and sprouting but IBA was more effective. Chander *et al.* (1996) studied the effect of auxins on the number of leaves per cutting and the length/age of the main branch. The cuttings, taken from a 8–10-year-old tree and treated with 500 ppm IBA and 2000 ppm of IAA, resulted in maximum increase in the number of branches and leaves.

**BIOTECHNOLOGY AND TISSUE CULTURE**

Tissue culture techniques are an excellent tool in neem research to obtain organogenesis and plant regeneration in a short time under controlled conditions. By using these techniques, we are not dependent on the germinability of the seed, which is of very short duration, and will be able to obtain pesticidal constituents which could not be synthesized so far (Tewari, 1992), or whose synthesis may not be economically viable.

Work on this aspect has been carried out for the past several years. Sanyal *et al.* (1981) studied *in vitro* hormone-induced chemical and histological differentiation in stem culture, while Ekundayo (1983) gave an account of the biosynthesis of nimbolides from (2–14C) mevalonate and (2–14C) acetate. Schulz (1984) obtained callus proliferation from various parts of the plant by improving the culture media. Pratap and Jaiswal (1985) tried plantlet regeneration for leaflet callus from 15-year-old trees. Data were collected on percentage root and shoot production under all treatments. Root formation from calli occurred wherever 1 naphthalene acetic acid (NAA) was used, whereas addition of a low concentration of NAA to a medium containing benzyladenine increased the frequency of shoot formation. The callus-derived shoots produced roots and developed into plantlets. Sarkar and Datta (1986) established the relationship between biosynthesis of nimbin and β-sitosterol in bark and bark-originated callus of increasing age and studied the effect of glycine on *in vitro* biosynthesis of nimbin and β-sitosterol in tissues. Sanyal *et al.* (1988) found that the cotyledons contained nimbin, glycine, other amino acids and β-sitosterol. The effect of glycine on the synthesis of nimbin and β sitosterol was investigated and it was observed that glycine affected the synthesis of both. Ramesh Kumar and Padhya (1988) isolated nimbin from the leaves and its callus culture growth. Indole acetic acid (IAA) and indole butyric acid (IBA) interaction showed a linear increase in nimbin content. Rao *et al.* (1988) also studied the growth of callus. For genetic transformation, Naina *et al.* (1989) induced tumors in the culture using *Agrobacterium tumefaciens*. The transformed plantlets were regenerated from the infected region of the seedling and also from tumor cells.
Ramesh Kumar and Padhya (1990) propagated plantlets from leaf disks. These workers (Ramesh Kumar and Padhya, 1993) worked further on this aspect and obtained 18–20 adventitious shoot buds within four weeks by variously treating epidermal cells, which gave rise to 12–15 plantlets within six months. These were later transferred to the plots, where stable lines were developed from them.

Gautam et al. (1993) studied the development of root and shoot in anther-derived callus. The callus originated in the microsporangial wall layer and connective tissue of anthers. After 13–15 weeks, a green nodular structure and prominent roots developed. A different culture medium induced multiple shoots. Nirmala Kumari et al. (1993) observed that the growth physiology of the basal region of hypocotyl appears to be fundamentally different from other shoot regions and holds good potential for multiplication through cuttings of seedlings and tissue culture propagation. The various explants differentiated for embryoid formation and produced leafy shoots. Stephen et al. (1993) produced callus without further differentiation from basal hypocotyl segments. Similar callus at the cut surface of the cotyledons produced roots. In a one-month-old basal hypocotyl segment, several lenticels were produced which gave rise to shoot buds. Joarder et al. (1993) cultured nucellus tissue of immature seed after fertilization. Five to seven plants were regenerated per culture. Immature cotyledons of the age group of 60–70 days were found to be the best for embryogenesis, but the majority of the cultures produced embryoids which did not germinate. Drew (1993) developed a technique for micro cuttings in vitro for clonal propagation of neem lines with a high yield of azadirachtin and free of genetic off-types.

According to Thiagarajan and Murali (1994), the optimum conditions for high frequency regeneration of plantlets can be obtained from a medium containing 3 percent sucrose supplemented with 0.1 mg/L NAA and 0.3 mg/L of 6-benzoylamino purine. The regenerated plants were successfully established outdoors. Allan et al. (1994) cultured embryos from fruits, collected thirty days after pollination. The culture yielded 0.0007 percent of azadirachtin on a dry weight basis. Newton et al. (1995) described somatic embryogenesis, embryogenie culture and culture maintenance in neem along with other woody plants.

Peroxidases isoenzyme of organogenesis were used by Preetha et al. (1995) as biochemical markers for differentiating regenerative callus lines in the early stages of differentiation. These authors gave the methods of callus induction, sample preparation and electrophoresis. The isoenzyme characters of the various types of cultures were discussed in detail. Clonal propagation of neem from non-woody explants was achieved by Joshi and Thengane (1996) by supplementing an MS medium with benzyladenine and kinetin. Two to three shoots per explant were observed after twenty days. The multiplication frequency of shoots per explant and the average shoot length were observed after 14–16 weeks. Rootings from these shoots developed, and regenerated plantlets were established in the soil.

**GENETIC IMPROVEMENT**

Neem is a heterogeneous group showing a great amplitude of variation. The wide distribution of neem and its growth under a variety of climatic conditions amply
demonstrate the genetic diversity present within the species, which can be utilised for genetic improvement, but traditional methods of breeding, including hybridization, can be of very limited use owing to their long generation time and the prevalence of outbreeding. The other methods which can be utilized for the above purpose, as suggested by Tewari (1992), are:

1. Selection of plus tree from variations within the wide population. The international neem network has also recommended investigation on the evaluation of genetic variability through provenance trials. A number of studies have been conducted in India in this direction.
2. Germplasm bank.
3. Seed production for mass multiplication for a selected superior clone.
4. Mutation breeding.
5. Polyploid breeding.
6. Modern techniques: biotechnology, genetic engineering, tissue culture, callus formation, somatic embryogenesis, direct multiple shoot induction from seedling explant, genetic transformation by *Agrobacterium tumefaciens*, which infects a large number of plant species and causes tumor formation, etc. may be utilized.

Chinnamani (1993) and Read and French (1993) suggested a strategy for neem improvement.

DISTRIBUTION IN INDIA

In the literature it is often repeated that this tree is distributed all over the Indian subcontinent, but it is a general statement. The tree does not prefer silty, saline, mica-rich or inundated soil. It cannot bear frost or high humidity or water-logged conditions. It is light demanding and is often cultivated near human habitation for its multifarious uses and also as an avenue or roadside tree.

Champion and Seth (1968) mentioned its occurrence in tropical dry deciduous thorn forest and in tropical dry evergreen forest as follows:

- Southern tropical dry deciduous forest
- Southern tropical thorn forest
- Tropical dry evergreen forest
- Northern tropical thorn forest

DISTRIBUTION ALL OVER THE WORLD

The naturalization of *A. indica* in the countries adjacent to the Indian subcontinent, which include Thailand, some parts of Malaysia, Java and Bali (now Indonesia) and probably east Africa, can be explained on the basis of the long cultural and commercial relations that these areas have had with India since ancient times.
In Asia, in addition to the Indian subcontinent the tree has been reported from upper Myanmar, drier parts of Sri Lanka (earlier known as Ceylon), south east Thailand, Vietnam, southern Malaysia, drier islands east of Java in Indonesia, the Philippines, Hainan in China, middle east Yemen, Qatar, and some parts of Saudi Arabia.

In the past two centuries it was introduced into other parts of the world by immigrants from India who settled in east Africa, the Caribbean islands, Fiji, Mauritius etc. It also grows in the islands of the South Pacific, the West Indies, Haiti, Surinam, the Dominican Republic, Cuba, Nicaragua, and in some areas of Mexico. It was introduced into selected pockets of California, south Florida (USA) and Queensland (Australia).

It adapted well to the savannah conditions south of the Sahara in Africa, so introduction has been very fast in some of the African countries. It grows well in Egypt, Sudan, Ethiopia, Somalia, Kenya, Uganda, Tanzania, Mozambique and Chad in the northern and eastern part of Africa and in Mauritania, Senegal, Mali, Ghana, Ivory Coast, Togo, Nigeria and Cameroon in the rest of Africa.

**Recent Introduction in Various Geographical Regions**

Cultivation from Haiti was reported by Lewis and Elvin-Lewis (1983), who recommended the large-scale introduction of this tree in other neotropical countries, as the tree tolerated the arid conditions. It was cultivated on a large scale in Upper Volta (Sieder, 1983). Ujhmura (1986) presented a case study of neem introduction in semi-arid zones of Nigeria.

Ahmed et al. (1989) reported an establishment of 50,000 trees in the plains of Arafat in Saudi Arabia. Fallata people who originated from West Africa and now live in Sudan have long experience of planting neem trees (Kismul, 1989). Silvicultural characteristics and a manual for the propagation and afforestation in Sudan have been given by Badi et al. (1989) and Branney (1989), while Oguntala (1989), after studying the wild forest fires in Nigeria, found that the survival of exotic species like neem after fire was higher as compared to indigenous ones. In the rain-fed zones of Indonesia, Tampabolon and Alrasyid (1989) discussed the prospects of neem in the rehabilitation of critical land. Spaak (1990) included neem as one of the trees in the afforestation program on Cape Verde in an arid and semi-arid zone which has extensively high annual rainfall. Adegbehin et al. (1990) gave an account of an agroforestry project near the shelter belt in the Savannah area of Nigeria.

In the northernmost province of Cameroon, 25 to 30 percent of the land is composed of sterile or degraded soil. In an attempt to rehabilitate this soil, neem, along with other trees, were planted by Matig (1989), and the results were very encouraging. There was spontaneous regeneration of the natural vegetation. On the west coast of Reunion, Roederer (1991) gave an account of forestry and agroforestry experiments with neem. The tree survived well but grew very slowly. In Chad also, neem was introduced to improve the forest resources (Thomossey, 1991). Chiu (1993) reported successful introduction in the Hainan Province of China.

**Dry Zone Afforestation**

Encouraged by the survival of the tree in near-drought conditions, experiments were conducted on the introduction of neem in areas having degraded and arid soil. For
degraded forest areas, direct sowing was more successful and cheaper, provided that protection and shelter were available to young seedlings. Direct sowing was by dibbling in bushes, broadcasting in lines on mounds and ridges, in trenches, and in sunken beds. Aerial seeding gave profuse germination on soil in Nigeria but all seedlings in direct sunlight died and only those in the shade of natural shrubs could survive during the dry season. Kalla et al. (1978) gave an account of the felling cycle of trees in the Indian desert for integration of forage forestry in an afforestation program. Neem, along with other trees, was found suitable for the degraded condition of forest areas in low rainfall regions by Raddi (1981). In the Indian desert three shelter beds were planted in 1973 by Muthana et al. (1984). The belts had three rows of trees each and they were laid out perpendicular to the wind direction. The outside rows contained *Acacia tortilis*, *Prosopis juliflora* and *Cassia siamea*, while the center rows were planted with *A. indica*, *Albizia lebbeck* and *Eucalyptus camaldulensis*. The shelter beds were easily established and grew well. Ahmed et al. (1985), on the other hand, planted neem trees in coastal sand, using highly saline water for irrigation. An optimum spacing of 140 m for neem in shelter bed establishment was recommended by Onyewotu (1985) in the Sahel and Sudan zones of Nigeria. Bose and Bandoyopadhyay (1986), during a study of the economics of energy plantation in alkaline soil near Delhi (India), found a mortality rate in neem quite low after 3 months of planting, but height was least as compared to the other trees. These authors felt that fuel wood farming in this area was economically feasible. Chaturvedi (1985), in a forest bulletin, gave an account of the firewood farming of 16 important trees on degraded lands in the Gangetic plains in India. Neem was also included in the list. The author discussed the ecology of sites on saline, sodic and alkaline user (infertile) soils, ravines and areas of brackish water, with growth and yield statistics. The salt tolerance of neem tree seedlings was studied by Gupta et al. (1987) by carrying out greenhouse experiments on sand, loam and clay forest soils to which calcium chloride had been added. Toxicity was lowest in the clay soils, and greatest in sandy soils.

An effective method of wasteland afforestation was developed by Kinhal (1968) by early planting and critical watering, in pits at a spacing of 3×2m. Martinez (1987) gave an account of the silviculture of neem under multi-purpose tree. Panchan et al. (1989) studied the effect of salinity on the growth of the neem tree, by applying different concentrations of salt (sodium chloride). Neem did not tolerate even 0.4 percent salt. Oguntala (1989) studied a forest fire climate in relation to fire incidents. After the fire, the survival and regeneration in neem was higher as compared to the indigenous species. A technique was developed by Mehta (1989) for planting trees on highly alkaline soil, by levelling and bunding the soil and applying gypsum over the whole surface, followed by the cultivation of crops for 4–5 years. The trees were planted afterwards.

Prasad and Dhuria (1989), for the reclamation areas mined for iron ore, carried out afforestation trials with neem by planting it in pits. Half of the pit was filled with original soil and the other half with humus-rich soil and manure. Biomass data was collected. Various anti-desertification techniques were suggested by Campolucci and Paolim (1990). These suggestions were (a) contour terracing, (b) half-moon-shaped bunds, (small dams), (c) furrows up to 80cm deep and (d) deep ploughing, followed by two central rows of container-grown trees of *Acacia* spp. and *Azadirachta indica*. 
CULTIVATION

...to act as a wind-breakers; flanking them were two rows of other trees and six rows of shrubs to act as a protective barrier against browsing damage.

Desh Raj (1990) developed land which was without any vegetation but had large patches of white salts. Neem did not grow well on this soil.

Mohan et al. (1990) observed that neem grew best in clay soil. To confirm this, the authors used various potting mixtures by using different doses of soil and fertilizers. The mixture of sand-clay-farmyard manure with 30ppm nitrogen and 20ppm phosphorus was found to be the best. Singh et al. (1991) gave uses of various soil-working techniques as follows: (a) pit planting, (b) ploughing to 20cm depth, and (c) digging V-shaped furrows 30cm deep. The ploughed areas gave the best results, while Ahmed and Puzari (1991), in a growth data trial, got their best results from neem trees at a livestock research station. The tree did not fare well in coastal areas. Gill and Abrol (1991) gave a summary of the results of field experiments carried out since 1971 at the Central Soil Salinity Research Institute in India. The studies showed that the alkali soil of the Indo-Gangetic plains of north India was toxic to neem because of an excess of sodium carbonate and bicarbonate, poor soil structure and moisture transmission. The soil mixed with gypsum and farmyard manure supported some trees, but neem did not show high tolerance to these conditions. The neem tree was found superior in enriching the sandy, loamy soil with calcium and in increasing soil pH by measuring litter quality and soil fertility (Drechsel et al., 1991). Verinumbe (1991) developed an agroforestry system for neem and the other trees on vertisol soil. In a wasteland reclamation project, neem was planted with other trees as a fuel tree by Chowdhury (1992). The project was a success. The survival rate of neem was 83.33 percent after 3 years. In other experiments, neem trees were planted in arid regions (Harsh et al., 1992) for control of soil erosion and fertility enhancement of the soil (Laskar and Datta, 1992). To find the effect of the neem tree on the yield of wheat in arid zones, neem trees were planted in the wheat field by Puri et al. (1995). The authors did not observe any significant difference in the yield of farm grain in the fields with and without neem trees. Dalal et al. (1992) also worked on this direction.

Shaikh (1992) suggested that in highly eroded and degraded soil, seedlings should be planted in pits with shoulder trenches, and that for water harvesting, micro catchment areas should be formed. Shaikh (1993) suggested a change in strategy for neem tree plantation in highly eroded and degraded areas. Neem was not found suitable for sodic soil aquic petrocalcic natrustalf soil having pH 9.7 (Sharma et al., 1993). Harikrishanan (1993) gave an account of nine species, including neem, for small marginal farmers.

The rooting pattern was studied by Toky and Bist (1992), with further research on the growth pattern and architectural analysis of neem along with other trees (Bist and Toky, 1993) in a six-year-old plantation. These authors observed that the period of growth, which began in the dry season, was completed in the rainy season that followed. Brenner et al. (1995) studied wind-break crop interaction behind two rows of neem trees in a field of millet.

In arid areas, proper moisture for the root system is essential, particularly at the initial stages of tree growth. Gupta (1992) studied the growth of nursery plants of neem as influenced by soil mixture and fertilizers. Meena et al. (1995) discussed various soil techniques on early growth in arid zones, while Gupta (1995) studied different systems/combinations of rain/water harvesting. The ridge and furrow method
was found to be the best and significantly improved the growth of trees. Gupta et al. (1995) carried out various experiments to find the most suitable water harvesting techniques for plants/saplings in (a) pits of normal size, (b) pits surrounded by a saucer-shaped depression, (c) pits surrounded by a ring-shaped depression, (d) pits with a sloping side and with a mound in the middle, (e) a trench and mound, (f) deep ploughing. The effect of spacing on sprout growth was studied by Raizada and Padmaiah (1995), after coppicing. The best results were obtained by 3×1m spacing.

Other Studies

For an agroforestry system, particularly in dry areas, the rate of water consumption and biomass production are important. Chaturvedi et al. (1985) gave an account of the farming of neem as a source of fuel wood. Chandrasekharaiyah and Prabhakar (1987) studied the harvestable biomass of 4- and 5-year-old trees. It was 15.77kg for neem, as compared to 70.1 kg for Dalbergia sisoo. There was more than 50 percent increase in dry matter production after 8 weeks when 2–3-month-old seedlings were kept with extra carbon dioxide maintained at the concentration of 8000 ppm, twice as much compared to the control (Suraminath et al., 1988). Lysimeter measurements for this were taken by Chaturvedi et al. (1988). Mulching using coir pith (a waste material from coconut processing) and fertilizer treatment improved N, P and K concentration in plants and significantly improved soil fertility and moisture content (Balwinder Singh et al., 1988). The effect of mulching under different irrigation levels on the height of tree seedlings was studied by Singh et al. (1989), while Gupta (1991) gave an account of the effect of fertilizer application on initial development. Singh et al. (1991) studied the response of leaf residues and irrigation on plants. Application of leaf mulch considerably improved growth, even at lower levels of irrigation treatment.

Biomass equation above ground, biomass in dry matter and nutrient content were studied by Bunlyavejchewin (1989), Bunlyavejchewin and Kiratiprayoon (1989) and Bunlyavejchewin et al. (1989). Pine needles, water management of transplanted seedlings in arid areas was studied by Burman et al. (1991). An average application of 46 liters, full-field capacity per nine-month-old plant at two-week intervals, led to maximum growth and biomass production with no mortality. Brenner et al. (1991) determined the daily transpiration rate for the tree in an unstressed wind-break. Marti et al. (1991) studied growth and biomass in the coastal saline wasteland of east India. Gupta (1994) described the effect of rain water on biomass production in neem in the Indian desert. In Nigeria, waste water was found to result in enhanced height growth with no apparent harm (Lacuali et al., 1995).

Arya et al. (1992) described a sweeping association of Azadirachta indica with Prosopis cineraria.

Jattan et al. (1995, 1995a) suggested various steps for the management of neem plantations; this included pruning the trees for floral initiation and to stop irregular bearing. When fruit shedding is expected, it can be prevented by proper irrigation and the supply of nutrients, by removing shade and shoot tips. Weeding and hoeing, which loosens the soil and helps aeration, have also been found useful.

Various diseases in neem can be controlled, as given by Tiwari (1992).
PLANT DISEASES

Various constituents from neem have been found effective against insect, fungal and even viral pathogens but neem itself is attacked by some of these. The attack may be on seed, on the young seedling, on the leaf, on the root and even on the whole tree.

Wealth of India (1948) mentioned the various fungal diseases of neem caused by *Cercospom leucosticta, C. subsessilis, Xylaria azadiractae, Fomes* and *Ployporus*. Schmutterer (1990) has given an account of observations on 20 species of insects and 12 species of arthropods pests, along with their biology and geographical distribution. Tewari (1992) and Schmutterer (1995) have given detailed accounts of all these diseases.

The important ones, organ wise, are the following.

**Whole Tree**

Bacterial wilt by *Pseudomonas solanacaerum* has been described by Diatloff *et al.*, (1993). It caused black to brown discoloration in the root and stem with collar rot developing in the advanced stage. Benge (1993) has given an account of the disease called “neem disorder” or “neem decline”. It has been seen in an area from Mali and the west to Chad and Cameroon in Africa. The disease is characterized by gradual loss of foliage, a general debilitation of the tree and sometimes death. This is probably caused by the *Verticillium* species. The other disease is pink disease caused by *Cortidium salmoniclor*. It is characterized by the formation of a pink incrustation and canker on the stem. Phomopsis twig blight is caused by *Phomopsis*, which attacks the twig (Tewari, 1992).

**Seed**

*Oryzaepbilus surinamensis* was found infesting the seed by Zongo (1990). The infested seed were entirely hollowed out. Uniyal and Uniyal (1996) collected neem seed from 24 different localities and found them heavily inhabited by seven fungal genera, *Fusarium, Aspergillus, Penicillium, Cephalosporium* spp., *Alternaria* spp., *Pythium* spp. and *Mycelia sterelia*. The dominating species were *Aspergillus niger, A. fumigatus* and *Fusarium*.

**Seedlings**

In a forest nursery, Raghunathan *et al.* (1982) saw thousands of fire ants (*Solenopsis* spp.) attacking the saplings, and completely defoliating them. These ants did not touch the other plants nearby. Sankaran *et al.* (1988) observed leaf spot disease caused by *Colletotrichum capsici* and *Cercospora subsessilis* in 3-month-old seedlings. Leaf blight and stem rot by *Sclerotium rolfsii*, and Web blight by *Rhizoctonia solani*, have also been observed. Twig blight has been reported by Kaushik *et al.* (1993). It caused minute, circular, black to brown lesions on the neck region of 2–3-month-old seedlings. The infection caused the death of seedlings. The causative organism was the *Colletotrichum* state of *Glomerella cingulata*.

**Root**

*Phytophthora dannamoni* has been found attacking the tree worldwide (Benge, 1993). Black to brown discoloration occurred in the root due to *Ganoderma applanatum*.
(Chakraborty and Kongers, 1995). Ganoderma root rot is caused by *G. lucidum*. Drying of leaves takes place after the root system is damaged (Tewari, 1992).

**Leaves**

The various diseases reported are leaf spot by *Xanthomonas azadirachti* (Moniz and Raj, 1967), powdery mildew by *Oidium azadirachtae* (Naryanasmy and Ramkrishnan, 1971) and bacterial leaf spot (Nayudu, 1972). Singh and Chohan (1984) observed that *Phoma joylana* caused severe twig canker and shot holes in the leaves. Sankaran *et al.* (1988) for the first time recorded the *Colletotrichum gloeosporioides* state of *Glomerella cingulata*. Jamaludin *et al.* (1988) described Cercospora blight disease, caused by *C. subsectalis* in winter. It started on the leaves, and spread to the other parts, leading to shedding of leaves and fruit. Seeds so produced were not viable.

Harsh *et al.* (1989) also observed neem spot caused by *Cercospora*. Pillai and Gopi (1990) mentioned that the drying up of the distal parts of the shoot and foliage in the months December to April has become a common phenomenon in south India, which is due to the tea mosquito bug. In addition to this, 16 insect species and one mite, *Caliptrimerus azadirachtae*, were identified. Mehrotra (1990) gave an account of foliar disease, leaf web blight, caused by *Rhizoctonia solani* in forest nurseries. The symptoms of leaf web blight disease are the development of greyish brown blotches, which grow with age and cover the whole leaflet. The infected leaflet remain joined by fungal hyphae, as in the case of a spider’s web. The disease appears during the high humidity months. The pathogen is borne in the soil, and climbs up on the leaves, infecting them one by one. Hiremath *et al.* (1991) recorded infection by *Cercospora leucosticta* and *Glomerella cingulata*. Colletotrichum leaf spotting blight and Alternaria leaf spotting blight have been described by Tewari (1992).

Karthikeyan *et al.* (1993) observed neem to be infested by 13 pests; the tea mosquito bug *tiellopeltis antonii* the mealy bug *Pseudococcus gilbertensis*, the scale insect *Parlatoria orientalis* and the leaf webber *Loboschiza koenigiana* are the important pests. *H. antonii* damaged the entire foliage from October to March.

Alam (1993) noticed that *Curvularia* spp. and *Exserohilum* spp. caused leaf spot patches, resulting in total rot of the leaf. Another dermataceous hyphomycetes was also recorded causing leaf spot.

**REVIEWS**

Shankar (1988) gave an account of silvipasture studies on neem in India, for forage and fuel purposes, while Siddique (1989) described neem as a multi-purpose species in Pakistan. Social forestry and rural economy have been discussed by Sharma (1989) for fuel, forage and the development of small-scale/cottage industry. The same approach has been given by Adegbehin *et al.* (1990) in Nigeria. Kamo (1990) and Lauridsen *et al.* (1991) studied it as a fast-growing species in Thailand. Hegde (1991) studied the silvopastoral system with a brief account of land utilization. Tewari (1992), in a monograph on neem, devoted a chapter to silviculture and management. Sidhu (1995) gave an account of neem in agroforestry, by combining the woody perennial...
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with annual crops and animals on the same unit of land management to obtain maximum productivity.

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5. PLANT RAW MATERIAL

The seed as a source of azadirachtin and the oil are the main raw material of the industry. To some extent, bark is used in Ayurvedic and Homoeopathic medicines and leaves as a herbal remedy. An account of these is given separately.

THE SEED

Collection

During field studies, it was observed that the neem tree has a sporadic distribution; there are no neem forests, consequently only a small quantity of seed could be collected from a particular area. Mostly seed is collected by picking ripe seed which falls on the ground by itself but sometimes, when the fruit turns yellow, the branches are shaken or beaten by the collector climbing the tree. By this act the fruit falls on the ground and may be collected on a sheet of cloth or made into a heap by sweeping the ground with a straw broom. The neem fruit matures in most places during the rainy months of monsoon, when there is not only high atmospheric humidity and water logging of the soil but water everywhere on the ground accompanied by flood and rivers in spate. This makes the job of collecting, drying and transporting good quality neem seed very difficult. In the absence of any known post-harvest technology, and in the above conditions, in earlier days most of the seed deteriorated during and after collection. A good infrastructure is required for the transportation of seed from rural areas to the markets or to collection centres in the towns.

Harvesting Methods

Mitra (1963) carried out a detailed literature survey and gave proper post-harvest technology for the seed, so that good quality raw material is obtained. It was seen that in the markets seed was brought in the following stages: (1) dry, (2) semi-dry, (3) dry but wet, (4) damaged and (5) fresh.

Drying Methods

For proper drying of the seed, Mitra (1963) suggested that they should be regularly moved for aeration. This can be done in the field by making four quadrants. The seed should be stored on racks of bamboo or on benches for movement of air. Every day one heap should be moved from one quadrant to another. When the seed are at the optimum level of moisture and there is no danger of fungal contamination after storage, they should be de-pulped.

The process of de-pulping involves soaking the seed for 4 to 5 days in water. When the pulp becomes soft, the seed should be fed into a revolving drum which has a central shaft with pedals. The inner side of the drum has baffles. After feeding the
seed, the drum can be rotated by hand or by other mechanical means so that the outer soft seed coat gets separated from the inner hard seed, leaving some pulp only.

The above treatment is for large-scale operations but for smaller quantities, the seed are de-pulped by macerating them by hand after the addition of wood ash. By this treatment the mucilaginous pulp loses its mucilaginous nature, and the removal of the seed coat becomes easier. In villages in south India, the seed coats from fresh seed are removed by collecting them in a heap on the dry ground and placing old jute bags on it. The seed are macerated by foot, by standing on the heap. By foot movement, pressure is exerted on the seed pulp which causes the separation of the seed coat. The mucilaginous seed pulp sticks to the jute bag.

For proper storage, it is essential that seed are free of pulp. The pulp not only absorbs moisture during storage, but acts as a medium for fungal growth, because of its nutrient nature. Seed can be cleaned of pulp by repeatedly washing them with water. In a method suggested by Mitra (1963), an inclined metallic sieve is used. The de-pulped seed are spread on it with a shovel and a jet of water is sprinkled on them by force. This separates the remnants of pulp on the seed, which escapes through the sieve. The clean seed settles down.

The seed are dried once more in dry air or in dryers.

Storage Conditions

As given in Chapter 4 on cultivation, quite a number of studies have been conducted on the proper storage of seed so that there is no deterioration. It has been observed that seed made free from pulp by repeatedly washing them with water have less chance of getting contaminated. Both high humidity and high temperature are conducive to fungal growth within and outside the seed. If the seed are stored in a dry atmosphere and with good aeration, the deterioration of seed can be prevented to a major extent.

Morphology of the Fruit and Seed

Fruit: It is an ovoid drupe, yellow to brown when ripe, epicarp thin, mesocarp with scanty mucilaginous sweetish pulp, endocarp hard enclosing the seed. The fruit gets darker in color and wrinkled on drying (Fig. 9A). When the seed coat is removed, seed with hard endocarp get separated (Fig. 9b1; b2). Seed exalbuminous, plano-convex, notched at the base (Fig. 9c1), very often the seed kernel is covered over by brownish fungal spores (Fig. 9c2) or is totally replaced by these spores. The seed length varied between 11 and 18mm, with width 4.5–8.5 mm and weight 100–530 mg.

THE BARK

Collection

It is collected mainly from the felled trees by stripping it off from the stem and dried in shade. It may also be obtained from the standing tree, but in that case the precaution is taken that only a small portion of the stem is exposed, otherwise the tree may die.
of dehydration. Where it is essential to remove most of the bark, the areas of the tree from where the bark has been removed are pasted with a plaster of clay or mud and kept moist for a few days.

Characteristics of the Bark

Bark on young stem or branches is smooth greenish to rusty green. In transverse section, it can be differentiated into three zones:

1. A narrow pink outer part.
2. A whitish, brown middle portion.
3. Thick innermost region of secondary bast. A few secretory cavities are found scattered in the phloem.

The older bark is with numerous scattered tubercles, grey or dark grey to grey black in colour, feebly fissured and exfoliating. The entire bark is comparatively thin, about 10 mm thick. The outer cork consists of nearly half the thickness of the entire tissue. The inner surface is pink brown and fibrous. It has very feeble smell when fresh and a slight bitter taste.

In the old bark, a well-defined outer rind is present, formed of alternate strips of cork layers and dead secondary bast. The cork layer is composed of secondary cells,
often with reddish brown contents. Phellogen is not very distinct and a secondary cortex is not normally present. The secondary bast is composed of groups of sclerenchyma which are polygonal, isodiametric with a thick unpitted wall and medullary rays. In still older barks, collapsed and compressed phloem tissue is present. The phloem parenchyma cells are packed with compound starch grains. Medullary rays are usually 2–5 serrate. Most of the cells contain starch. Cubical, rectangular and polyhedral crystals are quite common.

Root Bark

It has numerous oblong lenticels, 2–5 mm long, arranged regularly in longitudinal rows. The outer bark generally consists of thin walled cork cells, yellow to rusty brown in color. In older bark there may be alternate zones of thin hard crustaceous portions and soft cork tissue.

In older root in transverse section, a large number of pores and medullary rays are visible. Medullary rays are usually 2–5 serrate.

Leaves

These are collected from the tree, with the young twigs, chopped into small pieces and dried.

Characteristics of the Leaves

The detailed morphology is given in Chapter 2 on Plant Sources.

The micromorphological characters are: Leaf epidermis, adaxial cells elongated with straight or slightly undulating walls, stomata absent or a few. Abaxial cells at intercostal zone are with straight to slightly undulating walls. Stomata are common and oval in shape, anomocytic, 360–500 stomata per mm, each surrounded by 4–6 subsidiary cells and slightly raised on the general surface. In some cases stoma are plugged with thick resinous deposits. Costal cells are elongated and arranged in rows. Crystals in the form of druses, single or in groups, are present in the epidermal cells. Hairs unicellular to multicellular, 50–200 µm in length and 10–15 µm in width at the base. The leaf has a normal dicotyledonous structure, when seen in transection.

REFERENCES

6. QUALITY ASSURANCE OF PLANT RAW MATERIALS

Bark is official in the *Indian Homoeopathic Pharmacopoeia* (1971) while in the industry, neem oil and neem seed cake are used.

**BARK**

*Indian Homoeopathic Pharmacopoeia*

This describes both the macroscopical and microscopical character of the bark. As per the pharmacopoeia, the bark is dark grey to greyish black externally, while the inner surface is pinkish brown and fibrous. In old bark there is a well-defined outer rind formed of alternating strips of cork layers and dead secondary bast. The cork cells have reddish brown contents. Phellogen is not very distinct and a secondary cortex is normally not present.

**SEED OIL**

Azadirachtin in the samples can be estimated by the method given in Fig. 7 in Chapter 3 on chemical constituents. The physical standards of the oil are given by the Panel of the Ministry of Industry, the Government of India (Table 2), the Solvent Extractor Association of India (Table 3) and the Indian Standards Institute (Table 4).

**Composition of Oil**

In the earlier literature (*Wealth of India*, 1948), oil was reported to contain 0.2 percent myristic acid, 16.2 percent palmitic acid, 14.6 percent stearic acid, 3.4 percent archidic acid, 56.6 percent oleic acid and 9.0 percent linoleic acid. It is quite rich in tocopherols which may be up to 1.17mg/g, having equal amounts of α and γ but very little β tocopherols. Recently Milan Mehtra (1997) has given the composition of the oil as follows: myristic acid 2.6 percent, palmitic acid 13.6 to 14.9 percent, stearic acid 14.4 to 19.15 percent, oleic acid 49.1 to 61.9 percent and linoleic acid 7.5 to 15.8 percent. The percentage of glycerides is 0.6, fully saturated glycerides 22.0 percent,

| Table 2 Standard of neem oil by panel of Ministry of Industry, Government of India |
|-----------------|----------------------------------|
| Moisture and insoluble impurities | ≤0.9% |
| Saponification value | 180–205 |
| Iodine value (Wij’s) | 70–82 |
| Un saponifiable matter (by wt.) | ≤2% |
| Flash Point (Pensky Marten Method) | ≥120°C |

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tri-unsaturated glycerides 34.0 percent, stearodiolein 2.0 percent, palmitodiolein 26 percent, oleopalmitostearin 12 percent and oleodipalmein 5 percent.

SEED CAKE

Keeping in view the importance of this in agriculture, the Indian Standards Institute (Anonymous, 1977) has given specification No. 8558 for neem cake for manuring, which is as follows:

- Maximum moisture (percentage by mass) 10%
- Maximum water soluble organic nitrogen on moisture free basis (percentage by mass) 2.5%
- Maximum total ash (percentage by mass) 13%
- Maximum acid insoluble ash (percentage by mass) moisture free basis 5.0%

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7. WOOD FOR FUEL AND TIMBER

Neem in recent times has emerged as a tree of choice for afforestation projects and for meeting the fuel and timber needs of under developed countries (Kalla et al., 1978; Anonymous, 1980, 1983, 1986; Radwanski and Wickens, 1981; Grainger, 1982; Prasad, 1983).

CHARACTERISTICS OF NEEM WOOD

Sapwood greyish white, heartwood red, when exposed fading to reddish brown and then resembling dull to somewhat lustrous, aromatic with characteristic taste. The wood is hard, closed grained. According to Gamble (1902), annual rings are doubtful, the wood shows alternating bands with numerous and fewer pores, and pale concentric lines but Pearson and Brown (1932) mentioned that growth rings are distinct, sharply delimited by narrow brown concentric lines. Pores scanty, moderate sized and large, often oval and sub-divided.

Medullary rays numerous, white, prominent, bent outward where they touch the pores, the distance between the rays less than the transverse diameter of the pores. Rays oval to elliptical with linear and lenticular orifice, oil globules frequent, crystals not found. Gill et al. (1985) gave the morphological characters of treachery elements, i.e. vessels, fibers, rays, parenchyma and tracheids. Vessels form inconspicuous vessel lines along the grain, which are darker than the background and contain deposits of reddish brown gum. Vessel segment short (120–450 µ) abruptly or alternate tailed on one or both sides. The perforation is simple, horizontal to oblique, frequently occupied with gum. Nair (1987) reported Scanning Electron Microscopical (SEM) studies on the helical thickening covering the entire vessel elements. Parenchyma terminal paratracheal, paratracheal zonate and meta tracheal in cambiform rows. Fibers non-libriform or semi-libriform in radial rows, non-septate, 250–1620 µ in length and 22–28 µ wide.

Nair (1988) gave a revised account of wood anatomy and heartwood formation. According to the author, wood is of diffuse porous type. Axial parenchyma cells and sometimes the vessel and fiber of heartwood show the presence of extractives. The necrobiosis of the parenchyma cells occurs at the heartwood boundary. The death of parenchyma cells is associated with depletion of starch grains and accumulation of extractives.

Mechanical Properties of the Wood

The wood is moderately heavy, the average weight is 50–52 Ib per cubic feet (Gamble, 1902). Pearson and Brown (1932) gave the following mechanical properties of the wood:

- Compression parallel to grains Ib/sq inch=6680
- Shear parallel to grains Ib/sq inch=1326
- Module of elasticity or Young modulus=1,008,800
Sekhar and Gulati (1971) determined the strength parameters of neem wood with respect to teak (*Tectona grandis*) as 100, which are as follows:

- Weight at 12% moisture content=124
- Strength and stiff as beam=87
- Retention of shape=77
- Shear=129
- Surface hardness=131
- Refractoriness=113
- Nail and screw holding power=117

Koul *et al.* (1990) reported the specific gravity of wood as 0.65–0.85. According to these authors, the wood is straight grained and is more resistant to shock.

**WOOD AS TIMBER**

After extensive study the wood was recommended for general construction (Sekhar and Gulati, 1971; Rajput and Shukla, 1984). It was found to be naturally decay resistant (Rao, 1990), absorbing a minimum amount of water so good for outdoor use (Das *et al.*, 1993) and for timber house construction (Punhani, 1995) but Kossou (1992) did not find it suitable for the construction of granaries because of susceptibility to insect attack.

On the basis of the tests for glue adhesion, tensile strength, bending, compressive strength, amenability to preservatives and fireproofing treatment, Chauhan and Bist (1987) considered neem suitable for general-purpose plywood, while Shukla *et al.* (1990) studied its carving behavior. Punhani and Pruthi (1990) measured the lateral bearing strength using nails and shear strength with joints loaded parallel, perpendicular and at 30°, 40°, 60° to the grains. Pant *et al.* (1962) found overall performance of the wood better than teak (*Tectona grandis*) as far as working properties are concerned.

The earlier literature on the wood quality of neem was studied by Tewari (1992) in detail. It has low shrinkage and seasons well in air. The wood is durable, but impregnation with chemicals like preservative and adhesives is difficult because of the gummy deposits in the cells. The quality index based on the quality performance is 114 as compared to 100 for teak. It resembles mahogany, hence is sometimes called “Indian mahogany”, but lacks grain and smoothness. The compressed wood shuttle used in the textile industry, made from neem wood, was studied by Shukla and Bhatnagar (1988, 1993), but not found suitable because of lack of enough strength.

**WOOD AS FUEL**

Fuel wood characteristics, viz. density, calorific value, and content of C, H, N and O, were determined by Jain (1990). In Ghana, the first rotation of neem yields 108–137 m³ of fuel wood per hectare, but in northern Nigeria 19–69 m³ per hectare after 8 years (Tewari, 1992). Puri *et al.* (1994) determined the fuel wood value index for it,
taking into account calorific value and density as positive characteristics, and high water content and high ash values as the negative points. Neem had better fuel efficiency in comparison to *Prosopis tinereria*, but was less efficient when compared with other indigenous trees like *Acacia nilotica*.

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8. PROCESSING OF PLANT RAW MATERIAL

THE SEED

According to an estimate, there are 13–15 million neem trees in India alone. If one tree produces, say, 50 kg of seed and even a small percentage of the total yield is processed for the oil, the quantity of oil and seed cake so produced would be enormous. The industry based on neem (Fig. 10) will not only provide extra income to the rural poor for the collection of seed, but produce a fatty oil, which can be raw material for many industries. As a by-product, neem seed cake will be available for use directly as organic manure and indirectly as a nitrification inhibitor and pesticide.

Neem as a source of oil has been of interest to both social workers and scientists for a very long time in India, but the yield of oil by traditional methods was less; moreover, the oil obtained by bullock-driven expellers, called “Kohlu” or sometimes “Katchi Ghani”, was darker in color, fetid in odor and bitter in taste. It was not acceptable to consumers in any form and could not be put to any major industrial use. It was used by the lower strata of society to a limited extent for the skin diseases of man and farm animals, as a lubricant for farm implements, particularly for the leather straps, for prolonging their life and as an oil for use in clay lamps for illumination.

The earlier studies were confined to refining the oil by chemical means by removing the bitter principles. Sen and Banerjee (1931) studied these bitter principles. Child and Nathanael (1944) gave the fatty oil composition of neem, which was very similar to the other edible oils. Siddiqui (1942) presented a note on the isolation of three new bitter principles from neem oil, while Siddiqui and Mitra (1945) suggested various ways by which these principles could be utilized in the pharmaceutical industry. Quite a number of patents were also taken out on this aspect (Siddiqui and Mitra, 1945a, 1945b, 1945c), but these studies could not lead to the industrial utilization of neem.
oil, because the oil obtained was not of a desirable quality even after refining, and there was no major use to which the bitters could be put.

Collection

When the problem of not getting good quality oil even after refining was studied in detail, it was seen that it was mainly due to the low quality of the raw material used. The oil made from these heavily contaminated low quality seed was resinous and malodorous. The Indian Oil Seed Committee, then located at Hyderabad, looked into all the above aspects and Mitra (1963) came out with an excellent book, *Neem*, in which on the basis of field studies, experiments and trials, methods of obtaining a better quality of oil by drying the seed properly before storage were given. Mitra (1979) gave an account of technology, incorporating the other developments.

Decortication

The seeds are dried once more in dry air or in dryers and are subjected to decortication, i.e. removal of the kernel from the seed coat. It can be done in various ways. Mitra (1963) suggested double roller mills, with rollers moving in opposite directions. He used cast iron rollers with various types of grooves. These days, the rollers have been replaced by decorticators used for dehusking peanuts. These decorticators essentially consist of rubber rollers with a blower at the base of the chamber. The roller presses the seed coat and releases the seed kernel; the mixture of both seed and seed coat passes through a sieve into the chamber, where the lighter seed coat gets blown away, keeping the seed kernel in the chamber.

In improved decorticators (Figure 11B) now, a single roller rotates above a sieve, which presses the seed between them, causing separation of the seed coat and the

![Figure 11 Decorticators. A, an improvised technique using old tyres; B, flow diagram of improved machine. Abbreviations: BL=blower, CP=central pulley, HO=hopper, KE=kernel, SC=seed coat, SD= whole seed, SH=seed husk, SI=sieve, UT=used tyres](image)
kernel. On a small scale, the seed coat can be separated by taking two old car tyres (Fig. 11 A). These are fixed by a central shaft so that the two tyres are adjacent to each other. The whole seed is fed between the gaps of the tyres, which get pressed by their movement, causing the separation of the seed from the outer cover; the two can be separated from the mixture by a winnower. This winnower in earlier times was made from bamboo strips or reeds. It essentially consists of a broad plate, tapering at the broad end, and having a raised platform on the other. This winnower is moved left to right and up and down, after loading with the mixture, and by this action the heavy seed remain on the upper part of the winnower while the lighter chaff comes out at the open free end. An improvised winnower can now be easily made by cutting a big plastic Jerry can obliquely. The plate so obtained has a raised platform on one side, raised sides and a tapering free end.

Winnowing can also be done by utilizing the wind direction. The mixture of chaff and seed is placed on the winnower and is lifted by a person in a standing position. The winnower is shaken slowly by hand movement in the direction of the wind so that the mixture is allowed to fall on the ground. By this action the lighter chaff flies away because of the wind and the heavy seed kernel falls on the ground, nearly perpendicular to the standing person. This operation can also be performed by using a blower. The separated seed kernels so obtained can be used for oil extraction.

Extraction of Oil

Traditional Method

Oil extraction in India was a specialized profession and the people undertaking this job were called “Teli” or the oilmen, and the traditional oil extractor was called “Kohlu” (Fig. 12A). One well-known version of it, available now, is called “Katchi Ghani” or the cold process, in which heat is neither applied nor generated (Fig. 12B).

In earlier times, the dried whole fruit with seed coat intact was used for extraction in a kohlu at certain moisture levels, and no efforts were made to remove the seed coat or pulp, but the yield of oil was very low because the seed coat absorbed most of the oil liberated from the kernel. The Kohlu was made of wood in earlier days. It essentially consisted of a pestle and mortar connected to wooden plank. The pestle had a metallic cover at the base and the inner surface of the mortar was made rough with strips of wood or bamboo. For the extraction of oil, the seed were fed into the mortar. The wooden plank was rotated by blind folded oxen in a circular motion. This caused the movement of the pestle along the sides of the mortar, crushing the seed which came in between the two. Water was sprinkled on the seed from time to time to maintain a particular level of humidity. The oil, on extraction, settled down at the base of the mortar, and was collected by a bamboo pipe, inserted in a hole at the base of the mortar.

The present modified version of the above kohlu (Fig. 13A) consists of a rotating mortar installed on an electric motor. The pestle is stationary, and can be adjusted along the sides of the mortar by a spring and rod adjacent to it. After feeding the seed, the mortar is rotated, and the seed which come in between the pestle and the mortar get crushed, liberating the oil. Because of the low yield of oil, the use of above
cold process was discouraged by Indian government agencies and stress was laid on solvent extraction plants or on big mechanical expellers.

Much of the neem oil in small-scale industry is obtained by expellers made of iron, which is a screw-like press in a horizontal position. In improved versions, a boiling kettle is attached to the expeller which is heated by steam. The yield of oil is greater when using these mechanical devices, but the quality of oil is very poor, because of metallic contamination and the use of excessive heat. The oil obtained is black, tar-like, resinous and with a bad odor. The steam required by the industry is generated by waste-fired boilers, where neem seed coat is used as a fuel.

Since the seed coat is not an efficient fuel and has low combustibility, it has to be fed regularly to the furnace with a spade. To avoid the seed coat forming lumps, a staircase like structure is placed at the mouth of the furnace so that the seed coat gets well spread on the fire inside the furnace. If the seed coat is fed in a mass onto the fire, it extinguishes it.

**Filtration of Oil**

The oil obtained by traditional methods has an appreciable amount of suspended particles, which do not get filtered by coarse cloth or by simple filtration techniques. The best method for their removal is to let them settle by keeping the oil undisturbed for a few days. After that, the oil can be decanted off. The sediment which has settled
at the bottom is very rich in oil and can be used as such in place of oil or can be re-fed to the expeller along with the seed for the extraction of oil.

When the quantity of the oil to be filtered is large, the oil is subjected to a mechanical filter press. This filter press essentially consists of wooden or iron frames, and a long filter cloth (Fig. 13B). With the help of a motor, when the frames are pressed along with the filter cloth, the oil gets filtered. It is a common practice to filter the oil twice, and the oil so obtained is sold as “double filtered”.

If a very refined oil is required, the oil may be processed in a centrifuge or a filter aid may be used.

**Solvent Extraction**

Much of the oil in industry is obtained by solvent extraction, using hexane as a solvent. It is based on the principle of Soxhlet extraction, in which the seed is repeatedly extracted with the same solvent. In Fig. 14, a photograph of a pilot plant is given. Solvent extraction gives maximum yield of good quality oil.

In the case of neem, the starting material may be the whole kernel or the seed cake obtained after the cold process or from expellers. In this process, the seed or seed cake is crushed to a particular particle size and mixed with hexane which is used as a solvent. The oil dissolved in the solvent is filtered to leave the hexane-insoluble portion in the sieve. The mixture of hexane and oil is subjected to distillation which separates the volatile hexane from fatty oils. The solvent residue from this oil may be removed by passing steam through it or by de-odorization.

Because of the hazardous nature of hexane, some alternative solvents such as propyl alcohol have been suggested, but they are uneconomical.

**Refining of Oil**

In spite of the best efforts by chemists in India, neem oil free of bitter principles could not be obtained. This is because of the unique nature of these principles. These are so
intimately mixed with oil that their separation during the refining of oil was not possible for a very long time. Since 1945, all the well-known methods for refining oil have been tried, but the results were not encouraging. Ahuja et al. (1976) worked on miscella refining of oil. Ramachanderaiah et al. (1977) tried alcohol segregation. In one alternative process, neem oil was saponified and grained, while in the second process boiling was carried out after dilution with water and adding a higher concentration of caustic lye. The odoriferous compounds along with the free fatty acids were removed by distillative de-acidification or by splitting into odorless fatty acids (Milan Mehtra, 1997).

Figure 14 A pilot plant for solvent extraction. The neem seed are fed into the vessel, extracted with petroleum ether, hexane or any other suitable solvent, filtered and the filtrate is subjected to solvent recovery, to obtain fatty oils along with fractions effective as pesticide (Photo by author)
The Central Oil Technology Research Institute in India has recently developed a process for the cold extraction of neem oil, rich in 2000 ppm or more of azadirachtin (Ramkrishna et al., 1995). In this process, the temperature during the whole operation does not go beyond 40°–45°C.

An account of an improved technology for the extraction of oil and for the recovery of azadirachtin has been given by Waghray (1993), in which seed collected from low humidity areas are immediately de-pulped and dried in a continuous drier. Double twinroller decorticators are utilized with continuous winnowing for the separation of the kernel, and from it oil is extracted. The whole process is monitored throughout by High Performance Liquid Chromatography for azadirachtin.

Standards of the Oil

Keeping the economic importance in view, particularly for the soap industry, standards for neem oil have been developed by various organizations; these are given in Tables II, III and IV of Chapter 6.

PESTICIDE FORMULATIONS

Commercial

Mainly two types of neem-based products are now available. In the first case, a standardized quantity of azadirachtin is incorporated, along with other compounds, to make an azadirachtin-rich product having a long shelf life. In the other case, azadirachtin-rich neem oil is taken and fortified further with azadirachtin to a particular level. In the first case, pesticidal activity is due only to azadirachtin and there is every possibility that some of the insects may develop resistance to it. In the second case, in addition to azadirachtin the formulation has many compounds, naturally present in neem oil. This type of product may have a multi-pronged attack and even if the pests develop resistance to one active ingredient, there are other compounds in it for pesticidal effect.

On a Small Scale

In Wealth of India (1948), several recipes for the control of pests by small-scale farmers are given using neem-based formulations. All of these were developed before the unstable nature of the active constituent azadirachtin was known, and might not have been very effective. The simple formulations were painting the tree trunk with neem oil, or preparing a water in oil emulsion for spraying on crops, using household detergent or mixing a paste of neem kernel prepared with a stone grinder in water and filtering the emulsion through a coarse cloth. Lange and Feuerhake (1984) conducted experiments to see if synergist-like piperonyl butoxide could be used in this type of preparation. Rajasekaran and Kumaraswami (1985) studied other methods by which the efficacy of these extracts could be increased. Schmutterer (1985) successfully demonstrated the suitability and application of neem in the control of pests in the tropics.

From a crude neem kernel extract, Feuerhake and Schmutterer (1985) developed an inexpensive standardized formulation for small-scale farmers by using crude neem
kernel extract, solvent glycol ether and propyl gallate benzophenone. The activity was confirmed by bio-assay and chromatography.

Ketkar (1989) in the Neem Mission in India propagated several simple neem preparations for the use of small farmers, but in the absence of any direct anti-knocking agent of neem formulations as compared to synthetic chemical ones, these preparations did not become popular. It was difficult to convince a peasant that a common tree like neem can be effective against insects.

Olaifa et al. (1993) were aware of the unstable nature of azadirachtin, so they developed a technology to improve the shelf life of an emulsifiable concentrate from neem oil by using an aqueous extract of Tetrapkma tetrapera L., along with 0.1 percent propyl paraben and 0.1 percent octylgallate, which prevented microbial spoilage and oxidation of emulsion. Optiz (1991) developed methods which could be utilized by small-scale farmers, by using their own skills.

Vijayalakshmi et al. (1995) described a number of methods which a villager could apply without the use of modern technology, but incorporating modern knowledge about the degradability of azadirachtin in the environmental conditions. Heating was to be avoided in all cases; emulsion should be freshly prepared, and sprayed in the morning and evening, when there is very little sunshine. In summer the frequency of spraying should be greater. Spray could be prepared by soaking neem seed, leaves, etc. overnight and straining through a thick cloth in the morning. In the case of oily preparations, common household detergent may be added as an emulsifier. These authors did not mention the use of soap prepared from neem oil for spray, but this could prove very helpful for the non-cereals crop maturing in March-April, where attack by aphids is so wide-spread, particularly in mustard fields, that these insects could be seen suspended in air like dust particles, even in urban areas.

For Treatment of Bags

It is common practice in India to store the grains in used bags for economic reasons. These bags, or even new ones, harbor the worms or their eggs before use and attack the cereals stored in them. Much of the infestation of these grains by well-known worms can be prevented if these bags are treated with some pesticide before use. Vijayalakshmi et al. (1995) have recommended the soaking of jute bags in 10 percent neem kernel emulsion for 15 minutes and then drying them, before use.

In the villages, it is common practice to plaster hut or floor or small storage bins with clay or mud frequently. For this mud plastering, neem seed or oil emulsion can be incorporated in the mud mixture in place of water to keep the pests away.

NEEM OIL AND SOAP TECHNOLOGY

Before synthetic detergents became popular in India, soap making, particularly for laundry, was a cottage industry and any readily available, cheap fatty oil was incorporated in soap formulations. Neem oil was one of the ingredients, but was not
preferred because of the malodor and dark color of the soap so obtained. Conventional
industrial methods failed to refine the oil because the bitter principles are mixed with
the lipid constituents so intimately that it is difficult to separate the two. Mitra (1963)
gave an account of various steps taken for refining neem oil for the soap industry. In
due course of time, the soap manufacturers learnt that neem oil could be incorporated
up to 15 percent into the mixture of fatty oils along with cotton seed oil, coconut oil,
etc. The cotton seed oil was found to mask the odor of neem.

For the manufacture of soap on a small scale by the cold process, vegetable oils
are mixed with 13 percent caustic flakes and 25 percent water and stirred vigorously
until a thick paste is formed, which on keeping overnight becomes solidified and is
cut into cakes. Ketkar and Ketkar (1995) have also provided information about
other formulations for this type of soap making.

In an alternative method, on a small scale, neem oil with an excess of water and
caustic soda is heated and after boiling, allowed to cool. Many of the impurities
either escape in the air by heating or get dissolved in water. This mixture of neem oil
and caustic soda is further treated with other fatty oil to get a soap of the desired
composition.

In large-scale operation, for making toilet soap, one method suggested was to
prepare noodles as above, while the other method consisted of the separation of fatty
acids from crude neem oil. These pure fatty acids are then reacted with alkali to get
soap. Milan Mehtra (1997) gave a method for the production of quality soap by
saponifying the neem oil and graining it, where most of the smell and the color are
separated out. A second boiling is then carried out by diluting with water and adding
a higher concentration of caustic lye. The author has also suggested distillative de-
acidification. The distilled odorless oil so obtained is subjected to the usual splitting
and distillation, to get a light-colored odorless soap.

NEEM OIL IN THE LEATHER INDUSTRY

Common salt (sodium chloride) is commonly used for curing hides and skin, and is a
major pollutant in tanning effluents. In a process developed by the Central Leather
Research Institute in India, neem oil has been used in place of salt with good results
(personal communication).

NEEM OIL AS FUEL

Neem oil as a substitute for petroleum-derived fuel has been considered from time to
time. Mitra (1963) has given an account of all these developments. By pyrolytic
degradation a product, “pyronimin,” was obtained, which could be converted into
many industrial products. Munavu (1984) considered it a non-conventional vegetable
oil for fuel. Bansal and Juneja (1989) studied the performance of neem oil as a diesel
engine supplement fuel, while Renuka and Ramani (1989) tried neem cake as a source
of bio-gas.
OTHER USES

Neem can be used for the production of olein and high melting stearin after hydrogenation. Kane and Kulkarni (1954) carried out high pressure hydrogenation of oil. A process for edible neem oil has been developed by Lidert (1994) in which crude oil is treated with an alkaline solution or hydrogen peroxide and subjected to distillation or chromatography. The oil so produced has a very low sulfur content.

NEEM CAKE AS CATTLE AND POULTRY FEED

Neem cake is very rich in proteins and comparable to soya bean, peanut or even fish meal, and can be partly substituted for any of these in cattle and poultry feed (Ketkar, 1995). Neem seed cake is very rich in minerals and is easily converted into organic minerals by animals during digestion. If neem cake is supplemented in the diet, the animals do not require additional inorganic minerals in the diet. It is very rich in amino acids like methionine and lysine. Further details are given in Chapter 14.

Leaf extract can be used as a preservative in food to prevent it from bacterial infestation and aflatoxicosis. Neem oil in very low concentration may be used as an antimal for poultry and cattle feed.

RECENT DEVELOPMENTS

Earlier, the stress was on obtaining an odorless neem oil for non-edible purposes, but with the discovery of azadirachtin, processes have now been developed for the isolation of bitter principles, and a good quality oil is obtained as a by-product.

For the recovery of pesticide, Amata-Archachai and Wasuwanich (1986) suggested a method for the de-pulping of seed, storage and extraction. According to Hoschle-Zeledon and Keyserlingk (1993), a pilot plant for the production of a standardized extract of neem as an insecticide has been in operation in Myanmar (Burma) since 1987. In this process, the seed are extracted with methanol to get the crude extract rich in azadirachtin. The extract is formulated with an emulsifier and diluted with methanol. Waghray (1993) described a new technology for pesticides and oil. Ramakrishna et al. (1995) have developed a cold process for neem oil. Sivakumar et al. (1995) have described various machines for processing neem, while Dakshinamurthy (1995) has given an account of technology for the production of azadirachtin.

REFERENCES


9. TRADITIONAL USES

The therapeutic efficacy of neem must have been known to man since antiquity as a result of constant experimentation with nature. Ancient man observed the unique features of this tree: a bitter taste, non-poisonous to man, but deleterious to lower forms of life. This might have resulted in its use as a medicine in various cultures, particularly in the Indian subcontinent and later on in other parts of the world.

AYURVEDA

The word neem is derived from Sanskrit Nimba, which means “to bestow health”; the various Sanskrit synonyms of neem signify the pharmacological and therapeutic effects of the tree. It has been nicknamed Neta—a leader of medicinal plants, Pichumarda—anti-leprotic, Ravisambba—sun ray-like effects in providing health, Arishta—resistant to insects, Sbeetal—cooling (cools the human system by giving relief in diseases caused by hotness, such as skin diseases and fevers), and Krimighana—anthelmintic. It was considered light in digestion, hot in effect, cold in property.

In earlier times, patients with incurable diseases were advised to make neem their way of life. They were to spend most of the day under the shade of this tree. They were to drink infusions of various parts of the tree or stem sap, if available, when thirsty, eat tender leaves as salad and cooked leaves as a vegetable. Young twigs were to be used for oral hygiene and gum as lozenges for dryness of the throat and to allay thirst. Whenever mature, ripe fruits were available, they were to be sucked for their sweetish tasty pulp.

Occasionally the seed were also swallowed. While on neem therapy, patients were to avoid all products of animal origin, such as egg, flesh, milk and alcohol, which were considered “hot” in nature. It was considered that the diseases for which neem was specified were caused by hot conditions prevailing in the body, and they were mitigated by the cooling effect of neem and other herbs.

The above observations led to the widespread use of neem, from the era of Carak Samhita (200 BC–200 AD) to recent times in the Indian subcontinent and adjoining countries. It became so popular in ancient India that some scholars believe that it was an ingredient of up to fifty percent of Ayurvedic preparations. Physicians at that time advised Panchang of neem, i.e. five parts of the tree, the leaves, bark, fruit, flower and root. With further development in Indian medicine, the raw single-plant-part therapy was replaced by compound preparations. In these, many ingredients were incorporated; some had a complementary effect, some had a supplementary effect, some were considered antagonistic to the deleterious effects of certain herbs and some were nutritive.

Neem preparations were prescribed in various forms: Churn—powder, Kwath—decoction, Khand—mixed with sugar, Kshara—alkali, obtained by burning the plant part, Vatika—pills, Asav and Arishta—fermented decoctions, Gbritam—butter fat extract, Malham—ointment, Dhupan—fumigant, Lep—poultice and Nasayam—nasal drops.
Ayurvedic Pharmacopoeial Products

Some of the important polyherbal neem preparations of the Ayurvedic Pharmacopoeia and their main uses are:

- **Aparjith Dhttp** — fumigant for purification of air (air sterilizer)
- **Erhamanisthadi kwath** — skin diseases
- **Dhattur tailam** — oil for skin diseases and muscular pain
- **Jatyadi tailam** — oil for ulcer
- **jeevanti adi Kashyam** — for smallpox
- **Laghu Manjishtadi kwath** — decoction for skin diseases
- **Kandavadu Lepah** — poultice for itching
- **Maha tikatam ghritam** — butter fat for skin diseases
- **Maha tikatam kashyam** — a bitter tonic
- **Naryana tailam** — oil for rheumatic disorders
- **Nimbadi Kashyam** — for skin diseases
- **Nimbadijatam palit** — for baldness
- **Palit Nasyam** — nasal drops for alopecia
- **Panch titktam ghritam** — butter fat for latent fevers
- **Panch nimba churnam** — powder for skin diseases
- **Patoladi kvath** — decoction for fevers
- **Phaladi kvath** — decoction for expelling worms
- **Purnavadi kshyam** — for skin diseases
- **Purnavadi kvath** — decoctions for swellings
- **Sudarshan churnam** — powder for fevers
- **Thiktakam ghritam** — butter fat for skin diseases
- **Thiktakam kashyam** — for skin diseases
- **Varanejatayidi ghritam** — butter fat for ulcers
- **Yograjaguggulu** — for rheumatoid arthritis

All the above are polyherbal compounds, having many herbs and a special method of preparation in each case, which may not be of interest to all readers. As an example, details of **Nimbadi kashyam** are given here to give some idea of these products. This is prepared by making a decoction of 50 gm each of **Azadirachta indica**, **Tinospora cordifolia**, **Zingiber officinale**, **Curcuma longa**, **Adhatoda vasica**, **Trichosanthes dioica**, **Solanum indicum**, **Terminalia chebula**, **T. belerica**, and **Phyllanthus emblica**. This herbal mixture is boiled in 8L of water, until reduced to 1L. Usually 60ml of this decoction is taken twice daily on an empty stomach.

As an Antimalarial

In the nineteenth century, European physicians practicing in India and Indian practitioners of the orthodox system of medicine (allopathy) found neem bark an effective therapeutic agent for fevers, particularly for malaria, which was very common in some parts with a tropical or sub-tropical climate. The powdered bark and fresh leaves were made official in the *Pharmacopoeia of India* (Dey, 1896). It was also
included in *Practical Materia Medica* (Clarke, 1900). In due course of time, during the first world war or thereabout, cinchona or salts of quinine were introduced into India from England but these could not reach most of Indian population living in remote areas. Neem bark was tried as a substitute for cinchona; it did not have any direct effect on the malarial parasite, yet patients obtained relief in most cases.

The use of a decoction of bark as an antipyretic is well known, particularly for malaria. For making the decoction, 15gm of bark should be boiled in fifty times that amount of water, until it is reduced to 50 ml. It is strained and then 10 ml of this filtrate is given thrice daily.

**In Venereal Diseases**

Before the arrival of the Portuguese and other European colonizers, venereal diseases were not known in India. These diseases, particularly syphilis, were noticed in some cosmopolitan areas but local physicians could not diagnose them as there was no mention of the symptoms of these diseases in their texts and they nicknamed them “*Firang rog*”, or foreign diseases. As per the concept prevailing at that time, the evident symptoms were due to *hotness* in the body and *impure blood*, as for other skin diseases like leprosy, eczema, leucoderma, etc. and they prescribed neem as a *cooling agent* and a *blood purifier*.

In Ayurveda, as mentioned earlier, use of all five parts of the tree together was considered the best, but later, on the basis of experience, physicians used one or two parts together, so details of each part are given separately here. Recently Lok Parampara Samvardhan Samithi has given a data base for neem, and the interested reader may consult the data base, in addition to the following details.

**Uses of Plant Part**

**Bark**

When peeled it has two different zones; the outer one is dry, scaly, darker in color, and the inner one is smooth and brown in color. For medicinal purposes, the inner portion of the bark should be used, preferably when it is fresh. Root bark is considered better, but is now very difficult to get, because of the damage that may be caused to the tree during collection. The outer portion of the bark is rich in tannins and is astringent, whereas the inner region is rich in secondary metabolites. The inner bark contains a bitter principle of a resinous nature, which when moistened emits the smell of sulfur compounds like those found in garlic.

Bark exerts a strong antimicrobial and astringent effect due to the presence of phenolic compounds and tannins, which have a strong healing effect on the skin. For this reason it is an ingredient of preparations for pimples, piles, wounds, bleeding gums, etc.

In the case of pimples, a decoction of bark with the pod of *Cassia fistula* is applied, while for piles, 3gm of bark powder with 5gm of cane sugar is administered three times a day and neem oil is applied externally. A tooth powder made from a fine powder of bark and alum (sodium aluminium sulfate) is used for spongy gums, particularly when they are bleeding. This mixture also has a good antiinflammatory effect.
The decoction of bark was also prescribed for plague.
In the nineteenth century, European physicians in India considered neem bark to be a bitter tonic and prescribed it in place of gentian or quassia, in the form of an infusion or tincture.

*Sap from the Tree*

It is said that after the tree reaches a hundred years of age, on a day which cannot be predicted, it begins to exude a nectar or sap from the crevices of the bark. The sap is thick, sweet in taste and fetid in smell. Great virtues are ascribed to this sap and it is said to be a panacea, particularly for leprotic ulcers and skin diseases of various etiology, particularly those which, as per the Ayurvedic concept, are associated with heat in the body.

*Gum*

The gum is very much like gum acacia in physico-chemical properties but is darker in color. It is a demulcent and is used for sore throats.

*Wood*

A pestle and mortar made from neem wood is preferred for pounding herbs. The wood is rubbed on a wet stone to form a paste. This paste is used for dressing wounds.

*Leaves*

The tender leaves were cooked like spinach in ancient India and sometimes fried in butter oil (the fat obtained after heating butter and discarding the fat insoluble part). To remove the bitterness from the leaves, they were boiled in water, and the leaves so obtained were cooked with some sour fruit like those of *Embelica officinalis*. Leaves in the form of an infusion, a decoction or as a chutney by grinding with black pepper were also used. These preparations were often prescribed for skin diseases, inflammation etc. and recently for diabetes. In the case of smallpox and measles, a decoction of neem leaves was given to quench thirst and to prevent dehydration. To make the surroundings of the patient aseptic and humid, the head was placed on a pillow made from neem leaves and small branches were hung all around and on the windows. In serious cases, instead of a decoction, leaf juice was prescribed and the whole body was smeared with a paste of neem leaves with or without neem oil. When pustules appeared, neem with liquorice (*Glycyrrhiza glabra*) root was made into a paste and applied.

For hemorrhoids, a paste of the leaves of neem and *Nerium mdicum*, when applied, caused the shrinkage of inflamed tissues. For ulcers, neem paste acted the same way.

Neem was also used for snake bites, particularly the bite of a russel viper. It was said to destroy venom, but the effect appears to be due to delay in the clotting time of the blood.

In gynecological practice, the use of neem is quite popular in post-parturition disorders. It induces labor by uterine muscle contraction. As an anti-inflammatory
and antiseptic agent, a decoction of neem is used for washing and douching the vagina. It induces the formation of milk, so was given for three days after delivery before meals. The same practice is followed by the dairy industry in some places in India even now. The effect appears to be hormonal because a neem branch is said to attract fish during the spawning period.

For oral hygiene and dental care, young twigs are cut into pieces 15–20 cm long; one end of this twig is chewed to form a fibrous brush and the teeth are cleaned with it. During this process the juice of the twig gets mixed with the saliva in the mouth and the mixture clears the throat by irritating the mucus membrane and causing the expulsion of phlegm from the throat.

This practice is present even now, and it is a common sight to see fresh neem twigs being sold in the marketplaces in India (Fig. 15A), but this is proving disastrous for neem plantations, where even young plants are being destroyed for their twigs (Fig. 15B).

In the Siddha system (the traditional system practiced in some parts of south India), dried and cured neem leaves are used. The curing is said to make leaves more effective, more palatable and less toxic. These dry leaves are called *Vaipilla*, and a well-known Siddha medicated oil preparation is *Vaipilla tailam*.

**Flower**

Flowers as an item of diet were used in India and the other east Asian countries as a spinach and chutney. The flowers are said to expel worms, give relief from coughing and are considered good for the eyes. In cataracts, a suggested formulation is to make a very fine powder of equal parts of neem flower and potassium nitrate. The mixture is applied to the eyes.

**Figure 15** *Azadirachta indica*, neem twigs. A, young twigs are being cut to have about 15 cm long and 0.5 cm thick pieces for chewing; B, young neem tree destroyed for the twig. (Photo by Gurcharn Singh)
Fruit and Seed

Unripe fruits are used in the same ways as leaves, but are considered useful for bleeding piles, worm infestation and urinary disorders. The ripe fruit is considered a blood purifier and anthelmintic and it makes the bowel soft.

For premature greying of hair, neem juice boiled several times in the juice of Eclipta alba is used as a nasal drop.

Oil

It was a common practice in some parts of south India to give a few drops of oil orally to infants regularly to keep them fit.

For skin diseases, the oil has always been considered a drug of choice and often used along with that of karanj (Pongamia pinnata) oil. It was applied on pustules, hard abscesses, obstinate types of wounds, leprotic lesions, ringworm, eczema and itch. It is particularly recommended for hair care problems such as psoriasis and dandruff, for killing lice and for giving relief in itching.

The oil destroys worms and may be useful in anal itching in infants caused by nematodes like the pin worm. It is a stimulative cerebral tonic and is used as a massage oil for rheumatism and joint pains.

Well-known Preparations of Neem

As given earlier, neem preparations have been prescribed for a very long time, but were not very much liked by patients because of their bitter taste. Recently, in some herbal patent preparations, neem extracts in the form of gelatin capsules are being sold, but the classical preparations are manufactured and supplied in the traditional way. The important ones are:

Nimba ghrit—general tonic in debility
Nimba asav—for fevers, particularly in malaria
Nimbadi anjan—antisepctic for eyes
Nimba arisht—tonic for building resistance in the body
Nimbadi kwath—decoction for fevers
Nimba Haridra—with turmeric and sugar, for throat problems
Nimbadi tail—oil for massage in dry eczema, leucoderma, and rheumatism
Panchnimba churn—for skin diseases like white patches, ringworm, etc.
Panchnimba gutika or Panchamrit—in leprosy and white patches
Panchnimba avleh—for skin diseases of different etiology, headache, diabetes and obesity
Panchtikata ghrit—in chronic skin diseases
Panchtikata ghrit guggal—for obstinate diseases, respiratory and heart problems

NEEM IN UNANI TIBB

In the Greco-Persian system of medicine (Unani tibb), which was patronized by Muslim rulers in the medieval era in the Indian subcontinent, the leaves and fruit were in the
pharmacopoeia. As per this system, neem is cold 1°, dry 2°, a resolvent and blood purifier (Nadkarni, 1954). Neem leaves, called “burgh-i-neem”, are said to expel foul wind from the body and heal ulcers in the urinary passage; it is an emmenagogue and good for skin diseases.

**Well-known Unani Tibb Preparations**

Neem leaves, bark, seed and oil are incorporated in some of the Unani preparations (Said, 1970), which are as follows:

*Arq Gaz*—a distillate from all five parts of the neem tree, used for fevers due to inflammation of the spleen.

*Arq Harabhara*—a distillate from the seed coat, a tonic for the lungs

*Arq Murakkab Musaffa khus*—a distillate, has a cooling effect on the body, used for purification of blood in venereal disease

*Hab Musaffi Khus*—blood purifier for boils, itching etc.

*Hab Narkachur*—anti-inflammatory for children

*Hab Bauvisir Bad*—for bloodless piles

*Hab Siyah Chatham*—for application inside the eyelid in conjunctivitis.

*Majun Juzam*—blood purifier for venereal diseases, leprosy etc.

*Marham Bauvisir Jad*—ointment for external application on piles

*Roghan Neem*—neem oil for external application on sores and wounds, and for killing ectoparasites like lice

*Zimad Bauvisir*—whole fruit powder, for application on hemorrhoids

*Zimad Mobasa*—for application on pimples, and other minor skin eruptions

**NEEM IN HOMOEOPATHY**

Neem under the name *Melia azadirachta* or *Melia Azadirachta* is well known in Homeopathy for its bark, called Margosa. It was included in the *Pocket Manual of Homoeopathic Materia Medica* (Boericke, 1927). According to Ghose (c. 1930), the tincture of bark was introduced by Dr P.C. Majumdar, after proving by him and his pupil. A full report on this proving was published in the *Indian Homoeopathic Review*. Later on, two more provings were made and published. It was also included in *New, Old and Forgotten Remedies* (Anshutz, 1930), *Pathogenesis de Azadirachta indica* (Lamasson, 1968), *Indian Homoeopathic Pharmacopoeia* (1971) and *Materia Medica of New Homoeopathic Remedies* (Julian, 1979).

The *Indian Homoeopathic Pharmacopoeia* has described both the macroscopic and microscopic character of the bark. As per the pharmacopoeia, the bark is dark grey to greyish black externally, while the inner surface is pinkish brown and fibrous. In old bark there is a well-defined outer rind formed of alternating strips of cork layers and dead secondary bast. The cork cells have reddish brown contents. Phellogen is not very distinct and a secondary cortex is normally not present. The secondary bast is formed of sclerenchyma. Medullary rays are 2–5 seriate.
In the *Indian Homoeopathic Pharmacopoeia*, the drug has been introduced on the authority of *Drugs of Hindoostan* by Dr S.C. Ghose. The mother tincture (one liter) can be prepared by taking 125gm of fresh bark (25 gm moisture and 100 gm solid matter), distilled water 375ml, strong alcohol 635ml. The fresh bark is pounded to pulp and macerated into alcohol. Potency 2x with dilute alcohol, 3x and higher with dispensing alcohol.

**Findings of the Symptoms**

A brief summary of the findings of the symptoms, as given by Ghose (c. 1930) are:

Mind: oppressed, forgetfulness, dull and loss of memory  
Head: giddiness, headache  
Eye: burning, dull and heavy, painful, red  
Ear: buzzing in the ear  
Face: flushing  
Stomach: thirst, appetite very acute  
Abdomen: great uneasiness with flatulence  
Stool: insufficient bowels, constipation, stool hard  
Genitourinary organs: great excitement of sexual organs  
Respiratory organs: very troublesome cough, sputa white  
Pulse: quick and hard  
Extremities: numbness of the limbs, burning of hands and soles  
Sleep and dreaming: sleeplessness, dreams of quarrels  
Fever: glowing heat and burning, copious sweat, itching

Julian (1979) has also given the symptomatology, according to which neem acts in active, depressed and forgetfulness states, right-sided headache, insomnia, thirst, constipation, fevers, tendency to miscarriage. The clinical diagnoses are senile dementia, amnesia, hypochondria, acute articular rheumatism, acroparesthesia, typhoid fever, recurring of miscarriage and metritis of the cervix.

**REFERENCES**


10. THERAPEUTIC INDICATIONS AND PHARMACOLOGICAL STUDIES

The effects of bitters on human systems are well known. Much of the earlier therapeutic efficacy claimed for neem may be due to these bitters. It was included in the Indian Pharmacopoeial List as a bitter preparation which acted as a sialogogue and a stimulant of various gastric secretions due to reflex action of the body against the bitter taste, which in turn increased the appetite.

In *Potter’s New Cyclopaedia of Botanical Drugs and Preparations* (Wren, 1907), neem bark is mentioned as an anthelmintic, cathartic and emetic, used for children in the southern states of America. It was prescribed in the form of a decoction, followed by purgation by castor oil. These and other studies, as given in the section on Ayurveda, Unani-Tibb and Homeopathy and in folklore medicine, led to research on the pharmacological activity of neem preparations on the various systems of the human body, and human pathogens. The details of these studies are given here. Use in veterinary practice has been given separately.

ANTHELMINTIC ACTIVITY

Keeping in view the reputation of neem as an anthelmintic, Cauis and Mhaskar in 1923 administered leaf juice to patients in a dose of 4 drams (14.25 gm) for expelling intestinal worms, followed by purgation, but the treatment was ineffective (Nadkarni, 1954); on the other hand, an Ayurvedic compound (Tyagi *et al.*, 1977) containing neem and long pepper (*Piper longum*) gave good results in ankylostomiasis. Singh *et al* (1980) carried out further clinical evaluation of the anthelmintic activity of neem.

Recently a large number of studies in plants have confirmed the nematodocidal effect of neem and it should be effective against nematodes in the human body, like round worm (*Ascaris lumbricoides*), threadworm (*A. vermicularis*) and pin worm (*A. oxyrus*). The activity of neem against pin worm in particular is of interest because the worm causes irritation and itching in the anal region, mostly in infants and children, in many parts of the world. For symptomatic relief, sometimes mustard oil is applied in the anal region, which may have an antifeedant property. Neem oil appears to be a better substitute for mustard oil for this purpose.

Neem was tried against the nematode *Ascaridia galli* (Shilaskar and Parashar, 1989). An Ayurvedic antifilarial compound with neem as a major ingredient has given good results against *Acanthocheilonema vitae in vitro* (Comley *et al.*, 1990).

ANTIBACTERIAL

Neem preparations have been used as a disinfectant since ancient times. With the discovery of antibiotics from the lower organisms, there was a move to look into

Prasad et al. (1993) reviewed the earlier literature and concluded that the extracts can significantly inhibit pathogenic microorganisms, with the most potent effect against Salmonella typhi, S. paratyphi and Bacillus subtilis, in vitro. The leaf and bark extract showed activity against Staphylococcus epidermidis, S. aureus, Bacillus spp. and Proteus spp. The activity against Escherichia coli was moderate (Ahmad et al., 1995). Garg et al. (1995), while looking for the antimicrobial spectrum for a neem-based cream developed by them, enumerated the following bacteria, on which the cream had a bactricidal effect: Bacillus subtilis, B. antibracts, Corynebacterium spp., Escherichia coli, Micrococcus spp., Proteus vulgaris, Staphylococcus aureus, S. citrus, S. epidermidis, S. lactitis, Salmonella paratyphi A and B, and S. typhi.

ANTIDIABETIC

Neem is one of the indigenous Indian drugs used in the treatment of diabetes (Mukherji, 1957). Shukla et al. (1973) observed that oral administration of 5 g of an aqueous extract or equivalent amount of dried leaves in capsules enabled patients to reduce their dosage of insulin up to 30–50 percent. Luscombe and Taha (1974) tried a 10 percent aqueous extract of dry young tender leaves for hypoglycemie effect in rabbits and a marked fall in blood glucose concentration was observed. The same effect was seen in fasting rats and guinea pigs. It appeared that antidiabetic activity was dependent on the functioning pancreatic beta cells, since neem leaves did not produce hypoglycemia in totally pancreatectomized rats or animals made severely diabetic by alloxan. Neem extract appeared to behave the same way as sulphonylurease. Murty et al. (1978) on the other hand, when they administered intravenous aqueous leaf extract to dogs, observed a significant decrease in blood glucose level in both normoglycemic and adrenaline induced hyperglycemic animals (Anonymous, 1979).

The effect of oral doses of an aqueous leaf extract, seed oil and nimbidin (the bitter principles from neem) was investigated by Pillai and Santhakumari (1981 a, 1981 b). The aqueous extract was found to be inactive but seed oil and nimbidin exerted significant activity. Sharma et al. (1983) also studied blood sugar level in hyperglycemic and diabetic animals. Management of diabetes mellitus by neem was discussed by Upadhyay (1984). Chakraborty and Poddar (1984) observed that a water extract of neem leaves exhibited a blood sugar lowering effect and this was
statistically significant as compared to hyperglycemic activity induced by streptozotocin (STZ). Obaseki et al. (1985) suggested that an aqueous extract of neem may have a therapeutic action via interaction with membrane. There was a reduction in serum acid phosphate activity and an increase of 5’ nucleotidase. Neem oil was also found to have a significant effect on lowering the blood glucose level of and hyperglycemic rats by Dixit et al. (1986). Neem as a synergistic agent to antidiabetic drugs was suggested by Bhargava (1986, 1987), Bhargava et al. (1986). Chakraborty et al. (1989) evaluated the activity of neem leaf extract for hypoglycemic activity in rats. El-Hawary and Kholief (1990) carried out studies on the biochemical effect of neem leaf. Dixit et al (1992) found that neem oil reduced blood glucose levels in normal animals as well as in alloxan-induced diabetic rats.

Chattopadhyay et al. (1993, 1993a) observed that neem leaf extract failed to improve muscle glycogen content and did not influence the effect of exogenous insulin. It did not affect the hepatic glycogen content of normal rats but reduced it in the case of glucose-fed hypoglycemic rats.

Literature on the antidiabetic activity of neem has been reviewed by Handa et al. (1989), Nat et al. (1991) and Prasad et al. (1993). According to Nat et al. (1991), the anti-diabetic mechanism of action may be due to the release of indigenous insulin in a way similar to that of sulphonyl urea. Prasad et al. (1993) put forth the view that neem only produced moderate-grade hypoglycemia. Extracts induced a significant fall in glycemic levels in a model of glucose-induced hyperglycemia in rats, had less effect on moderate alloxan diabetic rats but was ineffective in severe alloxan diabetic rats, probably due to the complete destruction of β cells by alloxan.

**ANTIFERTILITY ACTIVITY**

During pharmacological studies, Murthy and Sirsi (1958a) observed estrogenic activity in neem oil and its derivatives. Two derivatives of neem oil, sodium nimbininate and sodium nimbidinate, were found to possess a spermicidal effect *in vitro* in human beings and in rats (Sharma and Saksena, 1959, 1959a). Deshpande et al. (1980) gave a preliminary report on the male antifertility effect of neem in mice. Oral administration of an extract of crushed neem leaves for one month produced reversible antifertility activity in animals without inhibition of spermatogenesis. Chaudhry and Haq (1980) also observed this activity. Sinha et al. (1984) reported antiflagellate activity of the sperm by neem oil. The sperm became immobile and were not able to fertilize the ovum. Undiluted neem oil was found to possess a strong spermicidal activity both *in vitro* and *in vivo* in both rhesus monkey and man. Once the sperm touched the margin of oil it got completely entangled, leading to further sluggishness culminating in immobility within thirty seconds. All the animals that mated after the application of neem oil showed no sign of implantation. The application of oil did not produce any irritation or side effect. In females, the application of oil increased uterine contraction and there was suppression of ovulation, with an antiimplantation effect (Sinha et al., 1984a).

Reddy et al. (1984a) observed that neem oil affected the estrous cycle in albino rats; it had an antiovulatory and antifertility effect (Reddy et al., 1984b). Salunkhe
and Adsale (1985) considered neem oil as one of the agents for population control. Lai et al (1986) confirmed the antifertility effect of neem oil in female albino rats by intravaginal and oral routes. To study the mechanism of the antifertility effect, a freshly prepared aqueous extract of crushed neem leaves was fed to rats by Mateenuddin et al (1986). There was no significant loss or gain of weight in rats due to this extract. The absence of an open vagina, epithelial cells in the vaginal smear and incomplete development of the uterus indicated that there is no oestrogenic activity in neem leaf extract.

Subcutaneous administration of 0.2 to 0.3ml/kg of neem seed oil for 1–5 days postcoitum terminated pregnancy in rats (Tewari et al., 1986). These authors, on the basis of further biochemical and histological studies of the reproductive organs in cyclic and ovaricotomosed rats, confirmed that the postcoital contraceptive effect of neem oil was non-hormonal. It appeared that the active compounds of neem exerted the antifertility effect by absorption by the vaginal tissue and later on by circulation in the blood (Riar et al., 1988).

The non-hormonal nature of neem oil was further confirmed by Prakash (1986) and Prakash et al. (1987, 1988). Sharma et al. (1987) also observed antiandrogenic properties of neem seed oil in both males and females. The testes of rats treated with neem oil showed that the oil caused a reduction in their weight, arrested spermatogenesis and brought about severe degenerative changes in the cauda epididymus with a decline in protein, acid phosphate, etc. The effect appeared to be antiandrogenic, without any estrogenic, anti-estrogenic or progestinol activity. It did not interfere with the action of progesterone. In females it acted by causing damage to the uterine histological structure, especially breakage of the endometrium (Prakash et al., 1988).

The antifertility effect of the oil appeared to be due to:

1. A spermicidal or ovicidal effect so that the zygote is not formed.
2. If the zygote is formed, neem oil is blastocidal; a dead blastocyte cannot get implanted in the uterus.
3. Even if the blastocyte is formed, it is not able to establish itself because of the denatured endometrial linings of the uterus.

Further studies were carried out by Prakash et al. (1991, 1991 a) to study the effects of the ethanolic extract of the seed on female rats. Bardhan (1991) tried oil in rhesus monkeys and found a good spermicidal activity after pre- and postcoital application of the oil intravaginally, while Garg et al. (1992) carried out research on spermatogenesis by solvent-extracted, water-washed neem seed cake.

Upadhyaya et al. (1990) put forth a concept of immunocontraception, according to which a single intrauterine treatment led to a long term antifertility effect in female rats. The effect lasted for a variable period of 107–180 days and was reversible in nature.

Nat et al. (1991), after a review of the literature, concluded that the antifertility activity reported both for neem seed oil and leaves pointed at limonoids, which occur in both plant parts and are the active constituent. The existing relationship between the structure of some steroids and triterpenoid of neem suggests hormonal regulation of fertility control, but the non-hormonal mode may also be playing a part.
Shaikh *et al.* (1993) also confirmed the antispermatogenic activity of neem but some other studies have shown that use of neem oil as a contraceptive is not safe but has serious side effects. Sampathraj *et al.* (1993) confirmed marked structural alterations in the testes by administration of neem oil. Spermatogenesis was drastically impaired with an adverse effect on the testicular function and also as the function and integrity of the epididymus. Parshad *et al.* (1994) studied the effect of neem by oral administration. It resulted in a significant decrease in the total testosterone, bilirubin, and K⁺ in serum. There was no cytotoxicity.

Keeping the above side effects in view, Garg *et al.* (1993), at the National Institute of Immunology in India, looked for the safe fractions from neem oil for contraception. Garg *et al.* (1994) investigated the hexane extract of neem seed. Paranjape and Paranjape (1993) tried neem oil suppositories for contraception at the pre-coitus stage. The preparation was not irritating to the vaginal mucosa or the male genital organ, and was very safe.

Kasturi *et al.* (1995) studied histological and biochemical changes in the caput and cauda didymus of albino rats treated with dry leaf powder in various doses for 24 days. The effect was dose dependent. The serum testosterone concentration in the animals treated with high doses decreased significantly, suggesting a possible antiandrogenic property. Neem seed oil was applied by 77 human fertile females before coitus. It covered 6341 menstrual cycles. There was no contraception in any case (Tyagi and Sahrawat, 1995). The safe fraction of neem oil developed at the National Institute of Immunology in India for contraception was tried by Mukherji *et al.* (1995, 1995a) by oral administration. All the embryos without exception were resolved. The treatment was well tolerated with no mortality or morbidity. Animals regained fertility in subsequent cycles. Talwar *et al.* (1995) in the same institute developed a polyherbal cream and pessary, containing extract of neem, quinine hydrochloride, and soap nut extract (*Sapindus mukrosii*). The formulation showed a good spermicidal action *in vitro* and a high contraceptive efficacy in rabbits and monkeys. An acute and sub-acute toxicology study of this cream was carried out and no change in any of the hematological and biochemical parameters compared to pre-treatment values has been observed. Autopsy of animals revealed no gross changes in the tissues (Garg *et al.*, personal communication). This cream is now being subjected to multi-centric clinical trials.

**ANTIFUNGAL**

In the case of infectious diseases, it is a common practice among some people to use neem with other ingredients as a fumigant. To find out the significance of this, Murthy and Sirsi (1957), Naryana (1965), David (1965) and Jain and Pathak (1970) studied anti-fungal activity. Upadhyay and Arora (1975–76) studied the sporostatic nature of neem smoke and its possible ecological influence on the air fungal flora. Some people hang leaves around the patients. The curative effect may probably be due to essential oils. To confirm this, Satyanaryana and Rao (1977) and Thind and Dahiya (1977) tried the essential oil obtained from neem leaves against keratinophilic fungi. Singh and Sharma (1978) and Chary *et al.* (1984) also studied antifungal activity.
An account of raw materials from neem, which can be used against fungi pathogenic to man, has been given by Khan and Wassilew (1987). and Tripathi (1987) confirmed that the bark and leaf extracts inhibited spore formation and were toxic against *Epidermopyton floccosum Microsporum canis*, and *Trichophyton mentagrophytes*, but Singh *et al.* (1987) failed to demonstrate any antifungal activity of two fractions of leaf, extracted with acetone, against a variety of human pathogens. Khan *et al.* (1988) also observed that neither dried neem material nor medicinal preparations containing neem or its oil had any effect on fungal growth. Petroleum ether extract showed some activity, which may be due to quercetin—a flavonoid.

Khan *et al.* (1991) also studied the effect of petroleum ether extract of neem on the fungi pathogenic to humans. Iyer and Williamson (1991) concluded that neem extract inhibited the protease activity of *Trichophyton* spp.

Prasad *et al.* (1993), on the basis of earlier reports, concluded that neem extract exerted varying degrees of fungal toxicity which may be due to volatile sulfurous compounds or the limonoid gedunin. Fabry *et al.* (1996) also studied fungistatic and fungicidal activity.

ANTIINFLAMMATORY AND ANTIPYRETIC

In Ayurveda, Nimbadi Kashyam (a decoction of neem with other herbs) is prescribed for various inflammatory conditions. Other neem preparations are also used as a poultice for external application in gout, rheumatism, arthritic pains, etc. Neem oil was found to be an antipyretic and effective against rheumatism by Murthy and Sirsi (1958). David (1969) studied the antipyretic activity of neem oil, while Bhargava *et al.* (1970) worked on its antiinflammatory activity. Lorenz (1976) tried neem bark extract in inflammatory stomatis and David (1978) the effect of neem oil and its constituents on cotton pellet inflammation. Shankarnaryana (1978) used two crystalline bitter principles, nimbin and nimbinin, obtained from oil and found that they were comparable to cortisone in their action. Crude neem oil, sodium nimbidinate and nimbidol also showed considerable antiinflammatory activity. Pillai *et al.* (1980) observed that nimbdin 100mg/kg showed a significant analgesic and antipyretic effect in mice and rats which was comparable to acetylsalicylic acid and pethidine hydrochloride. Further studies by Pillai and Santhakumari (1981) indicated that nimbidin reduces acute paw oedema significantly in rats and also suppressed formalin-induced arthritis of the ankle joint and fluid exudation in croton oil-induced granuloma in rats. The authors were of the view that bitter from the neem may be suppressing oedema formation by reduction in vascular permeability and was not mediated through adrenals.

An antiinflammatory (rat paw oedema) and fairly good antipyretic effect (pyrogen-induced hyperpyrexia) with 75 percent methanolic leaf and bark extract in rabbits was observed by Okpanyi and Ezeukwu (1981). Two patents were granted to Terumo Corporation of Japan for anti-inflammatory polysaccharides from the bark in 1983 and 1985. Fujiwara *et al.* (1982, 1984) also reported inhibition of caragenin-induced oedema after oral administration of polysaccharides isolated from bark. Khatak *et al.* (1985) obtained an antipyretic effect from various fractions of 90 percent ethanolic leaf and twig extract. Tidjani *et al.* (1989) observed antiinflammatory activity from
chloroform extract when administered orally and also applied topically. Tandon et al. (1990) reported that the ethersoluble fraction of ethanol extract of leaves showed a good analgesic activity in pain but did not have any anti-inflammatory effect. Handa et al. (1992) and Vohora and Dandiya (1992) have reviewed the literature on antiinflammatory activity. Chattopadhyay et al. (1994) have also discussed it.

According to Bray et al. (1990), relief due to neem preparations in inflammation, pain, and fevers may be due to interference with hormonal regulation, interaction with receptors and alternation of membrane permeability and integrity. Nat et al. (1987) postulated the view that the inhibition of calcium activation and chemiluminescence in vitro correlates with the antiinflammatory and antirheumatoid effects. While continuing this study, Nat et al. (1991 a) identified the compounds responsible for the inhibition of the chemiluminescence product by activating human polymorphonuclear leukocytes. These compounds are gallic acid (+) gallocatechins, (-) epicatechins, and (as a 2:1 mixture) (+) catechin and epigallocatechin. Commercial samples of gallic acid, (+) catechin and (-) epicatechin showed the same effect. Kroes et al. (1992) confirmed the antiinflammatory activity of gallic acid.

Khanna et al. (1995) studied the mechanism involved in the antinociceptive action of neem. The results suggested that both central and peripheral mechanisms and complex pathways of both opioid and non-opioid are involved.

ANTIPROTOZOAL

Neem has been tried against the malarial parasite and *Trypanosoma*.

**Malarial Parasite**

Neem preparations, particularly bark, have been esteemed for malaria for a very long time. Studies were conducted to see if it could be substituted for cinchona bark. A compound of margosic acid and its sodium salt was prepared from the oil and on the basis of clinical trials was found to have activity against protozoa, which was onetenth that of quinine sulphate (Nadkarni, 1954). Murthy and Sirsi (1958b) experimented on avian malaria. In recent times, neem has been investigated in detail for its anti-malarial activity but the results are conflicting. Tella (1976) found oral doses of neem preparations quite ineffective but Ekanem (1976) demonstrated the schizonticidal activity of these against *Plasmodium berghei berghei*. Nimboloid, a terpenoid lactone, inhibited *P. falciparum* in culture with a moderate potency (Rochanakij et al., 1985). Obaseki et al. (1985) studied the effect of aqueous extract in rats and suggested that it may elicit its therapeutic action via an interaction with membrane. Obih and Makinde (1985) administered neem extract subcutaneously into mice after inoculating with a chloroquinresistant strain of *P. berghei berghei*. There was appreciable suppression of parasitemia. Bray (1985) studied the antimalarial activity of some limonoids.

Abatan and Makinde (1986) screened solvent fractions obtained from leaves for anti-malarial action, using *P. berghei berghei* as a parasite in mice. Statistically significant suppression was seen after four days of oral dosing. Obaseki and Judge-Fadunsin (1986) administered aqueous extract of neem to albino mice. It proved effective against acute infection by *P. yoelli nigeriensis*. The flavonoids and limonoids,
gedunin and quercetin isolated from neem were found to be highly active against the malarial parasite (Khalid et al., 1986; Khalid and Duddeck, 1986).

When Iwu et al. (1986) studied levels of some enzymes in rat liver microsome after administering the water-soluble part of methanolic extract, a significant inhibition of nicotinamide adenine dinucleotide phosphate (NADPH) cytochrome C (P/450) reductase activity was observed. It also stimulated the oxidation of hemoglobin and glutathione. The leaf extract and seed extract of neem were found active against chloroquin-sensitive and resistant strains. The extracts were non-toxic.

The potential of natural plant products for treating major protozoal diseases, with reference to neem and other trees, was reviewed by Phillipson and O’Neill (1989). These limonoids were tested for their in vitro antimalarial resistant K1 strain of *P. falciparum*. Out of all these compounds, gedunin was three times more active than chloroquin. Vasanth et al. (1990) studied neem as one of the antimalarial plants, while Üdeinya (1993) looked into the antimalarial activity of neem leaves. Jones et al. (1994) observed that the sexual development of the malarial parasite is inhibited in vitro by neem extract, azadirachtin and its analogs. Iwalewa et al. (1995) studied cardiogenic glycosides and sterols from neem on *P. yoelii nigerensis* for antimalarial activity. These compounds exhibited high prophylactic, moderate suppressive and very minimal curative properties. Cardiac glucosides were more effective than sterols.

The antimalarial activity of neem appears to be due to other mechanisms exerted on the body, in addition to that on the malarial parasite. According to Okpako (1977), neem extracts have been shown to break the periodic sequence of recurring fevers because of its prostaglandin synthetase inhibitory action. The body of the patient mobilizes the immune system to suppress the parasite. Badam et al. (1987) and Benoit et al. (1996) found an antimalarial effect of both leaf and seed in vitro. Bray et al. (1985, 1990) also considered the beneficial effect in malaria due to anti-inflammatory and immunomodulator activity. Nat et al. (1987), after their study, concluded that it may be due to immunomodulator activity in bark and an antipyretic effect in other parts of the tree (Nat et al., 1991).

**Trypanosoma**

Neem was tested against Chagas’ disease, which spreads in endemic form in some parts of Latin America. García (1989) studied the effect of azadirachtin on the development of *Trypanosoma cruzi* a flagellate protozoa found in the blood, and a causative agent of Chagas’ disease and its vector *Rhodnius prolixus*. It affected the hormonal balance of host (Azambuja, 1991) and caused growth inhibition of the parasite. Nok et al. (1993) confirmed the trypanocidal effect of neem leaf extract.

Two other diseases, tsetse fly and sleeping sickness, are caused by *T. brucei* and *T. gambiensis* respectively. Neem should be active against these parasites also.

**ANTITUMOUR**

Some earlier reports (Chatterjee, 1961; Anonymous, 1980) indicated the anticancerous activity of neem. Mitotic inhibition activity by the leaf extract was observed by Yadav and Rathore (1976). In vitro activity against sarcoma 180 ascites tumor by intra-
peritoneal administration of polysaccharides from the bark was reported by Fujiwara et al. (1982, 1984), while Pettit et al. (1983) studied the effect of limonoids against the P-388 lymphocytic leukemia system. Out of the limonoids investigated, $1\beta, 2\beta$ diepoxyazadirdrone and 7-acetylneo-trichlenone appear to be active. Antitumor polysaccharide (code-name N9C1) which, when administered to mice, markedly inhibited the growth of sarcoma 180 was patented by Terumo Corporation in 1983 and 1985. In the first patent, organic solvents were used to get the extract but in the second, neem bark was treated with water at a temperature between 0° and 40°C. The residue was subjected to a purification process by alcohol precipitation.

In their review article on neem, Prasad et al. (1993) mentioned another patent for antineoplastic action. Its usefulness for mouth cancer and its cytocidal effect on malignant cells have been documented. In another report, it was mentioned that azadirachtin had an immediate effect on the rate of cell proliferation, including its intensity. This was due to rapid inhibition of RNA synthesis along with inhibition of the synthesis of protein and DNA. Takeya et al. (1996) and Cohen et al. (1996) have reported the cytotoxic effect of limonoids.

ANTIULCEROGENIC

Nimbardin, a mixture of neem bitters, was studied by Pillai (1978) for antigastric activity and was found to be effective. Pharmacological studies showed that nimbardin blocked the stimulatory effect of acetylcholine and also inhibited the stimulation produced by histamin. It partially blocked the action of nicotine on the arterial blood pressure of dogs under the effect of anesthesia. All these findings suggested the antigastric activity of nimbardin. Nimbardin, when given orally, significantly prevented Shay ulceration. These observations were confirmed by Pillai and Santhakumari (1984), and further studies showed the significant protective effect of nimbardin in a dose of 20–40mg/kg. It afforded a remarkable protection in duodenal lesions. Oral and intra-peritoneal administration of nimbardin did not produce toxic manifestation or foetal abnormality (Pillai and Santhakumari, 1984a).

Koley et al. (1994) studied the chronic ulcerogenic effect of alcohol extract of neem leaf on the gastric mucosa of rats. The lack of irritation and promising antiinflammatory effect suggested its use as an antiulcerogenic or as an antipeptic ulcer drug. Garg et al. (1993) and Devdas et al. (1995) also observed the gastric antiulcer effects of neem leaves.

Nimbatiktam, a bitter principle of neem containing terpenic ester (nimbdin), was administered at the rate of 30 mg daily. It showed a significant ulcer-healing effect. Endoscopy healing was 87 percent for duodenal ulcer and 100 percent for gastric ulcer, with significant improvement from the third week of medication. No side effects were observed (Pillai, 1995).

ANTIVIRAL

Rao et al. (1969) noted that ten percent water extract of tender leaves possessed antiviral activity against vaccinia and variola viruses. Rai and Sethi (1972) found that an aqueous extract of neem leaves did not kill the virus directly but inhibited the
multiplication of *Vaccinia* and fowl pox virus. They suspected that this effect may be due to limonoids or flavonoids. Babbar *et al.* (1982) also found antiviral activity in neem leaves. Reddy and Sethi (1984) studied, *in vivo* the antiviral effect against *Vaccinia* virus while Wagh (1988) used it in clinical trials in the case of viral hepatitis.

Out of 15 viruses studied by Gogate and Marathe (1989), only three, *Chikungunya* (634029), *measle* (ED/3) and *Vaccinia* virus, were inactivated by neem; others were affected to various degrees. Praneem polyherbal cream, which has been developed by the National Institute of Immunology in India and in which neem extract is the main ingredient used, has been found to be effective in the treatment of human papillom virus-16, which infects the female genital tract (Talwar *et al.*, 1995).

The above antiviral effect of neem in some cases justifies the use of neem leaves, particularly in smallpox in the Indian subcontinent.

**CARDIOVASCULAR EFFECTS**

Thompson and Anderson (1978) studied a crude extract of neem on the cardiovascular system. The effects included profound hypertension and a minimal negative chronotropic effect, which increased at higher doses. The rise in arterial blood pressure with low doses of the extract supports the suggestion that it may have a biphasic effect on arterial blood pressure. The vasodilation remained somewhat persistent. Ilesanmi *et al.* (1988) also found cardioactivity in rats with neem leaves.

**DIURETIC EFFECT**

Bhide *et al.* (1958) found diuretic activity in sodium nimbidinate. It was seen by Shah *et al.* (1958), who carried out clinical trials with it in nine cases of congestive heart failure, that patients obtained relief due to the diuretic action of neem preparations. There was no local discomfort, no toxic effect and the diuretic effect continued for a few days. Luscombe and Taha (1974) also confirmed that neem leaves have a mild diuretic property as observed in water-loaded rats. Singh *et al.* (1987), using oral administration of neem leaf, observed a significant reduction in blood pressure as well as heart rate but no diuretic activity.

**EFFECT ON CENTRAL NERVOUS SYSTEM**

Debelmas and Hache (1976) tested the aqueous extract on the central nervous system (CNS). No significant effect was observed. No anticonvulsant, anticholinergic, analgesic, or sedative effect could be demonstrated, but Pillai *et al.* (1980) showed an antistress property. Pillai and Santhakumari (1984a) observed a mild suppressive effect on the CNS functions of mice with nimbidin (the bitter principle of neem). Singh *et al.* (1987) noted that acetone extract of leaf had a variety of effects on mice which were attributed to CNS depression and to an effect on the autonomic nervous system. Dandiya (1990) also considered neem as one of the herbs acting on CNS.
In a continuation of their early study, Singh et al. (1987) worked on two fractions of acetone extract of neem leaves and observed a CNS-depressant activity. Jaiswal et al. (1994) mentioned an anti-anxiety effect (anxiolytic) of leaf extract in rats.

While summarizing the earlier findings, Nat et al. (1991) put forth the view that the active constituents from the leaf extract showing CNS activity may be limonoids, because most of these compounds are sufficiently lipophilic to cross the blood brain barrier. The acetone extract displayed a much stronger activity than an aqueous leaf extract. According to Nat et al. (1991), considering the antihistamine properties of nimbidin, interference with CNS neurotransmitters may be possible.

HEPATOPROTECTIVE EFFECT

Ayurvedic polyherbal preparation “Nimbawagadi kashyam”, containing neem bark and other herbs, is specific for jaundice. Oomacham et al. (1990) reported neem as one of the plants used for the treatment of jaundice. To find the hepatoprotective action of neem, Chattopadhyay et al. (1992) studied serum levels of glutamate oxaloacetate transaminase (GOT), glutamate pyruvate transaminase (GPT), acid phosphate and alkaline phosphate to elucidate liver damage. The levels of these enzymes were elevated after 24 hours of paracetamol treatment, but rats fed with neem extract prior to addition of paracetamol to their diet showed a much lower serum level. The hepatoprotective activity may be due to 6 flavanol-o-glycosides.

IMMUNOMODULATORY PROPERTY

The antistress property of nimbidin, a compound isolated from neem oil, was noted by Pillai (1978). In an initial screening of Sri Lankan plants in vitro neem bark was found to have an anticomplementary effect (Nat et al., 1987), which exerted a dose-dependent effect on immunomodulator activity. A dose-dependent decrease in the chemiluminescence of polymorphonuclear leukocyte was observed along with a dose-dependent increase in the production of the migration inhibition factor by lymphocyte. When the immunomodulator property of aqueous extracts on the human humoral and cellular defense mechanism was investigated, the extracts decreased both the classical and alternative C pathways. Labadie et al. (1989) reported the immunomodulator activity of a compound isolated from neem bark on the basis of data collected from various sources. Nat et al. (1989) characterized the two-polymer anticomplement compound as peptide glycans. The carbohydrate part consisted of glucose and arabinose. The protein content was 5.5 to 9.8 percent. Compounds responsible for the inhibition of chemiluminescence production by activated human polymorphonuclear leukocytes were isolated and identified from the crude aqueous bark extract. The compounds were gallic acid (+) gallocatechin, (-) epicatechin, (+) epicatechin and epigallocatechin (Nat et al., 1991a).

An Ayurvedic preparation, Nimbarishta—a potion prepared by fermenting Woodfordia fruticosa flowers, neem bark and the other herbs—was studied for its
immunological property by Kroes et al. (1990). These authors modified the preparation process of Nimbarishta to see the effect on complement activity and functioning granulocytes. It was seen that a long boiling time considerably reduced, whereas the addition of flowers of *Woodfordia fruticosa* considerably increased, its immunomodulatory activity (Kroes et al., 1993). Gopal Raj (1993) mentioned that neem boosts the body’s defense. Sen et al. (1993) observed the adaptogenic effect. Upadhyaya et al. (1993) studied extract of leaf bark and seed. The result indicated that neem was an immunostimulant; it selectively activated the TH-1 component of the lymphocyte population to elicit an enhanced cell-mediated immune response. *In vitro* HIV antieffects were observed. The addition of neem leaf extract significantly reduced the secretion of P-24 viral protein into culture supernatant. Bark extract was most effective. It also induced *in vitro* production of IL-1 interferon. Garg et al. (1993) felt that lipid terpenoid association was responsible for immunomodulation activity. Sen et al. (1993) evaluated neem on some biochemical parameters in normal and stressed rats and found changes comparable to the antistress agent dopamine. Upadhyaya and Dhawan (1994) observed that an aqueous extract of leaf, bark and seed enhanced the phagocytic activity of macrophages.

**USE IN SKIN DISEASES**

On the basis of the earlier studies, Nadkarni (1954) gave an account of various neem preparations used for skin diseases. One of these was sodium and potassium margosate, derived from margosic acid, isolated from neem oil. It was found to be a disinfecting agent for skin infections and was used for dressing in place of carbolic acid. Its major use was for external application in tetanus, leprosy, urtica, eczema, eryspelas, scrofula, ringworm, scabies, etc. but intramuscular administration was more effective. For leprosy, it was used alone or mixed with *chaulmoogra* oil (*Hydnocarpus kurzii*). Sodium margosate was administered subcutaneously and intramuscularly in the primary, secondary and tertiary stages of syphilis but it did not compare well with heavy metal preparations available at that time (Nadkarni, 1954).

Nimbudin, the bitter principle from neem, was found effective in furunculosis and arsenical dermatitis (Singhal and Mudgal, 1983).

An oil called Eth Ennei in the Sidha system of medicine, prepared by boiling nux vomica (*Strychnos nux vomica*) seed in neem oil, gave good results in eczema (Masilamani et al., 1993). An extract of dry leaves prepared with 70 percent alcohol was dissolved in propylene glycol. This preparation was applied in chronic skin diseases such as eczema, both acute weeping and acute chronic, ringworm and scabies. Most of the cases were of an obstinate type, treated earlier with well-known preparations like salicylic acid, benzoyl benzoate, sulfur, etc. Singh et al. (1979) got very encouraging results from this preparation.

A neem-containing formulation for hair was patented by Latiff (1979). The effect of nimbudin on psoriasis was studied by Rajasekharan et al. (1980). Shrivastava and Singh (1982) carried out research on the activity of neem against dermatophytes.

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*Tinea pedis* is a chronic fungal infection, occurring in between the toes and characterized by shedding macerated skin, and is caused by *Trichophyton mentagrophytes*. Rai and Upadhyay et al. (1988) investigated the effect of stem bark on this disease and observed a significant growth inhibition.

The effect of petroleum ether extract of neem leaves on fungi was reported by Khan et al. (1991) while Iyer and Williamson (1991) reported that neem inhibited the protease activity of *Trichophyton* spp. Charles and Charles (1992) tried a mixture of fresh neem leaves and turmeric powder in the proportion of 4:1 by weight for the treatment of scabies caused by *Sarcoptes scabei*. It was a very safe, economical treatment and 94 percent of the patients were cured by it. A preparation with neem bark as the main ingredient was tried for leprosy by Subramanian and Lakshmanan (1993). The activity was comparable with modern anti-leprotic drugs. Srimathi and Murthy et al. (1993) studied a herbal preparation for scabies.

Conrick (1994) gave a brief account of skin diseases where neem can be used. Nat et al. (1991) in their review article after studying the literature, concluded that the immunostimulating property of neem may possibly be the reason for the recovery of patients with skin diseases.

**USE IN VAGINITIS**

Inflammation of the vagina due to genital tract infections in females is fairly common both in the developed and underdeveloped countries. The symptoms may vary from mild irritation to profuse leukorrhea and treatment varies from douching with whey (yogurt) to a broad range of antibiotic and antifungal treatment.

Garg et al. (1995) studied a purified fraction of neem in a cream base to assess the clinical efficacy of leukorrhea caused by *Chlamydia trachomatis* by applying cream vaginally one hour before going to bed. The formulation proved highly effective in clearing infection from the cervico-vaginal region. Upadhyaya and Dhawan (1994), when they treated supernatants of mouse spleen cells with neem, observed an inhibition of the intracellular multiplication of *Chlamydia*. Another polyherbal preparation widely tested for its antifertility effect as a cream and as a pessary by Talwar et al. (1995) had an antimicrobial effect against a wide range of pathogens of the vaginal canal, including *Candida albicans* and *Gardnerella vaginalis*. The cream was effective in the treatment of human papilloma 16 virus infecting the female genital tract. The other creams based on neem used by Usha Rani et al. (1994, 1995) were also effective for vaginal infections.
WOUND-HEALING (MUSCLE REGENERATION) PROPERTY

A polyherbal cream containing neem bark was tried by Bhatt and Koro (1984) on burning wound sepsis, bacteria and fungi. Thaker and Anjaria (1986) and Bhargava et al. (1986a) also found a wound-healing effect of neem, which according to Tandon et al. (1988) may be due to increased cutaneous capillary permeability at the site of the wound. Goget (1991) treated traumatic wound patients with neem fumigation.

Sathyanaryana et al. (1994) evaluated the wound-healing activity of neem oil ointment after 24 hours of wounding. At the end of the fourteenth day, the animals treated with 1 and 2 percent neem oil ointment showed complete epithelization. Hair growth was faster as compared to the other treatments and control. Sohni et al. (1995) studied septicemia in rats by using a traditional preparation containing neem, which was prophylactic against *Salmonella typhi*. When studied for safety (Tandon et al., 1995), neem oil was without any side effects on the liver and kidney. It was non-irritating to skin and safe for external use. It showed very low toxicity when given orally, with no histopathological changes.

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11. IN VETERINARY PRACTICE

TRADITIONAL USE

It is a common practice to apply neem oil alone or along with cedar wood oil externally to cattle, for any type of skin disease of any pathogenicity and even on wounds. Sometimes the animal is also made to drink the oil. It is said that neem oil aids in healing the skin, and thus gives relief to infestation. While grazing in marshy areas, the hooves of cattle often get septic. In this case, the hoof is washed with a decoction of neem and dressed with neem oil; 20–30 ml of neem oil is administered daily.

The above use of neem oil has been found useful by modern veterinarians also, and experiments have been conducted with neem oil or its compound preparations.

FOR SKIN DISEASES

Vijayan et al. (1987) prepared Oil Bordeaux from copper sulfate, quick lime and neem oil. It was administered in doses of 4 ml by intramammary infusion for 7 days. Most of the cases of mastitis recovered. Neem oil was also tried in calves, experimentally infected with the protozoa Theileria annulata (Srivastava et al., 1987). Antimicrobial activity was observed in a veterinary herbal antiseptic cream containing neem (Pandya et al., 1991). Neem oil was found effective in healing wounds in calves (Bhargava et al., 1991) and in camels (Purohit and Chauhan, 1992). In camels the healing process was evaluated by clinical observation, percent healing, histopathological and histochemical examination and biochemical analysis of the biopsy of specimens. The dressing material containing neem enhanced tissue repair. Anil Kumar et al. (1993) studied similar tissue-repairing activity in buffaloes.

Neem preparations have been found effective for various ectoparasitic insects. In demcodectic mange of dogs, caused by a mite, a lotion with neem soap gave very good results (Tripathy et al., 1988). A compound herbal preparation containing Cedrus deodara, Azadirachta indica, and Embelia ribes was tried against common poultry lice, Mennopon gallinae and Liperus caponis. This preparation caused 100 percent mortality of lice (Das et al., 1993). Another commercial preparation with the same herbal ingredients controlled canine dermatitis caused by Demodex canis and Sarcoptes spp. Hair appeared after 24–28 days and there were no symptoms of toxicity (Das and Bhatia, 1993). It was also effective in canine demodecosis in dogs with severe cutaneous lesions around the ear, neck and head, skin encrustation and pruritus due to D. canis (Das, 1993). In sarcoptic mange of goat, when the same preparation was sprayed, no mites (Sarcoptes scabiei var. caprae) could be found (Das et al., 1994). Adverse effects of neem preparations were observed on ticks (Williams, 1993).

For non-conventional treatments, Heath et al. (1995) applied azadirachtin to a biting louse (Bovicola ovis) on sheep. Azadirachtin was found quite effective as compared to the other synthetic compounds. The treatment was cost-effective in
reducing louse members on the sheep for at least 40–50 days. The lack of persistence as compared with conventional insecticides was the only apparent drawback.

AS A CONTRACEPTIVE

One of the important potential uses of neem oil is its contraceptive effect. It is a spermicidal (Yao, 1993), antifertility agent and abortifacient, but could not be used in human populations because of its side effects; however, this oil can be of immense use in sterilizing stray animals by mixing in a bait for common domestic pests, rats, mice, etc. By using neem treatment we can get rid of unwanted mammals without cruelty.

REFERENCES


The use of neem in skin diseases lead to its application on preventive aspects also. Taking a bath in a decoction of neem leaves was a ritual in some societies. The anti-inflammatory properties of neem preparations made their use more popular.

As given in Chapter 9 on Traditional uses, the neem twig is well reputed for oral hygiene, neem oil, extract or fibers have been incorporated in some of the recent toothpastes and a floss has also been prepared. Neem soap is quite popular in India and its use is also spreading in the western world. Neem extract is an important ingredient of some herbal shampoo, and neem oil is used in hair oils, body lotions, creams and mosquito repellent preparations. Neem oil is said to prevent baldness and greying of hair, and has anti-lice and anti-dandruff effects. Patents for these products have also been taken out (Sawanbori et al., 1977; Latiff, 1979).

Neem has been incorporated in face packs. A typical formulation may have a very fine powder of leaves, bark and seed in clay. Milan Mehtra (1997) has given some formulations incorporating neem for face packs for oily skin, hair oil and cream for cracks on the back of the heel. In face packs, neem has been mixed with Carica papaya which contains papain and with liquorice. The author has suggested that a bath oil based on neem can be applied immediately after swimming to remove the last traces of chemicals and salts left on the body. Neem-based gels can possibly reduce the amount of clothes required by people living in a cold climate. The antiseptic and emollient properties of neem lotion can be useful for minor skin diseases. Neem along with Tulsi (Ocimum sanctum) has been incorporated in prickly heat powder and body talc.

In India, it was a common practice to apply coryllium (lamp black) along the side of the eye, particularly by young ladies as a beauty aid, to make eyes conspicuous. The common method of making lamp black was to take an earthen lamp, and put neem oil and a cotton wick in it. When ignited, the wick liberated copious smoke, from which lamp black could be collected by placing a brass cup, containing water for cooling, some distance away from the flame. The lamp black deposit was scraped from underneath the cup, and mixed with a small quantity of mustard oil to form a thick paste, called kajal.

This carbon black can also be used as a very safe, temporary hair dye, for concealing grey hair, by forming a very thin film on it using a hand glove.

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13. TOXICOLOGY

INTRODUCTION

In Ayurveda and in Indian traditions, consumption of neem in one way or another is often prescribed and chronic patients were advised to make neem their way of life. The leaves have been fed to cattle, goats and camels as fodder. Birds have been reported to swallow the fruit. All these indicate the safety of neem to warm-blooded animals, but in recent times some of the studies have indicated that it is toxic beyond certain limits, particularly the oil. Ayurvedic preparations are usually polyherbal and neem is only one of the ingredients, used much below the toxic limits. Moreover, the herbs are processed extensively during the preparation of the product, which may reduce the toxicity further.

EARLIER REPORTS ON TOXICITY

Nadkarni (1954) mentioned that neem is a narcotic poison if used in large doses. It produces giddiness, dimness of sight, mental confusion, stupor, dilated pupils and steror. It also acted as a gastrointestinal irritant producing vomiting and purging. Nayar (1954) also included neem among poisonous seed.

The earlier toxicological studies were conducted on the major ingredient of neem bitter, the amorphous compound nimbidin, nimbidinic acid (Siddiqui and Mitra, 1945) and sodium nimbidinate (Bhide et al., 1958) which were not found to be toxic.

RECENT STUDIES

While studying the traditional use of neem in malaria, Okpanyi and Ezeukwu (1981) found that some patients developed side effects, which may be due to the toxic nature of neem preparations, but Pillai and Santhakumari (1984) observed that nimbidin in doses up to 100mg/kg did not elicit any foetal abnormality or adverse effects on rearing performance.

Qadri et al. (1984) tested a commercial preparation with 30 percent neem oil for subdermal toxicity in albino rats and found scaling of the epidermis and hyperkeratosis of the stratum corneum. No other adverse effect, even with doses of 200–600mg/kg, was observed. Another commercial preparation with 3000 ppm of azadirachtin also had a very low toxicity in rats (Jacobson, 1989).

Obaseki et al. (1985) studied the biochemical effects after the oral administration of leaf extract and found a marked increase in the activity of 5’nucleotidase, leading to hepatobiliary toxicity, but Khatak et al. (1985) found no significant acute or sub-acute toxicity of various fractions up to 1.6gm/kg. Singh et al. (1990) also observed that methanolic fractions from aqueous leaf extract were not toxic even up to doses of 200mg/kg. Gandhi et al. (1988) noted that 5ml/kg of neem oil was very much tolerated.
by animals. After a 10 ml/kg dose, mild to moderate stuporous states were observed, followed by a tremulous gait, along with signs of severe respiratory distress, followed by death in some cases. Based on the observed 24 h mortality in the various treatment groups, the LD1 and LD50 estimated by the graphic method were found to be 4.9 and 20.0 ml/kg respectively. Gross examination of the dying rat showed that all organs except the lungs looked normal. The serum value of total bilirubin and SCOT were significantly higher in the test animals as compared to the control, suggesting an early damage of liver function. Lai et al. (1990), on the other hand, observed that small amounts of neem oil given to neonates and infants on a regular basis caused toxic encephalopathy. The usual features were drowsiness, tachypnoea, and recurrent generalized seizures. Leukocytosis and metabolic acidoses was also observed.

Studies by Komolafe et al. (1988) in the case of leaf extract indicated that mixed function oxidases (MFO) are involved in the hepatobiliary toxicity, since pre-treatment with MFO inhibitors were protective. Akah et al. (1992) studied the effect of aqueous extract of neem leaves on rabbit liver using enzyme indices of hepatic dysfunction. The results indicated that a high oral dose (2328 mg/kg) of the aqueous extract may have some hepatobiliary toxic effects. Chattopadhyay et al. (1993) carried out biochemical and toxicity studies with neem leaf extract. The 24h LD50 of the extract was 4.57 g/kg in mice. The leaf extract lowered serum cholesterol level significantly without altering serum protein, blood urea and uric acid levels.

While studying reproductive toxicity due to extract of seed in the adult cyclic female rat, Prakash et al. (1991) observed that the extract caused degenerative changes in the reproductive organs, leading to breakage and deterioration of the luminal epithelium. Total protein and glycogen contents decreased with the period of treatment. Badri Srimannarayana (1993) also concluded that a chronic administration of neem oil to adult rats for 8 days was not well tolerated and there were microscopic lesions both in the liver and the kidney. All the biochemical and histological parameters showed marked changes, indicative of the toxic effect of the oil.

The acute and sub-acute oral toxicity of azadirachtin-based pesticides has been determined by Mahboob et al. (1995) in rats. The study revealed that medium and high doses of azadirachtin caused alterations in the detoxification enzymes of various tissues, whereas low doses produced no such effects. In the case of high doses, the symptoms produced were reversible on cessation of treatment.

NEEM OIL AND REYE’S SYNDROME

Neem oil as a cause of Reye’s syndrome was first observed by Sinniah and Baskaran (1981). These authors investigated the oil further to find out if the poisoning is due to oil itself or due to aflatoxin. It has been observed that neem seeds are highly contaminated with fungal flora and rich in aflatoxin, when used for oil extraction. This is mainly because neem fruit matures in the rainy season, and it has to be dried in very humid conditions. Moreover, as the oil is used mainly for making soap and other non-edible purposes, no attention is paid to cleanliness and the oil is stored and transported in used iron drums, which may cause metallic contamination or even rancidity, or bring about some chemical changes in the fatty acid composition of the
oil, giving rise to toxicity. Sinniah et al., (1983) while pursuing the matter further, found that the aflatoxin B and C content of unrefined commercial oil varied between 40 and 100mg/kg. It is pertinent to note here that Gandhi et al. (1988), who obtained oil from a reliable source for their experiments, found that oil up to 5ml/kg was well tolerated by the animals. The toxic symptoms started after 10ml/kg, whereas the infants studied by Sinniah and Baskaran (1981) were given only a few drops of neem oil, say a maximum of 1ml. Considering the average weight of the child as 3kg, the dose comes out to be 0.33ml/kg, which is much below the toxic levels of the oil reported by Gandhi et al. (1988). The toxic symptoms observed by Lai et al. (1990) in infants are also different from those seen by Sinniah and Baskaran (1981). In another study, Sinniah et al. (1985) observed that intra-peritoneal injections of neem oil and seed extract may produce many of the abnormalities seen in Reye's syndrome, but there was no effect on hepatic enzymes and no evidence of cerebral oedema, which may be due to the contaminants in neem oil. Koga et al (1987) noted inhibition of mitochondrial functions with neem oil and seed extract in isolated rat liver, which could be reversed with coenzyme Q.

In a review article, Nat et al. (1991) concluded that extracts of leaf, bark and isolated limonoids show a very low toxicity, especially when taken orally, but the seed oil is toxic. The toxic principles in the oil may be fatty acids, limonoids or an unsaturated hydroxy aldehyde. The possible mode of action of these may be due to interference with hormonal regulation, inhibition of various enzymes, interaction with receptors and alteration of membrane permeability and integrity.

Prasad et al. (1993) reviewed the earlier literature on the toxicology of neem and concluded that there is no mutagenic toxicity, as studied in the Ames test using Salmonella typhimurium.

NEEM POLLEN ALLERGY

Allergy due to neem pollens was reported by Saha and Lalyanasundram (1962). Two free amino acids were isolated from the pollens by Chanda et al. (1975), while Karmakar and Chatterjee (1994) isolated two allergenically active compounds AlAl and AlAl IVb.

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14. NEEM IN AGRICULTURE

EARLIER REPORTS ON PESTICIDE ACTIVITIES

The activity of neem against locusts, though not well documented, has been well known to Indian farmers since very early times and some information about it is available in the earlier publications (Pruthi, 1937; Volkonsky, 1940; Pradhan et al., 1962; Mitra, 1963). It was mentioned that locusts avoided feeding on leaves sprayed with crude extracts of neem and China berry. It was Robert Larson of Vikwood Botanicals, USA, who during his many business trips to India, brought to the notice of American scientific workers the property of neem against insects. This was the era when the use of synthetic pesticides was widespread, and more and more health hazards about them were coming to light, but no alternative was in sight. There was a need for safer and effective biodegradable pest control compounds with greater stability.

THE PROBLEMS CREATED BY SYNTHETIC PESTICIDES

It was seen that the continuous and indiscriminate use of synthetic chemicals for the control of insects led to the following problems:

1. Environmental pollution, as the chemicals brought about biochemical changes in the various organisms.
2. Health hazards due to high residue levels.
3. Indiscriminate destruction of insects, without any consideration of their beneficial or harmful nature.
4. Poisoning of warm-blooded animals like birds, farm animals, fish and persons coming into direct contact with these.
6. Resurgence of certain major and minor pests, which were earlier being dominated by the insects, which were destroyed by the pesticide. With their disappearance there was less competition and new pests appeared.

PESTICIDES FROM PLANTS

Keeping the above point in view, a search for a phytotoxic pesticide started; it was seen that about 2500 plants had one or more activities against insects but only neem was found to be a highly effective, non-toxic, and environmentally friendly agent for controlling insects by acting as feeding inhibitor and growth regulator (Warthen, 1979), and it was projected as the insecticide of the future for protection against field pests (Jotwani and Srivastava, 1981). Thakur et al. (1981) published a bibliography on neem.
Kubo and Klocke (1982), while looking for limonoids as insect controlling agents, isolated and identified azadirachtin as an antifeedant. It was also observed that these limonoids prevented the completion of larval moulting by inhibiting the exuviae after the formation of new cuticle. These compounds did not kill the insects directly but lowered their growth rate and made them more vulnerable to other mortality factors. Jaipal et al. (1983) also noted juvenile hormone-like activities in the bark of neem, and observed that the metamorphosis of the insect was inhibited to varying degrees by these. The use of purified extract of neem was suggested for pest control. Swaminathan (1983) brought forward the potentiality of neem in pest control. Freeman and Andow (1983) described the role of neem as a tree for protection of other plants as an insect feeding deterrent. Jacobson (1986) gave details of its insecticidal activity.

With the above publications, the importance of neem became well known in the scientific world, and it became a topic of discussion at various international conferences. Schmutterer and Ascher (1986) edited the proceedings of a conference which had research papers on the pesticidal activity of neem. Saxena (1987) brought forward the use of neem as an antifeedant in pest management in the tropics and recommended quality control and standardization of its biological properties for introduction on a commercial scale. Kareem et al. (1987) meanwhile observed that with the use of neem oil mixed with custard apple (Anona reticulata) in rice fields, virus incidence was significantly less, and the yield of rice was higher. Singh and Singh (1988) also noted antiviral activity of leaf and bark extract of neem.

AZADIRACHTIN AS AN INSECTICIDE

Pest control aspects of neem were found to be useful in both developing and industrialized countries by Schmutterer (1988), who observed that azadirachtin and azadirachtin-containing neem extract acted as an antifeedant growth regulator and sterilant. The mode of action of azadirachtin may be due to interference with the neuroendocrine system controlling ecdysone and juvenile hormone synthesis and to inhibition of ecdysone release from the hormone-producing gland. In addition, azadirachtin causes inhibition of chitin synthesis. Azadirachtin was found to be an unstable compound, whose residual effect lasted for 4–8 days, but degradation may be hastened by ultra-violet light, rainfall and other environmental factors.

a detailed account of azadirachtin. Rovesti and Deseo (1990) further discussed the potentiality of neem in pest control. Arnason and Philogene (1991), in memoirs of the entomological society of Canada, gave an account of plant-derived substances in insect control. Isman et al (1991) studied variations in the azadirachtin content of twelve commercial samples of neem by their growth inhibition, antifeedant and moulting disruptive activity and concluded that the bioactivity of neem oil was dependent on its azadirachtin content. The possibility of a neem-based insecticide for Canada was discussed. Maramorosch (1991) reviewed the current status of research, while Remboldt and Raychaudhuri (1991) gave further details of the growth-inhibiting properties of azadirachtin. Champagne et al (1992) described the biological activity of limonoids from neem and the other members of the Rutales family. Mordue and Blackwell (1993) presented an update on azadirachtin. The potential and limitations of neem pesticide were reviewed by Soon and Bottrell (1994). The authors outlined the use of neem to control pests and its effect on nontarget organisms like the honey bee, earthworms, aquatic life, man and other warmblooded animals. In a workshop (Kleeberg, 1994), the production of neem ingredients, pheromones, and their effect on phytophagus insect pests, fresh water snails and pathogenic fungi were discussed. Remboldt (1994) gave a further account of azadirachtin and its mode of action.


The above exhaustive studies confirmed the application of neem in the fight against pests. It was found to act on eggs, larvae/nymph and adults.

**Mode of Action of Azadirachtin**

The various studies showed that the mode of action (Fig. 16) may be as follows:

1. **Antifeedant through mouth.**
   (a) Primary: it inhibits the activity of sensory receptors of mouth parts, distorts normal probing feeding and intake of food.
   (b) Ingestion of active ingredients through food leads to starvation and death.
2. **Dermal action:** it enters through the cuticle of the insects and inhibits chitin synthesis, thus causing desiccation and death.
3. Repellent effect: due to change in the locomotor and settling behavior of insects, in some cases mating as well as sexual communication is disrupted.

4. Growth-disruptive effect: by inhibition of the normal growth of the insect by interfering in the moultng cycle. It suppresses the activity of ecdysone so the larva does not moult, but remains at the young stage and dies.

5. Effect on survival and reproduction by oviposition deterrent action: when the female comes to an egg-laying period of her life cycle, the egg laying is prevented.

6. Effect on endocrine system: neem preparations are accumulated in the neurosecretory system and, by penetrating the blood brain barrier, are concentrated in the corpus cardiacum, resulting in reduced turnover of neurosecretory proteins.

Neem does not have an immediate knock-down effect like most of the synthetic chemicals and thus it is effective against those insects that have now become resistant to chemicals. It was also found effective against those pests that live concealed and well protected in the plant parts. Neem is not universal in its effect, which varies from insect to insect, lepidoptera being more sensitive to it, as compared to others.

**RECENT STUDIES**

Neem has also been found to have a nematode-suppressant activity; Siddiqui and Alam (1990), Sen and Dasgupta (1990) and Alam (1993) described this.

In 1995, quite a number of books were published on neem for its use on agricultural and domestic pests. In the book, *Neem—a user’s manual*, Vijayalakshmi et al. (1995) gave simple methods of neem preparation from fresh materials, for control of insects by marginal farmers in India. The authors gave a brief account of the mode of action of neem products on some of the selected pests. Line drawings of the insects were given, along with their zoological names. The method of application of these on various crops has also been mentioned. Mariappan (1995) edited a book *Neem for the Management of Crop Diseases*. The book deals with the application of neem in...
plant diseases caused by agents other than insects, such as fungi, bacteria, viruses and nematodes in various crops of India. The results of research in applied entomology by a working group in the last decade at Giessen University are given by Schmutterer (1995). The research was carried in Latin America and in African and South Pacific countries, with very encouraging results. In some cases, by application of neem, use of synthetic pesticides could be reduced considerably, with appreciable increase in yield.

Schmutterer (1995a) edited a very exhaustive book on neem. The book carries articles on nearly all aspects from specialists who worked on neem in the last decade or so. It has detailed information on integrated pest management. The various articles deal with the use of neem pest-wise and crop-wise and also in farms and orchards. Singh et al. (1995) edited the book Neem and Environment. The book contains articles presented at the world conference on neem held in 1993.

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15. NEEM SEED CAKE AS A MANURE AND NITRIFICATION INHIBITOR

AS A MANURE

The use of seed cake, obtained after the extraction of oil, particularly that of non-edible seed, as a manure is well known in agriculture. The use of neem cake has also been recommended for this purpose. Keeping in view the importance of this in agriculture, the Indian Standards Institute (Anonymous, 1977) has given specification No. 8558 for neem cake, for manuring, which is as follows:

- Maximum moisture (percent by mass) 10%
- Maximum water soluble organic nitrogen on moisture free basis (percent by mass) 2.5%
- Maximum total ash (percent by mass) 13%
- Maximum acid insoluble ash (percent by mass) moisture free basis 5.0%

Neem seed cake not only provides nutrition to the plant, but controls soil-borne pests, particularly nematodes; it also acts as a nitrification inhibitor, helps respiratory activity, increases the population of earthworms and produces organic acids, which help in removing the alkalinity of the soil (Korah and Shingte, 1968).

AS A NITRIFICATION INHIBITOR

When a nitrogenous fertilizer is applied to the soil, about half the nitrogen escapes into the air by the activities of nitrifying bacteria, such as *Nitrosomonas* and *Nitrobacter*. If a bacteriostatic like sulfur is used, it suppresses the growth of these bacteria and acts as a nitrification inhibitor (Rajendra Prasad *et al.*, 1971). The same results were obtained by Sahrawat and Parmar (1975) from an alcoholic extract of neem. Neem-coated urea increased the apparent recovery of nitrogen to 48.6 percent, which was 29.3 percent in the case of untreated urea (Ketkar, 1976). In a field experiment, 160kg urea with 40kg neem cake per hectare produced 6 quintals more rice, as compared to the field where only 200kg urea was applied. Ramababu *et al.* (1983) got similar results.

Watanabe *et al.* (1981) studied the effect of neem cake on the nitrogen fixation activity of blue green algae in a flooded rice field. Slangen and Kerkhoff (1984) presented a review of literature on nitrification inhibitors in agriculture and horticulture. Uma Singh and Gurumurti (1984) suggested neem cake as a potential fertilizer. In a study it was seen that after ten months, the root length, number of branches and dry weight of the plants increased as compared to the control (Uma Singh *et al.* 1986).

Naidu and John (1984) found that fermented neem cake, when used with other oil seed cakes, inhibited the fungal rice pathogens *Rhizocotonia solanii* and *Sclerotium oryza* (*Magnaporthe salvinii*). Korah and Shingte (1988) presented data to show that
nitrogen mineralization was greatest with neem as compared to other non-edible oil seed cakes. Nair and Sharma (1988) noted the difference in the performance of ordinary and neem-coated urea. The persistence of neem kernel powder and neem oil on phorate granules in the soil was studied by Dethe and Babtiwale (1989), and it was found that degradation was faster in unsterilized soil as compared to sterilized soil. AH and El-Sanousi (1989) tested the sensitivity of different aerobic bacterial species to azadirachtin, while Turkur et al. (1989) studied the effect of different nitrificaton inhibitors including neem on protein, non-protein, ammonical and nitrate content in the soil. Rajinder Prasad et al. (1990) also carried out this type of study.

Thampatti et al. (1992) applied muriate of potassium with urea neem cake blend and found that the exchangeable potassium was highest in all soil layers, as compared to the control.

Many studies, including the one by Ameta and Singh (1990), confirmed an increase in yield when urea coated with neem powder was applied to the field, but Das and Mukherjee (1990) noticed the adverse effects of neem cake on beneficial organisms present in the soil. Mukherjee et al. (1991) confirmed this and observed a lesser number of bacteria, fungi and actinomycetes. It was also observed that neem seed cake caused an accumulation of ammoniated nitrogen in the soil during inhibition of nitrification of urea and an increase in pH, which gives rise to several problems, including damage to young plants due to nitrogen toxicity.

The denitrification property of neem cake has also not been properly understood so far. It may be due to sulfur compounds which have a well-known antibacterial effect, or the lipids associated with the cake may be responsible for inhibition of bacterial growth. The activity may be due to tannins which inhibit the urease enzyme in the soil. Devakumar and Riar (1993) considered it due to bioregulators like meliacins, epinimbin, salin and azadirachtin.

For large-scale production of neem-coated urea, Vyas et al. (1991) developed a neem extract. This was an improvement on the earlier method, in which coal tar dissolved in kerosene oil was sprayed on urea as an adhesive, followed by a coating of fine powder of neem cake on each granule of urea. Suri (1995) suggested a commercially viable process in which an emulsion of neem oil was coated on urea during the production stage.

REFERENCES


16. POULTRY AND CATTLE FEED

In developing countries particularly, there is competition between man and other domestic animals for conventional food, leading to malnutrition for all. If regular cereals and legumes consumed by man are replaced partly by other food items in the diet of chicken, cattle, pig, etc. (Punj, 1988), it will release pressure on the food directly as well as indirectly. If there is less consumption of cereals by animals, more food will be available to the human population, and man will be saving cereals/legumes by eating a non-vegetarian diet.

Neem leaves are consumed by camels, goats and in drought by cattle also (Patel and Patel, 1957; Hentgen, 1985). Shukla and Desai (1988) suggested neem as a source of cattle feed. The seed is rich both in fatty oil and in protein, but it could not be used unprocessed for edible purposes on account of the deleterious effects it has on animals and birds because of the bitter principles contained in it. For the last fifty years, research has been conducted, and various feed trials on animals have been undertaken to utilize these seed as feed. An account of these activities is given here.

ANIMAL FEED

The Leaves

These are fed particularly to camels, which appear to relish them. Goats may also eat them. In Andhra Pradesh (India), leaves are fed to milk-yielding animals to increase the yield of milk after parturition.

Singh (1982) included neem among the fodder trees of India. (The neem tree was introduced in arid zones or on degraded soil on a large scale and the use of leaves as a forage was advocated). As a multi-purpose tree, the leaves were recommended as suitable for browsing and for incorporation into a concentrated ration after drying. It was reported that dry neem leaves have better nutritional quality than sorghum, which is the main dry season fodder (Zech and Weinstable, 1983; Webb, 1988). The leaves are said to be palatable to cattle and buffaloes; they are a good source of protein (15 percent) and carotene 185 µg/g, and contain most minerals except zinc. They have lower fiber content. Fallen leaves can also be fed but they are less palatable, containing about 8 percent protein and 12.71 percent ash. Conventional digestible trials by Patel and Shukla (1962) found that neem leaves contain 6.2 percent digestible protein and 52.5 percent total digestible nutrients. It was reported by Jayal (1963) that the digestible co-efficient of neem leaves was 51.96 percent, for protein 58.45 percent and for fiber 22.33 percent.

The approximate composition and nutritive value of neem leaves as given by Ranjhan (1980) are organic matter 92.25 percent, crude protein 16.12 percent, ether extract 3.40 percent, crude fiber 20.69 percent, nitrogen-free extract 52.06 percent, total ash 7.73 percent, digestible energy 2394Kcal/kg and metabolizable energy 1926Kcal/kg. Murugan and Kathaperumal (1987) and Murgan et al. (1987)
determined the macro and micro mineral content of leaves to see if these can be used as animal fodder.

When animals are fed with cereal and fodder, the danger of a negative nitrogen and mineral balance exists; the feeding of neem leaves help to alleviate these.

It is the fodder of choice during dry periods and drought. In the west Indian state of Gujrat, during a famine 15–20kg of neem leaves were fed to cattle and buffaloes daily for their survival (Ketkar, 1976).

The above use does not appear to be universally acceptable, as Singh and Pathak (1981) and Murugan and Kathaperumal (1987) observed that sheep fed on leaves or fruits, dry or meal or silage, lost body weight. Ali (1987), when he administered an aqueous suspension of dry leaves at a dose of 50 or 200 g/kg to goats over a period of up to 8 weeks, saw a progressive decrease in body weight, together with weakness and lack of appetite. Higher doses even produced tremors and ataxia, with drastic histopathological changes in various organs. In guinea pigs the toxic symptoms were less severe. Ibrahim et al. (1992) fed chicks with a diet containing 2–5 percent leaves. A decrease in body weight gain and efficiency of feed utilization was observed, with yellow discoloration of legs and combs. Many histopathological changes took place in the body.

Neem Oil

Because of bitters and odoriferous principles, the oil was not considered suitable for animal consumption. It was found to be toxic though its fatty oil composition is very near to other edible oils. Rukmani (1987) and Rukmani et al. (1991), to evaluate the nutritional value of the oil, extracted the seed with aqueous ethanol followed by hexane, and got oil free from bitters, odouriferous principles, coloring matter and free fatty acids. A toxicity study on neem was also carried out by Rukmini (1987), who found it to be a safe source of edible oil to be used as such or blended or hydrogenated. Reddy et al. (1988) concluded that oil treated with 10 percent potassium hydroxide (KOH) was detoxified, as was evident when fed to chicks. The oil, prepared by treating it with 5 percent KOH, without heat, was not completely utilized by chicks, but when given in a phased manner, it was comparable with the control diet. Vasishtha et al. (1992) studied solvent-extracted neem oil. Chinnasammy et al. (1993) carried out further studies on reproductive toxicology for three generations. There was no change in the various organs of the body or mutagenicity as compared to the control. The oil was found to be fit even for human consumption. The fatty oil composition of neem oil reported was myristic acid 0.2 to 2.6 percent, palmitic acid 13.6 to 16.2 percent, stearic acid 14.4 to 24.1 percent, oleic acid 49.1 to 61.9 percent, linoleic acid 2.3 to 15.8 percent, and archidonic acid.

Full Fat Seed Meal

Chicks, when fed a diet containing 2, 5, 10 percent ripe fruit of basic diet, from 7 to 35 days, showed a decrease in body weight gain, efficiency of feed utilization and hepatonephropathy (Ibrahim et al., 1992a). Full fat seed meal at 0, 25, 75 and 100g/kg was studied in broilers and rabbits at 0, 200, 300 g/kg for 8 weeks. In chicks,
there was a significant negative relation between neem meal, weight gain and feed conversion efficiency. The rabbits, when given up to 100g/kg of neem preparations, were superior to the control, but not at higher concentrations (Salawu et al., 1994).

Seed Cake

The easy availability of seed cake in large quantity at affordable price and high nutritive value, came to the notice of scientific workers a long time ago.

According to Ranjhan (1980), neem cake contained 82.40 percent organic matter, of which crude protein was 17.03 percent, ether extract 1.02 percent, crude fiber 42.12 percent, nitrogen-free extract 22.13 percent and total ash 17.60 percent. The percentage of amino acid profile of neem cake was aspartic acid 1.31, threonine 0.50, serine 0.38, glutamic acid 2.40, proline 0.84, glycine 1.08, cysteine 1.73, valine 0.76, methionine 0.70, isoleucine 0.60, leucine 0.95, tyrosine 0.26, phenylalanine 0.80, histidine 0.21, lysine 0.28 and arginine 0.57. Maitra and Duttagupta (1982) gave an analytical report of neem seed cake as follows: crude protein 21.9 percent, ether extract 8.6 percent, nitrogen-free extractives 44.2 percent and crude fibers 11.6 percent. It is very rich in organic minerals. In commercial cattle feed, inorganic minerals are usually added; cattle have to convert them into organic compounds and hence the availability of minerals is only up to 40 percent, the rest being lost in the process of digestion. Neem cake helps in mineral nutrition, as it is particularly rich in Ca, P, Fe, Cu and Zn. Rao (1987) carried out nutritional trials with debittered and defatted oil cake.

The chemical analysis by Reddy et al. (1988a) to establish its suitability for animal feed indicated that it compared well with peanut cake. It contained less crude protein, more fiber, fat-and nitrogen-free extractives, calcium and phosphorous. Carbohydrate content was 19.28 to 22.27 percent. True metabolizable energy value for decorticated expellerprocessed neem cake was 2.959 to 2.973Kcal/g and for undecorticated expeller-produced neem cake it was 2.279 to 2.823 Kcal/g with gross protein value 55 to 59 percent.

The only negative point with neem cake was the taste and odor, which are not palatable to animals. For removing the bitter principles, various methods were tried earlier; some of these are:

1. Treating with various organic solvents like petroleum, alcohol, etc. (Lehri et al., 1987). One of the methods suggested was to dry the seed properly, wash them with hexane to recover all the oil, and then to extract with ethanol.
2. Treating with various concentrations of acid and alkali.
3. Repeatedly washing with water until the bitter principles are removed.
4. Heating the neem cake to 150°C.

FEEDING TRIALS WITH ANIMALS

Cattle and Buffalo

Earlier work on the nutritive value of neem cake and its digestability by buffaloes was done by Arora et al. (1975) and Bedi et al. (1975). Pyne et al. (1979) studied the
composition of the milk of lactating buffaloes which were given neem cake in the feed. Maitra and Duttagupta (1982) heated untreated neem cake to 150°C. Heat treatment did not affect nutrient digestibility significantly. Maitra et al. (1982) also observed that the net effect of energy utilization was higher for a concentration of 10 to 15 percent, as compared to a concentration of 20 percent. Nath et al. (1989), when they treated neem seed kernels with water and dried them, found that these were palatable to cattle. The animals could digest them well and they did not affect the physiology or even some blood constituents. Agarwal et al. (1987) also observed that water-washed neem kernel feed had higher efficiency of utilization of digestible protein, as indicated by the greater nitrogen balance (Singhal and Mudgil, 1983).

In field trials, various efforts were made to make neem cake palatable to animals, by adding molasses and starch. There was some acceptability when double the amount of maize was added, but in these cases the protein metabolism was adversely affected. In buffalo, the addition of 10, 15 and 20 percent neem cake did not have any adverse effect on the milk composition or the health of the animals. Both red and white blood cells and hemoglobin were higher in the blood but serum protein was lower. The serum protein was decreased by increasing the neem seed cake in ration. Some factors affected hematopoiesis (blood formation). In the liver, it did not affect SCOT and SGPT. Calcium and phosphorus in the blood was also not affected (Gangopadhyay et al. 1979, 1981).

The feeding of water-washed neem kernel cake to cows at 400g/kg did not affect performance, blood constitution or reproductive ability (Nath et al., 1983, 1989). A higher nitrogen balance was caused by reduced excretion of nitrogen in urine and a decrease in blood urea nitrogen. Kumar et al. (1990, 1992), after the study, concluded that neem seed cake can replace up to 30 percent dairy concentrate, consisting of proteins, fats, minerals, etc. It did not affect nutrient digestibility or milk yield. In another study, concentrate was replaced by various proportions of neem cake, Pennisetum purpureum and mixed grass hay. The above feed did not affect milk yield or its composition. But when peanut cake was fully replaced by neem cake by Garg and Nath (1990), it depressed nutrient digestibility and also the nitrogen, calcium and phosphorus balance. When the neem seed kernel was fed with a balanced diet after soaking in a solution of sodium hydroxide and washing with water, there were no clinical signs of ill health and vital organs showed no pathological changes (Katiyar et al., 1993). It was concluded that alkali treatment totally detoxified the neem cake. Water-washed neem kernel powder, when included in feed by Mahendra et al. (1995) for four months, did not have any adverse effect on the daily milk yield, but lowered the feed cost considerably.

Pig

A diet which contained 5 percent neem cake decreased the value of feed efficiency (Thomas and Prasad, 1983), but when neem cake was replaced by water-washed neem kernel cake (40 percent crude protein), pigs grew faster and utilized the feed more efficiently with higher nitrogen retention. The feed cost was reduced by 11 percent (Sastry and Agarwal, 1992).
POULTRY AND CATTLE FEED

Poultry

Toxicity in the water-extracted neem cake was found by Singh et al. (1985). Expeller processed neem cake decreased the weight and feeding efficiency of birds (Reddy and Rao, 1988), as was the case with solvent-extracted neem cake, but when neem cake treated with acid or alkali was fed to birds, it improved growth which was comparable to the control (Reddy and Rao, 1988a, 1988b; Reddy et al., 1988). Chakravarty and Prasad (1991) studied the effect of neem leaf extract and other neem extracts on the performance of broiler chicks.

Rabbit

Fajinmi et al. (1990) studied neem seed in the diet of rabbits with encouraging results.

Rat

In feeding trials with rats also, the growth-suppressant compound was found (Vijjan and Parihar, 1983) to be water soluble and could be removed by washing neem cake (Vijjan, 1983). Prakash et al. (1991) observed reproductive toxicity in the case of adult females. Garg et al. (1991) tested this compound in males and observed a decline in growth rate, food consumption and testes weight.

Sheep and Goat

With feeding trials, weight loss and toxicity symptoms were evident in lambs. After 15 days there was severe gingivitis and sloughing of mucus membrane with foaming discharge from the mouth, after 25 days stomatitis, gastro-enteritis, and diarrhea were followed by death (Vijjan et al., 1982), but when a mixture containing 17 and 34 parts of neem was fed to sheep the symptoms were less severe. Gupta and Bhaid (1981) and Ramu et al. (1994) fed 10, 20, 30 percent water-washed neem seed cake to sheep and goats. They digested it well. These authors carried out further studies (Ramu et al., 1994a) and concluded that a complete ration can be formulated by incorporating 30 percent water-washed neem kernel seed. Evaluation of cooked meat did not reveal any bitter taste up to 20 percent when water-washed neem cake was added in the diet of sheep and goats in place of deoiled rice bran (Reddy et al., 1994). In another study, Verma (1995) and Verma et al (1996) also concluded that water-washed neem kernel can be incorporated in the diet of growing goats up to 25 percent without any deleterious effects on nutrient utilization and metabolism. This type of diet did not have any adverse effects on the weight of the goats, and sensory evaluation of cooked meat did not reveal any bitter taste.

REFERENCES


17. NEEM AND POLLUTION

Rapid industrialization, urbanization, and congestion of population in a few pockets, in most part of the world, are giving rise to pollution caused by emission of gases such as carbon monoxide, carbon dioxide, sulfur dioxide and nitrogen peroxide which may play havoc with the human population. In Indian culture, neem has been referred as an “air purifier” so it may be an avenue tree of choice in thickly populated areas, by its capacity to survive in adverse conditions, absorb some of the environmental pollutants, and act as an “air freshener” by releasing oxygen and mild odorous principles.

INDUSTRIAL POLLUTION

Tanneries

Tannery is one of the industries responsible for pollution of river water. In third world countries, in some areas, the cattle population exceeds that of humans, so an appreciable amount of animal hide is available which is treated with tanning materials to turn it into leather. The whole process requires repeatedly washing with water, so the water requirement is very high; after washing, this water becomes heavily contaminated and is drained back to the rivers.

Chaturvedi (1986) tested neem as one of the trees for tolerance to tannery waste water. The survival rate of the tree was 22–94 percent. Ramanuja and Misra (1986) did culture experiments over 12 weeks, with 8–9 month old seedlings. Plants were irrigated weekly with 2 liters of effluent from a tannery settling tank. Neem was found quite tolerant to this waste water.

The other major polluting industries are thermal plants and chemical factories, such as those for fertilizers and pesticides, which release carbon dioxide, sulfur dioxide and nitrogen peroxide, in addition to suspended particles like dust or fly ash.

Chemical Factories

Devi and Patel (1982) and Patel and Devi (1985) studied morphological variation in the vegetation around a fertilizer complex. The variation noted was reduced foliage or defoliation, cracking and peeling of bark, reduced leaf and leaflet area and petiole length, mutilation of leaves in various ways and the total absence of flowering.

In the case of leaves in normal plants, the cell wall was undulating but in the polluted area it was straight, with reduced stomatal frequency and some variation in stomatal width and pore area. Starch, insoluble polysaccharides and lipids also varied.

On 3 December 1984, large quantities of methyl iso-cyanate (MIC) escaped from a pesticide plant in central India. Ram Prasad and Pandey (1985) studied the effect of this poisonous gas on the neem tree. The tree was sensitive to MIC. There was defoliation and blackening of the foliage, but the tree revived two months after the injury with the emergence of new leaves. Farooq et al. (1988) studied the sulfur
dioxide resistance of trees and also visible symptoms of sulfur dioxide absorption. Seedlings of 12 species, including neem, were exposed to sulfur dioxide to various concentrations and it was found that neem could tolerate this gas to a major extent. Rao and Dubey (1990), in their study to find differential responses to sulfur dioxide, analyzed the neem tree along with others for stomatal conductance, sulphate, protein, superoxide dismutase and peroxides for one year in an ambient environment with varying concentration of sulfur dioxide. The results indicated that trees under sulfur dioxide stress developed phytotoxicity by undergoing certain biochemical changes.

To find out the effect of sulfur dioxide exposure to tree saplings, Krishnnayya and Bedi (1989) noted that this gas damaged chloroplasts and cytoplasm in palisade cells, followed by rupturing of the outer envelope of the chloroplasts and extrusion of plastoglobuli and starch into the cytoplasm. Thinma Raju et al. (1993) observed that a neem tree growing in a highly polluted area was not affected by various gases, whereas some other trees exhibited symptoms of defoliation, die-back, poor flowering and fruiting.

**Thermal Stations**

In one experiment, fly ash-induced injury to leaves and proline metabolism in plants growing at two different distances away from a thermal powerhouse was studied. Trees closer to the source of pollution had higher dust deposition, leaf injury and proline accumulation at a lower pH of cell sap as compared to neem trees away from the source. Proline accumulation was present at both sites, throughout the period of study of four months. It indicated the greater ability of neem to adapt to stress from exposure to air pollution (Chauhan and Varshney, 1989; Maini and Harapanahalli 1991; Anonymous, 1991).

Beg et al. (1990) studied the performance of neem trees at five selected sites. In the vicinity of a power station, gases like nitrogen peroxide and sulfur dioxide were below the permissible levels, but the total suspended particles were higher. Air pollution had a major effect on chlorophyll. It was seen in this study that there was some destruction of chlorophyll, showing thereby that neem is moderately sensitive when exposed to sulfur dioxide in various concentrations but could tolerate this gas to a major extent.

**Exhaust from Automobiles**

Pollution due to the exhaust from automobiles in some congested areas is fairly high. The effect of this exhaust on the survival of neem, if planted as an avenue tree, is very important. Keeping this point in view, Bhatti and Iqbal (1988) studied the leaf length area, dry weight, etc. of these trees. It was noted that neem tolerated this type of pollution very well and can be planted as a roadside tree in thickly populated areas.

When dust loading of leaves in an automobile polluted area was studied, it was seen by Satyanarayana et al. (1990) that the neem tree accumulated a thick, sticky crust. In the leaves, as compared to the control, the epidermal cell size was smaller.
and stomatal frequency was higher. Sharma and Roy (1995) also observed the same features on leaves when subjected to automobile exhaust.

Steel Industry

The phytotoxic effects of aerial discharge from the steel industry were evaluated for neem by Kumawat and Dubey (1988). In the neem trees growing in the area, the chlorophyll pigments, carotenoids and leaf pH level decreased, while the leaf injury index, leaf area/dry weight ratio, conductivity of leaf disk water, sulfate content and total chlorophyll: sulfate ratio increased. The authors observed that the pollution injury was maximum in winter, followed by summer and the rainy season.

Detection of Heavy Metals

The neem bark was tested to monitor heavy metals in polluted sites, and compared with those absorbed by moss. Lead, zinc and iron content in both materials were higher at most sites (Kakulu, 1993).

Water Purification

Johri et al. (1993) developed a process of flocculation of water pollutants by using extract of seeds of neem along with Moringa oleifera and Madhuca latifolia. This extract formed a floe with the contaminants and the suspended particles settled down, giving rise to clean water.

ENVIRONMENTALLY FRIENDLY

Bees

Before neem products could be used against insects and pests, quite exhaustive studies were carried out on the safety aspects, particularly with regard to warm-blooded animals. Concern was also expressed over its effect on honey bees. Schmutterer and Holst (1987) studied this aspect. The neem-treated flowers were not repulsive to bees and the neem preparations did not cause any serious damage to their systems. On the other hand, neem controlled the tracheal mites of the bee (Liu, 1995).

Neem-based products were not only found safe for earthworms, but for young salmon also (Wan et al, 1996).

TOXICITY

Neem trees in a grove are eco-friendly, but pollens have been known to cause allergy in some cases. Karmakar and Chatterjee (1994) isolated and characterized IgE-reactive proteins for pollens. They have shown that AlaI and AlaIVb are the major allergens. Amino acid analysis of these, the effect of pH on them and cross-activity has also been carried out by these authors.
REFERENCES


18. NEEM AND HOUSEHOLD PESTS

Various studies have now indicated that neem may be useful against household and kitchen garden pests, as follows.

ANT

The sterilizing effect of neem extract on the queen and workers of *Formica polyctena* were tested by feeding and fumigation. Fumigation increased egg production, but in feeding experiments the laying capacity of eggs was reduced with higher concentration of extract (Schmidt and Pesel, 1987).

BED BUG

Toxicity against *Cimex lectularius* has been found (Naqvi *et al.*, 1993).

BIRDS

Enormous loss to grains and young seedlings is caused by birds. Syamsunder Rao *et al.* (1993) tried various commercial neem formulations to find a non-lethal, environmentally safe product. The common birds showed feeding aversion and avoided consumption of neem-treated grains.

COCKROACH

A commercial preparation from neem seed extract was tried by Adler and Uebel (1985) against six species of cockroach, for toxicant, growth inhibitor and repellent action. Last instar nymphs showed increased mortality and retarded development but the other actions differed in different species. Topical application or an injection was toxic but the surface treated with neem extract was not effective in controlling them.

GOAT

The stray or wild herbivorous animal causes enormous loss to kitchen gardens and avenue trees, and protection of these can only be provided by fencing which is not only expensive but very often not effective. If some deterrent is applied on these, which is not toxic to vegetation and safe for the environment, most of these plants can be saved from these grazing animals. Gope *et al.* (1988) sprayed neem preparations
on tea bushes using hand sprayers; goats avoided the sprayed plants and after three months there was considerably less damage to the bushes.

This treatment may also be useful against deer, which in the USA not only cause extensive damage to plants but also spread Lyme disease.

HOUSE FLY

Ethanol extract of neem was used by Azmi et al. (1995) against Musca domestica. It increased mortality and was comparable to deltamethrin. Azadirachtin inhibited moulting in the larvae of the face fly M. autumnalis (Gaaboub and Hayes, 1984). A compound from neem extract caused morphogenic effects on various stages of fly, including weight reduction and abnormal development (Naqvi et al., 1995).

MOSQUITO

For control of malaria, Japanese encephalitis and dengue fevers, eradication of the mosquito is very important. The mosquito has spread to new areas because of the construction of dams, canals, and other irrigation facilities. The improper disposal of drainage water has provided further breeding grounds. The recently synthesized insecticides were indiscriminately used for the control of the mosquito, but some strains developed resistance against these chemicals. In due course of time, the polluting effects of these chemicals became well known, and a search started for some eco-friendly agent such as neem.

Neem is effective against the mosquito in two ways, as a larvicide, and as a repellent.

Larvicide

During and after the second world war, kerosene oil and tar were commonly used for larvicidal effect. It was a common practice to sprinkle these non-biodegradable products on standing water or in other mosquito breeding grounds in mosquito-infested areas. Recently, it has been observed that eco-friendly neem extract and neem oil can replace the above petroleum products (Attri and Ravi Prasad, 1980). Petroleum ether extract of neem was tested by Deshmukh and Renapurkar (1987) against third instar larvae of Culex quinquefasciatus. It totally suppressed larval development. The essential oil obtained by steam distillation had good activity against larvae of Anopheles stephensi (Kumar and Dutta, 1987).

Four azadirachtin-rich fractions also had this effect (Rao et al., 1988). Petroleum ether extract of dried leaves (total alkanes) was tried for larvicidal action. A one percent solution of extract gave 100 percent mortality. A mixture of purified extract was more effective than the crude leaf extract (Chavan and Nikam, 1988).

The larvicidal effect of neem seed bitters was also tested by Rao et al. (1989), who observed that 2000 ppm of the bitter produced 100 percent mortality in 72 hours. Reuben et al. (1990) discussed at length the use of neem extract for biological control of the mosquito. Tare and Sharma (1991) and Rao et al. (1992) experimented with
neem seed bitter principles at different concentrations against newly moulted fourth instar larvae of *C. quinquefastiatus* in flooded rice fields. The bitter principles were very effective, even at a concentration of 2000ppm. There was 100 percent mortality in 72 hours. The neem extract was applied to the rice field for the dual purposes of controlling the mosquito vector of Japanese encephalitis virus and enhancing the grain yield. The standardized extract of neem was found effective for both purposes.

In another study on larvicidal potential, *Aedes aegypti* was found to be more susceptible than *C. quinquefastiatus* (Monzon et al., 1994) to neem. When larvae of *A. aegypti* were reared in water containing different concentrations of a commercial preparation having 40 percent azadirachtin, it was seen by Boschitz and Grunewald (1994) that sensitivity to the product decreased with increasing age of the larvae. Female mosquitoes laid fewer eggs which were directly proportional to increasing concentration of the neem preparation to which they had been exposed during larval development.

The crude extracts of neem were effective against mosquito larvae but these extracts had a problem of stability and storability, when used on a large scale in the rice fields. Rao et al. (1995) tried a good quality neem product for control of larval cluicine mosquitoes. It also produced a slight but significant reduction in the population of anopheline pupae.

Mittal et al. (1995) tested six neem products against fourth instar larvae of *A. stephensi*, *C. quinquefastiatus* and *A. aegypti*. Larvae of *A. stephensi* were the most susceptible, those of *A. aegypti* the least. Jin Ping et al. (1995) studied toxicity and the growth-regulating activity of neem seed kernel extract in the larvae of *C. quinquefastiatus*. This extract induced prolongation of first instar larvae but caused death and morphogenetic aberrations of fourth instar larvae.

Neem oil and deoiled cake were also found to be promising larvicides by Amorose (1995). Third instar larvae were more susceptible than fourth instar larvae. Nagpal et al. (1995) soaked wooden balls in 5, 10, 20 percent neem oil diluted in acetone for the control of *Anopheles stephensi* and *Aedes egypti*. These balls were dipped in storage and overhead tanks. The balls soaked in 5 percent neem oil gave the best results. Similarly, pieces of cardboard dipped in oil may be used as a mosquito-repellent mat.

Repellent

Lemon grass oil in various forms has been used as a mosquito repellent. It has been used as an active ingredient of ointments or oils for external application on the exposed part of the body or in fumigants, coils, mats and candles, for fumigation. Lemon grass is a volatile oil, so preparations containing it are to be applied repeatedly to the body or the surroundings should be continuously fumigated. Recently, in mats, lemon grass oil has been replaced by synthetic pyrethroids and other compounds, because of their persistence nature, but some studies indicate that these are hazardous, particularly for infants and children in an enclosed atmosphere.

In India, for repelling insects or for so-called purification of air, premises are often fumigated with a mixture of neem leaves, oleo-gum resin, particularly that of *Commiphora wighti*, and sulfur. Pandian et al. (1989) and Pandian & Manoharan (1995) observed that with smoke from dry leaves of neem, the landing and biting rates of mosquitoes were reduced considerably.
Sharma *et al.* (1993, 1995) mixed 2 percent of neem oil in coconut oil and applied this mixture of oils to the exposed part of human volunteers. This provided complete protection against mosquito bites for 12 hours. In a field trial in a village in west India, a mixture of 0.5, 1 or 2 percent neem oil in coconut gave 79.65, 96.07, and 98.03 percent protection respectively against *Anopheles culicifacies* in an all-night biting test. Two percent neem oil provided 75 percent protection against other types of mosquito (Kant and Bhatt, 1994). Similar results were obtained by Mishra *et al.* (1995) with 1–4 percent neem oil in coconut oil, when applied to human volunteers in a tribal village. Dua *et al.* (1995) applied a neem cream to see if it can provide protection against mosquitoes. One application of the cream was effective in 68 percent of the population for four hours.

**SAND FLY**

Two percent neem oil mixed in coconut or mustard oil provided 100 percent protection against *Phlebotomus argentipes* throughout the night under field conditions. It was effective against *P. paptsi* for about seven hours only (Sharma and Dhiman, 1993).

**SNAIL**

The snails *Lymnaea acuminata* and *Indoplanorbis exustsus* are hosts of *Fasciola gigantica*, which causes fascioliasis. *Melania scabra* is also a vector snail (Muley, 1978). The various types of neem products alone or in a mixture (Bali *et al.*, 1985) or with *Cedrus deodara* and *Embelia ribes* have shown molluscicidal activity (Singh *et al.*, 1995) and may be effective against the diseases caused by the snails. The toxic effects of pure azadirachtin against snails are greater as compared to other neem products (Singh *et al.*, 1996).

**TERMITE**

Neem cake seems to act as a repellent barrier against *Odontotermes* spp. and *Microtermes obesi* (Gold *et al.*, 1989). A methanolic extract of neem oil with antifeedant activity was tried against the termite *Reticulitermes speratus* and was found effective (Ishida *et al.*, 1992).

**TICK**

Azadirachtin inhibited the onset of oviposition by only a few days in the case of the tick, *Amblyomina americanum* (Lindsay and Kaufman, 1988).

**REFERENCES**


19. PROTECTION OF FOOD MATERIALS

TRADITIONAL USES

It was a common practice, particularly in rural areas, to put dry leaves of neem between folds of dry cloth, or in stored grains and cereals to ward off various insects. In some parts of south India and Sri Lanka, fumigation with neem leaves was practiced instead. Vijayalakshmi et al. (1995) have described simple methods of treating gunny bags, storage bins and rooms with a solution of neem kernel powder or plastering the wall of bamboo baskets with a paste of cow dung and neem cake powder. Wealth of India (1948) gave a brief account of the use of neem in the protection of food items in daily use, including potato, without cold storage. It has been seen by experience that various treatments of neem products on cereals and legumes are harmless and do not effect palatability or cooking quality.

According to Islam (1993), neem may be used in Bangladesh for the protection of food materials from insects and worms during storage, in the following ways in the rural areas:

1. Fresh or dried leaves mixed with grains.
2. Spraying of neem oil in the storage area.
3. Rubbing a paste of seed or oil on the inside walls of the basket.
4. Plastering the hut with a mixture of clay, cow dung and neem seed powder.
5. Soaking the bags in the crude extract of neem seed.
6. Hanging of leaves or making a leaf mat in the storage room.
7. Fumigation with a mixture of neem leaves and oleo resins.
8. Washing or rinsing fresh fish with a water extract of neem.
9. Hanging bunches of leaves over the food or meat to repel flies.

Neem protects the stored grains and fresh food in various ways but it is not effective against all the causative organisms. It may be acting differently on different insects. In some insects it may prevent oviposition to varying degrees, may halt post-embryonic development completely and the emergence of the insects may be delayed for months. In others it deterred feeding and the worm died of starvation. In general, seed extract is more effective than the leaves. It suppressed the growth of fungi producing aflatoxins.

EFFECT ON CEREALS

In Nigeria, Ivbijaro (1983) mixed maize grain with dry ground seed of neem. It saved the maize from damage by the weevils *Sitophilus oryzae* for six months. Adult weevils placed on the maize treated earlier with neem had a very high mortality rate. Zhang and Zhao (1983) in China found that neem oil (5ml/kg), when mixed in stored rice, reduced the population growth of weevils considerably. In India, Jadhav and Jadhav (1984) observed that neem oil in various concentrations significantly inhibited the emergence of the pulse beetle *Callosobruchus maculatus* in chick pea (gram). Malik
et al., (1984) studied the anti-feedant and repellent properties of neem. Pandey et al. (1986) found the same effect with petroleum ether extract of neem on C. chinensis. Singh and Kataria (1986) experimented with the effect of deoiled neem kernel powder and leaf powder on the development of Trogoderma granarium. All the larvae died but the ethanolic extract showed higher toxicity as compared to the kernel powder; the neem leaf was least effective.

Singh et al. (1987) observed the activity of extract of neem against the insect Rhyzopertha dominica in stored grains. Gupta et al. (1988) tried neem and some other non-edible oils on storage of wheat seed and their germinability. Oils of neem and Butea frondosa at the rate of 5 ml/kg were most effective. The germination of seed was not affected by this treatment. Das (1989) showed that 1ml/100 gm neem was an effective surface protectant against C. chinensis in chick pea for six months. Cobbinah and AppiaiKwartang (1989) compared various neem products, along with others, against Sitophilus zeamais. The damage was less to maize treated with neem oil or neem wood ash as compared to the untreated one. But ash was less effective. Kossou (1989) also tried various plant parts like neem seed kernel, leaf, flower and bark against S. zeamais in maize.

The activity of neem extract of water and methylated spirit (commercial ethanol, mixed with methanol and the other solvents to make it unfit for human consumption) on C. maculatus S. oryzae, S. zeamais and Cylas punicollis for the protection of cow pea (Vigna unguiculata) and maize was studied. The extracts showed more activity in cow pea as compared to maize, and suppressed C. maculatus more than S. oryzae. There was no effect on C. punicollis (Makanjuola, 1989).

Jilani and Saxena (1990) tested the repellence of neem oil and neem based insecticide against R. dominica for eight weeks and found that these have a long range of effect because of persistence. Dabire (1992) tested the traditional claim about the effect of neem seed cake on the protection of cow pea during storage.

Kumar and Mehta (1993) found 4 percent neem leaves very effective against R. dominica in milled rice. High mortality of R. dominica and Tribolium castaneum by neem oil was also seen by Mohiuddin et al. (1993), while Kahare et al. (1993) tried 1 percent neem extract against C. chinensis and found considerable reduction in the number of eggs of this bruchid.

Dust of deoiled neem kernel was tested against three pests of wheat flour by Singh et al. (1993). The development of larvae of Tribolium castaneum was inhibited. The growth and development of first instar larvae of Trogonoderma granarium were completely arrested. It was effective against Corcyra cephalonica and Sitotroga cerealella. Chau (1993) also found neem preparations effective against insects of stored mung bean and rice and for the control of confused flour beetle (Tribolium confusum). Pandao et al. (1993) observed the efficacy of 5 percent neem extract against Exelastis atomosa in arhar (Cajanus cajan L.). Trung et al. (1993) described the development of neem pesticide for storage in Vietnam and Hu and Chiu (1993) in south China.

Belko (1994) mentioned that small farmers in Nigeria traditionally use neem leaves to control insects during the storage of cow pea. Neem oil at 1 percent concentration showed a significant reduction of egg hatching by the pulse beetle C. chinensis (Kachare et al., 1994). The toxic effects of acetone and methanol extracts of whole neem fruit
were tested by Majeed et al. (1994) on the pulse beetle *C. analis*, and these extracts were found to be quite effective, while Juneja and Patel (1994) treated green gram (*Vigna radiata*) with seed kernel powder against *C. analis* but it gave protection for three months only. Raju et al. (1994) observed the mortality of *Plutella xylostella* by commercial-grade neem oil, while Tabassum et al. (1994) tried neem preparations against *C. analis* and Naqvi et al. (1994) tried neem oil against it, and found them effective in controlling this pest.

Borker and Pawar (1995) found 1 percent neem seed powder to be a grain protectant against *C. chinensis*. Xie et al. (1995) tested three products containing different concentrations of azadirachtin and neem extract against three stored product insects, *S. oryzae*, *T. castamum* and *Cryptolestes ferrugineus*. The neem extract was more effective than azadirachtin, which showed that the activity was not due to azadirachtin only but because of other compounds in the neem also. In the case of azadirachtin, the activity was dose dependent, i.e. the higher the concentration, the greater the effect. Sharma (1995) observed that 10 percent neem seed kernel powder was effective in stored maize for all the insects but did not provide complete protection. Niber (1995) studied the protective effect of stored maize against *Prostephanous truncatus* Gangopadhyay (1995) studied the use of turmeric (*Curcuma longa*) and neem for safe storage of food grains. Singh et al. (1996, 1996a) found that extract of neem was effective against the lesser grain borer *R. dominica* in terms of lower fecundity, adult mortality and less grain damage.

**TREATMENT OF STORAGE AREA**

A herbal formulation from neem, *Pongamia pinnata* and *Vitex negundo* was tried by Dakshinamurthy (1993) for the storage of wheat and pulses free from worms like *S. oryzae*, *C. maculatus* and *Corcyra cephalonica* by impregnation of bags and surface treatment of storage bins. With this treatment, the level of infestation could be reduced considerably. Rajesh Kumar et al. (1994) observed the effect of soaking bags in a commercial neem preparation for the storage of rice. There was some protection.

**PRESERVATION OF FISH**

Okorie et al. (1991) placed Tilapia fish with neem seed powder and the insect *Dermestes maculatus*. The neem seed inhibited oviposition in the insect and killed the adults; most of the larvae in the fish did not develop and died within 30 days. Ward and Golob (1994) discussed various plant materials, including neem, to control insect infestation of cured fish.

**FUNGISTATIC ACTION**

**Dry Seed**

Seed stored in hot, humid and dark conditions often have fungal growth, sometimes consisting of *Aspergillus flavus* and *A. parasiticus*. These fungi produce a very highly
carcinogenic substance, aflatoxin. An aflatoxin content exceeding 20 parts per billion (ppb) in animal feed and 0.5 ppb in eggs and milk is not permitted in the United States. Neem seed very often harbour these fungi when fresh but neem leaf extract has been found to suppress the production of aflatoxin in groundnut (Gherwande and Nagaraj, 1987). Zeringue and Bhatnagar (1990) observed that in cotton bolls, fungal growth was unaffected by aqueous extract of neem but it caused 16 percent inhibition of aflatoxin production. Bansal and Sobti (1990) controlled the growth of Aspergillus spp. by soaking peanut seed in neem extract. Khan and Shah (1992) studied the antifungal activity of leaf extract of neem on seed mycoflora of neem. Fresh leaves blended with potassium phosphate solution, when added to the fungal growth medium, did not affect fungal growth but stopped the synthesis of aflatoxins (Anonymous, 1993). In vitro studies with fungi suggested that non-volatile neem leaf constituents inhibit aflatoxin biosynthesis in the early stages of biosynthetic pathways (Bhatnagar and Zeringue, 1993). Up to 92 percent suppression of aflatoxins by methanolic extract has been observed by Shankar Rao et al. (1994). There was no correlation between growth of A. flavus and aflatoxin production.

Fresh Fruit

Fresh fruits and vegetables are often spoiled during storage and transit, particularly in hot and humid conditions, due to fungal growth. Some studies have been conducted to see if the shelf life of these products can be prolonged by treatment with neem products. Hasabnis and D’Souza (1987) studied post-harvest storage by dipping Alphonso mango fruits in neem leaf extract and by lining bamboo packing baskets with neem leaves as cushioning material. Ali et al. (1992) evaluated neem oil, leaf extract and pericarp dust against isolates of Penicillium italicum, Alternaria alternata and Aspergillus niger from rotten fruit. Neem oil was as effective as thiabendazole in checking growth of these fungi in rotting tomatoes. Moline and Locke (1993) tested the antifungal properties of a hydrophobic seed extract against post-harvest mango and apple pathogens, Botrytis cinerea, Penicillium expansum, and Glomerella cingulata. Neem seed was as effective as calcium chloride. Apples dipped in 2 percent emulsion of a commercial product after harvesting and stored at 0°C for four months had 30 percent less decay than those kept at room temperature (Hohn et al., 1996).

REFERENCES


Toxicity determination of different plant extracts (saponin and juliflorine) and neem based pesticide Margosan OTM against stored grain pest Callosobruchus analis. *Proceedings of 14th Congress of Zoology held at the University of Karachi, Pakistan, 1–3 April, 1994.*


20. COMPOSITE PLANT FOR UTILIZATION OF NEEM

Neem fruit can be a very good raw material for a composite plant for the manufacture of various industrial and consumer products as given in Fig. 17. The properly dried decorticated seed will yield the seed coat and the kernel.

THE SEED COAT (HUSK)

It is very rich in cellulose, lignin, etc. and can possibly be put to the following uses.

(a) *As a fuel:* It has a low calorific value and does not burn easily, and is fed into the furnace of the boiler with a spade so that it does not form a lump but gets spread on the flame for easy combustibility. It can be used in places where slow heat for a long time is required, as in brick kilns.

(b) *Briquettes:* After powdering (Fig. 18A) it may be mixed in a mixing machine with other unwanted organic materials of the industry like discarded neem oil, seed cake or other agricultural wastes and pressed to form briquettes with the help of a machine (Fig. 18B). These briquettes can be used in the boiler for the steam required for heating purposes.

(c) *The ash:* Seed coat is very rich in inorganic matter, and yields a substantial amount of ash, which if not disposed of properly can not only create storage problem but may cause pollution by increasing the amount of suspended particles in the air. The best use of it can be as an ingredient in cement industry in place of fly ash obtained from thermal electric plants. It may be incorporated in clay for the brick industry.

(d) *Particle Board:* The seed coat has good mechanical strength and may be incorporated in the mixture of wood shavings used for making particle boards for thermal insulation by mixing with a synthetic resin, such as phenol formaldehyde, as adhesive. It may be powdered before use or treated with acid or alkali so that it forms a homogenous mass with the particle board mixture.

(e) *Manure:* It can be incorporated in soil amendment formulations, particularly for soil rich in clay or having nematodes. In the former case, it loosens the soil particles while in the latter case it may destroy nematodes by the traces of limonoids contained in it.

(f) *As a raw material:* Some new methods can be developed for other industrial products after chemical treatment.

SEED OIL

Earlier, neem oil on an industrial scale in India was obtained mainly by solvent extraction, but with the demand for pesticides, modifications to the technology have been suggested so that azadirachtin and allied compounds can also be recovered.
Depending on the requirement, the oil from seed kernel powder can be extracted either by solvents or by cold mechanical pressing.

**By Solvents**

As given in the process for the isolation of azadirachtin in Chapter 3 on chemical constituents, the seed is powdered to a specific particle size. If it is coarse, proper extraction will not take place; on the other hand, fine powder may clog the pipes and filtration of oil may be difficult. When seed is extracted with solvents, limonoids, other constituents and the oil get dissolved in it, leaving the seed cake, free of these, in the extractor. The solvent from this mixture can be recovered by distillation. For pesticide formulations, pesticides are removed from the oil by using other solvents, and standardized extract may be used in the desired products.

The fatty oil obtained may be used in the manufacture of soap by mixing with other oils, but for good quality soap, noodles are preferred which can be manufactured by treating the oil with sodium hydroxide. Only the fatty acids react with caustic soda, leaving most of the impurities, which affect the quality of the soap. These soap noodles are mixed with various ingredients like silicates, polyethylene glycols, surfactants, color and fragrances, and kneaded together. The mixture is fed into an extruder to get the soap cakes. The oil can be split further by various chemical treatments to get free fatty acids, which are very important raw materials in the detergent, cosmetic and rubber industries. Glycerine is an important by-product. The fatty acids may be converted into lubricants after hydrogenation or may be used for the manufacture of olein and high melting styrene.
Figure 18 A, Machine for making fine powder of neem seed coat and other agricultural waste; B, briquette-making machine. Abbreviations: AB=air balloon, GI= grinder, WS=waste stuff, PM=pressing machine.
Cold mechanical process

In this, the seed kernels are subjected to mechanical pressing, and no solvents, chemicals or heat is used. By this process, oil rich in limonoids is obtained but the yield is low. This oil may be further enriched by azadirachtin and the other compounds obtained from solvent extraction for pesticide formulations or may be used as such or in the form of an emulsion as a spraying agent or in mosquito-repellent products, by incorporating it in other oils or in a cream or paste base. The soap manufactured from this oil can be effective in skin diseases. The oil may be used as a lubricant for household and agricultural purposes.

SEED CAKE

The cake obtained after solvent extraction has a small amount of oil and limonoids and after washing with water can be incorporated into poultry and cattle feed, as given earlier, but the cake obtained from the cold process is rich both in oil and limonoids and may be extracted with ethanol for use in pesticide formulations and the extracted seed cake may be used as a denitrifying agent or as a manure. For a nematode suppressant formulation, neem cake mixed with karanj (Pongamia glabra) seed cake and tobacco waste may be used. The seed cake may be mixed with urea for denitrifying purposes in fertilizers. Neem cake manure has better customer acceptance for household purposes and indoor plants because it lacks the malodor of cattle manure and has good nutritive value for the vegetation.
21. PATENTS ON NEEM

EARLIER PATENTS

As given in the list of patents in Appendix 1 of this chapter, Indian workers were the first to isolate neem bitters in 1945–46 and they got patents for these, but the patents could not be effectively utilized by the industry because no product which might have good demand could be developed from these bitters. Later on, the insecticidal value of neem was observed by Indian scientists but they were not granted a patent for their discovery, because of the common use and common knowledge clause in the Indian Patent Act. The Indian Central Insecticides Board did not register neem products under the Insecticide Act 1968 (Vijayalakshmi et al., 1995). In England and Japan, a few bodycare products with neem as an ingredient were patented.

NEEM PATENT CONTROVERSY

The controversy started when a patent was granted to W.R. Grace and Co. for a neembased pesticide. The question arose whether a product of natural origin, the traditional use of which has been known since ancient times, can be patented. More than 200 organizations from 35 countries raised their voice, against this type of patent, which was called “Corporate Colonialism”, “Genetic Imperialism” and “Folk-wisdom Piracy”. Vijayalakshmi et al. (1995) have given details of these activities, as shown in the poster (Fig. 19).

W.R. Grace and Co. defended their patent right under the pretext that they had developed a process for the isolation of a stable form of azadirachtin and for increasing the shelf life of the product. In 1993, a Congressional Research Service (CRS) reported to the United States Congress put forth the view that synthetic form of natural products like azadirachtin may be patentable (Vijayalakshmi et al., 1995).

In an editorial in the magazine Nature (Anonymous, 1995), patents on neem were justified on the ground that companies have to spend a huge amount of money on research and development for this type of product, and have to protect this investment by taking out patents on their discoveries. The author agreed that in this way, general knowledge becomes a private commodity.

In reply to this, Balasubramanian (1995) under patents and indigenous lore mentioned that patents for natural products should be granted on their genuine originality and not to the extent they conflict with traditional knowledge systems, turning public goods into a private commodity. The author feared that large-scale purchase of the raw materials by multinationals, which have huge resources at their disposal, may take the price of unprocessed neem beyond the reach of farmers who may be forced to rely on the commercial product rather than on traditional recipes.
NO PATENTS ON NEEM PRODUCTS

Poster against granting patent rights to neem products

 Patents on neem products are based on a bias against non-Western knowledge systems which are treated as ‘obvious’ and ‘natural’.

Such patents will lead to monopoly production and undermine the natural resource and market base for the small scale sector.

The neem patents are a blatant case of intellectual piracy.

The Research Foundation for Science, Technology & Natural Resource Policy, A-60 Hauz Khas, New Delhi 110 016

Figure 19 Poster against granting patent rights to neem products
The role played by advanced technology in products developed from natural resources like neem have been discussed by Hoyle (1995). The author quoted an instance of India’s own Central Drug Research Institute which has transferred the technology for natural spermicide DK-1 to two companies. Crsepi (1995) suggested ways by which the patents on products developed by biotechnology can be defended.

The whole issue of patents for neem products has been discussed by Johnston (1995). According to the author, the Neem campaign feels that the utilization of traditional knowledge, which is an intellectual contribution of third world countries, for commercial gain, is a sophisticated name for modern piracy. The Foundation on Economic Trends in Washington DC requested the US Patent and Trademark office to overturn patents on neem and sought legislation to bar this type of patent in future (Anonymous, 1996).

APPENDIX 1

Patents on Neem (in Chronological Order)


PPG Inc. (1991) Concentrated matter in oil microemulsion that forms storage stable oil in water emulsion consisting of 50% to 90% by weight of neem oil containing the pesticide azadirachtin, among other ingredients. US Patent No. 5110591.


REFERENCES


