Botanical immunodrugs: scope and opportunities

Bhushan Patwardhan and Manish Gautam

Modulation of the immune system can be addressed through a variety of specific and non-specific approaches. Many agents of synthetic and natural origin have stimulatory, suppressive or regulatory activity. There is growing evidence that drugs or biological agents capable of modulating single pathways or targets are of limited value as immune-related therapies. Systems biology approaches are now gaining more interest compared with monovalent approaches, which can be of limited benefits with complications. This has stimulated interest in the use of ‘cocktails’ of immunodrugs to restore immunostasis. Botanicals are chemically complex and diverse and could therefore provide appropriate combinations of synergistic moieties useful in drug discovery. Here, the importance of traditional medicine in natural product drug discovery related to immunodrugs is reviewed.

The immune response requires the timely interplay of multiple cell types within specific microenvironments to maintain immune homeostasis. The selectivity and flexibility that is necessary to regulate cell traffic under homeostatic and diseased conditions is provided by the differential distribution and regulated expression of cytokines and their receptors. As a consequence, cytokines are responsible for the development of phenotypes and are, therefore, logical targets for therapeutic immune modulation [1]. Immunodrugs include synthetic organics, biologicals, such as cytokines and antibodies, and microbial and botanical natural products, which influence immunoregulatory cascades to bring about their specific stimulatory, suppressive or regulatory effect. Immune suppression has been widely studied for clinical applications [2]. Historically, botanicals have been a mainstay of drug treatments and are currently receiving attention as sources of synergistic combinations. Here, recent developments in botanical immunodrug discovery are reviewed.

Immunodrugs in cancer

Although the treatment of cancer using active immunotherapy has had limited success, passive immunotherapy with antibody and cytokine therapies brings new hope. The most widely studied approaches consist of whole-cell vaccines, dendritic cell-based immunotherapy and peptide vaccines. Many clinical studies have demonstrated safety but not necessarily the efficacy of such strategies [3]. Moreover, there is emerging consensus that the most efficacious therapy should activate several components of the immune system [4]. Cytokine therapy in cancer is another attractive approach; however, balancing optimal doses to avoid toxic reactions remains challenging [5]. Several cytotoxic drugs have immunomodulatory effects at relatively low doses and can exert immunity-dependent curative effects in animal models of cancer. Combination therapies involving low-dose anticancer agents and cytokines have demonstrated some benefits. For example, combinations of appropriate regimens of doxorubicin plus interleukin (IL)-2...
or tumour necrosis factor (TNF) could be curative and produced a life-long immunological memory in an EL4 lymphoma C57BL/6 mouse model. Many researchers have demonstrated that the induction of T-helper (Th) 1-promoting cytokines, using specific adjuvants, can enhance antitumour immunity and can prevent or reduce tumour growth. These trends substantiate the possibility of establishing combination regimens based on low-dose anticancer drugs, specific cytokines and immunological adjuvants [6].

**Immunodrugs in infection**

Immunomodulators could also have beneficial roles in the prevention and treatment of infectious disease. A diverse array of synthetic, natural and recombinant compounds are available. Of the synthetic immunomodulators, Levamisol, Isoprinosine, Pentoxifilie and Thalidomide are some of the more significant [7]. Microbial immunomodulators, such as bacille Calmette-Guérin, have been in use for years for non-specific activation of the immune system in some forms of cancer (bladder) and infectious disease [8].

Targeting cytokines is now considered to be one of the logical approaches for the prevention and treatment of infectious disease. Some of these substances, such as granulocyte colony-stimulating factor (G-CSF), interferons, IL-2, various chemokines, synthetic cytokine phosphate-guanosine (CpG) oligodeoxynucleotides and glucans are being investigated in preclinical and clinical studies [9]. Interferons are widely used for the treatment of chronic infections, particularly hepatitis B, D and C viruses [10]. PEG-interferon is a recently developed pharmaceutical preparation that has proven beneficial in non-responsive patients, particularly in those with cirrhosis or hepatitis C virus genotype 1 [11]. Other cytokines, such as IL-1, IL-2 and IL-17, have shown potential in augmenting immune responses in various infectious conditions and malignancies. The therapeutic effect of cytokine blockers is also reported in septic shock [12]. A recent study on hepatitis C has shown that interferon-γ (IFN-γ), in combination with ribavirin, induces a higher percentage of lymphocyte activation [13] than for IFN-γ alone. Such approaches are currently being examined for their potential to boost host immune response to fight infection.

**Immunomodulation and inflammation**

Many immune targets have been identified as having potential for the central control of inflammation. Targeting activated T-cell subsets was considered to be one of the most rational approaches and biologicals, such as monoclonal antibodies (mAbs) against CD4, CD5, CD7, CD25 and CD52, were evaluated in patients with autoimmune disorders. Limited clinical benefit and complications, such as the prevalence of opportunistic infection and/or malignancies, were observed and the hope of reprogramming the host immune response remained unfulfilled [14,15].

Studies on targets related to leukocyte infiltration, such as leukocyte-function associated antigen-1 (LFA-1), CD11a/CD18 (adhesion-receptor–counter-receptor pair) and intercellular adhesion molecule-1 (ICAM-1) CD54, are in progress; however, the development of humanized antibodies and their long-term safety evaluation have yet to be established [16]. Cytokine therapy, such as anti-TNF-α or IL-1, has been an attractive treatment option; however, an optimal treatment regimen with respect to dosage, interval and particularly long-term safety needs to be explored [17–19]. The use of the effector functions of Th cells is now considered to be one of the more promising innovative therapeutic strategies. Thus, the idea of switching Th1-dominated responses into Th2-mediated responses appears intriguing. This approach has been studied in various animal models of autoimmune diseases but clinical validation has not been achieved [20–22]. Newer targets central to innate and adaptive immunity, such as Toll-like receptors (TLR) and the complement system and nuclear factor-κB (NF-κB) activation, are being studied [23–25]. Combination approaches to the treatment of inflammatory diseases appear promising; in particular, combining methotrexate with TNF-α inhibitors has provided some encouraging results [26].

**Immunodrugs and vaccine adjuvants**

The combined use of vaccines and immunomodulators are innovative strategies in vaccine design and development. Many synthetic, biological and natural immunomodulators are under evaluation as vaccine adjuvants. Administration of cytokine genes along with DNA vaccines has been shown to achieve selective modulation of T-cell responses [27]. Moreover, innate immunity targets, such as TLR, and their modulation are currently being researched for their ability to provide effective adjuvant action. QS-21 and glucans are experimental adjuvants currently under clinical evaluation with different vaccines [28–30].

**Immunostasis: targets and regulation**

Th lymphocytes are divided into distinct phenotype sub-sets of Th1 (e.g. IFN-g, IL-2 and TNF-a) and Th2 (e.g. IL-3, IL-4, IL-5, and so on) effector cells. This classification is based on their functional capabilities and cytokine profiles. Th1 cells drive the cellular immunity to fight intracellular organisms, eliminate cancerous cells and stimulate delayed-type hypersensitivity reactions. By contrast, Th2 cells drive humoral immunity and upregulate antibody productions to fight extracellular organisms. T-cell homeostasis or immunostasis requires a fine balance between Th1–Th2 response and such agents could exhibit stimulatory, suppressive or regulatory activity [31]. Currently, much of the literature supports the view that Th1–Th2 is essential to immunostasis and many of the T-cell-directed therapies have provided modest clinical benefits [32]. Although this view of signal conversion looks relatively simplistic, mediators of signal transduction do not interact in a linear
manner but within a biochemical matrix. As a consequence, each cytokine has multiple functions and their modulation would be expected to result in diverse therapeutic functions. Such complex crosstalk between signalling networks imposes challenges for the discovery of optimal therapeutic interventions [33]. In cancer, researchers have arrived at a consensus that reconstitution of function is important, rather than reconstitution of cells or cytokines, and the most efficacious therapy will be one in which multiple responses of the immune system are activated. Trends indicate that future immunotherapy should involve cocktails of drugs that concurrently and/or simultaneously address vital components of the immune matrix [34]. This will require synergistic combinations for homeostatic regulation of the immune system. Historically, botanicals have facilitated and enriched the drug discovery process and, we propose, should be explored as sources for synergistic combinations.

**Botanical immunodrugs**

Many herbal preparations alter immune function and display an array of immunomodulatory effects. In various *in vitro* and *in vivo* studies, herbal medicines have been reported to modulate cytokine secretion, histamine release, immunoglobulin secretion, class switching, cellular co-receptor expression, lymphocyte expression, phagocytosis, and so on [35]. A recent study, using a transgenic mouse model of melanoma, showed that the anticancer effects of popular Kampo medicine were mediated via an enhanced antigen-specific antitumour cytotoxic T-lymphocyte response [36]. Botanicals produce a diverse range of natural products with antimicrobial and immunomodulating potential, including isoflavonoids, indoles, phytoestrogens, polyaccharides, sesquiterpenes, alkaloids, glucans and tannins. There are many immune-related conditions with a high unmet clinical need and this is particularly true in the case of new viruses and the phenomenon of increasing antibiotic

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**FIGURE 1**

Chemical structures of immunoactive leads from botanicals.
resistance. Botanical immunomodulating agents might be able to provide an alternative to costly immunotherapeutics.

**Natural product drug discovery**

There are two major ways of bioprospecting natural products for investigation. First, the classical method that relies on phytochemical factors, serendipity and random screening approaches. Second, the use of traditional knowledge and practices as a drug discovery engine: this is also known as an ethnopharmacology approach, which is time and cost effective and could lead to better success than routine random screening [37,38]. Traditional methods, for example, Chinese medicine, Japanese Kampo and Indian Ayurveda, are becoming important bioprospecting tools [39]. Ayurveda gives a separate class of immunomodulatory botanicals named Rasayanas. Several botanicals from these texts have been studied for their immunomodulatory properties and have the potential to provide new scaffolds for safer, synergistic, cocktail immunomods [40]. In subsequent sections, examples are provided of important botanicals based on traditional knowledge that have been studied for the bioprospecting of potential immunomods. These botanicals have an array of compounds with diverse activities directed at various targets of the immune matrix, in cancer and in infection and inflammation.

**Glycyrrhiza glabra**

The root *Glycyrrhiza glabra*, commonly known as liquorice, has been used since ancient times in Indian, Chinese, Egyptian, Greek and Roman medicine. It is prominent in Ayurveda as Rasayana with cytoprotective and demulcent effects and is a popular home remedy for minor throat infections. Biologically active substances in liquorice roots are studied for the bioprospecting of potential immunomods. These botanicals have an array of compounds with diverse activities directed at various targets of the immune matrix, in cancer and in infection and inflammation.

Many researchers have reported response-modifying activity in flavonoids and chalcones isolated from the root extract. Modulation of the Bcl-2/Bax family of apoptotic regulatory factors, by components of the root, has been suggested as a possible mechanism for its reported cytoprotective activity [46]. These activities include antioxidand, chemopreventive and antimicrobial activities [47].

Chemical modification of GL and GA has been tried and significant improvement in anti-inflammatory, antiallergic, and antiulcer activities was observed [48]. These observations indicate immune-modulating and biological response-modifier activities associated with GL [49].

**Uncaria tomentosa**

*Uncaria tomentosa*, also known as cat’s claw, is from the highlands of the Peruvian Amazon and has been used by natives for hundreds of years to treat immunologic and digestive disorders. It was found that two chomotypes of this plant occur in nature, each with different alkaloid patterns [50]: the roots of one type contain pentacyclic oxindoles and the other contains tetracyclic oxindoles, reported to have antagonistic activities. Tetracyclic oxindole alkaloids dose-dependently reduce the activity of pentacyclic oxindole alkaloids on human endothelial cells [51].

Aqueous extracts and mixtures of oxindole alkaloids have shown positive influence on IL-1, IL-6 and IFN-γ production, suggesting immunoregulatory activity. In one clinical study, extract exhibited immune adjuvant activity with pneumococcal vaccine resulting in enhanced lymphocyte/neutrophil ratio and persistent antibody titre responses towards 12 pneumococcal serotypes [52].

In *in vitro*, *in vivo* and gene expression studies on extracts of this plant indicated that anti-inflammatory activity is mediated through negation of NF-κB activation and suppression of TNF-α synthesis [53,54]. Randomized clinical studies on a purified extract, rich in pentacyclic alkaloids, demonstrated safer and moderate benefit in patients with active rheumatoid arthritis compared with those taking sulfasalazine or hydroxychloroquine [55].

Cytoprotection was observed in extracts devoid of alkaloids. The oxindole alkaloid-free fraction modulated apoptosis, tumour cell proliferation and DNA repair processes, leading to cytoprotection in chemically induced leucopenia rat model [56]. Extracts have also shown promising antitumour activity mediated via selective induction of apoptosis [57].

**Echinacea spp.**

The *Echinacea* plant is a member of the *Compositae* family; the three species of medicinal interest being *Echinacea angustifolia*, *Echinacea purpurea* and *Echinacea pallida*. Most uses of *E. purpurea* are based on its reported immunological properties. There are four types of constituents purported to be pharmacologically active molecules: phenolic caffeic acid derivatives, alkylamides and isobutylamides, polysaccharides and glycoproteins. Limited experimental evidence
of immunostimulatory activity exists for caffeic acid derivatives and alkylamides [58]. By contrast, *Echinacea* polysaccharides were found to directly activate non-specific immune cell types, such as monocytes, macrophages and natural killer (NK) cells [59]. This characterization of *Echinacea* polysaccharides is the best demonstration of in vitro bioassay activity yielding reproducible *in vivo* pharmacological effects [60,57]. Several randomized trials have reported health benefits of *Echinacea* extracts in upper respiratory tract infections [61].

*Withania somnifera*

*Withania somnifera* (WS) – known as ashwagandha, Indian ginseng and winter cherry – is also classified as Rasayana in Ayurveda. The major biochemical constituents of WS root are steroidal alkaloids and steroidal lactones known as withanolides. Much of WS pharmaceutical activity has been attributed to withaferin A and withanolide D. WS is reported to have immunomodulatory, antitumour, cytoprotective and antioxidant properties [62]. All these activities are thought to be involved in the overall immunoregulatory properties of WS.

Several preclinical studies have examined the cytoprotective potential of WS. In one study, WS exhibited myeloprotection in tumour model without compromising antitumour efficacy of cyclophosphamide, azathioprin or prednisolone [63]. In another instance, cytoprotection against experimental skin cancer was observed, where reduced levels of glutathione, superoxide dismutase, catalase and glutathione peroxidase returned to normal following WS administration [64].

WS exhibited modulatory effects on cytotoxic lymphocyte production leading to reduced tumour growth. Withaferin A was found to be better than doxorubicin in inhibiting growth of breast and colon cancer cell lines [65]. A recent study suggested that the increased production of inducible nitric oxide synthase was one of the possible mechanisms for the increased cytotoxic effect of macrophages exposed to WS extracts. Moreover, withaferin A in combination with radiotherapy increased the response to radio-resistant tumours [66].

WS treatment in normal and tumour-bearing mice showed a positive influence on NK cell activity resulting in enhanced cell killing [67]. Many studies, including our own, have demonstrated the immunomodulatory potential of WS, resulting in increased haemolytic titres, inhibition of delayed type sensitivities and an increase in phagocytic activity of macrophages [68,69]. We also observed that animals receiving WS showed immunoprotection to *Bordetella pertussis* infection, as evident by increased antibody titres and higher survival percentage [70]. In a recent study, Immun-21, a polyherbal formulation containing WS, exhibited immunomodulatory activity leading to modest clinical benefits in groups of HIV patients [71]. These observations suggest that WS could be used as an immunological adjuvant with multiple therapeutic benefits in cancer, infection and AIDS. In a comparative pharmacological investigation of WS and ginseng, the WS-treated group showed better anabolic and antigastress activity than ginseng with additional anti-inflammatory activity [72].

*Withania somnifera* and ginseng, the WS-treated group showed better anabolic and antigastress activity than ginseng with additional anti-inflammatory activity [72].

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**TABLE 1**

<table>
<thead>
<tr>
<th>Compound name</th>
<th>Source</th>
<th>Chemical class</th>
<th>Activity</th>
<th>Refs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ginsan</td>
<td><em>Panax ginseng</em></td>
<td>Polysaccharide</td>
<td>Anticancer</td>
<td>[85]</td>
</tr>
<tr>
<td>Triptolide</td>
<td><em>Tripterygium wilfordii</em></td>
<td>Diterpenoid triepoxide</td>
<td>Rheumatoid arthritis and leprosy</td>
<td>[86]</td>
</tr>
<tr>
<td>Mistletoe lectin</td>
<td><em>Viscum album</em></td>
<td>Lectin</td>
<td>Cytostatic and apoptotic</td>
<td>[87]</td>
</tr>
<tr>
<td>Piperine</td>
<td><em>Piper longum</em></td>
<td>Alkaloid</td>
<td>Antitumour and bioavailability enhancer</td>
<td>[88]</td>
</tr>
<tr>
<td>Matrine</td>
<td><em>Sophora alopecuroides</em></td>
<td>Alkaloid</td>
<td>Antiviral</td>
<td>[89]</td>
</tr>
<tr>
<td>Sinomenine</td>
<td><em>Sinomenium acutum</em></td>
<td>Alkaloid</td>
<td>Arthritis and rheumatoid arthritis</td>
<td>[89]</td>
</tr>
<tr>
<td>Artemisinin</td>
<td><em>Artemisia annua</em></td>
<td>Sesquiterpene lactone</td>
<td>Psoriasis and autoimmune disorders</td>
<td>[90]</td>
</tr>
<tr>
<td>ASP</td>
<td><em>Acanthopanax senticosus</em></td>
<td>Polysaccharide</td>
<td>Antitumour</td>
<td>[91,92]</td>
</tr>
<tr>
<td>Apocynin</td>
<td><em>Picrorhiza kurroa</em></td>
<td>Iridoid glycoside</td>
<td>Cytoprotective and antitumor</td>
<td>[93]</td>
</tr>
<tr>
<td>Shatavarin</td>
<td><em>Asparagus racemosus</em></td>
<td>Triterpenoid saponin</td>
<td>Immunostimulant and vaccine adjuvant</td>
<td>[94]</td>
</tr>
<tr>
<td>KRN7000™ (Kirin Brewery)</td>
<td>Sponge (Agelas mauritianus)</td>
<td>α-Galactosylceramide</td>
<td>Anticancer</td>
<td>[80]</td>
</tr>
</tbody>
</table>

Much of the work has been conducted on berberine, jatrorrhizine, tinosporaside and columbin. The possible mechanism of immunomodulatory activity was elucidated as activation of macrophages, leading to increases in granulocyte–macrophage colony-stimulating factor (GM-CSF), leading to leukocytosis and improved neutrophil function. TC is also reported to inhibit C3-convertase of the classical complement pathway [77]. In a recent study, a polysaccharide (α-D-glucan) derived from TC resulted in activation of NK cells, complement system, Th1-pathway cytokines, coupled with low nitric oxide synthesis [78]. Antitumour activity of TC was evaluated in cultured HeLa cells and revealed that the effect of extract was comparable and better than doxorubicin treatment. Hepatoprotective activity of TC is also reported to inhibit C3-convertase of the classical complement pathway [77]. In a recent study, a polysaccharide (α-D-glucan) derived from TC resulted in activation of NK cells, complement system, Th1-pathway cytokines, coupled with low nitric oxide synthesis [78]. Antitumour activity of TC was evaluated in cultured HeLa cells and revealed that the effect of extract was comparable and better than doxorubicin treatment. Hepatoprotective activity of TC against carbon tetrachloride-induced liver damage has also been reported [79].

Other immunomodulatory leads from botanicals sources, which have shown potential as cytoprotectives, antitumour, anti-infective and anti-inflammatory agents are given in Table 1 and Figure 1. These are only a fraction of botanical immunodrugs and the scope of these agents remains quite vast.

### Immunotherapy scenario

(i) Host immune response can be modulated for clinical benefit.
(ii) Monovalent therapeutic approaches might not be effective.
(iii) Combination approaches offer synergistic efficacy and safety.
(iv) Cytokine therapy remains to be optimized.

### BOX 2

**Ayurveda and Rasayana**

Ayurveda means science of life in Sanskrit (Ayur means life; veda means science) and aims at the holistic management of health and disease. It remains one of the most ancient medical systems widely practiced in the Indian subcontinent and has a sound philosophical, experiential and experimental basis. Charak samhita and Sushrut Samhita (100–500 BC) are main Ayurvedic classics, which describe over 700 botanicals along with their classification, pharmacological and therapeutic properties.

Rasayana therapy is one of the eight branches of Ayurveda and generally means nourishing and rejuvenating drugs with multiple applications for longevity, memory enhancement, immunomodulation and adaptogenic. Many researchers have supposed neuroendocrine immune axis theory to explain Rasayana action.

### Conclusion

Analysis of current immunomodulating strategies indicates that monovalent approaches in isolation are unlikely to restore immunostasis or attain status of complete therapy (Box 1). This complements emerging systems biology approaches, which holistically monitor operating biological processes as an integrated system. It is likely that multiple immune-modulating strategies will be necessary to achieve clinical success owing to complex interplay between pathways. We need designer drugs, which involve safer, curative and synergistic combinations. Bioprospecting will have a major role in identifying such combinations. The botanical immunodrugs discussed here have such potential because they offer additional therapeutic benefits to address associated conditions such as infection, inflammation and cancer. Botanical extracts provide cytoprotective, anticancer, anti-inflammatory and anti-infective activities in addition to immunoregulatory activity. For example, crude extract of TC has immunostimulant, anticancer and cytoprotective activities, where cytoprotective activities are attributed to polysaccharides and immunostimulant activity is attributed to diterpenoids [76]. Recently, researchers observed antimicrobial activity coupled with immune modulation [80]. Bryostatins isolated from the marine organism *Bugula neritina* have antineoplastic activity along with mimicking the effect of recombinant human GM-CSF [81]. Three polysaccharides (krestin, lentinan and schizophyllum) developed by Japanese scientists from mushrooms are in clinical use as adjuvants in chemotherapy. These polysaccharides have now been developed into multipurpose medicines with cytostatic, anti-inflammatory and antithrombotic activities [82]. Similarly, polysaccharides and oxygenated triterpenoids derived from *Ganoderma lucidum* (a fungus from traditional Chinese medicine) showed broad spectrum pharmacological functions [83]. These examples underline the importance of bioprospecting for newer synergistic combinations and pharmacological agents. Although many researchers have studied synergism and antagonism in botanicals, systematic scientific investigations on pharmacodynamics, kinetics, dosing...
and interactions are required. Traditional knowledge, particularly from the great traditions of, for example, Ayurveda and China, will have an important role in bioprospecting. Rasayanas are a selected class of botanicals from Ayurveda with putative capabilities to rejuvenate, promote longevity and health (Box 2). A golden triangle, with the integration of modern medicine, traditional medicine and the robust use of science and technologies with a systems biology approach, could open up new opportunities for immunodrugs and therapy (Figure 2) [84].

In short, botanical immunodrugs could provide a unique opportunity to bioprospect diverse and synergistic chemicals moieties, which in combination might act on multiple targets and improve the therapeutic spectrum. Traditional knowledge and practices bring experiential wisdom to provide a safer and more cost effective platform for newer scaffolds and immunodrugs discovery.

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