REVIEW

Insects: an underrepresented resource for the discovery of biologically active natural products

Lauren Seabrooks\textsuperscript{a,*}, Longqin Hu\textsuperscript{a,b,**}

\textsuperscript{a}Department of Medicinal Chemistry, Ernest Mario School of Pharmacy, Rutgers, The State University of New Jersey, Piscataway, NJ 08854, USA
\textsuperscript{b}Cancer Institute of New Jersey, New Brunswick, NJ 08901, USA

Received 6 March 2017; received in revised form 27 April 2017; accepted 2 May 2017

Abstract Nature has been the source of life-changing and -saving medications for centuries. Aspirin, penicillin and morphine are prime examples of Nature's gifts to medicine. These discoveries catalyzed the field of natural product drug discovery which has mostly focused on plants. However, insects have more than twice the number of species and entomotherapy has been in practice for as long as and often in conjunction with medicinal plants and is an important alternative to modern medicine in many parts of the world. Herein, an overview of current traditional medicinal applications of insects and characterization of isolated biologically active molecules starting from approximately 2010 is presented. Insect natural products reviewed were isolated from ants, bees, wasps, beetles, cockroaches, termites, flies, true bugs, moths and more. Biological activities of these natural products from insects include antimicrobial, antifungal, antiviral, anticancer, antioxidant, anti-inflammatory and immunomodulatory effects.

© 2017 Chinese Pharmaceutical Association and Institute of Materia Medica, Chinese Academy of Medical Sciences. Production and hosting by Elsevier B.V. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
1. Introduction

The chemical and structural diversity of natural products are unparalleled by any synthetic based library and have inspired a wealth of life-changing and -saving medicines for centuries. In 1803, the remarkable pain-relieving natural product, morphine, was isolated from the Papaver somniferum plant and in the 1870s served as the template for codeine. Two decades later, salicin, the medicinally active natural product from the willow bark, Salix alba, was isolated and characterized by Henri Leroux. From this acetylsalicylic acid, an anti-inflammatory agent known commercially as aspirin was synthesized by the Bayer scientist Felix Hoffman. Aspirin has been a commercial success for the Bayer Company since its market debut in 1899. Thirty years later penicillin was isolated from the fungus Penicillium notatum by Alexander Fleming and proven to be a groundbreaking antibiotic by Howard Florey and Ernest Chain. This discovery revolutionized medicine, earning Fleming, Chain and Florey the 1945 Nobel Prize in physiology or medicine for their efforts. Then in 1971, the most effective malarial treatment to date was discovered while scientists were reviewing ancient tomes on traditional Chinese medicine. It was within these texts that artemisinin was identified as a promising drug discovery lead. The head scientist, Youyou Tu, earned the Lasker prize in 2011 and the 2015 Nobel prize in physiology or medicine for this impressive discovery.

The aforementioned examples and the fact that 33% of drugs approved from 1981–2010 and a staggering 68% of all approved antibacterial drugs are natural products or derivatives thereof prove that nature is an important source of new target molecules. However, the focus of natural product medicine has been overly focused on terrestrial plants, various algae and fungi. An under-represented source of inspiration, entomotherapy, has been in practice for as long as and often in conjunction with medicinal plants. Entomotherapy is the use of insects as medicine and is an important alternative to modern therapy in many parts of the world including India, Mexico, Korea, China, Spain, Brazil, Argentina, Ecuador and various African countries today. Honey, a sweet bee byproduct, is valued as an antioxidant and antimicrobial agent, suitable for the battle against heart disease and skin disorders. Propolis, a gap filling “glue” used in the construction of the beehive, has been evaluated extensively for its potential antioxidant, antimicrobial, anti-inflammatory, cardioprotective, immunomodulatory and antiangiogenic properties. Venom, especially from bees, has been studied as an alternative medicine for inflammation and cancer therapy. Success has also been achieved in identifying biologically active isolates from insect bodies, excretions and secretions. Antimicrobial peptides, an innate component of insect immunity within the hemolymph (blood), have shown activity against fungi, parasites, viruses and most importantly antibiotic resistant bacteria. Insect toxin isolates pederin and cantharidin from defense secretions of the rove and blister beetles, respectively, have peaked scientific interest as potential anticancer therapies.

Insects as natural product resources have also been featured in reviews published from 2010–2015. However, the scope of the aforementioned articles often includes other arthropods, covers a limited selection of insects, focuses heavily on insect byproducts and incorporates uses outside of pharmaceutical applications. This review focuses on the continued cultural use of insects as medicine worldwide, validation of their medicinal properties and identification, characterization and synthesis of insect natural products and derivatives.

Like plants and other invertebrates, insects are hosts for many types of microbial endophytes that include bacteria and fungi. These endophytic microorganisms may be involved or working together with their insect hosts in the production of a natural product reported. The potential involvement of endophytes should be taken into account when we isolate the natural products from insects.

Insects belong to the kingdom Anamalia, phylum Arthropoda, and class Insecta. Members of this class are further divided into 29 orders though 81% belong to one of the following four orders: Coleoptera, Diptera, Hymenoptera and Lepidoptera. This review article is split into sections based on the different orders and occasionally further subdivided by families. Sections are titled with scientific nomenclature with common names in parentheses.

2. Hymenoptera (bees, wasps, ants and sawflies)

2.1. Symphyta (sawflies, wood wasps, and horntails)

The family Symphyta is the smallest of the hymenoptera representing only 7% (8000–10,000 species) of the order. Like ants, they feed on leaves or burrow into wood earning them the position as pests by most foresters. In 2013, researchers out of China and Australia found medicinal value in these pests. Two small molecules with strong antimicrobial activity (Fig. 1) were isolated from the methanolic extract of Australian sawfly larvae. The novel macrocarpal (1) and known grandinol (2) were evaluated against Bacillus subtilis and exhibited inhibition zones of 8.0 and 7.0 mm, respectively.

Additionally, the hemolymph of sawfly larvae has been found to be rich in phenolic compounds, such as novel flavonol oligoglycosides, flavonoid glycosides and naphthodianthrones, all with known potential health benefits that were not evaluated in the studies.

2.2. Formicidae (ants)

Ants are one of the most familiar members of the hymenoptera order. This is a social family with an impressive life span, the record being 29 years. In 2005, the number of valid described species was approximately 12,000. Ants and ant byproducts are widely used in traditional folk medicine across the globe. In southern India, mud from the interior portion of an anthill is applied topically for the treatment of scabies by the Panjiani tribe. In northern India, scabies, wounds and boils are treated through the topical application of a paste made from the crushed black ants, Bothroponera rufipes. Additionally, ground B. rufipes mixed with water is gurgled to relieve toothaches and daily consumption of 1–2 whole ants reportedly reduces blood
pressure. Weaver ants (Oecophylla smaragdina) have been used in the treatment of severe cough, cold and flu in Myanmar, Africa, Australia and India18,44,56,57. In Thai culture, they are used for detoxification of blood, arresting hemorrhage during miscarriages, restoration of uterus and removal of any aftermath from the uterine canal after childbirth, stimulating pulse and heartbeat, and dizziness. The leaf cutting ant, genus Atta, is used for the treatment of sore throat, stomachache, chest palpitations and heart disease in Latin America18. Dinoponera are Neotropical ants used in the treatment of asthma in Latin America and earaches, rheumatism, and back pain in northeastern Brazil14,53. In 2012, investigation into Dinoponera venom revealed antinociceptive activity supporting its use in pain treatment33. Furthermore, it showed neuroprotective effects when administered as an intraperitoneal treatment as measured by a pentyleneetrazole induced seizure model in mice34. This is an important proof of concept step in the search for new treatments for epilepsy.

Red ants of the ChangBai mountain (Tetramorium spp.), utilized in traditional Chinese medicine, were found to contain three novel coumarin compounds and two known amide alkaloids (3–7, Fig. 2)35. Each compound was evaluated for antibacterial activity against Gram-positive B. subtilis and Gram-negative Escherichia coli. Compound 6, known alkaloid N-(2-hydroxyethyl)-benzamide, showed strong activity against B. subtilis comparable to the positive control, penicillin G. The novel coumarins, compounds 3–5, were mildly active and compound 7, known alkaloid N-phenyl-3,4,5-trihydroxybenzamide, was inactive. None of the compounds exhibited activity against Gram-negative bacteria E. coli. Solenopisin A (8), are well-known antiangiogenic active alkaloids isolated from Solenopsis invicta and Solenopsis germinate fire ants. The key mechanism of action reportedly involves selective inhibition of protein kinase B (AKT), a molecular target of growing interest in cancer18,44,50,57. In 2012, three new alkaloids (9–11) were isolated from the alate queen ants of S. richteri, S.invicta and hybrid (S. richteri × S. invicta) and two years later an additional 36 alkaloids58,59. However, the biological activities of these alkaloids have yet to be evaluated.

The iron ant, Tetraponera rufonigra, is used in local tribes of northeastern India to treat foot and mouth disease in cattle11. One to two ants are crushed into a powder and added to fodder which is then fed to the cattle up to 3 times a day. Studies into the venom of Tetraponera ants revealed a unique class of neurotoxic bioactive compounds termed tetraponерines. Tetraponерines are tricyclic alkaloids isolated for the first time in 1987 from venom of the New Guinean pseudomyrmecine ant36. These molecules, designated T1–T8 (12–15, Fig. 3), are split evenly into two groups based on the grouping of cyclic rings, 5-6-5 or 6-6-560. In each group, further division exists based on the stereochemistry and length of the alkyl chain (pentyl or propyl). In 2001, additional investigation into Tetraponera from India, China and Africa revealed the presence and distribution of the tetraponёres are not equivalent across the species. An Indian ant (T. allaborans) possessed T2, T4 and T8, while the Chinese ant (T. binghami) contained T5, T6, T7 and T8. The remaining tested species, T. rufonigra (India), T. penzigi (Africa), T. clypeata (Africa) and T. sp. cf. emery (Africa) showed no evidence of tetraponёres61. The natural application of tetrapонёres as neurotoxic agents led to interest in their cytotoxic profiles. An evaluation of the anti-proliferative activity of T5–T8 against 4 different carcinoma cell lines highlighted the importance of C5 configuration on potency60. Furthermore, T7 exhibited an impressive cytotoxic profile on the breast carcinoma cell line MCF-7. A structure-activity relationship (SAR) study was conducted to analyze the effect of varying alkyl chain lengths and 6-6-6 tricyclic rings on cytotoxicity against human colorectal (HT29) cancer cells62. Results revealed the alkyl chain to have the greatest effect on cytotoxicity with a decrease in IC50 with increasing chain length, whereas the nature of the ring structure and stereochemistry had no effect.

Chinese black ants, Polyhachis dives, are traditionally used to treat numerous ailments, including rheumatoid and osteoarthritis, inflammatory diseases, and central nervous system (CNS) associated disorders63. Recently, several dopamine derivatives (Fig. 4) were isolated from an ethanol extract: (±)-polyrhadopamine A (16), (±)-polyrhadopamine B (17), polyrhadopamines C–E (18–20), trolline (21) and compounds 22–24. The compounds were evaluated for inhibition of Rho-associated protein kinase ROCK isoforms, ROCK1 and ROCK2, known targets for the potential treatment of cardiovascular, neurological, oncological and renal diseases. Compounds (±)-polyrhadopamine A (16), (±)-polyrhadopamine B (17a) and trolline (21) inhibited both ROCK isoforms. Compounds 22–24 inhibited only the ROCK2 isoform. Potential for the treatment of CNS disorders was further evaluated utilizing a neural stem cell (NSC) proliferation assay. Only (−)-polyrhadopamine B (17b) exhibited a potent dose-dependent promotion of NSC proliferation. Additionally, the immunosuppressive and anti-inflammatory effects of the compounds were studied to verify activity against rheumatoid and osteoarthritis.

One dopamine derivative, polyrhadopamine C (18), was found to have anti-proliferative effects against T-lymphocytes isolated from Kunming mice comparable to the control, cyclosporine A. Likewise, this same compound had comparable inhibitory effects on tumor necrosis factor (TNF)-α production in lipopolysaccharides (LPS)-induced RAW264.7 cells as the control indomethacin35. The combined immunosuppressive and anti-inflammatory activity is strong evidence that polyrhadopamine C (18) could be effective against rheumatoid arthritis. Last year, an additional thirteen nitrogen containing compounds were isolated from P. dives ants and evaluated for their anti-inflammatory, immunosuppressive, and renoprotective activities63.
The compounds 5-(3-indolylmethyl)-nicotinsaureamide (25, Fig. 5) and β-carboline-3-carboxamide (26) suppressed the proliferation of T-cells, key cellular targets for rheumatoid arthritis therapy.64,65 Additionally, compounds 25, 27, and 3-hydroxypyridine significantly reduced collagen IV, fibronectin, and interleukin-6 (IL-6) overproduction supporting the use of the ants as a tonic for kidney problems.64 Furthermore, compounds 25, 26, 29, and 30 led to moderate inhibition of TNF-α production, compound 29 inhibited cyclooxygenase-1, and compound 28 significantly inhibited janus kinase (JAK)-3 kinase all of which indicate anti-inflammatory activity.

2.3. Vespidae (paper, potter, and hover wasps; hornets; yellow jackets)

Wasps are another familiar group of the hymenoptera known for their wondrous paper and clay nests and the act of stinging as a defense mechanism. The Vespidae have approximately 5,000 species classified into six subfamilies: Vespinae, Polistinae, Stenogastrinae, Eumeninae, Masarinae and Euparagiinae. In Latin America and Korea, the Apoica, Brachygastra and Synoeca genera of the Polistinae (paper wasps) are valued traditional therapies for asthma and cough.12,17 Additionally, Apoica and Protonectarina are thought to cure mumps, hemorrhage, and menstruation problems.12 Polistes, the most diverse genus of the Polistinae, are used to treat colds and severe cough in northeastern India and parts of Latin America.11,12 Also, the buffer extract of Polistes mandarinus was cytotoxic against cervix cancer cell line, HeLa, at an equivalent level to the positive control drug mitomycin C.66 As eusocial insects wasps are known to produce antimicrobial substances to provide a collective immunity against microorganisms. Malaysian female wasp venom from various genera of the Stenogastrinae (hover wasps) subfamily were evaluated for antimicrobial activity.57 The venom was extracted with methanol and tested against Gram-positive bacteria, B. subtilis, Gram-negative bacteria, E. coli, and yeast, S. cerevisiae. Each extract produced inhibition zones approximately half the size of the positive controls, penicillin G and kanamycin, indicative of mild activity against the bacterial and yeast strains tested. The venom of the Brazilian social wasp, Synoeca cyanae (Polistinae), was found to inhibit bacterial growth of Gram-positive Enterococcus faecalis and Gram-negative E. coli bacteria by 93% at 100 μg.16 The bite or sting of the Vespa orientalis (Vespinae) is used to treat coughs, colds and stomach pains in the Galo and Nyishi tribes of northeastern India.11 Crude venom of the V. orientalis wasp was found to have strong activity against B. subtilis though not as potent as the positive control tetracycline.69 The venom was also shown to have mild activity against Staphylococcus aureus, E. coli and Klebsiella pneumonia. In addition to antibacterial activity venom is known to possess neurotoxic effects and in many cases is pharmacologically active against tumor cells.56,70 For example polybioside (31, Fig. 6), an isolate of Polybia paulista (Polistinae) venom, was found to be neuroactive.70 Its neuroactive effect was evidenced through the expression of c-Fos protein, a known biochemical marker for stimulated neurons, in various areas of a rat brain after intracerebroventricular (ICV) application.

2.4. Apidae and Halictidae (bees)

Bees are the most popular insect of the hymenoptera albeit for the honeybees (Apis) and bumblebees (Bombus) which ironically represent a tiny fraction of the 20,000 species. Likewise, the Apis,
numbering only 170 species, has been used extensively in medicine. In 2013, an Asiatic honeybee (Apis cerana) chymotrypsin inhibitor (AcC1) gene was identified to encode an 85 amino acid protein that includes a 20 amino acid signal sequence. The recombinant 65-amino acid mature peptide inhibited chymotrypsin with an IC₅₀ of 24.71 nmol/L and showed anti-elastolytic activity via the inhibition of human elastase (IC₅₀ = 38.50 nmol/L) and porcine elastase (IC₅₀ = 70.21 nmol/L). This unique mode of action provides a novel approach for anti-inflammatory therapies. The venom of A. cerana was evaluated for its anti-diabetic and anti-dandruff activities. Researchers assessed its antidiabetic properties by measuring blood sugar, cholesterol and triglyceride content in test subjects before and 30 min after bee stinging. Prior to being stung the blood sugar levels ranged from 91–120 mg/dL, cholesterol was 189–235 mg/dL and triglycerides were 110–184 mg/dL. After bee stinging values fell to 80–94, 178–199 and 84–138 mg/dL for blood sugar, cholesterol and triglyceride levels, respectively. Anti-diarrhoeal activity was investigated by testing the inhibitory effect of the venom on Malassezia furfur a known dandruff causing organism. Ketoconazole, a common active ingredient in anti-dandruff shampoos, was used as a positive control. Interestingly, at a concentration of 5 mg/mL the venom was more than twice as effective as ketoconazole. Last year, Indian Apis dorsata bee venom (ADBV) was found to possess anti-inflammatory, anti-nociceptive, and anti-arthritic properties. These properties were evinced through the venom’s ability to reduce joint inflammation, paw edema and arthritis index along with the reduction of pain perception parameters such as dorsal flexion, motility, and stair climb score. Furthermore, it significantly decreased the major pro-inflammatory mediators like TNF-α and IL-6 in rats induced with arthritis. Crude bee venom extracts from A. cerana, A. dorsata and Apis florea were tested against six Gram-negative bacteria and fungal strains to investigate antimicrobial properties with ampicillin as a control. The venom was as effective as ampicillin against E. coli, Salmonella typhimurium and Xanthomonas subtilis and were more active than ampicillin against Klebsiella pneumonia.

Portuguese propolis from the bee hive of Apis mellifera was recently assessed for total phenolic and flavonoid content, anti-inflammatory and antimicrobial activity. The propolis samples were collected from three regions in Portugal including: Bragança, Coimbra and Beja. The Bragança propolis extract had the strongest antioxidant profile with a total phenolic and flavonoid content of 277.17 and 142.32 mg/g, respectively. Anti-inflammatory activity was assessed by inhibition of hyaluronidase, an enzyme that degrades hyaluronic acid leading to bone loss, pain and inflammation. All extracts inhibited hyaluronidase activity by 50%–75% at a concentration of 25 mg/mL and not surprisingly, the Bragança extract was the most effective. Antimicrobial activity was evaluated against S. aureus (Gram-positive), P. aeruginosa (Gram-negative), E. coli (Gram-negative) and C. albicans (yeast). All extracts showed activity against each strain with the weakest activity being against C. albicans and the strongest activity against S. aureus. Propolis generated by the Brazilian stingless bee, Mellipona orbignyi, was recently evaluated for its chemical composition and antimicrobial, antioxidant and cytotoxic activities. The 80% ethanol extract was determined to contain aromatic acids, phenolic compounds, alcohols, terpenes and sugars in its chemical profile. Antioxidant activity was confirmed by its ability to scavenge free radicals, inhibit hemolysis and lipid peroxidation in human erythrocytes incubated with an oxidizing agent. The extract was active against S. aureus bacterium (Gram-positive) and the fungus C. albicans but not the E. coli (Gram-negative) bacterium. The cytotoxic activity of M. orbignyi ethanol extract was evaluated against the human K562 erythroleukemia cell line which showed a decrease in viability in contact with the extract which was attributed to cell necrosis. A phytochemical screening of the ethanolic extract of Brazilian red propolis indicated the presence of phlobaphene tannins, catechins, chalcones, auranones, flavonones, flavonols, xanthones, pentacyclic triterpenoids and guttiferones. Given the strong phenolic content, it is not surprising the extract exhibited strong antioxidant activity that was in fact far superior to the control Trolox, as measured by 2,2′-diphenyl-1-picrylhydrazyl (DPPH) sequestration. Additionally, the extract exhibited intense cytotoxicity against tumor cell lines SF-295 (human glioblastoma), OVCAR-8 (ovary) and HCT-116 (colon). Geopropolis from the Brazilian stingless bee, Scaptotrigona postica, was evaluated for chemical composition and effectiveness against antiherpes simplex virus-1 (HSV-1). The chemical composition was found to consist mostly of pyrroloidizine alkaloids and C-glycosyl flavones. Treatment with the extract reduced the presence of HSV-1 by 98% which was attributed to inhibition of viral replication and virus entry into cells. Propolis from the Brazilian stingless bee, Tetragonisca fiebrigii, was evaluated for antimicrobial, antioxidant and anti-inflammatory activity. The ethanol extract was analyzed for chemical composition utilizing gas-chromatography mass spectrometry (GC-MS). The major components of the extract, determined by GC-MS, were phenolic compounds, aromatic acids, alcohols, terpenes, and sugars. Antimicrobial activity was evaluated against various Gram-positive and Gram-negative bacteria and fungi. The extract was active against all strains tested but was most effective against Gram-positive bacteria. Antioxidant activity was also noted through scavenging free radicals, inhibiting hemolysis and lipid peroxidation in human erythrocytes incubated with an oxidizing agent. Furthermore, anti-inflammatory potential of the extract was confirmed via inhibition of the hyaluronidase enzyme and cytotoxic activity was concentration-dependent against K562 (leukemia) cells, with a predominance of death by necrosis.

The Indian propolis from the stingless bee, Trigona sp., was evaluated for its chemical composition and antimicrobial activity. The chemical composition of the ethanol extract contained 24 compounds belonging to the following classes: alkanes, thiopholic acids, aromatic acids, aliphatic acids, sugars, esters, and terpenes. The antimicrobial activity was tested against 6 standard bacterial strains (E. coli, S. typhimurium, S. aureus, etc.) several multi drug resistant strains (K. pneumoniae and A. baumanlui, etc.) and a standard yeast strain (C. glabrata). The extract proved potent against all strains tested with minimum inhibitory concentration (MIC) values ranging from 1.21–9.75 μg/mL. Although in general, MICs for Gram-positive bacteria were lower than that for Gram-negative bacteria. Methanol extracts of Nigerian insect propolis were recently evaluated for activity against the malarial parasite Plasmodium berghei. Mice were infected with P. berghei and evaluated for parasitaemia levels over several days. Survivornship of mice treated with the positive control, chloroquine, was not significantly higher than those treated with the extract. Thus, the extract presents a promising lead for new anti-malarial drugs.

The honey from A. mellifera is used in traditional Mexican medicine for the treatment of cough, bronchitis, asthma, rheumatism and diarrhea. A daily spoonful of honey is believed to cure coughs, fevers and stomach pains in the Galo and Nyishi tribes of northeastern India. Additionally, the honeycomb is rubbed on the skin to treat irritation. In northeastern India honeybee (A. cerana, A. florea, and A. mellifera) honey is thought to cure coughs, fevers, stomach pains and the comb, skin irritations. In parts of India honey from the honeybee, Apis indica, is also used to treat colds, headache, asthma and sore throat, indigestion and urinary
disorders. Honey is also an important staple in Argentinian homeopathy. It is taken orally to treat dyspepsia, constipation, colds, influenza and asthma. It is also applied topically to relieve ocular illness, insect bites, bruises, arthritis and muscular. Likewise, in Mexican traditional medicine honey is used as a topical anti-septic agent to treat cutaneous injuries and infections and as an anti-inflammatory agent against muscle fatigue and sprains.

Honey made from the Brazilian stingless bee, Melipona marginata, was evaluated for potential anti-inflammatory activity of honey extract on skin inflammation. High performance liquid chromatography-electrospray ionization-tandem mass spectrometry (HPLC-UV–ESI-MS) identified 11 phenolic compounds such as kaempferol and caffeic acid. Topical treatment using M. marginata honey extract confirmed its anti-inflammatory activity by reducing edema (54%), leucocyte migration, and production of ROS (55%) in the skin of mice after 12-O-tetradecanoylphorbol-13-acetate (TPA) application. Honey made from the Brazilian bee Melipona subnitida was evaluated for its chemical profile and antioxidant activity. The chemical profile consisted of gallic acid, vanillic acid, 3, 4-dihydroxybenzoic acid, cinnamic acid, isorhamnetin, naringenin, quercetin, trans-trans and cis-trans abscisic acid. Three different methods were employed to verify antioxidant activity of pure honey and various extracts including reduction of cooxidation in β-carotene/linoleic acid mixtures and the scavenging of DPPH and 2, 2′-azino-bis(3-ethylbenzothiazoline-6-sulphonic acid) (ABTS). The pure honey and all extracts prevented the bleaching of β-carotene in the carotene/linoleic acid mixtures and scavenged free radicals (DPPH and ABTS) with varying degrees of efficiency. Six honeys from the Republic of Mauritius were analyzed for antimicrobial and antioxidant potential. Antimicrobial properties were evaluated against five strains and antioxidant capacity was determined using eight assays: trolox equivalent antioxidant capacity (TEAC), ferric reducing antioxidant power (FRAP), iron (II) chelating activity, hydroxyl radical (OH) scavenging, DPPH free radical scavenging, hypochlorous acid (HOCl), nitric oxide (NO) and ABTS diammonium salt radical (ABTS –) scavenging. All honeys had similar antioxidant profiles and were found to have higher antibacterial activity than anti-fungal, supporting its reported effects in wound healing. Four Malaysian honey samples, acacia (A. mellifera), pineapple (A. mellifera), borneo (A. cerana) and tualang (A. dorsata) were evaluated for chemical composition and antioxidant activities. Total phenolic, flavonoid and protein content were measured. Tualang had the highest phenolic, flavonoid and protein content and strongest antioxidant activity as measured by the FRAP assay and DPPH radical scavenging activity. The seasonal effect on antimicrobial activity of honey from the Brazilian stingless bee, Melipona compressipes manaoensis, was recently evaluated. Honey harvested in November (rainy season) was only effective against E. coli and S. sonnei. However, honey harvested in March (dry season) was effective against S. aureus, E. coli, P. vulgaris, S. sonnei, and Klebsiella sp. Wild honey produced by four species of bees, A. cerana, A. andreaniformis, A. koschevnikovi and A. mluensis, in Malaysia were evaluated for antioxidant and acetylcholinesterase inhibitor potential. Antioxidant properties were assessed by radical scavenging activity as determined from DPPH assay, ferric reducing/antioxidant power (FRAP) assay, and ABTS decolorization assay. The 80% methanol extract of the wild A. cerana honey performed the best as measured by DPPH assay (84%), FRAP assay (37 μmol/L Fe (II)/1 g dry sample) and ABTS decolorization assay (8 mg AEC/1 g dry sample).

3. Coleoptera (beetles)

The coleopterans are the largest insect order with 350,000-386,000 species in 176 families occupying the land, freshwater and sea. Approximately 62% of beetles belong to the following six mega families, each of which have at least 20,000 members: Staphylinidae, Carabidae, Cerambycidae Curculionidae, Chrysomelidae, and Scarabaeidae. Although diverse and abundant, with the exception of the Scarabaeidae, mention of medicinal use for these mega families in the literature within the last 5 years is scarce. For example, the Chrysomelidae have approximately 38,000 species and the Curculionidae have nearly 60,000 species, yet the only mention of medicinal use was found in Latin American folk medicine. Adult Chrysomelidae, commonly known as leaf beetles, are taken for epilepsy and their caterpillars are further utilized for earache, stroke, swelling, wounds, seborrheic dermatitis, inflammation and thrombosis treatment. Snout beetles (Curculionidae) are leveraged for the relief of fever, headache and boils.

The Tenebrionidae, though not a mega family, have an extensive collection of 19,000 species. One such species is Blaps japanensis, a Chinese medicinal insect, which has been used for the treatment of fever, cough, rheumatism, cancer, and inflammatory disorders. As such, four new dimeric compounds, blapsols A–D, were identified from Blaps japanensis and isolated from Blaps japanensis were evaluated for their activity against cyclooxygenase (COX) enzymes: COX-1 and COX-2. Cyclooxygenases are responsible for catalyzing the conversion of arachidonic acid to prostaglandins, important mediators of pain, fever, and inflammation. The COX-1 isomer is expressed constitutively in tissues and is responsible for maintaining a homeostatic concentration of prostaglandins. The second, COX-2, is induced by stimuli including mitogens, hormones, cytokines, and growth factors. Selective inhibition of COX-2 is therefore desirable as disruption of COX-1 activity leads to adverse side effects like ulcers of the upper gastrointestinal tract.

All compounds inhibited COX-2 and only compound inhibited COX-1, thus supporting the use of these insects to treat pathogenic inflammation. The genotoxic and cytotoxic effects of darkling beetles, Ulomoides dermestoides (Tenebrionidae), used in
traditional Argentinian medicine to treat asthma, Parkinson’s disease, diabetes, arthritis, HIV and cancer, were recently evaluated[92]. The dichloromethane extract was cytotoxic against human lung carcinoma epithelial cells (A549) and normal mononuclear cells. Additionally, the extract induced significant DNA damage in both tumor and healthy cells. However, if targeted directly to tumor cells, there is still potential for anticancer treatment despite the deleterious side effects[92].

The Scarabaeidae make up about 8% of all coleopterans and have been mentioned frequently in the literature for their medicinal value. The large chafier beetle Holotrichia parallela (Scarabaeidae), traditionally used as drug in China and East Asia for the treatment of gout, tetanus, erysipelas and superficial infections, was recently evaluated for antioxidant activity[93]. Both water and ethanolic extracts were subjected to various in vitro assays including linoleic acid peroxidation inhibition, metal-chelating activity and DPPH radical scavenging activity. The ethanol extract had potent metal-chelating activity and inhibition of lipid peroxidation which was attributed to the presence of Catechin. The water extract, rich in proteins, proved to be an excellent oxidation which was attributed to the presence of Catechin. The ethanol extract including linoleic acid peroxidation inhibition, metal-chelating compounds including the

The extracts of rhinoceros beetle larvae, Allomyrina dichotoma (Scarabaeidae), were found to play a vital role in reactive oxygen species (ROS) scavenging in response to oxidative stress[96]. Antioxidant properties of various extracts, including methanol, water, chloroform, ethyl acetate and hexane, were measured through DPPH radical scavenging, superoxide anion radical scavenging and singlet oxygen quenching assays. The methanol extract had the most effective DPPH radical scavenging activity and was 1.7 times more efficient than the standard, ascorbic acid, at singlet oxygen quenching. Also, the extracts protected the biological systems in E.coli and lactate dehydrogenase against detrimental effects of \( ^1 \)O\(_2\) of type II photosensitization in vitro. Last year, A. dichotoma larvae were shown to have hepatoprotective and anticancer activities[97]. Oral administration of powdered larvae lessened diethylthiosulfamine (DEN)-induced hepatotoxicity in mice and an ethyl acetate extract showed cytotoxicity against various cancer cells through induction of apoptosis, necrosis, and autophagy.

In 2012, the antioxidant activity of Proteaia brevitas (Scarabaeidae) at various growth stages, larvae, pupae and imago, was evaluated[98]. The larvae and imago extracts had equivalent minimal morphological deformations (MMDC) values from ethanol and petroleum ether extracts of H. diomphalia larvae showed antifungal activities in vitro against P. oryzae by morphologically deforming hyphal growth from conidia[99]. Minimum morphological deformation concentration (MMDC) values from ethanol and petroleum ether extracts were 7.8 and 125 µg/mL, respectively.

The extracts of rhinoceros beetle larvae, Allomyrina dichotoma (Scarabaeidae), were found to play a vital role in reactive oxygen species (ROS) scavenging in response to oxidative stress[96]. Antioxidant properties of various extracts, including methanol, water, chloroform, ethyl acetate and hexane, were measured through DPPH radical scavenging, superoxide anion radical scavenging and singlet oxygen quenching assays. The methanol extract had the most effective DPPH radical scavenging activity and was 1.7 times more efficient than the standard, ascorbic acid, at singlet oxygen quenching. Also, the extracts protected the biological systems in E.coli and lactate dehydrogenase against detrimental effects of \( ^1 \)O\(_2\) of type II photosensitization in vitro. Last year, A. dichotoma larvae were shown to have hepatoprotective and anticancer activities[97]. Oral administration of powdered larvae lessened diethylthiosulfamine (DEN)-induced hepatotoxicity in mice and an ethyl acetate extract showed cytotoxicity against various cancer cells through induction of apoptosis, necrosis, and autophagy.

In 2012, the antioxidant activity of Proteaia brevitas (Scarabaeidae) at various growth stages, larvae, pupae and imago, was evaluated[98]. The larvae and imago extracts had equivalent singlet oxygen quenching ability to the control, ascorbic acid, as well as comparable DPPH and ABTS radical scavenging activity. Two years later the larvae was shown to also have liver protective and antineoplastic activity[99]. Oral administration of the powdered larvae reduced signs of acute and chronic liver injuries in DEN-induced hepatotoxic mice. Also, a hexane extract proved highly effective against a variety of cancer cell lines with minimal effect on healthy cells. The mechanism of action was determined to be apoptosis via the perturbation of mitochondrial membrane potential[100].
Scientific explorations into these beetles have revealed their bioactivity is attributable to terpene-related compounds. The monoterpenes, cantharidin (CTD, 49, Fig. 9), was the first to be isolated and characterized (c.1914) from blister beetles colloquially known as the ‘Spanish fly’39,46,88. The demethylated CTD analogue, (R)-(−)-palasonin (50), was discovered in the dried bodies of the South African Hycleus oculatus, H. tinctus, and H. lunata beetles and is the only CTD analogue found in plant species as well39,103,104. Imide analogues of CTD and palasonin, cantharimide (51) and palasonimide (52) respectively, in which the oxygen atom in the anhydride core of the molecule was replaced by a nitrogen atom were also found in the H. lunata beetle104. Within the last decade cantharimide, 5-hydroxy cantharimide (53) and norcantharidin (NCTD, 54) were isolated from the native Korean and Chinese blister beetle, Mylabris phalerate PALLAS39.

The earliest isolated compound, CTD, is the highest concentrated terpene in blister beetles and has been explored extensively to further understand its pharmacological and toxicological properties against cancer. Cantharidin is often reported as inducing apoptosis and cell cycle arrest through a variety of underlying pathways39,105. The primary mechanism is the inhibition of protein phosphatase 1 (PP1) and protein phosphatase 2A (PP2A)106. These two phosphatases are known to be involved in various cellular processes including DNA damage, cell cycle arrest, and apoptosis. Researchers from North-Eastern Hill University reported CTD treatment caused plasma membrane disintegration, the appearance of membrane vacuoles and blebbing on murine Ehrlich ascites carcinoma cells which are hallmarks of apoptosis107. Further evaluation revealed CTD acted via an intrinsic pathway involving disruption of the mitochondrial membrane, rerelease of cytochrome c and downstream activation of caspases directly involved in apoptosis108. Similar mechanisms for apoptosis were reported for in vitro tests on human skin and gastric cancer cells in addition to cell cycle arrest at the G2/M phase39,110. Cantharidin was recently reported to effect human lung cancer cells via the intrinsic pathway by increasing reactive oxygen species (ROS) and calcium concentration leading to disruption of the mitochondrial membrane or inhibition of the phosphatidylinositol 3-kinase/Akt signaling pathway111,112. This leads to the selective attenuation of one of the gelatinases, matrix metalloproteinase (MMP)-2, which can degrade components of extracellular matrix (ECM) and inhibits migration and invasion of human lung cancer cells111. Also an extrinsic pathway whereby the death receptor is activated by an extracellular response followed by caspase 8 and caspase 3 activation leading to apoptosis has been postulated111. Recently, it was discovered CTD also suppresses human umbilical vascular endothelial cell proliferation, migration, and tube formation, abrogates vascular endothelial growth factor (VEGF)-induced activation of signal transducer and activator of transcription 3 (STAT3) and suppressed the phosphorylation of JAK1, AKT and Extracellular signal–regulated kinase (ERK)113. All of which are markers of angiogenesis which is required for the large scale growth of tumor cells114. Suppression of angiogenesis further supports the promise of cantharidin as a potential anticancer drug.

Cantharidin has also been found to induce a special type of cell suicide in erythrocytes-termed eryptosis105. The erythrocytes of healthy subjects exposed to cantharidin were found to exhibit cell membrane scrambling and cell shrinkage, both hallmarks of apoptosis. Eryptosis eliminates infected or defective erythrocytes thus counteracting parasitemia in malaria and contributes to the pathophysiology of several clinical disorders including metabolic syndrome and diabetes, malignancy, cardiac and renal insufficiency, hemolytic uremic syndrome, sepsis, mycoplasma infection, malaria, iron deficiency, sickle cell anemia, thalassemia, glucose 6-phosphate dehydrogenase deficiency, and Wilson’s disease115. Cantharidin was recently evaluated for its potential immunomodulatory effects through regulation of dendritic cells (DC). Cantharidin modulates the differentiation and maturation of DCs with alteration of the expression of surface markers and cytokines secretion, causing deviation of standard DC differentiation toward a state of less maturation116. Thus, CTD could treat or prevent disorders involving unwanted immune responses, such as allergies, transplant rejection, or autoimmune diseases.

Though CTD has a variety of therapeutic uses, it has a number of toxic side effects, such as kidney and gastrointestinal damage and disruption of reproductive functions107,117,118. Its closely related naturally occurring analogue, norcantharidin (NCTD), has similar therapeutic uses but with lower toxicity risk17,118. As a chemo-preventive agent NCTD has inhibited tumor progression in non-small cell lung cancer (NSCLC), prostate cancer and leukemia, in vitro, through cell-cycle arrest and apoptosis119–121. Just last year NCTD was also linked to the suppression of the canonical wingless type (Wnt) signaling pathway which is known to cause a variety of human cancers including NSCLC119. Additional therapeutic benefits, cited in a recent review of NTCD, include the modulation of cancer stem cell self-renewal pathways, overcoming multidrug resistance and as radiation sensitizer118.

The presence of nitrogen in cantharimide and 5-substituted cantharimide improves their solubility and cytotoxicity against human hepatocellular carcinoma cell lines39. Given the therapeutic potential of CTD, NCTD and cantharimide, thousands of related analogues have been designed and synthesized with the aim of improving their anticancer potency and decreasing their toxicity. In 2014, a novel CTD analogue, N-benzylcantharidinamide (55, Fig. 10), was found to suppress the metastatic potential of Hep3B hepatocellular carcinoma cells through inhibition of MMP-9 expression122. Cancer cells produce proteolytic enzymes like MMPs to invade surrounding tissue such as extracellular connective tissues and the basement membrane. An additional twenty-three N-substituted NCTD analogues were
synthesized and evaluated for cytotoxicity against five human cancer cell lines that same year\textsuperscript{123}. Lipophilicity of the derivatives was altered through the N-substituents which included alkyl, alkoxy, tertenyl or terpenoxy groups. The lipophilicity of a molecule is an important factor in drug development as it affects solubility and permeability through membranes\textsuperscript{117}. Eleven of the molecules showed significant inhibition against all five cancer cell lines tested with a general trend of increased cytotoxicity with increasing side chain length\textsuperscript{123}. Thus, the lipophilicity of the N-position of norcantharimide is an important parameter affecting the biological activity of the compound. In fact, two compounds N-farnesilxylo-norcarntharimide (56) and N-farnesil-norcarntharimide (57) showed a 2–5-fold increase in cytotoxicity as compared to NCTD against liver carcinoma cells. Even more impressive neither had significant adverse effects on normal murine embryonic liver cells. Not surprisingly, microscopy of the treated cells showed typical morphological features of apoptosis, such as cell shrinkage, chromatin condensation and DNA fragmentation\textsuperscript{125}. The biological activity of norcantharimide analogues with terminal phosphate esters was evaluated against a panel of nine human cancer cell lines: human colon carcinoma cell lines (HT29 and SW480); human breast carcinoma cell line (MCF-7); human ovarian carcinoma cell line (A2780); human lung carcinoma cell line (H460); human skin carcinoma cell line (A431); human prostate carcinoma cell line (DU145); human neuroblastoma cell line (BE2-C); and human glioblastoma cell line (SJ-G2)\textsuperscript{125}. Multiple analogues were at least equipotent with NCTD and a number displayed about a 5-fold increase in cell death with the most potent being bis-trichloroethyl (58) and diphenyl (59) analogues.

In fact, the anti-proliferative activity of these analogues generally correlated with the relative ease of phosphate ester hydrolysis. A series of synthetic NCTD analogues were evaluated against a panel of nine human cancer cell lines including colon carcinoma (HT29 and SW480), breast carcinoma (MCF-7), ovarian carcinoma (A2780), lung carcinoma (H460), skin carcinoma (A431), prostate carcinoma (DU145), neuroblastoma (BE2-C) and glioblastoma (SJ-G2)\textsuperscript{125}. Multiple analogues were at least equipotent with NCTD and a number displayed about a 5-fold increase in cell death with the most potent being bis-trichloroethyl (58) and diphenyl (59) analogues.

4. Blattodea (cockroaches and termites)

Blattodea is a smaller insect order of approximately 7600 species comprised of cockroaches (4600) and termites (3000)\textsuperscript{89}. The giant cockroach, Rhyparobia maderae (Blaberidae), is used as an asthma treatment in Latin American folk medicine\textsuperscript{12}. Additionally, the cockroach Euryctis manni (Blattidae) is prescribed for headache relief. The American cockroach, Periplaneta americana (Blattidae), is used for a variety of conditions including: heartburn, asthma, stomach ache, intestinal colic, earache, alcoholism, epilepsy, vomit, boil, hemorrhage, bronchitis, diarrhea, gonorrhea, panaris, cancer, stroke, burns and menstrual cramps\textsuperscript{12}. In parts of India, it is cooked in a soup to treat postpartum, uterine problems and urinary obstructions\textsuperscript{10}. Additionally, the whole body is consumed to treat gangrene, dyspnea, asthma and tuberculosis and applied topically for tetanus and earache relief\textsuperscript{35,10}. In China, the ethanol extract of the dried whole body of P. americana have been used as traditional Chinese medicine for the treatment of blood stasis syndrome, acne and abdominal mass for hundreds of years. Given the varied therapeutic uses for the pest, the ethanol extract of the insect was screened for bioactive compounds resulting in the discovery of four new isocoumarins periplatinis A–D (70–73, Fig. 12) along with three known ones (74–76)\textsuperscript{28}. The compounds were tested against various cancer cell lines and compounds 72–74 showed significant cytotoxic activities against human liver (HepG2) and breast cancer (MCF-7) cells. In a small scale clinical study, sixty septic patients were treated with P. americana extract which successfully improved the condition and prognosis in patients with sepsis\textsuperscript{129}. Blatta orientalis (Blattidae) is used topically for the treatment of tetanus and ear pain in Mexican traditional medicine\textsuperscript{13}. Recently,
the anti-asthmatic and anti-anaphylactic activities of *B. orientalis* mother tincture was evaluated. The results indicated nonsel-ective anti-cholinergic and anti-histaminic activities were responsible for its anti-asthmatic activity. Furthermore, mast cell stabilization, lowering elevating IgE antibody and reducing eosinophil led to both passive and active anaphylactic activity.

The Chinese cockroach, *Eupolyphaga sinensis* (Corydidae), is a key ingredient in a number of traditional Chinese medicinal products including *Shu-Mai Tang*, *Dahuang Zhechong* pill, *Shen-song Yangxin* capsule, *Shen-Yuan-Dan*, *Yi Guan Jin* and *Yao-mother* products including *Shu-Mai Tang*, *Dahuang Zhechong* pill, *Shen-Yuan-Dan* is used for the treatment of cirrhosis. Shen-Yuan-Dan is used for the treatment of ischemic heart disease and was found to have a therapeutic effect against inflammation-induced myocardial fibrosis leading to the attenuation of left ventricular remodeling and improved cardiac function. The Dahuang Zhechong pill is clinically used for the treatment of hepatic diseases, gynecopathy, and atherosclerosis in China. Further studies into its role in athrosclerosis treatment revealed its ability to significantly inhibit proliferation of vascular smooth muscle cells through depressing PDGF expression in the cells, retarding the cell cycle and promoting cellular apoptosis. Shenong Yangxin capsule is prescribed for the treatment of arrhythmias in China and was recently found to have potential use in diabetic cardiopathy. The Chinese herb modified Yi Guan Jin is an effective regimen that is usually used in outpatients with chronic liver diseases such as fibrosis and cirrhosis. Shen-Yuan-Dan is used for the treatment of ischemic heart disease, proven to relieve angina and reduces cardiac dysfunction. Traditional Chinese medicine Yaotongning Capsule is a common remedy to treat rheumatism in China and possesses diverse biological activities including anti-inflammation.

Another traditional use of *Eupolyphaga sinensis* is the treatment of cancer. Researchers extracted the adult insect with petroleum ether, ethyl acetate and *n*-butanol and evaluated the *in vitro* activity of the extracts in ten cancer cell lines. All extracts had significant antiproliferative activity against all cell lines. An LC-MS evaluation of the ethyl acetate extract was analyzed further to isolate the potential bioactive compounds. Four compounds including two new isocoumarins were isolated but 

---

**Figure 13** Structures of isocoumarins and an alkaloid isolated from *Eupolyphaga sinensis*.

---

reduce tumor progression in human and murine chronic myeloid cells. The mechanism was believed to involve G2/M cell cycle arrest and inhibition of epidermal growth factor receptor (EGFR) phosphorylation in the EGFR signaling pathway of human and murine chronic myeloid tumor cells. This extract was also found to inhibit NSCLC progression through the inhibition of angiogenesis. The proposed underlying mechanism was interruption of the auto-phosphorylation of kinase insert domain receptor (KDR) and subsequently AKT and ERK1/2.

*Eupolyphaga plancyi* is a close relative of *Eupolyphaga sinensis* that is widely distributed in China, has also been used in Chinese traditional medicine to treat diseases such as amenorrhea and fracture and its extracts were shown to have anticancer and anti-inflammatory activities. A recent study of *P. plancyi* Bolivar isolated five new compounds (80–84, Fig. 14). Compounds 81 and 83 inhibit JAK3 kinase with IC50 values of 12.6 and 5.0 μmol/L, respectively. In addition, compound 83 inhibit DDR1 kinase with an IC50 value of 4.87 μmol/L. Both JAK3 and DDR1 are potential drug targets for cancer.

The winged termite, *Reticulitermes* (Rhinotermitidae), is roasted with sugar and consumed to treat eye diseases and diabetes in certain regions within India. In the Kerala state in southern India, the Kurichchan tribe consume a decoction of termites and *Vitex negundo* (a five-leaved Chinese chastetree shrub) twice daily to cure ulcers. The tribes of Irular and Mudugar dry and powder *Odontotermes formosanus* (Termitidae) which is then added as needed to milk for general health improvement and relief of body pain. These termites are also fried in oil, powdered and administered orally for rheumatic diseases. The termites of *Microcerotermes exigus* (Termiteidae) and *Nasutitermes macrocephalus* (Termiteidae) are used for asthma, bronchitis and flu in Latin American folk medicine. In Zambia, *Hodotermes mossambicus* (Hodotermitidae) is used to alleviate childhood malnutrition. In Somalia, *Macrotermes bellicosus* (Termiteidae) is used to suture wounds. In Nigeria, *Macrotermes nigeriensis* is leveraged for wounds and used to alleviate sickness in pregnant women. In Brazil, various genera of the family Termitidae are used for asthma, cough, flu, sore throat, sinusitis, tonsilisits and hoarseness. In India, *Odontotermes formosanus* of the Termitidae family is thought to cure ulcers, improve health, relieve body pain, rheumatism and anemia, and enhance lactation in women.

5. **Diptera (flies and mosquitoes)**

Dipterans, commonly called true flies, are a large and diverse species of 120,000–150,000 colonized on all continents and aquatic environments. Though often associated with forensic
Hemiptera (true bugs, aphids, cicadas and scale insects)

Hemiptera is the fifth largest insect order with 90,000 species described to date. Hemipterans are often cited in the literature as a healthy alternative food source and only more recently been explored for bioactive substances. The cochinellid insect, Dactylopius coccus (Dactylopiidae), whole body of which is used for nasal congestion and ear pain in holistic Mexican medicine. A major isolate of D. coccus, carminic acid, was determined to be a powerful free radical scavenging agent as ascorbic acid and stronger than trolox. Furthermore, it can protect against enzymatically induced β-carotene bleaching as well as quercetin.

The shell of the lac insect Kerria lacca (Kerriidae) is utilized for the treatment of diarrhea and indigestion in parts of India. Shellac, a natural polymer resin secreted from K. lacca, is valued as a treatment for measles, macula and scabies in traditional Chinese medicine. Recently, several novel sesquiterpene acids were isolated from this shellac including four α-cedrene types (Shellolic acid A–C, Laccishellolic acid, 85–88, Fig. 15), two R-curcumene types (Shellolic acid D–E, 89–90) and Shellolic acid F (91). The anti-tumor and antibacterial activities were evaluated revealing that none had cytotoxic activity and only shellolic acid A had antimicrobial activity against B. subtilis.

The stinkbug Encosternum delegorguei (Tessaratomidae), an edible treat in Zimbabwe and South Africa, was recently proven to contain significant amounts of protein, fat and minerals validating its use as a nutritional food source. In addition to being a healthy appetizer, the stinkbug has acclaimed traditional medicinal values, such as decreasing hypertension and curing asthma and heart diseases. Likewise, the Chinese stinkbug, Aspongopus chinensis (Pentatomidae), is a nutrient-rich edible insect which has been employed to relieve pain, and treat nephropathy and kidney disease in China. In recent years, much research has gone into identifying bioactive small molecules from this insect. Researchers evaluated petroleum ether, ethyl acetate and n-butanol A. chinensis extracts for cytotoxicity against a panel of human cancer cell lines including colorectal (HCT-116), lung (H-125), prostate (LNCaP), melanoma (MDA), pancreatic (PANC-1), breast (MCF-7), ovarian (OVC-5), leukemia (CEM), glioma (U251N) and murine normal cells (CFU-GM) as a control. Each exhibited significant activity against the cell lines and the ethyl acetate extract was analyzed further for active compounds. The compounds isolated from the ethyl acetate extract, however, all exhibited no or poor activity against the cancer cells. In 2014, a novel trimer of N-acetyldopamine (NADA) termed (±)-aspongamide A (92, Fig. 16) was isolated from A. chinensis along with a racemic mixture of NADA dimeric enantiomers. The compounds were tested as potential therapeutic agents to treat chronic kidney disease (CKD). Aspongamide A (92) promotes a significant decrease in...
expression and phosphorylation of Smad3 a key role in the TGF-B/Smad signaling pathway which regulates CKD pathogenesis. The dimeric NADA derivatives (7) were observed to cause an increase in collagen I and α-smooth muscle actin (α-SMA) expression, indicating benefit for restoring skin after wounds as collagen I and α-SMA are implicated in the process of fibrosis and wound healing.

That same year an additional 29 small molecules were isolated from the insect eight of which were novel bioactive compounds including four norepinephrine derivatives (aspongopusamides A–D, 93–96), three sesquiterpenoids (aspongnoids A–C, 97–99, Fig. 17) and one lactam (asponglaclam A, 100). A subset of the insect-derived substances was evaluated as renal protection agents in high-glucose-induced mesangial cells and COX-2 inhibition. Aspongopusamide A (103) caused a significant decrease in collagen IV, fibronectin, and IL-6 production in high-glucose-induced mesangial cells and inhibited COX-2 (IC₅₀ = 6.50 μmol/L). Aspongopusamide B also provided protective effect on diabetic nephropathy and aspongopusamide C (95) inhibited COX-2 with an IC₅₀ of 117 μmol/L. Last year, an additional 9 novel small molecules (101–109) were isolated from A. chinensis. With the exception of aspongadenine A (103) and aspongester A (109), all had the ability to promote the proliferation of neural stem cells. The most active compound was (+)-asponguanine B (100a).

7. Orthoptera (grasshoppers and crickets)

There are some 22,500 species of orthopterans described to date 46. Orthopterans are sometimes singled out as pests though that determination is based on select members. One notable family includes the acridids, plague locusts that at times enter gregarious stages forming swarms numbering in the millions. In Mexico, people eat the grasshopper, Schistocerca piceifrons (Acrididae), to relieve hiccups and the cricket, Gryllus spp., for stuttering 18. The mole cricket, Gryllotalpa Africana (Gryllotalpidae), is used in traditional Korean medicine for retention of urine, urolithiasis (stones in bladder), edema, lymphangitis and furuncles 17. The house cricket, Acheta domesticus (Gryllidae), is used for the treatment of scabies, asthma, eczema, lithiasis, carache, oliguresis, rheumatism, urine retention, urinary incontinence and ophthalmological problems in Latin America 12. The crickets, Paragryllus temulentus (Gryllidae) and Gryllus assimilis (Gryllidae), were additionally cited for rheumatism and warts, respectively. The desert locus Schistocerca gregaria (Acrididae) was found to be rich in sterols with therapeutic uses including cardiovascular protective and immune regulatory effects, anti-inflammatory and anticancer benefits 159. The known anticancer agents pancratistatin (110, Fig. 18), narciclasine (111) and an additional derivative, ungeremine (112), were isolated from the Texas grasshopper, Paragryllus temulentus (Gryllidae) and Gryllus assimilis (Gryllidae), were additionally cited for rheumatism and warts, respectively. The desert locus Schistocerca gregaria (Acrididae) was found to be rich in sterols with therapeutic uses including cardiovascular protective and immune regulatory effects, anti-inflammatory and anticancer benefits. The known anticancer agents pancratistatin (110, Fig. 18), narciclasine (111) and an additional derivative, ungeremine (112), were isolated from the Texas grasshopper,
| Family | Genus | Common name | Molecule | Biological activity | Ref. |
|---|---|---|---|---|---|
| **Hymenoptera (bees, wasps, ants and sawflies)** | **Pergidae** | Sawfly | Macrocarpal (1); grandinol (2) | Antimicrobial | 50 |
| | **Tetramorium** | Red ant | Compounds 3–7 | Antimicrobial | 55 |
| | **Solenopsis** | Fire ant | Solenopin A (8) | Antiangiogenic | 18, 44, 56, 57 |
| | **Solenopsis** | Fire ant | Compounds 9–11 | – | 58, 59 |
| | **Tetraponera** | Iron ant | Compounds 12–15 | Antiproliferative | 60, 62 |
| | **Polyrhachis** | Black ant | Polyhadopamines A–E (16–20); Troline (21); compounds 22–30 | Antiinflammatory; antiproliferative; renoprotective | 63–65 |
| **Vespidae** | **Polybia** | Wasp | Polybioside (31) | Neuroactive | 70 |
| **Coleoptera (beetles)** | **Tenebrionidae** | Blaps | Blapsols A–D (32–35); compounds 24, 36–39 | Antinflammatory | 90 |
| | **Scarabaeidae** | Dung beetle | Molossusamides A–C (40–42); compounds 38 and 43 | Antinflammatory | 100 |
| | **Bruchidae** | Seed beetle | Compounds 44–48 | Antioxidant | 101 |
| | **Meloidae** | Various | Blister beetle | Cantharidin (49) | – | 103, 104 |
| | | | (R)(+)-palasonin (50) | – | 104 |
| | **Meloidae** | **Hydeus** | Blister beetle | Cantharimide (51); palasonimide (52) | – | 104 |
| | **Meloidae** | **Mylabris** | Blister beetle | S-Hydroxy cantharidinamide (55); norcantharidin (NCTD, 54) | Antiproliferative | 122 |
| | **Synthetic** | – | – | N-farnesyl-norcantharimide (56); N-farnesyl-norcantharindimide (57) | Antiproliferative | 123 |
| | **Synthetic** | – | – | Compounds 58–59 | Antiproliferative | 125 |
| | **Synthetic** | – | – | Compounds 60–69 | Antiproliferative | 126, 127 |
| **Blattoidea (cockroaches and termites)** | **Blattidae** | **Periplaneta** | American cockroach | Periplatins A–D (70–73); compounds 74–76 | Antiproliferative | 128, 129 |
| | **Eupolyphaga** | **Chinese cockroach** | Cockroach | Compounds 77–78; eupolyphagin (79) | Antiproliferative | 137 |
| | **Polyphaga** | **Chinese cockroach** | Plancyamides A (80) and B (82); plancypyrazine A (81); plancyols A (83) and B (84) | Antiproliferative | 142 |
| **Hemiptera (true bugs, aphids, cicadas and scale insects)** | **Kerriidae** | **Kerria** | Lac insect | Shelloic acid A (85) | Antimicrobial | 152 |
| | **Kerriidae** | **Kerria** | Lac insect | Shelloic acids B–C (86–87); Laccishelloic acid (88); shelloic acids D–F (89–91) | – | 152 |
| | **Pentatomidae** | **Aspongus** | Chinese stinkbug | Aspongamine A (92) | Active against chronic kidney disease | 156 |
| | | | Chinese stinkbug | Aspongopamides A–D (93–96) | Antinflammatory | 157 |
| | | | Chinese stinkbug | Aspongopamides A–C (97–99); aspongactam A (100) | – | 157 |
| | | | Chinese stinkbug | Compounds 101–109 | Promote neutral stem cell proliferation | 158 |
| **Orthoptera (grasshoppers and crickets)** | **Romaleidae** | **Brachystola** | Texas grasshopper | Pancratistatin (110); narciclasine (111); ungeremine (112) | Antiproliferative | 160 |
| **Lepidoptera (butterflies and moths)** | **Papilionidae** | **Byasa** | Taiwan butterfly | Papilistatin (113) | Antiproliferative | 161 |
| | **Bombycidæ** | **Bombyx** | Silk moth | Compound 114 | Analgesic effects | 162 |

–Not applicable.
**Brachystola magna** (Romaleidae)\(^{10}\). The compounds were further tested against murine lymphocytic leukemia cells and a variety of human tumor cell lines and as expected all were active against the cell lines with narciclasine being the most potent on each cell line. The chitosan from two species of grasshoppers, *Calliptamus barbarus* (Acrididae) and *Oedaleus decorus* (Acrididae), were found to have potent antimicrobial activity against several strains of fish, clinical and food-borne pathogens\(^{105}\). Additionally, the chitosan was proven to have antioxidant activity as measured by DPPH radical scavenging activity and ferric reducing power. The nutritional value of the short-horned grasshopper, *Chondacris rosea* (Acrididae), and mole cricket, *Brachytrupes orientalis* (Gryllidae), were recently investigated\(^{104}\). These food staples of Arunachal Pradesh, India were found to possess a substantial amount of protein, minerals and energy.

8. **Lepidoptera (butterflies and moths)**

Lepidopterans, which include butterflies and moths, are one of the largest insect orders with anywhere from 155,208 to 174,312 members to date\(^{165}\). Moths especially have been leveraged at all life stages for medicinal purposes. In traditional Indian medicine, dog bites are treated by consummation and/or direct application of freshly laid eggs of *Diaecrisia obliqua* (Arctiidae) to the affected area\(^{83}\). Dried full grown larvae of *Stomphosistis thraustica* (Gracillariidae) are taken with herbs to treat common fever and to increase milk flow in lactating women. In Latin American folk medicine, the moths, *Oiketicus kirbii* (Psychidae), are given for the treatment of asthma, earache and haemorrhage\(^{172}\).

A novel small molecule, designated papilistatin (113, Fig. 19), was isolated from the wings of *Byasa polyeuctes termessa*, a Taiwan butterfly (Papilionidae)\(^{161}\). Papilistatin showed strong cytotoxicity against colon and pancreatic cancer cells and selective Gram-positive antibacterial activity. The cocoon of mulberry white caterpillar, *Rondotia menciana* (Bombycidae), was recently evaluated for the presence of flavonoids\(^{166}\). Researchers discovered flavonol galactosides quercetin 3-O-galactosyl-galactoside and kaempferol 3-O-galactosyl-galactoside, the first reported flavonoids conjugated with galactose in animal species\(^{169}\). The medicinal activity of these novel compounds was not evaluated however.

The cocoon crude extract of *Bombyx mori*, a close relative of the *R. menciana*, was found to have significant effect on hypercholesterolemia and atherosclerosis, as evinced through lipid-lowering capability and decreasing the extent of atherosclerotic lesions in rabbits, induced with hyperlipidemia and atherosclerosis\(^{167}\). Three years later, a detailed chemical profile of an aqueous extract of its cocoon revealed various flavonoids including quercetin 7-O-β-D-glucoside, kaempferol 7-O-β-D-glucopyranoside, coumaric acid glucoside, 2,5-dihydroxy-4-nonenedicarboxylic acid and 9,12-dihydroxy steaic acid\(^{168}\). Not surprisingly, this flavonoid rich extract exhibited free radical scavenging activity and cardioprotection via reduction of apoptotic factors, oxidative stress, cardiac enzyme activity and interleukin-6 (pro-inflammatory marker).

Even *B. mori* excreta is known to contain bioactive compounds useful in the treatment of diabetes, cancer, high cholesterol and high blood pressure, and is also commonly prescribed in traditional Chinese medicine to treat infectious diseases, headaches and abdominal pain\(^{162,169}\). For example, the excreta possess ecysterones (114), which are known to have analgesic effects in man\(^{162}\). In 2014, a novel hydroxyl fatty acid termed bombyxmiconic acid methyl ester was also discovered in *B. mori* droppings; however, it’s medicinal properties were not evaluated\(^{169}\). *B. mori* are not the only moths valued for the medicinal qualities of their feces\(^{170}\). Another Chinese tradition is the consumption of insect tea which is prepared from the feces of moths, such as *Aglossa dimidiata*, *Hydrillodes morosa* and *Nodaria niphona* and mentioned in the “Compendium of Materia Medica” as a potential treatment for otitis media. Additional efforts in the past two years have been made to validate new therapeutic properties of insect tea. In a study on human tongue carcinoma cells, insect tea significantly induced apoptosis in cancer cells by upregulating BAX, CASP3, CASP9 and down regulating BCL2\(^{171}\). This was likewise the case for human liver cancer cells evaluated last year\(^{172}\). Anti-inflammatory activity was also evidenced through the down regulation of genes encoding nuclear factor kappa-light-chain-enhancer of activated B cells (NFκB), inducible nitric oxide synthase (iNOS), and COX-2\(^{173}\). Furthermore, insect tea was found to be as effective at preventing gastric ulcers as the gastric ulcer drug, ranitidine and becull mucosa cancer preventative effects in *in vivo* murine models\(^{173,174}\).

9. **Conclusions**

Summary of small molecules isolated from insects and their biological activities is listed in Table 1. It is clear that Mother Nature has more to offer the world of medicine than just plants. Insects have a long and rich history in traditional medicine across the globe. Not to mention over one hundred small molecule isolates with bioactivity discovered within the past several years. Despite these discoveries, insect-derived products have yet to establish the recognition and market success of their plant derived counterparts. Undoubtedly, this is partly due to negative cultural attitudes towards insects, particularly in the West. However, public perception of insects should not deter scientific pioneers from advancing these largely untapped resources. Indeed, if given the proper attention, insect-derived substances hold great promise for the future of natural product drug discovery.

**References**

1. Guo ZR. The modification of natural products for medical use. *Acta Pharm Sin B* 2017;7:119–36.
2. Dias DA, Urban S, Roessner U. A historical overview of natural products in drug discovery. *Metabolites* 2012;2:303–36,
3. Miner J, Hoffhines A. The discovery of aspirin's antithrombotic effects. *Tex Heart Inst J* 2007;34:179–86.
4. Cragg GM, Newman DJ. Natural products: a continuing source of novel drug leads. *Biochim Biophys Acta* 2013;1830:3670–95.
5. Shen B. A new golden age of natural products drug discovery. *Cell 2015;163:1297–300.
6. Brown DG, Lister T, May-Dracka TL. New natural products as new leads for antibacterial drug discovery. *Bioorg Med Chem Lett* 2014;24:413–8.
7. Newman DJ, Cragg GM. Natural products as sources of new drugs over the 30 years from 1981 to 2010. *J Nat Prod* 2012;75:311–35.
8. Benitez G. Animals used for medicinal and magico-religious purposes in western Granada Province, Andalusia (Spain). *J Ethnopharmacol* 2011;137:1113–23.
9. Mahawar MM, Jaroli DP. Traditional zootherapeutic studies in India: a review. *J Ethnobiol Ethnomed* 2008;4:17.
10. Jaroli DP, Mahawar MM, Vyas N. An ethnozoological study in the adjoining areas of Mount Abu wildlife sanctuary, India. *J Ethnobiol Ethnomed* 2010;6.
11. Chakravorty J, Ghosh S, Meyer-Rochow VB. Practices of entomophagy and entomotherapy by members of the Nyishi and Galo tribes, two ethnic groups of the state of Arunachal Pradesh (North-East India). J Ethnobiol Ethnomed 2011;7:5.

12. Alves RRN, Alves HN. The faunal drugstore: animal-based remedies used in traditional medicines in Latin America. J Ethnobiol Ethnomed 2011;7:9.

13. Vallejo JR, González JA. The use of the head louse as a remedy for jaundice in Spanish folk medicine: an overview. J Ethnobiol Ethnomed 2013;9:52.

14. Młcek J, Borkovcova M, Bednarova M. Biologically active substances of edible insects and their use in agriculture, veterinary and human medicine—a review. J Cent Eur Agric 2014;15:225–37.

15. Chakravorty J, Ghosh S, Meyer-Rochow VB. Comparative survey of entomophagy and entomotherapeutic practices in six tribes of eastern arunachal pradesh (India). J Ethnobiol Ethnomed 2013;9:50.

16. Wilsanand V, Varghese P, Rajitha P. Therapeutics of Insects and insect products in south indian traditional medicine. Indian J Tradit Know 2007;6:563–8.

17. Pemberton PW. Insects and other arthropods used as drugs in Korean traditional medicine. J Ethnopharmacol 1999;65:207–16.

18. Alonso-Castro AJ. Use of medicinal fauna in Mexican traditional medicine. J Ethnopharmacol 2014;152:53–70.

19. Kim H, Song MJ. Ethnozoological study of medicinal animals on Jeju Island, Korea. J Ethnopharmacol 2013;146:75–82.

20. Kim H, Song MJ. Ethnomedicinal practices for treating liver disorders of local communities in the Southern Regions of Korea. Evid Based Complement Alternat Med 2013;2013:869176.

21. Kujawska M, Zamudio F, Hilgert NI. Honey-based mixtures used in traditional medicine. J Ethnopharmacol 2011;133:49–55.

22. Patricia V, Oliverio V, Triny L, Favián M. Meliponini biodiversity study of antimicrobial peptides: origins, functions, relative mechanisms and applications. Peptides 2012;37:207–15.

23. Ntwasa M, Goto A, Kurata S. Coleopteran antimicrobial peptides: prospects for clinical applications. Int J Microbiol 2012;2012:101989.

24. Maddocks SE, Jenkins RE. Honey: a sweet solution to the growing problem of antimicrobial resistance?. Future Microbiol 2013;8:1419–29.

25. McLoone P, Warnock M, Fyfe L. Honey: a realistic antimicrobial for reversed-phase high-performance liquid chromatography. J Ethnopharmacol 2014;149:41–5.

26. Medalla MD, Kapari L, Salminen JP. New types of flavonol oligosaccharides accumulate in the hemolymph of birch-feeding sawfly larvae. J Chem Ecol 2010;36:864–72.

27. Crockett SL, Bovee JL. Flavonoid glycosides and naphthodianthrones in the sawfly Tenthredo zonala and its host-plants, Hypericum perforatum and H. hirsutum. J Chem Ecol 2011;37:943–52.

28. Rastogi N. Provisioning services from ants: food and pharmaceuticals. Asian Myrmecol 2011;1:103–20.

29. Sousa PL, Quinet Y, Ponte EL, De Vale JJF, Torres AB, Havt A, et al. The effects of the Brazilian Ant Dinoponera quadriceps. J Ethnobiol Ethnomed 2011;7:315–21.

30. Daleprenje JB, Abdalla DS. Emerging roles of propolis: antioxidant, cardioprotective, and antiangiogenic actions. Evid Based Complement Alternat Med 2013;2013:75135.

31. Freires IA, De Alencar SM, Rosalen PL. A pharmacological perspective on the use of Brazilian Red Propolis and its isolated compounds against human diseases. Eur J Med Chem 2016;110:267–79.

32. Sforcin JM, Bankova V. Propolis: is there a potential for the development of new drugs?. J Ethnopharmacol 2011;133:253–60.

33. Bellik Y. Bee venom: its potential use in alternative medicine. Anti-Infect Agents 2015;5:3–16.

34. Ntwasa M, Safaieejad Z, Parivar K, Divsalar A, Nazari Z. Antineoplastic effects of honey bee venom. Zahedan J Res Med Sci 2013;15:1–5.

35. Orsolic N. Bee venom in cancer therapy. Cancer Metast Rev 2012;31:173–94.

36. Lee WK, Park KK, Pak SC. The protective effect of bee venom on fibrosis causing inflammatory diseases. Toxins 2015;7:4758–72.

37. Yi HY, Cho DW, Huang YD, Yu XQ. Insect antimicrobial peptides and their applications. Appl Microbiol Biotechnol 2014;98:5807–22.

38. Li Y, Xiang Q, Zhang Q, Huang Y, Su Z. Overview on the recent study of antimicrobial peptides: origins, functions, relative mechanisms and application. Peptides 2012;37:207–15.
61. Garrafão HM, Spande TF, Jain P, Kaneko T, Jones TH, Blum MS, et al. Ammonia chemical ionization tandem mass spectrometry in structural determination of alkaloids. II. Tetraponeres from pseudo-myrmecine ants. Rapid Commun Mass Spectrom 2001;15:1409–15.

62. Rouchaud A, Braekman JC. Synthesis of new analogues of the tetraponeres. Eur J Org Chem 2009;2009:2666–74.

63. Tang JJ, Zhang L, Jiang LP, Di L, Yan YM, Tu ZC, et al. Dopamine derivatives from the insect Polybarychis dives as inhibitors of ROCK1/2 and stimulators of neural stem cell proliferation. Tetrahedron 2014;70:8852–7.

64. Tang JJ, Fang P, Xia HL, Tu ZC, Hou BY, Yan YM, et al. Constituents from the edible black ants (Polybarychis dives) showing protective effect on rat mesangial cells and anti-inflammatory activity. Food Res Int 2015;67:163–8.

65. Cope AP, Schulze-Koops H, Aringer M. The central role of T cells in the social wasp (Hymenoptera: Apidae: Meliponini) in the anti-malarial activity of stingless bee (Hymenoptera: Apidae: Meliponini) in the municipality of Nocupetê, Micoahuan, Mexico. J Ethnobiol Ethnomed 2014;10:47.

66. Ahn MY, Ryu KS, Lee YW, Kim YS. Cytotoxicity and L-amino acid oxidase activity of crude insect drugs. Arch Pharm Res 2000;23:477–81.

67. Baracchi D, Mazza G, Turillazzi S. From individual to collective immunity: the role of the venom as antimicrobial agent in the stenostraginae wasp species. J Insect Physiol 2012;58:188–93.

68. Mortari MR, Do Couto LL, Dos Anjos LC, Mourão CB, Camargos TS, Camargos JA, et al. Pharmacological characterization of Synoeca cyanea venom: an aggressive social wasp widely distributed in the neotropical region. Toxicon 2012;59:163–70.

69. Jalaie F, Fazeli M, Rajaian H, Shekarforoush S. In vitro antibacterial effect of wasp (Vespa orientalis) venom. J Venom Anim Toxins Incl Trop Dis 2014;20:22.

70. Saidemberg DM, Da Silva-Filho LC, Tognoli LM, Tommena CF, Palma MS. Polybiocide, a neuroactive compound from the venom of the social wasp Polybia paulistia. J Nat Prod 2010;73:527–31.

71. Prakash S, Bhargava HR. Apis cerana bee venom: it’s anti-diabetic and anti-dandruff activity against Malassezia furfur. World Appl Sci J 2014;32:343–8.

72. Surendra NS, Jayaram GN, Reddy MS. Antimicrobial activity of crude venom extracts in honeybees (Apis cerana, Apis dorsata, Apis florea) tested against selected pathogens. Afr J Microbiol Res 2011;5:2765–72.

73. Silva JC, Rodrigues S, Feás X, Estevinho LM. Antimicrobial activity, phenolic profile and role in the inflammaton of propolis. Food Chem Toxicol 2012;50:1790–5.

74. Campos JF, Dos Santos UP, Macorini LF, De Melo AM, Balesieter JB, Paredes-Gamoer EJ, et al. Antimicrobial, antioxidant and cytotoxic activities of propolis from Melipona orbignyi (Hymenoptera, Apidae). Food Chem Toxicol 2014;65:374–80.

75. De Mendonça IC, de Moraes Porto IC, do Nascimento TG, de Souza NS, dos Santos Oliveira JMD, dos Santos Arruda RE, et al. Brazilian red propolis: phytochemical screening, antioxidant activity and efect against cancer cells. BMC Complement Altern Med 2015;15:357.

76. Coelho GR, Mendonça RZ, De Senna Vilar K, Figueiredo CA, Moutsou CB, Camargos TS, Camargos JA, et al. Pharmacological characterization of Synoeca cyanea venom: an aggressive social wasp widely distributed in the neotropical region. Toxicon 2012;59:163–70.

77. Campos JF, Dos Santos UP, Dos Santos Da Rocha P, Damião MI, Balesieter JB, Cardoso CA, et al. Antimicrobial, antioxidant, anti-inflammatory, and cytotoxic activities of propolis from the stingless bee Tetragonisca hirtigra (Jataí). Evid Based Complement Altern Med 2015;2015:926186.

78. Choudhuri MK, Punekar SA, Ranade RV, Paknikar KM. Antimicrobial activity of stingless bee (Trigona sp.) propolis used in the folk medicine of Western Maharashtra, India. J Ethnopharmacol 2012;141:363–7.

79. Olayemi KL. Therapeutic potentials of nigerian insect-propolis against the malarial parasite, Plasmodium berghei (Haemoporida: Plasmodiidae). Am J Drug Disc Devel 2014;4:241–7.

80. Vijayakumar S, Prabhu S, Yabesh JEM, Pragashraj R. A quantitative ethnozoological study of traditionally used animals in Pachamalai hills of Tamil Nadu, India. J Ethnopharmacol 2015;171:51–63.

81. Reyes-González A, Camou-Guerrero A, Reyes-Salas O, Argüeta A, Casas A. Diversity, local knowledge and use of stingless bees (Apidae: Meliponini) in the locality of Agua de Dios, Oaxaca, Mexico. J Ethnobiol Ethnomed 2014;10:47.

82. Borsato DM, Pradente AS, Boscadini PM, Borsato AV, Luz CF, Maia BH, et al. Topical anti-inflammatory activity of a monofloral honey of Mimosa scabrella provided by Melipona marginata during winter in Southern Brazil. J Med Food 2014;17:817–25.

83. Silva TM, dos Santos FP, Evangelista-Rodrigues A, da Silva EM, da Silva GS, de Novais JS, et al. Phenolic compounds, melissapolygalous, phistochemical analysis and antioxidant activity of jandaíra (Melipona subnitida) honey. J Food Comp Anal 2013;29:10–8.

84. Dor GO, Mahomoodally MF. Chemical profile and in vitro bioactivity of tropical honey from mauritius. Asian Pac J Trop Dis 2014;4:S1002–13.

85. Moniruzzaman M, Khalil MI, Sulaiman SA, Gan SH. Physicochemical and antioxidant properties of malaysian honeys produced by Apis cerana, Apis dorsata and Apis mellifera. BMC Complement Altern Med 2013;13:43.

86. Pimentel RB, da Costa CA, Albaqueque PM, Junior SD. Anti-microbial activity and rutin identification of honey produced by the stingless bee melipona compressa manaosensis and commercial honey. BMC Complement Altern Med 2013;13:151.

87. Philip Y,Mohd Fadzelly AB. Antioxidative and acetylcholinesterase inhibitor potential of selected honey of Sabah, Malaysian Borneo. Int J Food Res 2015;22:1953–60.

88. Bouchard P, Grebennikov VK, Smith ABT, Douglas H. Biodiversity of Coleoptera. In: Foottit RG, Adler PH, editors. Insect biodiversity: science and society. West Sussex: Blackwell Publishing Ltd. 2009. p. 265–301.

89. Hoffman KH. Insect molecular biology and ecology. Boca Raton: CRC Press; 2014.

90. Yan YM, Li LJ, Qin XC, Lu Q, Tu ZC, Cheng YX. Compounds from the insect Blaps nipponensis with COX-1 and COX-2 inhibitory activities. Bioorg Med Chem Lett 2015;25:2469–72.

91. Narsinghani T, Sharma R. Lead optimization on conventional non-steroidal anti-inflammatory drugs: an approach to reduce gastro-intestinal toxicity. Chem Biol Drug Des 2014;84:1–23.

92. Crespo R, Villaverde ML, Giorri JT, Güerci A, Juárez MP, De Bravo MG. Cytotoxic and genotoxic effects of defence secretion of Uonomides dermestoides on A549 cells. J Ethnopharmacol 2011;136:204–9.

93. Liu SF, Sun J, Yu LN, Zhang CS, Bi J, Zhu F, et al. Antioxidant activity and phenolic compounds of Holotrichia parallela motschusky extracts. Food Chem 2012;134:1885–91.

94. Dong QF, Wang Z, Liu HJ, Zhang CF, He DX, Wu G, et al. Flavonoid and other compounds from Holotrichia diaphomaia larvae. Chem Nat Compd 2011;47:114–5.

95. Dong QF, Wang J, Zhang SF, Wang Z, Zhang CX, Gao H, et al. Antifungal activity of crude extracts and fat-soluble constituents of Holotrichia diaphomaia larvae. Bioresource Technol 2006;99:8521–3.

96. Suh HW, Kim SR, Lee KS, Park S, Kang SC. Antioxidant activity of various solvent extracts from Allomyrina dichotoma (Arthropoda: Insecta) Larvae. J Photochem Photobiol B Biol 2010;99:67–73.

97. Lee JE, Jo DE, Lee AJ, Park HK, Youn K, Yun EY, et al. Hepatoprotective and anticancer activities of allomyrina dichotoma larvae. J Life Sci 2015;25:307–16.

98. Suh HJ, Kang SC. Antioxidant activity of aqueous methanol extracts of Protaetia brevitarsis Lewis (Coleoptera: Scarabaeidae) at different growth stages. Nat Prod Res 2012;26:510–7.

99. Lee JE, Jo DE, Lee AJ, Park HK, Youn K, Yun EY, et al. Hepatoprotective and antiinflammatory properties of Protaetia brevitar- sis larvae. Entomol Res 2014;44:244–53.

100. Lu J, Sun Q, Tu ZC, Lv Q, Shui PX, Cheng YX. Identification of N-acetyldopamine dimers from the dung beetle Catharsius molossus
and Their COX-1 and COX-2 inhibitory activities. Molecules 2015;20:15589–96.

101. Hirose Y, Ohta E, Kawai Y, Ohita S. Dorsamin-A’s, glycerolipids carrying a dehydroxyphenylalanine ester moiety from the seed-eating larvae of the bruchid beetle Bruchidius dorsalis. J Nat Prod 2013;76:554–8.

102. Sah HJ, Kim SR, Hwang JS, Kim MJ, Kim I. Antioxidant activity of aqueous methanol extracts from the lucanid Beetle, Serrographus platymerus castanicolor motschulsky (Coleoptera: Lucanidae). J Asia Pac Entomol 2011;14:95–8.

103. Mebs D, Pogoda W, Schneider M, Kauert G. Cantharidin and demethylcantharidin (palasonin) content of blister beetles (Coleoptera: Meloidae) from southern Africa. Toxicon 2009;53:466–8.

104. Dettner K, Schramm S, Seidl V, Klemm K, Gäde F, Fietz O, et al. Occurrence of terpene anhydride Palasonin and Palasoninimide in blister beetle Hycleus lunata (Coleoptera: Meloidae). Biochem Syst Ecol 2003;31:203–5.

105. Alzoubi K, Egler J, Briglia M, Fazio A, Faggio C, Lang F. Induction of interleukin-2 production. Evid Based Complement Alternat Med 2011;2011:777981.

106. Zhao J, Guan XW, Chen SW, Hui L. Synthesis and biological evaluation of norcantharidin analogues as protein kinase-1 inhibitors. Bioorg Med Chem Lett 2015;25:363–6.

107. Twu NF, Srinivasan R, Chou CH, Wu LS, Chiu CH. Cantharidin and norcantharidin inhibit caprine luteal cell steroidogenesis via suppressing VEGF-induced JAK1/STAT3, ERK and AKT signaling pathways. Arch Pharm Res 2014;37:282–9.

108. Verma AK, Prasad SB. Bioactive component, cantharidin from Malydris cichorii and its antitumor activity against Ehrlich ascites carcinoma. Cell Biol Toxicol 2012;28:133–47.

109. Verma AK, Prasad SB. Antitumor effect of blister beetles: anetho-medicinal practice in Karbi community and its experimental evaluation against a murine malignant tumor model. J Ethnopharmacol 2013;148:869–79.

110. Hsiao TC, Tsai CH, Wu PP, Hsu SC, Liu HC, Huang YP, et al. Cantharidin induces G2/M phase arrest by inhibition of CDC2/CDC25C and cyclin A and triggers apoptosis through the upregulated expression of the mitochondria-dependent pathway. J Cell Biochem 2014;135:658–68.

111. Zhang CJ, Chen ZT, Zhou XL, Xu W, Wang G, Tang XX, et al. Cantharidin induces G2/M phase arrest and apoptosis in human gastric cancer SGC-7901 and BGC-823 cells. Oncol Lett 2014;8:2721–6.

112. Kim YM, Ku MJ, Son YJ, Yun JM, Kim SH, Lee SY. Anti-metastatic effect of cantharidin in A549 human lung cancer cells. Arch Pharm Res 2013;36:479–84.

113. Hsia TC, Yu CC, Hsu SC, Tang NY, Lu HF, Huang YP, et al. Cantharidin induces apoptosis of H460 human lung cancer cells through the mitochondria-dependent pathway. Int J Oncol 2014;45:2393–402.

114. Zhang CJ, Chen ZT, Zhou XL, Xu W, Wang G, Tang XX, et al. Cantharidin induces G2/M phase arrest and apoptosis in human gastric cancer SGC-7901 and BGC-823 cells. Oncol Lett 2014;8:2721–6.

115. Lang E, Lang F. Triggers, inhibitors, mechanisms, and significance of cryptosis: the suicidal erythrocyte death. BioMed Res Int 2015;2015:513518.

116. Hsieh CH, Huang YC, Tsai TH, Chen YJ. Cantharidin modulates development of human monocye-derived dendritic cells. Toxicol in Vitro 2011;25:1740–7.

117. Twu NP, Srinivasan R, Chou CH, Wu LS, Chiu CH. Cantharidin and norcantharidin inhibit caprine luteal cell steroidogenesis in vitro. Exp Toxicol Pathol 2012;64:37–44.

118. Hsieh CH, Chao KSC, Liao HF, Chen YJ. Norcantharidin, derivative of cantharidin, for cancer stem cells. Evid Based Complement Alter Med 2013;2013:838651.

119. Xie JR, Zhang YP, Hu XM, Lv R, Xiao DJ, Jiang L, et al. Norcantharidin inhibits Wnt signal pathway via promoter demethylation of WIF-1 in human non-small cell lung cancer. Med Oncol 2015;32:145.

120. Shen B, He PJ, Shao CL. Norcantharidin induced DU145 cell apoptosis through ROS-mediated mitochondrial dysfunction and energy depletion. PLoS One 2013;8:e84610.

121. Liao HF, Chen YJ, Chou CH, Wang FW, Kuo CD. Norcantharin induces cell cycle arrest and inhibits progression of human leukemia Jurkat T cells through nitrogen-activated protein kinase-mediated regulation of interleukin-2 production. Toxicol In Vitro 2011;25:206–12.

122. Lee JY, Chung TW, Choi HJ, Lee CH, Eun JS, Han YT, et al. A novel cantharidin analog N-Benzylcantharimidine reduces the expression of MMP-9 and invasive potentials of Hep3B via inhibiting cytosolic translocation of HuR. Biochem Biophys Res Commun 2014;447:371–7.

123. Wu JY, Kuo CD, Chu CY, Chen MS, Lin JH, Chen YJ, et al. Synthesis of novel lipophilic N-substituted norcantharimide derivatives and evaluation of their anticancer activities. Molecules 2014;19:6911–28.

124. Armott JA, Planey SL. The influence of lipophilicity in drug discovery and design. Exp Op Drug Disc 2012;7:863–75.

125. Robertson MJ, Gordon CP, Gilbert J, McCluskey A, Sakoff JA. Norcantharimide analogues possessing terminal phosphate esters and their anti-cancer activity. Bioorg Med Chem 2011;19:5734–41.

126. Tarleton M, Gilbert J, Sakoff JA, McCluskey A. Synthesis and anticancer activity of a series of norcantharimide analogues. Eur J Med Chem 2012;54:573–81.

127. Shimizu T, Iruka M, Matsukura H, Hashizume D, Sodeoka M. Synthesis of optically pure cantharidin analogue NCA-01, a highly selective protein phosphatase 2B inhibitor, and its derivatives. Chem Asian J 2012;7:1221–30.

128. Luo SL, Huang HJ, Wang Y, Jiang RW, Wang L, Bai LL, et al. Isocoumarins from American cockroach (Periplaneta americana) and their cytoxic activities. Fitoterapia 2014;95:115–20.

129. Zhang HW, Wei LY, Zhang ZY, Liu SZ, Zhao G, Zhang J, et al. Protective effect of Periplaneta americana extract on intestinal mucosal barrier function in patients with sepsis. J Tradit Chin Med 2013;33:70–3.

130. Nimgulkar CC, Patil SD, Kumar BD. Anti-asthmatic and anti-anaphylactic activities of Blatta orientalis mother tincture. Homeopathy 2011;100:138–43.

131. Yin HQ, Wang B, Zhang JD, Lin HQ, Qiao Y, Wang R, et al. Effect of traditional Chinese medicine Shu-Mai-Tang on attenuating TNFα-induced myocardial fibrosis in myocardial ischemia rats. J Ethnopharmacol 2008;118:133–9.

132. Zhang YH, Liu JT, Wen BY, Liu N. Mechanisms of inhibiting proliferation of vascular smooth muscle cells by serum of rats treated with Dahuang Zhechong pill. J Ethnopharmacol 2009;124:125–9.

133. Shen NN, Li XG, Zhou T, Bilal MU, Du N, Hu YY, et al. Shensong Yangxin Capsule prevents diabetic myocardial fibrosis by inhibiting TGF-β1/Smad signaling. J Ethnopharmacol 2014;157:161–70.

134. Lin HJ, Tseng CP, Lin CF, Liao MH, Chen CM, Kao ST, et al. A Chinese herbal decoction, modified Yi Guan Jian, induces apoptosis in hepatic stellate cells through an ROS-mediated mitochondrial/caspase pathway. Evid Based Complement Alternat Med 2011:2011:459531.

135. Liu HX, Shang JJ, Chu FY, Li AY, Wu B, Xie XR, et al. Protective effects of Shen-Yuan-Dan, a traditional Chinese medicine, against myocardial ischemia/reperfusion injury in vivo and in vitro. Evid Based Complement Alternat Med 2013:2013:956397.

136. Ni LJ, Xu XL, Zhang LG, Shui WZ. Quantitative evaluation of the in vitro effect and interactions of active fractions in Yantongning-based formulae on prostaglandin E2 production. J Ethnopharmacol 2014;154:807–17.

137. Jiang HL, Luo XH, Wang XZ, Yang JL, Yao XJ, Crews P, et al. New isocoumarins and alkaloid from Chinese insect medicine, Eupolyphaga sinensis walker. Fitoterapia 2012;83:1275–80.

138. Ge GF, Yu CH, Yu B, Shen ZH, Zhang DL, Wu QF. Antitumor effects and chemical compositions of Eupolyphaga sinensis Walker ethanol extract. J Ethnopharmacol 2012;141:178–82.

139. Zhang YM, Zhan YZ, Zhang DD, Dai BL, Ma WN, Qi JP, et al. Eupolyphaga sinensis Walker displays inhibition on hepatocellular carcinoma through regulating cell growth and metastasis signaling. Sci Rep 2014;4:5518.
