A Review of Plants Used in South African Traditional Medicine for the Management and Treatment of Hypertension

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ABSTRACT

South Africa contains 9% of the world's higher plants, and despite its rich biodiversity, it has one of the highest prevalence of hypertension in Africa. This review provides information on medicinal plants embraced in South Africa for hypertension management, with the aim of reporting pharmacological information on the indigenous use of these plants as antihypertensives. This review not only focuses on the activity of antihypertensive medicinal plants but also reports some of its phytochemical constituents and other ethnopharmacological and therapeutic properties. Information obtained from scientific and or unpublished databases such as Science Direct, PubMed, SciFinder, ISTOR, Google Scholar, Web of Science, and various books revealed 117 documented antihypertensive plant species from 50 families. Interestingly, Asteraceae topped the list with 16 species, followed by Fabaceae with 8 species; however, only 25% of all plant species have demonstrated antihypertensive effects originating from both in vitro and in vivo studies, lending credence to their folkloric use. Only 11 plant species reportedly possess antihypertensive properties in animal models, with very few species subjected to analytical processes to reveal the identity of their bioactive antihypertensive compounds. In this review, we hope to encourage researchers and global research institutions (universities, agricultural research councils, and medical research councils), particularly those showing an interest in natural products, for the need for concerted efforts to undertake more studies aimed at revealing the untapped potential of these plants. These studies are very important for the development of new pharmaceuticals of natural origin useful for the management of hypertension.

ABBREVIATIONS

ACE	angiotensin-converting enzyme
CVD	cardiovascular diseases
CCD	congestive cardiac diseases
CHD	coronary heart diseases
HBP	high blood pressure
HTN	hypertension
KZN	KwaZulu-Natal
SA	South Africa

Introduction

HTN (HBP)–a silent killer that provides no symptoms, signs, or warnings–is defined according to the World Health Organization and Naish et al. [1] as a medical condition that results in a persistent rise in blood pressure (BP) within the arteries of an individual. HBP or HTN, expressed as systolic (maximum pressure) or minimum pressure (diastolic) measurements, may be a result of unclassified lifestyle or genetic factors (such as excessive salt intake, smoking, or alcohol), which account for about 90–95% of the total HBP [2], although there are less common secondary causes–for

example, narrowing of the kidney arteries [2] or the use of birth control pills (represented by unconfirmed hypotheses). The BP of a normotensive individual is in the range of 100–140 mmHg systolic and 60–90 mmHg diastolic pressures (although different values apply to children) [3]. Exceeding the normal or accepted values of these measurements has been linked to increased risk of CVD and many chronic complications that include but are not limited to CHD, stroke, cardiac arrest, CCD, renal insufficiency, and myocardial infarction [4, 5].

Globally, more than 1 billion people suffer from HTN, with an estimated 1.25 billion by 2025 [6]. In addition, HTN and its related diseases are gradually becoming major causes of disability [7]. To date, HBP has been regarded as a disease of developed countries; however, in recent years, the prevalence of HTN is speedily growing in low- and middle-income countries (LMIC) as well as continents like Africa [8], where poverty, exposure to infectious diseases (e.g., HIV, malaria, tuberculosis), and inadequate awareness of disease treatment and control are the order of the day. Today, 1 out of every 5 adults in LMIC suffers from HBP, and it has been projected that by 2025 this ratio will balloon to 3 out of every 4 adults [9]. Similarly, in sub-Saharan Africa, the incidence of HTN has increased tremendously, particularly during the last 20-30 y, with 40% (approximately 20 million) of the entire adult population living with the disease [10]. This translates to about 80 million people (of the overall 650 million populations) in 2000 (within the region), with the possibility of this number increasing to 150 million by 2025 [9]. Intriguingly, HTN constitutes a major health concern in SA, as HBP and its related conditions (CVD, diabetes, and stroke) remain the second highest cause of mortality after sexually transmitted diseases [11], with more than 6.3 million adult sufferers (particularly black South Africans) [12]. Moreover, further epidemiological information in terms of age and prevalence revealed the average incidence of HPB to be 59% among blacks, 55% among Indians, and 50% among whites, with females (11.2%)-most importantly those between the ages of 15 and 34-recording a higher prevalence than males (7%) of the 34-65 age bracket [12, 13].

The treatment or control of HBP aims to lower the maximum (systolic) and minimum (diastolic) pressures below 140/90 mmHg (normal BP). This is achieved through nonpharmacological (lifestyle modification) and pharmacological (drug therapy) approaches. Lifestyle modification rallies around maintaining body weight (< 25 kgm⁻²), regular exercise (30 min daily), restricting excessive salt consumption (limiting intake to < 2400 mg/d), and the regulation of alcohol intake (limiting intake to 2 standard drinks for men and 1 for women) [12]. It is noteworthy that a number of investigations buttressed the relevance of these modifications in HTN management [14, 15]. Pharmacologically, the use of orthodox drugs is encouraged to control HBP via nitric oxide, as well as neural and renal endocrine mechanisms when lifestyle modification fails to achieve the desired results [16]. Antihypertensive agents including diuretics (relaxation of blood vessels/expulsion of excess fluids/salt), beta blockers such as atenolol (inhibition of adrenaline action to control heart rate), calcium channel blockers (arterioles relaxation and heart beat regulation), ACE (an important enzyme within the renin-angiotensin cascade) inhibitors such as lisinopril (limiting the arteries production of angiotensin/altering narrowing of the arteries), and vasodilators such as aspirin (which relaxes the blood vessels for ease of blood flow within the circulatory system) are embraced individually or in combination to manage HBP and related conditions [7, 12]. Experimentally, HBP may be evaluated *in vitro* (ACEI assays) and/or *in vivo* (Dahl-salt stress assays). However, the adoption of these synthetic moieties in HTN management has presented a number of challenges in recent years, including serious financial implications, unavailability to buy, numerous side effects attributed to altered renal function, angioedema (severe swelling below the skin surface), dry cough, weakness, and headaches [17]. As such, there is an urgent call for an appropriate substitute in phytotherapy, which is adjudged to be safe, effective, inexpensive, and pose little or no side effects in HTN control [18].

Medicinal plants (MPs), otherwise known as medicinal herbs, are employed in herbal medicine for their usefulness in promoting health, affording brief relief of symptoms or providing therapeutic uses. They produce a number of compounds (phytoconstituents) that play important roles in their biological functions, such as protecting them from predation (e.g., insects, fungi, herbivores mammals). Herbal medicine or phytotherapy is an age-old tradition practiced globally and involves the use of whole plants or plant parts (leaves, flowers, fruits, stems, roots, or rhizomes) to maintain the general well-being of an individual [19]. Globally, 60% of the developed and 80% of the developing and underdeveloped populations of the world now rely on herbal therapy for their health care needs [20]. Of particular interest is Africa, where an estimated 60-90% of the entire population makes use of phytomedicine in the management of ailments, including HTN [21], based in part on the availability and affordability of these medicines as compared to popular or conventional biomedicines [22].

Besides being endowed with rich cultural biodiversity, SA accounts for 9% of the higher plants worldwide, with well over 30,000 plant species [23]. It is no surprise that over 3000 species of these plants are currently used by more than 200,000 traditional healers in the treatment, cure, or management of many diseases [24]. It is noteworthy that more than 60-80% of the entire population, particularly those of rural settlements, adopt the use of medicinal herbs in various forms as medicines for the maintenance of health. Research efforts examining the pharmacological activities of MPs have been ongoing for decades in SA. While a number of the potential effects of these plants against many diseases such as diabetes, cancer, malaria, and other infectious ailments have been scientifically validated, very few of them have been evaluated for their hypotensive as well as antihypertensive efficacies [12], despite their traditional use and continuous documentation. In line with the aforementioned, there is a need for more research efforts (vis-à-vis the rapid prevalence of HPB) from government, scientific/academic institutions, and research agencies toward affirming their efficacy and determining the safety profile of documented antihypertensive plants in order to stem the growing prevalence of HPB. Hence, this review has been prepared to report the antihypertensive or hypotensive MPs used in SA to encourage the financing of additional studies to explore those with unproven pharmacological or biological properties, as well as to evaluate and isolate their active compounds as potential drug candidates.

Results and Discussion

Antihypertensive MPs in SA

According to Watt and Brewer-Brandwijk [25], Hutchings et al. [26], van Wyk et al. [24], van Wyk and Gericke [23], Thring and Weitz [27], Mofette [28], Olorunnisola et al. [29], Moteetee and van Wyk [30], Semenya et al. [31], de Wet et al. [32], and David et al. [33], more than 100 MPs are traditionally used and ethnobotanically documented in SA for the management of HTN (> Table 1). It is noteworthy that, for the purpose of this review, random selection of these plants was considered. Moreover, some plants without formal documentation of their medicinal properties use are also included in the list based on their reported antihypertensive potential.

Adenopodia spicata (E.Mey.) C. Presl

A. spicata is an endemic plant of southern Africa that belongs to the Leguminoceae (or Fabaceae) family. There are 7 species of the genus Adenopodia (gymnantha, oaxacana, patens, rotundifolia, scelerata, schlechteri, and spicata) distributed in the northern neotropics, particularly Mexico (southern and western) and Central America. The prominent common names include stekel-splinterboontjie (Afrikaans), ibobo, umlungumabele, ubobo (Zulu), spiny splinter-bean (English), and umbambangwe, and they are traditionally used for treating chest or breast pain, syphilis, and HTN [26]. The prominent chemical components are saponins, and the in vitro antihypertensive activity of the plant against ACE (an enzyme responsible for the conversion of angiotensin I to angiotensin II that plays a role in HBP) has been reported, where it showed 97% and 72% inhibition of the enzyme using aqueous and ethanolic leaf extracts, respectively, each at a concentration of $25 \,\mu g/$ mL (> Table 2) [34].

Agapanthus africanus (L.) Hoffmanns.

A. africanus, a member of the Amaryllidaceae (or Agapanthaceae) family, is also called blue lily, Cape agapanthus, fynbos agapanthus (English), kleinbloulelei (Afrikaans), ubani (Zulu), and isicakathi (Khosa). The plant is widely distributed on the Cape of Good Hope (eastern), SA, and grows up to 25–70 cm long. Its folkloric usage includes treatment for chest pains, cough, and heart disease, as well as to induce or ease labor [24,26]. Oxytocic, *in vivo* antifungal, and antihypertensive activity of the aqueous and ethanolic leaf extracts (25 µg/mL) against ACE (► Table 2) have been reported with 63% and 44% inhibition, respectively [34]. Active constituents include spirostane (found to induce apoplastic peroxidase activity) and furostane, as well as several other saponins and sapogenins [24].

Agave americana (L.)

A. americana is a flowering plant of the Asparagaceae family indigenous to Mexico and the United States, particularly Texas, Arizona, and New Mexico, but is now grown in many continents and nations of the world such as Africa (Kenya, Nairobi, Western Cape, SA), Australia, China, India, and Thailand. Its common names include century plant, agave, agave cactus, American agave aloe, or American aloe (despite not being related to the genus *Aloe*), blougaring-

boom, amerikaanse aalwee, kaalgaarboom, amerikaanse aalwyn, blou-aalwee, gareboom, makaalwyn (Afrikaans), lekhala (Sotho), and xikwenga (Tsonga). It is traditionally used in the treatment of HTN by the Bapedi traditional healers [31], as well as having other ornamental uses. Its chemical constituents include but are not limited to isolated cantalsaponin-1 ((25R)-5 α -spirostan-3 β ,6 α ,23 α triol-3,6-di-O-β-D-glucopyranoside) and (25R)-3beta,6alpha-dihydroxy-5alpha-spirostan-12-one 3,6-di-O-beta-D-glucopyranoside [35], while pharmacological evidence of its anti-inflammatory, antileishmanial, antifungal, antitumor (in MTT assays), cytotoxic, antimicrobial, antioxidant, antiulcer, antibacterial, and irritant properties have been reported [35, 36]. Additionally, the antihypertensive effect of an ethanolic leaf extract (25 µg/mL) of the plant has been verified against ACE and found to inhibit the activity of the enzyme by 82%, in addition to a 72% inhibition by aqueous leaf extract (> Table 2) [34].

Alepidea amatymbica Eckl. & Zeyh.

A. amatymbica is included in the Apiaceae family, with local names such as larger tinsel flower (English), kalmoes (Afrikaans), iqwili (Xhosa), iKhathazo (Zulu), and lesoko (Sotho). There are close to 30 species of the genus *Alepidea* existing in southern Africa, with a few species distributed in Kenya and Ethiopia. Interestingly, 3 subspecies of *A. amatymbica* are available. Traditionally, the plant is used for the treatment of colds, cough, malaria, rheumatism, wound healing, asthma (when combined with *Cannabis sativa* L. [Cannabaceae]), influenza, and abdominal cramps [24], while its application as an antimicrobial, antihypertensive, diuretic [37], antioxidant, anti-inflammatory, antitrypanosomal, antibacterial, antimicrobial, antifungal, antihypertensive, and Afolayan reported on the cytotoxicity of the plant and its safety profile in a comprehensive review [38].

Allium sativum (L.)

A. sativum is a member of the Amaryllidaceae family native to Central Asia and Iran (northeastern region). Commonly referred to as garlic, it is the second most widely used plant in the genus Allium next to onion, Allium cepa L. A. sativum contains as its phytochemical constituents sulfur compounds (like allypropyl, aliin, allicin, s-allylcysteiin, ajoene, and vinyldithiins), peptides, terpenoids, phenols, saponins, amino acids including arginine, minerals such as selenium, and enzymes like myrosinase and peroxidase, as reported by Tesfeya and Mengesha [39]. The plant is indigenously used as a spice to flavor foods and to aid digestion, as well in the control of numerous ailments such as heart problems [25], fever, diabetes, breast cancer, stroke, and arthritis and as an immune system enhancer. Of particular interest is the thorough review of its pharmacological properties (antibacterial, antifungal, antiparasitic, antiviral, antihypertensive, antiatherosclerotic, antithrombotic, antioxidant, antimutagenic, antihyperglycemic, antilipidemic, antihelminthic, diuretic, digestive, hepatoprotective, radioprotective, cardioprotective, antiprotozoal, antimicrobial, anti-inflammatory, antitubercular, immunomodulatory, analgesic, and wound healing), as well as its cytotoxic and safety profile from different nations of the world including Spain, India, and Ethiopia [39-41], among others. Additionally, its biological importance as an antidiabetic

Table 1 South African MPs documented for HBP control.

Name	Family	Common name (Zulu/Afrikaans)	Plant type	Part(s) used	Evidence of documented HBP use	Ethnobotanical indications on HBP	References
Achyranthes aspera	Amaranthaceae	Isinama	Herb	Roots	[26]	NS	[26]
Acokanthera oppositifolia	Apocynaceae	inHlungunyembe	Shrub	Leaves, roots stem	[24]	Maceration	[24]
Adenopodia spinata	Fabaceae	Ubobo	Shrub	Leaves, roots	[26]	Maceration	[26, 34]
Agapanthus africanus	Amaryllidaceae	Ubani	Herb	Leaves, roots	[26]	Maceration, infusion	[26, 34]
Agathosma betulina	Rutaceae	Regteboegoe	Shrub	Leaves, stem	[29]	Infusion	[29]
Agave americana	Asparagaceae	Unknown	Shrub	Leaves	[31]	Decoction	[26, 31, 34]
Albertisia delagoensis	Menispermaceae	Umgandaganda	Shrub	Rhizome, leaves, stem, root	[32]	Decoction	[32]
Alepidea amatymbica	Apiaceae	iKhathazo	Shrub	Rhizomes, roots	[24]	NS	[24]
Allium sativum	Amaryllidaceae	Unknown	Herb	Flower bud	[25, 27]	NS	[27]
Aloe ferox	Asphodelaceae	iNhlaba	Shrub	Leaves/sap	[27, 28, 33, 110]	NS	[33]
Aloe marlothii	Asphodelaceae	inhlaba umhlaba	Herb	Leaves, roots	[32]	Decoction	[32]
Aloe striatula	Asphodelaceae	Unknown	Shrub	NS	[28]	NS	[28]
Amaranthus dubius	Amaranthaceae	Unknown	Herb	Leaves	FPR	Maceration	[18]
Amaranthus hybridus	Amaranthaceae	Unknown	Herb	Leaves	FPR	Maceration	[18]
Arachis hypogaea	Fabaceae	Amakinati	Herb	Leaves, seed	[32]	Decoction	[33]
Artemisia afra	Asteraceae	Mhlonyane	Shrub	Leaves	[27, 33]	Infusion	[27,33]
Asystasia gangetica	Acanthaceae	Isihobo	Herb	Leaves	FPR	Maceration	[18]
Ballota africana	Lamiaceae	Kattekruide	Shrub	Leaves	[24, 25, 33]	Infusion	[27,33]
Cadaba aphylla	Brassiceae	Bobbejaanarm	Shrub	Leaves, stem	[33]	Infusion	[33]
Canabis sativa	Cannabaceae	Nsangu	Herb	Leaves	[24,29-30,32]	Infusion, decoction	[29, 32]
Carpobrotus dimidiatus	Mesembryanthemaceae	Ikhambi Iamabulawo	Herb	Leaves, stem, fruit	[32]	Decoction	[32]
Catha edulis	Celastraceae	Umhlwazi	Shrub	Leaves	[26]	Maceration	[26, 34]
Catharanthus roseus	Apocynaceae	Unknown	Herb/ sub-shrub	Roots, flower, seeds	[32]	Decoction	[32]
Chrysocoma ciliata	Asteraceae	Kaalsiektebos	Shrub	Leaves, roots	[33]	Decoction	[33]
Cinnamomum camphora	Lauraceae	Uroselina	Tree	Gum	[24]	NS	[24]
Cissampelos capensis	Menispermaceae	Fynblaarklimop	Shrub	Roots	[24,29]	Infusion	[29]
Citrullus lanatus	Curcurbitaceae	Bitterwaatlemoen	Herb	Fruit, seed, leaves	[32]	Decoction	[32]
Citrus limon	Rutaceae	Ulamula	Shrub/ small tree	Peel of fruit, pulp, root	[32]	Decoction	[32]
Citrus maxima	Rutaceae	Upapamuzi	Tree	Fruit	[32]	N/A (taken raw)	[32]
Cladostemon kirkii	Capparaceae	umThekwini	Shrub/ small tree	Root, stem, bark	[32]	Maceration, decoction	[32]
Clausena anisata	Rutaceae	Umnukambhiba	Shrub	Leaves, roots	[26]	Maceration, decoction	[26, 34, 65]
Commelina africana	Commelinaceae	Idangabane	Herb	Whole plant	[26,30]	Decoction (T. capensis)	[30]
Commelina benghalensis	Commelinaceae	Idangabane	Herb	Whole plant	[26]	Poultice	[26] cor

Table 1 Continued

Name	Family	Common name (Zulu/Afrikaans)	Plant type	Part(s) used	Evidence of documented HBP use	Ethnobotanical indications on HBP	Reference
Convolvulus capensis	Convolvulaceae	Skaapklimop	Shrub	Bulb	[33]	Decoction	[33]
Conyza scabrida	Asteraceae	Umanzimnyama	Shrub	Leaves	[27]	Infusion	[24, 27]
Crinum macowanii	Amaryllidaceae	Umdube	Herb	Bulbs, leaves, whole plants	[24]	NS	[24]
Crassula muscosa	Crassulaceae	Skoenvetebos	Herb	Leaves, stem, roots, flowers	[33]	Decoction	[33]
Dicerothamnius rhinocerotis	Asteraceae	Ranosterbos	Small shrub	Leaves, stem	[33]	NS	[33]
Dicoma anomala	Asteraceae	Umuna	Herb	Leaves, roots	[24, 28]	NS	[24,28]
Dicoma capensis	Asteraceae	Koorsbossie	Herb	Leaves, twigs, roots	[24]	NS	[24]
Dietes iridioides	Iridaceae	isishuphe somfula	Herb	Leaves, roots, rhizomes	[26]	Maceration, infusion	[26, 34, 65
Diosma oppositifolia	Rutaceae	Bitterboegoe	Shrub	Leaves, stem, flowers	[33]	NS	[33]
Dipcadi brevifolium	Hyacinthaceae	Ikhakahkha	Herb	Bulb	[26]	Decoction	[26]
Dombeya rotundifolia	Malvaceae	iNhliziyonkhulu	Tree	Leaves, roots	[26]	Maceration	[26,34]
Drimia elata	Asparageceae	Undongana- zibomvana	Herb	Bulb	[26]	NS	[26]
Ekebergia capensis	Meliaceae	Essenhout	Tree	Leaves, bark	[26]	NS	[26]
Elephantorrhiza elephantina	Fabaceae	Intolwane	Sub-shrub	Leaves, rhizome	[26, 29, 30]	Infusion	[29]
Elytropappus rhinocerotis	Asteraceae	renosterbos	Shrub	Leaves	[27]	Infusion	[27]
Eriobotrya japonica	Rosaceae	Unknown	Shrub/ small tree	Leaves	[31]	Infusion	[31]
Eriocephalus africanus	Asteraceae	Kapokbos	Shrub	Leaves	[24, 25, 27, 30]	Infusion	[27]
Euclea undulata	Ebenaceae	Inkunzane	Shrub/ small tree	Bark, roots, whole plant	[24]	Infusion	[24]
Euryops abrotanifolius	Asteraceae	Bergharpuisbos	Shrub	Leaves, stems	[33]	NS	[33]
Galinsoga parviflora	Asteraceae	Unknown	Herb	Leaves	FPR	Maceration	[18]
Geranium incanum	Geraniaceae	Vrouetee	Grass or small herb	Leaves, stem	[29]	Decoction	[29]
Gethyllis spp	Amaryllidaceae	Koekoemakranka	N/A	Seeds, pod	[27]	Maceration	[27]
Helichrysum crispum	Asteraceae	Hotnotskooigoed	Herb	Leaves	[25, 27]	Infusion	[27]
Helichrysum odoratissimum	Asteraceae	Imphepho	Herb	Leaves, roots	[29]	Infusion	[29]
Hoodia gordonii	Apocynaceae	Bobbejaanghaap	Herb	Leaves, stem	[110]	NS	[110]
Hyphaene coriacea	Arecaceae	iLala	Tree	Roots	[32]	NS	[32]
Hypoxis argentea	Hypoxidaceae	Inongwe	Herb	Corm, tuber	[32]	Decoction	[32]
Hypoxis colchicifolia	Hypoxidaceae	Ilabatheka	Herb	Bulb	[29]	NS (chewed)	[29]
Hypoxis hemerocallidea	Hypoxidaceae	Inkomfe	Grasses or small herb	Corm, roots	[29,32]	Decoction	[29, 32]
Justicia flava	Acanthaceae	Impela	Herb	Leaves	FPR	Maceration	[18]
Lantana camara	Verbenaceae	Ubukhwebezane	Shrub	Roots	[31]	Infusion/decoction	[31] co

► Table 1 Continued

Name	Family	Common name (Zulu/Afrikaans)	Plant type	Part(s) used	Evidence of documented HBP use	Ethnobotanical indications on HBP	Reference
Leonotis leonurus	Lamiaceae	Umfincafincane	Shrub	Leaves, roots, flowers, whole plant	[27, 33]	Decoction	[27, 33]
Lessertia frutescens	Fabaceae	Umnwele	Shrub	Leaves, shoot	[29, 33, 110]	Decoction	[29]
Leucosidea sericea	Rosaceae	Umtshitshi	Small tree	NS	[28]	NS	[28]
Lichtensteinia lacera	Apiaceae	Kalmiswortel	Herb	Leaves, stem	[33]	Infusion	[33]
Lippia javanica	Verbanaceae	Umsuzwane	Shrub	Leaves	[32]	Decoction	[32]
Medicago sativa	Fabaceae	Unknown	Herb	Whole plant	[31]	Decoction	[28, 31]
Mentha aquatica	Lamiaceae	Unknown	Herb	Leaves, stem, seed	[28,30]	Infusion	[28, 30]
Mentha longifolia	Lamiaceae	ufuthana Iomhlanaga	Herb	Leaves, stem	[30,33]	Decoction	[33]
Mesembryanthe- mum spp.	Aizoaceae	Unknown	N/A	Leaves, stem	[26]	Maceration, decoction	[26, 34]
Momordica balsamina	Curcurbitaceae	Intshungu	Herb	Leaves	[32]	Decoction	[32,75]
Momordica charantia	Curcurbitaceae	Unknown	Herb	Whole plant	FPR	Maceration	[75]
Momordica foetida	Curcurbitaceae	Intshungu	Herb	Leaves, stem	[26]	NS	[26]
Musa acuminata	Musaceae	Ihliziyo	Herb	Flower bracts	[32]	Decoction	[32]
Ocimum basilicum	Lamiaceae	Unknown	Herb	Leaves, stem	[29]	Infusion	[29]
Oldenlandia affinis	Rubiaceae	Umampeshane	Herb	Root	[26]	Decoction	[26]
Olea europaea subsp. africana	Oleaceae	Umnquma	Tree	Leaves, roots	[23,24,26–29]	Decoction	[29]
Opuntia ficus	Cactaceae	Umthelekisi	Shrub/tree	Roots	[31]	Decoction	[31]
Oxygonum sinuatum	Polygonaceae	Unknown	Herb	Leaves	FPR	Maceration	[18]
Ozoroa engleri	Anacardiaceae	Isifico	Tree	Roots, bark, leaves	[32]	Decoction	[111]
Pentanisia prunelloides	Rubiaceae	Icimamlilo	Herb	Rhizome, corm, roots	[24, 26]	NS	[26]
Persea americana	Lauraceae	Unknown	Tree	Leaf, pulp, fruit, root	[31]	Decoction	[31]
Peucedanum galbanum	Apiaceae	Droedas	Shrub	Leaves	[24]	Infusion	[24]
Physalis viscosa	Solanaceae	Unknown	Herb	Leaves	FPR	Maceration	[18]
Protorhus longifolia	Anacardiaceae	Uzintlwa	Tree	Bark, leaves	[26]	Maceration	[26, 34]
Psidium guajava	Myrtaceae	Koejawel	Shrub/ small tree	Leaves, roots	[31,32]	Decoction	[31, 32]
Pyrenacantha kaurabassana	lcacinaceae	Unknown	Shrub	Bulb, tuber	[32]	Decoction	[32]
Ptaeroxylon obliquum	Rutaceae	umThathi	Tree	Roots	[24,32]	Maceration	[26, 32]
Rapanea melanophloeos	Myrsinaceae	iKhubalwane	Tree	Bark	[26]	NS	[26]
Rauvolfia caffra	Apocynaceae	um Hlambamanzi	Tree	Bark, stem, whole plant	[24]	NS	[24]
Rhoicissus digitata	Vitaceae	umNangwazi	Shrub	Tubers, bulb	[29]	Infusion	[29]
Rhus chirindensis	Anacardiaceae	Umhlabamvudu	Tree	Leaves, roots, bark, twigs, fruits	[26]	Maceration	[34]

Table 1 Continued

Name	Family	Common name (Zulu/Afrikaans)	Plant type	Part(s) used	Evidence of documented HBP use	Ethnobotanical indications on HBP	References
Ricinus communis	Euphorbiaceae	Umhlakuva	Shrub	Leaves	[32]	Decoction	[32]
Ruta graveolens	Rutaceae	Unknown	Shrub	Leaves	[24, 25, 27, 29]	Infusion	[27,29]
Salvia africana- caerulea	Lamiaceae	Wildesalie	Shrub	Twig, leaves	[28,33]	Infusion	[33]
Sarcophyte sanguinea	Balanophoraceae	Umvumbuka	Herb	Whole plant, Stem, root	[32]	Infusion, decoction	[32]
Sarcostemma viminale	Apocynaceae	Umbelebele	Shrub	Stem, aerial parts, twigs	[32]	Infusion	[32]
Sceletium tortuosum	Aizoaceae	Tandtrekbos	Herb	Leaves, roots	[33]	NS	[33]
Schkuhria pinnata	Asteraceae	Ruhwahwa	Herb	Whole plant	[29]	Infusion, decoction	[31]
Sclerocarya birrea	Anacardiaceae	Maroela	Tree	Bark, leaves, stem-bark, stem	[26]	Decoction	[26]
Scolopia mundii	Salicaceae	iHlambahlale	Tree	Bark	[26]	NS	[26]
Searsia burchellii	Anacardiaceae	Karookoeniebos	Shrub/ small tree	Leaves, stem, roots	[33]	Infusion	[33]
Senecio bupleuroides	Asteraceae	Isiqandamatshana	Shrub	NS	[26]	NS	[26]
Senecio inornatus	Asteraceae	Uhlabo	Herb/small shrub	Root	[26]	Decoction	[26]
Senecio serratuloides	Asteraceae	Ichazampukane	Herb	Leaves, stem	[32]	Decoction	[32]
Spermacoce natalensis	Rubiaceae	Umabophe	Herb	Bark, roots, leaves	[26]	NS	[46]
Stangeria eriopus	Zamiaceae	Umfigwani	Cycad	Root, leaves	[26]	NS	[34]
Strychnos madagascariensis	Strychnaceae	umKwakwa	Tree	Seeds, bark, roots, fruit	[32]	NS (crush to powder for swallowing)	[32]
Teedia lucida	Scrophulariaceae	Hlwenya	Herb/dwarf shrub	NS	[28]	NS	[28]
Tephrosia capensis	Fabaceae	Unknown	Small shrub	Root	[26]	Decoction	[28]
Tetradenia riparia	Lamiaceae	Ibozane	Shrub/tree	Leaves, seeds	[32]	Decoction	[32]
Trichilia emetica	Meliaceae	Umathuzini	Tree	Leaves, fruits, roots, bark, stem	[32]	Poultice, decoction	[32]
Trifolium africanum	Fabaceae	Wildeklawer	Herb	Whole plant	[28,30]	Infusion	[30]
Trifolium burchellianum	Fabaceae	Usithathi	Herb	Leaf, stem, roots	[28, 30]	Infusion (com- bined with <i>M. aquatica</i>)	[30]
Tulbaghia acutiloba	Alliaceae	Ishaladi Lezinyoka	Herb	Bulb, flower, Whole plant	[28,30]	Infusion, decoction	[30]
Tulbaghia violacea	Alliaceae	wilde knoffel	Herb	Rhizome, bulb, leaves, roots	[24, 26–29, 33]	Infusion, decoctions	[27, 29, 33]
Turraea floribunda	Meliaceae	Umadlozane	Tree	Bark, leaves, roots	[26]	Infusions	[26]
Urtica urens	Urticaceae	Unknown	Herb	Leaves	[30]	Decoction	[30]
Valeriana capensis	Valerianaceae	Wildebalderjan	Herb	Rhizomes, roots	[24]	NS	[24]
Vangueria infausta	Rubiaceae	umTulwa	Shrub/ small tree	Bark, leaves	[32]	Maceration, infusion	[32]
Warburgia salutaris	Canellaceae	Isibhaha	Tree	Bark, root	[110]	NS	[110]

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Name	Pharmacological activities	al activities		Assay	Method(s)/	Part(s) used	Medium for	Conc. tested	Province(s)	References
	Antioxidant,	Antidiabetic	Antihypertensive, cardiovascular	type(s)	inducing agent		extraction		or location	
Acokanthera oppositifolia	Antioxidant			In vitro	DPPH, ABTS, FRAP	Stem	Methanol	0.1 mg/mL	Eastern Cape	[112]
Adenopodia spicata			Antihypertensive	In vitro	ACEI assay	Leaves, root	Aqueous, ethanol	25 µg/mL	KZN	[34]
Aganpanthus africanus			Antihypertensive	In vitro	ACEI assay	Leaves, root	Aqueous	25 µg/mL	KZN	[34]
Agathosma	Antioxidant			In vitro	ABTS, DPPH	Leaves	Hot water, DCM	1, 3, 5, 10, 20 µg/mL Pretoria	Pretoria	[113]
betulina		Antidiabetic		In vitro	α-amylase, α-glucosidase					
	Antioxidant			In vivo	ORAC, TEAC, FRAP		Aqueous, ethanol, acetone		Cape Town	[114]
Agave americana			Antihypertensive	In vitro	ACEI assay	Leaves	Aqueous, ethanol	25 µg/mL	KZN	[34]
Alepidea amatymbica			Antihypertensive, cardiovascular, diuretic	oviv nl	Sodium thiopentone (rats model)	Rhizome	Hexane, methanol, DCM	20 mg/kg body weight	KZN	[38, 115]
Amaranthus dubius			Antihypertensive	In vitro	ACEI	Leaves	Aqueous, methanol	0.1 g/mL	KZN	[18]
Amaranthus hybridus			ACE inhibitory activity	In vitro	ACEI	Leaves	Aqueous, methanol	0.1 g/mL	KZN	[18]
Artemisia afra		Antidiabetic		In vivo	STZ-induced rat model	Leaves	Aqueous	50, 100, 200 mg/kg	Eastern Cape	[116]
			Cardioprotection	In vivo	ISP	Leaves	Aqueous	100, 200 mg/kg	Eastern Cape	[117]
Asystasia gangetica			Antihypertensive	In vitro and	ACEI	Leaves	Aqueous and methanol	0.1 g/mL	KZN	[18]
				In vivo	Sodium pento- barbitone		Aqueous	10, 25, 50, 100, 200, 400 mg/mL	Western Cape	[46]
Carpobrotus dimidiathus	Antioxidant			In vitro	DPPH, FRAC	Leaves	Aqueous methanol (50%)	10, 20, 30, 40, 50, 60, 70 µg/mL	KZN	[108]
Catha edulis			Antihypertensive	In vitro	ACEI assay	Leaves	Aqueous, ethanol	25 µg/mL	KZN	[34]
Citrus lemon	Antioxidant			In vitro	DPPH, nitric oxide, reducing power,	Peel of fruits	Ethanol, acetone	0.025, 0.05, 0.1, 0.2, Eastern Cape 0.5 ug/mL	Eastern Cape	[54] continued

References		[59]	[58]	[62–63]	[118]	[119]	[34]	[34]	[120]	[120]	[121]	[75]	[75]	[18]	continued
Province(s)	or location	Limpopo	KZN	Western Cape	Free State	Same	KZN	KZN	NZX	Gauteng (Pretoria)	Gauteng (Pretoria)	KZN	KZN	KZN	
Conc. tested		50, 100, 200, 400 mg/bw	100, 200, 400, 800 mg/kg bw	0.05, 0.10, 0.15, 0.20 mg/kg	125, 250, 500 mg/kg	same	25 µg/mL	25 µg/mL	18 mg/kg i.v. and 120 mg/kg p.o	50 µg/mL and 16–250 µg/mL GU	50, 100 mg/kg b.w	12.5, 25, 50, 100, 200, 400 mg/mL	25, 50, 100, 200, 400 mg/kg b.w	0.1 g/mL	
Ļ	extraction	Aqueous 4	Methanol	Aqueous (Aqueous	Aqueous	Aqueous	Aqueous	Ethanol	Acetone	Acetone	Aqueous	Same as above	Aqueous, methanol (
Part(s) used	•	Leaves	Roots	Whole plants	Root	Root	Leaves, root	Leaves, root	Leaves	Whole plant	Bark	Fresh corm	As above	Leaves	
Method(s)/	inducing agent	HRM	STZ- rat model	Normotensive rats	STZ-rat model	ISP-rat	ACEI assay	ACEI assay	Inactin- induced normoten- sive rats and Dahl salt stress rats	α-amylase, α-gluco- sidase, glucose uptake (GU)	STZ-rat model	Isolated muscle experiment	Sodium 5-ethyl- (1-methylbutyl)-2- thiobarbiturate- induced Dahl salt- stress rats	ACEI assay	
Assay	type(s)	In vivo	In vivo	In vivo	In vivo	In vivo	In vitro	In vitro	uivo	In vitro	In vivo	In vitro	oviv nl	In vitro	
	Antihypertensive, cardiovascular	Antihypertensive		Cardiovascular		Cardioprotection	Antihypertensive	Antihypertensive	Cardiovascular, vasorelaxant, bradycardia			Cardiovascular, anti- hypertensive		Antihypertensive	
activities	Antidiabetic		Antidiabetic		Antidiabetic					Hypoglycemic					
Pharmacological activities	Antioxidant,				Antioxidant										
Name		Clausena anisata		Crinum macowani	Dicoma anomala		Dietes iridioides	Domb <i>eya</i> rotundifolia	Ekeberghia capensis	Euclea undulata		Hypoxis hemerocallidea		Justicia flava	

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Table 2 Continued

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	Pharmacological activities	al activities		Assay	Method(s)/	Part(s) used	Medium for	Conc. tested	Province(s)	References
	Antioxidant,	Antidiabetic	Antihypertensive, cardiovascular	type(s)	inducing agent		extraction		or location	
Leonotis leonurus	Antioxidant			In vitro	DPPH, ABTS radical scavenging assays	Leaves	Acetone, Aqueous, methanol	0.02, 0.05, 0.1, 0.5, 0.75, 1 mg/mL	Eastern Cape	[122]
		Antidiabetic		oviv nl	STZ-rats model	Leaves	Aqueous	50, 100, 200, 400, 800 mg/kg b.w.	KZN	[123]
					As above			125, 250, 500 mg/ kg b.w.		
						Leaves	Aqueous	0.5–7.0 mg/kg	Eastern Cape	[123]
					Sodium pentobarbi- tone induced rats		Methanol	25, 50, 100, 200, 400, 800 mg/kg	Western Cape	
			Cardiovascular	In vitro, In vivo	Dahl salt hyperten- sive rats	Leaves				[124]
			Normotensive, antihypertensive	oviv nl	Sodium pentobarbi- tone induced rats	Leaves	Aqueous	1, 2 mg/mL	KZN	[123]
			Normotensive	In vivo		Whole plant	Decoction		Western Cape	[64]
		Hypoglycaemic		In vivo	STZ-induced rats	Shoot	Aqueous	50, 100, 200, 400, 800 mg/kg	KZN	[100]
				In vivo	High fat diet	Leaves	Hot water	0.1 g/mL	Eastern Cape	[125]
	Antioxidant			In vitro	DPPH, ABTS, FRAC	Leaves	Aqueous, Aqueous-methanol (70%)	1, 2, 3, 4, 5 mg/mL, 0.3 g/mL	Gauteng	[126]
Mesembryanthe- mum spp			Antihypertensive	In vitro	ACEI assay	Leaves, stem	Aqueous, ethanol	25 µg/mL	KZN	[34]
		Hypoglycaemic		oviv nl	STZ-induced rats	Whole plant	Aqueous	50, 100, 200, 400, 800 mg/kg	KZN	[75]
			Hypotensive	ln vivo	sodium 5-ethyl-(1- methylbutyl)-2-thi- obarbiturat-induced Dahl salt-stress rats	As above	As above	As above	As above	[75]
ida	Momordica foetida Antioxidant			In vitro	DPPH, ABTS, Reducing power, metal chelating	Leaves	Methanol	0.1 mg/mL	KZN	[127]
		Antidiabetic		Cell lines	Chang, 3T3 L1, C2Cl2 cells	Whole plant	DCM-methanol (1 : 1), Water	NS	Eastern Cape	[128]
			Antihypertensive	In vitro	ACEI	Leaves	Aqueous, methanol	0.1 g/mL	KZN	[18]
										continued

Name	Pharmacological activities	al activities		Assay	Method(s)/	Part(s) used	Medium for	Conc. tested	Province(s)	References
	Antioxidant,	Antidiabetic	Antihypertensive, cardiovascular	type(s)	inducing agent		extraction		or location	
Pentanissia prunelloides	Antioxidant			In vitro	DPPH, Superoxide anion, Hydroxyl radical, Metal chelating	Root	Aqueous, ethanol, hydro-ethanol (1:1), hexane	6.25, 12.5, 25, 50, 100 µg/mL	Free State	[129]
		Antidiabetic		In vitro	Alpha (amylase and glucosidase)					
Physalis viscosa			Antihypertensive	In vitro	ACEI	Leaves	Methanol, aqueous	0.1 g/mL	KZN	[18]
Protorhus Iongifolia	Antioxidant			In vitro	DPPH, ABTS, metal chelating, Reducing power	Bark	Hexane, chlorofum, ethyl acetate, methanol, aqueous	0–5 mg/100 mL	KZN	[89]
			Antihypertensive	In vitro	ACEI	Leaves	Aqueous, ethanol	25 µg/mL	KZN	[34]
Psidium guajava		Hypoglycemic	Hypotensive	oviv nl	STZ-induced rats and Dahl salt-stress rats	Leaves	Aqueous	50, 100, 200, 400, 800 mg/mL	KZN	[93]
Rauvolfia caffra	Antioxidant			In vitro	FRAP, DPPH,	Stem, bark	DCM, methanol, ethyl acetate, ethanol	1 mg/mL	Limpopo	[96]
			Antihypertensive	In vivo	Spontaneously hypertensive rats	Whole plants	DCM, methanol, ethyl acetate	15, 300, 500 mg/kg b.w		
Rhus chirindensis			Antihypertensive	In vitro	ACEI	Leaves	Aqueous, ethanol	25 µg/mL	KZN	[34]
Sclerocarya birrea		Hypoglycemic		oviv nl	STZ	Stem-bark	Aqueous	25, 50, 100, 200, 400, 800, 1600 mg/mL	KZN	[100]
			Antihypertensive	In vitro	ACEI	Leaves	Aqueous, ethanol	25 µg/mL	KZN	[34, 101]
				In vivo	Dahl salt sensitive rats	Stem bark	Aqueous	25, 50, 100, 200, 400 mg/mL	KZN	
Senecio serratulloides	Antioxidant			In vitro	DPPH, FRAC	Leaves	Aqueous methanol (50%)	10, 20, 30, 40, 50, 60, 70 µg/mL	KZN	[112]
Stangeria eriopus			Antihypertensive	In vitro	ACEI	Leaves	Aqueous, ethanol	25 µg/mL	KZN	[34]
Trichilia emetica	Antioxidant			In vitro	DPPH	leaves	Aqueous, methanol		Gauteng	[130]
										continued

Table 2 Continued

	Pharmacological activities	al activities		Assay	Method(s)/	Part(s) used	Medium for	Conc. tested	Province(s)	References
A	Antioxidant,	Antidiabetic	Antihypertensive, cardiovascular	type(s)	inducing agent		extraction		or location	
Tulbaghia Av violacea	Antioxidant			In vitro	DPPH, reducing power, H ₂ O ₂	Rhizome	Methanol	0.02, 10, 20, 30, 40, Eastern Cape 50 µg/mL	Eastern Cape	[29]
		Antidiabetic		In vivo	STZ	Rhizome	Aqueous	60, 120 mg/kg	KZN	[103]
					ACEI			0.1 g/mL	KZN	
			Antihypertensive	In vitro		Leaves	Methanol, aqueous	60 mg/kg		[18]
				In vivo	Human Angiotensin I, II acetate salt hydrate	Leaves	Methanol		Western Cape	[104]
					Normotensive rats	Leaves	Methanol	5, 10, 20, 40, 80, 160 mg/kg	Western Cape	[105]

agent against streptozocin-induced diabetic Wistar rats was reported when it lowered blood glucose levels after 14 d of administration at 3 concentrations (0.1, 0.25, 0.5 g/mg body weight) [42]. Furthermore, its antibacterial, analgesic, prophylactic, antioxidant, antimicrobial, phytotoxic, and anticoccidal properties, as well as its promotion of fertility and reduction of red blood cells' osmotic fragility, have been mentioned prominently in previous studies.

Amaranthus dubius Mart ex. Thell.

A. dubius is a member of the Amaranthaceae family, with the common name of spleen amaranth. The plant was domesticated in South America and lately instituted in Europe, Asia, and Africa. Medicinally, the plant is used as an antidote for kidney problems, anemia, fever, hemorrhage, stomach troubles [43], and HTN [18], while the potential active phytoconstituents in the plant include but are not limited to ascorbic acid, thiamine, riboflavin, β -carotene, iron, calcium, alkaloids, tannins, saponins, and glycosides. The methanolic extract of the plant showed good inhibition (67%) against ACE (**> Table 2**), indicating antihypertensive activity *in vitro* according to Ramesar et al. [18].

Amaranthus hybridus (L.)

A. hybridus is also a member of the Amaranthaceae family endemic to eastern North America, with distribution in other continents such as Asia (Thailand, India, Japan, Lebanon, Pakistan, Indonesia, and Jordan), Africa (SA, Ethiopia, Kenya, Tanzania, and Zambia), and South America (Argentina, Brazil, Peru, and Colombia). The common names (local and international) include green or slim amaranth (English), terere (Kenya), bledo, guelite (Spanish), amarante hybride (French), caruru-de-folha-larga (Portuguese), caruru-branco, caruru-roxo (Brazil), bastard-amarant, gruenaehriger amarant (Germany), honagaaogeito (Japan), and basterdamarant (Netherlands). Aside from being used medicinally as a cure for diarrhea, anemia, menstrual/intestinal bleeding, wound healing, and for burnt or itchy skin [43] with the presence of cardiac glycosides, steroids, flavonoids, and terpenoids as inherent active ingredients, the plant is also cultivated for its nutritive value (protein, vitamins, and minerals) and is consumed in SA and some other African countries as a leafy vegetable. Pharmacologically, A. hybridus has been reported to possess antimicrobial, hepatoprotective, anticarcinogenic, and antioxidant activities, as well as in vitro antihypertensive activity, when an aqueous extract of the plant was tested at a concentration of 0.1 g/mL against ACE in a South African study [18] (> Table 2).

Asystasia gangetica (L.) T. Anderson subsp. micrantha (Nees) Ensermu

A. gangetica, locally called creeping foxglove (English) or isihobo (Zulu), is associated with the Acanthaceae family (http://pza. sanbi.org/asystasia-gangetica). It is a fast-growing herbaceous groundcover plant that extends from 300–600 mm. Based on the information available about its distribution, it is located in tropical areas from Asia to SA (Eastern Cape, KZN, Limpopo, and Mpumalanga) and other southern African countries like Namibia, Botswana, and Swaziland. In fact, the South African subspecies differs from that of the Asian subspecies and has prominent pink flowers. Because of its nutritive value, the leaves are consumed

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as a spinach [43], while it is also used in traditional medicine for the management of asthma in many regions of Nigeria. Its active phytoconstituents include flavonoids, phenols, alkaloids, cardiac glycosides, tannins, and terpenoids [44], with 5, 11-epoxymegastigmane glucoside and megastigmane glucoside as some of the isolated compounds. In Nigeria, the antibacterial, antifungal, bronchospasmolylic, antihyperlipidaemic, and anthelmintic activities of *A. gangetica* have been reported. Other pharmacological activities include but are not limited to anti-inflammatory, antihyperglycemic, antioxidant, and cytotoxic [44, 45]. Interestingly in a SA study, *A. gangetica* inhibited the activity of ACE *in vitro* [18], while also reducing the BP level and heart rate of spontaneously hypertensive rats when varying concentrations (10– 400 mg/kg body weight) of the aqueous extract were administered orally [46] (**> Table 2**).

Catha edulis (Vahl.) Endl.

C. edulis, a member of the Celastraceae family, is a very attractive shrub with glowing green leaves that grows up to 10-m high. Widely distributed in tropical Africa including Ethiopia, SA (KZN, Eastern Cape, Mpumalanga, and Limpopo), and the Arabian Peninsula, the local names are bushman's tea (English), boesmanstee (Afrikaans), umhlwazi (Zulu), iggwaka (Khosa), and khat (Arabic). Besides its other uses as firewood, furniture, insect repellent, and fencing poles, it medicinally functions against respiratory diseases, relieving fatigue and sleeplessness, and is also useful as an antidote against cough, asthma, and other chest illnesses [24], as well as for stimulating the heart [26]. In SA, it is regarded as a drug because its active ingredient, cathinone (an alkaloid which is closely related to ephedrine and amphetamine), is listed in the Drug Act. Other polyphenolic compounds including tannins, flavonoids, terpenes, sterols, and essential oils are present [47], from which compounds such as cathidine, cathiduline, norephedrine, and cathinone are isolated. Similarly, the essential oil has also been found to contain carvotanaacetone, trans-pulegol, and 2,5-dimethoxy-p-cymene. Pharmacologically, its effect is similar to amphetamine, and it is thus referred to as a natural amphetamine [48]. Other properties include hyperthermia, analgesic, hyperactivity, anorexia, cardiovascular, antimicrobial, antifungal, antibacterial, cytotoxic, antimicrobial, anti-inflammatory, antioxidant, antiulcer, hepatotoxic, and nephrotoxic. In addition, its in vitro antihypertensive properties were established by a study conducted in KZN in which aqueous and ethanolic leaf extracts of the plant at $25 \,\mu g/mL$ inhibited the activity of ACE by 48% and 82%, respectively (see > Table 2) [34].

Catharanthus roseus (L.) G.Don

The common names of this Apocynaceae family plant include Madagascar periwinkle, rose periwinkle, Cape periwinkle, old maid, bright-eyes, and vinca (English). The plant is endemic to Madagascar but may be found in other areas of Africa, such as Kenya, Uganda, Tanzania, and SA [32] as an ornamental or a MP. The plant is traditionally used for treating diabetes, eye inflammation, rheumatism, malaria, and HBP [32]. Vinblastine and vincristine are alkaloids isolated from the herb and are chemotherapeutic agents used for managing various types of cancer [49]. The anti oxidant, antidiabetic, anthelmintic, antineoplastic, antidiarrheal,

antimicrobial, anticancer, antiulcer, wound healing, hypolipidemic, memory enhancing, and hypotensive activities of the plant were mentioned in a recent review by Nisar et al. [50]. Moreover, in an experiment to evaluate the antihypertensive effect, which was conducted by Naznin et al. using the ethanolic leaf extract of the plant [51], the results showed that the BPs of adrenaline-induced hypertensive rats were lowered when a dose of the extract (30 mg/155 g body weight) was administered to the animals daily for 1 wk, indicating its hypotensive effect. Furthermore, the safety profile of the plant has also been established [52].

Citrus limon (L.) Osbeck

C. limon is commonly named lemon (English) or ulamula (Zulu), and belongs to the Rutaceae family. It is a spiny, all-year-round shrub or small tree that may grow up to 6 m in height. The plant is thought to have originated in Asia and today is one of the most widely homegrown fruit trees. Aside from being rich in vitamin C, which assists in warding off infections, it is traditionally used to treat scurvy, sore throats, fevers, rheumatism, HBP, and chest pain [32]. Moreover, the detected phytochemicals include but are not limited to phenolic acids, flavonoids, triterpenoids, carotenoids, essential oils, and tannins [53], while a series of pharmacological benefits such as anticancer, hypoglycemic, anti-obesity, antimicrobial, antiviral, antihyperlipidemic, antihypercholesterolemic, hypotensive, anti-inflammatory, analgesic, and hepatoprotective [54] have been reported.

Citrus maxima (Burm.) Merr

C. maxima, another Rutaceae family plant is commonly referred to as pummelo (English) or upapamuzi (Zulu). C. maxima is a lowbranching evergreen tree 5–10 m in height, though it may sometimes grow as tall as 15 m. It is native to Southeast Asia but has distribution in tropical and subtropical regions such as Thailand, southern China, Japan, Indonesia, Malaysia, the Philippines, and the United States. The plant is a natural (nonhybrid) citrus fruit and is the largest of all the citrus species, bearing a great resemblance to the grapefruit, though it is larger, not bitter, and possesses a thicker rind. Traditionally, it is used in the treatment of cough, fever, gastric ailments, HTN, hemorrhage, and epilepsy [32, 55]. In a comprehensive review by Vijaylakshmi and Radha [55], its pharmacological properties including antioxidant, antidiabetic, analgesic, anti-inflammatory, antidepressant, anticonvulsant, antitumor, hypnotic, hepatoprotective, antibacterial, antihypercholesterolemic, and hypotensive have been reported. Interestingly, a report by Oboh et al. [56] established the antihypertensive effect of the juice extract of the plant, which inhibited ACE (> Table 2) by more than 50%. In addition, the juice reduces the plasma total cholesterol, triglycerides, and low-density lipoprotein-cholesterol levels of rats fed a high cholesterol diet, with subsequent increase in the high-density lipoprotein-cholesterol.

Clausena anisata (Willd.) Hook. f.ex Benth

C. anisata is a member of the Rutaceae family and is locally called isifudu, isifuthu, umsanka, umnukambhiba (Zulu), basternieshout, perdepis, baster-perdepis (Afrikaans), umnukambiba, umnukandiba (Xhosa), and horsewood, or false horsewood (English). The plant is not endemic to SA, but is widely distributed in Eastern

Cape, Free State, KZN, Limpopo, Mpumalanga, and Western Cape. Traditionally, it is used as a remedy for heart diseases [57], tape worm, rheumatism, fever, bad breath, as a blood purifier, and for diseases of the liver [26]. The active phytoconstituents are terpenoids, sesquiterpenes, fatty acids, alkaloids, coumarins, limonoids, essential oils, phenolics, tannins, and saponins [57], with numerous compounds (e.g., limonene, myrcene umbelliferone, clausanitin, and mupamine) isolated from various parts of the plant (roots, pericarp, root bark, and leaves). Five known compounds have been isolated from the essential oil: β -pinene (32.8%), sabinene (28.3%), germacrene-D (12.7%), estragole (6.4%), and linalool (5.9%), which are found to possess larvicidal action against Anopheles stephensi (Culicidae) Liston, Aedes aegypti (Culicidae) Linn., and Culex guinguefasciatus (Culicidae) Say. When different concentrations of the extract were administered to streptozotocin-induced diabetic rats, their blood glucose levels were restored, demonstrating the plant's hypoglycemic effects [58]. The antioxidant and anti-inflammatory activities of acetone extracts of the plant against lipoxygenase and nitric oxide synthase enzymes; the analgesic, antifungal, antiplasmodial, and antibacterial activity of clausenol (carbazole alkaloids) isolated from the plant; and the activity of crude plant extracts against bacterial and fungal isolates and HIV, as well as their antioxidant, cytotoxic, wound healing, antimalarial, anticonvulsant, anthelmintic, and antimicrobial properties, are available in scientific journals. Interestingly, C. anisata was reported to lower BP at a dose of 400 mg/kg in a hypertensive rat model in a 40-d experimental study [59], suggesting its antihypertensive effects.

Crinum macowanii Baker

C. macowanii is a deciduous, summer-growing bulb about 600-2500 cm long, with a fleshy perennial root. This Amaryllidaceae family plant with common names such as river crinum, river lily, sabie crinum, Cape coast lily (English), riverlelie, boslelie, sabielelie (Afrikaans), intelezi (Xhosa), and umdube (Zulu) is indigenous (with 19 other species) to southern Africa, particularly SA (Eastern Cape, Free State, KZN, Mpumalanga, Gauteng, North West, and Limpopo), despite being the most widely distributed of the Crinum species. The plant is traditionally embraced as a remedy for kidney and bladder infections, tuberculosis, fever, scrofula, micturition, rheumatic fever, itchy rashes, boils, acne, and venereal ailments [23], as well as for the stimulation of milk production in women and cattle. The active constituents are alkaloids, with more than 11 isolated compounds such as lycorine (major compound), macowine, crinamidine, powelline, crinine, buphanidrine, krepowine, cheryline, undulatine, 4a-dehydroxycrinamabine, and 1-epideacetylbowdensine [60], and it has been reported to have antitumor, analgesic, and hypotensive potential [61]. Other activities include cytotoxic, antimalarial, antiviral against exotic RNA viruses, antifungal, genotoxic, mutagenic, and nematicidal. Additionally, dose-dependent increases in heart rate, as well as systolic and diastolic pressures were observed when anaesthetized normotensive Wistar rats were pretreated with i.v. infusion of the aqueous extract of the plant either singly or in combination with cardioactive substances such as atropine, atenolol, reserpine, prazosin, and verapamil. The effects of the extract on the tested parameters were on par with those of the known cardioactive substances (except for verapamil in systolic and diastolic pressures), indicating the cardiovascular effects of the plant [62]. Similarly, hippadine (another compound isolated from the plant) at various doses (0.05, 0.10, 0.15, 0.20 mg/kg) has also been reported to lower systolic and diastolic BPs, as well as mean arterial BP, suggesting its antihypertensive [63] and cardiovascular [64] effects.

Dietes iridioides (L.) Sweet ex Klatt

The rhizomatous *D. iridioides* is an evergreen herb that extends up to 600 mm in height and is sword-shaped with dark-green leaves in a loose fan. It is a member of the Iridaceae family, with local names such as wood iris (English), indawo-yehlathi, isiqiki-sikato-koloshe, and isishuphe somfula (Zulu), and is widely spread from Eastern Cape through KZN to Gauteng in the green forests or cleared bush of SA and Ethiopia. Infusions of the inner contents of the rhizomes are orally taken to stop dysentery [26]. Similarly, the rhizomes are used during childbirth and for HTN control [65], while they may also serve as constituents of tonics used in goats and sheep. The *in vitro* antiprotozoal and antihypertensive effects were suggested by Duncan et al., who reported that it inhibited the activity of ACE when evaluated with other Zulu plants [34] (**► Table 2**).

Dombeya rotundifolia Hochst

Wild pear (English), drolpeer (Afrikaans), iNhliziyonkhulu (Zulu), mohlabaphala (Northern Sotho), motubane (Tswana), and nsihaphukuma (Tsonga) are local names for D. rotundifolia, a member of the Malvaceae family. It is a deciduous, fast-growing, and frostand drought-resistant tree that grows up to 10 m in height but is usually between 3 and 6 m. Medicinally, the inner bark is used to treat weakness of the heart. Other uses of various parts of the plant are to treat intestinal ulceration, stomach troubles, and sharp stomach pains, fever, diarrhea, palpitation, nausea, enemas for dyspepsia, and to induce labor [24, 26]. Its identified chemical moieties include but are not limited to saponins, tannins, and cardiac glycosides [66], with a few isolated compounds such as lauric, myristic, stearic, and palmitic acids with antibacterial activity. Similarly, reports of lupeol and β -sitosterol being isolated from the bark of the stem have been documented [67]. Reported biological activities are antibacterial, anti-inflammatory [66], anthelminthic, antidiarrheal, and abortifacient. Furthermore, Duncan et al. [34] suggested that ethanolic extracts of the plant leaf have antihypertensive effects based on their inhibition of ACE (83%) (> Table 2).

Ekebergia capensis Sparrm.

E. capensis is an evergreen tree in the Meliaceae family with the potential to grow as high as 15 m, existing in different habitats ranging from high altitude evergreen forests to riverine forests, and from sea level to approximately 1500 m above sea level. Commonly called Cape ash, dogplum (English), essenhout (Afrikaans), mmidibidi (Northern Sotho), umnyamatsi (Southern Sotho), and nyamaru (Tswana), *E. capensis* grows in southern Mozambique, Zimbabwe, Swaziland, Uganda, Ethiopia, Congo, and SA, especially Eastern Cape, KZN, and Limpopo. It is used to treat various disease conditions, such as dysentery, heartburn, cough, acute gastritis, headache, scabies, abscesses, boils, and acne. Moreover, the antimicrobial, antituberculotic, antitrypanosomal, cytotoxic,

antimicrobial, anti-inflammatory, mutagenic, antimutagenic, antimalarial, and uterogenic potential of the plant have been affirmed. Similarly, Kamadyaapa et al. [68] reported that acute intravenous (18 mg/kg) and subchronic oral (120 mg/kg) administration of ethanolic leaf extracts of the plant resulted in a hypotensive effect on anaesthetized normotensive rats, while it also prevented the onset of HBP in weanling Dahl salt stress rats [69], suggesting a vasorelaxing action mediated via endotheliumderived relaxing factor-dependent or independent pathways, as well as a hypotensive effect mediated by regulating the total peripheral resistance of the vascular smooth muscles [70]. The chemical constituents of E. capensis are glycosides, polyphenols, tannins, triterpenes (major), and saponins. Further efforts to isolate the photoactive compounds yielded oleanonic acid; 2,3,22,23-tetrahydroxy-2,6,10,15,19,23-hexamethyl-6,10,14,18tetracosatetraene; oleanolic acid, ekeberin A; proceranolide; kaempferol-3-*O*-β-D-glucopyranoside; 3-*epi*-oleanolic acid; quercetin-3-O-β-D-glucopyranoside and 2-hydroxymethyl-2,3,22,23tetrahydroxy-6,10,15,19,23-pentamethyl-6,10,14,18-tetracosatetraene from triterpenoids, some of which have been found to possess antimalarial, cytotoxic, hypoglycemic, and antihypertensive activities [71,72].

Galinsoga parviflora (Cav.)

G. parviflora is an herbaceous plant from the Asteraceae family. It is endemic to South America but is well distributed in tropical, subtropical, and temperate regions of the world. Locally called chickweed, gallant soldier, or kew weed (English), the herb is about 60-75 cm tall, with oppositely clustered leaves that are toothed at the margins. Isolated compounds identified within the plant are apigenin, 7- β -d-glucoside, luteolin, phytol, stigmasterol, β -sitosterol, fumaric acid, uracil emanating from flavonoids and so on [73]. It is used as flavoring in the preparation of ajiaco soup, a very common and popular soup to the people of Colombia, Cuba, and Peru, and is also used in fruit salads. Medicinally, it is used for curing sores, colds, and wound healing. Pharmacologically, the plant possesses antibacterial, anti-inflammatory, wound healing, antioxidant, and allelopathic effects, as well as having established antihypertensive activity, as a methanolic extract inhibited the activity of ACE by 56% in a study conducted by Ramesar et al. [18] (> Table 2).

Hypoxis hemerocallidea Fisch., C. A.Mey. & Ave-Lall

H. hemerocallidea, honored as the "miracle muti" and "wonder potato" by the majority of South African herbal users, is a very beautiful, tuberous, perennial member of the Hypoxidaceae family, with a yellow star-like flower that signifies the onset of spring and summer rains. Commonly referred to as starflower and other names [74], the plant is medicinally employed in SA as an antidote for various human and veterinary diseases and is likely the most highly researched plant in Africa. It is useful against numerous ailments such as heart weakness, stroke, HBP [29], cancer, testicular tumors, infertility (male and female), headaches, dizziness, urinary tract infections, psychiatric disorders, and burns [26], while the prominent phytoconstituents include but are not limited to phytosterol glucosides (β -sitosterol), diglucoside hypoxoside, aglycone rooperol, sterols, and sterolins [25,26]. Interestingly, several pharmacological indications such as antioxidant, antibacterial, anticonvulsant, antiviral, antidiarrheal, antinociceptive, anti-inflammatory, antidiabetic, uterolytic, anti-HIV, anticancer, and cardiovascular effects of the plant have been reported in our recent review and those of others [74]. The cardiovascular effects as reported by Ojewole et al. [75] were revealed when the aqueous corm extract (12.5–400 mg/mL) of the plant produced prominent inotropic and chronotropic responses on the myocardial contractility of excised guinea pig muscles. This finding was corroborated in an animal model when the extract (25–400 mg/kg) lowered the systemic arterial BP and heart rate of Dahl salt-stress hypertensive rats, indicating the antihypertensive effect of the extract (**> Table 2**).

Justicia flava (Vahl) Vahl

J. flava, a perennial herb or shrub, is a member of the Acanthaceae family locally called yellow justice (English), geelgarnaalbos (Afrikaans), and impela (Zulu), and is widely distributed from tropical east Africa to SA (North West, Eastern Cape, KZN, Limpopo, Gauteng). In folkloric medicine, it is used to treat cough, stomach pain, diarrhea, fevers, dysentery, and convulsions [26]. The active constituents are β -sitosterol, stigmasterol, campesterol, β -sitosterol- β -D-glucoside, and salicyclic acid. In addition, its isolated compounds such as lignans, justicinol, helioxanthin, (+)-isolariciresinol, docosanoic acid, podophyllotoxin and β -sitosterol- β -D-glucoside, orosunol, 8-demethyl-orsunol [26, 76] have been tested and shown to possess antiviral and antitumor activities. Similarly, the aqueous leaf extract of the plant was reported to inhibit the activity of ACE (by 53%) in an *in vitro* study conducted in KZN (**> Table 2**), suggesting the antihypertensive potential of the plant [18].

Mesembryanthemum species

The genus *Mesembruanthemum* in the family Aizoaceae comprises 25 species endemic to the northern hemisphere and southern Africa. The most common of the 25 species are *Mesembryanthemum crystallinum* (Linn.) and *Mesembryanthemum nodiflorum* (Linn.). The former is a robust succulent herb or dense prostrate shrub of up to 1 m wide. However, in terms of morphology, *nodiflorum* resembles *M. crystallinum*, except smaller in virtually all parts. It is used medicinally to treat bad dreams, heart weakness, HBP, stomach, and other gastrointestinal ailments, swellings, dropsy, and general body pain [26, 34]. The antihypertensive activity of the plant has been investigated following the work of Duncan et al. [34], in which the aqueous leaf and ethanolic stem extracts of the plant inhibited the activity of ACE by 90% and 57%, respectively (**> Table 2**).

Momordica balsamina (L.)

M. balsamina belongs to the Cucurbitaceae family, with balsam pear (English), laloentjie (Afrikaans), intshungu, intshungwana yehlathi (Zulu), and mohodu (Sotho) as its common names. The plant is a climbing perennial herb containing a prostrate or climbing stem (5 m high), and is native to tropical Africa, Asia, Arabia, India, and Australia, but may be found in most provinces in SA (except Western Cape), as well as Namibia, Botswana, and Swaziland among other Africa countries. Besides the fact that most of its parts are consumed as a food source, it is indigenously used to

treat stomach and intestinal ailments [26], as well as diabetes, headaches, boils, sores, HBP, burns, and wounds. [32,77]. The phytochemical screening of various parts of the plant have established the presence of major secondary metabolites such as alkaloids, flavonoids, saponins, terpenes, and steroids attributed to numerous medicinal activities including anti-HIV, hepatoprotective, antiplasmodial, antimicrobial, antidiarrheal, anti-inflammatory, antidiabetic, analgesic, antiseptic, antioxidant, antibacterial, and antihypertensive [77].

Momordica charantia (L.)

M. charantia is a tropical and/or subtropical plant belonging to the Curcurbitaceae family with common names such as bitter melon, bitter gourd, bitter squash, or balsam pear (English). The plant is indigenous to India but is widely distributed in Asia, Africa, and the Caribbean, owing to its edible fruits. It grows up to 5 m in height, bears simple, alternate leaves (4-12 cm) with 3-7 deeply differentiated lobes. Medicinally, it is used as a treatment for stomach ailments, diabetes, cough, respiratory diseases, skin diseases, wounds, ulcers, gout, rheumatism, HBP, menstrual problems, cancer, and HIV/AIDS [78], while its active constituents consist of triterpenes, protein, steroids, polyphenols, alkaloids, lipids, and inorganics [79]. The pharmacological implications of the plant as an antioxidant, antidiabetic, anticancer, antitumor, antiviral, antimicrobial, antigenotoxic, anthelminthic, antimalarial, antineoplastic, antiulcerative, and immunomodulatory were extensively reviewed in an updated report by Gupta et al. [80]. In a South African study, the hypoglycemic and hypotensive effects of M. charantia were presented by Oyewole et al. [75], wherein aqueous extract of the whole plant at concentrations of 50, 100, 200, 400, and 800 mg/kg were found to restore the normal plasma glucose level of streptozotocin-induced diabetic Wistar rats in a dosedependent manner, while similarly reducing the systemic BP and heart rates of Dahl salt-stress hypertensive rats, suggesting its antihyperglycemic and antihypertensive activities (> Table 2).

Ocimum basilicum (L.)

O. basilicum is a characteristic herb ascribed to the Lamiaceae family. Its local names are basil, great basil, Saint-Joseph's wort, royal herb, and king of herbs, and it is native to India. O. basilicum is a perennial herb that grows to a length of 0.5 m and has monoecious flowers (i.e., possessing both male and female reproductive organs). Medicinally, the plant is used for a plethora of ailments such as digestive and nervous system ailments, colic and indigestion, nausea, abdominal cramps, snake bites, skin diseases, gonorrhea, diarrhea, earache, dysentery, itching, malaria, and menstrual dysfunction [81]. The essential oil and other phytochemicals contain linalool, eugenol, α -cubebene, caryophyllene, rosmarinic, estragole, and methyl cinammate [82], which are bioactive as anticonvulsants, hypnotics, antimicrobials, and antifungals, as well as anti-sickling agents [82]. Interestingly, the radical scavenging, antioxidant, antiseptic, insecticidal, and anti-sickling activities of the extracts from O. basilicum are reported in previous studies. Additionally, an updated and exhaustive review of its pharmacological properties such as analgesic, anti-inflammatory, hypoglycemic, hepatoprotective, antihyperlipidemic, antiulcerative, cardioprotective, stimulatory, sedative, hypnotic, anticonvulsant, memory retention, stroke preventive, antimicrobial, antiviral, antimycobacterial, chemo-modulatory, immuno-modulatory, anticancer, anti-osteoporotic, antihypertensive, and antithrombotic, as well as a safety profile of the plant, have been reported by Miraj and Kiani [83].

Olea europaea (L.) subsp. africana (Mill.) P. S. Green

The Oleaceae family houses the genus Olea, of which the species europaea is a member, and has common names such as wild olive (English), olienhout (Afrikaans), mohlware (South Sotho, North Sotho), umnguma (Xhosa, Zulu, and Swati), mutilwari (Venda), and mothware (Tswana). It is a small to medium-sized tree growing between 5 m, and to as tall as 18 m. It is predominant in SA, where it was previously referred to as O. africana and is a smallfruited subspecies of the commercial olive [24]. While the plant is widespread in Eastern Cape, KZN, Western Cape, Mpumalanga, Northern Cape, Limpopo, North West, and Gauteng, it is medicinally used to lower BP [23, 26, 29], enhance renal function, alleviate colic, to act as diuretics, tonics, and antidiarrheals, and to treat sore throats [26]. The phytochemicals include but are not limited to isolated flavonoids, essential oils, flavone glycosides, biphenols, sugars, sterols, triterpenoids, secoiridoids, and phenolic compounds including oleuropein (major), α -pinene, rutin, ursolic acid, oleanoic acid, uvaol, apigenin, and luteolin, which have anticancer, antidiabetic, antimicrobial, antiatherosclerotic, antihypertensive, antioxidant, cardiovascular, antidysrhythmic, and antihyperlipidemic properties [71, 84]. Moreover, the pharmacological investigations on the various parts of the plant from different regions of the world were revealed in a complete review presented by Hashim et al., and included antidiabetic, anticancer, antimicrobial, antioxidant, antihypertensive, cardioprotective, anti-inflammatory, gastroprotective, and neutroprotective activities [84].

Opuntia ficus-indica (Mill)

Indigenous to Mexico, O. ficus-indica, the most common, domesticated, and studied member of the Cactaceae family [85], is locally referred to as Indian fig, mission prickly pear (English) boereturksvy or grootdoringturksvy (Afrikaans), and umthelekisi (Zulu). It is a succulent, branched shrub or tree growing up to 3 m with a sturdy trunk. Besides being widespread in other places such as Australia, southern Europe, Africa (SA, Uganda, and Kenya), and southern, northern, and central America, the plant is ethnobotanically used to cure gonorrhea and HBP by Bapedi traditional healers of the Limpopo Province [31] in SA, as well as diabetes, ulcers, burns, and asthma in different parts of the world. The chemical constituents include but are not limited to sugars (glucose and galactose), flavonoids, phenols, vitamins (ascorbic acid, vitamin A, B, K), and minerals (Ca, Mg, Na, P) from various parts of the plant [86], with compounds such as guercetin, kaempferol, piscidic acid, sinapoyl-glucose, rutin, and catechin [85] isolated to provide hypoglycemic, antiatherosclerotic, and antihyperlipidemic effects [85]. Pharmacologically, the antiulcer, anti-inflammatory, neuroprotective, anticancer, antiviral, antidiabetic, antioxidant, antihyperlipidemic, antihypertensive, antimicrobial, wound healing (cicatrising), antiatherogenic, antihyperinsulinemic, and hepatoprotective effects of the plant have been reviewed and reported by Kaur et al. [86], in addition to other available reviews.

Oxygonum sinuatum (Hochst. & Steud. Ex Meisn.) Dammer

O. sinuatum, otherwise known as Ceratogonum sinuatum among others names, belongs to the Polygonaceae family. It is a less erect, branched herb widely distributed in eastern and southern Africa including South Sudan, Angola, Ethiopia, Uganda, Kenya, Congo, and SA. Irrespective of its acceptance as an edible vegetable in some parts of the continent, it is employed for the treatment of boils, tonsillitis, and fungal infections (eyes, legs). Moreover, its antihypertensive activity has been reported by Ramesar et al., along with that of 16 other Zulu nutritive MPs [18], when 50 microliters (0.1 g/mL) of the aqueous leaf extract inhibited the activity of ACE by 59% (**> Table 2**).

Persea americana (Mill)

The Lauraceae family houses the genus Persea of which the species americana is a member. P. americana is referred to as avocado, with numerous synonyms including but not limited to Persea edulis (Raf.), Persea gigantea (L.O. Williams), and Persea gratissima (C.F. Gaertn.). P. americana, thought to have originated in Mexico and central America, is a fast-growing tree or shrub popular for its edible, green-fleshed fruits. It grows up to 40 m tall, but most are no taller than 20 m. Medicinally, the plant is used to treat ulcers, diabetes, and HTN [31], while the active constituents include but are not limited to terpenoids, glycosides, flavonoids, and coumarin [87]. The pharmacological relevance of the plants as anti-inflammatory, vasorelaxant, analgesic, hypotensive, anticonvulsant, antiviral, wound healing, antiulcer, antihepatotoxic, antioxidant, hypoglycemic, antimalarial, antithrombotic, antihypercholesterolemic, antiarthritic, cardiovascular, haemopoietic, and antihyperlipidemic, as well as its safety profile, have been reported by Yasir et al. [87] and Tcheghebe et al. [88] in their exhaustive reviews. Moreover, in addition to the above-mentioned pharmacological activities, the immunomodulatory, antimicrobial, cytotoxic, and antioxidant effects are also reported.

Physalis viscosa (L.)

P. viscosa is a rhizomatous perennial herb in the Solanaceae family with local names including starhair ground cherry, stellate ground cherry, grape ground cherry (English), and arrebenta-cavalo (Portuguese). Native to South America and widely distributed on other continents, *P. viscosa* is endowed with a hairy stem and grows up to 400 mm in length. Medicinally, it is used for urine suppression, fever, gout, wounds, anemia, and as a diuretic, while the antihypertensive effect is affirmed in a study conducted by Ramesar et al., when the methanolic leaf extract *in vitro* inhibited ACE by 60% [18].

Protorhus longifolia (Bernh. Ex C. Krauss) Engl.

P. longifolia is a tree in the Anacardiaceae family with common names such as red beech, purple currant, red Cape beech (English), rooiboekenhout, rooimelkhout (Afrikaans), ikhubalo, isifuce (Xhosa), umkomiso, and uzintlwa (Zulu). It is a fast-growing, single-stemmed, evergreen garden plant that grows up to 15 m in height, with the ability to withstand frost and drought. *P. longifolia* is indigenous to Swaziland and SA, with major distribution in Eastern Cape, KZN, Limpopo, and Mpumalanga. Traditionally, its use for the treatment of HBP, heartburn, internal bleeding, diarrhea

(both in humans and cattle), dysentery, lightning shock, cramps, and swollen legs has been documented [26,65]. The antiplatelet aggregation, antioxidant, cytotoxic, antimycobacterial and antimicrobial effects of the crude extract are reported [89], while a lanosteryl triterpene isolated from the plant has also shown antihyperlipidemic, antihyperglycemic, anticoagulant, anti-inflammatory, and cardioprotective effects [90]. Additionally, the antihypertensive property of aqueous (64%) and ethanolic (77%) leaf extracts against ACE has also been reported [34].

Psidium guajava (Linn.)

P. quajava is a member of the Myrtaceae family called Koejawel in Afrikaans. It is an evergreen shrub or tree with hairy branchlets growing up to 10 m high. It is endemic to the Caribbean, Central America, and South America, and is used medicinally for inflammation, diabetes, HTN, diarrhea, rheumatism, ulcers, wounds, pain, fever, and lung ailments [31, 91]. The chemical constituents include tannins, essential oils, sterols, flavonoids, phenolics, saponins, triterpenes, carbohydrates, fats, proteins, and vitamins, while the compounds isolated from various parts of the plant, such as α -pinene, limonene, β -pinene, β -sitosterol, ascorbic acid, thiamine, nerolidol, 3-caryophyllene, and so on [91], have been established. Similarly, Gutierrer et al. [91] submitted its antidiarrheal, antimicrobial, anti-inflammatory, antimalarial, antitussive, antioxidant, hepatoprotective, antigenotoxic, antimutagenic, antiallergic, anticancer, cardiovascular, hypotentensive, antihyperglycemic, analgesic, antinociceptive, and wound healing effects, as well as the safety profile of *P. quajava* in a comprehensive review, which has been updated recently by Díaz de Cerio et al. [92]. In another study, Ojewole [93] established the hypoglycemic and hypotensive effects of the plant when an aqueous extract, orally administered caused a dose-dependent (50-800 mg/kg body weight) normoglycemic and antihyperglycemic effect, as well as a significant reduction in systemic arterial BPs and heart rates in streptozotocin-induced diabetic and Dahl salt-stress hypertensive rats.

Ptaeroxylon obliquum (Thunb.) Radlk.

Sneezewood (English), nieshout (Afrikaans), and umThathi (Xhosa, Zulu) are the local given names of *P. obliquum*, a member of the Rutaceae family. Endemic to the southern African countries, including Zimbabwe, Mozambique, and SA, particularly Eastern Cape, Western Cape, Mpumalanga, KZN, and Limpopo, sneezewood trees are endowed with oppositely arranged leaves with 3– 8 leaflets clearly asymmetrical in shape. Medicinally, the plant is used to alleviate headaches, repel moths, cure heart diseases, and for wound healing (in both cattle and humans, particularly after circumcision), myiasis [24], cough, dysentery, malaria, itching, and chest pain, among other uses. Its antibacterial, broadspectrum immune modulation, antioxidant, antifungal, mosquitocidal, cytotoxic, and hypotensive [94] effects have been documented.

Rauvolfia caffra (Sond.)

R. caffra's common names include but are not limited to quinine tree (English), kinaboom (Afrikaans), umJelo (Xhosa), and umHlambamanzi (Zulu). It is a fast-growing, evergreen tree of

about 300 cm in height. In terms of distribution, roughly 60 species of the genus Rauvolfia exist, 7 of them are found predominantly in Africa, 3 in Madagascar, and caffra, which is found primarily in southern Africa, particularly Swaziland, and most provinces in SA (Eastern Cape, KZN, Mpumalanga, Limpopo, Gauteng, and North West). Its indigenous usages are wound healing and to treat coughs, diarrhea, stomach ailments, and HBP [95], while it also possesses alkaloids, terpenoids, flavonoids, and polyphenols as some of its chemical components [96], as well as reserpine, which has been demonstrated to have HBP-lowering effects [97]. Its various pharmacological properties such as antioxidant, antimycobacterial, antimicrobial, molluscicidal, anticancer, anti-inflammatory, antitrypanosomal, and antimalarial, as well as antihypertensive activity when various doses (15, 300, 500 mg/kg bw) of the plant extracts (methanolic, ethyl acetate, and dichloromethane) brought down the systolic and diastolic BP in spontaneously-hypertensive rats [96] are reported.

Rhus chirindensis (Baker) f.

R. chirindensis, otherwise referred to as Searsia chirindensis (Baker f.) Moffett, is an average-sized semi-deciduous tree (about 10 m in height) belonging to the Anacardiaceae family. The plant is native to SA and found along the coastal belt sparing from the Cape through KZN to other parts of southern Africa such as Swaziland, Mozambique, Tanzania, and Zimbabwe. Its common names include red currant (English), bostaaibos (Afrikaans), muvhadelaphanga (Venda), umhlabamvudu (Zulu), and umhlakothi (Xhosa). Traditionally, R. chirindensis has been applied in the treatment of measles, cough, chest pain, syphilis, convulsions, and epilepsy in most rural populations of SA, as well as in the management of HBP [26]. Pharmacologically, the anticonvulsant, analgesic, antiinflammatory, hypoglycemic, antioxidant, cytotoxic, anti-HIV, and antibacterial activities of the plant have been reported [98]. The ethanolic leaf extract of the plant has been confirmed to possess antihypertensive activity when it inhibits the activity of ACE by 85% [34].

Sclerocarya birrea (A.Rich.) Hochst. subsp. *caffra* (Sond.) Kokwaro

This member of the Anacardiaceae family is commonly referred to as jelly plum, cat thorn, cidar tree, marula (English), morula (Sotho), mufula (Tshivenda), mufuna, mupfura, mushomo (Shona), ukanyi (Tsonga), umganu (Ndebele), and maroela (Afrikaans), with distribution in the Miombo woodlands of southern Africa (KZN, Limpopo and Mpumalanga, Zimbabwe, Mozambique, and Ethiopia), the Sudano-Sahelian region of West Africa and Madagascar. It is a medium-large sized deciduous tree that grows as high as 12 m. Besides its major phytochemical components, such as phenolics, saponins, flavonoids, essential oils, and tannins [99], the traditional uses (such as treatment of HTN [26]) of various parts of the plant have been documented in various nations of Africa [99]. The bark is used for dysentery, diarrhea, rheumatism, malaria, hemorrhoids, and to determine the sex of an unborn child. The leaves are used for curing gonorrhea, while the fruits are edible-eaten raw or turned into a delicious jelly or used to make a local alcoholic drink, mukumbi (by the Venda people), which is now a commercialized liquor. Interestingly, numerous

pharmacological properties have been thoroughly reviewed by Ojewole et al. [99], including anticonvulsant, antihypertensive, antibacterial, antifungal, antihelminthic, antiplasmodial, hypoglycemic, analgesic, anti-inflammatory, antidiarrheal, antioxidant, as well as in the treatment of renal and skeletal muscle disorders. The aqueous extract of the stem-bark of the plant orally administered at various concentrations (25–1600 mg/mL) lowered the blood glucose level of diabetic streptozotocin Wistar rats in a dosedependent manner, with 800 mg/mL exhibiting optimal reduction, and reduced the systemic arterial BP as well as the heart rates of hypertensive rats [100, 101].

Stangeria eriopus (Kunze) Baill

A perennial member of the Zamiaceae family, it is locally referred to as Stanger's cycad, natal grass cycad (English), bobbejaakos (Afrikaans), imfingo (Zulu), and umfingwani (Xhosa). S. eriopus is a slow-growing perennial plant consisting of large fern-like leaves (0.25-2.0 m); a swollen, carrot-shaped tuberous root; and an upper stem that reveals branches with up to 10 heads. The plant is also monoecious (male and female reproductive organs existing on separate plants) and coniferous, with its cones giving rise to each point of growth. The male cone is cylindrical in shape and tapers at the tip reaching a diameter of 3-4 cm at maturity (yellowish-brown), while the female cone is 18×8 cm in diameter and is egg-shaped or ellipsoidal to conical with a rounded tip, turning a dark-green color at maturity. It is endemic to the east coast of SA (KZN, Eastern Cape) and southern Mozambique, and is used to treat headaches, internal parasites in cattle, and HTN [24, 26, 34], with macrozamin (methylazoxymethanol glycoside) as its major phytoconstituent [102]. Pharmacologically, the activity of its aqueous leaf extract against ACE has been reported in the work of Duncan et al. [34], establishing its in vitro antihypertensive effect with 55% inhibition.

Tulbaghia violacea Harv.

T. violacea is a member of the Alliaceae family, with common names such as wild garlic (English), wildeknoflok, and wilde knoffel (Afrikaans), is a rapid-growing, bulbous plant extending to 0.5 m in height (25 cm wide). This plant, among the other 29 Tulbaghia species, is endemic to southern Africa particularly SA (Eastern Cape, KZN, Limpopo) and Zimbabwe, with various indigenous applications. It is used as an aphrodisiac, as well as to treat sinus conditions, headaches, cough, colds, asthma, tuberculosis (pulmonary), intestinal worms, HTN, and cancer of the esophagus [24, 27, 29, 30, 33], owing to its numerous pharmacological and biological properties. Besides the fact that it possesses sulfur-containing compounds (e.g., alliin), showing much resemblance to Allium ursinum (Amaryllidaceae) Linn. in odor, the leaves and flowers are consumed as a leafy green vegetable, and it is worth noting that it may also be used as a seasoning. The antioxidant, cytotoxic, antiparasitic, antimicrobial, antibacterial, anthelminthic, antiamoebic, antidiabetic, hypolipidemic, anti-HIV, antithrombotic, androgenic, anticancer, and antihypertensive properties [18, 29, 34, 103] of the plant have been reported, as has the safety profile. In fact, in a study by Raji et al. [104, 105], a methanolic leaf extract of the plant at various concentrations (5, 10, 20, 40, 60, 80, 160 mg/kg b.w.) lowered the BP (systolic, diastolic), mean arterial pressure and heart rate of ageing male normotensive and adult male spontaneously hypertensive rats in a dose-dependent manner, suggesting its antihypertensive effects. Similarly, Ramesar and others [18] reported the inhibition of ACE using aqueous (68%) and methanolic (71%) extracts of the plant in an *in vitro* study. Additionally, Duncan et al. [34] submitted that aqueous extracts of the leaf and bark inhibited ACE by 72% and 49%, respectively, while ethanolic leaf extracts inhibited the activity of the enzyme by 61%.

Turraea floribunda Hochst.

Honeysuckle tree, wild honeysuckle tree (English), kanferfoelieboom (Afrikaans), umdlozana (Swazi), and umadlozane (Zulu) are the common names of T. floribunda, a member of the Meliaceae family. It is a fast-growing, slender, small to medium-sized deciduous tree that grows to between 10–15 m in height and is single or multi-stemmed and loosely branched. Found in rocky areas in woodland and forested ravines, its global distribution includes countries like Congo, Uganda, Mozambique, Kenya, Zimbabwe, Malawi, Tanzania, Swaziland, Uganda, and SA, especially Eastern Cape, KZN. Medicinally, it can be used to manage rheumatism, dropsy, heart disorders, malaria, including its use as a purgative, enema, and emetic (use in trance by the Sangomas) [106]. Besides the presence of limonoids, turraflorins, tetranortriterpenoids, and phytoconstituents [107, 108], pharmacologically, the antitrypanosomal, larvicidal, antioxidant, anti-inflammatory, anticholinesterase, mutagenic, antimalarial, and antibacterial properties of the plant have been reported. Notwithstanding the above properties, an aqueous leaf extract of the plant showed weak inhibition (45%) of ACE in a study conducted by Duncan et al. [34], thus established its weak antihypertensive effect.

Discussion

The use of MPs and or phytotherapy against a variety of human diseases is no longer considered an archaic practice, as its global acceptance continues to rise. This is attributed to their assumed safety, efficacy, cost-effectiveness and minimal side effects [17, 18]. Interestingly, their role in the treatment of CVD, including HTN, have contributed greatly to cardiovascular studies [109], owing to the challenging prevalence and burden of HTN, as well as the ineffectiveness and/or expense of antihypertensive single drug therapy, among others. The information in this review has been gathered from numerous sources and includes over 100 MPs whose various parts (particularly leaves and roots) or whole plants are prepared (mostly by maceration or decoction) and applied on a regular basis for the management of HBP by traditional healers residing, practicing and/or hailing from different tribes (Zulu, Xhosa, Sotho) across the republic of SA. It is noteworthy that 27 of these plants have been identified as having BP-lowering effects through either in vitro or in vivo studies or evaluated with both, despite the evaluation of almost half of all plants reviewed reflecting that 17 of these plants inhibit the activity of ACE using in vitro assays, others have been further confirmed in animal (Dahl-salt stress and or sodium thiopentone-induced) studies as potential antihypertensives. Additionally, most of the assays described and corroborated the use of polar solvents-primarily

water, methanol and ethanol (\triangleright **Table 2**)-by indigenous traditional healers as the most effective medium for extracting active components from plants. Similarly, the detection of saponins, tannins, flavonoids and alkaloids in most of the plants with reported antihypertensive effects suggests that these phytochemicals may be responsible for their BP-reducing effects. Moreover, the isolation of alkaloids having reported antihypertensive effects from these plants-including but not limited to hippane, lycorine (*C. macowanii*) and reserpine (*R. caffra*)-supports the latter submission.

Notwithstanding the *in vitro* evaluations of the plants, it is imperative to submit that such testing cannot be used to confirm the pharmacological activity of a MP unless it is buttressed by an *in vivo* study. It is therefore pertinent that more efforts be made by stakeholders (researchers, scientists, medicinal practitioners, institutes, and pharmaceutical companies) toward confirming the pharmacological effect of these indigenously used antihypertensive MPs. In addition, those that have remained unassessed should be evaluated *in vitro* to investigate their pharmaceutical properties and gain additional insights into their potential pharmaceutical uses, leading to the development of effective, safe, and perhaps novel moieties or formulations to curb the prevalence of HBP.

Toxicity studies provide the assessment of safety profiles for MPs or drug substances (formulations), and may be acute, subacute, chronic, or subchronic. However, despite the pharmacological activities of many MPs and the general assumption that they are safer than conventional medicines in the treatment and/or management of diseases, there are compelling arguments regarding their safety, as some of them are understood to cause illeffects in experimental animals. Intriguingly, plants such as *Aloe ferox, Hoodia gordonii, C. roseus*, and *Artemia afra* have not only established their potential in alleviating HBP, but are also reported to be nontoxic [52, 116], suggesting that they may be studied in clinical trials, thus presenting an excellent prospect for the development of pharmaceuticals to control HTN.

Future studies

Based on our observations in the present review, the following studies are proposed:

- Intensified efforts to subject traditional antihypertensive plants to initial *in vitro* and *in vivo* pharmacological activity screening.
- Isolation and elucidation of bioactive compounds against HTN to confirm that they are responsible for the elicited effects. Thus far, most of the MPs with BP-lowering effects have not been subjected to isolation processes, an area requiring urgent attention.
- The global or general acceptance of MPs as an alternative therapy to orthodox medicine requires that the drug moieties discovered should be able to pass all the rigorous processes of drug development. Hence, there is need for all the plants already demonstrated as safe and effective in reported studies to be subjected to rigorous clinical trials for subsequent drug formulation and eventual commercialization.

In conclusion, HTN is a disease attributed to an increased risk of many chronic complications such as CHD, CCD, stroke, cardiac arrest, renal insufficiency, myocardial infarction, and so on. However, the current management of HBP with synthetic antihypertensives is characterized by various side effects such as dizziness, blurred vision, skin rash, muscle cramps and tiredness, while the use of phytotherapy has continued to receive wider interest and publicity owing to their therapeutic properties and minimal side effects. This review highlights that many documented antihypertensive MPs have not been adequately explored or pharmacologically proven. Hence, there is a need for the government to sponsor additional research to establish the efficacy and safety of these plants as alternative options to curb or manage HBP.

Methodology

Exhaustive literature search was used for this review on Google Scholar and research articles from scientific databases such as Science Direct, PubMed, JSTOR, and Medicine were adopted. Most of the articles used centers between 2000 to 2018 and more than 700 journals, over 20 dissertations/theses, and 5 books were retrieved, with emphasis being placed on the antihypertensive activities of the MPs when keywords such as SA, MPs, HTN, hypotension, HBP, and antihypertensive as well as languages particularly Zulu, Xhosa, Sotho, and Afrikaans were provided to browsing engine. Information on other pharmacological activities of the documented plants aside their antihypertensive properties (► Tables 1 and 2) were sourced from 458 peer-reviewed articles.

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Conflict of Interest

The authors declare no conflicts of interest.

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