Natural Products Modulating Angiotensin Converting Enzyme 2 (ACE2) as Potential COVID-19 Therapies

Murtala Bello Abubakar1,2*, Dawoud Usman1,2, Gaber El-Saber Batiha3*, Natália Cruz-Martins4,6,7, Ibrahim Malami2,7, Kasimu Ghandi Ibrahim1,2, Bilyaminu Abubakar2,8, Muhammad Bashir Bello2,8, Aliyu Muhammad10, Siew Hua Gan11, Aliyu Ibrahim Dabai2,12, M Alblihed13, Arabinda Ghosh14, Reem H. Badr15, Devarajan Thangadurai16 and Mustapha Umar Imam2,17

1Department of Physiology, Faculty of Basic Medical Sciences, College of Health Sciences, Usmanu Danfodiyo University, Sokoto, Nigeria, 2Centre for Advanced Medical Research and Training, Usmanu Danfodiyo University, Sokoto, Nigeria, 3Department of Pharmacology and Therapeutics, Faculty of Veterinary Medicine, Damanhour University, Damanhour, Egypt, 4Faculty of Medicine, University of Porto, Porto, Portugal, 5Institute for Research and Innovation in Health (i3S), University of Porto, Porto, Portugal, 6Laboratory of Neuropsychophysics, Faculty of Psychology and Education Sciences, University of Porto, Porto, Portugal, 7Department of Pharmacognosy and Ethnopharmacy, Faculty of Pharmaceutical Sciences, Usmanu Danfodiyo University, Sokoto, Nigeria, 8Department of Pharmacology and Toxicology, Faculty of Pharmaceutical Sciences, Usmanu Danfodiyo University, Sokoto, Nigeria, 9Department of Veterinary Microbiology, Faculty of Veterinary Medicine, Usmanu Danfodiyo University, Sokoto, Nigeria, 10Department of Biochemistry, Faculty of Life Sciences, Ahmadu Bello University, Zaria, Kaduna State, Nigeria, 11School of Pharmacy, Monash University Malaysia, Bandar Sunway, Malaysia, 12Department of Microbiology, Usmanu Danfodiyo University, Sokoto, Nigeria, 13Department of Microbiology, College of Medicine, Taif University, Taif, Saudi Arabia, 14Microbiology Division, Department of Botany, Gauhati University, Guwahati, India, 15Department of Botany and Microbiology, Faculty of Science, Alexandria University, Alexandria, Egypt, 16Department of Botany, Karnatak University, Dharwad, India, 17Department of Medical Biochemistry, Faculty of Basic Medical Sciences, College of Health Sciences, Usmanu Danfodiyo University, Sokoto, Nigeria

The 2019 coronavirus disease (COVID-19) is a potentially fatal multisystemic infection caused by the severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2). Currently, viable therapeutic options that are cost effective, safe and readily available are desired, but lacking. Nevertheless, the pandemic is noticeably of lesser burden in African and Asian regions, where the use of traditional herbs predominates, with such relationship warranting a closer look at ethnomedicine. From a molecular viewpoint, the interaction of SARS-CoV-2 with angiotensin converting enzyme 2 (ACE2) is the crucial first phase of COVID-19 pathogenesis. Here, we review plants with medicinal properties which may be implicated in mitigation of viral invasion either via direct or indirect modulation of ACE2 activity to ameliorate COVID-19. Selected ethnomedicinal plants containing bioactive compounds which may prevent and mitigate the fusion and entry of the SARS-CoV-2 by modulating ACE2-associated up and downstream events are highlighted. Through further experimentation, these plants could be supported for ethnobotanical use and the phytomedicinal ligands could be potentially developed into single or combination therapies.

Abbreviations: ACE2, Angiotensin converting enzyme two; ADME/T, Absorption, Distribution, Metabolism, Excretion/and Toxicity; DPP4, Dipeptidyl peptidase four; mAb, monoclonal antibody; NA, not available; PK, Pharmacokinetic; S-RBD, Spike glycoprotein receptor binding domain; TMPRSS2, Transmembrane protease serine two; Wi-A, Withaferine-A; Wi-N, Withanolone.
INTRODUCTION

By the end of December 2020, less than a year after it was declared an outbreak, the severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) had infected over 80 million people and claimed almost two million lives (European Center for Disease Prevention and Control, 2020). This places its mortality score over a thousand-fold greater than that of its homolog, the 2002/2003 SARS-CoV (WHO, 2004; de Wit et al., 2016). To date (30th, January, 2021), based on a MEDLINE search, due to its severity, over 90,000 papers (and possibly more) have addressed the 2019 coronavirus disease (COVID-19). COVID-19 develops after SARS-CoV-2 gains entry into host cells by binding to angiotensin converting enzyme 2 (ACE2) receptors (Hoffmann et al., 2020). This crucial first step initiates a cascade of events that results in a vicious cycle of virion to cell infection and replication that includes attachment, penetration, uncoating, translation, replication, assembly and release. The fusion to ACE2 receptor and internalization causes down-regulation of the ACE2 receptor, halting its traditional protective functions over the renin-angiotensin system (RAS) and potentiating the pathophysiological sequelae of COVID-19 (Verdecchia et al., 2020).

In this way, finding solutions to COVID-19 is at the core of solving this global pandemic. At present, available pharmaceuticals face numerous limitations in the treatment of COVID-19. The challenges range from safety and side effects to poor efficacy which warrants the search for better treatment modalities (Gupta and Misra, 2020; Nittari et al., 2020). Like with SARS-CoV, persistent complications, such as lung damage, make preventive strategies the best option (Kuba et al., 2005). In this pursuit, the development of vaccines and therapeutic approaches conceptually linked to ACE2 rank high for preventing COVID-19 (Bourgonje et al., 2020). Vaccines are either undergoing trials (with some concerns) or facing issues of distribution, acceptance or storage, and are a long shot from herd immunity. Meanwhile, in the search for ACE2-centered therapeutics, scientists have uncovered an ACE2 receptor decoy called human recombinant ACE2 (hrsACE2; APN01) (Monteil et al., 2020). These synthetic approaches may prove expensive and may take longer to verify across people and conditions.

Natural products with some degree of biological benefits are regarded as bioactive compounds. Phytochemicals include a broad spectrum of plant bioactive compounds which are majorly proven to be safe. In recent times, these phytochemicals obtained from herbs have gained prominence in ethnomedicine as attractive choices for antiviral therapies (Utomo et al., 2020). Some polyphenols, a subset of phytochemicals, have been suggested to inhibit the fusion and entry of SARS-CoV-2 (Paraiso et al., 2020). So far, reports indicate that most African countries bear lesser COVID-19 burden compared to some Northern and Western nations of the globe (Figure 1). To explain this, several reasons have been proposed, including demography, climate, genetic variations, cross immunity, and antimalarial usage (Lalaoui et al., 2020).

The African and Asian regions are famous for their ethnomedicinal herbs (Figure 2) and a close comparison reveals a previously unseen correlation between countries having no ethnomedicinal policies and a higher burden of

![Figure 1](https://example.com/figure1.png)

**FIGURE 1** | Geographical distribution of COVID-19 infection per 100,000 of population across 14-days. Redrawn from source: European Center for disease Prevention and Control (ECDC). Data for September 26, 2020 (European Center for disease Prevention and Control, 2020).
COVID-19 (compare Figures 1, 2). Ethnomedicinal policies are developed following the demanding use of ethnomedicines. In turn, these policies offer goals and objectives to governments for the development and support of ethnomedicines and may include guideline principles related to safety and efficacy (Oyebode et al., 2016; WHO, 2019). The uniform rise in incidence of COVID-19 in regions of lower ethnomedicinal policies hints to a possible link between ethnomedicinal use and COVID-19 incidence. Despite the number of reasons put forward to explain the relatively mild emergence of COVID-19 in regions like Africa compared to America and Europe (Lalaoui et al., 2020), no involvement of ethnomedicinal use has been suggested until now. Possibly, ethnomedicinal plants contain bioactive compounds that contribute to prevent or mitigate SARS-CoV-2 invasion via interactions with ACE2-mediated viral tropism (fusion and entry).

Given the global burden and challenge posed by the pandemic and the urgent need for the rapid development of efficient, safe, cost-effective, and readily available prevention and treatment modalities, we turned to phytomedicine. Following the spike of converging evidences, we sought to review evidences related to ACE2 that may suggest the use of phytochemical remedies. Here, we also take a closer look at the SARS-CoV-2 and the indirect phytomedicinal targeting of ACE2 mediated tropism. This aims to provide an indirect evidence for the use of ACE2 interactive phytochemicals, singly or combined, for preventing and mitigating COVID-19, its symptoms and clinical comorbidities. Notably, the influence of ethnomedicines on ACE2 modulation is not an exclusive factor in determining the rate of spread of COVID-19, but a highlight that identifies what logical next steps can be developed based on existing research. This will benefit investigators in quest for remedies from phytomedicine.

Morphology and Genomic Organization of SARS-CoV-2
Coronaviruses are taxonomically grouped into the order of Nidovirales, family Coronaviridae and subfamily Orthocoronavirinae. The subfamily is further divided into alpha, beta, delta, and gamma coronavirus genera. The beta coronavirus genus comprises SARS-CoV, MERS-CoV and the recently emerged SARS-CoV-2 (Chen et al., 2020c). Morphologically, all coronaviruses are enveloped with crown-like particles enclosing a positive sense RNA genome (Figure 3). The lipid bilayer envelope which surrounds the viral particle is derived from the host cell membrane, and embeds three structural proteins, namely: 1) membrane (M), 2) envelope (E), and spike (S) glycoproteins. The other viral structural protein, nucleoprotein (N), is internally located and intimately associated with the viral genomic RNA (Kumar et al., 2020b).

The SARS-CoV-2 genomic RNA has approximately 30 kb in size and is made up of 13–15 (12 functional) open reading frames (ORFs) that are arranged in 5′–3′ order of appearance. The first ORF from the 5′ end comprises approximately 67% of the genome and encodes several non-structural proteins. On the other hand, the remaining ORFs encode both accessory and structural proteins, including the major S, E, M, and N proteins (Naqvi et al., 2020). The M and E proteins are required for virus morphogenesis, assembly and budding, while the S glycoprotein is a fusion viral protein comprised of two subunits; S1 and S2. The S1 subunit is made up of signal peptide, N-terminal domain (NTD) and a receptor-binding domain (RBD) (Watanabe et al., 2020). The S2 subunit contains two heptad-repeat regions known as HR-N and HR-C, which form the coiled-coil structures surrounded by the protein ectodomain. Importantly, the S protein in its native...
form is coated with polysaccharide molecules, a property that partly enables SARS-CoV-2 to evade surveillance of the host immune system during entry (Shah et al., 2020).

Pathophysiology of COVID-19

Although coronaviruses target the respiratory tract, they can also affect the gastrointestinal (GI) tract, kidney, central nervous (CNS) and cardiovascular (CV) systems (Peiris et al., 2003; To et al., 2004). Their systemic effects manifest as lesions and multiple organ dysfunctions especially in immunocompromized patients and the elderly with already existing history of chronic ailments like cancer, CNS disorders, and diabetes. Similarly, the organs that are predominantly affected by SARS-CoV-2 express high levels of ACE2, meaning that such organs, due to ACE2, have high affinity to SARS-CoV-2 reception (Letko et al., 2020). The ACE2 can catalyze the hydrolysis of angiotensin II (a vasoconstrictor peptide) into angiotensin (1–7) which is a vasodilator that lowers the blood pressure (Verdecchia et al., 2020). Eventually, the invasion of SARS-CoV-2 deviates the ACE2 functionality, causing increase in blood pressure, disruption of blood vessels, and inflammation of organs (Letko et al., 2020).

Notably, the SARS-CoV-2 preferentially targets the respiratory tract. Alveolar type II cells (AT2) of the lungs express high levels of ACE2, making them the primal targets of SARS-CoV-2 (Verdecchia et al., 2020). In the native alveoli, oxygen is essentially absorbed, whereas carbon (IV) oxide is released. However, due to the presence of SARS-CoV-2, ACE2 is prevented from modulating the protein angiotensin II (ANG II), thereby causing further damage to the blood vessels of the lungs, overall preventing oxygen and carbon (IV) oxide exchange. The resultant effect is difficulty in breathing and improper functioning of the lungs, which worsen the damaging effect to the lung parenchyma, ending up in bronchitis, edema, alveolar collapse and acute respiratory distress syndrome (Huang et al., 2020; Tan et al., 2021).

Should the invasion continue, the virus may gradually infiltrate and infect the circulating immune cells which carry it to other organs like the digestive tract, (i.e. stomach, duodenum, colon, and ileum) displaying inflammatory changes by attacking the epithelial cells of the mucous membrane of ileum and colon (Shi et al., 2005). In a broad view, ACE2 is expressed in abundance within the small intestine. Autopsy under high-resolution electron microscopy has shown that the virus replicates rapidly in intestinal tissue, thereby preventing the ACE2 from carrying out its functions in amino-acid balance, intestinal inflammatory response, homeostasis of the intestinal microbiota and regulatory expression of antimicrobial peptides. Consequently, a secondary infection by bacteria may also affect the weakened intestine, since ACE2 has been hindered from

FIGURE 3 | Schematic illustration of SARS-CoV-2 virion and its genome.
regulating the intestinal microbiota (Wang et al., 2020a). Due to low movement of bowels, an early sign of digestive tract invasion by SARS-CoV-2 could be diarrhea.

Circulating coronavirus in the blood stream can migrate to the heart and affect the cardiac tissue (Hamming et al., 2004). Accordingly, underlying cardiac injury elevates the risk of disease severity in COVID-19 due to heightened risks of myocarditis (To et al., 2004; Chen et al., 2020a). Kidney dysfunction is also a common feature of COVID-19 (Li et al., 2020c), as evidenced by reports of increased blood urea nitrogen, serum creatine and urea (Fan et al., 2020; Yang et al., 2020b). Urinary system failure, genital invasion and possibly death are also possible outcomes (Fan et al., 2020; Zhang et al., 2020b).

The most detrimental of all possibilities is damage to the CNS, when the SARS-CoV-2 nucleic acid contaminates the cerebrospinal fluid of the spinal cord and brain stem. Using the neuronal pathway, i.e., motor and sensory neurons, it streams via the nerves of the olfactory lobe of the nasal cavity to the brain. In fact, the use of motor proteins, kinesins and dynein, prominent to these neuronal pathways is essential for viral migration. The olfactory lobe, a part of the brain that translates smell responses, is the starting point of dysfunction. This partly explains the early signs of inability to smell, i.e. anosmia (Sun and Guan, 2020; Wu et al., 2020). Additionally, since the brain stem controls vital functions including blood pressure, heart beat and reflex actions like respiration, viral perforation of the brain stem eventually results in multiple organ dysfunctions. The development of intracranial infection may result in confusion, headache and epilepsy. Also, more serious complications, such as obstruction of cerebral blood flow, brain swelling, and coma could also occur. These are mostly mediated by interleukins (such as IL-6, IL-12, IL-15) and tumour-necrosis factor (TNF) in the brain, indicating an inflammatory response (Li et al., 2020b; Wu et al., 2020).

Notably, several organs could also deviate from proper functioning, such as the spleen and liver, but those highlighted above are the most susceptible. Furthermore, aside factors such as patient’s age, patient’s immune system, early diagnosis and treatment, the level of ACE2 expression also affects the COVID-19 pathophysiology. These pathophysiologic features of COVID-19 as highlighted, could be quite detrimental and chronic; therefore, necessitating the need for more research into effective therapies for the prevention and/or treatment of COVID-19.

**COVID-19 and ACE2**

One of the key signaling pathways that acts as a homeostatic regulator for the cardiovascular system is the RAS. Irrespective of an individual’s health status, RAS maintains a dynamic control of vascular function. This is achieved through different regulatory components and effector peptides such as the carboxypeptidase, ACE2, which converts angiotensin II to a vasodilator, angiotensin (1–7) through the ACE2/Ang one to seven axis (Donoghue et al., 2000; Tikellis and Thomas, 2012). ACE2 counteracts the effect of ACE/RAS pathway (de Kloet et al., 2010). ACE2 is an integral membrane protein present in the lungs, liver, heart, kidney, and endothelium (Donoghue et al., 2000). More importantly, ACE2 receptors are abundant in the epithelial cells of the surfaces of the nostrils, mouth, and lungs (Chen et al., 2020b). Sequence analysis demonstrated that ACE and ACE2 have over a 40% homology of their respective amino acids (Donoghue et al., 2000). The 40 kb ACE2 gene located on chromosome Xp22 consists of 18 exons and 20 introns and codes for a protein of 805 amino acids (Turner et al., 2002; Marian, 2013).

Unlike other coronaviruses that explore several receptors, like aminopeptidase N and dipeptidyl peptidase four in addition to ACE2 in gaining entry into cells during infection, SARS-CoV-2 exploits only the ACE2 protein for cell entry and subsequent viral replication (Zhou et al., 2020). The N-terminal region of the ACE2 receptor is anchored by the spike glycoprotein (S1) of SARS-CoV-2. Following receptor binding, SARS-CoV-2 employs some of the host proteases for spike protein priming. These include cathepsin L, cathepsin B, trypsin, factor X, elastase, furin, and transmembrane protease serine 2 (TMPRSS2) (Gheblawi et al., 2020). Being the major point-of-entry for SARS-CoV-2, the ACE2 protein expression on respiratory epithelial cell surfaces is critical for the pathogenesis of COVID-19 in individuals exposed to the virus. The vulnerability of the lungs to SARS-CoV-2 infection is premised on two facts; the large surface area of the lungs and the vast expression of ACE2 proteins on the type 2 pneumocytes of the alveoli in the lungs (Zhao et al., 2020). It is also noteworthy to state that the GI tract, especially the small intestine and colon could be critical points of entry since previous studies have demonstrated a high expression of ACE2 in these organs (Hamming et al., 2004; Li et al., 2020a). Also, the multiple organ failure in individuals with complicated COVID-19 could be linked to the vast distribution of ACE2 receptors in these organs (Guan et al., 2020; Huang et al., 2020) indicating the important roles of ACE2.

**ACE2 as a Therapeutic Target**

Though, a focus on the ACE2 activity accompanying COVID-19 may be considered commonplace, it is arguably one of the significant direct/indirect targets for therapeutics in COVID-19. ACE2 physiology is important in hypertension, cardiac, kidney, and lung physiology (Tikellis and Thomas, 2012). The apparent relevance of ACE2 physiology in COVID-19 has come in two ways, both supporting immense clinical focus on its modulation: 1) as a receptor for SARS COV2 (ClinicalTrials.gov identifiers: NCT04335136) and 2) as an enzyme for the generation of ANG 1–7 (ClinicalTrials.gov identifiers: NCT04311177, NCT04312009, NCT04338009, NCT04338009, NCT04394117, and NCT04394117) (Bhalla et al., 2020). The unfortunate neglect of ACE2 therapeutic potential (and lessons learned) following SARS-CoV outbreak strongly necessitated and still advocates for the recent efforts to develop ACE2-centered therapeutics for the current SARS COV2 (Bhalla et al., 2020). Thus, efforts (till date) to find an ACE2-centered therapeutic are re-emerging with a steep focus after its rediscovery as a receptor for SARS COV 2 (Bhalla et al., 2020).

To date, ACE2 receptor is the major known entry point for SARS-CoV-2 into human cells. The virus uses its spike protein in attaching to ACE2 receptor of susceptible cells. This would mean
that a COVID-19 vaccine could be developed based on the spike protein sub-unit of SARS-CoV-2. In this regard, Tan and Colleagues have recently described a new vaccine candidate immunogenic against SARS-CoV-2 spike RBD, with demonstrated stability at ambient temperature; reducing cold-chain dependence (Tan et al., 2021). Alternatively, the development of small molecules or antibodies that are competitive antagonists for ACE2 receptor have been explored (Glasgow et al., 2020). These agents have shown superior neutralizing efficiency to convalescent sera (Glasgow et al., 2020; Tan et al., 2021).

Furthermore, Kuba and colleagues demonstrated that a downregulation of ACE2 protein by SARS-CoV leads to severe lung injury in laboratory mice (Kuba et al., 2005). By implication, this could mean that excessive administration of soluble ACE2 proteins would not only counteract the scenario of cell viral entry but also maintain the physiological activity of ACE2, including protection against lung injury via negative RAS regulation (Imai et al., 2005; Yu et al., 2016). This is the basis for recombinant ACE2 decoys (Monteil et al., 2020). Based on the clinical trials done so far, ACE2 protein that is synthesized through recombinant DNA technology, has shown to be efficacy in healthy and ARDS patients (Haschke et al., 2013; Khan et al., 2017). Recombinant ACE2 sequesters circulating viruses, thereby preventing the interaction between SARS-CoV-2 spike and endogenous ACE2 proteins. This effect allows endogenous ACE2 protein to negatively regulate the RAS, thereby preventing tissue injury as seen in COVID-19 (Zhang et al., 2019). These reports highlight a potential of recombinant ACE2 decoys for treating COVID-19 patients.

Natural Products and ACE2
Natural products have for long been considered as important sources of therapeutic agents. These agents are remarkably low molecular weight molecules capable of eliciting enzyme activities due to their complex nature (Malami et al., 2016). The secondary bioactive agents/plant metabolites, which are able to reduce or wholly inhibit enzyme catalytic activities, can be used as ACE2 modulators. Available evidence from previous studies supports the inhibitory properties of natural products toward ACE2 enzyme activity. Moreover, these natural products are considerably used in traditional medicine and are extensively found as part of the human diet. Here, we describe some of the common natural products that have been previously reported with ACE2 modulatory activity prior to the COVID-19 era (Table 1).

Flavonoids are widely found in a variety of fruits and vegetables, including citrus fruits and tomatoes (Tutunchi et al., 2020). Among the different classes of natural products combined, flavonoids represent the largest group of ACE2 inhibitors. In fact, several studies have previously implicated some flavonoids to possess inhibitory properties against ACE2 activity. For instance, a previous in vivo study showed that apigenin upregulates the expression of the ACE2 gene in spontaneously hypertensive rats (Sui et al., 2010). Wei and co-workers also demonstrated that baicalin, a natural flavone, attenuates angiotensin-II induced endothelial dysfunction by modulating ACE2 expression both at mRNA and protein levels (Wei et al., 2015). Naringenin, a flavanone found in grape fruit, can also attenuate hypertensive reno-vascular damage in vivo by downregulating the ACE2 expression (Wang et al., 2019). These reports highlight a potential of flavonoids in mitigating SARS-CoV-2 viral infection via modulation of ACE2. Notably, as is highlighted later in this review, flavonoids like naringenin, naringin, nobiletin, hesperidin, hesperetin, neo-hesperidin, pinocembrin, quercetin, myricetin, and kaempferol have shown potential ACE2 binding affinity toward the residues that contact S protein of SARS-CoV-2 (Chen and Du, 2020; Cheng et al., 2020; Güler et al., 2020; Ngwa et al., 2020; Omar et al., 2020; Yang et al., 2020a).

Another class of natural products with ACE2 modulatory properties are the steroids and steroid glycosides. Ginsenosides are naturally occurring steroid glycosides commonly found in ginseng rhizome. An in vivo study demonstrated that ginsenoside Rg3, a tetracyclic triterpenoid saponin, induces the upregulation of ACE2 levels and attenuates Ang II-mediated renal injury (Liu et al., 2019). Following the outbreak of SARS-CoV-2, a recent

| Natural products | Bioactive compound | Method of assessment | References |
|------------------|-------------------|---------------------|------------|
| Flavonoids       | Naringenin        | In vivo             | (Sui et al., 2010; Wang et al., 2019) |
|                  | Apigenin          | In vivo             |            |
| Steroids and steroid glycosides | Ginsenoside Rg3 | In vivo             | (Wei et al., 2015) |
| Coumarins        | Osthole           | In vivo             | (Shi et al., 2013; Hao and Liu, 2016) |
| Alkaloids        | Nicotiamine       | In vivo             | (Takahashi et al., 2015) |
|                  | Emodin            | In vitro            | (To et al., 2007) |

- Table 1: Some natural bioactive compounds shown to modulate ACE2 prior to the COVID-19 era.
virtual screening revealed that the steroid, arundoin, and some steroid glycosides (azukisaponin I, 20(S),24(R)-ocotillol and ginsenoside Rg6) can modulate ACE2 activity (Zi et al., 2020). In this study, the bioactive compounds demonstrated promising activity with a lower percentage inhibition based on an ACE2 kinase inhibition assay. Additionally, glycyrrhizin, another steroid glycoside found in the roots Glycyrrhiza glabra L., has been predicted to latch on to ACE2 with an estimated binding energy of −9 kcal mol⁻¹, potentially preventing viral entry (Chen and Du, 2020).

Furthermore, coumarins, which are naturally occurring phenolic compounds, have also shown ACE2 modulatory properties. An in vitro and in vivo study showed that the prenylated coumarin, osthol, mitigates inflammation and acute lung injury in mice by preventing ACE2 and Ang (1–7) down-regulation, resulting in a lowered release of proinflammatory cytokines (TNF-α and IL-6) (Shi et al., 2013). This protective influence of osthol was proposed to occur due to the counter effects ACE2/Ang (1–7) has over the ACE/AngII axis, and was confirmed by an abrogation effect following the use of an ACE2 inhibitor (Shi et al., 2013). In a related study, Hao and Liu demonstrated that osthol attenuates pulmonary fibrosis and inhibits lung inflammation by modulating ACE2 in rat models of bleomycin-induced pulmonary fibrosis (Hao and Liu, 2016).

With regards to alkaloids, a recent study has highlighted that cepharanthine, a naturally occurring alkaloid, can bind to spike protein and inhibit the viral interaction with ACE2 protein in silico (Ohashi et al., 2020). In another study, the non-proteinogenic alkaloid, nicotianamine, exhibited a similar effect on ACE2 (Chen and Du, 2020), which agrees with a previous in vitro demonstration by Takahashi et al. (2015). Also, evidence had shown that a natural anthraquinone, emodin, significantly inhibits the infectivity of S protein-pseudotyped retrovirus to Vero E6 cells by binding to ACE2 protein and blocking the SARS-CoV S protein from binding to ACE2 at a 50% minimum inhibitory concentration (IC₅₀) of 200 μM (Ho et al., 2007).

Functional Foods and ACE2
Apart from providing nutrients and energy, functional foods also modulate the body’s physiological functions (Nicoletti, 2012). The potentially numerous health benefits of functional foods to humans have been explored (Muhammad et al., 2018). Among the several important beneficial effects of functional foods include their potential to prevent and treat COVID-19. Here, we present functional foods that have been shown to modulate ACE2 (Table 2).

Fatty Acids (FA)
Naturally occurring fatty acids have been previously reported to have health benefits in several disease states. Omega-3 FA, fixed oils, oils obtained from nuts, coconut, soy, sesame, and fruits are common sources of naturally occurring FA. Omega-3 FA are a class of polyunsaturated fatty acids (PUFA) that are known to have beneficial effects in the prevention and management of various diseases (Muhammad et al., 2018). The two major biologically active types of omega-3 FA are eicosapentaenoic acid and docosahexaenoic acid (DHA) (Daak et al., 2020). The ACE2 activities of these bioactive agents have been demonstrated in previous studies. Ulu and co-workers have shown that diet rich in omega-3 FA attenuated inflammation in angiotensin-II dependent hypertension by up-regulating the ACE2 activity (Ulu et al., 2013). In another study, Gute. (2011) revealed that omega-3 FA upregulated the expression of ACE2 mRNA in adipocytes (Gute, 2011). As we shall discuss, following the emergence of COVID-19, several essential oils from functional foods have also been identified as ACE2 modulators (Senthil Kumar et al., 2020; Thuy et al., 2020).

Food-Derived Peptides
Previous indications suggest food-derived bioactive peptides can regulate the body’s physiological functions. These food-derived peptides have been demonstrated to exert potential health benefits to humans and thus serve as functional foods. In relation to this, soy proteins isolated from soybeans (Glycine max L.) were demonstrated to have ACE2 inhibitory properties (Table 2). In the first ACE2 inhibitory study using food products, Takahashi et al. (2015) demonstrated the ACE2 inhibitory activity of soybean (Glycine max L.) and its isolated bioactive protein. In their study, the bioactive protein, nicotianamine, intensely inhibited the ACE2 activity at 84 nM concentration. Similarly, bioactive peptides from rapeseed, Leu-Tyr (LY), Arg-Ala-Leu-Pro (RALP) and Gly His-Ser (GHS), have been shown to modulate the ACE2 activity in spontaneously hypertensive rats (He et al., 2019). In this study, ACE2 expression markedly increased following oral administration of the peptides (30 mg/kg) both at the gene and protein levels. A related study had also demonstrated that administration of bioactive peptides (10 mg/kg/day) from blue-green algae, Spirulina platensis (Arthrospira platensis), upregulates the expression of ACE2 in spontaneously hypertensive rats (Zheng et al., 2017). The fact that
these specific peptides modulate ACE2 expression and subsequent activity hints to their bioactivity. However, the pharmacokinetics and mechanisms behind these physiological effects remain largely unknown and may require extensive investigations before becoming practically applicable. Additionally, food components interact with peptides in different ways that may affect the availability of peptides within the matrix (Chakrabarti et al., 2018). As such, to recommend foods rich in ACE2 modulatory peptides, further research is needed to determine the release, stability and mechanisms of bioactive peptides following normal digestion. Furthermore, hydrolysates from pre-digestion when subjected to normal digestion may produce different peptide sets (Chakrabarti et al., 2018). In this case, investigations are needed to demonstrate if the identified physiological effects are abolished, unabosled or changed.

Furthermore, evidences have indicated that the egg-white-derived antihypertensive peptide Ile-Arg-Trp induces the expression of ACE2 and decreases proinflammatory gene expression in mesenteric arteries of spontaneously hypertensive rats (Majumder et al., 2015). A similar in vivo study had further demonstrated that the tripeptide IRW elicits vasorelaxation and abolishes vascular inflammation by increasing the levels of circulatory ACE2 and Ang (1–7), potentiating its protective activity. (Liao et al., 2019). Notably, IRW is less-susceptible to peptide modifications by digestive enzymes (Bejjani and Wu, 2013). Findings from these studies demonstrate the previously established beneficial effects of food-derived proteins and other components in modulating ACE2 activity that subsequently influences the body physiology in a way that could antagonize the development of COVID-19. Though these peptides for intake may be largely acceptable due to natural source, a minimal tissue bioavailability of most peptides limits theirs use as therapeutic agents (Campos et al., 2011). To address this, approaches like microencapsulation, use of enzyme inhibitors, may be needed to achieve effective oral delivery. Further in vivo studies are also needed to understand the immune response to peptides and human trials may become necessary to validate efficacy. However, the production of such synthetic bioactive peptides faces limitations in expense and requires newer low-cost innovations.

### Phytomedicinal Plants, COVID-19 and the Effects on ACE2

At this time, several claims have been made on herbal-based traditional medicines or phytomedicines as remedies to COVID-19. These herbs and products include COVID organsics (Madagascar), claims on Andrographis paniculata (Burm.f.) Nees, Tinospora crispa (L.) Hook. f. and Thomson, and Gymnanthemum amygdalinum (Delile) Sch. Bip. (DR-Congo), turmeric (Curcuma longa L.), a Sri Lankan herbal drink, and some Tanzanian claims about recipes from lemon grass, ginger, and neem leaves. Notably, like synthetic agents for COVID-19 such as remdesivir, dexamethasone and chloroquine, some of these herbs and products are recently being supported by preliminary evidence and have been suggested for further empirical assessments (Borkotoky and Banerjee, 2020; Enmoozhi et al., 2020; Rakib et al., 2020). With a focus on identifying and supporting the development of plant products having some level of scientific evidence in the context of ACE2-centered SARS-CoV-2 fusion-entry, we describe three classes of phytomedicinal agents, based on the target used to modulate viral tropism (virion fusion and entry) along the Spike-RBD/TMPRSS2/ACE2 axis. These phytomedicinal agents include those that may 1) interfere with host cell surface proteins (ACE-2 receptor and TMRSS2 protease), 2) interface with viral spike glycoprotein, or 3) interrupt spike-RBD/ACE2 interaction (Figure 4).

#### The Spike-RBD/TMPRSS2/ACE2 Axis

The SARS-CoV-2 fusion and entry, which is characteristic of COVID-19, marks a phytomedicinal target for COVID-19 therapy. Following SARS-CoV-2 spike-RBD attachment to the host cell ACE2 receptor, the membrane bound activator protease, TMRSS2, cleaves the viral spike protein and facilitates its fusion with the host cell membrane (Hoffmann et al., 2020). This constitutes the Spike-RBD/TMPRSS2/ACE2 axis and the cleavage (or priming) that occurs at the junction between subunit one and two of the spike protein precedes internalization. The Spike-RBD/TMPRSS2/ACE2 axis presents several interventional targets for natural products at the 1) receptor binding motif (RBM) of the receptor binding domain (RBD) of the spike glycoprotein, 2) ACE2 receptor active site that recognizes the RBD and 4) TMPRSS2 protease (Figure 5).

#### Interference With ACE2 Activity–ACE2

Several plant products have shown potential in interfering with the activity of ACE2 (Chen and Du, 2020; Joshi et al., 2020; Poochi et al., 2020). For example, the essential oils from Allium sativum L (garlic), Ammoides pusilla (Brot.) Breistr., Melaleuca cajuputi Maton and Sm. ex R. Powell (cajeput) and Piper spp (pepper) like Piper nigrum L. and Piper retrofractum Vahl, can potentially latch onto ACE2 and competitively inhibit viral binding and entry (Abdelli et al., 2020; Gutierrez-Villagomez et al., 2020; My et al., 2020; Thuy et al., 2020). These inhibitory effects have been attributed to the presence of certain ligands (Table 3). For instance, Senthil Kumar et al. (2020) demonstrated that essential oils from geranium and lemon prove to be effective against ACE2 by downregulating its mRNA and protein expressions in epithelial cells. Further gas chromatography mass spectrometry (GC-MS) analysis identified citronellol, geraniol and neryl acetate in geranium oils, and limonene in lemon oils as the major ligands (particularly citronellol and limonene) responsible for the downregulation of ACE2 expression in HT-29 cells. Same ligands were also identified to inhibit TMPRSS2 (Senthil Kumar et al., 2020). Notably, not all ligands that bind with ACE2 inhibit its enzymatic activity (Williamson and Kerimi, 2020). The fact that some of these essential oils bear several molecules with binding capacity hints to some possible synergistic effect that may account for the overall efficacy of essential oils. Interestingly, some of these interactive ligands can be synthesized (Gutekunst et al., 2012; Liu et al., etc.)
indicating a vastly unexplored potential. Additionally, in a novel attempt, Rattanapisit and co-workers used a plant expression system (*Nicotiana benthamiana* Domin) to develop and characterize the rapid production of recombinant RBD as a biopharmaceutical capable of neutralizing viral entry by specifically binding to ACE2 (Rattanapisit et al., 2020). Such use of plants as platforms for recombinant protein production, offers benefit over animal-based platforms in terms of cost and is also advantaged in its flexibility, rapid scalability and safety. The recombinant RBD possibly exhibit in vivo neutralization activity similar to recently developed capsid-like particle-based vaccines against SARS-CoV-2 invasion (Fougeroux et al., 2021), and may represent another turning point for COVID-19 therapeutics.

### Interference With ACE-2 Activity - ACE2 and Angiotensin

ACE2 is known to counter the vasoconstrictive and other detrimental effects of the Angiotensin-I/ACE/Angiotensin-II/AT-1 Receptor axis which is a component of the RAS. ACE2 converts angiotensin-1 and -II into angiotensin one to seven, which acts on Mas receptors. This shunt represents the renin angiotensin system suppressive axis called “ACE2/ Ang 1–7/Mas receptor axis,” which produces vasodilatory and other beneficial effects that represent the protective effect of ACE2 on the body (Furuhashi et al., 2020). As stated earlier, the fusion of SARS-CoV-2 to ACE2 and its subsequent entry downregulates the levels of ACE2 thereby potentiating the detrimental activity of RAS. This makes the RAS an important marker to look at when testing possible COVID-19 therapies. Currently, a Tomeka® Prevention Trial (TPT) is underway in DR Congo to assess the effect of using Tomeka® regimen for COVID-19 on markers of the RAS such as angiotensin-II and -(1–7) (ClinicalTrials.gov identifier: NCT04537585). Tomeka® is an herbal mixture made from soy (from soybean; *Glycine max* L.), sorghum (*Sorghum bicolor* L.), maze (*Zea mays* L.) and mushrooms (*Agaricus bisporus* L.). Nicotianamine had been highlighted as the bioactive phytoconstituent of soy responsible for its potent inhibition of ACE2 (Figure 5) and it was hence, called “soybean ACE2 inhibitor - ACE2iSB” (Chen and Du, 2020).

### Interface With the Viral Spike Glycoprotein and Its RBD

For COVID-19, it is more logical to find phytomedicinal agents that specifically target the interaction of RBD at the Spike/ACE2 complex and not particular to ACE2. This caution
is because agents solely targeting ACE2 may interfere with ACE2 enzyme activity and its complex protective functions (Kai and Kai, 2020). Natural products with significant RBD binding affinity compared to ACE2 can dominate over ACE2 in latching onto RBD, thereby blocking viral attachment, fusion and entry.

Preliminary data based on molecular docking analyses predict several natural plant products as RBD interface binding compounds (Chikhale et al., 2020a; Chikhale et al., 2020b; Sinha et al., 2020a; Sinha et al., 2020b; Table 3). Most data come from “conceivable therapeutics” through in silico molecular analysis of available structural macromolecules. An advantage to the preliminary in silico screening of substances is that pharmacokinetic properties, such as safety could be easily established, shifting priority to the in vitro confirmation of efficacy. Thus, preliminary evidences necessitate further in vitro and in vivo investigations. Notably, Wang and co-workers have recently identified AXL, a transmembrane tyrosine-protein kinase, as a supportive pulmonary entry receptor for the SARS-CoV-2 which may also interact with the spike glycoprotein, but at its NTD region (Wang et al., 2021). This represents another potential target for plant based therapeutic investigations.

Interrupting the Spike-RBD/ACE2 Interaction

Studies suggest that 1) resveratrol from grape skins, 2) rippladib from the leaf extracts of the native bryoni, and 3) withanolone from Indian ginseng, can selectively bind and inhibit both RBD and ACE2 (Alexpandi et al., 2020; Balkrishna et al., 2020; Wahedi et al., 2020). These ligands can potentially inhibit the Spike-RBD/TMPRSS2/ACE2 axis simultaneously at the RBD and ACE2. Moreover, multiple ligand activity on the said axis indicates a higher likelihood of getting some potent alternative for COVID-19. Ho et al. (2007), in an in vitro study, established that emodin, from Rheum sp. and Polygonium sp. abolish the interaction between 2002/2003 SARS-CoV spike protein and ACE2 in a dose-dependent manner. Given the level of homology between the two coronaviruses, an investigation into the influence of emodin against the interaction of the SARS-CoV-2 spike and ACE2 is necessary.

Notably, the components of some plants may differ based on the plant part used (Kumar et al., 2020a). For instance, the Citrus spp. peel represents the most abundant store of hesperidin, another possibly potent inhibitor of the RBD and ACE2 interaction (Utomo et al., 2020). Consequently, the hesperidin ligand is currently scheduled for a randomized double-blind controlled trial in Egypt (ClinicalTrials.gov identifier: NCT04452799) (Haggag et al., 2020; Okasha, 2020). Knowledge of such should guide research and may afford some better protection against SARS-CoV-2 invasion.

Interfering With the Host Membrane Enzymes; TMPRSS2, Furin and ADAM-17

The protease cleavage site between the S1/S2 subunits of the spike protein bears an arginine-arginine-alanine-arginine (RRAR) sequence (Wang et al., 2020b) (as shown in Figure 4). What
### TABLE 3 | Plant derivatives with potential effect on the Spike-RBD/TMPRSS2/ACE2 axis.

| Target | Plant (bioactive phytoconstituent/product) | Study type | Efficacious dose(s) | Biological test | Mechanistic effect | Findings | References | Comment |
|--------|------------------------------------------|------------|---------------------|-----------------|-------------------|----------|------------|---------|
| Interfere with ACE2 activity | Nicotiana benthamiana domin (recombinant RBD) | In silico and In vitro (vero E6 cells) | NA | Recombinant mAb RBD production in a plant expression system. Neutralization efficiency against positive sera | Gene expression profile (mRNA and protein) | Downregulates the expression of ACE2 and TMPRSS2 | Suggests binding to the SARS-CoV-2 receptor and its neutralization | Rattanapisit et al. (2020) | Suggested consideration of this plant derived recombinant RBD for the development of vaccines and viral detection/diagnostic reagents |
| Pelargonium graveolens L’Hér. (Citronellol) and Citrus × limon (L.) osbeck (Limonene) | In vitro (HT-29 cell line) | 50 μg/ml geranium oil and 25 μg/ml lemon oil | NA | Molecular docking and in silico ADME predictions methods | Docking to ACE2 | Significant inhibition of ACE2 and TMPRSS2 in epithelial cells to protect against SARS-CoV-2 invasion | Senthi Kumar et al. (2020) Citronellol and limonene were the most potent of eight ACE2 inhibitory oil extracts and had dose dependent effects |
| Momordica dioica roxb. ex wild (catechin, quercetin, hederagenin and oleanolic acid) | In silico and In vitro | NA | Molecular docking and molecular dynamics | Docking to ACE2 | | | Sakshi et al. (2021) Constituent flavonoids have better affinities than standard remdesivir, favipiravir and hydroxychloroquine |
| Valeriana jatamansi jones ex roxb. (Hesperidin), Oroxylum indicum (L.) kurz (chrysin), Rheum australe D.Don (emodin) | In silico | NA | Molecular docking and molecular dynamics | Docking to ACE2 | | | Basu et al. (2020) Spike inhibitory capacity similar to that of docked chloroquine and hydroxychloroquine |
| Artemisia absinthium L. (anabsinthin, absinthin, dicafeoylquinic acids), Syzygium aromaticum (L.) merr. and L.M.Perry (3-0-cafeoylquinic),Phaseolus vulgaris L. (quercetin 3-glucuronide-7-glucoside, quercetin 3-vicianoside, isosakuranetin 7-O-neohesperidoside) and Inula helenium L. (Quercetin-7-O-galactoside, 3,5-dicafeoylquinic acid, 3,4,5-tricafeoylquinic acid) | In silico | NA | Molecular modeling/ docking and dynamic simulations | High affinity binding to pocket of the active site of ACE2 | | | Joshi et al. (2020) Compounds demonstrated good intestinal and brain permeability. Also showed no carcinogenic tendency |
| Allium sativum L. (diallyl tetrasulfide and | In silico | N/A | Molecular modeling/ docking | Binds to ACE2 receptor | | | Thuy et al. (2020) | Ligands are the two most potent of seventeen inhibitors of SARS-CoV-2 |

(Continued on following page)
| Target | Plant (bioactive phytoconstituent/product) | Study type | Efficacious dose(s) | Biological test | Mechanistic effect | Findings | References | Comment |
|---------|------------------------------------------|------------|---------------------|-----------------|--------------------|----------|------------|---------|
| trisulfi de, 2-propenyl propyll Piper sp. (pipercyclobutanamide B, a and nigramide Q) like Piper nigrum L. and Piper retrofractum vahl | In silico NA | Molecular modeling/docking and dynamic simulations | Docks closely to active site of ACE2 | These dimeric piperamides of essential oil could possibly inhibit ACE2 mediated entry of SARS CoV2 | | Gutierrez-Vilagomez et al. (2020) | ACE2 gotten from essential oil of plant. Exhibit potential drug likeness based on ADME. Pipercyclobutanamide B (most potent) docked along duct to ACE2 active site. | |
| Ipomoea obscura (L.) ker gawl. (Urso-deoxycholic acid) | In silico N/A | Molecular modeling/docking | Bind to ACE2 receptor | Could inhibit viral entry and infectivity | | Poochi et al. (2020) | This is the most potent of five possibly bioactive ACE2 inhibitors from ethanolic extract of the plant. | |
| Ammoides verticillata (dist.) briq. -From Algeria (isothymol) | In silico NA | Molecular modeling/docking | Binds to ACE2 receptor | High affinity and can inhibit ACE2 better than captopril and chloroquine drugs | | Abdelli et al. (2020) | Suggest oil is one of the richest natural sources of isothymol. Good ADMET. | |
| Melaleuca cajuputi maton & sm. ex R. Powell (terpineol, guaiol and linalool) | In silico NA | Molecular modeling/docking and dynamic simulations | Latch to the active site of ACE2 | Lots of convergence points and the inhibitory intensity of these compounds on ACE2 could prevent viral invasion | | My et al. (2020) | Out of ten inhibitory substances, these three have the most potent effect on ACE2. Guaiol is also present in guaiacum and cypress pine oils. | |
| Scutellaria baicalensis georgi (baicalin), Erigeron breviscapus (vaniot) hand-mazz. (Scutellari), Citrus × aurantium L. and Citrus reticulata blanco (hesperetin), Liquorice, Glycyrrhiza uralensis fisch. ex DC. (glycyrrhizin) and soybean; Glycine max (L.) merr. (Nicotianamine) Gancao; Glycyrrhiza spp. and chaihu; Bupleurum spp. (glaucisperin F and isorhamnatin) | In silico NA | Molecular modeling/docking and dynamic simulations | Latch to the active site of ACE2 | Potential to bind ACE2 and hinder viral entry | | Chen and Du (2020) | Nicotianamine is “soybean ACE2 inhibitor” (ACE2iSB). | |
| Interface with the viral spike glycoprotein and its RBD | In silico NA | Molecular docking | Bind to ACE2 Latch onto site 1 and site 2 of ACE2 | | | Ren et al. (2020) | Suggest the ligands, glyasperin F and isorhamnetin, account for strong binding affinity. | |
| Liquorice; Glycyrrhiza glabra L. (glycyrrhizic acid) | In silico NA | Molecular modeling/docking and dynamic simulations | Bind to cavity of prefusion spike glycoprotein | High binding affinity to spike protein may block viral fusion to ACE2. May inhibit viral entry by interfering with virion-receptor binding and | | Sinha et al. (2020a) | High protein-ligand stability. Most potent of six interactive ligands. | |
| Bupleurum spp., Heteromorpha spp. and Scrophularia scorodonia L. | In silico NA | Molecular modeling/docking and dynamic simulations | Binds to RBD and cleavage site of the spike glycoprotein | | | Sinha et al. (2020b) | Out of 23 saikosaponins, had the most potent latching affinity to active site of | |

(Continued on following page)
| Target | Plant (bioactive phytoconstituent/product) | Study type | Efficacious dose(s) | Biological test | Mechanistic effect | Findings | References | Comment |
|--------|------------------------------------------|------------|--------------------|-----------------|-------------------|----------|------------|---------|
| (saikosaponins U and V) | Indian ginseng: Withania somnifera (L.) dunal (withanoside X and quercetin glucoside) | *In silico* | NA | Molecular modeling/docking and dynamic simulations | Binds receptor binding domain of prefusion spike protein from SARS-CoV-2 to block viral fusion | Protease cleavage, favourable interaction with receptor binding motif (RBM) of RBD | Chikhale et al. (2020a) | Spike glycoprotein (i.e. RBD) Ligands were potent (out of 17) inhibitors of SARS-CoV-2 spike glycoprotein |
| Asparagus racemosus wild. (Asparoside D and C) | | *In silico* | NA | Molecular modeling/docking and dynamic simulations | Bind to spike RBD | Good affinity and stable docking of spike RBD | Chikhale et al. (2020b) | Higher binding affinity than remdesivir (standard drug) |
| Interrupting the spike-RBD/ACE2 interaction | Withania somnifera (L.) dunal (withanone) | *In silico* | NA | Molecular modeling/docking | Interrupts at the junction between ACE2 receptor and viral S-RBD | Decreased binding free energies, destabilized salt bridges, hence blocks and weakens SARS CoV2 entry and infectivity | Balkrishna et al. (2020) | Suggest plant as first choice herb in curbing COVID-19 |
| Diplocyclops palmatus (L.) leaf extract (ripladib) | | *In silico* | NA | Molecular modeling/docking | Interrupts at the RBD of Spike-ACE2 complex | Predicted strong binding at RBD interface to block Spike-ACE2 interactions and viral entry | Alexpandi et al. (2020) | ADMET analysis indicate good pharmacokinetic properties |
| Citrus spp. (hesperidin) | | *In silico* | NA | Molecular modeling/docking | Binds to both RBD and ACE2 receptor | The flavonoid can possibly interrupt RBD/ACE2 interface to abrogate entry | Utomo et al. (2020) | The peel of Citrus sp. represents the most abundant methoxy flavonoid (hesperidin) store of the plant |
| Grape skin: Vitis vinifera L. (resveratrol) | | *In silico* | NA | Molecular modeling/docking and dynamic simulations | Bind tightly and interfere with viral S protein/ACE2 receptor complex | Highly stable binding and selectivity to viral protein/ACE2 receptor complex. Disrupts the spike protein | Wahedi et al. (2020) | Most potent of four stilbene-based natural compounds |
| Interfere with the host membrane protease | Aframomum melegueta K.Schum. (quercetin, apigenin) | *In silico* and *In vitro* | 30, 10, 3, 1, 0.3 mg/L. From fruit (with seed) | Docking, *In vitro* inhibition of recombinant soluble human furin. Immuno-blotting Molecular modeling/docking | Disrupts spike glycoprotein/receptor interaction by inhibiting furin cleavage | Metabolites inhibited furin dependent pre-glycoprotein processing by possibly blocking furin recognition site | Omotuyi et al. (2020) | Suggested bioactive influence of flavonoids (like quercetin, apigenin). Good ADMET sores |
| Withania somnifera (L.) dunal (withanone and Withaferine-A) | | *In silico* | NA | Molecular modeling/docking | Binds to TMPRSS2 catalytic site (Wi-N > Wi-A), alters allosteric site and also downregulates | Predicted multiple action in blocking SARS CoV2 cell entry and propagation by | Kumar et al. (2020c) | Possible drug-able agents for prevention and therapeutics |
TABLE 3 (Continued) Plant derivatives with potential effect on the Spike-RBD/TMPRSS2/ACE2 axis.

| Target                  | Plant (bioactive phytoconstituent/product) | Study type | Efficacious dose(s) | Biological test | Mechanistic effect | Findings | References | Comment |
|-------------------------|-------------------------------------------|------------|---------------------|------------------|-------------------|----------|------------|---------|
| TMPRSS2 transcription   | inhibiting TMPPRSS2                        |            |                     |                  |                   |          |            |         |

*These compounds were found to be the most potent of several others.

Abbreviations: ACE2, Angiotensin converting enzyme two; ADME/T, Absorption, Distribution, Metabolism, Excretion/and Toxicity; DPP4, Dipeptidyl peptidase four; mAb, monoclonal antibody; NA, not available; PK, Pharmacokinetic; S-RBD, Spike glycoprotein receptor binding domain; TMPRSS2, Transmembrane protease serine two; WI-A, Withaferine-A; WI-N, Withanone.

is now known is that furin, a proprotein convertase enzyme, performs cumulative role with TMPRSS2 by pre-activating this site, aiding viral entry (Bestle et al., 2020; Shang et al., 2020). One molecular simulation study suggests that saikosaponins interact significantly with the arginine residues of the cleavage site to prevent S1/S2 protein cleavage, hindering the virion-cell fusion and viral entry (Sinha et al., 2020b). Indeed, the furin pre-activation of spike protein facilitates viral entry particularly in cells with low levels of expressed TMPRSS2 (Lippi et al., 2020; Shang et al., 2020). Omotuyi et al. (2020) demonstrated in a conformational and in vitro study, that processed fruit (with seed) from *Aframomum melegueta* K.Schum., an African resource, and its secondary metabolites mitigates SARS-CoV-2 entry by inhibiting furin. Indeed, some proteinaceous plant products (peptides) and metabolites (flavonoids such as rutin, naringin, methylviosperdin and baikalin), have exhibited anti-protease potencies and have been implicated for disease treatment (Majumdar et al., 2010; Hel linger and Gruber, 2019). In addition, ADAM-17, a membrane metallopeptidase opposes viral entry by cleaving off the extracellular domain of ACE2 receptor. The ACE2 domain that is shed off retains some enzyme activity and can convert angiotensin II to angiotensin 1–7 (Palau et al., 2020; Williamson and Kerimi, 2020). Hence, a closer look into the ability of plant-based ligands to upregulate ADAM-17 represents a potential angle for exploration, which may achieve satisfactory outcomes.

SAFETY

Phytomedicinal agents investigated against the Spike-RBD/TMPRSS2/ACE2 axis have displayed beneficial pharmacokinetic properties when subjected to in silico analysis of Absorption, Distribution, Metabolism, Excretion and Toxicity (ADMET) (Abdelli et al., 2020; Alexpandi et al., 2020; Gutierrez-Villagomez et al., 2020). These preliminary findings hint to their safety and drug-like potential. However, in view of the limitation of virtual screening, further in vitro and in vivo experiments are needed to verify results and provide experimental basis for the research and development/promotion of natural antiviral drugs/nutrition. The cytotoxic status of a plant is also an important safety concern. An in vitro study showed that geranium and lemon essential oils inhibited the expression of ACE2 and TMPRSS2 at mRNA and protein levels with no cytotoxic effects on human cell (Senthil Kumar et al., 2020). Additionally, most plant products highlighted are very promising agents keeping in view their pre-reported biological benefits. Thus, further experiments are warranted based on previous research findings and the reported biological safety of these natural compounds. Moreover, these natural compounds may be synthesized and made readily available for immediate testing.

Formalizing the Use of COVID-19 Phytomedicines

Following the wide use of phytomedicines as alternatives for preventing and treating COVID-19 in Africa, the Africa Center for disease Control and Prevention (Africa CDC) recommended the development of state-based herbal registries. The mobilization of funds, technical research support and proper communication for the development of such proclaimed remedies have also been emphasized (Africa CDC, 2020). These activities can help guide the establishment of botanical assets in the management of COVID-19.

Similar to what was introduced during the 2002/2003 SARS-COV outbreak, a Chinese traditional medicine program challenged its experts with the development of preventive and treatment strategies for COVID-19. A similar global effort to introduce scientifically backed herbal remedies to combat COVID-19 alongside conventional treatment will prove helpful. Though these herbs are promising options in decreasing virion entry and the course of infection, caution must be established and tests must be conducted to avoid toxic and non-ef ficacious regimens. Undeniably, clinically proven therapeutics remain the gold standard for treatment. As such, efforts to engage preliminary and pre-clinically supported herbs of minimal side effects and cost with randomized controlled trials should be considered.

CONCLUSIONS AND FUTURE PERSPECTIVES

Natural products are of prominence and have been adopted for traditional use across many African and Asian countries. Aside policies and a bundle of other factors, the relatively lesser COVID-19 burden observed in regions of Africa and Asia could be influenced by certain potentiators of self-care such
as cost of medication, interest, beliefs, dissatisfaction in synthetic medications and taking responsibility of one’s health, which push toward ethnobotanical use for COVID-19; a scenario where “the kitchen cabinet becomes thy medicine cabinet.” Of concern, the undeniably increasing demand for plant products is occurring globally at a time threatened by plant extinctions; particularly of wild plants due to excess harvest, habitat degradation, specie invasion, low growth, low abundance, and/or susceptibility to disease. This coupled with the fact that at least one potential drug candidate is lost biennially emphasizes the need for conservation and resource management strategies (Chen et al., 2016). This is most possible given the current state of biotechnology and must be considered particularly in developing countries where about 80% of the populace adopt herbal products for basic healthcare (Moyo et al., 2015; Chen et al., 2016). The overall indicators show that a number of plants and their products possess multimodal influence on viral tropism. Beside the use of plant decoctions, some phytoconstituents previously highlighted, indicate potentially mitigating effects against viral entry by directly or indirectly modulating the ACE2 activity at the Spike-RBD/TMPRSS2/ACE2 axis. This perspective has implications for traditional medicine (TM) and we believe will support and inform ongoing and upcoming investigations on the management of COVID-19. More so, some of the highlighted ligands have been tested in the past and documented to be beneficial across comorbidities which may accompany COVID-19. Perhaps, this lightens the need for further studies before clinical trials. Other benefits including availability, affordability and safety, support the development of these phytomedicinal therapies; a step toward attaining the third united nations sustainable developmental goal - “good health and wellbeing for all.”

**AUTHOR CONTRIBUTIONS**

All authors contributed equally to this work. All have read and agree with the submission of this manuscript.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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