Review Article

Paving Plant-Food-Derived Bioactives as Effective Therapeutic Agents in Autism Spectrum Disorder


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Autism spectrum disorder (ASD) is a neurodevelopmental disorder, where social and communication deficits and repetitive behaviors are present. Plant-derived bioactives have shown promising results in the treatment of autism. In this sense, this review is aimed at providing a careful view on the use of plant-derived bioactive molecules for the treatment of autism. Among the plethora of bioactives, curcumin, luteolin, and resveratrol have revealed excellent neuroprotective effects and can be effectively used in the treatment of neuropsychological disorders. However, the number of clinical trials is limited, and none of them have been approved for the treatment of autism or autism-related disorder. Further clinical studies are needed to effectively assess the real potential of such bioactive molecules.

1. Introduction

Autism spectrum disorder (ASD) is a set of behavioral and neurodevelopmental diseases featuring social and communication deficits accompanied by increased repetitive and/or restrictive behaviors [1]. The most commonly registered behaviors under ASD are presented in Figure 1. In the last years, the prevalence of ASD has dramatically risen [2], and as per the World Health Organization (WHO), it is estimated that 1 in 160 children in the world has ASD [3]. Although the etiology and pathogenesis of this disorder are not fully understood, several environmental and genetic factors have been proposed as mediators, thus, limiting key molecular mediators’ identification as well as possible neurological and biochemical mechanisms. However, recent evidences are pointing to alterations in hormones, amino acids, and several biochemical markers as possible mediators in autistic individuals [4, 5]. Furthermore, unifying etiology involving immune dysfunction, abnormal lipid metabolism, glutamatergic dysfunction, and raised susceptibility to oxidative stress has been proposed for ASD [6], with studies increasingly linking mitochondrial dysfunction to ASD [1, 7, 8]. Indeed, mitochondrial dysfunction and oxidative metabolism defects are characteristic of many chronic illnesses, such as bipolar disorder, multiple sclerosis, Parkinson’s disease (PD), schizophrenia, depression, ASD, and chronic fatigue syndrome [9]. Since mitochondria is an integral part of many cell processes, it becomes susceptible to many insults that could affect its integrity. Furthermore, evidences are pointing at immune dysregulation in ASD etiology [9]. Several studies have revealed the presence of immune abnormalities in those suffering from autism [10, 11], with markers of autoimmunity, abnormal cell immunity, aberrant expression of cytokines, and other soluble immunity mediators being evident in ASD children [10–13].

Nonetheless, faced by data scarcity on ASD etiology and underlying causes, the ability to develop and mobilize effective treatments is still limited. Hence, efforts have been done on alleviating only comorbid manifestations of the disorder [1]. However, as a result of the limited treatment options available to improve ASD symptoms, financial challenges, and drugs side effects, dietary and nutritional approaches are becoming popular components of ASD management [14, 15]. In this sense, the present review is aimed at providing a brief overview on ADS and related pathophysiology, as well as on the promissory therapeutic abilities evidenced by plant-food bioactives.

2. Autism Spectrum Disease: A Brief Overview

ASD is an increasingly frequent disorder found among children. To date, several risk factors (prenatal, perinatal, and postnatal) that are either genetic or environmental have been stated [16]. Approximately, 15% cases of ASD are linked to genetic disorders (fragile X syndrome, neurofibromatosis, tuberous sclerosis, and Rett syndrome) [17], and various genetic alterations, such as de novo mutations in coding regions, copy number variations, and chromosomal alterations, are the most frequent variants that have been associated with ASD development [18, 19]. The significance of genetic factors in ASD development could be easily explained in twin studies, with results from these studies indicating that the ASD concordance rates could be around 30–99% in monozygotic twins, 0–65% in dizygotic twins, and 3–30% in siblings, respectively [20–23], while a recent study showed a lower ASD concordance rate in monozygotic twins (50%) [24].

However, some evidences have indicated controversial results in terms of environmental factors and ASD development. These studies suggest that environmental factors, such as vaccination, advanced parental age, maternal smoking, pregnancy and birth complications, thimerosal exposure, deficiency of vitamin D, and reproductive technologies have strongly correlated with ASD [16, 24, 25], but few studies showed no relation with ASD risk [24]. On the other hand, few studies also suggested that both genetic and environmental factors act in a synergistic manner, and epigenetic modifications could be mediators in the gene/environment interface [26, 27]. Various mechanisms have been proposed in such way, including DNA methylation, MECP2 mutation, folate-methionine pathway enzymes, histone acetylation, and chromosome remodeling for tracking of epigenetic changes [28].

Additionally, while few research groups reported that oxidative stress has been involved in ASD [29], also genetic abnormalities in glutathione (GSH) metabolism may be related to altered mitochondrial function [30]. Indeed, GSH, the key intramitochondrial reactive oxygen species (ROS) neutralizer, is decreased in ASD [31, 32]. Also, changes in glutamate metabolism have been implicated in ASD, with glutamate being postulated as a likely cause of mitochondrial dysfunction and selective Purkinje neuron degeneration stated in ASD [8].

3. From the Historical Perspective to Current Clinical Practice

Autism was first identified by Kanner [33] as an “inborn autistic disturbance of affective contact.” Then, many researchers tried to identify the definition and diagnostic criteria of autism [34]. The criteria for the diagnosis of childhood autism was established in 1993 [35]. Lobotomy was one of the initial ASD treatments [36, 37]. Psychotherapy
and holding therapy [39] were also practiced. The first medical drug that was approved to be used in autism was risperidone [40].

During the last decade, the frequency rate of the disease has been progressively rising [41]. Neurological dysfunctions such as autism substitute for a high impact in societies via the world. Though the indications resulting from those illnesses are popular, the mechanisms and reasons are composite and rely on multiple parameters [42]. ASD contains multiple dysfunctions with different stages of verbal skills, logical functioning, and several genetic etiologies [43].

Neurobiological systems being important for social functioning are debatably the most hopeful signaling pathways for ASD biomarker and therapeutic target detection. Two such nominees are the arginine vasopressin (AVP) and oxytocin (OXT) signaling pathways [44].

Apart from the earlier clinical studies, traditional medicine has been commonly used in ASD treatment [45]. Nevertheless, the negative effects of these medications or their adverse interaction between other medications are questionable, and the parents of the autistic children receiving treatment should be given information about the risks [46].

Japanese traditional herbal medicine, Kami-shoyo-san and Yokukansan, can be helpful in improving behavioral problems in autism [47, 48]. Moreover, Ayurvedic, Siddha, and homeopathic treatments are commonly used in India [49]. For instance, Panchagavya gritha was suggested as an effective Ayurvedic drug for autism [50]. In addition, acupuncture is also widely used in China [51, 52]. Positive effects of electroacupuncture on children with ASD were reported [53, 54].

It was suggested in several studies that ASD can be associated with nutritional and gastrointestinal problems, such as food selectivity, nutritional deficiencies, food allergies, intolerances, diarrhea, and constipation [55–59]. Thus, dietary approaches, including ketogenic diets, gluten-free and casein-free diets, high-fat diets, probiotic use, food additives, camel milk consumption, and other diets, have been frequently investigated [57]. Multivitamin/mineral supplementations are also frequently prescribed [60]. Some food components seem to cause both behavioral and gastrointestinal symptoms in children having ASD [61]. Autoimmunity may rise out of enhanced immune response to potential cross reactivity and dietary proteins to proteins in the brain or gut. Remarkably, children on specific protein-restricted diets display lower stages of activated underlying lamina propria lymphocytes (CD3+TNF-α+) known to be rich in immune and colonic intraepithelial lymphocytes compared to children on unrestricted diets [62].

When it comes to digestive capacity, if it is impaired in a subgroup of ASD children, another treatment option is the use of digestive enzyme supplementation, precisely with a full panel of protease enzymes, despite the limited number of researches carried out to assess the effectiveness of digestive enzyme supplementation on behavioral and gastrointestinal conditions [63].

On the one hand, although probiotic therapy has been advised in several reviews as a potential cure for children with ASD and gastrointestinal indications [64], it has been
explained that as of 2009, only one-fifth of physicians were inspired to use probiotics in children with ASD [65]. One research showed that *L. acidophilus* management (5 × 10⁹ CFU/day, twice a day for 2 months) expressively decreased a marker of offensive pathogenic candidiasis in children with ASD and gastrointestinal indications [66], although half of the children in the researches were on limited diets and the research design did not have a control group [66]. In 2014, Rossignol and Frye [67] systematically reviewed a large number of studies on Alzheimer’s medications in ASD including donepezil, galantamine, rivastigmine, tacrine, and memantine. The results of some medications showed encouraging evidence for effectiveness against treating core and associated ASD symptoms, but clinical trials are also needed to confirm their efficacy for treating ASD individuals.

4. Bioactive Molecules: An Upcoming Key in Autism Spectrum Disease

Even though ASD is a lifelong neurodevelopmental disorder with no cure, there are some pharmacological treatments available to suppress symptoms, like irritability and suppression, and to treat other psychiatric problems that accompany ASD, such as depression, bipolar disorder, and anxiety [68]. Several researchers have reported that ASD is associated with nutritional disorders. Thus, the importance of diet and food consumption is huge in ASD treatment [69, 70]. Certain bioactive molecules have been suggested to support the treatment of autism. The effects of plant-based bioactive materials on autism are summarized in the following subsections.

4.1. The Role of Bioactive Molecules in Autism Spectrum Disease. Bioactive molecules that have neuroprotective effects can be used as natural agents in the treatment of neuropsychological disorders, such as ASD, depression, and bipolar disorder [42, 71, 72]. Curcumin, luteolin, and resveratrol are the frequently investigated plant-based active components (Figure 2).

Curcumin, which is a bioactive compound of turmeric (*Curcuma longa* L.), has the potential to be used in the treatment of neuropsychiatric disorders including autism [71]. Even though clinical trials are yet to come, there are a couple of promising studies on rodents. Bhandari and Kuhad reported that after curcumin treatment at a daily dose of 50, 100, and 200 mg/kg for 4 weeks in rats with propionic acid- (PPA-) induced autism, curcumin restored the core and associated symptoms of autistic phenotype by suppressing oxidative-nitrosative stress, mitochondrial dysfunction, and TNF-α and MMP-9 in PPA-induced autism in rats [73]. Curcumin could be developed as a potential pharmacotherapeutic adjuvant for ASD. In addition, Al-Askar et al. [74] reported the postnatal therapeutic role of curcumin in improving most of the impaired parameters in valproic acid- (VPA-) induced rodent models with persistent autistic features.

It was reported that luteolin can reduce maternal immune activation-induced neural abnormalities, such as ASD [75]. In addition, Asadi and Theocharides [56] reported that it can be utilized in ASD treatment, as luteolin inhibited mast-cell activation due to allergy augmentation and mitochondria stimulation. Moreover, dietary luteolin supplementation resulted in the reduction of serum interleukin-6 and tumor necrosis factor levels in children with ASD [76]. Dietary supplementation (1 capsule per 10 kg body weight) containing luteolin (100 mg/capsule), quercetin (70 mg/capsule), and rutin (30 mg/capsule) in olive kernel oil for 26 weeks enhanced adaptive functioning and behavioral disorders in children with ASD [77]. Moreover, Theocharides et al. [78] reported that a dietary supplement, NeuroProtek®, that contains luteolin, quercetin, and rutin, improved symptoms related to gut and brain inflammation in children (aged 4–12 y) with ASD.

Resveratrol has shown therapeutic and anti-inflammatory effects on neurological disorders [79]. For instance, prenatal resveratrol treatment of rats that were also given VPA during

![Figure 2: The chemical structure of some potential phytochemicals for ASD treatment.](https://example.com/figure2.png)
the prenatal stage to induce autism, with daily 3.6 mg/kg injections for 13 days, resulted in enhancement in behavioral changes with low interaction between resveratrol and VPA [80]. In another study, the restoration of the autism-related dysfunctions, such as neurological, behavioral, sensorial, biochemical, and molecular changes, were reported in rats with PPA-induced autism after daily oral resveratrol treatment of 5, 10, and 15 mg/kg for 4 weeks [81].

Korean red ginseng supplementation improved social interaction of mice with autism-like disorder [82]. Gonzales et al. [83] reported an improvement in autism-related behavioral problems with no effects on motor coordination ability due to a daily Korean red ginseng oral dose of 100 or 200 mg/kg in mice with VPA-induced autism (prenatal exposure).

Niederhofer [84] suggested that Ginkgo biloba can be added to the treatment of autism patients, as they observed improvement on the “aberrant behavior” and symptom checklist after Ginkgo biloba treatment of three patients. However, Ginkgo biloba extract was used in adjunction to standard autism medication, risperidone, for treatment of autistic children, and no significant effect was observed [85].

Isothiocyanates such as sulforaphane that is found in high amounts in broccoli sprouts has been found to ameliorate ADS symptoms [72]. Moreover, capsules of sulforaphane-rich broccoli sprout extracts (50 μmol/capsule) were given to male patients with ASD (aged 13-30 y) in doses according to body weight (one capsule for <100 lb, two capsules for 101–199 lb, and three capsules for >150 lb), and improvement in ASD-related problems, such as oxidative stress, weakened GSH synthesis, lower mitochondrial function, decreased oxidative phosphorylation, lipid peroxidation, and neuroinflammation, were reported [86].

All these studies show that some natural plant-bioactive compounds can be promising tools in the control of ASD (Table 1).

Other formulations of bioactives can also be used in the treatment of autistic disorders. For instance, folinic acid...
several studies [99] activity, irritability, repetition, self-injury, and social with-
behavioral disorders (hyperactivity, irritability, sleep prob-
memantine [113], which were approved to treat Alzheimer
drugs and food supplements at the same time, and the
Made several other drugs are being investigated and new
drugs are also being developed by scientists to treat
Antipsychotic agents, such as risperidone and aripipra-
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4.2. Drifting from Natural to Synthetic Drugs. Plant-bioactive
molecules demonstrated promising results in ASD treatment.
However, the number of clinical trials is limited, and none of
them have been approved for ASD or ASD-related disorder
treatment. Thus, bioactive compounds and vitamin/mineral
supplements are used as an adjunct to the synthetic drugs
used to treat autism. The medications of autism generally
target behavioral problems, developmental disorders, physio-
logical destructions, and other cooccurring psychiatric and
medical conditions, such as sleep and gastrointestinal prob-
lems, hypertension, and diabetes [96, 97].

Antipsychotic agents, such as risperidone and aripipra-
zo, are commonly used for treatment of ADS. DeVane
et al. [98] reported that risperidone or aripiprazole treatments
improved the Aberrant Behavior, Checklist-Irritability scores,
behavior, and weight gain of patients with autistic disorder.
The positive effects of risperidone on several problems associ-
at with autism, such as Aberrant Behavior Checklist scores,
and behavioral problems (e.g., aggression, explosivity, hyper-
activity, irritability, repetition, self-injury, and social with-
drawal) with a side effect of weight gain, were reported in
several studies [99–103]. Aripiprazole also gave similar results
in the treatment of ASD patients, such as improvements in
behavioral disorders (hyperactivity, irritability, sleep prob-
lems, etc.) often with side effects, such as weight gain, and
higher risk for sedation and tremor [104–107].

Although risperidone or aripiprazole are the only medi-
cations approved by the US Food and Drug Administration
for ASD treatment [98, 108, 109], there have been several
studies that some drugs developed to treat other mental dis-
orders can be used in the treatment of ASD [67]. For
instance, donepezil [110, 111], galantamine [112], and
memantine [113], which were approved to treat Alzheimer’s
disease, were reported to improve autistic disorders, such as
behavioral, communication, and social impairments. For
instance, guanfacine developed for hypertension treatment
was found effective in improving disorders in people with
pervasive developmental disorders (PDD), such as ASD
[114]. However, there have been several studies showing that
metformin can be helpful in controlling weight gain in auto-
sic patients [115–118].

Several other drugs are being investigated and new
drugs are also being developed by scientists to treat
autism. For instance, a single dose of suramin (20 mg/kg)
improved behavioral problems due to autism and schizo-
phrenia in mice models [119]. In addition, Naviaux et al.
[120] reported that 20 mg/kg intravenous infusion (one
dose) of suramin resulted in improvements in autism syn-
dromes such as behavior and language in autistic children.
Ong et al. [121] reviewed the use of phospholipase A2 inhib-
itors in the treatment of neurological disorders including
autism.

Methylphenidate [122, 123], sertraline [124, 125], ato-
moxetine [126–128], and alpha agonists [129] were also
reported to have potential to be used in autism treatment,
even though they may have some side effects, such as mood
change, irritability, and gastrointestinal disturbance.

Apart from central drugs such as the antidepressants or
antipsychotics mentioned above, drugs such as propranolol,
memantine, d-cycloserine, and oxytocin can be used [130].
One of the most important is propranolol, a beta-
adrenergic antagonist that inhibits anger in higher doses
improving various neuropsychiatric disorders [131, 132].
Trials have shown that propranolol improves altered emo-
tional states that occur with anxiety, aggressiveness, self-
arm, and hypersexuality [133].

Furthermore, a dysregulation in the hypothalamico-
pituitary-adrenal axis and consequently in cortisol levels
has been observed among people with autism spectrum dis-
order (ASD) [134]. Therefore, the regulation of steroid sig-
naling has been suggested as a potential therapeutic route
for the treatment of ASD and other disorders of the central
nervous system [135].

Food selectivity and picky eating habits in patients with
ASD are common, and the patient may also suffer from
nutrient deficiencies even though the deficiency itself ele-
vates the symptoms. Thus, food supplements are generally
viewed as essential elements of ASD treatment. As vitamin
and mineral deficiencies are common in ASD patients [55,
136–139], vitamin-mineral supplements are frequently
investigated [140, 141]. Improvements in several ASD
symptoms due to vitamin and mineral supplementations
were reported [60, 142–145]. On the other hand, Stewart
et al. [146] suggested that dietary supplementation, includ-
ing vitamin and mineral supplements, should be done care-
fully, as they may not provide sufficient amounts of the
deficient nutrient or may cause excessive intake. On the
other hand, ASD patients are often treated with several
drugs and food supplements at the same time, and the
interactions between them are not well known and can be
undesired [147]. In addition, further comprehensive
investigations on the side effects of these medications are
still required.
5. Plant-Food-Derived Bioactive Studies in Autism Spectrum Disease

5.1. In Vitro Studies. Mitochondrial function is critical to CNS as evident in its role in brain energy turn-over [148] and maintenance of ionic gradients critical to neurotransmission and plasticity [149], and its involvement in neural stem cell proliferation, differentiation, and maturation as well as formation of dendritic processes, developmental and synaptic plasticity, and cell survival and death [1, 150, 151] reinforces the importance of mitochondrial dysfunction in ASD etiology. Several phytochemicals have been revealed to protect or restore mitochondrial function and chief among them are the polyphenols with potent antioxidant properties.

Quercetin, a polyphenol broadly distributed in many plants and vegetables, exhibits strong antioxidant properties and could prevent oxidative stress. Research has shown that quercetin and epigallocatechin-3-gallate are bioaccumulated in the mitochondria in its active form, and this could be responsible for their mitochondria protective effect via ROS-scavenging mechanism in vivo [152, 153]. Studies have revealed the modulatory effect of polyphenols such as resveratrol, quercetin, and hydroxytyrosol on the mitochondrial biogenesis process via the stimulation of several coactivators and transcription factors such as the proliferator-activated receptor coactivator-1α (PGC-1α) [153]. Furthermore, several polyphenols have been shown to activate silent information regulation 2 homolog 1 (SIRT1) in vitro; thus, they have been increasingly searched as potential inducers of mitochondrial biogenesis through PGC-1α deacetylation-mediated activation [154]. Recently, resveratrol has been reported to stimulate the SIRT1/PGC-1α-dependent effect on mitochondrial biogenesis in cultured endothelial cells [155]. Furthermore, resveratrol has been revealed to positively influence mitochondrial performance in C2C12 cells via evoking AMP-dependent protein (AMPK) activation and increased mitochondria biogenesis [156]. Sulforaphane (isothiocyanate) preserved mitochondrial functions in ischemia- or toxin-induced damages in normal noncancerous cells [157] and also stimulated mitochondrial biogenesis [158]. Quercetin, rutin, and resveratrol have been shown to prevent the ATP drop and indomethacin-induced alteration in mitochondrial membrane potential in Caco-2 cells [159].

Glutamate dysregulation and toxicity have been observed in ASD, and there is ample evidence that phytochemicals could counteract the effect of altered glutamate metabolism. Polyphenols from green tea have been shown to attenuate glutamate excitotoxicity via antioxidative and antiapoptotic pathways in cultured cortical neurons [160]. In 2003, Lee et al. revealed the protective effect of polyphenols baicain, baicalin, and wogonin isolated from Scutellaria baicalensis Georgi against glutamate/glucose-induced neurotoxicity in primary cultured rat central neurons via an increase in the cell viability, attenuation of increased intracellular calcium ions and nitric oxide production with baicain being the most effective [161]. Furthermore, chlorofucofuroeckol isolated from brown algae species improves glutamate-induced neurotoxicity through modulation of oxidative stress-mediated mitochondrial dysfunction in PC12 cells [162].

All these in vitro studies with plant-food-derived bioactives in autism spectrum disease are summarized in Table 2.

5.2. In Vivo Studies. Given the rapidly growing disabilities triggered by ASD, characterized by social deficits, communication impairment, and cognitive flexibility deficits, McKinnell et al. evaluated in 2021 the changes in rodent behavior (social and anxiety) and cognitive flexibility in the VPA model of autism and control. The results indicated that VPA rats showed loss in performing the set-shifting task. In other words, females with ASD displayed unique behavioral profiling compared to males with ASD [163]. In 2021, Rebolledo-Solleiro et al. examined the effect of bisphenol A (BPA) on behavior, neurodevelopment, and neurodegeneration through a systematic review. As their main findings, the authors underlined that BPA modulates the normal functioning of the reproductive system, metabolism, and brain functions, while triggering the development of few neurodevelopmental disorders including ASD [164].

Similarly, an increasing amount of naturally occurring bioactive compounds have been used for ASD management. Among them, Camellia sinensis (green tea) is an important dietary source of polyphenols, specifically flavonoids, with catechins, such as epigallocatechin-3-gallate, epigallocatechin, epicatechin-3-gallate, and epicatechin being the dominant ones. However, the presence of gallic, chlorogenic, and caffeic acids and flavonol derivatives, like kaempferol, myricetin, and quercetin have been stated [165, 166]. Significant improvement in behavioral assessments and neuroprotection and lowered oxidative stress were observed in autistic mice treated with flavonoid extract from green tea [166]. Furthermore, histological findings revealed the presence of a distinct Purkinje layer and cells after treatment with green tea, thus suggesting its neuroprotective effect [166]. The positive effect of plant extracts (green tea and black pepper) in the management of autistic behavior in rat models is illustrated in Figure 3.

Also, piperine, the major alkaloid found in Piper longum L. and Piper nigrum L. (black pepper), has been reported to possess antioxidant, neuroprotective, anxiolytic, and cognition-enhancing effects [167–169]. A previous study has revealed ameliorative effects of piperine on behavioral alterations and oxidative stress markers in ASD murine model as evident by improved/restored motor deficits and decreased reorientation time, due to its capability to mitigate sodium valproate-induced cerebellar damage [170]. Piperine treatment also brought on cerebellum integrity restoration via a decrease in Purkinje cell number, which is connected with the cerebral cortex and the limbic system [170]. For these properties and anti-inflammatory, antioxidant, and neuroprotective effects, resveratrol could also be relevant in ASD treatment [171, 172].

A study in an experimental murine model revealed that resveratrol improves social skills in valproic acid-induced autistic rats. It regulates and activates sirtuins, members of the class-III histone deacetylases, and also exerts neuroimmunomodulatory effects by regulating transcription factor signaling, decreasing proinflammatory molecules (IL-6 and TNF-α) on dopaminergic neurons, inhibiting NF-κB...
activation, and suppressing T cells [80, 173]. Another study reported that resveratrol suppresses neuroinflammation, mitochondrial dysfunction, oxidative/nitrosative stress, and TNF-α expression in propanoic acid-induced autistic rats [81], thus making it a potential therapeutic agent to ameliorate ASD’s neurobehavioral, and biochemical changes [81].

The impact of turmeric on neurodegenerative diseases and neuropsychiatric disorders has also been documented. Curcumin (diferuloyl methane), the major curcuminoid present in turmeric, a relatively nontoxic and permeable compound to the blood-brain barrier [174, 175], has been reported to have positive effects on the treatment of autistic rats as it targets several cell signaling pathways. For instance, it increases intracellular GSH levels, reduces inflammatory components, and mitigates mitochondrial dysfunction and oxidative/nitrosative stress as well as protein aggregation [73, 176]. This study has also shown that curcumin can alleviate autistic phenotype-associated symptoms via suppressing oxidative-nitrosative stress and mitochondrial dysfunction in propanoic acid-induced autistic rats [73]. In addition, curcumin is able to ameliorate delayed brain maturation and brain toxicity in an autistic animal model via restoration of altered neurological, behavioral, biochemical, and molecular changes related to ASD phenotype [73, 74].

Bacosides are medicinal substances widely used by Indian tribes and are the main bioactive compounds extracted from

<table>
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<tr>
<td>Quercetin</td>
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<td>Mitochondria protective effect</td>
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<tr>
<td>Resveratrol</td>
<td>Endothelial cells</td>
<td>Stimulate SIRT1/PGC-1α and increase of mitochondria biogenesis</td>
<td>[155]</td>
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<tr>
<td>Resveratrol</td>
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<td>AMPK activation and increase of mitochondria biogenesis</td>
<td>[156]</td>
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<tr>
<td>Sulforaphane</td>
<td>Ischemia induced in normal noncancerous cells</td>
<td>Preserve mitochondrial functions and increase of its biogenesis</td>
<td>[157, 158]</td>
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<tr>
<td>Quercetin, rutin, and resveratrol</td>
<td>Indomethacin-induced Caco-2 cells</td>
<td>Prevent the ATP</td>
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<td>Green tea polyphenols</td>
<td>Cortical neurons</td>
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<td>[160]</td>
</tr>
<tr>
<td>Baicalin, baicalein, and wogonin</td>
<td>Primary culture rat central neurons</td>
<td>Antioxidative properties, increased cell viability, reduced intracellular calcium ions and nitric oxide production</td>
<td>[161]</td>
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<tr>
<td>isolated from S. baicalensis</td>
<td>Glutamate-induced toxicity in PC12 cells</td>
<td>Improvement of mitochondrial dysfunction</td>
<td>[162]</td>
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Figure 3: Positive effect of plant extracts (green tea and black pepper) in the management of autistic behavior in rat model.
Bacopa monnieri (L.) Wettst [177]. This plant is traditionally known for its intellect- and cognition-improving properties as well as being a nerve tonic [178, 179]. The pharmacological properties of B. monnieri have been attributed to its constituent alkaloids, saponins, and sterols [180]. B. monnieri has significantly improved behavioral alterations, decreased oxidative stress markers, reduced pain threshold, and normalized locomotor deficiencies as well as anxiety in a murine model of autism. The improvement in locomotive activity was attributed to the anxiolytic properties of B. monnieri and its ability to decrease accumulated glutamate and restore cerebellum architecture [181].

Long-time, daily oral administration of ginsenoside-rich extract has been found to improve social interaction, repetitive behaviors, locomotor activity, and other ASD-related behaviors in mice models [83]. Thus, ginsenosides may be viewed as a possible drug candidate for ASD-associated phenotypes and symptom management as well as neurobehavioral deficits. Ginsenosides are Korean red ginseng (Ginkgo biloba L.) phytoconstituents known for their therapeutic properties, such as improvement in cerebral blood flow, stimulation of neuronal plasticity, learning, memory, and cognition improvement as well as CNS-associated disease treatment [182–184]. In addition, this plant has been reported to possess antistress, neuroprotective, anti-inflammatory, and antioxidant properties [85, 185].

Furthermore, the effect of the ultramicroized lipid molecule, N-palmitoylethanolamide (PEA) with luteolin (co-ultra-PEA-LUT), in an ASD murine model revealed a reduction in proinflammatory molecules, such as nitrotyrosine and nuclear factor kappa B (NF-κB), an improvement in neuroplasticity and neurogenesis, and the modulation of the apoptotic mechanism in several brain regions (cerebellum and hippocampus) following treatment with co-ultra-PEA-LUT [186]. Another study evaluated PEA on the autistic behavior of BTBR T+tf/J mice and shed light on the contributing mechanisms [187]. PEA improved the behavioral phenotype of BTBR mice dependent on PPAR-α activation. PEA restored the hippocampal BDNF signaling pathway and mitochondrial dysfunction and reduced the general inflammatory status of the mice, reducing the expression of proinflammatory cytokines at the hippocampal, serum, and colonic levels. It also improved intestinal permeability by increasing the expression of the tight junctions of the colon and the composition of the intestinal microbiota.

All these in vivo studies with plant-food-derived bioactivities in ASD are summarized in Table 3.

<table>
<thead>
<tr>
<th>Phytocompounds</th>
<th>In vivo model</th>
<th>Results</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bisphenol A</td>
<td>Different ASD animal models</td>
<td>Modulates the function of the reproductive system, metabolism, and brain functions</td>
<td>[164]</td>
</tr>
<tr>
<td>Green tea</td>
<td>ASD mice model</td>
<td>Neuroprotective and antioxidant properties, and improvement of behavior</td>
<td>[166]</td>
</tr>
<tr>
<td>Piperine</td>
<td>ASD murine model</td>
<td>Antioxidant, neuroprotective, anxiolytic, and cognition-enhancing effects</td>
<td>[167–169]</td>
</tr>
<tr>
<td>Resveratrol</td>
<td>VPA-induced ASD rats</td>
<td>Activates sirtuins and decreases IL-6, TNF-α, NF-κB, and T cells</td>
<td>[80, 173]</td>
</tr>
<tr>
<td>Resveratrol</td>
<td>Propanoic acid-induced ASD rats</td>
<td>Reduces neuroinflammation, mitochondrial dysfunction, and oxidative/nitrosative stress</td>
<td>[81]</td>
</tr>
<tr>
<td>Curcumin</td>
<td>ASD rats</td>
<td>Increases GSH levels and reduces inflammation, mitochondrial dysfunction, and oxidative/nitrosative stress</td>
<td>[73, 176]</td>
</tr>
<tr>
<td>B. monnieri</td>
<td>ASD murine model</td>
<td>Improvement of behavior and antioxidant, anxiolytic, and analgesic properties</td>
<td>[181]</td>
</tr>
<tr>
<td>Ginsenoside-rich extract</td>
<td>ASD mice model</td>
<td>Improvement of behavior and locomotor activity</td>
<td>[83]</td>
</tr>
<tr>
<td>Co-ultra-PEA-LUT</td>
<td>ASD murine model</td>
<td>Reduces proinflammatory markers (nitrotyrosine and NF-κB), improves neuroplasticity and neurogenesis, and modulates apoptosis</td>
<td>[186]</td>
</tr>
<tr>
<td>PEA</td>
<td>BTBR T+tf/J mice</td>
<td>Improvement of behavior, PPAR-α activation, Microbiota-gut-brain axis regulation</td>
<td>[187]</td>
</tr>
</tbody>
</table>

5.3. Clinical Studies. Given the huge impact of ASD on the health, wellbeing, and quality of life of patients and facing the increasing evidences of the role of diet in both prevention and management of several diseases, Hartman and Patel reviewed a large number of reports on food approaches to the ASD management and their relation with gastrointestinal, behavioral, neurological, and immune functions through food supplements (fatty acid and pro- and prebiotics, vitamins, minerals, phytochemicals, and hormones) [188]. As their main statements, gastrointestinal issues were correlated with a number of behavioral and neurological deficits, with food approaches being able to improve the lives of patients with ASD [188]. In another study, Gogou and Kolios
reported on the impact of food administration during pregnancy on the risk of ASD offspring. In this study, authors included both clinical and experimental studies, and food supplement (i.e., folic acid, iron, vitamins, choline, vitamin D, and docosahexaenoic acid) studies were also included [189]. Food supplements including choline, folic acid, and multivitamins had a significant impact on the expression of ASD-related genes, while iron had no effect. Thus, a suitable amount of multivitamins, vitamin D, and docosahexaenoic acid can help in reducing the risk of ASD in offspring [189]. Infante et al. reported a case study of a 23-year-old young adult male with ASD. Omega-3 and vitamin D combination therapy showed beneficial effects to the patient [190]. Also, Sivamaruthi et al. assessed the role of the microbiome, food supplements, and probiotics in the development of ASD and underlined that the maternal diet and lifestyle are greatly associated with the development of ASD [191].

In 2017, Wink et al. described the long-term impact of metformin on antipsychotic-associated weight gain in youth with and without ASD using brief ASD mealtime behavior inventory (BAMBI) and the behavior pediatric feeding assessment scale [118]. The authors stated that metformin treatment stabilized BMI z-score, and BAMBI demonstrated good consistency, test-retest reliability, and criterion-related reliability. In 2001, Ahearn et al. reported on the feeding behavior in children (ages 3–14 years; exposed to 12 food items in 6 sessions) with ASD and pervasive developmental disorder-not otherwise specified (PDD-NOS) [192]. Different parameters, including food acceptance, expulsion, and disruptive behavior were recorded on a trial-by-trial basis, with data obtained clearly indicating that some children were sensitive to food texture or category, while others were indifferent.

In addition, and when looking at clinical evidence reporting the use of plant-derived bioactive molecules on ASD, the anti-inflammatory, antioxidant, antiallergy, and neuroprotective effects of different biomolecules have been well documented. Mostafavi and Gaitanis analyzed a data set including preclinical and clinical data regarding the use of cannabis and cannabidiol in the treatment of ASD [193]. The results of the analysis suggested that both compounds revealed promising therapeutic benefits in some persons with ASD.

In the case of specific compounds, luteolin and quercetin extracted from Chamomile sp. and Sophora sp. leaves were used in children with ASD, and marked improvements in ASD symptoms were stated. Most of the patients (75%) reported a significant improvement in gastrointestinal features, such as in stool shape, smell, form, and color. In about 50% of children, habits were improved within a period of 2 to 3 weeks and “allergic-like” skin symptoms were also markedly reduced. Eye contact and attention also improved in about 50% of patients. In addition, 30%–50% of patients showed learned tasks and social interactions and about 10% of children started speaking words or short sentences. However, no improvements were recorded for hyperactivity or aggressiveness [194]. In 2013, Taliou et al. reported an improvement in adaptive functioning and in overall behavior of about 26.6%–34.8% of the autistic children when treated with luteolin, quercetin, and rutin extracted from Chamomile sp. and Sophora sp. leaves [77]. In another clinical study, co-ultra-PEA-LUT led to an improvement in ASD-related symptoms and reduced motor stereotypes [186]. Another study with PEA showed beneficial effects of this treatment in two child patients with ASD [195]. These subjects showed significantly improved cognitive aspects, expressive language, and sociability, as well as the general severity of autism. Furthermore, slight improvements were observed in irritability, hyperactivity, eye contact, and speech following treatment with a flavonoid component extracted from G. biloba leaves [84].

All these clinical studies with plant-food-derived bioactives in ASD are summarized in Table 4.

### Table 4: Summary of clinical studies.

<table>
<thead>
<tr>
<th>Phytochemicals</th>
<th>Subjects</th>
<th>Results</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Food supplements including choline, folic acid, and multivitamins</td>
<td>Pregnant patient</td>
<td>Reduced expression of ASD-related genes</td>
<td>[189]</td>
</tr>
<tr>
<td>Omega-3 and vitamin D</td>
<td>Case study of ASD patient</td>
<td>Reduced ASD symptom</td>
<td>[190]</td>
</tr>
<tr>
<td>Metformin</td>
<td>ASD adult patients</td>
<td>Stabilized BMI z-score</td>
<td>[118]</td>
</tr>
<tr>
<td>Cannabis and cannabidiol</td>
<td>ASD adult patients</td>
<td>Reduced ASD symptom</td>
<td>[193]</td>
</tr>
<tr>
<td>Luteolin and quercetin</td>
<td>Children with ASD</td>
<td>Improvement in gastrointestinal features, eye contact, and attention social interactions</td>
<td>[194]</td>
</tr>
<tr>
<td>Luteolin, quercetin, and rutin</td>
<td>Children with ASD</td>
<td>Improvement in adaptive functioning and in overall behavior</td>
<td>[77]</td>
</tr>
<tr>
<td>Co-ultra-PEA-LUT</td>
<td>ASD adult patients</td>
<td>Reduced motor stereotypes, anxiety, and worsening in social skills</td>
<td>[186]</td>
</tr>
<tr>
<td>PEA</td>
<td>Children with ASD</td>
<td>Improvement of cognitive aspects, expressive language, and sociability</td>
<td>[195]</td>
</tr>
<tr>
<td>Flavonoid component extracted from G. biloba</td>
<td>ASD adult patients</td>
<td>Slight improvements in irritability, hyperactivity, eye contact, and speech</td>
<td>[84]</td>
</tr>
</tbody>
</table>
6. Conclusions and Upcoming Perspectives

Plant-bioactive molecules demonstrated promising results in ASD treatment. However, there is a scarce number of clinical trials available so far, and none of them have been approved for the treatment of ASD or ASD-related disorder. Thus, further preclinical and clinical studies are needed for a more in-depth understanding of plant-derived bioactives as drug discovery candidates on ASD treatment.

Conflicts of Interest

The authors declare no conflict of interest.

Authors’ Contributions

All authors contributed to the manuscript. J.S.-R., N.C.-M., and W.C.C. were involved in conceptualization. All authors were involved in validation, investigation, resource acquisition, data curation, and writing of the manuscript. C.Q., L.-C.B., M.B., F.S., N.C.-M., J.S.R., V.L., M.M.A., W.C.C., and F.L. were involved in reviewing and editing. All authors have read and agreed to publish this version of the manuscript.

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