



## Endocrine disruptors also function as nervous disruptors and can be renamed endocrine and nervous disruptors (ENDs)

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### ABSTRACT

Endocrine disruption (ED) and endocrine disruptors (EDs) emerged as scientific concepts in 1995, after numerous chemical pollutants were found to be responsible for reproductive dysfunction. The World Health Organization established in the United Nations Environment Programme a list of materials, plasticizers, pesticides, and various pollutants synthesized from petrochemistry that impact not only reproduction, but also hormonal functions, directly or indirectly. Cells communicate via either chemical or electrical signals transmitted within the endocrine or nervous systems. To investigate whether hormone disruptors may also interfere directly or indirectly with the development or functioning of the nervous system through either a neuroendocrine or a more general mechanism, we examined the scientific literature to ascertain the effects of EDs on the nervous system, specifically in the categories of neurotoxicity, cognition, and behaviour. To date, we demonstrated that all of the 177 EDs identified internationally by WHO are known to have an impact on the nervous system. Furthermore, the precise mechanisms underlying this neurodisruption have also been established. It was previously believed that EDs primarily function via the thyroid. However, this study presents substantial evidence that approximately 80 % of EDs operate via other mechanisms. It thus outlines a novel concept: EDs are also neurodisruptors (NDs) and can be collectively termed endocrine and nervous disruptors (ENDs). Most of ENDs are derived from petroleum residues, and their various mechanisms of action are similar to those of “spam” in electronic communications technologies. Therefore, ENDs can be considered as an instance of spam in a biological context.

### 1. Introduction

Endocrine disruption (ED) or endocrine disruptors (EDs) emerged as scientific concepts in 1995 [Colborn [\[1\]](#); Lindström et al. [\[2\]](#); Ginsburg [\[3\]](#)] after numerous chemical pollutants were found to be responsible for reproductive dysfunction. This was first posited three decades prior [Carson [\[4\]](#)]. EDs were reviewed more recently in a book [Seralini [\[5\]](#)], which advanced the understanding of the molecular bioaccumulation of identified xenobiotics, as well as their combined and long-term effects on the whole physiology of one or several generations. They have been identified in organisms at all levels of the ecosystem and are also ubiquitously found in the food chain.

The World Health Organization established in the United Nations Environment Programme [WHO [\[6\]](#)], a list of 176 compounds comprising materials, pesticides, and various pollutants ([Table 1](#), columns 1–3) impacting not only reproduction, but also hormonal functions—directly or indirectly—primarily in mammals, including humans.

This has enabled numerous countries to establish regulatory policies for the production and use of these chemicals or to manage contamination in food, air and water. Numerous political debates have been raised around regulatory thresholds, based on the effects of ENDs demonstrated on a population or a subpopulation of animals or humans, and published at epidemiological and molecular levels. The herbicide Roundup has been added as the 177th compound due to its widespread usage as a pesticide, combined with the relatively recent demonstration of its ED effects [Richard et al. [\[7\]](#)].

Epidemiology is not technically adapted to solve the questions on combined and long-term effects of molecules or mixtures on mammalian or human health [Mesnage et al. [\[8\]](#)]; this becomes further complicated when epigenetic and transgenerational impacts are studied [Skinner and Anway [\[9\]](#)]. For instance, pesticide accumulation is rarely measured in organs after death in order to ascertain whether they can be used as markers to correlate their levels with pathologies. Instead, the understanding of endocrine disruption may be aided by advances in the

**Abbreviations:** ED, Endocrine disruptors; ENDs, Endocrine and nervous disruptors; ND, Nervous disruptors; WHO, World Health Organization.

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combined knowledge of biochemical, cellular, organic and environmental effects in experimental animal models, farm animals, wildlife observations in contaminated areas, and occupational medicine in factories producing the chemicals in question.

The endocrine system is not limited to the control of sexual reproduction and development. Endocrine disruptors may affect the thyroid as well as the glucocorticoid axis, adrenal and pancreatic systems, adipose tissue and immune or neuroendocrine targets [Laessig et al. [10]; Masuo and Ishido [11]; Weiss [12]; Leon-Olea et al. [13]]. They even possess cognitive effects [Schantz and Widholm [14]], particularly via various neuromediator interferences.

Cells communicate via either chemical or electrical signals that are transmitted within the endocrine or nervous systems. Generally, in endocrinology, hormones may have a biphasic action dependent on the receptors' availability and concentration, resulting in time-, dose- and sex-dependent effects that vary according to the targeted tissue and the specific organism. It is thus reductionist to say, for instance, that a hormone such as estradiol "stimulates ovulation", since it can inhibit this function when used at pharmacological doses as a pill or during embryonic or foetal life. Endocrine disruptors may thus possess the potential of exerting similar biphasic ambivalent effects.

Thus, to determine whether hormonal disruptors may also interfere directly or indirectly with neural development [Grandjean and Landrigan [15]] or functioning even in adults, either through a neuroendocrine or a more general mechanism, this study examined the scientific literature to ascertain the effects of ENDs on the nervous system—including neurotoxicity, cognition, and behaviour—of the major internationally identified (according to WHO) endocrine disruptors.

## 2. Materials and methods

Each compound was numbered (Nb, Table 1) out of 176 known endocrine disruptors [WHO [6]], plus Roundup. Its name was associated with the keyword "nervous" or "neurotoxicity" or "cognitive" or "behavio(u)r" on the PubMed data bank, and eventually on Google Scholar. When the references were too numerous, "or" was excluded in order to directly associate the keywords. If more than 20 references were found to be published, "review" was added to the keywords and cited. Finally, a maximum of five references were indicated, focusing on the most recent research in humans or mammals, without excluding other models. The mechanisms were documented (Table 1, column 4) as direct effects on the neurons or the nervous system, or as indirect effects, including thyroid regulation.

## 3. Results and discussion

All of the 176 internationally identified EDs (as per WHO guidelines) were studied in the international bibliography published between 1931 and 2021, in order to assess their effects on the nervous system (Table 1). It was observed that every single one of the EDs induced neurodisruption or clear neuromodulation, while even stimulation occurred in certain rare cases. Thus, 100 % of the EDs studied were known neuro-modulators, either directly or indirectly. Previously, it was generally believed that endocrine disruptors with a neuronal impact functioned via the thyroid gland, which is classically known to control nervous system development. However, this study presents substantive evidence that at least 79.1 % of the EDs are, in fact, also NDs through various other mechanisms (Table 1). 37 compounds out of the total were implicated to possess a thyroid-dependent mechanism. Therefore, the emergence of this overarching concept compelled us to propose the introduction and use of a collective abbreviation "ENDs" for endocrine and nervous disruptors.

Therapeutic hormones or pharmaceutical products (see column 3 in Table 1) that may exist as pollutants in rivers [Arya et al. [16]; Saussereau et al. [17]; Gouille et al. [18]] often comprise ENDs—such as certain natural phytoestrogens that are more biodegradable than stable petrochemistry-based xenobiotics—often resulting in bioaccumulation within organisms. Mixtures of chemicals such as Agent Orange or dioxins have also been cited as ENDs. In addition, formulants of pesticides—such as POEA (polyoxyethylene tallow amine)—have been discovered to be EDs more recently [Gasnier et al. [19]; Defarge et al. [20]] and may also possess nervous effects [Malhotra et al. [21]; Sato et al. [22]]. In some models, glyphosate alone [Martinez et al. [23]; Coullery et al. [24]], known as the declared active ingredient of a major pesticide formulation used across the world, has been shown to be less toxic or disruptive to the nervous system than its equivalent formulants present in Roundup [Mesnage et al. [8]; Aitbali et al. [25]; Gallegos et al. [26]]. This also proves to be true for non-glyphosate-based herbicides containing non-declared polycyclic aromatic hydrocarbons and heavy metals that are individually identified as EDs [Seralini and Jungers [27]].

The impacts of ENDs on physiological functions and pathologies appear to be less specific than expected, especially since they interact with several intracellular and intercellular communications simultaneously. A majority of them are derived from petroleum residues, and their mechanisms of action can be likened to those of "spam" in electronic communications technologies. Thus, ENDs can be considered as a widespread instance of spam in a biological context.

**Table 1**

Endocrine disruptors (according to WHO, 2013) also function as nervous system disruptors, and may thus be collectively termed as Endocrine and Nervous Disruptors (ENDs).

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
1	Acetochlor	Herbicide	Roman [28]: antithyroid agents Helbing et al. [29]: thyroid hormone receptor gene expression in the brain Zafeiridou et al. [30]: compound action potential of the sciatic nerve Goldner et al. [31]: hypothyroidism
2	Alachlor	Herbicide	Seok et al. [32]: central nervous system symptoms Lo et al. [33]: severe neurological and cardiovascular outcomes after acute poisoning Doïcheva [34]: higher irritability, lack of coordination and orientation Sirohi et al. [35]: specific binding to lactoperoxidase
3	Amitrole	Herbicide	Chilumuri et al. [36]: inhibit neuroprotection against amyloid peptides Pan et al. [37]: reduction of thyroid-stimulating hormone receptors Roman [28]: hypothyroxinemia
4	Anthracene	PAH	Brucker-Davis [38]: thyroid disruption in utero or direct neurotoxicity Palanikumar et al. [39]: neurotoxicity by inhibition of acetylcholinesterase Vieira et al. [40]: increase catalase activity and superoxide dismutase, glutathione reductase and peroxidase. Mucio-Ramírez et al. [41]: decrease somatodendritic vasopressin release
5	Aroclor 1254	PCB mixture	Wei et al. [42]: oxidative stress in the brain Coburn et al. [43]: Inhibition of vasopressin release from magnocellular neuroendocrine cells

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
6	Arsenic (As)	Heavy metal, pesticide	<p>Majumdar et al. [44]: induces oxidative stress in the brain</p> <p>Tilson et al. [45]: increases translocation of protein kinase C and decreases Ca<sup>2+</sup>-buffering in the brain</p> <p>Liu et al. [46]: cognitive impairment</p> <p>Webb et al. [47]: chronic neurological disease</p> <p>Preciados et al. [48]: influence nuclear respiratory factor 1 by genomic and epigenomic networks, this contributes to the development of complex chronic human brain disorders</p> <p>Karri et al. [49]: pro-oxidant elements dominate antioxidants factors and leads to cognitive dysfunction</p> <p>Tyler et al. [50]: alters epigenetics and hippocampal function, glutamatergic, cholinergic and monoaminergic signaling; altered adult neurogenesis; and increases Alzheimer's-associated pathologies</p> <p>Ma et al. [51]: reduces dopamine levels in the substantia nigra and corpus striatum in the midbrain</p> <p>Zhang et al. [52]: severe dopamine neuron degeneration</p>
7	Atrazine	Herbicide	<p>Lin et al. [53]: increases in striatal dopamine; decreases in perirhinal cortex serotonin</p> <p>Ottinger et al. [54]: impact on arginine vasotocin, catecholamines and gonadotropin releasing hormone.</p>
8	Benz(a)anthracene	PAH	<p>Dayal et al. [55]: most notable symptoms are neurologic</p> <p>Slotkin et al. [56]: impairs neurodifferentiation and neurite formation, development of dopamine and acetylcholine phenotypes</p> <p>Sarma et al. [57]: induces neuronal cell damage involving oxidative stress through mitochondria-mediated apoptosis pathway</p>
9	Benz(a)pyrene	PAH	<p>Yang et al. [58]: cause disruption of glutamate (Glu) neurotransmitter transmission by decreasing the level of Glu, reducing the expression of Glu receptors</p> <p>Chepelev et al. [59]: binds to the aryl hydrocarbon receptor, modulate the transcription of glutamate receptor subunits, decrease long-term potentiation, learning and memory</p> <p>Niu et al. [60]: reduces neurobehavioral function and monoamine, amino acid and choline neurotransmitter levels</p>
10	BB-153	PBB	<p>Jacobson et al. [61]: associations of PBBs and PCBs with thyroid disease and thyroid hormone levels.</p> <p>Tilson et al. [62]: chronic exposure produces behavioral or neurological toxicity</p> <p>Morgenstern et al. [63]: impairs thyroid function in preschool children</p>
11	Benzyl butyl phthalate	Phthalate	<p>Min et al. [64]: attenuation of the effects of cAMP Response Element-Binding Protein downstream, oxidative damage and impaired behavioral performance</p> <p>Betz et al. [65]: changes in amygdala protein s related on synaptic plasticity</p> <p>Kasuya M [66]: inhibited the outgrowth of nerve fibers and glial cells from cerebellar explants</p> <p>Chen et al. [67]: activation of NMDA receptors inhibits the expression of phosphodiesterases, favoring apoptosis induction and induces neurotoxicity.</p>
12	BDE-209	PBDE	<p>Li et al. [68]: the mRNA expressions of synaptobrevin 2, syntaxin 1A, SNAP-25, and synaptophysin are significantly decreases in the hippocampi of rat exposed to it</p> <p>Sun et al. [69]: during rat pregnancy increases hippocampal autophagy, decreases neuron viability</p> <p>Vuong et al. [70]: large prospective human cohorts demonstrates that prenatal and postnatal exposure adversely impacts externalizing behavior (e.g., hyperactivity and conduct problems)</p> <p>Zhuang et al. [71]: induces neurotoxicity and cognitive impairment by upregulation of nuclear TAR DNA-binding protein 43 in the hippocampus provoking neuronal apoptosis.</p>
13	BDE-47	PBDE	<p>Chen et al. [67]: inhibits axonal growth via ryanodine receptor-dependent mechanisms</p> <p>Zhai et al. [72]: prenatal exposure inhibits neurodevelopmental function and behavior with an increase of luteinizing hormone levels.</p>
14	BDE-99	PBDE	<p>Dach et al. [73]: reduces expression of myelin associated genes like HMBP due to oligodendrocyte reduction.</p> <p>Ding et al. [74]: prenatal exposure is associated with lower developmental quotients in young children.</p> <p>Roze et al. [75]: transplacental transfer is associated with worse fine manipulative abilities, and worse attention of children at school age.</p> <p>Bahadar et al. [76]: exposure can lead to aberration of vital systems in the body like nervous, cardiovascular, and respiratory.</p>
15	Benzene	Aromatic solvent	<p>Manto M. [77]: in humans, cerebellum is a main target of environmental toxins such as toluene/benzene derivatives.</p> <p>Ritchie et al. [78]: a number of published studies report acute or persisting neurotoxic effects of hydrocarbon fuels.</p>
16	Benzylidene camphor	UV filter	<p>Ruszkiewicz et al. [79]: potential neurotoxicity</p> <p>Faass et al. [80]: reduces preoptic and receptive behaviors and specific gene expression in ventromedial hypothalamic nucleus and medial preoptic area.</p> <p>Ejaredar et al. [81]: exposure in childhood is associated with higher levels of anxiety, depression, hyperactivity, inattention, and conduct problems.</p>
17	Bisphenol A	Plastics monomer	<p>Zhou et al. [82]: affects neuron numbers in different regions of the hippocampus altering learning and memory ability of adolescent mice.</p> <p>Inadera H. [83]: in a review, it has detrimental effects on neurological development.</p> <p>Masuo et al. [11]: locus ceruleus is enlarged in treated male rats and neurodegenerative disorders.</p>
18	Bisphenol A diglycid ether	Plastics monomer	<p>Hutler et al. [84]: the main abnormalities in amphibians larvae related to neurotoxicity.</p> <p>Ohtani et al. [85]: exposure alters offspring behavior, resulting in increases in anxiety and depressive state in mice.</p>
19	Bisphenol F	Plastics monomer	<p>Rosenfeld [86]: induces neurobehavioral disruptions.</p> <p>Castro et al. [87]: affects genes in the juvenile female rats prefrontal cortex.</p>
20	Bisphenol S	Plastics monomer	<p>Wu et al. [88]: triggers oxidative stress in the nervous system in vivo and in vitro in humans.</p> <p>Castro et al. [87]: affects genes expression and dopamin serotonin system in the prefrontal cortex of juvenile female rats.</p>
21	Bromacil	Herbicide	<p>Lombardi et al [89]: exposure through residential proximity to agricultural applications during pregnancy may increase the risk of childhood central nervous system tumors.</p>
22	Butylate	Herbicide	<p>Lulla et al. [90]: for Ziram: selectively toxic to dopaminergic neurons in vivo, and this toxicity is synuclein-dependent.</p>

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
23	Butylated hydroxyanisole	Antioxidant for long preservation of food products	Miyazaki et al. [91]: quinone reductase inducer which significantly and dose dependently blocked methamphetamine-induced elevation of quinoprotein, and ameliorated methamphetamine-induced cell death. Katsuki et al. [92]: abolishes neurotoxic action of arachidonic acid. Raciti et al. [93]: affects epigenetics with negative consequences on the development of the nervous system. Jacobo-Estrada et al. [94]: induces toxicity in fetus on the central nervous system. Zhang et al. [95]: induces autophagy in neurons promoting neurodegenerative disorders. Sanders et al [96]: exposure may be associated with poorer cognition. Bo et al. [97]: neurotoxic on the long-term. Al-Rubai et al. [98]: reduces neurosphere size and cell migration at high doses. Reduction in glial fibrillary protein and tubulin III.
24	Cadmium (Cd)	Heavy metal	Hansen et al. [99]: induces encephalopathy with hyperammonemia, intrinsic effects on cerebral receptors. Gualtieri et al. [100]: affects tests of memory, psychomotor speed, cognitive flexibility, and attention. Lee et al. [101]: inhibits classically acetylcholinesterase in the nervous system; induces cognitive impairments by disturbed neurodevelopment.
25	Carbamazepine	Pharmaceutical anti-epileptic	Freeborn et al. [102]: affects electroencephalogram by decreasing theta area and delta frequency, increases beta frequency. Wang et al. [103]: acetylcholinesterase is inhibited by high dose and damages the sciatic nerve.
26	Carbaryl	Insecticide	Cocco et al [104]: chronic exposure to PCBs affects the development and function of the nervous system. Lovato et al [105]: a mixture of PCB can induce functional deficits and altered behavioral threat in zebrafish.
27	CB-15	PCB	Ozcan et al. [106]: dioxin-like and non-dioxin-like PCB congeners are equally potent in causing cognitive decrements seen in children exposed prenatally to PCBs.
28	CB-77	Coplanar PCB	Howard et al. [107]: binds the aryl hydrocarbon receptors with high affinity. Brucker-Davis et al. [108]: negative impact on neurocognitive development, negatively correlated on motor and expressive language in children.
29	CB-118	PCB	Doi et al. [109]: four-month-olds children with a low-level of prenatal exposure exhibits a preference for the upright biological motion, impairs the development functioning and brain development.
30	CB-126	Planar PCB	Cauli et al. [110]: impairs motor coordination at 2 months in males but not in female rats, reduces locomotor activity in females.
31	CB-132	PCB	Uwinana et al. [111]: does not appear to affect dopaminergic cells in cultures or levels of dopamine. To be further studied.
32	CB-138	PCB	Boix et al. [112]: exposition activates metabotropic glutamate receptors and that increases dopamine in females and reduces it in males. The opposite changes are observed for glutamate, in rat nucleus accumbens. Campagna et al. [113]: Ca <sup>2+</sup> homeostasis and androgen receptor signaling pathways are primarily disrupted in cerebellum proteome, contributing toward a premature ageing and neurotoxicity.
33	CB-153	PCB	Naert et al. [114]: birds bioaccumulate in brain and the central nervous system. Enayah et al. [115]: neurotoxic, and affects dopamine turnover in vitro. Cauli et al. [110]: many motor alterations and induces hyperactivity at adulthood in rats.
34	CB-169	Planar PCB	Gascon et al. [116]: deleterious effects on neuropsychological development which are mainly attributable to prenatal exposure.
35	CB-180	PCB	Morse et al. [117]: local hypothyroidism occurs in the brains of fetal and neonatal rats exposed by increase in type II thyroxine 5'-deiodinase in the brain. Boix et al. [112]: affects motor activity in rats; increased glutamate release in nucleus accumbens following activation of metabotropic glutamate receptors would be involved in reduced dopamine release.
36	Chlordane	Organochlorine insecticide	Naert et al. [114]: birds bioaccumulate in brain and the central nervous system. Kilburn [118]: it is suggested that it causes protracted neurotoxicity in patients. Kilburn and Thornton. [119]: exposure is associated with protracted impairment of neurophysiological and psychological functions. The central nervous system is the most important target. Grutsch et al. [120]: the characteristic signs of acute toxicity are hypothermia, hyperexcitability, tremors and convulsions. In human, signs of acute toxicity are tremors and convulsions.
37	Chlordibromo-methane	Trihalo-methane	Villanueva et al. [121]: Minor associations observed between exposure during gestation and child neuropsychological development. Balster et al. [122]: effect on operant behavior in mice.
38	Chlorinated Paraffins	Flame retardants, lubricants, plasticizers	Liu et al. [123]: exposure could alter gene expression in the hypothalamic-pituitary-thyroid axis. Mariussen et al. [124]: neurobehavioral effects, indicating adverse effects on the central nervous system: alteration of neurotransmitter functions, Ca <sup>2+</sup> homeostasis processes, induction of protein kinase C and phospholipase A2 mobilization, and oxidative stress. Eriksson et al. [125]: significant decrease of presynaptically sodium-dependent choline uptake in mice. Burke et al. [126]: acute exposure of humans irreversibly inhibit acetylcholinesterase, and chronic exposure induces neurological deficits that range from cognitive impairments to tremors in childhood
39	Chlorpyrifos	Insecticide	Yamada et al. [127]: inhibit neural induction via mitochondrial fusion protein mitofusin 1-mediated mitochondrial dysfunction in human stem cells. Sogorb et al. [128]: seems able to induce neurodevelopmental alterations in animals Lee et al. [101]: affects protein levels in the mice developing brain and induces persistent adult behavior and cognitive impairments; neurotoxic effects. Yeo et al. [129]: disability is improved.
40	Citalopram	Antidepressant	Gaanderse et al. [130]: induces dyskinesia of the tongue. Sprowles et al. [131]: selective serotonin reuptake inhibitor, alters spatial learning and memory, anxiety, depression in rats.
41	Clofentezine	Pesticide Acaricide	Hurley [132]: induces thyroid follicular cell tumors in rodents; disrupts thyroid-pituitary homeostasis.

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
42	Coumaphos	Pharmaceutical	<b>Abdelsalam</b> [133]: inhibition of brain of acetylcholinesterase, inhibition of brain neurotoxic esterase, plus delayed neurotoxicity. <b>Abou-Donia et al.</b> [134]: degeneration of axons and myelin in the spinal cord.
43	Coumestrol	Phytoestrogen	<b>Jantaratnotai et al.</b> [135]: suppression of interferon regulatory factor-1 and phosphorylated STAT1 expression in lipopolysaccharide-activated microglia.
44	D4 Cyclic siloxane		<b>Andreou et al.</b> [136]: unusual constriction of the isolated sciatic nerve, death of nerve fibers.
45	D5 Cyclic siloxane	Material	<b>Fuzzard et al.</b> [137]: due to silicone implants, myalgias, chronic fatigue, cognitive impairment.
46	D6 Cyclic siloxane		<b>Yu et al.</b> [138]: perinatal exposure enhances estrogen receptor alpha expression in several brain regions such as stria terminalis, arcuate hypothalamic nucleus, and central amygdaloid nucleus. <b>Zeng et al.</b> [139]: significant effects on locomotor activity, mood and social behavior after long-term consumption.
47	Daidzein	Isoflavones	<b>Jin et al.</b> [140]: the neurotoxic effect of daidzein could be due to the inhibition of the GABA(A) receptor resulting in further enhancement of excitation by glutamate and leading to cellular damage in primary rat neuronal cultures.
48	Dibromochloropropane	Pesticides	<b>Román GC</b> [28]: inhibits thyroperoxidase that catalyzes iodination and thyroid hormone biosynthesis <b>Teitelbaum</b> [141]: the product is central nervous system depressant.
		Herbicide	<b>Liu et al.</b> [46]: neuroendocrine disruptor that impacts the expression of neurotoxicity-related genes such as Ache, Gap43, Gfap, Syn2a, Shha, Mbp, Elavl3, Nestin and Ngn1 in early developmental stages of zebrafish.
49	Desethylatrazine	Metabolite	<b>Gunderson et al.</b> [142]: Strong positive correlation between CYP19 and SF-1 transcript abundance in exposed tadpoles during brain development. <b>Hossain et al.</b> [143]: decreases striatal dopamine levels and in synaptosomes in rat. <b>Yi et al.</b> [144]: Increases various neurologic diseases; systemic atrophies affecting the nervous system, including spinal muscular atrophy, Alzheimer disease, and peripheral polyneuropathies.
50	2,4-D	Herbicide agent orange	<b>Bortolozzi et al.</b> [145]: changes in various neurotransmitter systems, such as serotonin (5-HT) and dopamine (DA), were proposed to mediate some of the behavioral effects in rats <b>Evangelista de Duffard et al.</b> [146]: Increases sensitivity in dopamine D2-like brain receptor from 2,4-dichlorophenoxyacetic acid (2,4-D)-exposed and amphetamine challenged rats.
51	2,4-Dichlorophenol	Chlorophenol	<b>Krieg</b> [147]: At low concentrations, it may act at acetylcholine and $\gamma$ -aminobutyric acid synapses in the central nervous system to modify neurobehavioral test performance.
52	3-Diltiazem	Pharmaceutical	<b>Stevens et al.</b> [148]: Calcium channel blocker.
53	2,4'-DDD (o,p'-DDD)	Organochlorine	<b>Heilmann et al.</b> [149]: Some central nervous disorders were observed. <b>Lanser et al.</b> [150]: Neuropsychologic and neurologic side effects. <b>Du Rostu et al.</b> [151]: neurological symptoms and neurotoxicity both central and peripheral. <b>Kajita et al.</b> [152]: depressive-like symptoms after prenatal exposure possibly by DNA hypomethylation and estrogen receptor signaling pathway.
		Insecticide and pharmaceutical mitotane	<b>Zhang et al.</b> [153]: effects on glucocorticoid-receptors on the nervous system. <b>Eskenazi et al.</b> [154]: Prenatal exposure to DDT, and to a lesser extent DDE, was associated with neurodevelopmental delays during early childhood.
54	2,4'-DDT (o,p'-DDT)	Insecticide	<b>Halldin</b> [155]: altered the development of the neural system and resulted in demasculinization of male quail. <b>Fry</b> [156]: impaired differentiation of the nervous system through mechanisms of hormonal mimicking of estrogens.
55	4,4'-DDD (p,p'-DDD)	Organochlorine	<b>Al-Saleh et al.</b> [157]: low parent's evaluation of developmental status of infants was significantly associated with DDD in breast milk.
		Insecticide	<b>Wnuk et al.</b> [158]: stimulation of retinoid X receptor $\alpha$ and retinoid X receptor $\beta$ -mediated intracellular signaling plays an important role in the propagation of DDE-induced apoptosis during early stages of neural development.
56	4,4'-DDE (p,p'-DDE)	Organochlorine	<b>Cartier et al.</b> [159]: p,p'-DDE exposure, both pre- and postnatally, during early childhood is associated with visual processing impairment later in life. <b>Eskenazi et al.</b> [154]: Prenatal exposure to DDT, and to a lesser extent DDE, was associated with neurodevelopmental delays during early childhood.
		Insecticide	<b>Kajita et al.</b> [152]: depressive-like symptoms after prenatal exposure possibly by DNA hypomethylation and estrogen receptor signaling pathway.
57	4,4'-DDT (p,p'-DDT)	Organochlorine	<b>Zhang et al.</b> [153]: effects on glucocorticoid-receptors on the nervous system. <b>Eskenazi et al.</b> [154]: prenatal exposure to DDT, and to a lesser extent DDE, was associated with neurodevelopmental delays during early childhood
		Insecticide	<b>Parent et al.</b> [160]: DDT is involved in neuroendocrine disruption of the reproductive axis.
58	Di-(2-ethylhexyl) adipate	Plasticizer	<b>Lee et al.</b> [161]: inappropriate expression of granulin and/or p130 genes in the brains of male and female neonatal rats by perinatal exposure may exert permanent effects on the hypothalamus, thereby decreasing sexual behavior after maturation. <b>Arbo et al.</b> [162]: neuroactive steroid that modulate neuronal and astroglial function and have neuroprotective effects.
59	Dehydroepi-androsterone	Natural hormone	<b>Woda et al.</b> [163]: synthesized in nervous cells, neuroactive local factor in the central nervous system and the periphery. <b>Yu et al.</b> [164]: induced depression-like behavior in polycystic ovary syndrome mice, possibly through down-regulation of brain monoamines and/or their metabolites. <b>Li et al.</b> [165]: provides robust ischemic neuroprotection but also exerts neurotoxicity when administered during ischemia and early reperfusion. <b>Coplan et al.</b> [166]: glucocorticoid-induced neurotoxicity. <b>Yu et al.</b> [164]: protracted disruption of mental functions.
60	Dexamethasone	Synthetic steroid	<b>Lopes et al.</b> [167]: affects the phosphorylation state of glutamate AMPA receptors in the human limbic system. <b>Feng et al.</b> [168]: decreased both body and brain weight gain. Tapering and repeated doses increased caspase-3 activity. Impaired learning and memory capability at juvenile age.

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
61	Dibutyl phthalate	Phthalate	<p>Uno et al. [169]: induced degeneration and depletion of the hippocampal pyramidal and dentate granular neurons in the brains of primate fetuses.</p> <p>Wójtowicz et al. [158]: Aryl hydrocarbon receptor is involved in dibutyl phthalate induced apoptosis and neurotoxicity, while the estrogen receptors and peroxisome proliferator-activated receptor gamma signaling pathways are impaired by the phthalate.</p> <p>Farzanehfard et al. [170]: could reduce total distance movement, impair memory function and induce anxiety in mice. Significant nuclei size reduction and condensation in dentate gyrus cells</p> <p>Yan et al. [171]: link between oxidative stress and anxiety-like behavior produced by dibutyl phthalate at high doses.</p> <p>Chantong et al. [172]: potentiation of oxidative stress and pro-inflammatory cytokine expression in microglia cells</p> <p>Tsuji et al. [173]: Dibutyltin is neurotoxic and poly-L-lactides toxicity increases with the increase in tin concentration.</p>
62	Dibutyltin	Plastics stabilizer	<p>Jenkins et al. [174]: developmental neurotoxicant; the incidence of apoptotic cell death, was increased in the neocortex and hippocampus.</p> <p>Kobayashi et al. [175]: synaptic parameters modulations; tributyltin metabolites inhibit various parameters of cholinergic activity with a potency ranking of tributyltin &gt; dibutyltin &gt; monobutyltin.</p> <p>Evangelista de Duffard et al. [146]: has effects on motor, sensory, or cognitive functions.</p> <p>Lessenger et al. [176]: case report, neurological injury, cognitive and emotional difficulties persisted over an 18-mo period.</p>
63	Dicofol	Organochlorine Insecticide	<p>Cowie et al. [177]: disrupts proteins related to oxidative respiration and mitochondrial stress in the central nervous system.</p> <p>Schmidt et al. [178]: induced neurotoxicity by impaired mitochondrial bioenergetics and endoplasmic reticulum stress in rat dopaminergic cells.</p>
64	Dieldrin	Organochlorine Insecticide	<p>Babot et al. [179]: Long term exposure reduces gamma-aminobutyric acid type A and N-methyl-aspartate receptor function in primary culture of mouse cerebellar granule cells.</p> <p>Evangelista de Duffard et al. [146]: motor sensory or cognitive function effects.</p> <p>Luu et al. [180]: regulate microRNAs in a sex-specific manner which may interfere with proper hippocampal development in males and preserve hippocampal development in females.</p> <p>Preciados et al. [48]: induced brain health deficits by NRF1 regulated gene networks.</p> <p>Park et al. [181]: sex-dependent effect on anxiety proneness in childhood.</p>
65	Diethyl hexyl phthalate	Phthalate	<p>Quinnies et al. [182]: transgenerational modifications in the expression of several pituitary hormones involved in the hypothalamic-pituitary-adrenal axis and in stress hormones.</p> <p>Huang et al. [183]: prenatal exposure was associated with decreased cognitive development in the young children.</p> <p>Télez-Rojo et al. [184]: prenatal exposure creates sex specific neurodevelopmental effects.</p> <p>Doherty et al. [185]: prenatal associations between urinary phthalates in aged mothers and brain performances in young children.</p>
66	Mono-2-ethyl-hexyl phthalate	DEHP Hydrolysis product	<p>Mao et al. [186]: induce spatial cognitive deficits through altering the expression of apoptosis-related protein.</p> <p>Won et al. [187]: increased exposure exhibited supralinear associations with social, thought and attention problems in children.</p> <p>Tomihara et al. [188]: developmental deficits may stem from both in utero toxicity and aberrant maternal care.</p>
67	Mono-n-butyl phthalate	DBP Hydrolysis product	<p>Frye et al. [189]: effects on the aryl hydrocarbon receptor, the peroxisome proliferator-activated receptor and the retinoid X receptor, signal transduction pathways, and on calcium influx and/or neurotransmitter receptors.</p> <p>Sato et al. [190]: marked influence on synaptogenesis and neuronal vulnerability through mechanisms other than through estrogen receptors.</p> <p>Ma et al. [191]: cause cognitive deficits and anxiety.</p>
68	Diethylstilbestrol	Synthetic estrogen	<p>Peng L [192]: oral exposure of mice induced brain damage, and oxidative stress, inflammation, and apoptosis.</p> <p>Boberg et al. [193]: behavioral effects, spatial learning effects in perinatally exposed rats.</p> <p>Kim et al. [194]: Inhibitory effects on proton currents in microglial cells.</p>
69	Diisonyl phthalate	Plasticizer	<p>Mansfeld et al. [195]: reduced attention and increased self-reported drowsiness.</p> <p>Wilken et al. [196]: caused significant decrements in vigilance and cognitive functioning.</p> <p>Vaswani et al. [197]: alterations of opioid neuropeptides such as beta endorphin, meth-enkephalin and dynorphin levels</p>
70	Diphenhydramine	Antihistamine	<p>Jang et al. [198]: induced acute neurotoxicity via induction of oxidative stress and pro-inflammatory responses.</p>
71	Dimethyl-benz(a)anthracene	PAH	<p>Caudle WM [199]: can alter the normal development and potential function of neurotransmission in the frontal cortex</p>
72	Endosulfan (alpha/beta)	Organochlorine Insecticide	<p>Silva et al. [200]: neurotoxicity and developmental effects in the zebrafish.</p> <p>Silva et al. [201]: effects on brain biogenic amine levels. Developmental reproductive toxicity or endocrine disruption occurs only at doses causing neurotoxicity.</p> <p>Bagchi et al. [202]: induced lipid peroxidation and DNA damage in brain and regional distribution of catalase activity in rat brain.</p>
73	Endrin	Organochlorine Insecticide	<p>Gray et al. [203]: alteration of central nervous system function in rats and hamsters even though endrin produces gross morphological defects only in hamsters.</p> <p>Li et al. [204]: anxiety disorders, augmentation of vulnerability factors associated with anxiety disorder development; and facilitation of the maintenance of anxious symptoms post-development.</p> <p>Preciados et al. [48]: influences NRF1 regulated gene networks in the development of complex human brain diseases.</p>
74	Estradiol	Natural hormone	<p>Perez-Alvarez et al. [205]: neuroprotective role after ischemic injury.</p> <p>Rossetti et al. [206]: neurosteroid bind specific receptors to promote essential brain functions.</p>
75	Estrone	Natural hormone	

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
76	Ethinylestradiol	Synthetic hormone	<p><b>Mahmoud et al.</b> [207]: may influence adult hippocampal neurogenesis, with a focus on cognitive function and mood regulation.</p> <p><b>Grimm et al.</b> [208]: may act upon neuronal bioenergetics in a delicate balance with an age-related effect that might be involved in mitochondrial dysfunction underlying neurodegenerative disorders.</p> <p><b>Porseryd et al.</b> [209]: alteration in expression of genes involved in synaptogenesis and synaptic function. In female brains, produced significant effects on pathways connected to the circadian rhythm, cytoskeleton and motor proteins and synaptic proteins. In male brains effects on pathways related to cholesterol biosynthesis and synaptic proteins.</p> <p><b>Preciados et al.</b> [48]: influences NRF1 signaling pathways, and epigenomic multiple networks.</p> <p><b>Zaccaroni et al.</b> [210]: very low doses during development can affect key behavioral traits that are modulated by anxiety.</p>
77	Ethylene thiourea	Herbicide	<p><b>Wang et al.</b> [211]: induced abnormal innervation patterns in the anorectum of fetal rats</p> <p><b>Debbbarh et al.</b> [212]: neurotoxic in utero, increases sensitivity to genetic and environmental risk factors for cell death and apoptosis.</p>
78	Ethylparaben	Antifungal Preservative	<p><b>Merola et al.</b> [213]: provoked behavioral changes including trembling of head, pectoral fins and spinal cord of zebrafish.</p> <p><b>Lynch et al.</b> [214]: displayed significant fear generalization in rats.</p> <p><b>Alward et al.</b> [215]: this aromatase inhibitor reduced the motivation to sing as well as song acoustic stereotypy.</p>
79	Fadrozole	Pharmaceutical	<p><b>Xing et al.</b> [216]: dopamine neuron degeneration and aromatase activity inhibition could be respectively achieved in vivo with treatments with the product in female goldfish.</p> <p><b>Langlois et al.</b> [217]: induced female- and male-biased sexual development on <i>Silurana tropicalis</i> brain mRNA levels, and reduced brain aromatase activity in frogs.</p>
80	Fenbuconazole	Fungicide Organophosphate	<p><b>Hurley et al.</b> [132]: disrupts thyroid hormone excretion.</p> <p><b>Geraldi et al.</b> [218]: affected the acquisition and, mainly, the retention of instrumental conditioning in rats.</p>
81	Fenitrothion	Insecticide	<p><b>Groszek et al.</b> [219]: High concentration of the pesticides was found in adipose tissue and also in the brain. Respiratory failure was the syndrome; and inhibition of acetylcholinesterase activity persisted even for 30 days from poisoning.</p> <p><b>Ram et al.</b> [220]: Neurobehavioral changes in freshwater fish exposed</p>
82	Fenoxy carb	Insecticide	<p><b>Lenkic et al.</b> [221]: allatostatin may be one of the effectors in the brain by which the pesticides inhibits juvenile hormone biosynthesis in cockroach.</p> <p><b>Fertig et al.</b> [222]: permanent sexual dysfunction and mood changes (fatigue, anxiety, depression and suicidal ideation) during treatment with this 5-alpha-reductase inhibitor.</p>
83	Finasteride	Pharmaceutical	<p><b>Traish et al.</b> [223]: Also non-sexual adverse effects such as diabetes, psychosis, depression, and cognitive function.</p> <p><b>Ganzer et al.</b> [224]: sexual libido, ejaculatory disorders, disorders of the penis and testes, cognitive symptoms, and psychological symptoms</p> <p><b>Godinho et al.</b> [225]: toxic interactions with the central nervous system of mammals and lead to memory impairment by modulating the GABAergic system.</p> <p><b>Park et al.</b> [181]: Progressive loss of nigrostriatal dopaminergic neurons induced by inflammatory responses to the pesticide.</p>
84	Fipronil	Insecticide	<p><b>Magalhães et al.</b> [226]: acts on maternal aggressive behavior through GABA(A) receptors.</p> <p><b>Simon-Delso et al.</b> [227]: disrupting neural transmission in the central nervous system of invertebrates, inhibits neuronal receptors.</p> <p><b>Marrs et al.</b> [228]: 4-Aminobutyric acid (GABA) and glycine are inhibitory neurotransmitters and their antagonist, fipronil, is excitatory.</p> <p><b>Golub et al.</b> [229]: provoked greater dendritic spine synapse density in prefrontal cortex of monkeys.</p> <p><b>Hong et al.</b> [230]: induced predominant sympatho-excitation and depressed parasympathetic activity leading to mild hypertension, tachycardia, and impairment of baroreflex function.</p>
85	Fluoxetine	Pharmaceutical	<p><b>Sprowles et al.</b> [131]: Differential effects of perinatal exposure to antidepressants on learning and memory, acoustic startle, anxiety, and open-field activity in rats.</p> <p><b>Yamada et al.</b> [127]: the effects of postnatal treatment on brain masculinization were observed by analysis of male sexual behavior.</p>
86	Flutamide	Pharmaceutical	<p><b>Svensson</b> [231]: induced anxiolytic-like behavior in castrated rats</p> <p><b>Zhang et al.</b> [232]: Effects of neonatal treatment on hippocampal neurogenesis and synaptogenesis correlate with depression-like behaviors in preadolescent male rats.</p>
87	Fonofos	Organophosphate Insecticide	<p><b>Ahmadiani et al.</b> [233]: Anticonvulsant effects on seizures involvement of benzodiazepine receptors.</p> <p><b>GK Sidhu,</b> [234]: known to inhibit acetylcholinesterase activity, not only in insect, but in aquatic and terrestrial organisms leading to nervous abnormalities among others.</p> <p><b>Liu et al.</b> (2018) [235]: Acute formaldehyde exposure induced early Alzheimer-like changes in mouse brain. Provoked the permeability of the blood-brain barrier, activation of astrocyte and microglia, oxidative stress and inflammation.</p> <p><b>Li et al.</b> [236]: effects on anxiety, depression-like behavior and cognition ability which may be associated with alterations in hippocampal glucocorticoid receptors and brain tyrosine hydroxylase levels.</p>
88	Formaldehyde	Solvent	<p><b>Zendehdel et al.</b> [237]: Its neurotoxic effect depend on acetylcholinesterase activity; provoked cholinergic signal reduction in cases of cognitive dysfunction.</p> <p><b>Tulpule et al.</b> [238]: contribute to the impaired cognitive performance and neurodegeneration in diseases.</p>
89	Furan	Solvent	<p><b>Songur et al.</b> [239]: neurotoxic characteristics; neurological diseases.</p> <p><b>Johnston et al.</b> [240]: exhibits a peculiar mode of attack on the central nervous system</p>
90	Galaxolide	Synthetic musk	<p><b>Ayuk-Takem et al.</b> [241]: neurotoxicity may be associated with the inhibition of cellular; polyisoprenylated methylated protein methyl esterase activity; significant risk to individuals predisposed to developing degenerative disorders.</p>
91	Genistein	Isoflavone	

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
92	Hexabromocyclodo-decane	Flame retardant	<p><b>Patisaul HB</b> [242]: involve action at nuclear estrogen receptors, effects on vasopressin, innervation of the lateral septum and other brain regions.</p> <p><b>Luo et al.</b> [243]: pretreatment significantly increased cell viability and protein kinase C activity, decreased the levels of intracellular calcium, and blocked caspase-3 activity in induced cells.</p> <p><b>Román GC</b> [28]: Transient in utero hypothyroxinemia is related to maternal flavonoid ingestion during pregnancy and may provoke autism.</p> <p><b>Lee et al.</b> [244]: mimic the actions and functions of estrogens on brain, two putative pathways; an estrogen receptor-mediated pathway and via the inhibition of tyrosine kinase.</p> <p><b>Lephart et al.</b> [245]: The specific influence of dietary soy phytoestrogens is identified on consumptive, learning and memory, and anxiety-related behaviors.</p> <p><b>Pham-Lake et al.</b> [246]: Impairment in the mesohippocampal dopamine circuit following exposure.</p> <p><b>Wang et al.</b> [247]: main metabolic pathways perturbed, nervous system damage, and developmental disorders.</p> <p><b>Maurice et al.</b> [248]: Short-term effects of a perinatal exposure in rats provoked impairments of early locomotor activity and sensory development.</p> <p><b>Al Mousa et al.</b> [249]: inhibiting reticulum Ca(2+)ATPase in human neuroblastoma cells and induced cells death possibly causing neurological disorders.</p> <p><b>Lilienthal et al.</b> [250]: Effects on dopamine-dependent behavior and brainstem auditory evoked potentials in rats.</p> <p><b>Fu et al.</b> [251]: exhibits through its metabolite a neurotoxic effect by inducing oxidative stress-mediated inflammatory responses.</p> <p><b>Kyriklaki et al.</b> [252]: High exposure during pregnancy reduction in working memory score and reduced cognitive development at preschool age.</p> <p><b>Reed et al.</b> [253]: exposure involved systemic impairment, as well as on nervous system.</p> <p><b>Li et al.</b> [254]: can induce enhanced lipid peroxidation on rats, and the oxidative stress plays an important role in the mechanism of neurotoxicity.</p> <p><b>Goldey et al.</b> [255]: behavioral teratogen, and suggests that human fetuses and suckling infants may be at risk because of the neurotoxic effects of the chemical.</p> <p><b>Nyffeler et al.</b> [256]: neural crest cell migration was inhibited by this toxicant disturbing a key neurodevelopmental process.</p> <p><b>Christen et al.</b> [257]: Strong and dose- dependent inhibition of neurite outgrowth was induced developmental neurotoxicity.</p> <p><b>Hong et al.</b> [258]: induced nigral dopaminergic neuronal loss and Parkinsonism-like movement deficits in mice.</p> <p><b>Moser et al.</b> [259]: perinatal exposure produced neurochemical and persistent neurobehavioral changes, including alterations in spatial learning and memory.</p> <p><b>Kirby et al.</b> [260]: toxic effects of heptachlor epoxide may be responsible for loss of maximal dopamine uptake</p> <p><b>Yamaguchi et al.</b> [261]: effects on calcium mediated transmitter release from brain synaptosomes of rats.</p>
93	Hexachlorobenzene	aromatic	<p><b>Badaeva et al.</b> [262,263]: neurotoxic effects in the postnatal period of ontogeny in the rats.</p> <p><b>Murzakaev</b> [264]: small doses affected central nervous activity.</p> <p><b>Chen et al.</b> [265]: synaptic plasticity and neuro-immune system may be two principal affected areas.</p> <p><b>Kimura et al.</b> [266]: over-activation of aryl hydrocarbon receptor following perinatal dioxin exposure, perturbs neuronal migration and morphological development in mammalian cortex, supporting previous observations of impaired dendritic structure, cortical dysgenesis, and behavioral abnormalities</p>
94	Heptachlor		<p><b>Heptachlor</b> [267]: Neonatal exposure impairs early learning and retention of active avoidance in the rat.</p> <p><b>Heptachlor</b> [268]: neurotoxic profile of tremor.</p> <p><b>Heptachlor</b> [269]: effect on the development of behavioral and/or neural function.</p> <p><b>Andrade et al.</b> [270]: can induce dyshomeostasis, potentially triggering neurodegenerative disorders, such as Alzheimer's disease and Parkinson's disease. Additionally, changes in heme synthesis have been associated with neurodegeneration.</p> <p><b>Chen et al.</b> [271]: exposure in the early stages of neurodevelopment results in long-lasting alterations that ultimately cognitive function and behavior. The prime targets of lead toxicity are the multipotent neural stem cells</p>
95	Heptachlor epoxide		<p><b>Heptachlor epoxide</b> [261]: effects on calcium mediated transmitter release from brain synaptosomes of rats.</p> <p><b>Heptachlor epoxide</b> [262]: neurotoxic effects in the postnatal period of ontogeny in the rats.</p> <p><b>Heptachlor epoxide</b> [263]: small doses affected central nervous activity.</p> <p><b>Heptachlor epoxide</b> [264]: over-activation of aryl hydrocarbon receptor following perinatal dioxin exposure, perturbs neuronal migration and morphological development in mammalian cortex, supporting previous observations of impaired dendritic structure, cortical dysgenesis, and behavioral abnormalities</p>
96	Hexachlorobutadiene	Solvent	<p><b>Hexachlorobutadiene</b> [265]: synaptic plasticity and neuro-immune system may be two principal affected areas.</p> <p><b>Hexachlorobutadiene</b> [266]: over-activation of aryl hydrocarbon receptor following perinatal dioxin exposure, perturbs neuronal migration and morphological development in mammalian cortex, supporting previous observations of impaired dendritic structure, cortical dysgenesis, and behavioral abnormalities</p>
97	Heptachloro-dibenzodioxin	Dioxin	<p><b>Heptachloro-dibenzodioxin</b> [267]: over-activation of aryl hydrocarbon receptor following perinatal dioxin exposure, perturbs neuronal migration and morphological development in mammalian cortex, supporting previous observations of impaired dendritic structure, cortical dysgenesis, and behavioral abnormalities</p>
98	HPTE	Methoxychlor	Not specifically studied (see Methoxychlor):
99	Iodine (I)	Metabolite	<p><b>Iodine (I)</b> [268]: Iodine deficiency as a cause of autism.</p> <p><b>Iodine (I)</b> [269]: effects on motor, sensory, or cognitive function; developmental neurotoxicant.</p>
100	Kepone	Organochlorine	<p><b>Kepone</b> [270]: effects on calcium mediated transmitter release from brain synaptosomes of rats.</p> <p><b>Kepone</b> [271]: neurotoxic profile of tremor.</p> <p><b>Kepone</b> [272]: effect on the development of behavioral and/or neural function.</p> <p><b>Kepone</b> [273]: can induce dyshomeostasis, potentially triggering neurodegenerative disorders, such as Alzheimer's disease and Parkinson's disease. Additionally, changes in heme synthesis have been associated with neurodegeneration.</p> <p><b>Kepone</b> [274]: exposure in the early stages of neurodevelopment results in long-lasting alterations that ultimately cognitive function and behavior. The prime targets of lead toxicity are the multipotent neural stem cells</p>
101	Lead	Heavy metal	<p><b>Lead</b> [275]: wide spectrum of toxic effects, a real threat to the public health, including on the central nervous system</p> <p><b>Lead</b> [276]: lead to imbalance between the pro-oxidant elements and the antioxidants, and induced cognitive dysfunction.</p> <p><b>Lead</b> [277]: The central nervous system is particularly vulnerable. The brain accumulates metals.</p> <p><b>Lead</b> [278]: induces a centrally-mediated sensitization of both autonomic and hypothalamic-pituitary-adrenal (HPA) axis.</p> <p><b>Lead</b> [279]: in combination with ethinyl estradiol reduced brain-derived neurotrophic factor mRNA in the hippocampus resulting in a decline in learning and memory.</p> <p><b>Lead</b> [280]: Long-term administration decreased allopregnanolone levels and altered GABA(A) receptor subunit expression and anxiety-like behavior.</p> <p><b>Lead</b> [281]: block the chloride channels of the GABA-A receptor.</p> <p><b>Lead</b> [282]: has neurotoxic potentials after both acute and chronic exposure.</p> <p><b>Lead</b> [283]: has effects on motor, sensory, or cognitive function modifying behavior.</p>
102	Levonorgestrel	Synthetic Estrogen	
103	Lindane	Organochlorine	
		Insecticide	

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**Table 1** (continued) disruptor

		Class or use	Mechanisms of nervous disruption
104	Linuron	Herbicide	<b>Quintaneiro et al.</b> [278]: adverse effects on neurotransmission and energy production, and interference with hypothalamic-pituitary-thyroid and -adrenal-axis. <b>Lichtensteiger et al.</b> [279]: in antiandrogenic mixtures impacted genes encoding for components of excitatory glutamatergic synapses and genes controlling migration and pathfinding of glutamatergic and GABAergic neurons, as well as genes linked with increased risk of autism spectrum disorders. <b>Schinn et al.</b> [280]: in mixture inhibited swimming activity of juvenile rainbow trout. <b>Richendrfer &amp; Creton</b> [281]: cause abnormalities in behavior and brain size during development, zebrafish larvae had significantly smaller forebrain and hindbrain regions. <b>Salama et al.</b> [282]: affect proliferation, differentiation and viability of cultured neurospheres, Hashjin et al. [283]: induced chronic toxicity and anxiety-like behavior in the male adult mouse. <b>Rastogi et al.</b> [284]: In mixture provoked neurologic self-reported symptoms, headache, watering in eyes, and burning sensation in eye/face, cholinergic symptoms, such as insomnia, headache, muscle cramps, weakness, and anorexia, in children. High frequency of neurologic symptoms may be due to parasympathetic hyperactivity. <b>Valvassori et al.</b> [285]: affects the central nervous system by inhibiting acetylcholinesterase, leading to an increase of acetylcholine in the synaptic cleft, and subsequent activation of cholinergic muscarinic and nicotinic receptors, and impairs aversive-memory retention but not non-associative memory, without affecting anxiety-related behaviors. <b>de Joode et al.</b> [286]: poorer verbal learning outcomes in children, may affect their neurodevelopment. <b>Brody et al.</b> [287]: behavioral dysfunction, notably serotonin-mediated egg-laying behavior in <i>Caenorhabditis elegans</i> .
105	Malathion	Insecticide	<b>Li et al.</b> [288]: potentiation on KCNQ2 potassium channels might be the possible mechanism of this product toxicity in the nervous system. <b>Domico et al.</b> [289]: acute exposure to high doses produces equipotent toxic effects in both dopamine and GABA neurons. <b>Kimura et al.</b> [290]: nerve conduction velocities and postural sway seem to be sensitive indicators of the effects on the central and peripheral nervous system. <b>Lucchini et al.</b> [291]: essential metal that plays a fundamental role for brain development and functioning. Environmental exposure may lead to accumulation in the basal ganglia and development of Parkinson-like disorders. <b>Peres et al.</b> [292]: Various neurotransmitter systems may be impaired, especially dopaminergic, but also cholinergic and GABAergic. <b>Tarale et al.</b> [293]: epigenetic mechanism in product-induced neurotoxicity, development of Parkinson's disease. <b>Zhang et al.</b> [293]: overexposure amplified the role of autophagy in the mechanisms of common neurodegenerative disorders.
106	Mancozeb	herbicide	<b>Wnuk et al.</b> [158]: apoptotic action during early stages of neural development with crucial involvement of retinoid X receptors. <b>Torres-Sánchez et al.</b> [294]: prenatal exposure impaired early child neurodevelopment. <b>Zhang et al.</b> [153]: showed remarkable GR antagonistic properties, disruption of glucocorticoid-responsive genes.
107	Manganese	Heavy metal	<b>Martini et al.</b> [295]: perinatal exposure has an organizational effect on hippocampus-dependent memory and emotional behaviors. <b>Schuh et al.</b> [296]: inhibited brain mitochondrial respiration and increases hydrogen peroxide production and CREB phosphorylation. <b>De Souza et al.</b> [297]: acute and chronic progressive neurologic injury: seizures, myoclonus, ataxia or cerebral oedema, defective neurotransmitter function and abnormal oxidative phosphorylation. <b>Kim &amp; Kang</b> [298]: chronic toxic encephalopathy. <b>Yang et al.</b> [299]: Sub chronically and chronically, principal target PAHsite appears to be the central nervous system. <b>Anger et al.</b> [300]: produce slight neurotoxic effects in fumigation, reduced performance on all cognitive tests.
108	Methylsulfonyl-DDE	DDE Metabolite	<b>Moshitzky et al.</b> [301]: neural inhibition from the brain ( <i>drosophila</i> ) act before farnesoic acid, a precursor of the product.
		Organochlorine	<b>Prestwich et al.</b> [302]: is secreted by the mandibular organs of crustaceans, role partially known. <b>DeLeo et al.</b> [303]: Effect on thyroid hormone action and stress in frog and mammalian culture systems. <b>Ruszkiwicz et al.</b> [79]: potential neurotoxicity
109	Methoxychlor	Insecticide	<b>Broniowska et al.</b> [304]: affected the viability of nerve cells, most likely by enhancing the process of apoptosis.
110	Methyl bromide	Fumigant, Pesticide	<b>Li et al.</b> [305]: reduction of neuronal and muscular development in zebrafish embryos <b>Faass et al.</b> [80]: effect on female sexual behavior and gene expression in sexually dimorphic brain regions after pre- and postnatal exposure in rats. <b>Maerkel et al.</b> [306]: Sex- and region-specific alterations of progesterone receptor mRNA levels and estrogen sensitivity in rat brain <b>Singh et al.</b> [307]: induces neurotoxicity in developing neurons derived from human stem cells by activation of aryl hydrocarbon receptor.
111	Methyl farnesoate	Juvenile Hormone	<b>Puertas et al.</b> [308]: showed a decrease in working memory in children. The deficit found in intellectual function during early childhood suggests that prenatal exposure may have a significant impact on school performance.
112	Methyl triclosan	Triclosan metabolite Product	<b>Shankland</b> [309]: enhanced the release of neurotransmitters. Direct evidence is available on cholinergic and glutaminergic junctions, but other kinds of junctions may be affected. <b>Foran et al.</b> [310]: Auditory hindbrain atrophy and anomalous calcium binding protein expression after neonatal exposure.
113	Methylbenzylidene camphor	UV filter	<b>Sadek et al.</b> [311]: induced neurotoxicity by cholinergic dysfunction, Bcl-2/Bax balance, and antioxidant enzymes gene transcripts in rats <b>Sasaki-Hamada et al.</b> [312]: Changes in hippocampal synaptic functions and protein expression in obese mice.
114	Methylcholanthrene	PAH	
115	Mirex	Organochlorine	
		Insecticide	
116	Monosodium glutamate	Food Additive	

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
117	n-Butylbenzene	Chemical Synthesis Intermediate	Lau et al. [313]: glutamate excitotoxicity has also been linked to chronic neurodegenerative disorders such as amyotrophic lateral sclerosis; evidence for the product excitotoxicity in acute neurologic diseases Chalansonnet et al. [314]: a decrease in the concentrations of free malondialdehyde in brain structures was observed after acute administration of this product. England et al. [315]: exposure during pregnancy and adolescence may contribute to cognitive and behavioral deficits in later life. Exposure during adolescence is associated with deficits in working memory, attention, and auditory processing, as well as increased impulsivity and anxiety. Ferreira et al. [316]: Neuroprotective and neurotoxic effects, modifications of cholinergic transmission. Ciani et al. [317]: neurotoxicity on rat habenulo-interpeduncular cholinergic neurons.
118	Nicotine	Alkaloid	Lee et al. [318]: chronic exposure to low doses linked to the risk of developing cognitive impairment in elderly. Kim et al. [319]: key role in the development of hypertension-related cognitive impairment. Litwa et al. [320]: RXR $\alpha$ , PXR and CAR xenobiotic receptors mediate the apoptotic and neurotoxic actions of the product in mouse hippocampal cells.
119	Nonachlor	Organochlorine Insecticide	Tabassum et al. [321]: potential risk of cognitive, neurochemical and histopathological perturbations; induced toxicity in frontal cortex and hippocampus of rat brain. Jie et al. [322]: inhibited neuronal development and differentiation as indicated by the reduction of the neurotrophic factor GAP-43. Coudere et al. [323]: perinatal exposure induced behavioral and neuro-developmental impairments. Pinna et al. [324]: selective brain steroidogenic stimulant, reduced post-traumatic stress disorder -like behavior in mice
120	Nonylphenol	Formulant	Pinna et al. [325]: facilitated GABA(A) receptor neurotransmission and effectively ameliorate emotional and anxiety disorders and depression. Matsumoto et al. [326]: non-serotonergic mechanism of action in mood and anxiety disorders. Pinna et al. [327]: stereo specifically and selectively increase brain neurosteroid content.
121	Norfluoxetine	Pharmaceutical	Tawara et al [328]: fetal growth may be influenced by maternal total exposure to dioxins
122	Octachlorodibenzo-p-dioxin	Dioxin	Chu et al. [329]: 90-day toxicity in the rat: effects on thyroid. Chu et al. [330]: long-term toxicity in the rat: effects on thyroid.
123	Octachlorostyrene	Chlorinated Aromatic	Ruszkiewicz et al. [79]: neurotoxic effect of active ingredients in sunscreen products.
124	Octyl-methoxycinnamate	UV filter	Axelstad et al. [331]: effects on auditory and neurological development of rat offspring. Bianco et al. [332]: greater accumulation in the cerebral cortex, more accumulation in the cerebellum compared to the mesencephalus and thalamus, with consequences to neural behaviour.
125	Octylphenol	Formulant	Ghisari et al. [333]: negative impact on fetal brain development, resulting in cognitive dysfunctions. Shikimi et al. [334]: promote Purkinje dendritic growth during neonatal life, may be mediated by estrogen receptor in the Purkinje cell.
126	Oxychlordane	Chloridane Metabolite	Kim et al. [335]: role of background exposure in the development of dementia should be explored Kim et al. [336]: greater cognitive decline with aging among elders with high serum concentrations Jain [337]: total serum thyroxine levels had an inverse association with the product.
127	Parathion	Organophosphate Insecticide	Slotkin et al. [56]: produced a net increase in norepinephrine emerged over the course of development in brain region. Liu et al. [338]: effects on endocannabinoid and endocannabinoid-like lipid metabolites in rat striatum. Beard et al. [339]: positively associated with depression in male private pesticide applicators in the agricultural health study.
128	Pendimethalin	Herbicide	Lerro et al. [340]: long-term exposure may alter thyroid function among male pesticide applicators. Campillo et al. [341]: Biomarkers indicative of neurotoxicity and physiological stress in caged clams exposed to a contaminated water containing the product. Pan et al. [342]: thyroglobulin decreased in rats thyroid cells after exposure.
129	Pentachlorobenzene	Chlorinated Aromatic	Den Besten et al. [343]: severe effects on rats thyroid.
130	Pentachloro-nitrobenzene	Herbicide	Hurley PM [132]: disrupted thyroid-pituitary homeostasis Cheng et al. [344]: affected the timing and coordination of development in the central nervous system.
131	Pentachlorophenol	Herbicide, fungicide	Krieg [147]: may act at acetylcholine and $\gamma$ -aminobutyric acid synapses in the central nervous system. Roze et al. [75]: worse coordination, less sensory integrity, worse attention, and worse visuomotor integration. Jorens et al. [345]: increased risk for nasal carcinoma Steinmaus et al. [346]: affected thyroid hormone production during pregnancy and fetal neurodevelopment.
132	Perchlorate	Oxidizer	Brent GA [347]: exposure in pregnancy impacted cognitive outcomes in children Gilbert et al. [348]: developmental exposure altered synaptic transmission in hippocampus of the adult rat. Hossain et al. [349]: may directly activate microglial cells and may contribute to neurodegeneration. Nasuti et al. [350]: decreased levels of dopamine in the striatum, loss of dopaminergic neurons in the substantia nigra pars compacta and cognitive impairments. Motor coordination defects appeared at adult age after early life exposure.
133	Permethrin	Insecticide	Zakirova et al. [351]: persistent neuroinflammation, neurobehavioral and neuropathological cognitive impairment in mouse. Yang et al. [352]: significant effects on the central nervous system.
134	Perfluorodecane sulfonic acid	Perfluoroalkyl substance	Ren et al. (2016) [353]: Binding interactions with thyroid hormone transport proteins and potential toxicological implications.
135	Perfluorohexane sulfonic acid	Perfluoroalkyl substance	Oulhote et al. [354]: High serum concentrations at ages 5- and 7-years, but not prenatally, were associated with parent-reported behavioral problems at age 7. Ren et al. [353]: Binding interactions with thyroid hormone transport proteins and potential neurotoxicological implications.
136	Perfluorononanoic acid	Perfluoroalkyl substance	Jantzen et al. [355]: males exposed showed a reduction in total distance traveled and time of immobility, and an increase in thigmotaxis behavior, aggressive attacks, and preference for the bright.

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
137	Perfluoroctanoic acid	Perfluoroalkyl substance	Acute, embryonic exposure resulted in significant biochemical and behavioral changes in young adult zebrafish. Lien et al. [356]: prenatal exposure was found to associate with neurobehavioral symptoms related to attention deficit hyperactivity disorder among Asian seven-year-old children. Oulhote et al. [354]: sex-dimorphic associations between concentrations and strengths and difficulties. Jantzen et al. [355]: embryonic exposure resulted in significant biochemical and behavioral changes in young adult zebrafish. Oulhote et al. [354]: significant associations were found in regard to hyperactivity, peer relationship, and conduct problems, as well as internalizing and externalizing problems and autism. Ge et al. [357]: could significantly reduce the cell viability and mediate cell apoptosis in HAPI microglia cells of rat.
138	Perfluorooctane sulfonate	Perfluoroalkyl substance	Jantzen et al. [355]: embryonic exposure resulted in significant biochemical and behavioral changes in young adult zebrafish. Oulhote et al. [354]: significant associations were found in regard to hyperactivity, peer relationship, and conduct problems, as well as internalizing and externalizing problems and autism.
139	Perfluorooctanesulfonyl fluoride	Perfluoroalkyl substance	Ren et al. [353]: Binding interactions with thyroid hormone transport proteins and potential neurotoxicological implications.
140	Phorate	Organophosphate	Starks et al. [358]: associated with better verbal learning and memory
		Insecticide	Vandana et al. [359]: obvious effect on cholinesterase enzyme profile of olfactory bulb of mice after systemic administration of low doses for long terms.
141	Picloram	Herbicide	Reddy et al. [360]: decreased neuronal branching and degenerating neurons, probably through a mitochondrial pathway. Meshchakova et al. [361]: provoked functional disorders risk connected with cardiovascular and nervous diseases.
142	Polyvinylchloride	Polymer; PVC	Podoll et al. [362]: acute intoxication resulted in vertigo, nausea and headache up to a narcotic effect. In patients with chronic occupational exposure, neurological disturbances included sensory-motor polyneuropathy, trigeminal sensory neuropathy, slight pyramidal signs and cerebellar and extrapyramidal motor disorders. Psychiatric disturbances present as neurasthenic or depressive syndromes. Sleep disorders and disorders of sexual functions are frequently encountered. Urmann et al. [363]: neurodifferentiating potential of the product.
143	8-Prenylnaringenin	Prenylflavonoid	Bagatin et al. [364]: panicolytic effects in rats with generalized anxiety and panic disorders. Oberbauer et al. [365]: promote neuronal differentiation and neurite outgrowth and are neuroprotective.
144	Prochloraz	Fungicide	Vinggaard et al. [366]: agonizes the aryl hydrocarbon receptor and inhibits aromatase activity. Ghisari et al. [333]: inhibitory effect on rat pituitary cell growth increasing the risk or a negative impact on fetal brain development, resulting in cognitive dysfunctions.
145	Procymidone	Fungicide	Xiang et al. [367]: potential to disrupt thyroid homeostasis, agonistic effects.
146	Prodiamine	Herbicide	Radio et al [368]: selectively increased neurite outgrowth. Gilbert et al. [369]: an impaired capacity for hippocampal neurogenesis may contribute to impairments in synaptic plasticity and cognitive deficits
147	Propylthiouracil	Thyroid inhibitor	Koromilas et al. [370]: inhibition of hypothalamic, pontine and cerebellar NaK-ATPase; a major marker of neuronal excitability and metabolic energy production as well as a regulator of important systems of neurotransmission. Koromilas et al. [371]: impairs neurochemical mechanisms that could be involved in the way clinical hypothyroidism could affect optimal neurodevelopment and, ultimately, cognitive function. Vanek et al. [372]: induced central nervous system vasculitis presenting as confusion. Yang et al. [58]: Benzo[a]Pyrene (BaP) exposure caused the disruption of glutamate (Glu) neurotransmitter transmission by decreasing the level of Glu, reducing the expression of Glu receptors, enhancing the level of SNAP-25, and neurotoxicity. Chepelev et al. [59]: BaP correlates with impaired learning and memory in adults, and poor neurodevelopment in children. Neurotoxic endpoints and DNA damages are more sensitive than cancer endpoints. Chen et al. [373]: behavioral impairments resulting from postnatal BaP exposure are potentially long-lasting in rats.
148	Pyrene	Polycyclic Aromatic Hydrocarbon	Wormley et al. [374]: neurobehavioral deficits; gestational exposure to BaP and dioxin reduced specific indices of learning and memory, including hippocampal-based synaptic plasticity mechanisms. Takeda et al. [375]: the fetal exposure of mice to diesel exhaust affected the emotional behaviors associated with the serotonergic and dopaminergic systems in the brain
149	Pyrimethanil	Fungicide	Hurley PM [132]: disrupt thyroid-pituitary homeostasis only
150	Pyriproxyfen	Juvenile hormone analog	Truong et al. [376]: induced craniofacial defects in zebrafish, and adverse behavioral effects. Fourrier et al. [377]: changes in social integration, acceptance by nestmates and social behaviors performance in bees.
151	Resorcinol	Disinfectant, Chemical intermediate	Motonaga et al. [378]: inhibit thyroid peroxidase to cause developmental toxicity and neurotoxicity. Román [28]: transient maternal hypothyroxinemia resulting from dietary and/or environmental exposure to this antithyroid agent.
152	Roundup	Main herbicide worldwide	Defarge et al. [20]: its formulates decrease aromatase activity below toxic levels. Gress et al. [379]: the product altered locomotor activity in rats. Modesto et al [380]: it inhibits acetylcholinesterase in fish brain.
153	Sertraline	Psychotropic	Lee et al. [381]: the product is used for trauma-focused psychotherapies Frölich et al. [382]: selective serotonin reuptake inhibitor, which has demonstrated efficacy on neuropsychiatric behavioral symptoms in general.
154	Short chain chlorinated paraffins	Flame retardant; plasticizer	Liu et al. [123]: exposure could alter gene expression in the hypothalamic-pituitary-thyroid axis and thyroid hormone levels. Wyatt et al. [383]: potent peroxisome proliferators; high dose shows a depressed plasma thyroxine level, with increase in thyroid stimulating hormone
155	2,4,5-T (in Agent Orange)	Herbicide	

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
156	Tamoxifen	Pharmaceutical	Gunderson and Daroff [384]: epilepsy and later on all effects of brain injury and post-traumatic stress disorders. St Omer et al. [385]: together with 2,4D, increased significantly the concentration of norepinephrine in whole developing brain and increased dopamine. Yi et al. [144]: increased the prevalence of endocrine disorders, especially in the thyroid and pituitary gland; and increased various neurologic diseases. Denk et al. [386]: granular neurons of the olfactory bulb and dentate gyrus, vascular cells and ependymal cells throughout the brain, and peripheral sensory neurons are modified by this treatment. Boeke et al. [387]: Cognitive domains that rely on verbal abilities (verbal memory and fluency) seem to be at risk for deterioration after treatment. Park et al. [388]: induced the loss of both zebrafish neuromasts and hair cells in the rat cochlea in a dose-dependent manner. Chen et al. [389]: induced apoptotic cell death, delayed cranial motor neuron development, inhibited primary motor neuron development and loosed muscle fiber during the early development in zebra fish. Jarema et al. [390]: may have developmental or pharmacological effects on the vertebrate nervous system. Wojtowicz et al. [391]: decreased the expression of PPAR-γ protein in neocortical neurons; and the mechanism of action also induced apoptotic and neurotoxic effects. Holmes et al. [392]: testosterone increased the expression of COX2 and apoptosis in dopamine neurons, increased incidence of Parkinson's disease in men compared with women.
157	Tetrabromo-bisphenol A	Flame retardant	Cunningham et al. [393]: induces dopaminergic neurotoxicity via caspase-3-dependent activation of protein kinase C-delta. Xu et al. [394]: this dioxin-like compound suppresses acetylcholinesterase activity via transcriptional downregulation in vitro. Pelclova et al. [395]: neurological and neurophysiological findings in workers with chronic intoxication 50 years after exposure. Xu et al. [394]: this dioxin suppresses acetylcholinesterase activity via transcriptional downregulations in vitro. Sánchez-Martín et al. [396]: aryl hydrocarbon receptor-dependent induction of apoptosis by the product in cerebellar granule cells from mouse.
158	testosterone	Natural hormone	Kato et al. [397]: reduction of thyroid hormone levels by different mechanisms. Chen et al. [398]: inhibition of UDP-glucuronosyltransferases.
159	Tetrachloro-dibenzofuran	Chlorinated dioxin	Dos Reis-Lunardelli et al. [399]: can alter animal behavior and learning and memory in rats. Zamoner et al. [400]: reorganizes the cytoskeleton of glial cells through Gfap phosphorylation and Rhoa-dependent mechanisms.
160	Tetrachloro-dibenzo-p-dioxin	Chlorinated dioxin	Hurley PM [132]: disrupts thyroid-pituitary homeostasis only. Calcium et al. [401]: congeners products showed a strong inhibitory effect on the otic system development.
161	PCB methyl sulfones	PCB metabolite	Waritz et al. [402]: increases the occurrence of two thyroid tumors and increased excretion of thyroid hormones. Kodavanti et al. [403]: inhibits calmodulin activated adenylate cyclase in rat brain.
162	Tetraiodothyronine	Natural hormone *	Brunström. [404]: affected the growth of the chicks and had neurotoxic effects.
163	Thiazopyr	Herbicide	Leonetti et al. [405]: accumulate in the placenta and potentially alter thyroid hormone function in a sex-specific manner.
164	Toxaphene	Organochlorine	Ishihara et al. [409]: induces oxidative neuronal injury.
		Insecticide	Frye et al. [189]: effects through the aryl hydrocarbon receptor, the peroxisome proliferator-activated receptor and the retinoid X receptor, signal transduction pathways, calcium influx and/or neurotransmitter receptor.
165	2,4,6-Tribromophenol	BFR, Natural product	Kotake [410]: neurotoxic, induces behavioral abnormalities and toxic to the developing central nervous system through AMPA receptor subunit. Yeung [411]: neurotoxicity inducing anxiety in man.
166	Trenbolone	Anabolic steroid	Da Broi et al. [412]: produces pleasant inebriating effects with rapid dissipation, followed by central nervous system depression, coma.
167	Tributyltin	Fungicide	Kang et al. [413]: provokes chronic central nervous system disorders and peripheral neuropathy. Chiu WA et al. [414]: carcinogenic to humans by all routes of exposure and toxic to the central nervous system.
168	Trichloroethylene	Chlorinated solvent	Bale et al. [415]: interacts directly with several different classes of neuronal receptors by generally inhibiting excitatory ions receptors/channels and potentiating the function of inhibitory receptors/channels.
169	Trichlorophenol	Fungicide	Xu et al [416]: urinary levels increased risk of attention deficit hyperactivity disorder among school-aged children.
170	Triclocarban	Antibacterial agent	Dong et al. [417]: altered expression of proteins involved in nervous system development. Barros et al. [418]: modified chronically female amphipod Gammarus behavior.
171	Triclosan	Antibacterial agent	Wu et al. [419]: inhibited iodide uptake, but had differential effects on the expression of thyroid hormone synthesis-related genes and the activity of thyroid peroxidase. Ruszkieiewicz et al. [79]: few neurotoxic effects.
172	Tri-iodothyronine	Natural thyroid hormone	Szychowski et al. [420]: activates aryl hydrocarbon receptor-dependent apoptosis and affects Cyp1a1 and Cyp1b1 expression in mouse neocortical neurons. Wu et al. [419]: inhibited iodide uptake, but had differential effects on the expression of thyroid hormone synthesis-related genes and the activity of thyroid peroxidase.

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**Table 1 (continued)**

Nb	Endocrine disruptor	Class or use	Mechanisms of nervous disruption
173	Triphenyl phosphate	Flame retardant	<p>Noda [421]: impairs glial function as well as neuronal function and thus disturb the brain, which may cause mental disorders.</p> <p>Zhang et al. [422]: affects feeding behavior.</p> <p>Ortiga-Carvalho et al. [423]: regulates by negative feedback the hypothalamus pituitary-thyroid axis.</p> <p>Sun et al. [424]: Developmental neurotoxicity in early life stages of Japanese medaka.</p> <p>Kim et al. [425]: up-regulated the expression of the genes related to the metabolism, transport, and elimination of thyroid hormones.</p> <p>Jarema et al. [390]: produced behavioral effects; may have developmental or pharmacological effects on the vertebrate nervous system.</p> <p>Tanaka et al. [426]: induced shrinkage and increased packing densities, axonal and terminal degenerations in the developing visual system of the ferret.</p> <p>Kotake [410]: potent inhibitor of mitochondrial ATP synthase, neurotoxic and induces behavioral abnormality.</p> <p>Karpiak et al. [427]: disrupts components of glutamate homeostasis in rat astrocyte cultures.</p> <p>Lin et al. [428]: triphenyltin acetate may cause cellular dysfunction of brain without structural damage and provoked demyelinated neuropathy.</p> <p>Attahiru et al. [429]: produced significant central nervous system and respiratory depressions and produced brain congestion.</p> <p>Lehotzky et al. [430]: crosses the blood-brain barrier, and induces diminished plasticity of the central nervous system.</p> <p>Belovicova et al. [431]: stressed rat dams had lowered hippocampal neurogenesis, while the product treatment reversed this lowering.</p> <p>Singh et al. [432]: induced ROS-mediated apoptotic neurodegeneration in fetal neocortex, and neurobehavioral sequelae in rat offspring.</p> <p>Pinzani et al. [433]: it is an antidepressant that selectively inhibits serotonin reuptake and is a norepinephrine inhibitor.</p> <p>Gillette et al. [434]: Debilitating effects were seen at all levels of the phenotype, including physiology, behavior, brain metabolism, gene expression, and genome-wide transcriptome modifications in specific brain nuclei, and transgenerational effects.</p> <p>León-Olea et al. [13]: can lead to severe and widespread neuroendocrine disruptions in discrete brain regions, including the hippocampus, amygdala, and hypothalamus, resulting in behavioral changes in a wide range of species.</p> <p>Skinner et al. [435]: Transgenerational epigenetic programming of the brain transcriptome and anxiety behavior are modified by the product.</p> <p>André et al. [436]: learning deficits are expressed following perinatal exposure.</p> <p>Makowska et al. [437]: affects calcitonin gene related peptide-like immunoreactive neurons in the pig enteric nervous system.</p> <p>Obremski et al. [438]: induced changes in the lymphoid tissue and mucosal nerve fibers in the porcine ileum.</p> <p>Venkataramana et al. [439]: showed a marked suppressive effect on the neuronal gene expression; oxidative stress is the main upstream signal leading to increased neurotoxicity due to the product.</p>
174	Triphenyltin	Fungicide	<p>Attahiru et al. [429]: produced significant central nervous system and respiratory depressions and produced brain congestion.</p> <p>Lehotzky et al. [430]: crosses the blood-brain barrier, and induces diminished plasticity of the central nervous system.</p> <p>Belovicova et al. [431]: stressed rat dams had lowered hippocampal neurogenesis, while the product treatment reversed this lowering.</p> <p>Singh et al. [432]: induced ROS-mediated apoptotic neurodegeneration in fetal neocortex, and neurobehavioral sequelae in rat offspring.</p> <p>Pinzani et al. [433]: it is an antidepressant that selectively inhibits serotonin reuptake and is a norepinephrine inhibitor.</p> <p>Gillette et al. [434]: Debilitating effects were seen at all levels of the phenotype, including physiology, behavior, brain metabolism, gene expression, and genome-wide transcriptome modifications in specific brain nuclei, and transgenerational effects.</p> <p>León-Olea et al. [13]: can lead to severe and widespread neuroendocrine disruptions in discrete brain regions, including the hippocampus, amygdala, and hypothalamus, resulting in behavioral changes in a wide range of species.</p> <p>Skinner et al. [435]: Transgenerational epigenetic programming of the brain transcriptome and anxiety behavior are modified by the product.</p> <p>André et al. [436]: learning deficits are expressed following perinatal exposure.</p> <p>Makowska et al. [437]: affects calcitonin gene related peptide-like immunoreactive neurons in the pig enteric nervous system.</p> <p>Obremski et al. [438]: induced changes in the lymphoid tissue and mucosal nerve fibers in the porcine ileum.</p> <p>Venkataramana et al. [439]: showed a marked suppressive effect on the neuronal gene expression; oxidative stress is the main upstream signal leading to increased neurotoxicity due to the product.</p>
175	Venlafaxine	Psychotropic	<p>Singh et al. [432]: induced ROS-mediated apoptotic neurodegeneration in fetal neocortex, and neurobehavioral sequelae in rat offspring.</p> <p>Pinzani et al. [433]: it is an antidepressant that selectively inhibits serotonin reuptake and is a norepinephrine inhibitor.</p> <p>Gillette et al. [434]: Debilitating effects were seen at all levels of the phenotype, including physiology, behavior, brain metabolism, gene expression, and genome-wide transcriptome modifications in specific brain nuclei, and transgenerational effects.</p> <p>León-Olea et al. [13]: can lead to severe and widespread neuroendocrine disruptions in discrete brain regions, including the hippocampus, amygdala, and hypothalamus, resulting in behavioral changes in a wide range of species.</p> <p>Skinner et al. [435]: Transgenerational epigenetic programming of the brain transcriptome and anxiety behavior are modified by the product.</p> <p>André et al. [436]: learning deficits are expressed following perinatal exposure.</p> <p>Makowska et al. [437]: affects calcitonin gene related peptide-like immunoreactive neurons in the pig enteric nervous system.</p> <p>Obremski et al. [438]: induced changes in the lymphoid tissue and mucosal nerve fibers in the porcine ileum.</p> <p>Venkataramana et al. [439]: showed a marked suppressive effect on the neuronal gene expression; oxidative stress is the main upstream signal leading to increased neurotoxicity due to the product.</p>
176	Vinclozolin	Fungicide	<p>André et al. [436]: learning deficits are expressed following perinatal exposure.</p> <p>Makowska et al. [437]: affects calcitonin gene related peptide-like immunoreactive neurons in the pig enteric nervous system.</p> <p>Obremski et al. [438]: induced changes in the lymphoid tissue and mucosal nerve fibers in the porcine ileum.</p> <p>Venkataramana et al. [439]: showed a marked suppressive effect on the neuronal gene expression; oxidative stress is the main upstream signal leading to increased neurotoxicity due to the product.</p>
177	Zearalenone	Mycotoxin	<p>Obremski et al. [438]: induced changes in the lymphoid tissue and mucosal nerve fibers in the porcine ileum.</p> <p>Venkataramana et al. [439]: showed a marked suppressive effect on the neuronal gene expression; oxidative stress is the main upstream signal leading to increased neurotoxicity due to the product.</p>

Each chemical compound or pollutant has been numbered (Nb) out of 177 known endocrine disruptors; its name was associated with the key word “nervous” or “neurotoxicity” or “cognitive” or “behavio(u)r” on PubMed data bank, or eventually on Google Scholar. When the number of references per compound were too numerous, “or” was excluded in order to directly associate the keywords. If more than 20 references were found to be published, “review” was added to the keywords and cited as a reference. Finally, a maximum of five references were indicated, focusing on the most recent research in humans or mammals, without excluding other models. The mechanisms of nervous disruption could be direct, on the neurons or the nervous system, or indirect, through endocrine disruption interfering with neurodevelopment or nervous system functioning, including thyroid regulation. PAH, polycyclic aromatic hydrocarbon; PCB, polychlorobiphenyl; PBB, polybromobiphenyl; PBDE, polybrominated diphenyl ether; PFAS, perfluoroalkyl substances.

### Declaration of Competing Interest

The authors Seralini & Jungers declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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