

# 1 Persistent Organic Pollutants- A Silent Threat to the Agro-ecosystem and Surrounding 2 Environment

3

## 4 ABSTRACT

5 Persistent organic pollutants (POPs) are a class of hazardous, bio-accumulative, biomagnified, and  
6 extremely persistent chemicals with plausible extended off-site mobility in the environment or agro-  
7 ecosystem. These constitute a wide variety of chemicals, their sources may be either natural or  
8 anthropogenic. These are contaminants of emerging concerns for researchers as well as environmentalists.  
9 These contaminants are supposed to cause toxicity in terms of their carcinogenic, genotoxic, **neurotoxic**,  
10 and endocrine disruptors nature. Therefore, it is very important to discuss various kinds of POPs present  
11 in the environment and their detection methods. Various extraction and analytical techniques have been  
12 strategized to determine the POPs in soil and food samples either quantitatively or qualitatively. The  
13 literature lacks a comprehensive review over various POPs present in the environment, different  
14 techniques of their extraction and analysis, along with their toxicity in agro-ecosystem. The present  
15 review is expected to fulfill this gap by considering all the necessary aspects discussed above.

16

17 **Keywords:** Persistent organic pollutants, Agro-ecosystem, Pollutants, Carcinogenic, extraction  
18 techniques

19

## 20 1. Introduction

21 With the rapid urbanization and industrialization, a plethora of environmental contamination issues have  
22 been unveiled and have become increasingly serious around the world [1,2]. A variety of pollutants  
23 present in the environment and identified by the scientific community are considered highly toxic to  
24 living bodies and environment. Some of these are resistant to environmental degradation (physical,  
25 biochemical and biological phenomena) and can exist for a very long time [3]. Persistent organic  
26 pollutants (POPs) are the pollutants that are existing in our environment for an extended period of time.  
27 POPs include polychlorinated biphenyls (PCBs), polychlorinated dioxins and dibenzofurans (PCDFs- as a  
28 single entry), organochlorine insecticides, polycyclic aromatic hydrocarbons (PAHs), and many more.  
29 These pollutants have been present for a long time and travel from one location to another. They have  
30 also been reported in places where they have never been used, like the earth's **Polar Regions** [4].

31 Many reports are available in the literature which justified that POPs can disrupt the endocrine  
32 system and are often called as endocrine disruptors. Different hazardous outcomes have been observed  
33 such as population declines, reproductive impairment, embryonic deformities, and metabolic and  
34 behavioral changes in various species including eagles, alligators, and cormorants during exposure to

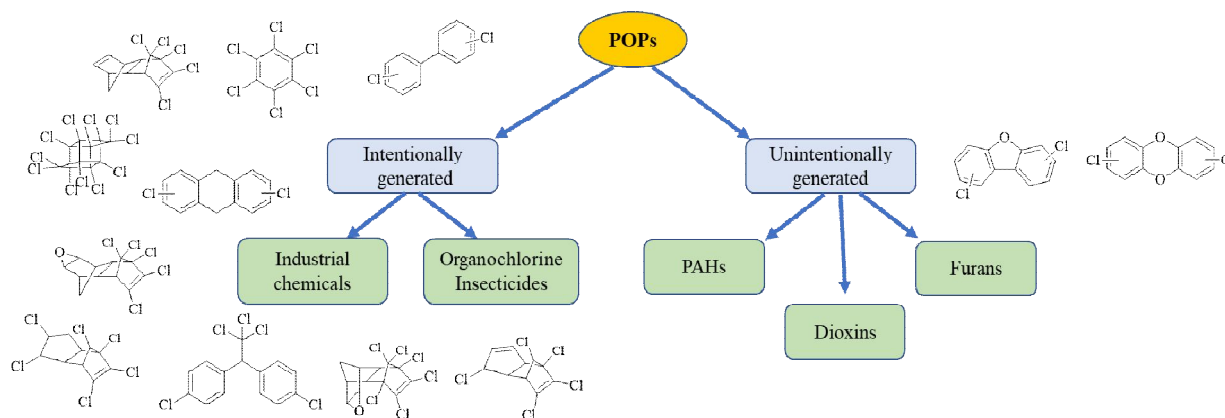
35 POPs. Therefore, there is a big challenge for the scientific community to successfully detect POPs in  
 36 agro-ecosystem and environmental samples, so that better strategies could be designed to remove them  
 37 from the environment. Hence, the present review will focus all types of POPs observed till date, their  
 38 toxicity behaviors, extraction techniques and the methods of their analysis.

## 39 2. Persistent organic pollutants

40 Various pollutants are constantly contaminating the earth's ecosystem and primarily agro-ecosystem.  
 41 POPs have drawn significant attention over the past several decades. These are mainly featured by their  
 42 abilities to resist degradation, persist longer and bioaccumulate. These molecules travel great distances,  
 43 withstand degradation, and bioaccumulate. **Owing** to their unique chemical structures and  
 44 physicochemical properties. Twelve POPs (commonly called the original 'dirty dozens' or 'legacy POPs')  
 45 were initially identified during the Stockholm Convention and proposed ban on their usage or source [5].  
 46 Later, certain non-chlorinated compounds were also added to the list such as perfluorooctanesulfonate  
 47 (PFOS), per and poly-fluoroalkyl substances (PFASs), brominated flame retardants (BFRs), and other  
 48 perfluorinated **compound** (PFCs).

## 49 3. Classification and types of POPs

50 There are two main categories of POPs on the basis of the source from which they originate. One is  
 51 intentionally produced and the other one is unintentionally produced (Fig. 1). Considering the main group  
 52 of unique 12 POPs include 10 intentionally generated pollutants i.e. aldrin, chlordane, DDT, dieldrin,  
 53 endrin, heptachlor, hexachlorobenzene (HCB), mirex, toxaphene, and polychlorinated biphenyls (PCBs)  
 54 and 2 unintentionally generated contaminants- polychlorinated dibenzo-p-dioxins (PCDDs) and  
 55 polychlorinated dibenzofurans (PCDFs) [6].



65 **Fig. 1: Types of POPs with examples**

### 67 3.1. Intentionally generated POPs

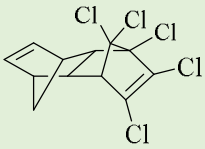
68 These are mostly chlorinated compounds and have highly lipophilic organic moieties linked with chlorine  
 69 atoms. These compounds are generally known as organ chlorine compounds (OCs) and are produced as a  
 70 result of different chemical reactions that involve chlorine. For example, dichlorodiphenyltrichloroethane  
 71 (DDT) and polychlorinated biphenyls (PCBs). They can be classified into two types that are  
 72 organochlorine pesticides and industrial chemicals [7].

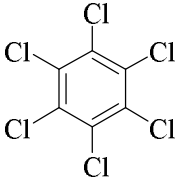
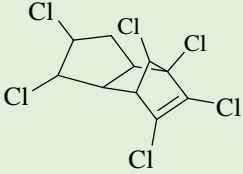
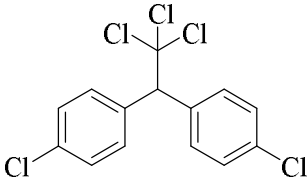
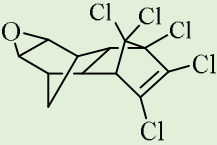
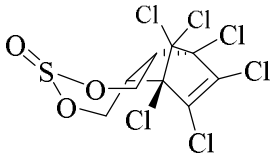
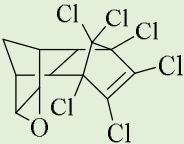
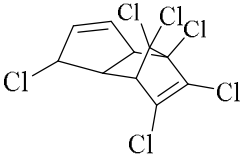
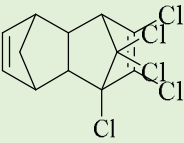
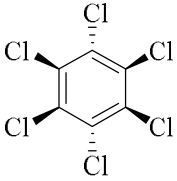
### 73 3.1.1. Organochlorine pesticides (OCPs):

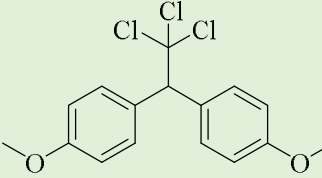
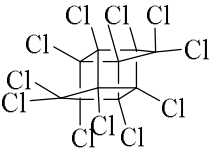
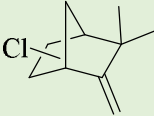
74 OCPs are a class of substances that are commonly employed as pesticides and due to their prolonged  
 75 persistence, these are also categorized under POPs. They may be insecticides, fungicides, herbicides,  
 76 rodenticides, nematocides, and acaricides. For example, aldrin, dieldrin, heptachlor, and endrin. They are  
 77 mostly man-made synthetic organic compounds but some are natural say for example nicotine. Some  
 78 OCPs along with their chemical structure, usage, toxicity values and WHO recommendations based on  
 79 the toxicity values are listed in Table 1. These are considered as the most widely used pesticides in  
 80 developing countries of Asia [8].

81 Approximately 80% of pesticides used in agriculture are released into the environment by  
 82 volatilization, runoff, infiltration, transit down the food chain, and other ways. Therefore, the use of OCPs  
 83 has been prohibited in many countries for a considerable period, but their residues are still causing a  
 84 significant impact on the environment and the ecosystems. These may be neurotoxic, genotoxic, and  
 85 carcinogenic in nature. OCPs when applied in the fields, get strongly adsorbed to the soil particles as  
 86 these are hydrophobic in nature with very high adsorption coefficients. Their residues can be detected on  
 87 the surface layers of soil after adsorption without draining into the soil profile, allowing them to survive  
 88 in the soil for months to years [9].

89 **Table 1: Common OCPs along with their chemical structures, usage, half-life, toxicity values, and**  
 90 **recommendations by WHO**

Sr No	Chemical Name	Structure	Use	Persistence Half-life	Toxicity in rats (mg/kg)	WHO Recommendation on the basis of LD50
1	Aldrin		Insecticide	High 4–7 years	Oral: 39 to 60 Dermal: 100	Highly hazardous

2	Benzene hexachloride (BHC)		Acaricide Insecticide Rodenticide	High 3 – 6 years	Oral: 10,000	Moderately hazardous
3	Chlordane		Insecticide	Very High 10 years	Oral: 200-700 Dermal: 530-690	Moderately hazardous
4	DDT		Acaricide Insecticide	High 2-15 years	Oral: 113-130 Dermal: 2510	Moderately hazardous
5	Dieldrin		Insecticide	Moderate 9 months	Oral: 46 Dermal: 50-120	Highly hazardous
6	Endosulfan		Insecticide	Moderate 35-150 days	Oral: 18 to 220	Highly hazardous
7	Endrin		Avicide insecticide	Moderate- High 1-12 Years	Oral: 3 Dermal: 15	Highly hazardous
8	Heptachlor		Insecticide	High 2 years	Oral: 40- 220 Dermal: 119-320	Highly – Moderately hazardous
9	Isobenzan		Insecticide	High 2.8 years	Oral: 4.8	Highly hazardous
11	Lindane		Rodenticide Acaricide Insecticide	Moderate 15 months	Oral: 88 – 270	Moderately hazardous

12	Methoxychlor		Insecticide	Moderate < 120 Days	Oral: 5000– 6000	Less hazardous
13	Mirex		Insecticide	Very High 10 years	Oral: 600– 740	Less hazardous
14	Toxaphene (Camphechlor)		Acaricide Insecticide	Very High 11 Years	Oral: 80– 293	Slightly hazardous

91 Source: Pesticides properties database (PPDB)

### 92 3.1.2. Industrial chemicals:

93 Thousands of chemicals are being produced in the industries and are used worldwide in large quantities  
 94 for various preparations such as dyes, resins, plasticizers, antioxidants, surfactants, food preservatives,  
 95 sanitizers, and detergents. As a result, these harmful compounds are releasing in the environment  
 96 continuously, especially in wastewater discharged from industries or domestic outlets. Till date, these  
 97 pollutants have been reported in surface water and wastewater. Some of these have been grouped into the  
 98 priority pollutants as they cause serious health issues to aquatic organisms. Polychlorinated biphenyls  
 99 (PCBs) and per and poly-fluoroalkyl substances (PFASs) are the most important types of industrial  
 100 chemicals found extensively in the agro-ecosystem and causing health hazards.

#### 101 3.1.2.1. Polychlorinated biphenyls (PCBs)

102 PCBs have been discovered in water, sediments, avian tissue, and fish tissue all across the world. These  
 103 substances are classified as special wastes because they include 2-10 chlorine atoms linked to the  
 104 biphenyl molecule. This class also includes monochlorinated biphenyls (biphenyl molecules with one  
 105 chlorine atom added). The basic chemical skeleton of chlorinated biphenyls and some commonly found  
 106 PCBs are shown in Table 2. PCBs were having numerous industrial applications and were used  
 107 extensively around 1960-80 as insulating condensers, heat exchangers, plasticizers, flame retardants, fire-  
 108 resistant transformers, papers, asbestos, etc. Their remains are still present at industrial sites and have  
 109 been found in building doors and windows, as well as in paints used in swimming pools and garages [10].  
 110 It is believed that over 200 commercial PCB-containing products are now accessible on the market.

111 **Table 2: Some previously reported PCBs along with their chemical structures**

PCB 10 <sup>a</sup>	PCB 11 <sup>b</sup>	PCB 24 <sup>a</sup>
PCB 30 <sup>a</sup>	PCB 62 <sup>a</sup>	PCB 65 <sup>a</sup>
PCB 77 <sup>b</sup>	PCB 95 <sup>b</sup>	PCB 116 <sup>a</sup>
PCB 118 <sup>b</sup>	PCB 126 <sup>b</sup>	PCB 153 <sup>b</sup>

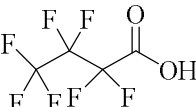
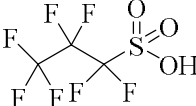
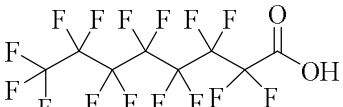
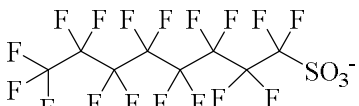
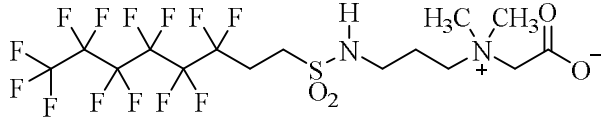
112 a[11], b[12]

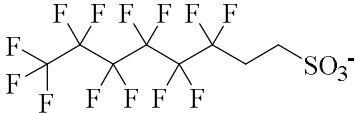
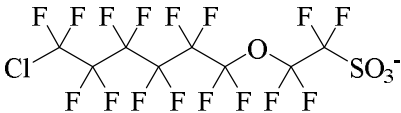
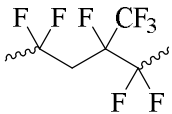
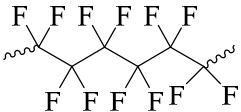
### 113 3.1.2.2. Per and poly-fluoroalkyl substances

114 For more than 50 years, PFASs have been produced and are widely present in the environment. These are  
 115 non-stick, non-wetting, non-toxic, fire-resistant, and heat resistant. As a result, they have been used in  
 116 nearly every aspect of labor, from cookware to firefighting foams, paper coatings, and textiles, and so on  
 117 [13]. These are organ fluoride compounds made up of oligomers and polymers in which the hydrogen  
 118 atoms in the hydrocarbon skeleton have been replaced by fluorine atoms. Per fluorinated sulfonic acids  
 119 (PFSA), perfluoro carboxylic acids (PFCAs), high-molecular-weight fluoropolymers, and low-  
 120 molecular-weight perfluoro alkanolamines are the four categories. Out of these, PFSA and PFOA, have  
 121 received the most attention. The most commonly discovered PFASs are mentioned in Table 3. They are  
 122 both hydrophobic and lipophobic in nature, and they contain a CF bond, which is one of the strongest  
 123 chemical connections and provides these molecules with extremely high chemical and thermal stability.  
 124 Due to these reasons, they are resistant to the typical environmental degradation processes like photolysis,  
 125 hydrolysis, and atmospheric photooxidation, thereby persisting and bioaccumulating in the environment

126 for years [14]. The National Toxicology Program has also conducted research on PFOA, other PFCAs  
 127 and PFSA's in order to better understand their toxicity and persistence in human blood [15]. PFOA-  
 128 contaminated water has been shown to have a negative impact on mammary gland development in mice  
 129 [16]. Another study found that PFOS exposure can influence the nervous and endocrine system in rats;  
 130 however, their mechanisms are yet to be established [17].

131 **Table 3: Different types of well-known PFASs, their chemical structures, and sources.**

Sr No.	Molecular Structure	Sources	References
1	 <p>Perfluorobutanoic Acid (PFBA)</p>	Stain repellants, food packaging carpets	[18]
2	 <p>Perfluorobutane sulfonic Acid (PFBS)</p>	Fire retardant foams, metal coating, mist suppressant	[19]
3	 <p>Perfluorooctanoic acid (PFOA)</p>	Fire retardant foams, ceramic glaze	[20]
4	 <p>Perfluorooctanesulfonic acid (PFOS)</p>	Water purification, non- stick utensils, textile industry	[21]
5	 <p>6:2 Fluorotelomer sulfonamide alkyl betaine (FTAB)</p>	Cleansing products, metal coating, textile industries	[22]

6	 <p>6:2 Fluorotelomer sulfonate (FTSA)</p>	Chemically resilient components, rug coverings	[23]
7	 <p>Chlorinated polyfluorinated ether sulfonate (F53-B)</p>	General waterproofing lubricants, gaskets power-utility stoppers	[24]
8	 <p>Viton A/B</p>	Fire retardant foams, metal coating, mist suppressant	[25], [26]
9	 <p>Polytetrafluoroethylene (PTFE)</p>	Fire retardant foams, metal coating	[27]

132

### 133 3.2. Unintentionally produced POPs

134 These are mostly undesired by-products of combustion or other chemical reactions that take place in the  
 135 presence of chlorine or chlorinated substances. Polycyclic aromatic hydrocarbons (PAHs), dioxins, and  
 136 furans are the three primary categories.

#### 137 3.2.1. Polycyclic aromatic hydrocarbons (PAHs)

138 Both natural and anthropogenic sources release PAHs into the environment [28]. The phenomena  
 139 including incomplete burning of fuels, garbage, some organic substances such as tobacco and plant  
 140 material, forest fires, and volcanic eruptions lead to the exposure of PAHs to the environment. This is a  
 141 large class of chemical compounds that include two or more fused aromatic rings, ranging from two-ring  
 142 naphthalene and its derivatives to complex structures with up to 10 rings. PAHs with up to six rings are  
 143 known as ‘small’ PAHs, while those having more than six aromatic rings are called ‘large’ PAHs [29].  
 144 Some of the PAHs are posing a serious threat to human health as these can be carcinogens, mutagens, and  
 145 teratogens as well. Because of their physicochemical properties, PAHs are highly mobile in the

146 environment, allowing them to spread through air, soil, and water bodies, where their presence is  
 147 ubiquitous. Some of the commonly found PAHs and their chemical structures are shown in Table 4.

148 **Table 4: Pyrogenic and petrogenic PAHs reported in the literature**

149

Pyrogenic PAHs			
Naphthalene	Acenaphthylene	Acenaphthene	Fluorene
Anthracene	Phenanthrene	Fluoranthene	Pyrene
Chrysene	Benz[a]anthracene	Benzo[b]fluoranthene	Benzo[k]fluoranthene
Benzo[a]pyrene	Indeno[1,2,3-cd]pyrene	Benzo[g,h,i]perylene	Dibenz[a,h]anthracene
Pyrogenic PAHs			
1-Methylphenanthrene	9-Ethylphenanthrene	1-Methyl-7-isopropyl phenanthrene	5-Methylchrysene

150

6-Ethylchrysene      Acenaphthenequinone      1,4-Anthraquinone      9,10-Phenanthrenequinone

151      The US Environmental Protection Agency (US EPA) has designated 16 PAHs as priority  
 152 pollutants. Benzo[a]pyrene is a well-known human carcinogen among these key PAHs and is frequently  
 153 employed as an indication of PAH exposure [30]. Furthermore, these are categorized based on their  
 154 origins as pyrogenic PAHs, which are produced as a result of fossil fuel combustion, and petrogenic  
 155 PAHs, which are peculiar to crude oil and polluting water following an oil spill [31]. Petrogenic PAHs  
 156 differ from pyrogenic PAHs in that they are either heavily alkylated or oxygenated to produce PAH  
 157 quinones.

### 158 **3.2.2. Dioxins and dibenzofurans**

159 Chlorinated dibenzofurans (furans) and chlorinated dibenzo-p-dioxins (dioxins) have similar chemical  
 160 properties and toxic effects, thereby being a major concern for decades. 2,3,7,8-TCDD; 2,3,7,8-TCDF and  
 161 PCB-126 are examples of these kinds of substances. Nowadays, there has been a new group called dioxin-  
 162 like PCBs (DLPCBs) which includes a specific subgroup of PCBs. Polychlorinated dioxins (PCDDs) and  
 163 dibenzofurans (PCDFs) are released from anthropogenic sources and activities rather than natural  
 164 processes [32]. These have been reported to be carcinogenic, as well as have been found to put adverse  
 165 impacts on the reproductive system, immune system, and other human health risks. Treatment of PCDDs  
 166 and PCDFs includes the low-temperature thermal degradation facility which has resulted in the efficient  
 167 decomposition of these toxic substances from the municipal solid waste incineration fly ash [33].  
 168 Recently, significant reductions have been reported in the atmospheric levels of PCDDs and PCDFs  
 169 which might be because of some technical improvisations like waste incinerators, smelters, cement kiln  
 170 plants, modifications of motor vehicles and also substituting coal/diesel in domestic heating with the  
 171 natural gas controls [34]. Further, some air control pollution devices were helpful in the significant  
 172 removal of PCDD/Fs from hazardous waste disposals with 93.1% removal efficiency, consequently  
 173 resulting in high sulfur content in the hazardous waste [35].

174      Furthermore, their structural analogs in which bromine substitutes all of the chlorine atoms i.e.  
 175 polybrominated dioxins and dibenzofurans (PBDD/Fs) have similar physicochemical properties but their  
 176 half-lives are reported to be longer than that of the chlorinated ones [36]. It has been found that the  
 177 thermal discharge of waste electronic items releases PBDDs and PBDFs up to 50–500 times in  
 178 comparison to that of PCDD/Fs [39] (Wu et al., 2020). Soils in e-waste burning areas have been found to

179 be rich in PBDFs (88% of the total contaminants) which showed a major concern for the  
180 environment[38,39].

#### 181 **4. Extraction methods of POPs in the environmental samples**

182 Extraction of POPs from solid environmental samples has frequently been done using various organic  
183 solvents with or without addition of heat. The basic extraction techniques are liquid/liquid extraction and  
184 soxhlet extraction. A number of extraction techniques along with the analytical methods used for various  
185 POPs have been summarized in Table 5. Different methods have different recoveries depending on the  
186 type of analyte and the sample type. For instance, some reports have suggested that sediments and soils be  
187 freeze-dried [40]; while others have reported lower recoveries of PCBs from freeze-dried sediments [41].

##### 188 **4.1. Soxhlet extraction**

189 This method involves extractors in which a high amount of sample is mixed with the organic solvents and  
190 extracts the component without filtering. The major limitations of this method were large amounts of  
191 solvents (about 500 mL), and high extraction time (up to 2 days). Hence, some amendments have been  
192 made to the traditional soxhlet extraction like automated, focused microwave-assisted, and high-pressure  
193 soxhlet extraction. For example, the analysis of PCBs in fish and sediment samples achieved the best  
194 recoveries [42,43]. From sediment samples of the baltic sea, PCBs were extracted using accelerated  
195 solvent extraction (ASE) with solvent and toluene for soxhlet extraction [44]. Furthermore, 58 POPs  
196 including PAHs, OCPs, PCBs were detected in different atmospheric samples using Soxhlet extraction  
197 with 1:1 acetone: n-hexane for isolation purposes and solid phase extraction (SPE) cartridges. The  
198 average recoveries obtained were 67-114% for three different kinds of POPs [45]. Similarly, a modified  
199 soxhlet extraction was adopted for the detection of 8 different classes of POPs including PCDD/Fs,  
200 PBDEs, PBDD/Fs, PCBs, OCPs, polychlorinated naphthalenes (PCNs), short-chain chlorinated paraffins  
201 (SCCPs) and Dechlorane Plus (DP) in sediment and fish samples with 45 to 120% average recoveries.

##### 202 **4.2. Liquid-liquid extraction**

203 Liquid-liquid extraction (LLE) was found to be very tedious and labor intensive, hence solid-phase  
204 extraction (SPE) was its possible replacement. It offers less solvent exposure, quick processing, and easy  
205 handling as compared to the LLE [46,47]. Solid-phase extraction was used for the extraction of POPs  
206 from water and wastewater samples, even with high loadings of analytes which resulted in the reduction  
207 of the solvent usage and the analysis time [48]. OCPs and PCBs were successfully extracted from human  
208 serum samples with this method by using n-hexane-DCM mixture with recoveries of 99 to 120% for  
209 PCBs and 88 to 115% for OCPs [49]. Recently, 18 different POPs (mostly OCPs and PCBs) were  
210 extracted from the marine trout samples using LLE with average recoveries ranging between 73-112%

211 [50]. Meanwhile, several POPs were extracted from the soil leachate samples using LLE method [51]  
212 Some unique samples require solid phase microextraction (SPME) techniques and headspace sampling.

### 213 **4.3. Supercritical fluid extraction**

214 Supercritical fluid extraction (SFE) is another efficient method for removing PCBs from the soil,  
215 sediments, and other solid samples. The primary determinants of recovery rates in this approach are  
216 optimizing temperature, pressure, and flow rate. For example, PCBs were removed from sediment  
217 samples using this process at 200°C and 150-650 atm pressure with a 95% recovery rate [52]. PCBs were  
218 extracted from fatty acid samples using this approach with a basic alumina combination as an alternative  
219 to silica/silver nitrate (1:10, w/w) [53]. But poor repeatability is the major drawback of this method which  
220 has not been improved yet.

### 221 **4.4 Passive sampling**

222 This is also an effective method for assessing water samples utilizing semi-permeable membrane devices  
223 (SPMDs). SPMD sampling is a type of passive sampling for water, air, or sediments. Some other passive  
224 sampling techniques have been recently reviewed in the literature which is mainly based on the free flow  
225 of analytes from the sampling medium to the collecting medium [54]. Furthermore, some samples  
226 necessitate cleanup processes, which include the removal of co-extracted lipids from such samples prior  
227 to analysis. This will be determined primarily by the type of analyte. Because PCBs are highly stable in  
228 acidic conditions, it will be an effective strategy to conduct out PCB cleanup processes employing  
229 sulfuric acid or acid-impregnated silica columns for the elimination of lipids. Following that, sulfur must  
230 be removed from the samples in order to reduce co-extractive interferences and lengthen the detector's  
231 lifetime. Sulfur can thus be eliminated by introducing copper granules during the extraction process [55].

### 232 **4.5. Solid-phase extraction and microextraction**

233 The solid-phase microextraction method is an effective sample preparation technique for integrating  
234 several operations such as sample collection, extraction, analyte enrichment, and isolation from sample  
235 matrices, and has been used to extract analytes from gaseous, liquid, and solid samples [56]. A novel  
236 SPME was developed for estimations of various POPs (BTEX, PAHs, and PCBs) in which SPME fibers  
237 were used that were almost 180 times more sensitive than commercial fibers [57]. SPME was employed  
238 for the determination of PCBs from water samples with detection limits of 0.08-0.89 ng/L in treated  
239 sewage samples [58]. Dispersive micro-solid phase extraction (D- $\mu$ -SPE), in which the solid extracting  
240 phase is suspended in the liquid sample has been found to be more advantageous over the conventional  
241 SPE in terms of operational simplicity, speed, recovery, and handling of large sample volumes [59, 60].

## 242 **5. Analytical techniques for POPs**

243 Different types of chromatographic techniques have been employed for POPs analysis, such as gas  
244 chromatography (GC), ultra-high-performance liquid chromatography/or ultraperformance liquid  
245 chromatography (UHPLC/UPLC), high-performance liquid chromatography (HPLC), liquid  
246 chromatography (LC), and capillary liquid chromatography (CLC). A number of analytical methods used  
247 for the detection and quantification of various POPs have been summarized in this study.

248 The traditional HPLC is regularly used for the analysis of ionic PFAS, specially PFCA and PFSA,  
249 whereas GC is used for the analysis of volatile and semi-volatile PFAS. Particles collected on C18  
250 extraction disks can be extracted quantitatively without Soxhlet or PLE extraction. PFAS have been  
251 extracted from sediments collected from different lakes by sonicating them in methanol (recoveries: 88-  
252 102%) and analyzed by UPLC/ESI-MS/MS and obtained the total concentrations ranging from 0.61 ppb  
253 to 26 ppb [61]. LC-MS/MS method was also developed for the analysis of hydroxylated PCBs with a  
254 LOD value of 0.1 ng/mL after applying suitable cleanup to them [62, 63,64]. In addition to this, liquid  
255 chromatography (LC) with tandem mass spectrometry (MS/MS) or high-resolution mass analyzer  
256 (HRMS) like time of flight (TOF) or Orbitrap have been used mostly for the analysis of POPs in food  
257 samples [65].

258 One of the most important analytical methods for the analysis of POPs is gas chromatography  
259 (GC). Pressure or temperature-programmed injection techniques, combined with increased injection  
260 volume, have been proven to be quite effective without having any detrimental effects [66]. The boiling  
261 temperatures of the compounds and their interactions with the stationary phase of the column determine  
262 the efficiency of GC-based separation. Most POPs are semi-volatile, with polarities ranging from mild to  
263 non-polar, hence well adapted to GC-MS measurement because of their physicochemical features, with  
264 the exception of PFAS-related compounds, which are always tested using the LC-MS/MS technique [67].  
265 In the case of OCPs, it has been reported that both GC-MS and LC-MS based methods gave significantly  
266 similar results except for dichlorodiphenyltrichloroethane (DDT) and its metabolites, where GC-MS/MS  
267 was found to give better results than LC-MS/MS [68]. Detection of PCBs and PBDEs in serum samples  
268 of livestock in the US, the higher concentration of PCBs and PBDEs were detected in the livestock's  
269 serum, but their origin is still unclear [69]. GC-MS-based methods have been used to confirm the  
270 presence of PCBs in the transformer oils and detected in air samples too [70]. Several GC-ECD methods  
271 have also been reported for the analysis of PCBs from sediments, serum, fish tissue, and indoor air  
272 samples in addition to the GC-MS methods [71]. In addition to this, recently atmospheric pressure  
273 chemical ionization associated with GC-MS as an ionization source has detected a large number of  
274 molecular ions which provided a wide scope of screening. GC-APCI-MS methods are found to be highly  
275 sensitive and selective for the analysis of various POPs in environmental as well as biological samples at  
276 trace levels with LOD values around 10 to 100 times lower than other traditional GC-MS methods [72].

277 Recently, PBDEs, PCBs, PCDD/Fs, PBDD/Fs, furans, and other THs (total L-thyroxine (TT4), total  
 278 3,3',5-triiodo-L-thyronine (TT3), and total 3,3',5'-triiodo-L-thyronine (TrT3) has been detected and  
 279 determined in human breast milk using GC-high resolution mass spectrometer (GC-HRMS) with three  
 280 different columns [73].

281 Recently, a new rapid on-site detection approach has been developed which was highly sensitive  
 282 with LOD values in parts-per-trillion levels i.e. less than 5.25 ng/L in aquatic samples with approximately  
 283 30 minutes of total analysis time without the use of vials. This approach was developed by coupling a  
 284 portable GC-MS with an on-site pre-equilibrium solid phase microextraction (SPME) sampling method  
 285 and has been recently used for the detection of three kinds of POPs i.e. PCBs, OCPs, and PAHs) [74]

286 **Table 5: Determination of POPs in different sample matrices using various extraction methods and**  
 287 **analytical techniques**

Sr No	Analytes	Matrix	Extraction Method	Analytical Techniques	LOD values	References
1	PCBs	Urine	SPE	LC/MS/MS	0.01 ng/mL	[75]
2	PCBs	Plasma	SPE	LC/MS/MS	0.01 ng/mL	[76]
3	EPPs, PCBs	Sediment	LLE	LC/MS/MS	0.01-0.4 ng/g	[77]
4	PCBs	Water	LLE	RP-HPLC	0.25 mg/mL	[78]
5	PCBs	Mammals	LLE	RP-HPLC	2.5 mg/kg	[79]
6	PCBs, PCDFs	Fish	LLE	RP-HPLC	2-100 ng/g	[80]
7	PCBs	Herring oil	LLE	RP-HPLC	9-50 ng/g	[81]
9	PFBS, PFOS	Ground, river, and tap water	SPE	LC/(-)ESI- MS/MS	25,000 ng/L (LOQ)	[82]
10	PFOA, PFOS	Surface water	SPE	LC/(-)ESI-MS	0.2–13 ng/L	[83]

						(LOQ)
11	PFOS, PFOA, PFNA, PFDA,	Rain water	SPE	LC/(-)ESI-MS	0.04–7.2 ng/L	[84]
12	HCB, HCH, PCBs, OCPs	Fish	SE, LLE	GC/ECD, HRGC/ECD	-	[85]
13	PBDEs, PCBs, OCPs	Dolphin	GPC	GC/MS	-	[86]
14	OCPs, PCBs, toxaphene	Bird	LLE, SE	GC/ECD, GC- ECNI-MS	-	[87],[88]
15	PCDD/F, HCB	Air	Swipe/biofilm s, passive sample	HRGC/HRMS	-	[89]
16	PCBs, HCH, OCPs, HCB, DDT	Human	LLE, SPE, SE, LSE, hot SE	SPE-HPLC	-	[90]

288

## 289 **6. Toxicity behavior of POPs**

290 The ecotoxicological effects of POPs on the environment, biota, and human health have raised a big  
 291 concern in the past few years which has led to their minimal usage or complete ban in many countries.  
 292 These are resistant to almost all biodegradation processes, hence are highly persistent and toxic. It has  
 293 been recently addressed that in children, POPs can cause cancer and tumors in many places, immune  
 294 system illnesses, reproductive issues, decreased ability to fight disease, slowed growth, and irreversible  
 295 cognitive function damage. As a result, POPs are a suspected carcinogen [91].

296 Considering the contamination of the aquatic environment, PCBs and OCPs have emerged as the  
 297 most toxic among the wide range of POPs. Moderate to high exposure to PCBs (0.25-2 mg/L) has  
 298 induced abnormalities in developing zebrafish retinas, resulting in visual impairment [92]. PCB 126  
 299 exposure has also resulted in heart malformations in the embryos of *Danio rerio* [93]. Many species of  
 300 *Daphnia* are used as models or indicator organisms for the toxicity determination of PCBs mainly in  
 301 freshwater [70]. PFASs have also been found to be toxic to the soil, wastewater, animals, and humans  
 302 [94,95,96,97].

303 Certain PAHs, when exposed to animals through food for a prolonged time were found to cause  
304 stomach cancer from ingestion, lung cancer from inhalation, and skin cancer from skin contact. For  
305 instance, benzo[a]pyrene, benzo[a]anthracene, and chrysene were responsible for significant chromosome  
306 aberrations in rodents [86]. Among PAHs, the most common PAH to cause cancer in animals and was the  
307 first chemical carcinogen to be discovered is Benzo(a)pyrene [98].

## 308 **7. Conclusion**

309 The present compilation of various POPs and their toxicities concludes that new candidate POPs are  
310 adding to the list day by day. The ‘dirty dozens’ are no more dozen in number. Thus, novel research must  
311 be focused in developing new analytical approaches for trace estimation of POPs. Moreover, their highly  
312 toxic nature clearly indicates that they should be excluded from the environment as soon as possible.  
313 Various removal strategies have been proposed for this purpose such as photocatalysis, bio  
314 nanocomposites, MOF-based adsorbents or other porous absorbents etc. More work should be done for  
315 the development of environmentally benign moieties for the adsorptive removal of these POPs for the  
316 well-being of the ecosystem. Future research must also focus on estimation of toxicity/health hazards of  
317 POPs to various non-target organisms and associated environment or agro-ecosystem.

318

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320 **Authors’ contributions** This review has been prepared in collaboration among all authors. All authors  
321 read and approved the final manuscript.

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