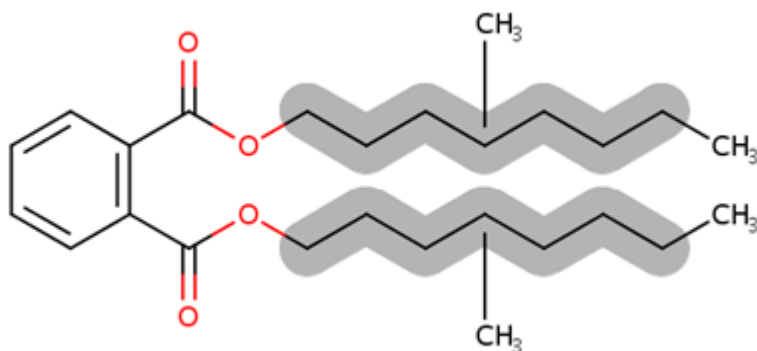




6  
7  
8  
9 **Draft Fate Assessment for Diisononyl Phthalate (DINP)**

10  
11 **Technical Support Document for the Draft Risk Evaluation**

12  
13 **CASRN: 28553-12-0 and 68515-48-0**



14  
15 (Representative Structure)

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23  
24  
25  
26 *May 2024*

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69

70 **ABBREVIATIONS AND ACRONYMS**


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BAF	Bioaccumulation factor
BCF	Bioconcentration factor
BMF	Biomagnification factor
BSAF	Biota-sediment accumulation factor
CASRN	Chemical Abstract Service registry number
DOC	Dissolved organic carbon
dw	Dry weight
DW	Drinking water
DWTP	Drinking water treatment plant
EPA	Environmental Protection Agency
ESI	Electrospray Ionization
FID	Flame ionization detector
FPD	Flame photometric detector
GC	Gas chromatography
HLC	Henry's Law constant
HPLC	High-performance liquid chromatography
ISO	International Organization for Standardization
K <sub>oa</sub>	Octanol-air partition coefficient
K <sub>oc</sub>	Organic carbon-water partition coefficient
K <sub>ow</sub>	Octanol-water partition coefficient
L/d	Liters per day
LOD	Limit of detection
LOQ	Limit of quantification
lw	Lipid weight
M	Molarity (mol/L = moles per Liter)
mL/min	milliliters per minute
mM	millimolar
MDL	Method Detection Limit
MRL	Method Reporting Limit
MS	Mass spectrometry
n	Sample size
N/A	Not applicable
ND	Non-detection
nm	nanometers
NR	Not reported
OECD	Organisation for Economic Co-operation and Development
·OH	Hydroxyl radical
OPE	Organophosphate ester
pg/L	picograms per liter
ppm	parts per million

QSAR	Quantitative structure activity relationship
RSD	Relative standard deviation
SI	Supplemental information
SIM	Selected ion monitoring
SPE	Solid phase extraction
STP	Sewage treatment plant
TMF	Trophic magnification factor
TOC	Total organic carbon
TOF	Time of flight
UPLC	Ultra-performance liquid chromatography
U.S.	United States
UV (UV-Vis)	Ultra-violet (visible) light
ww	Wet weight
WWTP	Wastewater treatment plant

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---

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76

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84 Health Protection (OCHP), Office of General Counsel (OGC), Office of Research and Development  
85 (ORD), and Office of Water (OW).

86

### 87 **Docket**

88 Supporting information can be found in the public docket, Docket ID ([EPA-HQ-OPPT-2024-0073](#)).

89

### 90 **Disclaimer**

91 Reference herein to any specific commercial products, process or service by trade name, trademark,  
92 manufacturer, or otherwise does not constitute or imply its endorsement, recommendation, or favoring  
93 by the United States Government.

94

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98

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100

101 **This report was reviewed and cleared by OPPT and OCSPP leadership.**

102 **SUMMARY**

103 This technical document is in support of the *Draft Risk Evaluation for Diisononyl Phthalate (DINP)*  
104 ([U.S. EPA, 2024b](#)). DINP is a common chemical name for the category of chemical substances that  
105 includes the following substances: 1,2-benzenedicarboxylic acid, 1,2-isononyl ester (CASRN 28553-12-  
106 0) and 1,2-benzenedicarboxylic acid, di-C9-11-branched alkyl esters, C9-rich (CASRN 68515-48-0).  
107 Both CASRNs contain mainly C9 dialkyl phthalate esters. See the draft risk evaluation for a complete  
108 list of all the technical support documents for DINP.

109  
110 In this document, EPA evaluated the reasonably available information to characterize the environmental  
111 fate and transport of, the key points are summarized below. Given the consistent results from numerous  
112 high-quality studies, there is robust evidence that DINP

- 113 • Is expected to undergo significant direct photolysis and will rapidly degrade in the atmosphere  
114 ( $t_{1/2} = 8.5$  hours) (Section 3.3).
- 115 • Is expected to degrade rapidly via direct and indirect photolysis (Section 3.3).
- 116 • Is not expected to appreciably hydrolyze under environmental conditions (Section 3.2).
- 117 • Is expected to have environmental biodegradation half-life in aerobic environments on the order  
118 of days to weeks (Section 3.1).
- 119 • Is not expected to be subject to long range transport.
- 120 • Is expected to transform in the environment via biotic and abiotic processes to form  
121 monoisononyl phthalate, isononanol, and phthalic acid (Section 3).
- 122 • Is expected to show strong affinity and sorption potential for organic carbon in soil and sediment  
123 (Sections 5.2.2, 5.3.2).
- 124 • Will be removed at rates greater than 93 percent in conventional wastewater treatment systems  
125 (Section 6.2).
- 126 • When released to air, will not likely exist in gaseous phase, but will show strong affinity for  
127 adsorption to particulate matter (Sections 4 and 5).
- 128 • Is likely to be found in, and accumulate in, indoor dust (Section 5).

129  
130 As a result of limited studies identified, there is moderate evidence that DINP

- 131 • Is not expected to biodegrade under anoxic conditions and may have high persistence in  
132 anaerobic soils and sediments (Sections 3.1, 5.2.2, 5.3.2).
- 133 • Is not bioaccumulative in fish in the water column (Section 7).
- 134 • May be bioaccumulative in benthic organisms exposed to sediment with elevated concentrations  
135 of DINP proximal to continual sources of release (Section 7).
- 136 • Is expected to be removed in conventional water treatment systems both in the treatment process,  
137 and via reduction by chlorination and chlorination byproducts in post treatment storage and  
138 drinking water conveyance (Section 6.3).

139 **1 INTRODUCTION**

---

140 DINP is considered ubiquitous in various environmental media to due to its presence in both point and  
141 non-point source discharges from industrial and conventional wastewater treatment effluents, biosolids  
142 and sewage sludge, stormwater runoff, and landfill leachate ([Net et al., 2015](#)). As an isomeric mixture,  
143 the fate and transport properties of DINP can be difficult to classify. However, the following sections of  
144 the fate and transport analysis of DINP are present the general fate and transport characteristics of  
145 DINP.

146 **2 APPROACH AND METHODOLOGY**

147 Reasonably available environmental fate data—including biotic and abiotic biodegradation rates,  
 148 removal during wastewater treatment, volatilization from lakes and rivers, and organic carbon:water  
 149 partition coefficient (log  $K_{OC}$ )—are the parameters used in the current draft risk evaluation. In assessing  
 150 the environmental fate and transport of DINP, EPA considered the full range of results from data  
 151 sources that were rated high-quality. Information on the full extracted dataset is available in the file  
 152 *Draft Risk Evaluation for Diisononyl Phthalate (DINP) – Systematic Review Supplemental File: Data*  
 153 *Quality Evaluation and Data Extraction Information for Environmental Fate and Transport* ([U.S. EPA,](#)  
 154 [2024c](#)). Other fate estimates were based on modeling results from EPI Suite™ ([U.S. EPA, 2012](#)), a  
 155 predictive tool for physical and chemical properties and environmental fate estimation.

156  
 157 Table 2-1 provides a summary of the selected environmental fate data that EPA considered while  
 158 assessing the fate of DINP and were updated after publication of *Final Scope of the Risk Evaluation for*  
 159 *Di-isononyl Phthalate (DINP) CASRN 28553-12-0/68515-48-0* ([U.S. EPA, 2021](#)) with additional  
 160 information identified through the systematic review process.

161 **Table 2-1. Summary of DINP’s Environmental Fate Information**

Parameter	Value	Source(s)
Octanol:Water (Log $K_{OW}$ )	8.8	<a href="#">ECHA (2016)</a>
Organic Carbon:Water (Log $K_{OC}$ )	5.5 (estimated; MCI method); 5.7 (estimated; $K_{OW}$ method)	<a href="#">U.S. EPA (2017b)</a>
Adsorption Coefficient (Log $K_d$ )	2.97 (suspended particulate matter/water)	<a href="#">Li et al. (2017a)</a>
	3.27 (sediment/water)	<a href="#">Li et al. (2017a)</a>
Octanol:Air (Log $K_{OA}$ )	11.9 (estimated)	<a href="#">U.S. EPA (2017b)</a>
Air:Water (Log $K_{AW}$ )	-2.20 (estimated)	<a href="#">Lu (2009)</a>
	-2.43 (estimated)	<a href="#">Cousins and Mackay (2000)</a>
Aerobic primary biodegradation in water	32 to 67.8% in 24 hours >90% in 5 days >99% in 28 days	<a href="#">(O’Grady et al., 1985; SRC, 1983; Monsanto, 1978)</a>
Aerobic ready biodegradation in water	57 to 81% in 28 days	<a href="#">(ECJRC, 2003b)</a>
Aerobic ultimate biodegradation in water	57 to 84% in 28 days	<a href="#">(HSDB, 2015; Monsanto, 1983)</a>
Aerobic biodegradation in sediment	0.54% in 14 days 1.11% in 28 days	<a href="#">(Johnson et al., 1983)</a>
Anaerobic biodegradation in sediment	0% in 100 days	<a href="#">(Ejlertsson et al., 1996)</a>
Aerobic biodegradation in soil	No significant change in concentration after 2 years	<a href="#">(ECJRC, 2003b)</a>
Hydrolysis	152 days at pH 8 and 25 °C, and 4.2 years at pH 7 and 25 °C	<a href="#">(U.S. EPA, 2017a)</a>
Photolysis	$t_{1/2}$ (air) = 5.36 to 8.5 hours $t_{1/2}$ (water <sub>pH=7</sub> ) = 140 days	<a href="#">(U.S. EPA, 2017a;</a> <a href="#">Lertsirisopon et al., 2009;</a> <a href="#">Peterson and Staples, 2003)</a>



Parameter	Value	Source(s)
Environmental degradation half-lives (selected values for modeling)	5.36 hours (air) 10 days (water) 20 days (soil) 90 days (sediment)	( <a href="#">U.S. EPA, 2017a</a> )
WWTP Removal	>93%	( <a href="#">U.S. EPA, 2017a</a> )
Aquatic Bioconcentration (BCF)	<3 L/kg wet weight (rainbow trout; <i>Oncorhynchus mykiss</i> ) 5.2 L/kg wet weight (upper trophic Arnot-Gobas estimation)	( <a href="#">U.S. EPA, 2017a</a> ; <a href="#">EC/HC, 2015a</a> )
Aquatic Bioaccumulation (BAF)	68 (75 ug/kg wet weight in mussel from field study in Seine estuary, France) 21 L/kg wet weight (upper trophic Arnot-Gobas estimation)	( <a href="#">U.S. EPA, 2017a</a> ; <a href="#">ECJRC, 2003b</a> )
Aquatic Food web Magnification Factor (FWMF)	0.46 (Experimental; 18 marine species)	( <a href="#">Mackintosh et al., 2004</a> )
Terr. Bioconcentration (BCF)	0.01 - 0.02 Experimental; earthworms ( <i>Eisenia fetida</i> )	( <a href="#">ECJRC, 2003b</a> )
Terr. Biota-sediment accumulation factor (BSAF)	0.018 OECD Test Guideline 207 ( <i>Eisenia fetida</i> )	( <a href="#">EC/HC, 2015a</a> )

## 2.1 EPI Suite™ Model Inputs and Settings

163

164 The approach described by ([Mackay et al., 1996](#)) using the Level III Fugacity model in EPI Suite™  
165 (LEV3EPI™) was used for this Tier II analysis. LEV3EPI is described as a steady-state, non-equilibrium  
166 model that uses a chemical's physical and chemical properties and degradation rates to predict  
167 partitioning of the chemical between environmental compartments and its persistence in a model  
168 environment ([U.S. EPA, 2017a](#)). A Tier II analysis involves reviewing environmental release  
169 information for DINP to determine whether further assessment is warranted for each environmental  
170 medium. Environmental release data for DINP was not available from the Toxics Release Inventory  
171 (TRI) or Discharge Monitoring Reports (DMRs); however, between 250 million and 550 million lb of  
172 CASRN 28553-12-0 and between 100 million and 1,000 million lb of CASRN 68515-48-0 were  
173 produced annually from 2016 to 2019 for use in commercial products, chemical substances or mixtures  
174 sold to consumers, or at industrial sites according to production data from the Chemical Data Reporting  
175 (CDR) 2020 reporting period. DINP is used as a plasticizer in polyvinyl chloride (PVC) and non-PVC  
176 products ([U.S. EPA, 2020](#); [EC/HC, 2015a](#)). DINP may be released to the environment during  
177 production, distribution, processing in PVC and non-PVC polymers, use of products such as paints and  
178 sealants, disposal or recycling, wastewater treatment, and disposal of solid and liquid waste ([ECJRC,](#)  
179 [2003b](#)).

180

181 Environmental release information is also useful for fugacity modeling because the emission rates will  
182 predict a real-time percent mass distribution for each environmental medium. Environmental  
183 degradation half-lives were taken from high and medium quality studies that were identified through  
184 systematic review to reduce levels of uncertainties. Based on DINP's observed and calculated  
185 environmental half-lives, partitioning characteristics, and the results of Level III Fugacity modeling (see  
186 Figure 4-1 below), DINP is expected to partition primarily to soil and sediment—regardless of the  
187 compartment of the environmental release. The LEV3EPI™ results were consistent with environmental  
188 monitoring data. Further discussion of DINP partitioning can be found in Section 4.

189 The following inputs parameters were used for the Level III Fugacity model in EPI Suite™:

- 190 • Melting point = -48.00 °C
- 191 • Vapor pressure =  $5.40 \times 10^{-7}$  mm Hg
- 192 • Water solubility =  $6.10 \times 10^{-4}$  mg/mL
- 193 • Log  $K_{OW}$  = 8.8
- 194 • SMILES: CCCCCCC(C)COC(=O)c1cccc1C(=O)OCCCCC(C)C(C)C (representative
- 195 structure)

### 196 3 TRANSFORMATION PROCESSES

---

197 DINP released to the environment will transform to the monoester form (monoisononyl phthalate) via  
198 abiotic processes such as photolysis (direct and indirect) and hydrolysis of the carboxylic acid ester  
199 group ([U.S. EPA, 2023](#)). Biodegradation pathways for the phthalates consist of primary biodegradation  
200 from phthalate diesters to phthalate monoesters, then to phthalic acid, and ultimately biodegradation of  
201 phthalic acid to form CO<sub>2</sub> and/or CH<sub>4</sub> ([Huang et al., 2013](#)). The monoisononyl phthalate is both more  
202 soluble and more bioavailable than DINP. It is also expected to undergo biodegradation more rapidly  
203 than the diester form. EPA considered DINP transformation products and degradants qualitatively but  
204 due to their lack of persistence we do not expect them to substantially contribute to risk; thus, EPA is not  
205 considering them further in this draft risk evaluation. Both biotic and abiotic routes of degradation for  
206 DINP are described in the sections below.

#### 207 3.1 Biodegradation

---

208 DINP can be considered readily biodegradable under most aquatic and terrestrial environments. The  
209 EPA extracted and evaluated fourteen data sources containing DINP biodegradation information in  
210 water, soil, and sediments under aerobic and anaerobic conditions (Table 3-1). Eight of the sources were  
211 classified as overall high-quality, five as overall medium-quality and one as overall low-quality data  
212 sources. DINP is considered an isomeric mixture, and certain components of DINP might biodegrade  
213 more readily than others ([ECJRC, 2003b](#)). DINP's aerobic primary biodegradation in water has reported  
214 to be 32 to 67.8 percent in 24 hours ([O'Grady et al., 1985](#); [Monsanto, 1978](#)), greater than 90 percent in 5  
215 days ([O'Grady et al., 1985](#)), and greater than 99 percent in 28 days ([U.S. EPA, 2019](#)) with half-life of  
216 1.5 to 5.31 days under acclimated conditions ([SRC, 1984, 1983](#)) and 7 to 40 days under unacclimated  
217 conditions ([EC/HC, 2015a](#)). Several studies evaluating the readily biodegradability of phthalate esters in  
218 water have reported DINP's half-life of 10.3 days ([ExxonMobil, 2010](#)) and 57 to 81 percent DINP  
219 removal in 28 days by CO<sub>2</sub> evolution ([HSDB, 2015](#); [ECJRC, 2003b](#)). The required 60 percent  
220 degradation during the 10-day pass window was met only in two of the four available studies ([ECJRC,](#)  
221 [2003b](#)). However, DINP in water has been reported to completely biodegrade into its basic elements by  
222 57 to 84 percent after 28 days (based on the available ultimate biodegradation information) ([EC/HC,](#)  
223 [2015a](#); [HSDB, 2015](#); [Monsanto, 1983](#); [SRC, 1983](#)). In contrast to the rapid biodegradation of DINP in  
224 aerobic environments, available information suggests that DINP is expected to have very low  
225 biodegradation potential under low oxygen conditions ([Ejlertsson et al., 1996](#)) and could remain longer  
226 in subsurface sediments and soils ([Kickham et al., 2012](#); [ECJRC, 2003b](#); [Johnson et al., 1984, 1983](#)).  
227

228 **Table 3-1. Summary of DINP’s Biodegradation Information**

Environmental Conditions	Degradation Value	Half-Life (days)	Reference	Overall Data Quality Ranking
Aerobic primary biodegradation in water	32% in 24 hours	ND	( <a href="#">Monsanto, 1978</a> )	High
	94 to 96% in 9 days	1.5 days (average; 1–1.9 days)	( <a href="#">SRC, 1984</a> )	High
	67.8% in 24 hours greater than 90% in 5 days	ND	( <a href="#">O’Grady et al., 1985</a> )	High
	greater than 99% in 28 days	5.31 days	( <a href="#">SRC, 1983</a> )	High
	91 to 100% in 7 days	7 to 40 days	( <a href="#">EC/HC, 2015a</a> )	Medium
	greater than 95% in 12 days	ND	( <a href="#">HSDB, 2015</a> )	Medium
	90 to 100% in 5 to 28 days 68% in 1 day	ND	( <a href="#">U.S. EPA, 2019</a> )	Medium
Aerobic ready biodegradation in water	70.5% in 28 days	ND	( <a href="#">ECJRC, 2003b</a> )	Medium
	57% in 28 days	ND		
	81% in 28 days	ND		
	74% in 28 days	ND	( <a href="#">HSDB, 2015</a> )	Medium
	ND	10.3 days	( <a href="#">ExxonMobil, 2010</a> )	Low
Aerobic ultimate biodegradation in water	61.5% in 28 days	ND	( <a href="#">SRC, 1983</a> )	High
	84% in 28 days	ND	( <a href="#">Monsanto, 1983</a> )	Medium
	57 to 71% in 28 days	ND	( <a href="#">HSDB, 2015</a> )	Medium
	56.6 % in 29 days	ND	( <a href="#">EC/HC, 2015a</a> )	Medium
	67.5% in 28 days	ND		
	74% in 28 days	ND		
Aerobic biodegradation in sediment	0.54% in 14 days 1.11% in 28 days	ND	( <a href="#">Johnson et al., 1983</a> )	High
	ND	12,000 days	( <a href="#">Kickham et al., 2012</a> )	High
	0.7% at 12 °C 1.2% at 22 °C 2.2% at 28 °C	ND	( <a href="#">Johnson et al., 1984</a> )	High
Anaerobic biodegradation in sediment	0% in 100 days	ND	( <a href="#">Ejlertsson et al., 1996</a> )	High
Aerobic biodegradation in soil	No significant change in concentration after 2 years	ND	( <a href="#">ECJRC, 2003b</a> )	Medium

229

### 230 **3.2 Hydrolysis**

231 Traditionally accepted methods of testing for abiotic hydrolysis (OECD Guideline Test 111) are not  
 232 viable for DINP due to the low aqueous solubility ([ECJRC, 2003a](#)). Therefore, hydrolysis rates of DINP  
 233 are difficult to accurately measure experimentally ([ECJRC, 2003a](#)). EPI Suite™ was utilized to estimate  
 234 the hydrolysis half-lives of DINP at 152 days at pH 8 and 25 °C, and 4.2 years at pH 7 and 25 °C ([U.S.](#)

235 [EPA, 2017a](#)) indicating that hydrolysis is a possible degradation pathway of DINP under more caustic  
236 conditions. Lertsirisopon (2009) reported the hydrolysis half-lives of 720 days (pH = 5), 1,200 days (pH  
237 = 6), negligible (pH = 7), 1,000 days (pH = 8), and 460 days (pH = 9), at average temperature of 10.8  
238 °C. However, this study received a low data quality ranking in the systematic review process due to  
239 poorly documented and variable test conditions.

240

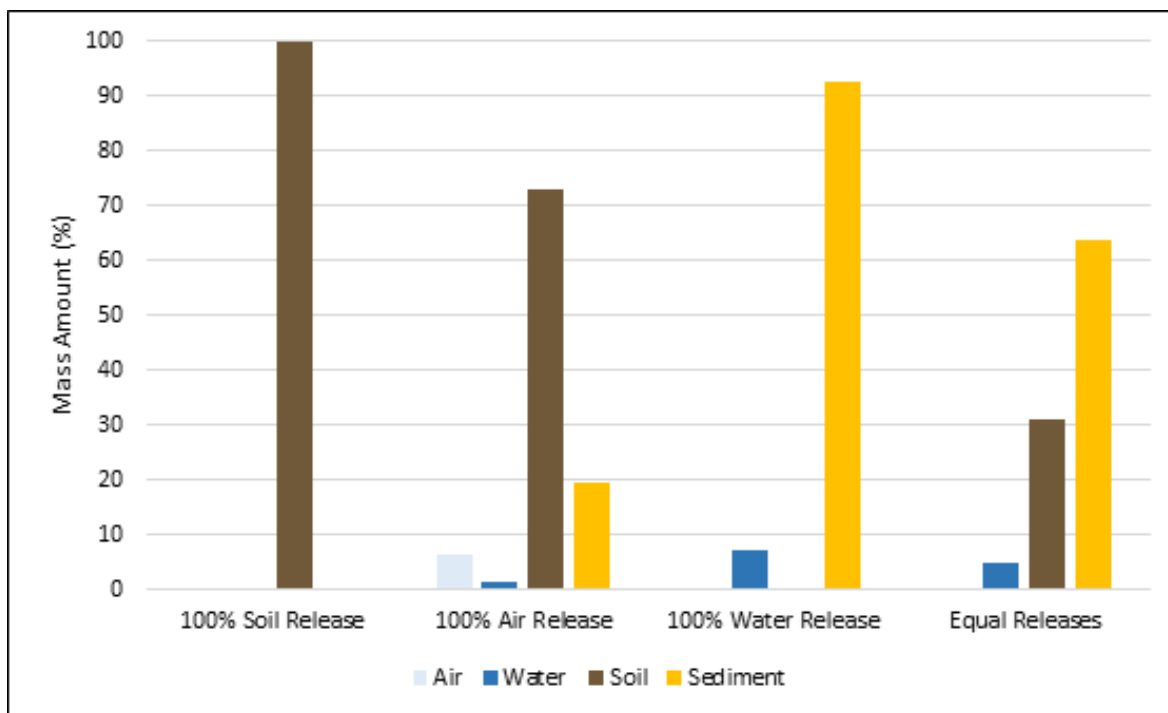
241 When compared to other degradation pathways, it is not expected that hydrolysis is a significant source  
242 of degradation under typical environmental conditions. However, the higher temperatures, variations  
243 from typical environmental pH, and chemical catalysts present in the deeper anoxic zones of landfills  
244 may be favorable to the degradation of DINP via hydrolysis ([Huang et al., 2013](#)). This is discussed  
245 further in Section 5.3.3.

### 246 **3.3 Photolysis**

247 DINP contains chromophores that absorb light at greater than 290 nm wavelength ([NCBI, 2020](#)),  
248 therefore, direct photodegradation is a relevant degradation pathway for DINP released to air. Modelled  
249 indirect photodegradation half-lives indicated a slightly more rapid rate of degradation, estimating a  
250 half-life of 0.22 days (5.36 hours) ( $\cdot\text{OH}$  rate constant of  $2.39 \times 10^{-11} \text{ cm}^3 / \text{molecule-second}$  and a 12-  
251 hour day with  $1.5 \times 10^6 \text{ OH/cm}^3$ ) ([U.S. EPA, 2017a](#)). Similarly, Peterson (2003) reported a calculated  
252 DINP photodegradation half-life of 0.35 days (8.5 hours) ( $\cdot\text{OH}$  rate constant of  $2.35 \times 10^{-11} \text{ cm}^3$   
253  $/\text{molecule-second}$  and  $1 \times 10^6 \text{ OH/cm}^3$ ). DINP photodegradation in water is expected to be slower than  
254 air, due to the typical light attenuation in natural surface water. The aquatic direct photodegradation  
255 half-lives of 32, 52, 140, 61 and 36 days were observed at pH 5, 6, 7, 8, and 9, respectively, when  
256 exposed to natural sunlight in artificial river water at 0.4 to 27.4 °C (average temperature of 10.8 °C)  
257 ([Lertsirisopon et al., 2009](#)).

## 258 4 PARTITIONING

259 Environmental release data for DINP was not available from the Toxics Release Inventory (TRI) or  
260 Discharge Monitoring Reports (DMRs), therefore DINP releases to the environment could not be  
261 estimated. The approach described by (Mackay et al., 1996) using the Level III Fugacity model in EPI  
262 Suite™ (LEV3EPI™) was used for this Tier II analysis. LEV3EPI is described as a steady-state, non-  
263 equilibrium model that uses a chemical's physical and chemical properties and degradation rates to  
264 predict partitioning of the chemical between environmental compartments and its persistence in a model  
265 environment (U.S. EPA, 2017a). DINP's physical and chemical properties were taken directly from  
266 Section 2.1 of *Draft Physical Chemistry Assessment for Diisononyl Phthalate (DINP)* (U.S. EPA,  
267 2024a). Environmental release information is useful for fugacity modeling because the emission rates  
268 will predict a real-time percent distribution for each medium. Environmental degradation half-live in  
269 water of 10 days was selected in this draft risk evaluation to represent the range of identified primary  
270 biodegradation half-life values (Section 3.1) from high and medium quality studies to reduce levels of  
271 uncertainties. EPA used environmental degradation half-lives of 5.36 hours in air (based on  
272 AEROWIN™ predicted values, an atmospheric fate prediction model within EPI Suite™), 20 days in  
273 soil (double the half-life in water), and 90 days in sediment (9 times the half-life in water) as  
274 recommended for EPIWIN estimations (U.S. EPA, 2017a). Based on DINP's environmental half-lives,  
275 partitioning characteristics, and the results of Level III Fugacity modeling, DINP is expected to be found  
276 predominantly in water, soil, and sediment (Figure 4-1). The LEV3EPI™ results were consistent with  
277 environmental monitoring data. Further discussion of DINP partitioning can be found in Sections 5.1,  
278 5.2, and 5.3.  
279



280  
281

**Figure 4-1. EPI Suite™ Level III Fugacity Modeling Graphical Result for DINP**

## 282 5 MEDIA ASSESSMENTS

---

283 DINP has been reported to be present in the atmosphere, aquatic environments, and terrestrial  
284 environments. Once in the air, DINP will be most predominant in the organic matter present in airborne  
285 particles and expected to have a short half-life in the atmosphere. Based on the physical and chemical  
286 properties, DINP is likely to partition to house dust and airborne particles and is expected to have a  
287 longer half-life compared to ambient (outdoor) air. DINP present in surface water is expected to mostly  
288 partition to aquatic sediments. DINP is expected to have an aerobic biodegradation half-life between 14  
289 and 28 days. In terrestrial environments DINP has the potential to be present in soils and groundwater  
290 but is likely to be immobile in both media types. In soils, DINP is expected to be deposited via air  
291 deposition and land application of biosolids. DINP in soils is expected to have a half-life on the order of  
292 days to weeks, have low bioaccumulation potential and biomagnification potential in terrestrial  
293 organisms. DINP is released to groundwater via wastewater effluent and landfill leachates, expected to  
294 have a half-life of 14 to 56 days, and not likely to be persistent in most groundwater/subsurface  
295 environments.

### 296 5.1 Air and Atmosphere

---

297 DINP is a liquid at environmental temperatures with a melting point of  $-48\text{ }^{\circ}\text{C}$  ([Haynes, 2014](#); [O'Neil,](#)  
298 [2013](#)) and a vapor pressure of  $5.40 \times 10^{-7}$  mmHg at  $25\text{ }^{\circ}\text{C}$  ([NLM, 2015](#)). Based on its physical and  
299 chemical properties and short half-life in the atmosphere,  $t_{1/2} = 5.36$  hours ([U.S. EPA, 2017a](#)), DINP was  
300 assumed to not be persistent in the air. The AEROWIN<sup>TM</sup> module in EPI Suite<sup>TM</sup> estimated that a large  
301 fraction of DINP could be sorbed to airborne particles and these particulates may be resistant to  
302 atmospheric oxidation. DINP has not been detected in ambient air; however, studies have detected DINP  
303 in settled house dust, indoor air samples and in indoor particulate phase air samples ([NCBI, 2020](#);  
304 [Kubwabo et al., 2013](#); [ECJRC, 2003b](#)).

#### 305 5.1.1 Indoor Air and Dust

---

306 In general, phthalate esters are ubiquitous in the atmosphere and indoor air. Their worldwide presence in  
307 air has been documented in the gas phase, suspended particles and dust ([Net et al., 2015](#)). Most of the  
308 studies reported DEHP (di-ethylhexyl phthalate) to be the predominant phthalate esters in the  
309 environment. Despite the limited information on the presence of DINP on the atmosphere, similar trends  
310 to those reported for DEHP could be expected based on their similar vapor pressure ([ECHA, 2013](#)).  
311 Limited studies have reported the presence of particle bound DINP on indoor and outdoor settings  
312 ([Gupta and Gadi, 2018](#); [Hasegawa, 2003](#); [Helmig et al., 1990](#)). Once in indoor air, DINP is expected to  
313 partition to organic carbon present on indoor airborne particles. Under indoor environments, DINP is  
314 expected to be more persistent in indoor air than in ambient (outdoor) air due to the lack of natural  
315 chemical removal processes, such as solar photochemical degradation.

316  
317 The available information suggests that the concentration of DINP in dust under indoor environments to  
318 be higher than outdoors dust and to be associated with the presence of phthalate-containing articles and  
319 the proximity to manufacturing facilities ([Kubwabo et al., 2013](#); [Wang et al., 2013](#); [Abb et al., 2009](#)).  
320 [Kubwabo \(2013\)](#) monitored the presence of 17 phthalate compounds in vacuum dust samples collected  
321 in 126 urban single-family homes. The study reported that DEHP, DIDP, and DINP were detected in all  
322 the collected dust samples accounting for 88 percent of the median total concentration of phthalates in  
323 dust. [Wang \(2013\)](#) evaluated the presence of phthalates in dust samples collected from indoor and  
324 outdoor settings in two major Chinese cities. The study reported the total phthalates concentration of the  
325 collected indoor dust samples were 3.4 to 5.9 times higher than those collected outdoors. The aggregate  
326 concentration of DEHP, DINP, and DIDP in indoor dust samples accounted for 91 to 94 percent of the  
327 total phthalate's concentration. The study revealed that the aggregate concentration of phthalates was

328 higher in the commercial and industrial areas with heavy production of textiles, costumes, and toys. Abb  
329 (2009) evaluated the presence of phthalates in indoor dust samples collected from 30 households in  
330 Germany. The study revealed the presence of DEHP, DIDP and DINP in all the collected samples.  
331 Samples collected from households containing a high percentage of plastics (greater than 50 percent  
332 plastic content) resulted with higher aggregate concentration of phthalates in dust. The aggregate  
333 concentration of DEHP, DIDP and DINP accounted for 87 percent of the total phthalate concentration in  
334 dust.

335  
336 Similarly, recent studies monitoring the presence of phthalates in dust from USA households have  
337 revealed DEHP and DINP to be detected in 96 to 100 percent of the collected samples (Hammel et al.,  
338 2019; Dodson et al., 2017). Hammel (2019) and Dodson (2017) reported the presence of phthalate esters  
339 on indoor air and dust samples collected in USA homes. Hammel (2019) reported that DINP accounted  
340 for close to 83 percent of the total concentration of phthalates found in indoor dust. Dodson (2017)  
341 evaluated the presence of phthalate esters in air samples of U.S. homes before and after occupancy  
342 reporting increased presence of DINP after occupancy due to daily anthropogenic activities that might  
343 introduce phthalate containing products into indoor settings. Increasing trends could be expected for  
344 DINP with its increased use on household's construction materials or consumer products.

## 345 **5.2 Aquatic Environments**

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### 346 **5.2.1 Surface Water**

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347 DINP is expected to be released to surface water via industrial and municipal wastewater treatment plant  
348 effluent, surface water runoff, and, to a lesser degree, atmospheric deposition. DINP and other phthalate  
349 esters have been detected in surface waters, although at lower frequencies than some other phthalate  
350 esters (Wen et al., 2018). The principal properties governing the fate and transport of DINP in surface  
351 water are water solubility, organic carbon partitioning coefficients, and volatility. Due to its Henry's  
352 Law constant ( $9.14 \times 10^{-5}$  atm·m<sup>3</sup>/mol at 25 °C) of DINP, volatilization is not expected to be a significant  
353 source of loss of DINP from surface water. A partitioning analysis of DINP released to the environment  
354 is described in Section 4 above. The analysis estimates that during releases to surface water bodies,  
355 greater than 92 percent of DINP released to surface water will partition to both suspended and benthic  
356 sediments.

357  
358 DINP has a low water solubility of 0.00061 mg/L, but is likely to form a colloidal suspension and may  
359 be detected in surface water at higher concentrations (EC/HC, 2015b). Based on DINP's water solubility  
360 and partitioning coefficients, DINP in water will partition to suspended organic material present in the  
361 water column. DINP is expected to be readily biodegradable in water (Section 3.1). In addition, total  
362 seawater samples concentrations of DINP measured in False Creek ranged from 61 to 135 ng/L; the  
363 dissolved fraction concentrations ranged from 29 to 64 ng/L and the suspended particulate fraction  
364 concentration ranged from 14,700 to 50,400 ng/g dry weight (dw) (EC/HC, 2015a; Mackintosh et al.,  
365 2006). Concentrations of DINP above the aqueous solubility of 0.00061 mg/L are not uncommon in  
366 monitoring studies proximal to releases of DINP to surface water (Wen et al., 2018).

### 367 **5.2.2 Sediments**

---

368 Based on the water solubility (0.00061 mg/L) and affinity for sorption to organic matter (log K<sub>oc</sub> = 5.5  
369 to 5.7), DINP will partition to the organic matter present in soils and sediment when released into the  
370 environment. Once in water, DINP is expected to be readily biodegradable and the Level III Fugacity  
371 Model in EPI Suite™ (U.S. EPA, 2017a) predicts that greater than 92 percent of the DINP will partition  
372 to and remain in sediments (Section 4). The available information suggests that DINP could persist  
373 longer in subsurface sediments and soils than in water. In terrestrial and aquatic environments, DINP has



374 potential to accumulate in sediments at areas of continuous release, such as a surface water body  
375 receiving discharge from a municipal wastewater treatment plant.  
376

377 Due to the strong sorption to organic carbon, DINP is expected to be found predominantly in sediments  
378 near point sources, with a decreasing trend in sediment concentrations downstream. This is consistent  
379 with monitoring information for phthalate esters from Sweden and Korea. One study reported the  
380 presence of DINP in only one sediment sample near a point source in Sweden that recently have  
381 replaced DEHP with DINP in their production processes ([Parkman and Remberg, 1995](#)). The presence  
382 of DINP has been documented in urban sediments at concentrations ranging between 130 to 3,200 µg/kg  
383 total solids with a 62 percent detection frequency ([Cousins et al., 2007](#)). In a similar study, Kim ([2021](#))  
384 evaluated the presence of plasticizers in sediments from highly industrialized bays of Korea. DINP was  
385 detected in all surface sediment samples. The study revealed a gradual decreasing trend in the overall  
386 concentration of phthalates toward the outer region of the bays farther away from industrial activities.  
387 The findings of this study suggest industrial activities to be the major contributor of phthalates in  
388 sediments within the area.  
389

390 Monitoring data from the Rhine River and the Neckar River in Germany detected DINP concentrations  
391 in sediment samples of 30, 220, 650, and 1,460 ppb and 430, 570, 1,050 ppb (3 sites), respectively  
392 ([NCBI, 2020](#)). DINP was also detected in sediment from 21 locations in the Netherlands at  
393 concentrations up to 6.16 mg/kg dry weight ([ECJRC, 2003b](#)).

## 394 **5.3 Terrestrial Environments**

---

### 395 **5.3.1 Soil**

---

396 DINP is expected to be deposited to soil via two primary routes: application of biosolids and sewage  
397 sludge in agricultural applications or sludge drying applications as well as atmospheric deposition.  
398 Based on DINP's Henry's Law constant of  $9.14 \times 10^{-5}$  atm·m<sup>3</sup>/mol at 25 °C and vapor pressure of  
399  $5.40 \times 10^{-7}$  mmHg, DINP is not likely to volatilize from soils.  
400

401 DINP shows an affinity for sorption to soil and its organic constituents (log K<sub>OC</sub> = 5.5–5.7; log K<sub>d</sub> of  
402 2.55–3.27 ([Li et al., 2017b](#); [Li et al., 2017a](#); [U.S. EPA, 2012](#)) and an estimated log K<sub>OW</sub> of 10.21 ([U.S.](#)  
403 [EPA, 2017a](#)) Given that these properties indicate the likelihood of strong sorption to organic carbon  
404 present in soil, DINP is expected to have low mobility in soil environments.  
405

406 Under aerobic conditions, DINP is expected to have a half-life in soil of 20 days. This aerobic  
407 biodegradation half-life for soil was estimated by doubling the experimentally derived half-life of DINP  
408 in water as very limited soil biodegradation data for DINP identified in the systematic review process  
409 ([SRC, 1983](#)).  
410

411 Under anaerobic conditions that may be present in some soil profiles, there is very little evidence to  
412 support that DINP appreciably biodegrades ([ECJRC, 2003b](#); [Ejlertsson et al., 1996](#)). One study found  
413 that 0 percent degradation had occurred under anaerobic conditions after 100 days by CH<sub>4</sub> evolution and  
414 no transformation reported based on the concentrations of methane and test substance with gas  
415 chromatographic analysis in municipal solid waste samples with an anaerobic microflora inoculum  
416 ([Ejlertsson et al., 1996](#)). Furthermore, another study reported less than 1 percent DINP degradation in  
417 anaerobic sediments after 28 days ([Johnson et al., 1984](#)).  
418

419 In general, DINP is not expected to be persistent in soil as long as the rate of release does not exceed the  
420 rate at which biodegradation can occur, but continuous exposure to DINP in soil proximal to points of

421 releases may be possible if the rate of releases exceeds the rate of biodegradation under aerobic  
422 conditions. Under anaerobic conditions in soil, DINP is assumed to be persistent, and continuous  
423 exposure is likely.

### 424 **5.3.2 Biosolids**

425 Sludge is defined as the solid, semi-solid, or liquid residue generated by wastewater treatment processes.  
426 The term “biosolids” refers to treated sludge that meet the EPA pollutant and pathogen requirements for  
427 land application and surface disposal and can be beneficially recycled (40 CFR Part 503) ([U.S. EPA,  
428 1993](#)). Typically, chemical substances with very low water solubility and high sorption potential are  
429 expected to be sorbed to suspended solids and efficiently removed from wastewater via accumulation in  
430 sewage sludge and biosolids.

431  
432 There is limited information about the presence and biodegradation of DINP in biosolids. As described  
433 in Section 6.2, DINP in wastewater has been reported to be mainly removed by particle sorption and  
434 retained in the sewage sludge. In general, greater than 93 percent of the DINP present in wastewater is  
435 expected to be accumulated in sewage sludge and discharged into biosolids. Once in biosolids, DINP  
436 could be transferred to soil during land applications. DINP will be strongly sorbed to organic matter on  
437 soils and to be more persistent in soil profiles with anaerobic conditions ([ECJRC, 2003b](#)). Due to its  
438 strong sorption to soils, land applied DINP is not expected to be bioavailable; thus, exposures to  
439 environmental organisms and people are negligible. In addition, based on the water solubility and  
440 hydrophobicity of DINP, will have low bioaccumulation, biomagnification appear to be of minimal  
441 concern. Additionally, terrestrial species have been reported to have the capacity to metabolize phthalate  
442 substances ([Bradlee and Thomas, 2003](#); [Gobas et al., 2003](#); [Barron et al., 1995](#)) and DINP is expected to  
443 have low bioaccumulation potential and biomagnification potential in terrestrial organisms (Section 7).

### 444 **5.3.3 Landfills**

445 For the purpose of this assessment, landfills will be considered to be divided into two zones: (1) an  
446 “upper-landfill” zone, with normal environmental temperatures and pressures, where biotic processes  
447 are the predominant route of degradation for DINP; and (2) a “lower-landfill” zone where elevated  
448 temperatures and pressures exist, and abiotic degradation is the predominant route of degradation for  
449 DINP. In the upper-landfill zone where oxygen may still be present in the subsurface, conditions may  
450 still be favorable for aerobic biodegradation; however, photolysis and hydrolysis are not considered to  
451 be significant sources of degradation in this zone. In the lower-landfill zone, conditions are assumed to  
452 be anoxic and temperatures present in this zone are likely to inhibit biotic degradation of DINP.  
453 Temperatures in lower-landfills may be as high as 70 °C. At temperatures at and above 60 °C, biotic  
454 processes are significantly inhibited, and are likely to be completely irrelevant at 70 °C ([Huang et al.,  
455 2013](#)).

456  
457 DINP is deposited in landfills continually and in high amounts from the disposal of consumer products  
458 containing DINP. However due to its strong sorption to soils and low water solubility, small  
459 concentration of DINP is likely to be present in landfill leachate. DINP is likely to be persistent in  
460 landfills due to the apparent lack of anaerobic biodegradation and unfavorable conditions for  
461 biodegradation in lower-landfills. Some aerobic biodegradation in upper-landfills may occur. In lower-  
462 landfills, there is some evidence to support that hydrolysis may be the main route of abiotic degradation  
463 of phthalate esters ([Huang et al., 2013](#)).

464  
465 Despite the expected persistence of DINP in landfills, it is not expected to be bioavailable and mainly  
466 sorbed to organic matter in soils due to the low water solubility of DINP (0.00061 mg/L), and its high  
467 sorption to organic carbon (log  $K_{oc}$  = 5.5–5.7). Although DINP may be present at small concentrations

468 in landfill leachate, it is unlikely to migrate to, or be mobile in groundwater proximal to landfills, and  
469 would not be expected to be transported distally from landfills via groundwater.

#### 470 **5.3.4 Groundwater**

---

471 There are several potential sources of DINP in groundwater, including wastewater effluents and landfill  
472 leachates, which are discussed in Sections 5.3.3 and 6.2. Further, in environments where DINP is found  
473 in surface water, it can enter groundwater through surface water/groundwater interactions. Diffuse  
474 sources include storm water runoff and runoff from biosolids applied to agricultural land.

475  
476 Given the strong affinity of DINP to adsorb to organic matter present in soils and sediments ( $\log K_{OC} =$   
477  $5.5-5.7$ ) ([U.S. EPA, 2012](#)) DINP is expected to have low mobility in soil and groundwater  
478 environments. Furthermore, due to the insoluble nature of DINP (0.00061 mg/L), high concentrations of  
479 DINP in groundwater are unlikely. In instances where DINP could reasonably be expected to be present  
480 in groundwater environments (proximal to landfills or agricultural land with a history of land applied  
481 biosolids), limited persistence is expected based on rates of biodegradation of DINP in aerobic  
482 environments, DINP is not likely to be persistent in groundwater/subsurface environments unless anoxic  
483 conditions exist.

## 484 6 PERSISTENCE POTENTIAL OF DINP

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485 DINP is not expected to be persistent in the environment, as it is expected to degrade rapidly under most  
486 environmental conditions, with delayed biodegradation in low-oxygen media. In the atmosphere, DINP  
487 is unlikely to remain for long periods of time as it is expected to undergo photolytic degradation through  
488 reaction with atmospheric hydroxyl radicals, with estimated half-lives of 5.36 hours. DINP is predicted  
489 to hydrolyze slowly at ambient temperature, but it's not expected to persist in aquatic media as it  
490 undergoes rapid aerobic biodegradation (Section 5.2.1). DINP have the potential to remain for longer  
491 periods of time in soil and sediments, but due to the inherent hydrophobicity ( $\log K_{OW} = 8.8$ ) and  
492 sorption potential ( $\log K_{OC} = 5.5-5.7$ ) DINP is not expected to be bioavailable for uptake. Using the  
493 Level III Fugacity model in EPI Suite<sup>TM</sup> (LEV3EPITM) (Section 4), DINP's overall environmental half-  
494 life was estimated to be approximately 34 days ([U.S. EPA, 2012](#)). Therefore, DINP is not expected to be  
495 persistent in the atmosphere or aquatic and terrestrial environments.

### 496 6.1 Destruction and Removal Efficiency

---

497 Destruction and removal efficiency (DRE) is a percentage that represents the mass of a pollutant  
498 removed or destroyed in a thermal incinerator relative to the mass that entered the system. DINP is  
499 classified as a hazardous substance and EPA requires that hazardous waste incineration systems destroy  
500 and remove at least 99.99 percent of each harmful chemical in the waste, including treated hazardous  
501 waste (46 FR 7684) ([U.S. EPA, 1981](#)).  
502

503 Currently there is no information available on the DRE of DINP. However, the DEHP annual releases  
504 from a Danish waste incineration facility were estimated to be 9 percent to air and 91 percent to  
505 municipal land fill ([ECB, 2008](#)). These results suggest that DINP present during incineration processes  
506 will be very likely to be released to landfills and the remaining small fraction released to air. Based on  
507 its hydrophobicity and sorption potential, DINP released to landfills is expected to partition to waste  
508 organic matter. Similarly, DINP released to air is expected to be rapidly react via indirect photochemical  
509 processes within hours ([U.S. EPA, 2017a](#)) and partition to soil and sediments as described in Section 4.  
510 DINP in sediments and soils is not expected to be bioavailable for uptake, and highly biodegradable in  
511 its bioavailable form ([Kickham et al., 2012](#)).

### 512 6.2 Removal in Wastewater Treatment

---

513 Wastewater treatment is performed to remove contaminants from wastewater using physical, biological,  
514 and chemical processes. Generally, municipal wastewater treatment facilities apply primary and  
515 secondary treatments. During the primary treatment, screens, grit chambers, and settling tanks are used  
516 to remove solids from wastewater. After undergoing primary treatment, the wastewater undergoes a  
517 secondary treatment. Secondary treatment processes can remove up to 90 percent of the organic matter  
518 in wastewater using biological treatment processes such as trickling filters or activated sludge.  
519 Sometimes an additional stage of treatment such as tertiary treatment is utilized to further clean water  
520 for additional protection using advanced treatment techniques (*e.g.*, ozonation, chlorination,  
521 disinfection).  
522

523 Limited information is available in the fate and transport of DINP in wastewater treatment systems. The  
524 EPA selected two high quality sources reporting the removal of DINP in wastewater treatment systems  
525 employing aerobic and anaerobic processes. One study reported 98.0 percent DINP removal efficiencies  
526 in a municipal wastewater treatment facility in France, employing a combined decantation and activated  
527 sludge tank ([Tran et al., 2014](#)). Like other phthalates esters with long carbon chains and high  $\log K_{OW}$ ,  
528 DINP was reported to be mainly removed by particle sorption and retained in the sewage sludge. This  
529 finding is supported by STPWIN<sup>TM</sup>, an EPI Suite<sup>TM</sup> module that estimates chemical removal in sewage

530 treatment plants. The model predicts greater than 93 percent removal of DINP during conventional  
531 wastewater treatment by sorption to sludge with the potential of increased removal via rapid aerobic  
532 biodegradation processes ([U.S. EPA, 2012](#)). In addition, the treatment of wastewater final solids via  
533 aerobic digestions processes have been reported to achieve 41.1 and 85.9 percent reduction on DINP  
534 concentration from the digestion effluents ([Armstrong et al., 2018](#)). In addition, the same study reported  
535 anaerobic solids digestion to be not effective in the removal of DINP. In general, the available  
536 information suggest that aerobic processes have the potential to help biodegrade DINP from wastewater  
537 in agreement with the expected aerobic biodegradation described in Section 3.1.

538

539 Overall, DINP has a high log  $K_{OW}$  and remains in suspended solids and efficiently removed from  
540 wastewater via accumulation in sewage sludge ([Tran et al., 2014](#)), partially removed during aerobic  
541 solids digestion processes ([Armstrong et al., 2018](#)), and ineffectively removed under anaerobic solids  
542 digestion conditions ([Armstrong et al., 2018](#)). Biodegradation and air stripping are not expected to be  
543 significant wastewater removal processes. Therefore, greater than 93 percent of the DINP present in  
544 wastewater is expected to be accumulated in sewage sludge and released with biosolids disposal or  
545 application, with the remaining fraction sorbed to suspended solids in the wastewater treatment effluent  
546 and discharged with surface water ([Tran et al., 2014](#); [U.S. EPA, 2012](#)).

### 547 **6.3 Removal in Drinking Water Treatment**

---

548 Drinking water in the United States typically comes from surface water (*i.e.*, lakes, rivers, reservoirs)  
549 and groundwater. The source water then flows to a treatment plant where it undergoes a series of water  
550 treatment steps before being dispersed to homes and communities. In the U.S., public water systems  
551 often use conventional treatment processes that include coagulation, flocculation, sedimentation,  
552 filtration, and disinfection, as required by law.

553

554 Very limited information is available on the removal of DINP in drinking water treatment plants. No  
555 data was identified by the EPA for DINP in drinking water. Based on the water solubility and Log  $K_{OW}$ ,  
556 DINP in water it is expected to mainly partition to suspended solids present in water. This is supported  
557 by the Level III Fugacity model in EPI Suite<sup>TM</sup> (Section 4) which predicts 92.7 percent of DINP released  
558 to water partitioning to sediments ([U.S. EPA, 2012](#)). The available information on the DEHP removal  
559 efficiency of flocculants and filtering media, suggest that DINP could potentially be partially removed  
560 during drinking water treatment by sorption into suspended organic matter. This data source reported  
561 58.7 percent reduction on drinking water DEHP concentration from a conventional drinking water  
562 treatment effluent in China and 78 to 86 percent loss of DINP during storage of treated drinking water  
563 effluent after 48 hours in Taiwan using chlorine for disinfection prior to distribution ([Kong et al., 2017](#);  
564 [Yang et al., 2014](#)). These findings suggest that conventional drinking water treatment systems may have  
565 the potential to partially remove DINP is present in drinking water sources via sorption to suspended  
566 organic matter and filtering media and the use of disinfection technologies.

**567 7 BIOACCUMULATION POTENTIAL OF DINP**

568 The presence of DINP in several marine aquatic species in North America suggest that the substance to  
569 be bioavailable in aquatic environments ([Mackintosh et al., 2004](#)). However, based on the very low  
570 water solubility and high hydrophobicity, DINP is expected to have low bioaccumulation potential,  
571 biomagnification potential, and low potential for uptake. EPA selected three overall high quality data  
572 sources and two overall medium quality data source reporting the aquatic bioconcentration, aquatic  
573 bioaccumulation, aquatic food web magnification, terrestrial biota-sediment accumulation, and  
574 terrestrial bioconcentration of DINP (Table 7-1). The available data sources discussed below, suggest  
575 that DINP has low bioaccumulation potential in aquatic and terrestrial organisms ([EC/HC, 2015a](#);  
576 [Solbakken et al., 1985](#); [Chemical Manufacturers, 1984](#)), and no apparent biomagnification across trophic  
577 levels in the aquatic food web ([Mackintosh et al., 2004](#)).

578  
579 Several studies have investigated the aquatic bioconcentration and food web magnification of DINP in  
580 several marine species. Solbakken ([1985](#)) evaluated the bioconcentration of DINP in Arca zebra in a 24-  
581 hour exposure study, followed by a 14-day depuration period. The study reported DINP BCF values of  
582 8.2, 183.8, 13.6, and 9.3 dpm/ $\mu$ L during the 24-hour exposure period on Arca zebra muscle,  
583 hepatopancreas, gills, and blood respectively (Table 7-1). The study reported a 92 to greater than 99  
584 percent decrease on BCF values during the 14-days depuration period. A similar study evaluating the  
585 presence of phthalates on estuaries reported a mussel BAF of 68 and DINP content of 75  $\mu$ g/kg wet  
586 weight ([ECJRC, 2003b](#)). A DINP exposure studies on rainbow trout have reported BCF lower than 3  
587 L/kg wet weight and biomagnification factor lower than 0.1 ([EC/HC, 2015a](#)). The reported low BCF  
588 values suggest that DINP has low potential to bioaccumulate in aquatic organisms. In the other hand, the  
589 Chemical Manufacturers Association ([1984](#)) reported a higher predicted DINP aquatic BCF of 1,155  
590 using a regression model based on the substance water solubility. Despite of the different range of  
591 reported BCF values, the empirical rapid BCF decrease during a 14-day depuration period ([Solbakken et  
592 al., 1985](#)), the empirical aquatic TMF of 0.46 ([Mackintosh et al., 2004](#)), modeled upper trophic BCF of  
593 5.2 L/kg wet-wt and upper trophic BAF of 21 L/kg wet weight ([U.S. EPA, 2017a](#)) helps support that  
594 DINP will have low bioconcentration potential and low biomagnification potential across trophic levels  
595 in the aquatic food web.

596  
597 There is very limited information on the bioconcentration and bioaccumulation of DINP in terrestrial  
598 environments. Based on DINP's strong sorption organic matter ( $\log K_{oc}$  5.5–5.7) ([U.S. EPA, 2017a](#))  
599 and water solubility (0.00061 mg/L) ([Letinski et al., 2002](#)), DINP is not expected to be bioavailable in  
600 soils. This is supported by the reported low BCF values of 0.1 to 0.2 on earthworms (*Eisenia foetida*)  
601 ([ECJRC, 2003b](#)). Therefore, DINP is expected to have low bioaccumulation potential, biomagnification  
602 potential in terrestrial organisms.  
603

604 **Table 7-1. Summary of DINP's Bioaccumulation Information**

Endpoint	Value	Details	Reference	Overall Quality Ranking
Aquatic Bioconcentration (BCF)	8.2 at day 0 4.6 at day 1 1.3 at day 4 0.03-0.01 at day 14 (dpm/mg)	Experimental; Muscle BCF; 14-C DINP 24 hours exposure followed by 14-day depuration period; <i>Arca Zebra</i> (mollusk)	<a href="#">(Solbakken et al., 1985)</a>	High
	183.8 at day 0 125.2 at day 1 64.5 at day 4 14.4 at day 14 (dpm/mg)	Experimental; Hepatopancreas BCF; 14-C DINP 24 hours exposure followed by 14-day depuration period; <i>Arca Zebra</i> (mollusk)		
	13.6 at day 0 12.4 at day 1 6.5 at day 4 0.8 at day 14 (dpm/mg)	Experimental; Gills BCF; 14-C DINP 24 hours exposure followed by 14-day depuration period; <i>Arca Zebra</i> (mollusk)		
	9.3 at day 0 5.6 at day 1 4.4 at day 4 0.1 at day 14 (dpm/μL)	Experimental; Blood BCF; 14-C DINP 24 hours exposure followed by 14-day depuration period; <i>Arca Zebra</i> (mollusk)		
	0.46 at day 0 0.45 at day 1 0.26 at day 4 0.13 at day 14 (dpm/mg)	Experimental; BCF; 14-C DINP 24 hours exposure followed by 14-day depuration period; <i>Diploria Strigosa</i> (coral)		
	1155	Predicted; $\log \text{BCF} = (0.542 \times \log K_{ow}) + 0.124$ ; calculated using Kow values that were calculated from water solubility; $\log K_{ow} = 5.2 - 0.68 \times \log (\text{micromolar WS})$ .	<a href="#">(Chemical Manufacturers, 1984)</a>	High
	<3 L/kg wet weight	experimental; rainbow trout; <i>Oncorhynchus mykiss</i> ; elimination rate: 1.16/day; tissue elimination half-life: <1 day; BMF: <0.1	<a href="#">(EC/HC, 2015a)</a>	Medium
Aquatic Bioaccumulation (BAF)	68	Experimental; preliminary study; Field study; Mussel; Collected from Seine estuary, France; 75 ug/kg wet weight in mussel from field study in Seine estuary, France	<a href="#">(ECJRC, 2003b)</a>	Medium

Endpoint	Value	Details	Reference	Overall Quality Ranking
Aquatic Food web Magnification Factor (FWMF)	0.46	experimental; 18 marine species, representing four trophic levels; trophic dilution, predominantly absorbed via the diet and depurated at a rate greater than the passive elimination rate via fecal egestion and respiratory ventilation, due to metabolism; FWMF (food web magnification factor) = 0.44;	<a href="#">(Mackintosh et al., 2004)</a>	High
Terr. Bioconcentration (BCF)	0.01 - 0.02	Terrestrial BCF; Experimental; earthworms ( <i>Eisenia fetida</i> ); steady state may not have been achieved.; 14 days	<a href="#">(ECJRC, 2003b)</a>	Medium
Terr. Biota-sediment accumulation factor (BSAF)	0.018	experimental; other: OECD Test Guideline 207 (Earthworm, acute toxicity; OECD 1984a); earthworm; <i>Eisenia fetida</i> ;	<a href="#">(EC/HC, 2015a)</a>	Medium

605



## 606 **8 OVERALL FATE AND TRANSPORT OF DINP**

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607 The inherent physical chemical properties of DINP govern its environmental fate and transport. Based  
608 on DINP's aqueous solubility, slight tendency to volatilize, and strong tendency to adsorb to organic  
609 carbon this chemical substance will be preferentially sorbed into sediments, soils, and suspended solids  
610 in wastewater treatment processes. Soil, sediment, and sludge/biosolids are predicted to be the major  
611 receiving compartments for DINP as indicated by its physicochemical and fate properties, partitioning  
612 analysis, and verified by monitoring studies. Surface water is predicted to be a minor pathway, and the  
613 main receiving compartment for phthalates discharged via wastewater treatment processes. However,  
614 phthalates in surface water will sorb strongly to suspended and benthic sediments. In area where  
615 continuous releases of phthalates occur, higher levels of phthalates in surface water can be expected,  
616 trending downward distally from the point of releases. This also holds true for DINP concentration in  
617 both suspended and benthic sediments. While DINP undergoes relatively rapid aerobic biodegradation  
618 but it is persistent in anoxic/anaerobic environments (sediment, landfills) and like other phthalates it is  
619 expected to slowly hydrolyze under normal environmental conditions.

620  
621 If released directly to the atmosphere, DINP is expected to adsorb to particulate matter. It is not  
622 expected to undergo long-range transport facilitated by particulate matter due to the relatively rapid rates  
623 of both direct and indirect photolysis. Atmospheric concentrations of DINP may be elevated proximal to  
624 sites of releases. Off gassing from landfills and volatilization from wastewater treatment processes are  
625 expected to be negligible releases in terms of ecological or human exposure in the environment due to  
626 its low vapor pressure. DINP released to air may undergo rapid photodegradation and it is not expected  
627 to be a candidate chemical for long range transport.

628  
629 Under indoor settings, air released DINP is expected to partition to airborne particles at concentrations  
630 three times higher than in vapor phase ([ECJRC, 2003a](#)) and is expected to have extended lifetime as  
631 compared to outdoor settings. The available information suggests that DINP's indoor dust  
632 concentrations to be associated with the presence of phthalate containing articles and the proximity to  
633 the facilities producing them ([Kubwabo et al., 2013](#); [Wang et al., 2013](#); [Abb et al., 2009](#)) as well as daily  
634 anthropogenic activities that might introduce DINP containing products into indoor settings ([Dodson et  
635 al., 2017](#)).

636  
637 DINP has a predicted average environmental half-life of 35 days. In situations where aerobic conditions  
638 are predominant, DINP is expected to degrade rapidly and be more persistent under anoxic/anaerobic  
639 conditions. In some sediments, landfills, and soils, DINP may be persistent as it is resistant to anaerobic  
640 biodegradation. In anaerobic environments, such as deep landfill zones, hydrolysis is expected the most  
641 prevalent process for the degradation of DINP.

642 **9 WEIGHT OF SCIENTIFIC EVIDENCE CONCLUSIONS FOR**  
643 **FATE AND TRANSPORT**

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644 **9.1 Strengths, Limitations, Assumptions, and Key Sources of Uncertainty**  
645 **for the Fate and Transport Assessment**

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646 Given the consistent results from numerous high-quality studies, there is a robust confidence that DINP

- 647
- 648 • is expected to undergo significant direct photolysis (Section 3.3);
  - 649 • will partition to organic carbon and particulate matter in air (Sections 4);
  - 650 • will biodegrade in aerobic surface water, soil, and wastewater treatment processes (Sections 5.2.1, 5.3.1, and 6.2);
  - 651 • does not biodegrade in anaerobic environments (Section 5.2 and 5.3);
  - 652 • will be removed after undergoing wastewater treatment and will sorb to sludge at high fractions, with a small fraction being present in effluent (Section 6.2);
  - 653 • is not bioaccumulative (Section 7);
  - 654 • is not expected to biodegrade under anoxic conditions and may have high persistence in anaerobic soils and sediments (Sections 3.1, 5.2.2, and 5.3.2);
  - 655 • may show persistence in surface water and sediment proximal to continuous points of release (Sections 3.1, 5.2.2, and 5.3.2); and
  - 656 • is expected to transform to monoisononyl phthalate, isononanol, and phthalic acid in the environment (Section 3).
- 657
- 658
- 659
- 660

661 As a result of limited studies identified, there is a moderate confidence that DINP

- 662
- 663 • is expected to be removed in conventional drinking water treatment systems both in the treatment process, and via reduction by chlorination and chlorination byproducts in post treatment storage and drinking water conveyance (Section 6.3); and
  - 664 • showed no significant degradation via hydrolysis under standard environmental conditions but hydrolysis rate was seen to increase with increasing pH and temperature in deep-landfill environments (Section 5.3.3).
- 665
- 666
- 667

668 Findings that were found to have a robust weight of evidence supporting them had one or more high-  
669 quality studies that were largely in agreement with each other. findings that were said to have a  
670 moderate weight of evidence were based on a mix of high and medium-quality studies that were largely  
671 in agreement but varied in sample size and consistence of findings.

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