# Fluorescent Brightener 71 and related chemicals

**Evaluation statement** 

**14 December 2023** 



# **Table of contents**

# Contents

AICIS evaluation statement	4
Subject of the evaluation	4
Chemicals in this evaluation	4
Reason for the evaluation	4
Parameters of evaluation	4
Summary of evaluation	5
Summary of introduction, use and end use	5
Environment	5
Conclusions	6
Supporting information	7
Rationale	7
Chemical identity	7
Relevant physical and chemical properties	10
Introduction and use	10
Australia	10
International	10
Existing Australian regulatory controls	11
Environment	11
International regulatory status	11
United Nations	11
OECD	11
United States of America	11
Environmental exposure	12
Environmental fate	13

Predicted environmental concentration (PEC)	15
Environmental effects	18
Effects on Aquatic Life	18
Effects on sediment dwelling life	22
Effects on terrestrial Life	22
Predicted no-effect concentration (PNEC)	23
Categorisation of environmental hazard	23
Persistence	24
Bioaccumulation	24
Toxicity	24
Environmental risk characterisation	24
References	26

# **AICIS** evaluation statement

# Subject of the evaluation

Fluorescent Brightener 71 and related chemicals

# Chemicals in this evaluation

Name	CAS registry number
Benzenesulfonic acid, 2,2'-(1,2- ethenediyl)bis[5-[[4-(4-morpholinyl)-6- (phenylamino)-1,3,5-triazin-2-yl]amino]-, disodium salt	16090-02-1
Benzenesulfonic acid, 2,2'-(1,2-ethenediyl)bis[5-[[4-[bis(2-hydroxyethyl)amino]-6-[(4-sulfophenyl)amino]-1,3,5-triazin-2-yl]amino]-, sodium salt (1:4)	16470-24-9
Benzenesulfonic acid, 2,2'-(1,2-ethenediyl)bis[5-[[4-[bis(2-hydroxyethyl)amino]-6-(phenylamino)-1,3,5-triazin-2-yl]amino]-, disodium salt	4193-55-9

# Reason for the evaluation

Evaluation Selection Analysis indicated a potential environmental risk.

# Parameters of evaluation

This evaluation considers the environmental risks associated with the industrial uses of Fluorescent Brightener 71 (FB-71) (CAS RN 16090-02-1), Fluorescent Brightener 220 (FB-220) (CAS RN 16470-24-9) and Fluorescent Brightener 28 (FB-28) (CAS RN 4193-55-9).

Chemicals in this group have been assessed for environmental risk according to the following parameters:

- Australian introduction volumes of 100–1000 t/year.
- Industrial uses listed in the 'Summary of use' section.
- Expected emission to sewage treatment plants (STPs) following consumer and commercial use.
- Expected emission to STPs or surface water following industrial use.

These chemicals have been assessed as a group because they are structurally similar and have similar use patterns.

# Summary of evaluation

### Summary of introduction, use and end use

Chemicals in this group are used as fluorescent brightening agents (FBAs) in the following products according to Australian and international use data:

- laundry and dishwashing products
- fabric, textile and leather products
- paper products
- paint and coating products.

Available information indicates that chemicals in this group are used in high volumes in Australia and worldwide.

#### **Environment**

#### Summary of environmental hazard characteristics

Based on the information presented in this evaluation and according to the environmental hazard thresholds stated in the Australian Environmental Criteria for Persistent, Bioaccumulative and/or Toxic Chemicals (DCCEEW n.d.), the chemicals are:

- Persistent (P)
- Not Bioaccumulative (Not B)
- Not Toxic (Not T).

#### **Environmental hazard classification**

The chemicals FB-220 and FB-28 do not satisfy the criteria for classification for environmental hazards according to the Globally Harmonised System of Classification and Labelling of Chemicals (GHS). All experimental toxicity endpoints exceed minimum effect levels for classification (UNECE 2017). This evaluation does not consider classification of physical and health hazards.

FB-71 satisfies the criteria for classification as follows:

Environmental Hazard	Hazard Category	Hazard Statement
Hazardous to the aquatic environment (acute / short-term)	Aquatic Acute 2	H401: Toxic to aquatic life

#### Summary of environmental risk

Chemicals in this evaluation are present in a range of household and commercial products. These chemicals are expected to be released to wastewater due to use of the products. The main sources of emissions are expected to be from their use in laundry detergents and from paper and textile processing. The environmental releases are expected to affect surface water, sediment and soil.

Chemicals in this group are persistent (P), not bioaccumulative (Not B) and not toxic (Not T). They are not expected to pose a significant risk to the Australian environment as RQs in water and soil fall well below the level of concern (RQ < 1). Although the risk to sediment-dwelling organisms cannot be ruled out (RQs in this compartment were not determined), international monitoring data indicates declining site specific sediment concentrations of fluorescent brightening agents (FBA), which may reduce the risk.

# Conclusions

The Executive Director is satisfied that the identified risks to the environment from the introduction and use of the industrial chemicals can be managed.

#### Note:

- 1. Obligations to report additional information about hazards under *Section 100* of the *Industrial Chemicals Act 2019* apply.
- 2. You should be aware of your obligations under environmental, workplace health and safety and poisons legislation as adopted by the relevant state or territory.

# Supporting information

# Rationale

This evaluation considers the environmental risks associated with the industrial use of Fluorescent Brightener 71 (FB-71), Fluorescent Brightener 28 (FB-28) and Fluorescent Brightener 220 (FB-220). Chemicals in this group have similar structures, physico-chemical properties, industrial uses and environmental hazards. Therefore, these chemicals are suitable for group assessment.

The evaluation selection analysis (ESA) for chemicals in this group highlighted a high volume of use globally and potential persistence and ecotoxicity, which indicate a high concern for the environment.

# Chemical identity

Chemicals in this group are characterised by a stilbenesulfonic acid core joined to substituted triazinyl groups via amine linkages. They are formally salts of sodium, but commercial products include aqueous slurries or powders that may contain added salts, dispersants and dedusting agents (OECD 2001; 2005a; 2005b). The stereochemistry of the internal alkene is not specified in the chemical names of this group. However, the cis-stilbene configuration is not fluorescent and commercial forms are therefore available in the trans-stilbene configuration. These configurations are denoted the (Z)- and (E)-isomers, respectively.

Chemical name Benzenesulfonic acid, 2,2'-(1,2-ethenediyl)bis[5-[[4-(4-

morpholinyl)-6-(phenylamino)-1,3,5-triazin-2-yl]amino]-,

disodium salt

**CAS RN** 16090-02-1

**Synonyms** Fluorescent Brightener 71 (FB-71)

Fluorescent Brightener 260

FWA 1

Benzenesulfonic acid, 2,2'-(1,2-ethenediyl)bis[5-[[4-(4-morpholinyl)-6-(phenylamino)-1,3,5-triazin-2-yl]amino]-,

disodium salt

Disodium 4,4'-bis(2-anilino-4-morpholino-1,3,5-triazin-6-

ylamino)-2,2'-stilbenedisulfonate

Molecular formula C40H38N12O8S2.2Na

Molecular weight (g/mol) 924.91 g/mol

SMILES (canonical) [Na].[Na].O=S(=O)(O)C1=CC(=CC=C1C=CC2=CC=C(C=

C2S(=O)(=O)O)NC3=NC(=NC(=N3)N4CCOCC4)NC=5C=

CC=CC5)NC6=NC(=NC(=N6)N7CCOCC7)NC=8C=CC=CC8

#### Structural formula

Chemical name Benzenesulfonic acid, 2,2'-(1,2-ethenediyl)bis[5-[[4-[bis(2-

hydroxyethyl)amino]-6-[(4-sulfophenyl)amino]-1,3,5-

triazin-2-yl]amino]-, sodium salt (1:4)

**CAS RN** 16470-24-9

Synonyms Fluorescent Brightener 220 (FB-220)

C.I. 40623

Benzenesulfonic acid, 2,2'-(1,2-ethenediyl)bis[5-[[4-[bis(2-hydroxyethyl)amino]-6-[(4-sulfophenyl)amino]-1,3,5-

triazin-2-yl]amino]-, tetrasodium salt

Tetrasodium 4,4'-bis[[4-[bis(2-hydroxyethyl)amino]-6-(4-

sulfonatoanilino)-1,3,5-triazin-2-yl]amino]-2,2'-

stilbenedisulfonate

Molecular formula C40H40N12O16S4.4Na

Molecular weight (g/mol) 1168 g/mol

**SMILES (canonical)** [Na].[Na].[Na].[Na].O=S(=O)(O)C1=CC=C(C=C1)NC=2N=

C(N=C(N2)N(CCO)CCO)NC3=CC=C(C=CC4=CC=C(C=C4S(=O)(=O)O)NC5=NC(=NC(=N5)N(CCO)CCO)NC6=C

C=C(C=C6)S(=O)(=O)O)C(=C3)S(=O)(=O)O

#### Structural formula

Chemical name Benzenesulfonic acid, 2,2'-(1,2-ethenediyl)bis[5-[[4-[bis(2-

hydroxyethyl)amino]-6-(phenylamino)-1,3,5-triazin-2-

yl]amino]-, disodium salt

**CAS RN** 4193-55-9

**Synonyms** Fluorescent Brightener 28 (FB-28)

Fluorescent Brightener 113

Disodium 4,4'-bis[6-anilino-[4-[bis(2-hydroxyethyl)amino]-

1,3,5-triazin-2-yl]amino]-2,2'-stilbenedisulfonate

Molecular formula C40H38N12O8S2.2Na

Molecular weight (g/mol) 924.91 g/mol

**SMILES (canonical)** [Na].[Na].O=S(=O)(O)C1=CC(=CC=C1C=CC2=CC=C(C=

C2S(=0)(=0)0)NC3=NC(=NC(=N3)N(CC0)CC0)NC=4C

=CC=CC4)NC5=NC(=NC(=N5)N(CCO)CCO)NC=6C=CC =CC6

Structural formula

# Relevant physical and chemical properties

Physical and chemical property data were retrieved from the registration dossiers for FB-71, FB-220 and FB-28 submitted under the Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH) legislation in the European Union (EU) (REACH n.d.-a; n.d.-b; n.d.-c).

Chemical	FB-71	FB-220	FB-28
Physical form	Solid	Solid	Solid
Melting point	Decomposes at 300°C (exp.)	Decomposes at 360°C (exp.)	Decomposes at 300°C (exp.)
Water solubility	1.9 g/L at 20°C (exp.)	` ' '	48.2 g/L at 20°C (exp.)
Ionisable in the environment?	Yes	Yes	Yes
log K <sub>ow</sub>	-1.58 (exp.)	-3.9 (exp.)	-3.5 (calc.)

Experimentally derived dissociation constants (pKa) were not identified for chemicals in this group. Based on data submitted to the former National Industrial Chemicals Notification and Assessment Scheme (NICNAS) for an analogous chemical, the sulfonate groups upon protonation will have pKa values of around 2 while protonated secondary and aromatic ring nitrogens will have pKa values of around 5–6 (NICNAS 1995).

### Introduction and use

#### Australia

Based on information reported to NICNAS under previous mandatory and/or voluntary calls for information, chemicals in this group are imported into Australia in volumes of between 100–1000 t/year.

No Australian uses were identified for chemicals in this group. However, FB-71 is reportedly used in laundry detergents (Kramer 1992).

#### International

Chemicals in this group are used in significant volumes worldwide (OECD n.d.). Global production volumes were estimated at 10,000–50,000 t/year of FB-71, 35,000 t/year of FB-220 and 5,000–10,000 t/year of FB-28 between 1999 and 2005 (OECD 2001; 2005a; 2005b). Current use volumes in specific international jurisdictions are 1–1,000,000 t/year per chemical:

Chemical	Europe (2023)	USA (2016–2019)	Japan
FB-71	1,000–10,000	454–4,540	101 (2014) 117 (2013)
FB-220	10,000–100,000	4540–22,700	1,000–2,000 (2021) 10,000–20,000 (2020)

Chemical Europe (2023)		USA (2016–2019) Japan	
FB-28	100,000–1,000,000	454–9,070	1-1,000 (2014-2021)

All units are in t/year. European volumes were retrieved from the REACH registration dossiers (REACH n.d.-a; n.d.-b; n.d.-c). Volumes from the United States of America (USA) were retrieved from the 2020 Chemical Data Reporting (CDR) database (US EPA 2020). Volumes for Japan were retrieved from the Japan CHEmicals Collaborative Knowledge database (J-CHECK) (NITE n.d.).

The chemicals in this group are fluorescent brightening agents (FBA) derived from condensation reactions of 4,4'-diaminostilbene-2,2'-disulfonic acid (CAS RN 81-11-8) with cyanuric chloride (CAS RN 108-77-0) and various primary and secondary amines (Kramer 1992). They absorb light in the near-UV region (300–400 nm) and rapidly re-emit in the blue region via fluorescence (Kramer et al. 1996). In paper, textiles and plastics, this fluorescence causes a whitening effect against the natural yellowish hue of these materials.

FB-71 is mainly used in laundry detergents (>90%), but has secondary uses in papermaking and textile finishing (<10%) (OECD 2005a). FB-220 and FB-28 are primarily used in papermaking, textile finishing and plastics manufacture, with some indicated use in laundry detergents (DeLima Associates n.d.; OECD 2001; 2005b). FB-71 and FB-220 are reportedly used in coatings and paints (REACH n.d.-a; n.d.-b). FB-28 is marketed as a laboratory chemical for use in fungal staining (NCBI n.d.).

# Existing Australian regulatory controls

#### **Environment**

The industrial uses of chemicals in this group are not subject to any specific national environmental regulations.

# International regulatory status

#### **United Nations**

Chemicals in this group are not currently identified as Persistent Organic Pollutants (POP) (UNEP 2001), ozone depleting substances (UNEP 1987), or hazardous substances for the purpose of international trade (UNEP & FAO 1998).

#### **OECD**

Chemicals in this group were independently sponsored by Germany under the Cooperative Chemicals Assessment Programme (CoCAP). FB-71 was recommended as a candidate for further work during the 21<sup>st</sup> Screening Information Data Set (SIDS) Initial Meeting Assessment (SIAM 21) in 2005 (OECD 2005a). FB-220 was recommended as a candidate for further work during SIAM 13 and FB-28 was recommended as a low priority for further work during SIAM 20 (OECD 2001; 2005b).

#### United States of America

Chemicals in this group are listed on the US EPA Safer Chemical Ingredients List (US EPA n.d.). This list identifies chemicals deemed to be safer alternatives to other chemicals

employed for the same functional use. The criteria for adding chemicals to the list cover a broad range of human health and environmental toxicological effects.

# Environmental exposure

Chemicals in this group are expected to be found as FBAs in household and commercial products available for use in Australia. Formulated products on the Australian market are assumed to be similar to those available internationally. According to international data, concentrations of FB-71 in laundry detergents are 0.05–0.35% (OECD 2005a), while concentration ranges of FB-220 and FB-28 for paper and textile whitening are 0.05–0.5% and 0.1–0.3%, respectively (OECD 2001; 2005b).

Chemicals in this group will be released into the environment when products containing these chemicals are used. Water, sediment and soil are expected to be the main compartments affected by the use of chemicals in this group. FBAs strongly adsorb to sludge during treatment in STPs. The remainder is released to surface water in sewage outfall or industrial effluents, leading to sediment exposure. Application of STP biosolids to agricultural land will lead to soil exposure, as will re-use of treated effluent for irrigation.

These chemicals (FBAs) are emitted to wastewater as a normal part of their use in laundry detergents. Between 20–95% of FBAs contained in detergents adhere to fabrics during washing (Kramer 1992). The remainder is discharged to sewers and received by STPs for treatment. End-product containers may be rinsed of residual detergent before disposal to landfill.

Chemicals in this group (FBAs) are contained in wastewater and paper sludge from industrial papermaking and recycling (OECD 2009). FBAs are emitted to wastewater during industrial use in textile finishing (OECD 2004). Industrial wastewater is typically reclaimed, discharged to sewers or managed *in situ* before release to sewers, agricultural land, or surface water. Paper sludge is landfilled, applied to agricultural soil, or incinerated for energy recovery.

The National Pollution Inventory (NPI) requires reporting of a number of pollutants, but FBAs are not one of them (NPI n.d.). However, reported emissions of other pollutants reflect how wastewater and sludge are disposed of in Australia. According to NPI data, there are currently three facilities from the *Pulp, Paper and Converted Paper Product Manufacturing* industry reporting pollutant emissions to water or land and seven facilities transferring pollutants to off-site sewerage, treatment, landfill or recycling destinations (NPI n.d.). Facilities emitting to water operate on-site treatment plants or partially discharge trade waste to off-site STPs (NSA 2022; Plant et al. 2014; Scholes 2021). There are currently no facilities from the *Textile, Leather, Clothing and Footwear Manufacturing* industry reporting pollutant emissions to water and only four facilities transferring pollutants to on-site immobilisation or off-site sewerage and landfill destinations (NPI n.d.).

Paper and textile products containing FBAs are exported, landfilled or recycled following use. More than half of all paper fibre used in Australia is derived from recycled material (DAFF n.d.). Post consumer printing and communication paper containing FBAs is recycled into office products (including office paper) and tissue paper, as well as some packaging grades (Industry Edge 2019). Except for tissue paper, which is disposed to sewers, waste paper is ultimately composted, recycled or disposed to landfill (Industry Edge 2019). Textiles are largely imported as finished products, the bulk of which (87.5%) are disposed to landfill following use (Khan et al. 2023).

The use of chemicals in this group in plastics, paints and coatings, or as laboratory chemicals, is not expected to lead to significant environmental release compared to other uses.

#### **Environmental fate**

#### Dissolution, speciation and partitioning

Chemicals in this group (FBAs) exist as anions in water but may form zwitterions at low environmental pH. They bind with dissolved cations in the environment, forming lipophilic ion pairs and clusters. FBAs are non-volatile from water but readily adsorb to suspended solids. They have low mobility in sediment and soil.

In water, FBAs are expected to be anionic as sulfonate groups are deprotonated at environmental pH (Poiger 1994). However, secondary and aromatic ring nitrogens may become protonated at low pH (pH < 6), resulting in the formation of zwitterions. Sulfonate groups will bind with dissolved cations in the environment, especially Na $^+$ , K $^+$ , Mg $^{2+}$  and Ca $^{2+}$ , forming ion pairs and clusters with higher partition coefficients (log K $_{OW}$ ). For example, the ion pair of FB-71 with calcium has a log K $_{OW}$  of 0.7–1.8, more than 2 orders of magnitude greater than the native FBA (Poiger 1994).

Chemicals in this group undergo rapid and reversible *E–Z* photo-isomerisation in the environment. In water, photo-stationary equilibrium is achieved within minutes and normally favours the non-fluorescent (*Z*)-isomer (Canonica et al. 1997). However, isomerisation rates decline in the presence of suspended solids, which shift the *E:Z* ratio at equilibrium toward the more strongly adsorbing (*E*)-isomer (Poiger 1994).

These chemicals (FBAs) rapidly adsorb to suspended solids in aqueous media. The adsorbed fraction can exceed 90% in activated sludge, leading to significant removal from wastewater in STPs (Poiger 1994). However, the adsorbed fraction in surface water is likely to be <5% based on available monitoring data for estuarine and coastal water in Japan (Hayashi et al. 2002; Managaki and Takada 2005). Sedimentation is nonetheless an important removal process for FBAs in the environment. For example, between April 1995 and April 1996, removal via sedimentation accounted for 27% of all FB-71 received by a small urbanised lake in Switzerland (Stoll 1997).

Chemicals in this group have low mobility in soil and sediment and are unlikely to leach from landfill or diffuse through soils via groundwater. Adsorption coefficients ( $K_{OC}$ ) in sands, sandy loams and loamy sands are 860–2,240 L/kg for FB-71 and 2,470–10,043 L/kg for FB-220 (OECD 2001; 2005a). Adsorption coefficients in sediments collected from the Glatt River in Switzerland are 1,025 and 4,186 L/kg, for the (Z)- and (E)-isomers of FB-71, respectively (Poiger 1994). These values are consistent with low mobility in the environment.

#### Degradation

Chemicals in this group (FBAs) have low potential for degradation in the environment. Based on available data, FBAs are neither readily nor inherently biodegradable in water. Chemicals in this group undergo rapid photo-degradation but this is likely to lead to persistent degradants. They are not expected to degrade in soil or sediment.

Experimental data were retrieved from the substance REACH dossiers for FB-71, FB-220 and FB-28 (REACH n.d.-a; n.d.-b; n.d.-c). No abiotic degradation data were identified for FB-28 and available biodegradation data is limited to an inherent biodegradation test study.

The chemical, FB-71 is not expected to biodegrade in water. The chemical is not readily biodegradable according to available test data. In a 28 day study conducted according to OECD TG 301 A, the chemical underwent only 10–20% degradation by removal of dissolved organic carbon (DOC). In a separate study conducted according to OECD TG 302 B, 98.8% degradation by removal of DOC was reported at 28 days, indicating inherent biodegradability. However, this was attributed to adsorption of the test substance to inoculum, as DOC removal reached 89.6% after only 3 hours duration. Therefore, FB-71 is neither readily nor inherently biodegradable in water.

The chemical, FB-220 is not expected to biodegrade in water. Only 1.2% degradation by removal of DOC was reported in a 28 day ready biodegradability test conducted according to OECD TG 301 A, while 0% degradation by O<sub>2</sub> consumption was reported in a separate 30 day test conducted according to OECD TG 301 D. In a 28 day inherent biodegradability test conducted according to OECD TG 302 B, degradation by DOC removal rose to 20.9% after 5 days but was approximately sustained throughout the remainder of the test. This behaviour is consistent with a slow adsorption process or the formation of persistent degradants. Therefore, FB-220 is neither readily nor inherently biodegradable in water.

The chemical, FB-28 is not expected to biodegrade in water. A study conducted according to OECD TG 302 B was terminated after 24 hours as 83.6% of test substance was removed by adsorption. As no reliable data were identified, biodegradation was read across from FB-220. The structures of FB-220 and FB-28 are similar, with identical backbones and sulfonate substituents on their aromatic rings, differing only by the presence of two additional sulfonates on FB-220. On this basis, they are expected to have similar degradation pathways. Therefore, FB-28 is conservatively assessed to be neither readily nor inherently biodegradable in water.

Photo-degradation may be a significant dissipation pathway for chemicals in this group (FBAs), but it is likely to lead to the formation of persistent degradants. The mechanism is thought to involve photo-oxidative cleavage of the internal stilbene, yielding alcohols and aldehydes, with further degradation leading to derivatives of melamine, which is persistent in the environment (AICIS 2022; AISE and Cefic 2004). The half-life of FBAs to direct photolysis in sunlit water is 3–6 hours according to laboratory studies of FB-71 and FB-220 (Kramer et al. 1996; Managaki and Takada 2005). However, dissipation rates in the environment will be seasonal and depth dependent, reaching their highest levels in the upper photic zone during summer but easing into winter (Poiger et al. 1999; Stoll and Giger 1997; Yamaji et al. 2010).

Chemicals in this group will not hydrolyse in the environment as they do not possess readily hydrolysable groups. In experimental studies conducted according to OECD TG 111, the half-lives of FB-71 and FB-220 to hydrolysis exceed one year under acidic (pH 4) and basic conditions (pH 9) (REACH n.d.-a; n.d.-b).

Based on available exposure and monitoring data, chemicals in this group are not expected to degrade in sediment or soil. Sediment core samples obtained from a Swiss lake in the 1990s contained a historical record of FBAs in layers deposited as far back as the 1950s (Poiger 1994; Stoll 1997). In an exposure study undertaken in Switzerland, soil plots amended with STP biosolids were reportedly contaminated with FB-71 throughout the 45 month test period (AISE and Cefic 2004). Available data are not sufficient to establish lifetimes in sediment and soil but support conservative half-lives exceeding 6 months.

#### Bioaccumulation

Chemicals in this group are not expected to bioaccumulate in organisms.

Measured partition coefficients of chemicals in this group, while dependent on speciation, are below than the level expected to cause significant bioaccumulation in aquatic and terrestrial life (log  $K_{OW} < 4.2$  and log  $K_{OW} < 2.0$ , respectively).

Available data for FB-71 indicates that chemicals in this group will not bioaccumulate in fish. According to an experimental bioaccumulation study conducted according to OECD TG 305 C, bioconcentration factors (BCF) for FB-71 in Eurasian carp (*Cyprinus carpio*) are 6.4–28 and 1.4–4.7 L/kg at dosage levels of 0.02 and 0.2 mg/L, respectively (NITE n.d.; REACH n.d.-a). These are well below the threshold for a bioaccumulation hazard to aquatic life (BCF < 2000).

#### **Environmental transport**

Chemicals in this group are not expected to undergo long range transport in the environment. Riverbed sediments containing FBAs will eventually migrate into coastal and marine environments. However, these sediments are unlikely to travel significant distances (Hayashi et al. 2002; Managaki and Takada 2005; SERI 2011).

### Predicted environmental concentration (PEC)

No Australian monitoring data were identified for chemicals in this group. International monitoring data and standard exposure modelling have been used to estimate reasonable worst-case PECs in affected compartments.

Considering river flows can consist entirely of treated effluent in some drier parts of Australia, a reasonable worst case PEC for chemicals in this group is 6.6  $\mu$ g/L per compound in rivers impacted by sewage outfall and 39  $\mu$ g/L FB-220 in rivers impacted by the pulp and paper industry. These values were derived from the highest reported concentrations in secondary sewage effluent and industrial effluent, respectively (SERI 2011; Yamaji et al. 2010).

In sediments affected by sewage outfall, a reasonable worst case PEC for chemicals in this group is 4.4 mg/kg per compound. This value was derived from recent monitoring data for sediments collected near effluent discharge sites in Stockholm, Sweden (SERI 2011).

The calculated PEC for chemicals in this group in Australian agricultural soil amended with biosolids is 0.86 mg/kg soil/year per compound. This was calculated using the highest reported biosolids concentration of 112 mg/kg per compound (Poiger et al. 1998), typical biosolids application rates and a soil bulk density of 1,300 kilograms per cubic metre (kg/m³) (EPHC 2009; Langdon et al. 2010):

Parameter	Value	Comment
PEC <sub>soil</sub>	0.86 mg/kg soil/year per compound	$PEC_{soil} = \frac{C_{biosolids} \times BIOSOLIDS_{land}}{SOILMIX_{depth} \times SOIL_{density}}$
C <sub>biosolids</sub>	112 mg/kg	Highest reported concentration in STP biosolids
BIOSOLIDS <sub>land</sub>	1 kg/m²/year	Default value
SOILMIX <sub>depth</sub>	0.1 m	Default value
SOILdensity	1,300 kg/m³	Default value

### **Monitoring data**

STP influent reportedly contains 10–100  $\mu$ g/L of FBAs (Kramer 1992). Based on available monitoring data, influent concentrations are <1–23  $\mu$ g/L per compound, removal rates in municipal STPs are 15–95% and residual concentrations in secondary effluent are <10  $\mu$ g/L, or <0.05–6.6  $\mu$ g/L per compound. FBA concentrations in STP biosolids are 56–169 mg/kg or 2.8–112 mg/kg per compound:

Chemical	Location	Ref	STP influent (µg/L)	Secondary effluent (µg/L)	Removal in STPs (%)	Biosolids (mg/kg)
FB-71	Sweden	1	≤1.0	0.059–0.49	>73	2.8–11 (4.5)
FB-71	Switzerland	2	7.1	2.6–4.5	(89)	55–105 (72)
FB-71	Switzerland	3	6.6–12.9 (10.5)	1.8–2.8 (2.4)	-	86– <b>112</b> (100)
FB-71	Switzerland	4		~2.9–5.7	-	<u> </u>
FB-71	Switzerland	5	-	1.9–3.6	-	-
FB-71	Germany	6	6.1–7.5	0.63 - 3.9	-	10.2–72
FB-71	Japan	7	2.9–8.2	0.68–4.9 (2.5)	15–79	-
FB-71	Japan	8	6.6-17.8	0.6– <b>6.6</b>	62.7-92.3	-
FB-71	Taiwan	9	-	0.042	-	-
FB-220	Sweden	1	~10	0.25 - 3.6	>90	15-62 (41)
FB-220	Germany	6	8.2-14.9	1.6-3.9	-	2.0-2.6*
FB-28	Sweden	1	~10	0.089-0.91	>90	35-83 (55)
FB-28	Switzerland	2	-	-	-	8–11 (9)
FB-28	Taiwan	9	-	1.58	-	-
Total FBAs	Sweden	1	20–24 (22)	0.43–5.1	-	56–160 (110)
Total FBAs	Switzerland	2	-	(9.6)	53–98	85–169 (118)

Overall averages are indicated in brackets.

References: [1] SERI (2011); [2] Poiger (1994); [3] Poiger et al. (1998); [4] Poiger et al. (1999); [5] Stoll (1997); [6] Van de Plassche et al. (1999); [7] Hayashi et al. (2002); [8] Yamaji et al. (2010); [9] Chen et al. (2006).

According to recent data obtained from the Swedish pulp and paper industry, average FBA concentrations in industrial effluent are <4  $\mu$ g/L per compound, excluding one outlier containing 39  $\mu$ g/L of FB-220 (SERI 2011).

Based on available data, FBA concentrations in surface water are <2  $\mu$ g/L per compound. This excludes three outliers measured in rivers during the 1970–80s in Sweden (8.3  $\mu$ g/L), the USA (40  $\mu$ g/L) and Japan ( $\leq$ 45  $\mu$ g/L):

Chemical	Location	Ref	River water (μg/L)	Lake water (μg/L)	Estaurine and Coastal water (µg/L)
FB-71	Sweden	1	<0.15	<0.02	<0.02
FB-71	Switzerland	2	0.036-0.439	-	-
FB-71	Switzerland	3	<0.6	-	-
FB-71	Switzerland	4	0.006-0.986	0.047-0.130	-
FB-71	Switzerland	5	-	0.053-0.098	-
FB-71	Germany	6	0.006-0.95	-	-
FB-71	Japan	7	0.1-0.4	-	~0.01–1
FB-71	Japan	8	-	-	0.001-1.3
FB-71	Japan	9	0.096–0.137 (0.123)	0.002-0.004	-
FB-71	USA	4	0.06-0.7	-	-
FB-71	USA	6	40 (0.7)	-	-
FB-71	Taiwan	10	<loq< td=""><td>-</td><td>-</td></loq<>	-	-
FB-220	Sweden	1	<0.3	<0.04	<loq< td=""></loq<>
FB-220	Germany	6	<0.002-0.152	-	-
FB-28	Sweden	1	<0.05	<0.02	<0.02
FB-28	Taiwan	10	0.1-0.145	-	-
Total FBAs	Sweden	1	<0.45	0.01-0.05	<0.03
Total FBAs	Sweden	6	≤8.3	-	-
Total FBAs	Japan	2, 6	≤45	-	-
Total FBAs	USA	4, 6	≤0.6	-	-

Overall averages indicated in brackets.

References: [1] SERI (2011); [2] Poiger (1994); [3] Poiger et al. (1999); [4] Stoll (1997); [5] Stoll and Giger (1997); [6] Van de Plassche et al. (1999); [7] Hayashi et al. (2002); [8] Managaki and Takada (2005); [9] Yamaji et al. (2010); [10] Chen et al. (2006).

Sediment concentrations vary by location and sample depth. Based on available data in upper sediment layers (0-5 cm), concentrations of chemicals in this group are expected to be  $\leq 4.4 \text{ mg/kg}$  dw per compound:

Chemical	Location	Ref	Depth (cm)	River sediment (mg/kg dw)	Lake sediment (mg/kg dw)	Coastal sediment (mg/kg dw)
FB-71	Sweden	1	0–2	-	≤0.43	<0.1, <b>4.4</b> *
FB-71	Switzerland	2	0–50	-	≤0.3	-
FB-71	Switzerland	3	0–5	=	0.41-1.41	-
FB-71	Switzerland	3	5–10	-	0.43-1.79	-
FB-71	Switzerland	3	10–15	-	<0.23-3.60	-
FB-71	Switzerland	3	15–20	-	0-1.98	-
FB-71	Switzerland	4	0–5	=	0.65-1.42	-
FB-71	Japan	2	-	0.1–3.4	-	-

Chemical	Location	Ref	Depth (cm)	River sediment (mg/kg dw)	Lake sediment (mg/kg dw)	Coastal sediment (mg/kg dw)
FB-71	Japan	5	0–2	0.02-1.55	-	0.02-0.37
FB-71	Japan	6	0–6	-	<0.1	-
FB-220	Sweden	1	0–2	-	<0.3	<0.1, 1.6*
FB-28	Sweden	1	0–2	-	<0.2	<0.1, 1.2*
Total FBAs	Sweden	1	0–2	-	<0.9	<0.2, 7.2*

<sup>\*</sup>Measured near an effluent discharge site.

References: [1] SERI (2011); [2] Poiger (1994); [3] Stoll (1997); [4] Stoll and Giger (1997); [5] Managaki and Takada (2005); [6] Yamaji et al. (2010).

Very little monitoring data for soil were identified. A regional PEC of 0.4 mg/kg soil/year FB-71 has been estimated in Switzerland based on a long-term exposure study, but this is unlikely to be representative of Australia (AISE and Cefic 2004).

## **Environmental effects**

## Effects on Aquatic Life

The toxicity of FBAs to aquatic organisms is well studied. No specific modes of action have been established. Aquatic toxicity tests are typically conducted under periodic exposure to a light source, indicating some potential for photo-degradation of the test substance. However, measured concentrations of FBAs, where available, were maintained during tests.

### **Acute toxicity**

The following measured median lethal and effect concentrations (LC50 and EC50, respectively) for freshwater model organisms across 3 trophic levels were retrieved from the REACH dossiers for chemicals in this group (REACH n.d.-a; n.d.-b; n.d.-c).

#### Fish

Chemical	Endpoint	Method
FB-71	96 h LC50 > 319 mg/L*	Danio rerio (Zebrafish) Measured concentrations Static 12 light and 12 dark hours/day OECD TG 203
FB-220	96 h LC50 > 1,000 mg/L*	D. rerio Nominal concentrations Static 12 light and 12 dark hours/day OECD TG 203
FB-28	96 h LC50 > 1,000 mg/L*	<ul><li>D. rerio</li><li>Nominal concentrations</li><li>Static</li><li>12 light and 12 dark hours/day</li><li>OECD TG 203</li></ul>

<sup>\*</sup>No lethal effects observed at the highest test concentration.

#### **Invertebrates**

Chemical	Endpoint	Method
FB-71	48 h EC50 = 6.85 mg/L	Ceriodaphnia dubia Immobilisation Nominal concentrations Static 16 light and 8 dark hours/day Australian NSW EPA test guideline
FB-220	48 h EC50 > 113 mg/L*	Daphnia magna Immobilisation Measured concentration (limit test) Static 16 light and 8 dark hours/day OECD TG 202
FB-28	48 h EC50 > 100 mg/L	<ul><li>D. magna</li><li>Immobilisation</li><li>Nominal concentration (limit test)</li><li>Static</li><li>12 light and 12 dark hours/day</li><li>OECD TG 202</li></ul>

<sup>\*</sup>No lethal effects observed at the highest test concentration.

# Algae

Chemical	Endpoint	Method
FB-71	72 h EC50 = 82.5 mg/L	Raphidocelis subcapitata Growth rate Nominal concentrations Static Continuous illumination OECD TG 201
FB-220	72 h EC50 > 1,000 mg/L	Desmodesmus subspicatus Growth rate Nominal concentrations Static Continuous illumination OECD TG 201
FB-28	72 h EC50 > 123 mg/L*	R. subcapitata Growth rate Nominal concentrations Static Continuous illumination OECD TG 201

<sup>\*</sup>No lethal effects observed at the highest test concentration.

# **Chronic toxicity**

The following measured lowest and no observed effect concentrations (LOEC and NOEC, respectively) were retrieved from the REACH registration dossiers for chemicals in this group (REACH n.d.-a; n.d.-b; n.d.-c):

### Fish

Chemical	Endpoint	Method
FB-71	30 d NOEC = 2.5 mg/L	D. rerio Survival Nominal concentrations Flow-through 16 light and 8 dark hours/day OECD TG 210
FB-28	35 d LOEC = 11.2 mg/L	<ul><li>D. rerio</li><li>Survival</li><li>Measured concentrations</li><li>Flow-through</li><li>16 light and 8 dark hours/day</li><li>OECD TG 210</li></ul>

### Invertebrates

Chemical	Endpoint	Method
FB-71	21 d NOEC = 11.3 mg/L	<ul><li>D. magna</li><li>Reproduction</li><li>Nominal concentrations</li><li>Semi-static</li><li>16 light and 8 dark hours/day</li><li>OECD TG 211</li></ul>
FB-220	21 d NOEC = 6.59 mg/L	<ul><li>D. magna</li><li>Reproduction</li><li>Measured concentrations</li><li>Semi-static</li><li>16 light and 8 dark hours/day</li><li>OECD TG 211</li></ul>
FB-28	21 d NOEC = 3.22 mg/L	<ul><li>D. magna</li><li>Reproduction</li><li>Measured concentrations</li><li>Semi-static</li><li>16 light and 8 dark hours/day</li><li>OECD TG 211</li></ul>

### Algae

Chemical	Endpoint	Method
FB-71	72 h NOEC = 11.7 mg/L	R. subcapitata Growth rate Nominal concentrations Static Continuous illumination OECD TG 201
FB-220	96 h NOEC = 500 mg/L	D subspicatus Growth rate Nominal concentrations Static Continuous illumination OECD TG 201
FB-28	72 h NOEC = 30.8 mg/L	R. subcapitata Growth rate Nominal concentrations Static Continuous illumination OECD TG 201

# Effects on sediment dwelling life

No sediment toxicity data were identified for chemicals in this group.

# Effects on terrestrial Life

The following effect concentrations for earthworms were retrieved from the REACH dossiers for chemicals in this group (REACH n.d.-a; n.d.-b; n.d.-c):

Chemical	Endpoint	Method
FB-71	14 d LC50 > 1,000 mg/kg	Eisenia fetida Mortality Nominal concentrations Continuous illumination OECD TG 207
FB-220	14 d LC50 > 10,000 mg/kg	E. fetida Mortality Nominal concentration (limit test) Photoperiod: not specified OECD TG 207
FB-28	14 d LC50 > 5,000 mg/kg*	E. fetida Mortality Nominal concentration (limit test) Photoperiod: not specified OECD TG 207

<sup>\*</sup>No sublethal effects observed.

## Predicted no-effect concentration (PNEC)

Available data are sufficient to determine PNECs for chemicals in this group in water and soil, but not sufficient to determine PNECs in sediment.

### **Aquatic compartment**

Aquatic PNECs for FB-71 and FB-28 were derived from measured chronic test data for fish and an assessment factor of 10. This assessment factor was selected as reliable chronic test data are available for three trophic levels (EPHC 2009). An assessment factor of 50 was used to calculate a PNEC FB-220, as reliable chronic test data are limited to aquatic invertebrates and algae:

Chemical	Pivotal endpoint (mg/L)	Endpoint	Assessment Factor	PNEC (µg/L)
FB-71	2.5	Fish NOEC	10	250
FB-220	6.59	Invertebrate NOEC	50	132
FB-28	3.22	Invertebrate NOEC	10	322

#### Sediment compartment

Sediment PNECs for chemicals in this group were not determined as no experimental toxicity data for sediment dwelling organisms were identified (EPHC 2009).

### **Terrestrial compartment**

Soil PNECs for chemicals in this group were derived from available data for earthworms and an assessment factor of 1000 (EPHC 2009). This assessment factor was selected as acute data are limited to one terrestrial macro-organism, per compound:

Chemical	Pivotal endpoint (mg/kg)	Assessment Factor	PNEC (mg/kg)
FB-71	>1,000	1,000	>1
FB-220	>10,000	1,000	>10
FB-28	>5,000	1,000	>5

# Categorisation of environmental hazard

The categorisation of the environmental hazards of the assessed chemicals according to Australian Environmental Criteria for Persistent, Bioaccumulative and/or Toxic Chemicals (DCCEEW n.d.) is presented below:

#### Persistence

Persistent (P). Based on available degradation data in water and environmental monitoring data for sediment and soil, chemicals in this group are categorised as persistent.

#### Bioaccumulation

Not bioaccumulative (Not B). Based on low measured bioconcentration factors (BCF) in fish and low octanol-water partition coefficients (log  $K_{\text{OW}} < 4.2$ ), chemicals in this group are categorised as not bioaccumulative.

### **Toxicity**

Not toxic (Not T). Based on acute toxicity endpoints exceeding 1 mg/L and chronic toxicity endpoints exceeding 0.1 mg/L, chemicals in this group are categorised as not toxic.

# Environmental risk characterisation

Based on the PEC and PNEC values determined above, the following Risk Quotient (RQ=PEC ÷ PNEC) have been calculated for release of chemicals in this group into surface water and soil.

#### FB-71:

Compartment	PEC	PNEC	RQ
Surface water (sewage outfall)	6.6 µg/L	250 μg/L	0.03
Soil	0.86 mg/kg soil/year	>1 mg/kg	<0.86

#### FB-220:

Compartment	PEC	PNEC	RQ
Surface water (sewage outfall)	6.6 µg/L	132 µg/L	0.05
Surface water (industrial effluent)	39 μg/L	132 µg/L	0.30
Soil	0.86 mg/kg soil/year	>10 mg/kg	<0.09

#### FB-28:

Compartment	PEC	PNEC	RQ
Surface water (sewage outfall)	6.6 µg/L	322 μg/L	0.02
Soil	0.86 mg/kg soil/year	>5 mg/kg	<0.18

For water and soil, a worst-case RQ of less than 1 indicates that chemicals in this group are not expected to pose a significant risk to the environment based on estimated emissions, as environmental concentrations are below levels likely to cause harmful effects.

A RQ for sediment could not be determined for chemicals in this group, due to a lack of information about toxicity to sediment dwelling organisms. The highest FBA levels occur near effluent discharge sites in urbanised catchments, where concentrations can exceed 1 mg/kg dw per compound. However, available monitoring data indicate that concentrations rapidly decline away from these sites (SERI 2011; Stoll 1997). Given these generally low sediment concentrations, and in the absence of evidence of significant toxicity to sediment-dwelling life, these chemicals are not anticipated to pose a significant risk to the ecosystems in the sediment compartment.

#### Uncertainty

This evaluation was conducted based on a set of information that may be incomplete or limited in scope. Some relatively common data limitations can be addressed through use of conservative assumptions (OECD 2019) or quantitative adjustments such as assessment factors (OECD 1995). Others must be addressed qualitatively, or on a case-by-case basis (OECD 2019).

The most consequential areas of uncertainty for this evaluation are:

- Insufficient information is available to characterise the terrestrial and sediment toxicity
  of chemicals in this evaluation. The outcomes of the evaluation may change if
  additional information becomes available.
- No Australian monitoring data are available for chemicals in this evaluation and
  overseas information is used as a surrogate. The outcomes of this evaluation may
  change if new monitoring information become available to indicate that environmental
  concentrations of these chemicals in Australia are higher than currently assessed.

# References

AICIS (Australian Industrial Chemicals Introduction Scheme) (2022) <u>Evaluation statement:</u> <u>1,3,5-Triazine-2,4,6-triamine (melamine)</u>, AICIS website, accessed 14 June 2023.

AISE and Cefic (Association for Soaps, Detergents and Maintenance Products, European Chemical Industry Council) (2004) *Fluorescent Brightener FWA-1*, AISE and Cefic, accessed 28 February 2023.

Canonica S, Kramer JB, Reiss D and Gygax H (1997) 'Photoisomerization kinetics of stilbene-type fluorescent whitening agents', *Environmental science & technology,* **31**(6), pp 1754-1760, doi:doi.org/10.1021/es960748a.

Chen H-C, Wang S-P and Ding W-H (2006) 'Determination of fluorescent whitening agents in environmental waters by solid-phase extraction and ion pair liquid chromatography–tandem mass spectrometry', *Journal of Chromatography A,* **1102**(1-2), pp 135-142, doi:doi.org/10.1016/j.chroma.2005.10.025.

DAFF (Department of Agriculture, Fisheries and Forestry) (n.d.) <u>Australia's Wood and Paper</u> <u>Industry</u>, DAFF website, accessed 19 May 2023.

DCCEEW (Department of Climate Change, Energy, the Environment and Water) (n.d.) <u>Australian Criteria for Persistent, Bioaccumulative and/or Toxic Chemicals,</u> DCCEEW website, accessed 28 August 2023.

DeLima Associates (n.d.) <u>Consumer Product Information Database</u>, DeLima Associates website, accessed 11 July 2023.

EPHC (Environment Protection and Heritage Council) (2009) <u>Environmental Risk</u> <u>Assessment Guidance Manual for Industrial Chemicals</u>, EPHC, accessed 11 July 2022.

Hayashi Y, Managaki S and Takada H (2002) 'Fluorescent whitening agents in Tokyo Bay and adjacent rivers: Their application as anthropogenic molecular markers in coastal environments', *Environmental science & technology*, **36**(16), pp 3556-3563, doi:doi.org/10.1021/es011352o.

Industry Edge (2019) <u>Assessment of Australian paper & paperboard recycling infrastructure</u> and 2018-19 exports, including to China, DCCEEW website, accessed 8 August 2023.

Khan MI, Wang L and Padhye R (2023) 'Textile waste management in Australia: A review', *Resources, Conservation & Recycling Advances*, pp 200154, doi:doi.org/10.1016/j.rcradv.2023.200154.

Kramer JB (1992) 'Fluorescent Whitening Agents', in: Oude NT (eds) *Detergents:* . *Anthropogenic Compounds, vol 3 / 3F*, Springer Berlin Heidelberg, Berlin, Heidelberg.

Kramer JB, Canonica S, Hoigné J and Kaschig J (1996) 'Degradation of fluorescent whitening agents in sunlit natural waters', *Environmental science & technology,* **30**(7), pp 2227-2234, doi:doi.org/10.1021/es950711a.

Langdon KA, Warne MSTJ and Kookana RS (2010) 'Aquatic hazard assessment for pharmaceuticals, personal care products, and endocrine-disrupting compounds from

biosolids-amended land', *Integrated Environmental Assessment and Management*, **6**(4), pp 663-676, doi:doi.org/10.1002/ieam.74.

Managaki S and Takada H (2005) 'Fluorescent whitening agents in Tokyo Bay sediments: molecular evidence of lateral transport of land-derived particulate matter', *Marine chemistry*, **95**(1-2), pp 113-127, doi:doi.org/10.1016/j.marchem.2004.08.007.

NCBI (National Center for Biotechnology Information) (n.d.) <u>PubChem Compound Summary for CID 6434593</u>, <u>Fluorescent Brightener 28</u>, National Library of Medicine (US), NCBI, accessed 28 February 2023.

NICNAS (National Industrial Chemicals Notification and Assessment Scheme) (1995) <u>Full Public Report: FPC-159</u>, NICNAS, accessed 9 August 2023.

NITE (National Institute of Technology and Evaluation) (n.d.) <u>Japan CHEmicals Collaborative Knowledge Database (J-CHECK)</u>, NITE, accessed 28 February 2023.

NPI (National Pollutant Inventory) (n.d.) NPI data, NPI website, accessed 23 August 2023.

NSA (Norske Skog Paper Mills (Australia) Ltd.) (2022) Norske Skog Boyer, <u>Annual Review</u> for the Boyer Mill, NSA, accessed 16 August 2023.

OECD (Organisation for Economic Co-operation and Development) (1995) <u>Guidance document for aquatic effects assessment</u>, OECD, Paris.

OECD (Organisation for Economic Cooperation and Development) (2001) SIDS Initial Assessment Report for SIAM 13, *Fluorescent Brightener* 220, OECD, accessed 23 February 2023.

OECD (Organisation for Economic Cooperation and Development) (2004) Emission Scenario Documents, *Emission Scenario Document on Textile Finishing Industry*, OECD, accessed 15 August 2023.

OECD (Organisation for Economic Cooperation and Development) (2005a) SIDS Initial Assessment Report for SIAM 21, *Fluorescent Brightener FWA-1*, OECD, accessed 23 February 2023.

OECD (Organisation for Economic Cooperation and Development) (2005b) SIDS Initial Assessment Report for SIAM 20, <u>C.I. Fluorescent Brightener 28/113</u>, OECD, accessed 8 March 2023.

OECD (Organisation for Economic Cooperation and Development) (2009) Emission Scenario Documents, *Emission Scenario Document Pulp, Paper and Board Industry*, OECD, accessed 15 August 2023.

OECD (The Organisation for Economic Co-operation and Development) (2019) <u>Guiding Principles and Key Elements for Establishing a Weight of Evidence for Chemical Assessment, Series on Testing and Assessment No. 311, Environment, Health and Safety Division, Environment Directorate, OECD, accessed 14 June 2023.</u>

OECD (Organisation for Economic Cooperation and Development) (n.d.) <u>OECD Existing Chemicals Database.</u> OECD, accessed 28 March 2023.

Plant EL, Smernik RJ, van Leeuwen J, Greenwood P and Macdonald LM (2014) 'Changes in the nature of dissolved organics during pulp and paper mill wastewater treatment: a multivariate statistical study combining data from three analytical techniques', *Environmental Science and Pollution Research*, **21**, pp 4265-4275, doi:doi.org/10.1007/s11356-013-2351-0.

Poiger T (1994) <u>Behavior and fate of detergent-derived fluorescent whitening agents in sewage treatment</u>, [PhD thesis], ETH Zurich, accessed 15 April 2023.

Poiger T, Field JA, Field TM, Siegrist H and Giger W (1998) 'Behavior of fluorescent whitening agents during sewage treatment', *Water Research*, **32**(6), pp 1939-1947, doi:doi.org/10.1016/S0043-1354(97)00408-9.

Poiger T, Kari FG and Giger W (1999) 'Fate of fluorescent whitening agents in the River Glatt', *Environmental science & technology*, **33**(4), pp 533-539, doi:doi.org/10.1021/es9804969.

REACH (Registration, Evaluation, Authorisation and Restriction of Chemicals) (n.d.-a) REACH registration dossier for Disodium 4,4'-bis[(4-anilino-6-morpholino-1,3,5-triazin-2-yl)amino]stilbene-2,2'-disulphonate (CAS RN 16090-02-1), REACH, accessed 27 February 2023.

REACH (Registration, Evaluation, Authorisation and Restriction of Chemicals) (n.d.-b) REACH registration dossier for Tetrasodium 4,4'-bis[f4-[bis(2-hydroxyethyl)amino]-6-(4-sulphonatoanilino)-1,3,5-triazin-2-yl]amino]stilbene-2,2'-disulphonate] (CAS RN 16470-24-9), REACH, accessed 22 February 2023.

REACH (Registration, Evaluation, Authorisation and Restriction of Chemicals) (n.d.-c) <u>REACH registration dossier for Disodium 4,4'-bis[6-anilino-[4-[bis(2-hydroxyethyl)amino]-1,3,5-triazin-2-yl]amino]stilbene-2,2'-disulphonate (CAS RN 4193-55-9), REACH, accessed 24 April 2023.</u>

Scholes ER (2021) <u>The causes of variability in the membrane bioreactor treatment of pulp and paper wastewater</u>, [PhD thesis], Monash University, accessed 13 July 2023.

SERI (IVL Swedish Environmental Research Institute Ltd.) (2011) Results from the Swedish National Screening Programme 2010, <u>Subreport 2. Fluorescent Whitening Agents</u>, SERI, accessed 11 May 2023.

Stoll J-MA (1997) *Fluorescent whitening agents in natural waters*, [PhD thesis], ETH Zurich, accessed 5 April 2023.

Stoll J-MA and Giger W (1997) 'Determination of detergent-derived fluorescent whitening agent isomers in lake sediments and surface waters by liquid chromatography', *Analytical chemistry*, **69**(13), pp 2594-2599, doi:doi.org/10.1021/ac9701952.

UNECE (United Nations Economic Commission for Europe) (2017) <u>Globally Harmonized System of Classification and Labelling of Chemicals (GHS), Seventh Revised Edition,</u>
UNECE, accessed 24 April 2023.

UNEP (United Nations Environment Programme) (1987) <u>The Montreal Protocol on Substances that Deplete the Ozone Layer,</u> UNEP, Ozone Secretariat, accessed 6 June 2023.

UNEP (United Nations Environment Programme) (2001) <u>The Stockholm Convention on Persistent Organic Pollutants</u>, UNEP, Secretariat of the Stockholm Convention, accessed 6 June 2023.

UNEP & FAO (United Nations Environment Programme & Food and Agriculture Organization of the United Nations) (1998) *Rotterdam Convention on the Prior Informed Consent Procedure for Certain Hazardous Chemicals and Pesticides in International Trade,* UNEP & FAO, accessed 6 June 2023.

US EPA (United States Environmental Protection Agency) (2020) <u>2020 CDR Industrial Processing and Use,</u> US EPA. Downloaded.

US EPA (United States Environmental Protection Agency) (n.d.) <u>TSCA Safer Chemical Ingredients List</u>, US EPA, accessed 11 July 2023.

Van de Plassche EJ, Bont PFH and Hesse JM (1999) <u>Exploratory Report Fluorescent Whitening Agents (FWAs)</u>, National Institute of Public Health and the Environment, Bilthoven, Netherlands.

Yamaji N, Hayakawa K and Takada H (2010) 'Role of photodegradation in the fate of fluorescent whitening agents (FWAs) in lacustrine environments', *Environmental science & technology*, **44**(20), pp 7796-7801, doi:doi.org/10.1021/es100465v.

