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#### Submission to Science of the Total Environment

# Pesticide use in the wheat-maize double cropping systems of the North China Plain: Assessment, field study, and implications

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#### 1 Abstract

In the North China Plain (NCP), rising inputs of pesticides have intensified the environmental
impact of farming activities in recent decades by contributing to surface water and
groundwater contamination. In response to this, the Chinese government imposed stricter
regulations on pesticide approval and application, and better monitoring strategies are being
developed. However, sufficient and well-directed research on the accumulation and impact of
different pesticides is needed for informed decision-making.

In this study, current pesticide use, and recent and current research on water contamination by pesticides in the NCP are reviewed and assessed. Additionally, a small-scale field study was performed to determine if residuals from currently-used pesticides in the NCP can be detected in surface water, and in connected shallow groundwater. The contaminants of interest were commonly used pesticides on winter wheat-summer maize fields (the dominant cropping system in the NCP), such as 2,4-D and atrazine. Sampling took place in May, July, and October 2013; and March 2014.

Results from our literature research showed that sampling is biased towards surface water monitoring. Furthermore, most studies focus on organic chlorinated pesticides (OCPs) like the isomers of dichlorodiphenyltrichloroethane (DDT) and hexachlorocyclohexane (HCH), which were banned in China in 1983. However, currently-used herbicides like 2,4-D and atrazine were detected in river water and groundwater in all samplings of our field study. The highest concentrations of 2,4-D and atrazine were found in the river water, ranging up to 3.00 and 0.96 µg/L, respectively.

The monitoring of banned compounds was found to be important because several studies
indicate that they are still accumulating in the environment and/or are still illegally in use.
However, supported by our own data, we find that the monitoring in groundwater and surface

- 25 water of currently permitted pesticides in China needs equal attention, and should therefore be
- 26 increased.
- 27 **Keywords**: North China Plain; Surface water groundwater interaction; Pesticides; Atrazine;
- 28 2,4-D

#### 29 1. Introduction

30 The North China Plain (NCP) has an output of the national maize and wheat production of 31 about 60% and 40%, respectively (Zhao and Guo, 2013), which makes it a crucial area for 32 China's domestic food supply. It is therefore also referred to as China's bread basket. In 33 recent decades, the total cultivated area in the NCP has decreased due to competition for land 34 use from industry and urbanization. To continue meeting the food demands of the rapidly growing Chinese population, higher production efficiencies needed to be achieved, so that the 35 loss of production areas could be compensated. This goal was successfully met by introducing 36 37 modern agricultural practices in the 1970s and 1980s (such as the use of fertilizers and 38 pesticides) and by increasing irrigated agriculture. On the down-side, pesticide overuse has nowadays become common and poses a risk to valuable surface water and groundwater 39 resources (Zhang et al., 2011). This negative effect of the agricultural activities on potential 40 41 drinking water is especially problematic in the provinces of the NCP because of its high 42 population density and its relatively low amounts of available water resources (Varis and 43 Vakkilainen, 2001). Therefore, further decline in water quantity and quality is an imminent challenge for the area (van Oort et al., 2016; Sun et al., 2012; Zheng et al., 2015). 44

45 To address the problem of agricultural water pollution by pesticides, studies in recent years have dealt with the optimization of applied pesticide doses (Zhang et al., 2013), the need to 46 47 develop a systematic risk assessment approach for water pollution by pesticides (Zhao and Pei, 2012), and the enhancement of pesticide management in general (Hamburger, 2002; Wei 48 49 et al., 2007; Zhen et al., 2005). Further studies on pesticide residues in waterways and 50 agricultural soils have repeatedly detected banned organochlorine insecticides, such as 51 hexachlorocyclohexane (HCH) and dichlorodiphenyltrichloroethane (DDT) (Feng et al., 2011; Sun et al., 2010). Risk mapping of these compounds, and the detection of potentially 52

53 still active application and bioaccumulation, are important tasks. However, other disputed pesticides such as atrazine (which has been banned in the EU since 2004, but is legally used 54 55 in China) may potentially accumulate in the environment and cause negative environmental 56 impacts, too. Therefore, monitoring of currently permitted and applied pesticides should also 57 be considered. This includes obtaining knowledge about pesticide choices, means of application and disposal, environmental loads, and potential environmental impacts. Based on 58 59 this information, upcoming problems can be better identified, and pesticides can be chosen 60 and applied more wisely by the farmers. However, a comprehensive overview and review of 61 commonly applied pesticides on Chinese winter wheat-summer maize rotations (and their 62 loads in regional water resources) are very limited, and the availability of detailed statistics on pesticide use in China is extremely low (World Bank, 2010). 63

Upon this background, the aim of this study was (a) to review the current use of pesticides within the NCP, (b) to assess the recent and current research and monitoring of pesticides in water resources in the NCP, (c) to perform a small-scale field study on the occurrence of currently-used pesticides in surface water and connected groundwater, (d) to assess (supported by our findings in the field study) if currently used pesticides might cause environmental problems and if this is sufficiently addressed in the current monitoring efforts , and (e) to give recommendations for future study focus within the region.

The paper includes a comprehensive review part based on publicly available information and a field investigation at Fu River (Hebei Province) in order to investigate the current pesticide use and occurrence in the NCP. The agricultural cultivation at the field site is typical for this region, and although the study is limited in time, the findings presented illustrate the limitations in the current monitoring programs and are thereby an important contribution to the recommendations and conclusions of this paper.

#### 77 2. Material and methods

#### 78 **2.1. Field site**

79 A field site located at Fu River, the main incoming river to the Baiyang Lake area in Hebei Province, China, was selected as an example for the wheat-maize double cropping system in 80 81 the NCP (Figure 1a-b). The vicinity of the field site to surface water was chosen in order to 82 investigate surface water-groundwater interactions and pollution exchange between the environmental compartments. A monitoring network (consisting of four shallow groundwater 83 wells of which three were used for pesticide sampling) was set up on an agricultural field 84 85 (about 70x240 m) that extends to about 5 m to the river bank (Figure 1c). The agricultural plot (owned by one household) is surrounded by other fields with different ownership, but the 86 87 same cropping system.



Fig. 1 (a) Location of the study site, (b) aerial view of the field site, and (c) cross-section of the sampling wells
 (GW1 only used for ion-analysis and general water chemistry) SINGLE-COLUMN COLOUR FIGURE

91 The area around Fu River is characterized by pressures from agricultural, domestic, and

92 industrial water users. Numerous paper mills, and chemical, battery and petrochemical plants

- 93 are located in the upstream part of Fu River (Hu et al., 2010a), and it receives an approximate
- 94 amount of over 100 000 m<sup>3</sup> of sewage and treated wastewater from Baoding city per day (Qi
- 95 et al., 2012). Previous studies show that the point and non-point pollution sources along the
- 96 flow path of Fu River increase levels of organic chlorinated pesticides (OCPs),

97 polychlorinated biphenyls (PCBs), and polycyclic aromatic hydrocarbons (PAHs) in the river—especially near industrial areas (Hu et al, 2010b). Studies at the river mouth (entering 98 99 into Baiyang Lake) show extremely high loads of the nutrients ammonium and potassium 100 (e.g. Zhao et al., 2012). This indicates contamination from poorly treated sewage that is led 101 into the river (Xu et al., 2012), or possibly from direct discharge of animal manure as 102 observed in other regions of China (Strokal et al., 2016). Temporal nitrate peaks that occur 103 simultaneously with increased sediment loads in the river imply contamination by surface 104 runoff of agricultural pollutants (possibly also pesticides) from adjacent field areas after 105 rainstorms (Yan et al., 2005).

#### 106 2.2. Field site management and pesticide application

107 The field site was managed according to the local customs of the area (described in more 108 detail in Brauns et al., 2016). The successive winter wheat-summer maize rotation 109 encompassed the growing season for wheat from October 2012 to June 2013 and that of 110 maize from June 2013 to October 2013. Irrigation with surface water took place three times 111 during the growing period of winter wheat. Towards the end of the study period, the region 112 from about 1 -2 km upstream of the field site until the entrance of Fu River into the Baiyang 113 Lake system was expected to become flooded due to unusually high summer rainfalls, which 114 exceeded the 10-year average by 57%. Therefore, the farmers along this section of the river 115 refrained from planting winter wheat, and the land became fallow (and flooded) during the 116 remaining study period.

Based on interviews with the farmer, pest and weed management at the sampling site itself was done via chemical crop protection which took place twice in the winter wheat growing season (with tribenuron-methyl and dimethoate) and twice during the growing season of the summer maize (paraquat and acetochlor) via knapsack spraying. More detailed information on

121 planting times, irrigation, and times and amounts of pesticide application can be found in 122 Appendix A Table A.2. The pesticide application in the study year was not identical to the 123 one in previous years, where atrazine had been used instead of acetochlor, and additional 124 insecticides had been applied in the fall. Application rates in the study year were mostly in 125 accordance to the recommendations on the labels of the pesticide containers. According to 126 statements from the farmer however, it seemed that the exact application amount was 127 sometimes based on avoiding leftovers in pesticide packages/bottles, rather than on judging 128 the actual needed application rate based on observation in the field, e.g. the amount of weed 129 sprouting.

Additional farmer interviews indicated that treatment with other substances in neighboring fields was likely. For example, either 2,4-D or atrazine is often also used as post emergence pesticide on winter wheat in early spring (March to April). The discovery of empty containers from other substances (e.g. triadimefon) in the ambient environment of the study field also indicated that additional compounds were used, at least in the surrounding area (Figure 2).



136

Fig. 2 (a) Improperly disposed insecticide/fungicide bags that were found at the field site (tribenuron-methyl and triadimefon, respectively), (b) pollution in the adjacent river, and (c) empty insecticide bottle (8% dimethoate and 2% cyhalothrin) found in the river next to the sampling site. SINGLE-COLUMN COLOUR FIGURE

#### 140 **2.3. Field sampling and chemical analysis**

- 141 1 L water samples were obtained from river and from groundwater wells at a distance of 6,
- 142 11, and 41 m from the river (corresponding to GW2, GW3, and GW4 on Figure 1c) in May,

143 July and October 2013, and in March 2014. The sampled water was filled to the rim into 144 brown glass bottles, immediately cooled to 4°C, insulated, and shipped by courier to the 145 laboratory in Denmark (Eurofins, Glostrup). The samples were analyzed for a standard 146 package of different pesticides, which included some of the most universally used herbicides 147 in the NCP (and at the field) site as identified in section 3.2. (a full list of analytes is given in 148 Appendix A Table A.2). Analysis was done via gas chromatography mass spectrometry 149 (GC/MS) for chlorphenols and dichlobenil, and via liquid chromatography tandem mass 150 spectrometry (LC-MS/MS) for all other pesticides. The detection limit was 0.01 µg/L for all 151 compounds.

#### 152 **3. Results and discussion**

#### 153 **3.1. Detected pesticides in the field study**

154 Detected pesticides and their degradation products (or, in the case of 2,6-dichlorphenol,

155 intermediates) in the field study included mainly 2,4-D, 2,4-dichlorphenol (metabolite of 2,4-

156 D), 2,6-dichlorphenol (intermediate to 2,4-D), atrazine and the atrazine-metabolites

157 hydroxyatrazine, deethylatrazine, and deisopropylatrazine, terbutylazine, MCPA, bentazone,

158 DNOC, BAM (metabolite of chlorthiamid, dichlobenil and fluopicolide), and dichlorprop (see

159 Table 1).

- 160 Table 1 Detected concentrations in ug/l of selected herbicides in river water (RW) and groundwater (GW) samples
- 161 (see Figure 1 for the location of the sampling points), and drinking water limits (DWL) for China and for the USA
- 162 (for comparison, the standard of the European Drinking Water Directive is 0.1µg/L for each individual pesticide,
- 163 and 0.5µg/L for the sum of all pesticides detected in the sample). Data for the Chinese standard is according to the
- 164 guideline GB 5749-2006.

<b>Pesticide</b> (all values in µg/L)	29 May 2013	22 July 2013	07 October 2013	25 March 2014	DWL	
	RW GW2GW3GW4	RW GW2GW3GW4	RW GW2GW3GW4	RW GW2GW3GW4	CN USA	
2,4-D and related compounds						
2,4-D <sup>1</sup>	<b>3.00</b> 0.41 <b>2.30</b> 0.70	0.27 0.09 0.10 0.31	0.51 0.03 0.03 0.01	0.65 0.01 0.05 0.02	30 70*	
2,4-dichlorophenol (M)	0.09 0.14 0.09 0.14	$0.02 \ 0.10 \ 0.09 \ 0.07$	0.06 0.03 0.04 0.03	0.01 ND ND ND	- 20**	
2,6-dichlorophenol (I)	ND 0.02 ND 0.01	$0.01 \ 0.03 \ 0.16 \ 0.02$	0.01 ND ND ND	0.01 ND ND ND	- 4***	
Atrazine and related compounds						
Atrazine	0.96 0.33 0.91 0.40	$0.32 \ 0.29 \ 0.29 \ 0.49$	$0.49 \ 0.19 \ 0.20 \ 0.20$	0.54 0.12 0.19 0.15	2 3*	
Hydroxyatrazine (M)	<b>9.70</b> 0.46 0.37 0.47	$0.31 \ 0.59 \ 0.54 \ 0.49$	<b>3.00</b> 0.55 0.39 0.48	<b>1.60</b> 0.26 0.33 0.20		
Desethylatrazine (M)	$0.07 \ 0.04 \ 0.05 \ 0.04$	$0.17 \ 0.09 \ 0.07 \ 0.16$	$0.03 \ 0.01 \ 0.02 \ 0.04$	$0.05 \ 0.03 \ 0.04 \ 0.08$		
Deisopropylatrazine (M)	$0.07 \ 0.05 \ 0.05 \ 0.04$	$0.04 \ 0.16 \ 0.08 \ 0.09$	ND ND ND 0.01	0.13 0.01 0.02 0.02		
Other compounds						
Terbuthylazine	ND ND 0.02 0.01	ND ND ND 0.02	ND ND ND ND	ND ND ND ND		
MCPA <sup>2</sup>	0.06 0.03 0.05 0.02	0.03 ND ND 0.01	ND ND ND ND	0.04 ND ND ND	- 10**	
Bentazone	0.03 0.02 ND 0.02	0.01 ND ND ND	0.03 0.08 0.07 0.07	0.01 0.03 0.03 0.04	300 20**	
DNOC <sup>3</sup>	0.06 ND ND ND	0.04 ND ND ND	0.02 ND ND ND	0.11 ND ND ND		
BAM <sup>4</sup>	0.06 0.01 ND 0.04	ND ND ND ND	ND ND ND ND	ND ND ND ND		
Dichlorprop	ND ND ND ND	0.02 ND ND ND	0.04 ND ND ND	ND ND ND ND		

Values >1  $\mu$ g/L are highlighted in bold; ND = Not detected (< 0.01  $\mu$ g/L); M = Metabolite; I = Intermediate 1 = 2,4-Dichlorophenoxyacetic

165 166 167 168 acid; 2 =2-methyl-4-chlorophenoxyacetic; 3= Dinitro-ortho-cresol; 4=2,6-dichloro-benzamide; \*=National standard; \*\*=Federal guideline; \*\*\*State guideline

169 The pre and post emergence herbicides 2,4-D, atrazine, and its metabolite hydroxyatrazine

170 were the most abundantly detected species. They were found throughout all four samplings

171 and in all of the samples, with peak values of 3.00  $\mu$ g/L (2,4-D), 0.96  $\mu$ g/L (atrazine), and

172 9.70 µg/L (hydroxyatrazine). The range of our measured concentrations is similar to results

173 from a study in the Beijing area, where herbicides in surface water were detected in

174 concentrations up to 5.1  $\mu$ g/L (Deuerlein et al. 2001a). All of the observed peak

175 concentrations occurred in river water in May 2013. This peak time in May might be due to

176 runoff from field irrigation after the spring application of herbicides (see Figure 3 for

177 irrigation times, rainfall events, and herbicide application times), and possibly from spray

178 drift during application. Discarded containers adjacent to—or even directly into—the river

179 might be an additional cause of the rather surprising peak in river water before the actual

180 raining season (when a stronger effect of surface runoff due to heavy rainfall events could be

181 expected).



183 Fig. 3 Detections of atrazine and 2,4-D and selected degradation products and isomers for river water (RW) and 184 groundwater (GW) samples, application periods of the herbicides, and precipitation from April 2013-April 2014. 185 Precipitation is indicated as inverted bars on the secondary Y-axis. SINGLE-COLUMN COLOUR FIGURE 186 During fall and early spring, 2,4-D concentrations in river water were mostly one order of 187 magnitude higher than in the sampled groundwater. Generally, the 2,4-D concentrations in 188 surface remained relatively high in all samplings, and did not drop below 0.27  $\mu$ g/L (July 189 2013). A sharp decrease in concentration was however observed after the peak detected in 190 May  $(3.00 \,\mu g/L)$ . This drop was likely caused by the high dilution of the river water that 191 occurred just before the sampling in July (20% of the annual rainfall). Atrazine and 192 hydroxyatrazine also peaked in river water, but the concentrations in groundwater and river 193 water samples were generally more similar to each other than in the case of 2,4-D, meaning 194 that 2,4 removal along the flow path was slightly more rapid than for atrazine and its metabolites. This might indicate a higher degradation rate of 2,4-D. 195

196 Another interesting finding was that the atrazine degradation product hydroxyatrazine 197 exceeded that of the mother compound atrazine in nearly all samples (see Figure 1 for 198 comparison in µmol/L); even in groundwater. Other studies have found this dominance of 199 hydroxyatrazine only in surface water, where chemical hydrolysis dominates the atrazine degradation process (Ren and Jiang, 2002a), while close to no hydroxyatrazine is usually 200 201 detected in groundwater (Geng et al., 2013). Furthermore, the molar ratio of deethylatrazine 202 over atrazine, also called DAR, in the groundwater samples was much lower than 1.0 (only 203 0.24 on average), which indicates that no transport through the unsaturated zone has taken 204 place (Adams and Thurman, 1991). We therefore conclude that the much of the groundwater 205 contamination originated from the polluted river.

206 Compared to the distinct peak concentrations observed after the first application period, 207 pesticide concentrations after the summer application did not peak as considerably. This is 208 likely due to the higher dilution factor caused by the summer rainfalls. The fact that no 209 application of pesticides took place from about 1-2 km upstream of the study site (due to 210 flooding of the lower reach of Fu River) might also have reduced the signal slightly. 211 However, the fallow stretch upstream of the site only accounts for a very small proportion of 212 the expected source area for pesticide contamination along the river, so that it is unlikely that 213 this caused high differences in surface water concentration compared to a year with regular 214 farming activities. Higher vertical input of pesticides from transport through the unsaturated 215 zone under normal field conditions could have been possible, but as the lateral inflow of the 216 river water seemed to be the dominant influence at this site, this also would only have 217 changed the study results to a minor extend.

In summary, the high concentration of most of the detected pesticides in river waterdemonstrates that there is a high potential for groundwater contamination by polluted surface

water — despite the high dilution factor in rivers. The peak concentrations in the river water
are mostly observed after the pesticide application on sparse vegetation before the beginning
of the raining season, indicating pollution by spray drift during application (especially during
application on sparse vegetation in the spring time), surface runoff, and possibly also by point
sources, such as improperly disposed containers.

#### 225 **3.2. Agriculture and pesticide use in the NCP**

226 The NCP (extending over much of the provinces Beijing, Tianjin, Hebei, Henan, Shandong, 227 Jiangsu, and Anhui in the Eastern part of China) supports 14% of China's arable land and 11% of its population with only 2% of the nation's total groundwater resources (Frenken, 228 229 2012). The relatively fertile alluvial soils of the NCP and its mild winters with long sunshine 230 hours allow for double-cropping systems, such as the cultivation of winter wheat (October-231 June) and summer maize (June-October). This is the dominant agricultural activity in this area 232 (Li et al., 2015). However, the strong seasonality of rainfall—of which 70-80% occurs from 233 July to September—makes irrigation a necessity to sustain a good harvest of the winter wheat 234 (but is usually not necessary for the summer maize). Furthermore, the intense use of the soils 235 requires a major input of fertilizers to sustain high yields (Gao et al., 2014).

Due to the heavy promotion of agrochemicals by the government in the 1980s, and the increase in time constraints of farmers due to new job opportunities outside of the agricultural sector, a major shift from traditional hand weeding to the use of pesticides took place in recent decades. At present, about 800 pesticide products are approved for use in China, of which herbicides had an increasing share from 20% in 1960 to 56% in 2009 (Devi et al., 2009; Zhang et al., 2011). This relative increase in the share of applied herbicides from the total applied pesticides has previously been observed in the United States (US). Here, the percentage of applied herbicides rose from 18% to 76% between 1960 and 2008 (Fernandez-Cornejo et al., 2014).

245 In the provinces of the NCP, a particularly strong trend of increasing pesticide utilization can 246 be observed. According to data from representative regions in the NCP, 93.5% of the sown 247 area is treated with pesticides, which is well above the national average of 70.4% (Löw, 248 2003). In the specific case of winter wheat-summer maize fields, it is estimated that about 249 80% of the farmers in the NCP are using herbicides, and that only little weeding is still done 250 by hand (Menegat, 2013). Average application rates of pesticides for wheat-maize systems in 251 the NCP at the turn of the century were reported to be 2.82 kg a.i./ha/year (Li and Zhang, 1999), but more recent publications report current application rates of 5.87 kg a.i./ha/year 252 253 (Wang et al., 2014).

#### 254 **3.3.** Pesticide handling, application, occupational risk, and farmer's knowledge

A common method of pesticide application in China (and also in the NCP) is via spraying 255 256 from knapsack sprayers; often while wearing little protective clothing (Yang et al., 2014). 257 Consequently, a survey of 270 households in the NCP showed that 20% of the interviewed 258 farmers experience health impacts such as headache, dizziness, nausea, stomach ache, skin 259 rashes, fatigue, and increased visits to the doctor during times of pesticide application (Zhen et al., 2005). On the country level, 300-500 farmers die each year because of pesticide 260 261 poisoning (Devi et al., 2009). A follow-up study on registered intoxications with the herbicide paraquat found that-despite the widespread use of the chemical-its hazard to health is not 262 263 fully conceived by farmers, and many believe the compound to be less toxic than it is (Yin et 264 al., 2013). This stresses that the occupational risk from pesticides still needs to be clearly 265 understood.

266 Furthermore, it is reported that pesticides are applied in very high doses, and "over application is a common phenomenon in China" (Zhao, 2013). In a study on pesticide overuse 267 268 on cotton, about 87% of the surveyed farmers used double or more than the recommended 269 doses even though they claimed to have read the instructions in the manual (Jin et al. 2015). 270 Another example is that many farmers that grow genetically modified Bt (*Bacillus* 271 thuringiensis) cotton still add pesticides for pests, against which the Bt cotton is resistant 272 (Chen et al., 2013). This overuse of pesticides has been related to a variety of reasons. 273 Hamburger (2002) argues that mistrust into the exact composition and effectiveness of the 274 compounds, and the misconception that higher application doses will give more protection 275 against pest outbreaks are major causes for the observed over applications. Other studies 276 imply that untimely spraying (when pests and diseases had already broken out), and a strong 277 tendency to do as the neighbor does might be the cause (Liang et al., 2011). Overall, the over 278 application indicates a lack of farmer education. Indeed, a study of 64 farming household in 279 Hebei province discovered that 86% had never received any agricultural training (Kühl et al. 280 2009).

Another observed phenomenon is that emptied pesticide containers are often disposed of by simply leaving them in the field, or even in ditches and waterways. According to the China Crop Protection Industry Association (CCPIA, 2014), more than 5-10 pesticide packages per mu of farmland (1 mu =0.06ha) can be found on average. This may create an additional source of pesticide pollution in soil and water, and current studies propose that an enhanced farmer education as well a better infrastructure for proper disposal are needed to change the situation (Geng and Ongley, 2013; Jin et al. 2015).

#### 288 **3.4. Recent and current research focus on pesticides in the NCP**

A literature review was undertaken to assess the recent and current research and monitoring of

290 pesticides in the NCP. As of today, no public national database on systematic governmental

291 monitoring of pesticides in surface and/or groundwater could yet be found, and recent

292 publications point out the need for the establishment of a detailed governmental

environmental monitoring and risk assessment system (Han and Jin, 2016; Zhao and Pei,

294 2012).

In 1997, some data on sites contaminated with persistent organic pollutants (POPs), such as

296 DDT and HCH, was published in *The National Implementation Plan for the Stockholm* 

297 Convention on Persistent Organic Pollutants (The People's Republic of China, 2007).

However, all data from mainland China was derived from published scientific studies only.

299 The plan therefore highlights that a national monitoring system for persistent organic

300 pollutants (POPs) needs to be established. The Chinese Geological Survey (CGS) conducted a

301 groundwater quality survey from 2005-2015. Over 36,000 samples were taken and more than

302 70 components (including a wide range of banned and currently-used pesticides, listed in

303 CGS, 2008) were measured over an area about 4,400,000 km<sup>2</sup>. However, this data is not

304 publicly accessible. For routine measurements of groundwater quality, about 4000 monitoring

305 wells (operated by CGS and the provincial government monitoring institutes) are currently in

306 use, but pesticides are not considered for most of the wells. A more comprehensive national

307 groundwater monitoring network (about 20,000 professional monitoring wells) is constructed

308 at present, and over 80 components including pesticides will be measured (Hao Aibing,

309 personal communication December 13, 2016).

310 Results from our literature study on non-governmental pesticide research in the NCP (Table 2,

311 sorted by sampling year), show that ten out of seventeen studies focused on surface water

312 pollution by HCH and DDT. These compounds had been the most widely applied pesticides

in China between the 1950s and 1980s (Dai et al. 2011), but have been banned for most uses

- 314 (including agriculture) by the Chinese government since 1983. A strong research focus on
- HCH and DDT between 1970 and 1980 has previously been observed by Li and Zhang
- 316 (1999). Only six of our reviewed studies primarily concentrated on pesticides that are
- 317 currently in use. Out of these six studies, four are limited to the analysis for atrazine residues
- only, and all the six studies have been published before or in 2004 (Deuerlein et al., 2001a;
- 319 Domagalski et al., 2001; Ren and Jiang, 2002a; Ren and Jiang, 2002b; Ren and Jiang, 2004;
- 320 Ye et al., 2001).

321 **Table 2** List of recent studies (1995-2013) on pesticide pollution in surface water (SW) and groundwater (GW) in 322 the North China Plain. The studies are sorted by sampling year. Concentrations highlighted in bold exceeded the

the North China Plain. The studies are sorted by sampling year. Concentrations highlighted in bold exceeded the
 Chinese drinking water limit (according to guideline GB 5749-2006).

Pesticide SCPOP Study region Sampling time GW - sampling Study SW - sampling GW Author/vear depth Range (year/month) Mean Range Mean (m)  $(\mu g/L)$  $(\mu g/L)$  $(\mu g/L)$  $(\mu g/L)$ 1 DDT (I) 1995/06 0.100 Dou and Zhao, 1997 Baiyangdian х ----BHC (I) " 0.300 \_ х \_ 2 Various1 Tangshan 1996 ND ND Domagalski et al., 2001 no data -3 Atrazine (I) Baiyangdian 1998 1.33 0.40-3.29 15-50 Ye et al., 2001 4 Ren and Jiang, 2002b 1999-2000 ND-26.1 30-300 Atrazine (I) Yanghe -ND-0.69 -130 & 5 2000/12 ND-0.72 Atrazine (I) Yanghe 6.7 \_ 0.36 Ren and Jiang, 2002a 380 6 Various<sup>2</sup> Beijing 2001 < 0.1-5.1 ND ND 100 Deuerlein et al., 2001a 7 Atrazine (I) Guanting\* 2002/03 3.0 2.0 - 4.8Ren and Jiang, 2004  $0.005^{G}$ 8 Aldrin (I) Huai'an 2002/07 0.002-0.033 Wang et al., 2009 х DDT (I) -2003/03  $0.011^{\,G}$ 0.003-0.081 \_ х --" Dieldrin (I) х "  $0.001^{G}$ < 0.001-0.002 \_ -\_ " ...  $0.003^{G}$ " < 0.001-0.010 Endosulfan (I) х \_ -" " "  $0.002^{\,G}$ Endrin (I) 0.001 - 0.064\_ \_ \_ х " " "  $0.003^{G}$ HCB (F) < 0.001-0.001 х ---" " " HCH (I)  $0.003^{\,G}$ 0.001-0.009 --х " " " Heptachlor (I)  $0.008^{G}$ < 0.001-0.003 -\_ х " " "  $0.003^{G}$ 0.001-0.013 Methoxychlor (I) \_ 9 2003/09&11 0.001 Acetochlor (H) ND-0.002 -Guanting\* \_ \_ Xue and Xu. 2006 & " Alachlor (H) 0.002 ND-0.006 2004/05&08 " Aldrin (I) 0.023 0.002-0.032 --х " " " Chlordane (I) 0.010 0.001-0.039 \_ х " " " Chlorpyrifos (I) 0.002< 0.001-0.002 \_ \_ -" " " 0.001 ND-0.002 Cypermethrin (I) \_ \_ -" " " DDT (I) 0.093 0.006-0.364 \_ \_ \_ х " " " Deltamethrin (I) 0.004 ND-0.006 --" " " Dicofol (I) 0.001 ND-0.003 --" " " 0.001 ND-0.005 Dieldrin (I) х -" " " Endosulfans3 (I) 0.025 0.005-0.077 \_ -х " " " Endrins4 (I) 0.009 ND-0.0019 \_ -х " " " Fenvalerate (I) ND-0.003 0.002 " " " 0.012 HCB (F) 0.001-0.027 х \_ \_ \_ " ... " HCH (I) 0.071 0.007-0.051 \_ х " " " Heptachlor (I) х 0.005 0.002-0.021 ---" " " Methoxychlor (I) 0.004 ND-0.022 " " " Metolachlor (H) 0.023 0.016-0.027 \_ \_ \_ " " " Nitrofen (H) 0.001 ND-0.002 ---" Trifluralin (H) 0.005 0.003-0.005 ---10 DDT (I) 2007/03 0.004-0.034 Huaihe 0.011 --\_ Feng et al., 2011 х HCH (I) 0.004 ND-0.013 \_ \_ х DDT (I) 0.002 11 2007/07 -Hu et al., 2010a х Baiyangdian -HCH (I) х 0.002 --12 ND DDT (I) 2007/10 ND Wang et al., 2013 х Baiyangdian -HCH (I) " 0.002 0.001-0.008 " х 13 2008/07 0.011 0.004-0.021 \_ Dai et al., 2011 DDT (I) х Baiyangdian -\_ " 0.003-0.011 HCH (I) х 0.006 \_ -" 14 2009/06 0.096 0.035-0.157 Ge et al., 2010 DDT (I) х Weishan Lake -HCH (I) " 0.036 0.013-0.195 -" х 2010/03-12 15 0.024 0.006-0.054 Dai et al., 2013 HCH (I) х Baiyangdian \_ --2010/03-12 16 DDT (I) х Baiyangdian 0.008 0.002-0.034 -Dai et al., 2014 17 DDT (I) х Baiyangdian 2012/10 0.006 0.003-0.017 \_ \_ Guo and Feng, 2014 \_ HCH (I) " 0.005 0.003-0.012 х

x indicates that the compound is listed in the SCPOP (Stockholm Convention on Persistent Organic Pollutants); Guanting\* = Guanting Reservoir; BHC = Benzene hexachloride; DDTs = Dichlorodiphenyltrichloroethanes; HCB = Hexachlorobenzene; HCHs = Hexachlorocyclohexanes; H = Herbicide; I = Insecticide; F = Fungicide; G = Geometric mean; ND = Not detected; 1 = Complete list of analytes not known, but included the herbicides atrazine, cyanazine, simazine, alachlor, metolachlor, dicamba; 2 = Analysis for 31 pesticides. Detected in SW: Butachlor (H), metolachlor (H), molinate (H), chlorpyrifos (I), diazinon (I), and dichlorvos (I). No fungicides were detected in GW; 3 = Endosulfan I & II + Endosulfan sulfate; 4 = Endrin + Endrin aldehyde 331 From 2002 onward, only studies that focused primarily on HCH and DDT were found-332 despite the fact that detected concentrations were continuously by at least one order of 333 magnitude below the Chinese drinking water limit (1  $\mu$ g/L). One reason for the observed 334 sampling bias might be that DDT and HCH were listed as persistent organic pollutants (POPs) 335 by the Stockholm Convention on Persistent Organic Pollutants (SCPOP) in 2004, which could 336 have incentivized scientists to primarily publish studies on this topic. Additionally, there 337 seems to be a stronger focus on surface water than on groundwater sampling. In fact, the 338 authors could not find a published study on pesticides in groundwater in the NCP after 2002 -339 despite the fact that Ye et al. had detected atrazine concentrations in groundwater samples that 340 exceeded the national threshold for drinking water of  $2 \mu g/L$  in 2001. This indicates that 341 pesticide research in the NCP has not only narrowed down to studies related to the SCPOP, 342 but also that sampling in the easier accessible surface water is preferred over groundwater 343 sampling.

#### **344 3.5.** Applied pesticides in the NCP and their potential environmental impact

At present, statistics on pesticide use in China are neither easily obtainable nor greatly
detailed (World Bank, 2010). The authors therefore identified the most commonly applied
pesticides for wheat-maize cropping in the NCP by means of a literature search, in which ten
studies were considered that referenced to specific pesticides as being "the most typically
applied" in the region (Table 3).

- 350 351 Table 3 Most commonly applied pesticides in the North
- China Plain (NCP) for winter wheat and summer maize.
- 352 The presented pesticide list is based on the identification
- 353 of different herbicides, insecticides, and fungicides in recent

354 literature as most abundantly used pesticides in the

355 wheat-maize systems of the NCP.

Pesticide	Citation	Applied on	Emergence			
			Pre	Post		
Herbicides						
2,4-D	1, 2, 4, 6, 10	W / M		х		
Acetochlor	1, 3, 8,9	М	х			
Atrazine	1, 5, 7, 8,9	W / M	х	х		
Butachlor	1,8	W / M	х			
Chlortoluron	1	W / M		х		
Glyphosate	1	W / M		х		
Iodosulfuron	6	W		х		
MCPA	1	W / M		х		
Mesotrione	8	М	х	х		
Metolachlor	8	М	х			
Metribuzin	8	М	х	х		
Molinate	1	W / M	х	х		
Nicosulfuron	8	М		х		
Paraquat	1	W / M		х		
Tribenuron-methyl	6	W		х		
Trifluralin	1	W / M	х			
Topramezone	8	М		х		
Insecticides						
Acephate	7	W				
Carbofuran	3	М				
Chlorpyrifos	5	М				
Dichlorvos	3	W / M				
Dimethoate	2, 3, 4,10	W / M				
Emamectin benzoate	5	М				
Omethoate	4	М				
Fungicides						
Carbendazim	4	М				
Chlorothalonil	7	W				

- 356 357 W = Wheat; M = Maize; 1 = Deuerlein et al., 2001b; 2 = Hou et al., 2012; 3 = Huang et al., 2013; 4 = Li and Zhang, 1999; 5 = Liu et al., 2012;
  - 6 = Menegat et al., 2013; 7 = Wang et al., 2014; 8 = Ye et al., 2001;

358 359 9 = Zhang et al., 2013; 10= Huang et al., 2015

360	The results show that the different studies observed a relatively broad range of different
361	pesticides of being used in the same wheat-maize cropping systems in the NCP. Within the
362	group of herbicides, 2,4-D, acetochlor, and atrazine seem to be the most widely applied
363	compounds for weed control. This is similar to the five most used herbicides in the US:
364	glyphosate, atrazine, acetochlor, metolachlor, and 2,4-D (Fernandez-Cornejo et al., 2014).
365	Potential pesticide pollution of surface water and groundwater depends on the means and rate
366	of application environmental factors, the physicochemical properties of the pesticide, and its
	of approximation, en en of the properties of the posterior, and its
367	degradability in aquatic systems and soil. For example, high adsorption coefficients ( $K_{OC}$ ) of
367 368	degradability in aquatic systems and soil. For example, high adsorption coefficients ( $K_{OC}$ ) of insecticides like chlorpyrifos lead to accumulation in soil and river sediments (McKnight et

al., 2015; Rasmussen et al., 2015). Many herbicides on the other hand have a higher potential 369

for leaching and runoff – not only because of their different chemical properties, but also due to their early application when little vegetation is present (insecticides and fungicides are normally applied on foliage). This can lead to a higher impact of spray drift during, and surface runoff after the application. Accordingly, the majority of detected pesticides in largescale investigations are usually herbicides (e.g. Kolpin et al., 1998; Steele et al., 2002).

### 375 **3.6. Transport, fate, and expected impact of 2,4- D, acetochlor and atrazine**

Table 4 summarizes the main physicochemical properties of 2,4-D, acetochlor, and atrazine; their environmental behavior that derives from these properties; and their main degradation products. Acetochlor binds strongly to organic matter in the soil (Hiller et al, 2008), which makes it unlikely to leach into groundwater. Additionally, studies have shown that acetochlor is readily degradable in topsoil and in the subsurface (Janniche et al., 2010). Accordingly, it has only rarely been detected in surface water and groundwater (Barbash et al., 1999; Kolpin et al. 1996).

## **Table 4** Physicochemical properties of the three herbicides 2,4-D, acetochlor, and atrazine, derived behavior in aquatic and soil systems, and main degradation products.

· · · · · · · · · · · · · · · · · · ·	2.4-D	Acetochlor	Atrazine
	Z,4 D Cl	CH <sub>3</sub>	Cl
		COCH <sub>2</sub> Cl	
	CH <sub>2</sub> COOH—O—	CH <sub>2</sub> OCH <sub>2</sub> CH <sub>3</sub>	Ŋ_ ≥Ņ
		CH <sub>2</sub> CH <sub>3</sub>	CH_CH_NH N NHCH(CH_)
Physicochemical properties <sup>1</sup>			
Molecular formula	$C_9H_8Cl_2O_3$	$C1_4H_20CINO_2$	$C_8H_{14}ClN_5$
Molar weight (g/mol)	221.0	269.8	215.7
Solubility (mg/L)	23 180 (pH 7)	282	33
Vapor pressure (mPa) at 25°C	$1.9  { m E}^{-2}$	4.6 E <sup>-2</sup>	3.6 E <sup>-2</sup>
log K <sub>ow</sub> at 20°C	2.58-2.83	4.14	2.5
Ecosystem behavior and mobility <sup>2</sup>			
Potential for bioaccumulation	Low	Low	Low
Persistency in aquatic systems	High	High	Low
Persistency in soil systems	Low	Low	Medium
Soil mobility	Mobile	Moderately mobile	Moderately mobile
Leachability into groundwater	Low/moderately	Low	Possible
Main degradation products <sup>3</sup>	2,4 DCP Cl	ESA COCH <sub>2</sub> SO <sub>3</sub> H N COCH <sub>2</sub> CO <sub>4</sub> CH <sub>2</sub> CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> CH <sub>3</sub>	
	4-CP OH—O—	$\overset{OA}{\underset{CH_3}{\overset{CH_3}{\overset{OOOH}{\overset{OOCH_3}{\overset{OOOH}{\overset{OOCH_3}{\overset{OOC}}{\overset{OOCH_3}{\overset{OOCH_3}{\overset{OOCH_3}{\overset{OOCH_3}{\overset{OOCH_3}{\overset{OOCH_3}{\overset{OOCH_3}{\overset{OOCH_3}{\overset{OOC}}{\overset{OOCH_3}{\overset{OOC}}{\overset{OOCH_3}}{\overset{OOC}}{\overset{OOCH_3}{\overset{OOCH_3}}{\overset{OOCH_3}{\overset{OOCH_3}}{\overset{OOCH_3}}{\overset{OOCH_3}}{\overset{OOCH_3}{\overset{OOCH_3}{\overset{OOCH_3}}{\overset{OOCH_3}}{\overset{OOCH_3}{\overset{OOCH}$	

<sup>2,4-</sup>DCP = 2,4-Dichlorophenol; 4-CP = 4-Chlorophenol; ESA = Acetochlor ethanesulfonic acid; OA = Acetochlor oxanilic acid; HA =
Hydroxyatrazine; DEA = Deethylatrazine; DIA = Deisopropylatrazine; 1 =Tomlin, 2009; 2 = IUPAC, 2015; 3 = Modified from Reitzel et al.
2004 (2,4-D), Janniche et al., 2010 (Acetochlor), and Vonberg et al., 2014 (Atrazine).

2,4-D is expected to show higher mobility in soil than acetochlor (Hiller et al., 2008), but

389 generally has good degradability in soil, unsaturated zone and aquifer conditions (Johnson et

al. 1995; Tuxen et al., 2000; Willems et al. 1996). The persistence of 2,4-D in surface water is

391 relatively high due to its non-volatility and limited chemical and photochemical degradation

392 (Aly et al., 1964; Loos et al., 2009; Zepp et al., 1975). 2,4-D has been sparsely detected in

393 surface water and groundwater (Domagalski et al., 2001, Rawn et al. 1999).

- leaching (Barbash and Resek 1996; Graymore et al. 2001). This is in accordance with many
- 396 studies worldwide that detected atrazine and atrazine metabolites in surface water and
- 397 groundwater (Crocker et al., 2002; Domagalski et al., 2001; Maloschik et al., 2007; Sparling

<sup>394</sup> Atrazine has a moderate mobility, but higher persistence in soil, and is prone to groundwater

398 et al., 2010; Székács et al. 2015). The degradation process in the subsurface is slow, so that 399 atrazine residues can occasionally still be detected at the 0.1 µg/L threshold even 20 years 400 after the last application (Vonberg et al., 2014). In groundwater and in the vadose zone, the 401 degradation products hydroxyatrazine, deethylatrazine, and deisopropylatrazine usually occur 402 in concentrations of deethylatrazine > deisopropylatrazine > hydroxyatrazine (Bayless, 2001). 403 In surface water, hydroxyatrazine is usually detected in higher concentrations than 404 deethylatrazine and deisopropylatrazine, and often also in higher concentrations than the 405 parent compound atrazine (Lerch et al., 1998; Ren and Jiang, 2002a). Under loosing stream 406 conditions, the hydroxyatrazine concentrations in the water entering the aquifer is expected to 407 be quickly lowered because hydroxyatrazine has been shown to readily sorb to soil and 408 organic matter in several studies (Brouwer et al., 1990; Mandelbaum et al., 1993; Mersie and 409 Seybold, 1996).

410 The pesticides atrazine and 2,4-D are regulated in both the US and the Chinese Drinking 411 Water Guidelines. In case of long-term exposure above the maximum contamination level 412 (MCL), atrazine is expected to cause cardiovascular and reproductive problems, and 2,4-D 413 may cause damage to kidney, liver, adrenal glands (EPA, 2009). The two chlorinated 414 metabolites of atrazine (deethylatrazine and deisopropylatrazine) have similar properties as 415 atrazine, and are likewise suspected endocrine disruptors (Stanko et al., 2010; Stoker et al., 416 2002) with slight ecotoxicity. In terms of non-human impact, atrazine residues have been 417 recognized as a threat to consecutive crops in double cropping systems and wild plants due to 418 the phytotoxicity of the pesticide residues (Ren and Jiang, 2002b). This risk may be increased 419 if surface water or groundwater used for irrigation contains atrazine residues. However (as 420 discussed in 3.4.), only little monitoring of atrazine has recently taken place in the NCP.

#### 421 **3.7. Implications for pesticide handling, management and monitoring**

422 Combined findings from our literature review and our own observations at our field site imply 423 that little knowledge of active ingredients in pesticide products and improper handling of 424 pesticides is common in many farming households in China. Point sources from carelessly 425 disposed containers remain a problem, and can be a source for soil, surface water, and 426 groundwater pollution. Therefore, more educational activities more easily accessible disposal 427 sites for pesticide containers should be offered.

428 Our field study has further shown that some of the currently applied pesticides can be 429 detected throughout different times of the year in both surface water and groundwater. 430 Polluted surface water that infiltrates into groundwater can be a source of pollution, and it is advisable to treat surface water and groundwater as a joint resource by using appropriate 431 432 monitoring in both environmental compartments. Compared to the results from our literature 433 research, there seems to be a lacking scientific focus on currently used pesticides. We 434 therefore see a need for comprehensive monitoring program of surface water and 435 groundwater. The program should include both currently applied and banned pesticides, and 436 should be able to capture temporal variation.

437 Shallow groundwater wells and river water are often used for irrigation in the NCP. The main 438 irrigation period is typically from March to May, which coincides with our detected peak 439 concentrations of pesticides in river. Ren and Jiang (2002a) reported a decline in crop yield 440 (caused by phytotoxicity) when fields were irrigated with groundwater from 130m depth, 441 where deethylatrazine dominated as degradation product, and the total residue of atrazine and 442 chlorinated atrazine metabolites (atrazine+deethylatrazine+deisopropylatrazine) was about 9 443  $\mu$ g/L. In our field investigation, the total chlorinated residue was comparatively low (with a 444 maximum concentration of 1.1 µg/L in surface water in May 2013) because hydroxyatrazine was the main degradation product. Both surface water and shallow groundwater may 445

therefore still be suitable for irrigation (at least in terms of atrazine and atrazine residues), but monitoring of atrazine and 2,4-D is nevertheless advisable. Furthermore, the most crucial time for monitoring of surface water is after the spring application of herbicides. This is in accordance with the results on herbicide study in the Midwestern United States, which concluded that the spring flush of herbicides in surface water might be a major contributor to alluvial groundwater pollution (Thurman et al., 1991).

#### 452 **4. Conclusion**

453 Our findings show that the three most widely used herbicides on winter wheat-summer maize 454 fields in the NCP are 2,4-D, acetochlor, and atrazine. All of these compounds are suspected to 455 have potential negative impacts on the environment, and two of them (acetochlor and 456 atrazine) have been banned for this reason in Europe. In our field study, 2,4-D, atrazine, and 457 atrazine metabolites could be detected in river water and groundwater throughout the year, 458 with peak values of  $3.00 \,\mu\text{g/L}$ ,  $0.96 \,\mu\text{g/L}$ , and  $9.84 \,\mu\text{g/L}$ , respectively. Though our study has 459 limited repetition in time and space, we still see it as a relevant indication for the need to 460 monitor these compounds.

Indeed, we found that the monitoring gap on herbicides in groundwater systems is currently closing by the on-going development of a national groundwater monitoring network, which will also be used for the monitoring of pesticides (including compounds such as atrazine and 2,4-D). Nevertheless, there is a current research bias on surface water pollution by the legacy pesticides DDT and HCH, which were banned in China in the 1980s. Even though it is important to monitor if these compounds are still accumulating in the environment and/or are still illegally in use; we think that more research on currently-used pesticides should also be 468 encouraged. This research should ideally include both surface water and groundwater469 monitoring, and should be able to catch temporal variations.

In addition to this, overuse and improper handling of pesticides have been observed in China for a long time, and studies have been pointing out since the 1980s that better management needs to be implemented. Farmer education is still poor in many areas, and infrastructure for proper pesticide disposal is missing. Better farmer education, enhanced quality control, and a sufficient disposal system for pesticides are highly recommended to avoid occupational risks, pesticide overuse, and environmental pollution.

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### 756 Appendix A – Supplementary information

	Date	Crop a	ctivities	Irrigation	Pesticide appli	cation (g/ha of acti	ve ingredient)	
		Seeding	Harvest	(mm)	Type of pesticide	Application	Recommended application*	
Winter wheat Summer maize	10 Oct 2012 12 Apr 2013 20 Apr 2013 01 May 2013	+		40 50	Tribenuron-methyl (H)	18	9-18	
	30 May 2013 03 June 2013 10 June 2013 12 June 2013	+	+	60	Dimethoate (I)	75	50-100	
	07 June 2013				Paraquat (H)	1000	600-900	
	05 Oct 2013		+		Acetochlor (H)	2100	1800-3750	

**Table A.1** Crop activities, irrigation, and pesticide application during wheat and maize cultivation in 2012/2013.

\*recommendation as stated on the pesticide container

**Table A.2** Full set of analyzed compounds from the four field campaigns for river water (RW) and groundwater 761 (GW) samples. All detection limits were 0.01  $\mu$ g/L.

<b>Pesticide</b> (all values in $\mu$ g/L)	<b>de</b> (all values in µg/L) <b>29 May 2013</b>			22 July 2013			07 October 2013			25 March 2014						
	RW	GW2	GW3	GW4	RW	GW2	GW3	GW4	RW	GW2	GW3	GW4	RW	GW2	GW3	GW4
Triazine-herbicides																
Atrazine	0.96	0.33	0.91	0.40	0.32	0.29	0.29	0.49	0.49	0.19	0.20	0.20	0.54	0.12	0.19	0.15
- Desethylatrazine (M)	0.07	0.04	0.05	0.04	0.17	0.09	0.07	0.16	0.03	0.01	0.02	0.04	0.05	0.03	0.04	0.08
- Deisopropylatrazine (M)	0.07	0.05	0.05	0.04	0.04	0.16	0.08	0.09	ND	ND	ND	0.01	0.13	0.01	0.02	0.02
- Hydroxyatrazine (M)	9.70	0.46	0.37	0.47	0.31	0.59	0.54	0.49	3.00	0.55	0.39	0.48	1.60	0.26	0.33	0.20
Cyanazine	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Simazine	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Terbutylazine	ND	ND	0.02	0.01	ND	ND	ND	0.02	ND	ND	ND	ND	ND	ND	ND	ND
Phenoxy-herbicides																
2,4-D <sup>1</sup>	3.00	0.41	2.30	0.70	0.27	0.09	0.10	0.31	0.51	0.03	0.03	0.01	0.65	0.01	0.05	0.02
- 2,4-dichlorphenol (M)	0.09	0.14	0.09	0.14	0.02	0.10	0.09	0.07	0.06	0.03	0.04	0.03	0.01	ND	ND	ND
- 2,6-dichlorphenol (I)	ND	0.02	ND	0.01	0.01	0.03	0.16	0.02	0.01	ND	ND	ND	0.01	ND	ND	ND
MCPA <sup>2</sup>	0.06	0.03	0.05	0.02	0.03	ND	ND	0.01	ND	ND	ND	ND	0.04	ND	ND	ND
- 4-chlor-2-methylphenol (M)	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Dichlorprop	ND	ND	ND	ND	0.02	ND	ND	ND	0.04	ND	ND	ND	ND	ND	ND	ND
Mecoprop	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Other herbicides																
Bentazon	0.03	0.02	ND	0.02	0.01	ND	ND	ND	0.03	0.08	0.07	0.07	0.01	0.03	0.03	0.04
Dichlobenil	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
- BAM <sup>3</sup> (M)	0.06	0.01	ND	0.04	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Dinoseb	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
DNOC <sup>4</sup>	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Hexazinone	0.06	ND	ND	ND	0.04	ND	ND	ND	0.02	ND	ND	ND	0.11	ND	ND	ND
Isoproturon	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Metamitron	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Pendimethalin	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND

M = Metabolite; I = Intermediate; ND = Not detected (<0.01 µg/L); 1 = 2,4-Dichlorophenoxyacetic acid; 2 = 2-methyl-4-

chlorophenoxyacetic; 3= Dinitro-ortho-cresol; 4=2,6-dichloro-benzamide