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Residues of organochlorine pesticides (OCPs) in aquatic environment and risk assessment along Shaying River, China

Ying Bai · Xiaohong Ruan · J. P. van der Hoek

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Abstract Organochlorine pesticides (OCPs) are pesticides with global scale ubiquity, persistence and bioaccumulation, which leave long-term residuals in the water body. OCPs' high toxicity poses significant threats to human health and aquatic biodiversity, making assessment of OCPs' impact on aquatic ecology and human health urgently necessary. In this research, the presence of 16 OCPs in surface water and groundwater along Shaying River, China, as well as OCPs concentration correlations, was investigated at 24 selected sampling sites. At the same time, the ecological risk and human carcinogenic risk were also analyzed by risk quotient method and USEPA's Risk Assessment Guidance, respectively. Results showed that the total concentration of OCPs ranged from 21.0

to 61.4 ng L⁻¹ in groundwater, and 12.3–77.5 ng L⁻¹ in surface water. Hexachlorocyclohexane (HCHs) and heptachlor were the prominent contaminants in groundwater, which indicated their use in the recent past and confirmed their persistence. The α -HCH/ γ -HCH ratios in groundwater confirmed that γ -HCH (lindane) was used as main substitute of technical HCH in the study area. The correlation analysis illustrated that δ -HCH and γ -HCH played a dominant role in HCHs residue. Heptachlor and α -HCH, as well as endosulfan and heptachlor epoxide, had a strongly significant positive correlation, suggesting an associated usage of the two pair OCPs. An extremely high ecological risk for aquatic organism was observed for γ -HCH, heptachlor and dieldrin, while the carcinogenic risks posed by the selected OCPs in surface water and groundwater were all acceptable.

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Keywords Organochlorine pesticides · Surface water and groundwater · Risk assessment

Abbreviations

OCPs	Organochlorine pesticides
HCHs	Hexachlorocyclohexane
DDT	Dichlorodiphenyl trichloroethane
RSD	Relative standard deviation
IDLs	Instrument detection limits
MDLs	Method detection limits
RQ	Risk quotient
C	Measured concentration of OCPs

PNEC	Predicted no-effect concentrations for a particular OCP
AF	Assessment factor
RAGS	Risk Assessment Guidance for Superfund
TLCR	Total carcinogenic risk
PCA	Principal component analysis
USEPA	United States Environment Protection Agency

Introduction

Persistent organic pollutants (POPs) are ubiquitous, toxic, anthropogenic substances that have been widely used in agriculture and manufacturing industries since the 1930s (Buah-Kwofie and Humphries 2017). Organochlorine pesticides (OCPs), as one of the widely used POPs with persistence and bioaccumulation, left long-term residuals in the water body at global scale (Jayaraj et al. 2016). Five OCPs were firstly recorded in Stockholm Convention, saying dichlorodiphenyltrichloroethane (DDT), aldrin, dieldrin, toxaphene and HCH, which were also banded by Chinese government in 1983. Technical HCHs, as the most extensively concerned OCPs compounds, were produced and used in China from 1952 to 1983. Some other OCPs' production and usage begins from the 1990s, like lindane and endosulfan, which may redistribute and change the composition of OCPs residuals in the environment. OCPs may reach the surface riverine water via agricultural runoff, direct applications, spray drift, aerial spraying and erosion (Chakraborty et al. 2016). The chemical properties of OCPs, such as low water solubility, stability to photo-oxidation and low vapor pressure, are the main reasons for their persistence in the environment.

Shaying River, the largest tributary of the Huaihe River, contributed large amounts of, was the main source of industry pollutants along the Huaihe River Basin, China. At the same time, rural area and farmland took over 80% of total basin area, on which pesticide applications were rising under increasing demand for grain production. From 1990 to 2008, the average pesticide application on farm land of Shaying River Basin increased from 7.43 to 22.63 kg km⁻², although the efficient proportion on crop disease prevention is barely above 30% (Guzzella et al. 2006;

Kolpin et al. 1998; Papastergiou and Papadopolou-Mourkidou 2001). Investigation shows that most of the drinking and sanitary water were taken from shallow aquifer (over 90% well with depth above 15 m) in the rural area of Shaying River Basin (<https://www.boxun.com>), which was under severe safety risk from agriculture organic chemicals (Lacorte and Barcelo 1996). The problem of high cancer incidence in some parts of Shaying River Basin is very prominent, with up to 300/100,000 in some villages, which are mostly located along the Shaying River (Chen et al. 2016). A certain relationship between high incidence rate of cancer and water environment pollution in this area was reported (Gonghuan and Zhong 2013).

The available researches paid more attention to OCPs residues in soil (Chakraborty et al. 2015; Yu et al. 2014), surface water and the sediment (Agarwal et al. 1986; Turgut 2003). For example, it was reported that the levels of OCPs range from 1.01 to 791.0 ng L⁻¹ in the surface water of the Yangtze River (Tang et al. 2008), Guanting Reservoir, China (Xue et al. 2006), and Gomti River, India (Malik et al. 2008). The OCPs concentration in sediment was reported as 0.4–813.6 ng g⁻¹ in Guanting Reservoir (Xue et al. 2006), Huangpu River (Hu et al. 2005), Huanghe River, China (Sun et al. 2007), and Gomti River, India (Malik et al. 2008). Riverine runoff was reported as the major mode carrying OCPs from the River Basin (Guan et al. 2009). While the OCPs residues in groundwater, as well as the correlation between OCPs concentration in surface water and groundwater were neglected. OCPs' high toxicity poses significant threats to human health and aquatic biodiversity (Qu et al. 2015), while there is no information on the ecological risk associated with the presence of pesticides including OCPs in various environmental compartments in Shaying River Basin, neither are there data available on the human carcinogenic risk associated with the consumption of pesticide-contaminated water in this area. Therefore, assessment of OCPs' impact on aquatic ecology and human health in this area is urgently necessary.

The objective of this paper was to study the occurrence, sources, possible migration evidence and risk of OCPs in the aquatic environment of Shaying River Basin. The residue concentrations and distribution pattern of 16 OCPs were firstly investigated in this area, and the correlations of individual OCPs

component in groundwater and surface water were analyzed. Besides, ecological and carcinogenic risks posted by OCPs were assessed based on a modified method, in order to depict the impact of these pesticide residues on aquatic ecology and human health in this area.

Materials and methods

Site description and sampling

Shaying River Basin is one of the largest tributary catchments of Huaihe River Basin, plain area of which is about 25,600 km². Climate of this area is continental with an annual average temperature of 14–16 °C and annual average rainfall of 769.5 mm mostly distributed in high flow period (June to August). The main soil type of Shaying River Basin is cinnamon soil and lime concretion black soil, and main land use type is farmland, accounting for 87% of the total area.

Sampling sites along Shaying River were selected equably to cover the research area. Fourteen surface water samples and 10 groundwater samples (0.5 m below water surface) were collected with pre-cleaned 1-L dark glass bottles using cylinder samplers in November 2013 (Fig. 1). All samples were placed in icebox after sampling and transported to the laboratory immediately for later analysis.

Used chemicals

The standard OCPs including α -HCH, β -HCH, γ -HCH, δ -HCH, *p,p'*-DDT, *o,p'*-DDT, *p,p'*-DDE, *p,p'*-DDD, aldrin, heptachlor, endosulfan I, endosulfan II, methoxychlor, dieldrin, endrin, heptachlor epoxide and internal standards (pentachloronitrobenzene) were purchased from Accustandard (USA). The solvent (petroleum ether) and concentrated sulfuric acid used for sample processing and analysis were analytical grade, and the former was re-distilled in a full-glass distilling appliance. Anhydrous sodium sulfate (analytical grade) was baked in a furnace oven at 500 °C for 8 h and kept in sealed desiccators prior to use.

Sample pretreatment

The pretreatment experimental process referred to gas chromatography determination method of HCHs and

DDTs in Chinese water quality standard (China 1987). The sample pretreatment process was as follows. 0.6 L of water samples was taken for oscillation in 80 mL of petroleum ether for 10 min under liquid–liquid extraction method. Then, organic phase samples were purified with 10 mL liquid sulfuric acid and with 2% sodium sulfate solution acid washing. After liquid anhydrous, the organic phase was dehydrated with sodium sulfate (drying under 500 °C for 8 h) and then concentrated to 1 mL for later gas chromatography test.

OCPs quantification and quality control

An Agilent 7890 GC equipped with a micro-cell electron capture detector (μ ECD) was used for OCPs quantification. The capillary column used was HP-5MS (30 cm $\times$ 0.32 mm $\times$ 0.25 μ m). The carrier gases were nitrogen for μ ECD at a flow of 2.0 mL min⁻¹ under the constant flow mode. The inlet was heated to 250 °C, and the nitrogen purge time was 0.75 min. The GC column temperature was programmed as follows: initially at 80 °C (equilibrium time 2 min), increased to 180 °C at the rate of 15 °C min⁻¹ and held for 2 min, before reaching at 280 °C at the rate of 5 °C min⁻¹ and then held for 2 min. The temperature of μ ECD was held at 300 °C.

The linear correlation coefficient of standard calibration curves for target OCPs was all above 0.999, with a relatively high repeatability. The relative standard deviation (RSD) was between 3 and 20%, among which larger ones only appear in low-concentration samples (Guan et al. 2009).

All analytical operations were conducted under strict quality control guidelines. Procedural blanks and spiked samples with standard were used to monitor procedural performance and matrix effects. All experiments were carried out in duplicate. The instrument detection limits (IDLs) of all OCPs ranged between 0.062 and 0.858 ng mL⁻¹, and the method detection limits (MDLs) ranged from 0.10 to 1.43 ng L⁻¹. The surrogate recovery in this method was 89–97%. Values for OCPs lower than the method detection limits (< MDL) were substituted with zero prior to statistical analysis.

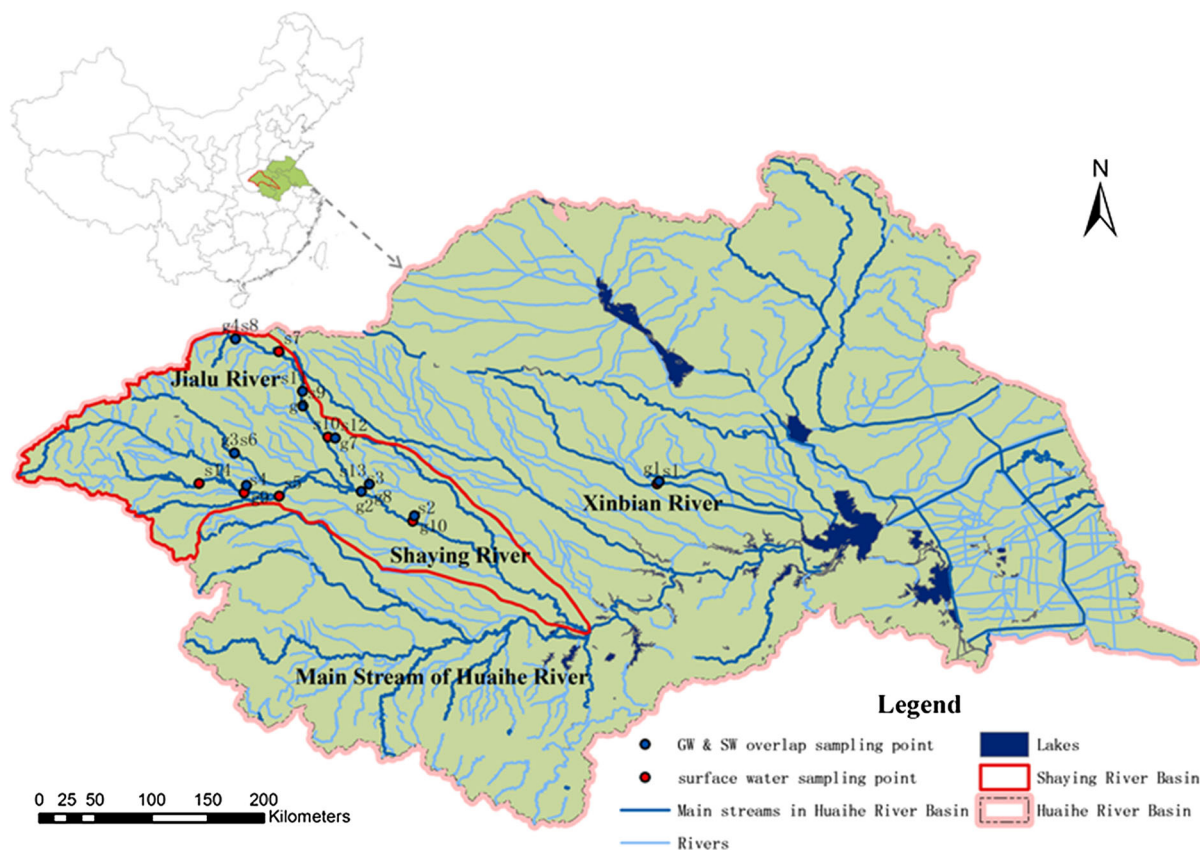


Fig. 1 Research area and sampling sites

Statistical analysis

The Pearson correlation test ($p < 0.01$) of different OCP components in surface water and groundwater of Shaying River Basin, and the principal component analysis (PCA) of OCPs in surface water were conducted using the software SPSS version 21. ArcGIS version 10.2 was used for spatial distribution analysis and graph making.

Ecological risk assessment

The ecological risk of OCPs to aquatic organisms was assessed using the risk quotient (RQ) method (WHO 2001), which was performed by calculation of RQ for the detected OCPs.

$$RQ = \frac{C}{PNEC}, \quad (1)$$

where C is the measured concentration and PNEC is the predicted no-effect concentrations for a particular OCP, respectively. The PNEC data were obtained from ECOTOX database (<http://www.epa.gov/ecotox>) and some articles (Hu et al. 2015; Ogbeide et al. 2015; Vryzas et al. 2011), where the value was calculated by multiplying the laboratory toxicology data with an assessment factor (AF) which takes into account the uncertainty between laboratory toxicity tests with real environment. The classification of ecological risks was different according to various assessment standards and management purposes. Generally, RQ above 1 represents that the environmental concentration is higher than predicted no-effect concentrations. Subsequently, the ecological risks were classified into three levels based on RQ value in the prior literature: 0.01–0.1 represents relatively low risk, 0.1–1 represents a medium risk, and above 1 indicates high risk (Peng et al. 2014).

Carcinogenic risk assessment

The United States Environment Protection Agency (USEPA) recommended that toxicity data and exposure data are combined to assess chemical's carcinogenic risk (USEPA 1989). In the present study, human carcinogenic risks associated with OCPs exposure were based on the approach from USEPA's Risk Assessment Guidance for Superfund (RAGS) methodology (USEPA 1997). The potential exposure to OCPs was calculated by two exposure routes, ingestion and dermal contact, which represent water exposure for drinking and bathing, as described in the USEPA screening-level equations for preliminary remediation goals (USEPA 2010). For different exposure pathways, the increasing trend in cancer risk for OCPs is determined as follows: inhalation < dermal contact < ingestion, thus comparably, inhalation of OCP particulates is almost negligible (Qu et al. 2015) and is not taken in this research. Total carcinogenic risk (TLCR) was obtained by summing individual risks calculated for the two exposure routes.

$$\text{TLCR} = \text{CR}_{\text{ingestion}} + \text{CR}_{\text{dermal}}, \quad (2)$$

$$\text{CR}_{\text{ingestion}} = \frac{C \times \text{CF} \times \text{IR} \times \text{EF} \times \text{ED}}{\text{BW} \times \text{AT}}, \quad (3)$$

$$\text{CR}_{\text{dermal}} = \frac{\sqrt{6\tau \cdot \text{TE}/\pi} \cdot C \cdot \text{CF} \cdot k \cdot A_{\text{sb}} \cdot \text{EF} \cdot \text{FE} \cdot \text{ED}}{500 \cdot \text{BW} \cdot \text{AT} \cdot f}. \quad (4)$$

In these equations, C is the detected concentrations of OCPs in the water (mg L^{-1}); IR is the average daily ingestion of water (2 L day^{-1}); CF is the conversion coefficient ($1 \times 10^{-6} \text{ kg mg}^{-1}$); EF is the exposure frequency ($365 \text{ day year}^{-1}$); ED is the exposure duration (30 year); BW is the body weight (60 kg for Chinese people); AT is the average lifespan ($70 \text{ years} \times 365 \text{ days year}^{-1}$); TE is the contact time (0.4 h); τ is lag time for each OCP in the body (1 h); k is a skin permeability parameter (0.001 cm h^{-1}); A_{sb} is body surface area ($16,600 \text{ cm}^2$); FE is bathing frequency (0.3 times day^{-1}); and f is the intestinal absorption ratio (1) (Schriks et al. 2010, Wei et al. 2015, WHO 2001).

In some studies, the carcinogenic risk was considered non-ignorable if the TLCR is above 1.0×10^{-6} , and higher than 1×10^{-5} indicated a 'not negligible' risk (Butler-Jones 2010, Li et al. 2015a). USEPA

(USEPA 1989) takes the range from 1.0×10^{-4} to 1.0×10^{-7} as acceptable carcinogenic risk. Qu et al. evaluated TLCR in five levels: Values lower than 10^{-6} represent very low risk; values between 10^{-6} and 10^{-4} represent low risk; values between 10^{-4} and 10^{-3} indicate moderate risk; values between 10^{-3} and 10^{-1} indicate high risk; and values higher than 10^{-1} represent very high risk (Qu et al. 2015). Generally, by integrating the above risk-rating standards, we took 1×10^{-6} as threshold value of potential carcinogenic risks for humans in this research.

Results and discussion

Residues of 16 OCPs were detected in surface water and groundwater along Jialu River and Shaying River (Table 1). The concentration of $\sum \text{OCPs}$ ranged from 21.0 to 61.4 ng L^{-1} in groundwater, and from 12.3 to 77.5 ng L^{-1} in surface water, with the mean concentration of 44.6 and 30.6 ng L^{-1} , respectively. The concentration of $\sum \text{HCHs}$ ranged from 8.6 to 26.7 ng L^{-1} in groundwater, and from 2.5 to 23.9 ng L^{-1} in surface water, with the mean concentration of 18.7 and 14.3 ng L^{-1} , respectively. Levels of HCHs and DDTs in surface water of upper reach of Huaihe River (Feng et al. 2011), Lake Baiyangdian (Guo and Feng 2014) and Yellow River Estuary (Li et al. 2015b) were 0.9–62.2 and 1.5–35.5 ng L^{-1} , respectively, while in River Yamuna, India, the levels were with higher concentration of 12.8–593.5 and 66.2–723.0 ng L^{-1} (Kaushik et al. 2008). Concentration of total OCPs ranges from 1.0 to 46.5 ng L^{-1} in Yangtze River (Tang et al. 2008), Pearl River Delta (Guan et al. 2009), Chao River (Yu et al. 2014), and 0.8–64.5 ng L^{-1} in underground river of southwest China (Hu et al. 2011). OCPs concentration in surface water was with higher concentration of 2–1200 ng L^{-1} in other Asia countries such as River Brahmaputra and River Hooghly, India (Chakraborty et al. 2016), and River Chenab, Pakistan (Eqani et al. 2012). Individually, dieldrin, heptachlor, heptachlor epoxide were with high detection frequency in river water of Korea, whose concentration range was 3–5410 ng L^{-1} (Cho et al. 2014). Generally, the concentrations of OCPs were low in Shaying River Basin compared with other areas in China and Asia countries.

Table 1 Level of OCPs in surface water and groundwater in Shaying River Basin (ng L⁻¹)

Compounds	Surface water (ng L ⁻¹)		Groundwater (ng L ⁻¹)	
	Conc.	Freq. (%)	Conc.	Freq. (%)
α -HCH	2.0 (0.7,3.3)	85.7	3.4 (0.8,10.4)	100.0
β -HCH	3.1 (1.0,7.5)	92.9	4.3 (1.3,10.5)	60.0
γ -HCH	5.2 (1.8,9.3)	100.0	6.4 (2.8,10.1)	90.0
δ -HCH	6.2 (1.8,12.5)	85.7	7.0 (2.7,14.2)	100.0
$\sum$ HCHs	14.3 (2.5,23.9)	100.0	18.7 (8.6,26.7)	100.0
Heptachlor	8.3 (1.3,17.9)	92.9	11.2 (1.1,24.4)	100.0
Aldrin	2.3 (1.0,3.8)	92.9	2.2 (1.0,5.2)	50.0
Heptachlor epoxide	1.4 (0.6,1.9)	28.6	1.6 (1.4,1.8)	20.0
Endosulfan I	8.5 (2.1,40.7)	57.1	5.5 (5.5,5.5)	10.0
Dieldrin	4.6 (4.6,4.6)	7.1	–	0.0
Endrin	6.2 (1.0,11.3)	14.3	–	0.0
$\sum$ OCPs	30.6 (12.3,77.5)	100.0	44.6 (21.0,61.4)	100.0

p,'-DDT, *o*,'-DDT, *p*,'-DDE, *p*,'-DDD, endosulfan II, methoxychlor were also analyzed although the results were below the detection limits

Occurrence of OCPs in Shaying River Basin

The results showed that the detection frequency of HCHs was the highest, indicating it as the prominent contaminants group of OCPs in Shaying River (Figs. 2a, 3a). Comparing with the centralized drinking water standards of China (China 2006), no residues were above the limits (0.005 mg L⁻¹ for HCHs, 0.002 mg L⁻¹ for lindane and 0.001 mg L⁻¹ for DDTs). The concentration level of HCHs in Shaying River was higher than Sao Paulo State Rivers in Brazil (Rissato et al. 2006), surface water in Jiangsu China (Li et al. 2012) and Bosumtwi Lake in Ghana (Darko and Acquah 2008), similar to Paranoa Lake in Brazil (Caldas et al. 1999), Yanzi River in China (Tang et al. 2008), but lower than most of other surface water (Kaushik et al. 2008; Singh et al. 2005; Turgut 2003; Wang et al. 2007; Zhang et al. 2003; Zhou et al. 2008).

All of the samples were detected with HCHs and heptachlor, indicating them as the prominent OCP contaminants in groundwater of Shaying River (Figs. 2b, 3b). No exceeding of concentration was observed based on centralized drinking water standards of China.

Principal component analysis (PCA) was performed to further determine the possible sources and composition in surface water of Shaying River Basin on four most frequently detected OCPs (Table 2-1). The contribution rate of each OCP to components can be taken as the main accordance of determination of main contaminant (Table 2-2).

The contribution rate of first main component F1 was 47.513%, which was highly associated with endosulfan I and heptachlor. Endosulfan I was firstly used in China from 1994, and the total consumption reached 25.7 thousand tons. On the 2011 'Stockholm Convention' of United Nations fifth conference of the parties, 127 countries, including China, finally agreed to add the highly toxic pesticide endosulfan to the prohibited substance of the Convention, but because of the historical large usage and difficulty of degradation, endosulfan became the main component of organochlorine pesticide contaminant in the water of the research area. The contribution rate of the second main component F2 was 33.047%, with high positive load of HCHs and heptachlor, indicating a similar pollution source. HCHs and heptachlor have been prohibited in the 1980s, but are still highly detected in water body, which suggested that the historical inputs in surface water cannot be easily deciphered.

Chemical fingerprints including conventional isomer ratios and enantiomeric compositions were usually used for indicating the varied sources of OCPs in the environment. During the 1970s and early 1980s, China was the largest producer and consumer of technical HCH (60–70% α -HCH, 5–12% β -HCH, 10–12% γ -HCH and 6–10% δ -HCH) (Willett et al. 1998) in the world with an increasing production from zero (1952) to nearly 300 thousand tons per year (Li et al. 1998) because of the intense food production pressure, and was officially banned in China in the middle 1980s. As a substitute, lindane (> 99% γ -

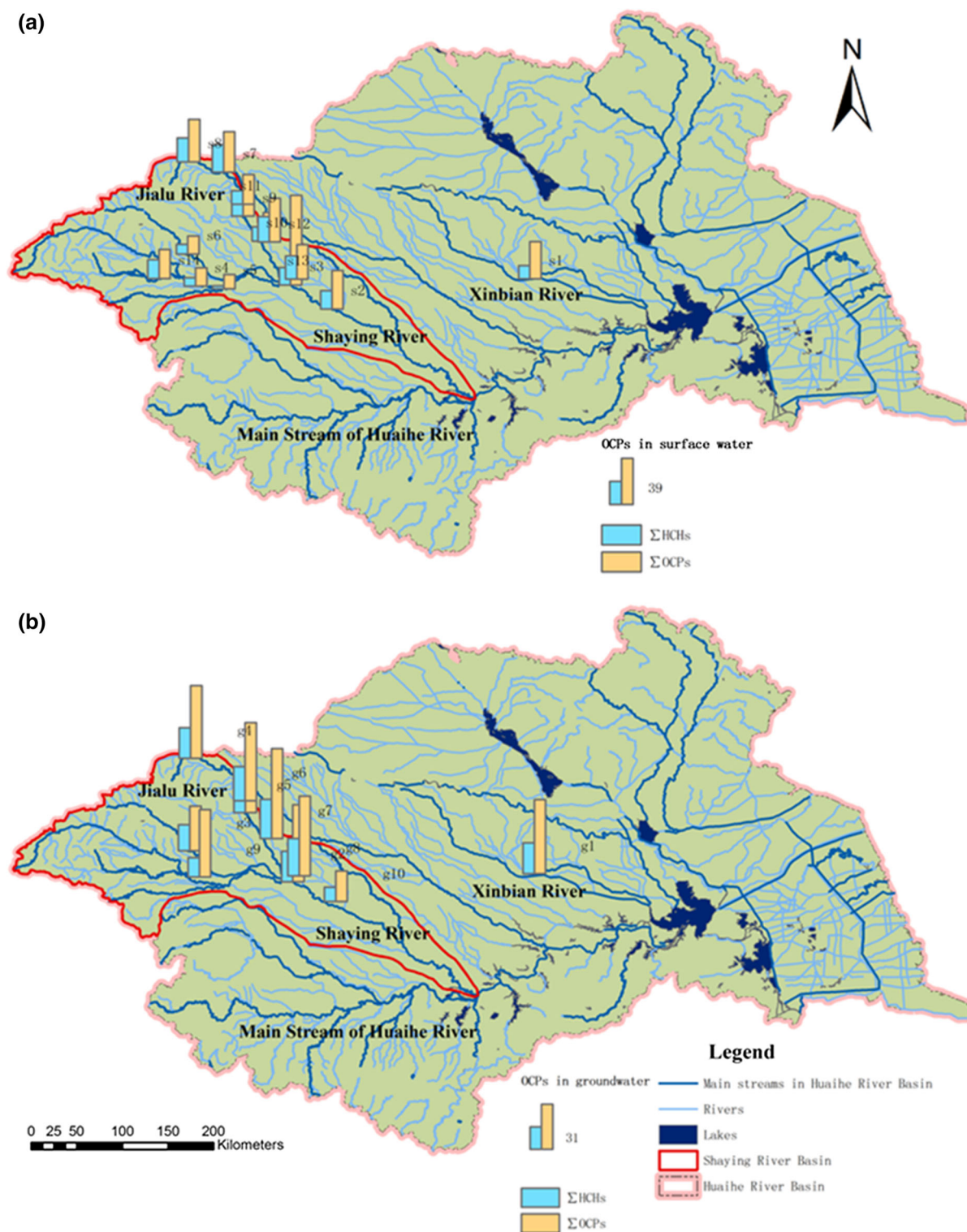


Fig. 2 OCPs residue spatial distribution in surface water (a) and groundwater (b) of Shaying River (ng L^{-1})

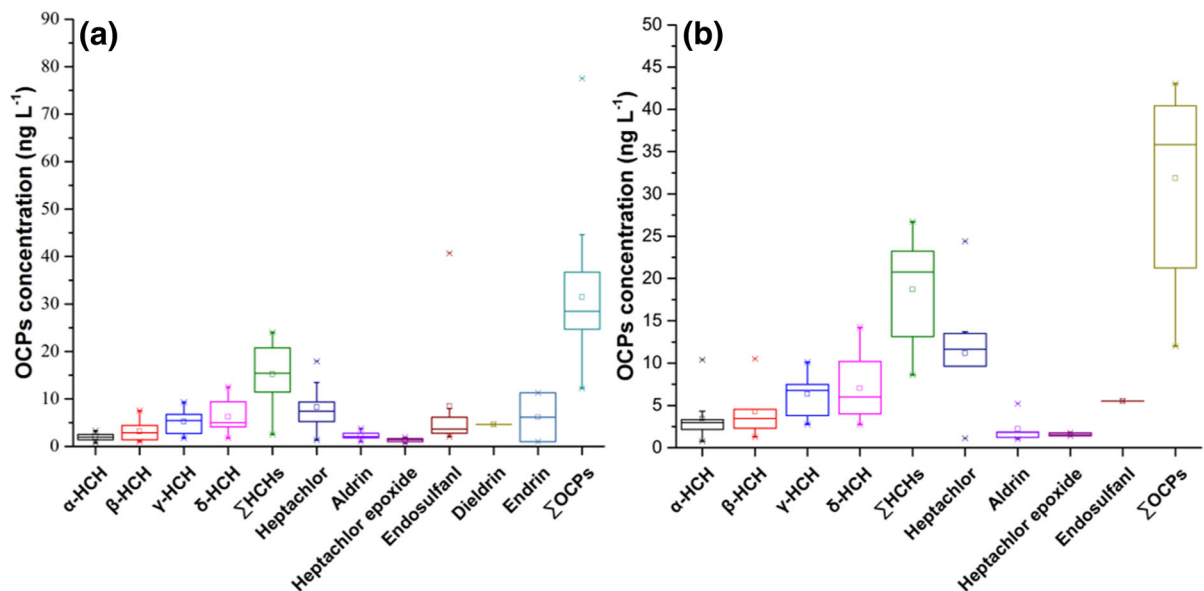


Fig. 3 OCPs concentration in surface water (**a**, $n = 14$) and groundwater (**b**, $n = 10$) of Shaying River

Table 2 (1) PCA results of surface water samples, (2) component matrix of PCA

Principle component	Eigen value and contribution rate		
	Eigen value	Contribution rate %	Cumulative contribution rate %
(1)			
1	1.901	47.513	47.513
2	1.322	33.047	80.561
3	0.516	12.908	93.469
4	0.261	6.531	100.000
Principal component	Factor		
	F1	F2	
(2)			
ΣHCHs	0.237	0.859	
Heptachlor	0.695	0.538	
Aldrin	− 0.748	0.490	
Endosulfan I	0.895	− 0.236	

The principal factor expression:

$$F1 = 0.237 * \sum HCHs + 0.695 * \text{heptachlor} - 0.748 * \text{aldrin} + 0.895 * \text{endosulfan I}$$

$$F2 = 0.859 * \sum HCHs + 0.538 * \text{heptachlor} + 0.490 * \text{aldrin} - 0.236 * \text{endosulfan I}$$

HCH) has been used in agriculture since 1990 (Li et al. 2001). Therefore, the α -HCH/ γ -HCH (α/γ) ratio is in the range of 3–7 for technical HCH, whereas low α/γ ratio depicts the usage of lindane (Li et al. 1998; Vu Duc et al. 2009). In this study, α/γ ratios of

groundwater samples along Jialu River ranged from 0.24 to 0.81 with an average of 0.46, which confirmed the use of lindane as the major source of HCHs contamination in the study area (Figs. 4, 5). Compared with other studies, the HCHs concentration is similar

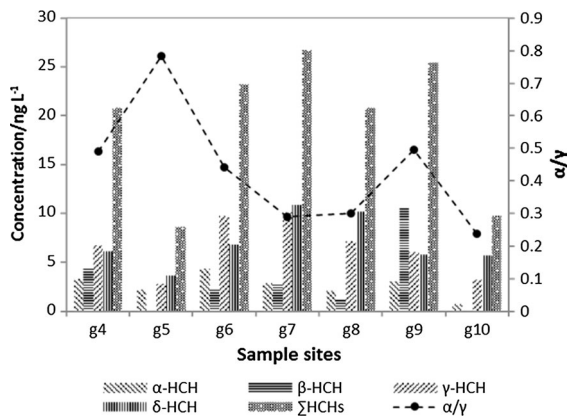


Fig. 4 OCPs concentration distribution in groundwater along Shaying River

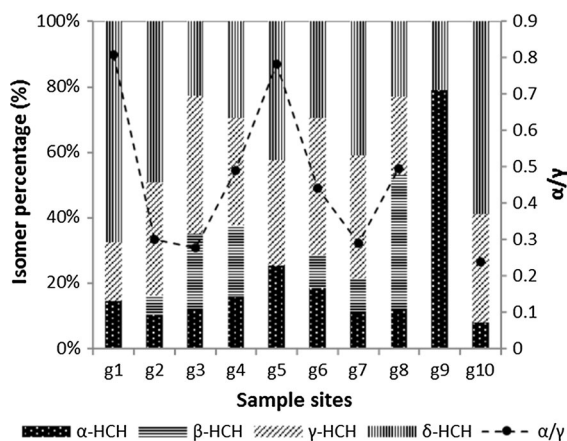


Fig. 5 Compositions of HCH isomers in groundwater of research area

with the groundwater in karst area, Chongqing, China (Hu et al. 2011), and the γ -HCH, δ -HCH also took the major proportion of HCHs (Fig. 4). However, it was opposite to that of Qiantang River, Ganges River and Pearl River estuary (Yu et al. 2008, 2009; Zhou et al. 2006). The results may also due to the deduced γ -HCH (lindane) input in Shaying River Basin and its possible transformation to δ -HCH under anaerobic condition in groundwater aquifer (Li et al. 2006). Another possible explanation was that α -HCH has a higher vapor pressure and Henry's law constant, which is approximately twice as that of γ -HCH (Walker et al. 1999) and 20 times higher than β -HCH (Li et al. 2004), indicating that α -HCH is more likely to partition to the air than to the water body (Li et al. 2014).

Correlation analysis of OCP components

Correlation analysis of the OCPs concentration in both surface water and groundwater was conducted to find out the relationship between the individual OCPs component in the research area and their possible migration evidence in the aquatic environment (Table 3). It can be seen that δ -HCH and γ -HCH showed a strongly significant positive correlation with Σ HCHs ($P < 0.05$), which illustrated their high residual in the water environment and their similarity of residual pattern with Σ HCHs. In addition, heptachlor and α -HCH, as well as endosulfan and heptachlor epoxide, have a strongly significant positive correlation, which may be an indication for the associated usage of these two pairs of OCPs.

Risk assessment of OCPs in Shaying River Basin

Ecological risk assessment

The results of ecological risk assessment of OCPs in surface water and groundwater in Shaying River Basin are shown in Table 4. RQ values between 0.01 and 0.1 were observed for α -HCH, β -HCH, δ -HCH, heptachlor epoxide, aldrin and endrin, suggesting that these OCP individuals post little ecological risk to selected aquatic organism in the research area. RQ value slightly above 0.1 was observed for β -HCH, while RQ values of dieldrin and endosulfan I were between 0.1 and 1.0 only in surface water, which indicated a generally moderate and potential ecological risk. Particularly, RQ value exceeding 1.0 was observed for γ -HCH (lindane) and heptachlor, revealing its extremely high ecological risk on aquatic organism in the research area which needs more concern. The average RQ value of γ -HCH (lindane) and heptachlor in surface water was 5.24 and 9.59, respectively, and that in groundwater was 5.72 and 13.97, respectively, resulting from the high detected concentration and their high toxicity on aquatic organism. Generally, OCPs post higher ecological risks to groundwater than surface water in the research area. Even though the selected target aquatic species may not appear in groundwater environment, but considering the possible groundwater discharging to surface water, it may provide reasonable reference to the ecological assessment and risk management practice.

Table 3 Correlations coefficient of OCPs in surface water and groundwater

Correlation coefficient	β -HCH	γ -HCH	δ -HCH	$\sum$ HCHs	Heptachlor	Aldrin	Heptachlor epoxide	Endosulfan I
α -HCH	0.143	0.529*	− 0.126	0.186	0.773**	− 0.083	0.4	− 0.069
β -HCH		0.207	− 0.245	0.353	− 0.074	0.503	− 0.575	− 0.568
γ -HCH			0.215	0.777**	0.212	0.084	− 0.096	− 0.425
δ -HCH				0.638**	0.224	− 0.157	0.167	0.45
$\sum$ HCHs					0.365	0.161	0.049	0.012
Heptachlor						− 0.053	0.581	0.106
Aldrin							− 0.521	− 0.633
Heptachlor epoxide								1**

*Significant correlation, $r > 0.5$, $P < 0.05$ **Strongly significant correlation, $r > 0.6$, $P < 0.01$ **Table 4** Parameters of RQs estimation

OCPs	Species	End point ($\mu\text{g L}^{-1}$)	AF	PNEC (ng L^{-1})	Average surface water RQs	Average groundwater RQs
α -HCH	Insects ^a	2d LC ₅₀ (150) ^a	1000 ^{ac}	150 ^a	0.01 (< 0.01–0.02) ^g	0.02 (0.01–0.07)
β -HCH	Fish ^a	84d NOEC(32) ^a	1000 ^a	32 ^a	0.09 (< 0.01–0.24)	0.08 (< 0.01–0.33)
γ -HCH	Crustaceans ^a	3d NOEC(0.1) ^a	100 ^a	1 ^a	5.24 (1.79–9.33)	5.72 (2.78–10.10)
δ -HCH	Fish ^a	4d LC ₅₀ (700) ^a	1000 ^c	70 ^a	0.08 (< 0.01–0.18)	0.10 (0.04–0.20)
$\sum$ HCHs					5.42 (1.79–9.52)	5.92 (0.11–10.36)
Heptachlor	Fish ^b	96h LC ₅₀ (0.8) ^b	100 ^e	0.8 ^b	9.59 (< 0.01–22.38)	13.97 (1.38–30.50)
Aldrin	Crustaceans ^c	2d EC ₅₀ (32) ^c	1000 ^c	1200 ^c	< 0.01	< 0.01
Heptachlor epoxide	Fish ^b	96 h LC ₅₀ (5.3) ^b	100 ^e	5.3 ^b	0.07 (< 0.01–0.36)	0.06 (< 0.01–0.33)
Endosulfan I	Fish ^d	NOEC(0.0001) ^d	100 ^e	10 ^d	0.49 (< 0.01–4.07)	0.06 (< 0.01–0.55)
Dieldrin	Fish ^b	96 h LC ₅₀ (0.62) ^b	100 ^e	0.62 ^b	0.53 (< 0.01–7.47)	Nd ^f
Endrin	Fish ^c	4d LC ₅₀ (2800) ^c	100 ^c	1000 ^c	< 0.01	Nd.
$\sum$ OCPs					16.10 (4.87–29.92)	20.01 (6.14–31.16)

^aHu et al. (2015)^bCho et al. (2014)^cUSEPA, ECOTOX^dVryzas et al. (2011)^eOgbeide et al. (2015)^fNd. not detected^gAverage value (min value–max value)

The RQ value of 4 HCH isomers in surface water ranged from 1.79 to 9.52 with an average of 5.42, which was similar with that in groundwater. With regard to the contributions of the different OCPs to the relative concentrations of the mixture, γ -HCH played

a dominant role in the comprehensive ecological risk. Guo and Feng (2014) found that α -HCH and β -HCH were the dominant factors for carcinogenic and non-carcinogenic risks in shallow freshwater lakes in China that are used as drinking water source. This

difference might be a result of the divergence of residue concentration in these two regions. As the α/γ ratio of HCH in Guo's research was above 1, while that of this paper is below 1, it can be deduced that there was a higher historical input of γ -HCH in Shaying River Basin, which could also result in the low proportion of risk posted by α -HCH and β -HCH.

Carcinogenic risk assessment

The calculated TLCRs using the RAGS method are shown in Table 5. All the TLCR values of OCPs in surface water and groundwater were between 10^{-6} and 10^{-4} , suggesting that the carcinogenic risks posed by these contaminants were acceptable, but still needed considerably concern. Particularly, the TLCR value of γ -HCH and heptachlor was obviously higher than other OCPs both in surface water and groundwater, indicating their high carcinogenic risk potential. The result also indicated that the carcinogenic risk posted by OCPs in surface water and groundwater was low for dermal contact (bathing) and acceptable for drinking.

As from the literature, ingestion is evaluated as the main intake pathway for OCPs to the human body, the risks related to drinking water produced from surface

water and groundwater are much higher than those of dermal contact, which can be seen in Table 5 and is accordance with other studies (Wei et al. 2015). The carcinogenic risk value of HCHs in this research was similar with that of edible fish from Wuhan, China (Cui et al. 2015), higher than that of drinking water sources from middle China (Guo and Feng 2014), underground river waters in Southeast China (Hu et al. 2011) and surface waters in Sichuan Basin China (Liu et al. 2015), which was around 10^{-7} – 10^{-9} . In river water of Korea, there is a higher carcinogenic risk of 10^{-6} associated with OCPs (Lee et al. 2014). In this study, the OCPs in surface water and groundwater in Shaying River have much higher carcinogenic risks on human than those in tap water (with estimated carcinogenic risk of 10^{-8} – 10^{-10}) (Li et al. 2012), which indicated drinking water treatment as an efficient and necessary procedure for water resource utilization.

The carcinogenic risk of OCPs in groundwater is higher than that of surface water, which is probably because of the direct leaching of irrigation and rainfall through the agriculture soil. However, the calculations may have over- or underestimated the risk because of the uncertainty of the residual concentration and intake rate. Besides, the research about assessing the

Table 5 Carcinogenic risk of OCPs in surface water and groundwater of Shaying River Basin

OCPs	Surface water			Groundwater		
	CR _{ingestion} ^b	CR _{dermal} ^c	TLCR ^{ab}	CR _{ingestion} ^b	CR _{dermal} ^c	TLCR ^{ab}
α -HCH	1.8	1.2	1.8	3.2	2.1	3.2
β -HCH	2.9	2.0	2.9	4.0	2.6	4.0
γ -HCH	4.9	3.3	4.9	5.9	4.0	5.9
δ -HCH	5.8	3.9	5.8	6.6	4.4	6.6
$\sum$ HCHs	15	10	15	20	13	20
Heptachlor	7.7	5.1	7.7	10	7.0	10
Aldrin	2.1	1.4	2.1	2.1	1.4	2.1
Heptachlor epoxide	1.3	8.4	1.3	1.5	1.0	1.5
Endosulfan I	7.9	5.3	7.9	5.1	3.4	5.1
Dieldrin	4.3	2.9	4.3	Nd ^e	Nd.	Nd.
Endrin	5.7	3.8	5.7	Nd.	Nd.	Nd.
$\sum$ OCPs	45	30	45	39	26	39

^aTLCR = CR_{ingestion} + CR_{dermal}

^bOne per ten thousand (10^{-5})

^cOne per billion (10^{-10})

^dNd. not detected

carcinogenic risk of surface water and groundwater in the same region simultaneously was limited. Thus, more work should be done before accurate conclusion could be drawn.

Conclusions

- In the research area, the concentration of $\sum$ OCPs ranged from 21.0 to 61.4 ng L⁻¹ in groundwater, and from 12.3 to 77.5 ng L⁻¹ in surface water, with the mean concentration of 44.6 and 30.6 ng L⁻¹, respectively. The concentration of $\sum$ HCHs ranged from 8.6 to 26.7 ng L⁻¹ in groundwater, and from 2.5 to 23.9 ng L⁻¹ in surface water, with the mean concentration of 18.7 and 14.3 ng L⁻¹, respectively.
- According to the results of main components analysis, the prominent OCPs contaminant in surface water of Shaying River Basin was HCHs and endosulfan I. Among the HCH isomers, γ -HCH and δ -HCH were the dominant composition. HCHs and heptachlor were with the highest concentration in groundwater of Shaying River Basin.
- The α -HCH/ γ -HCH ratios were all less than 3, which confirmed the use of lindane as the major source of HCHs contamination in the research area. Heptachlor and α -HCH, as well as endosulfan and heptachlor epoxide, had a strongly significant positive correlation, suggesting an associated usage of the two pair OCPs.
- An exceeding RQ value was observed for γ -HCH (lindane), heptachlor and dieldrin, revealing its extremely high ecological risk on aquatic organism in the research area. All the TLCR values of the selected 10 OCPs in surface water and groundwater were between 10⁻⁶ and 10⁻⁴, suggesting that the carcinogenic risks posed by these contaminants were acceptable based on the evaluation standard applied in this research.

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Compliance with ethical standards

Conflict of interest The authors have declared no conflict of interest.

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