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1 **Use of feathers to assess polychlorinated biphenyl and organochlorine
2 pesticide exposure in top predatory bird species of Pakistan**

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22

23 **Abstract**

24 Little is known about the levels of organochlorines (OCs) in predatory bird species from Asia or
25 the factors governing their concentration. This study is the first report on concentration of
26 polychlorinated biphenyls (PCBs) and organochlorine pesticides (OCPs) in predatory birds of
27 Pakistan. The concentrations of PCBs and OCPs were investigated using tail feathers of ten
28 different species of predatory birds. In addition, concentration differences among body, tail,
29 primary and secondary feathers were investigated for six individuals of black kite (*Milvus*
30 *migrans*). Ranges of concentrations were highest for dichlorodiphenyldichloroethylene (*p,p'*-
31 DDE: 0.11-2163 ng g⁻¹ dry wt.) followed by dichlorodiphenyltrichloroethane (*p,p'*-DDT: 0.36-
32 345 ng g⁻¹ dry wt.), hexachlorobenzene (HCB: 0.02-34 ng g⁻¹ dry wt.), Σ PCBs (0.03-16 ng g⁻¹
33 dry wt.) and *trans*-nonachlor (TN; 0.01-0.13 ng g⁻¹ dry wt.). CB 118, 153, 138, and 180 along
34 with *p,p'*-DDE were found as most prevalent compounds. Σ PCBs and Σ DDTs were significantly
35 different among species (both $p<0.01$) and omnivorous, scavengers, carnivorous and piscivorous
36 trophic guilds (all $p<0.03$). Whereas only Σ PCBs were significantly different ($p<0.01$) among
37 different families of birds. Values of stable isotopes ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$) differed significantly (all
38 $p<0.01$) among species, families, trophic guilds as well as terrestrial and aquatic habitat but not
39 between nocturnal and diurnal predators ($p=0.22$ for $\delta^{13}\text{C}$; $p=0.50$ for $\delta^{15}\text{N}$). Concentrations of
40 Σ PCBs, Σ DDTs and *trans*-nonachlor, but not HCB ($p=0.86$), were significantly different among
41 different feather types (all $p<0.01$). Trophic and taxonomic affiliation as well as dietary carbon
42 sources ($\delta^{13}\text{C}$) for species were identified as the variables best explaining the observed variation
43 in exposure to the studied compounds. The significance of contributing factors responsible for
44 OC contamination differences in predatory birds should be further elucidated in future studies.

45

46 **Key words:** trophic guild, feathers, habitat, POPs, $\delta^{13}\text{C}$, $\delta^{15}\text{N}$

47 **1. Introduction**

48 During the past few decades, prolific discharge of legacy persistent organic pollutants (POPs)
49 from industrial, urban and agricultural sources have remained a cause of many environmental
50 concerns particularly related to their toxic effects in humans and wildlife (Letcher et al., 2010).
51 These chemicals are persistent, bioaccumulative, toxic and travel large distances through long-
52 range transport (Vorkamp and Rigét, 2014). Two major classes of POPs, i.e. polychlorinated
53 biphenyls (PCBs) and organochlorine pesticides (OCPs) have been introduced after the
54 industrial revolution in 1920 and are still widespread in the environment (Lohmann et al., 2007)
55 despite being legally mitigated on a worldwide scale (UNEP, 2011). PCBs were used in a wide
56 array of substances as a coolant or additives and escape into the environment during their usage,
57 packaging and storage as well as through leaching from landfills (Covaci et al., 2006). On the
58 other hand, OCPs are chlorine-containing organic pesticides which were predominately used as
59 insecticides (Ali et al., 2014). Among a variety of compounds, dichlorodiphenyltrichloroethane
60 (DDT) was the mostly heavily used pesticide after World War I and its production was banned
61 from USA, Europe, China and Japan after 1972 when its toxic effects became established
62 (Tanabe et al., 1998). Hexachlorobenzene (HCB) was found among the most prevalent OCPs
63 because of its use as insecticide as well as an industrial by-product (Corsolini et al., 2006).
64 Besides these, metabolites of chlordanes (CHLs) were also found as compounds of concern
65 because of their exacerbated use as insecticides and their reported adverse effects upon wildlife
66 (Letcher et al., 2010). These OCPs mainly get their way into the environment during their
67 production, application and storage and are eventually dispersed through runoff and air currents
68 (Guan et al., 2009).

69 Ever since the toxicological significance of POPs was suspected, predatory birds have been
70 successfully used as sentinels to assess the levels of these compounds in the environment (et al.,
71 1993; Dauwe et al., 2005; Jaspers et al., 2006). However, sampling of predatory birds often
72 encounters various practical and ethical impediments. Use of non-destructive tissues, such as
73 blood, feathers and preen oil, is usually recommended as a preferable choice in case of
74 predatory birds. Among these, the use of feathers has become more and more applicable
75 because it is less invasive, comes along with easy collection and storage, and provides a
76 valuable assessment of internal body burdens of POPs (Jaspers et al., 2006). Some of the recent
77 studies have also emphasized to evaluate different types of feathers i.e. body, tail, primary and

78 secondary which could best represent the level of the studied compounds (Eulaers et al.,
79 2014b; García-Fernández et al., 2013; Jaspers et al., 2011).

80 Levels of POPs in avian tissues are influenced by a multitude of biological, spatial and
81 ecological factors (Eulaers et al., 2013; Lavoie et al., 2010). Trophic levels/feeding guilds and
82 taxonomic affiliation of species, locational and dietary exposure well as individual condition
83 factors such as gender, age and reproductive status may significantly influence the concentration
84 of POPs in birds (Eulaers et al., 2013; Behrooz et al., 2009). In general, POP concentrations at
85 higher trophic level species mainly stem from dietary intake, which can be quantified using
86 ratios of stable nitrogen and carbon isotopes (SIs; Eulaers et al., 2014a). The ratio of heavier $\delta^{15}\text{N}$
87 to lighter $\delta^{14}\text{N}$ ($\delta^{15}\text{N}$) provides information about the trophic level of an individual because it
88 enriches with each trophic level (Huang et al., 2013). The ratio of carbon SIs ($\delta^{13}\text{C}$: $^{13}\text{C}/^{12}\text{C}$) is
89 used as an indicator for dietary origin because of the varying degree of depletion of ^{13}C stable
90 isotopes in primary producers from different habitats (Boecklen et al., 2011; Jardine et al., 2006).
91 Although the use of SIs has shown promising to investigate trophodynamics of POPs, it has had
92 less focus in predatory birds, particularly those from the Asian continent.

93 Predatory birds of the southern Asian region are particularly exposed to a high magnitude of
94 legacy POPs because of their historical and current use in this region (Ali et al., 2014). Levels of
95 POPs have been documented in biotic as well as abiotic components of the environment from
96 South Asia (Sarkar et al., 2008; Yadav et al., 2015), but predatory birds have received less
97 attention (Abbasi et al., 2016). Contamination of birds with POPs has only been reported in eggs
98 of little (*Egretta garzetta*) and cattle egret (*Bubulcus ibis*) from Pakistan (Khan et al., 2014;
99 Malik et al., 2011; Sanpera et al., 2003). Seeing this scarcity of exposure data, the present study
100 was designed to investigate the current concentration levels of different OC compounds using
101 feathers of multiple predatory bird species of Pakistan. Further, we evaluated the importance of
102 various factors governing interspecific variation of OC exposure including intraspecific
103 variations through carbon and nitrogen SI values. Lastly, the suitability of different feather types
104 to characterize OC exposure was evaluated by comparing body, tail, primary and secondary
105 feathers from black kites (*Milvus migrans*).

106

107 **2. Methodology**

108 *2.1. Sample collection*

109 Feather samples ($N=76$) from ten different species of predatory birds were collected between
110 June 2012 to September 2014 (Fig. 1). Species selected for this study included black kite
111 ($N=13$), Eurasian sparrowhawk (*Accipiter nisus*, $N=10$), common kestrel (*Falco tinnunculus*,
112 $N=4$), red-necked falcon (*Falco chicquera*, $N=2$), Indian vulture (*Gyps indicus*, $N=9$), white-
113 rumped vulture (*Gyps bengalensis*, $N=12$), spotted owlet (*Athene brama*, $N=10$), little owl
114 (*Athene noctua*, $N=6$), great cormorant (*Phalacrocorax carbo*, $N=4$) and grey heron (*Ardea*
115 *cinerea*, $N=6$). Tail feathers were obtained from all these species. In addition, tail, body,
116 primary and secondary feathers were collected from six individuals of black kite to investigate
117 concentration differences among feather types. Sampling details of each site are summarized in
118 Table S1 (supplementary information). Predatory birds were sampled mainly from different
119 towns and cities and their outskirts in Punjab province, which is considered a hub of agricultural
120 activities of the country. Samples of black kite and spotted owlet were also collected from two
121 metropolitan cities, i.e. Lahore and Rawalpindi, with higher expected anthropogenic input than
122 other sites. Samples of both the vulture's species were obtained from their isolated and remotely
123 located colonies (S1&S2) at Nagar Parker, Sindh Province. Grey heron was the only species
124 sampled from northern regions at Lulusar Lake (S13), which is a remote waterbody. Each
125 species was sampled from one location except black kite, Eurasian sparrowhawk and spotted
126 owlet, which were sampled from two different locations (Table S1). Black kites and spotted
127 owlet were sampled around the outskirts of Lahore, which is a metropolitan city with extensive
128 agricultural activities in its suburbs, and Islamabad, which is a relatively smaller city with very
129 small scale agricultural activities in its premises. The third species, Eurasian sparrowhawk, was
130 sampled from Mianwali and Khaniwal, which are both small cities with extensive agricultural
131 lands around. Further, species are discussed under various categories based on their taxonomic
132 affiliation (families; accipitridae, ardeidae, falconidae, phalacrocoracidae, strigidae), trophic
133 guilds (Omnivorous, scavenger, carnivorous, piscivorous, habitats (terrestrial or aquatic) and
134 feeding regimes (diurnal or nocturnal). All the samples used in this study were taken from birds
135 captured in the framework of other studies. A special permit from CITES authorities in Pakistan
136 was acquired for shipping and transport of the samples of the two critically endangered vulture
137 species. After collection, feathers were kept in zipped plastic bags and stored at -20°C until
138 chemical analysis.

139 2.2. Quantification of PCBs and OCPs

140 The procedure for cleanup and extraction of POPs was adapted from previous described
141 methods (Dauwe et al., 2005; Jaspers et al., 2006). Feathers were thoroughly washed with
142 deionized water to remove exogenous dust particles and other unwanted depositions. After
143 washing, feathers were covered with standard laboratory paper and dried overnight at ambient
144 temperature. Dried feathers were cut into pieces of ~1 mm, weighed and transferred to
145 analytical glass recipients. Initially, feather samples were spiked with the internal standard CB
146 143 (50 μ L of 200 pg μ L $^{-1}$) and incubated overnight at 45°C in HCl (4M) and
147 hexane:dichloromethane (4:1; v:v). From the incubated mixture, analytes were liquid-liquid
148 extracted using hexane:dichloromethane (4:1; v:v). Cleanup of the resulting extract was
149 performed on acidified silica (800 mg; 44% H₂SO₄) topped with anhydrous Na₂SO₄ (400 mg),
150 and analytes were eluted with hexane:dichloromethane (4:1; v:v). Finally, the cleaned-up
151 extracts were concentrated using a gentle flow of Nitrogen gas, reconstituted in 80 μ L *iso*-
152 octane, and transferred to injection vials. The whole process of clean-up and extraction was
153 performed at the Bird ecotoxicology laboratory, Norwegian University of Science and
154 Technology (Trondheim, Norway), whereas the concentrations of PCBs and OCPs were
155 quantified at the Toxicological Center, University of Antwerp (Wilrijk, Belgium).

156 The concentrations of PCBs and OCPs were quantified using a mass spectrometer (Agilent MS
157 5973, Palo Alto, CA, USA) operated in electron-capture negative ionization mode to a gas
158 chromatograph (Agilent GC 6890, Palo Alto, CA, USA). A total of 19 PCB congeners (CB 105,
159 118, 146, 153, 138, 187, 183, 128, 174, 177, 171, 156, 180, 170, 199, 196/203, 194, 206, 209),
160 HCB, *trans*-nonachlor (TN), *cis*-nonachlor (CN), oxychlordane (OXC), and DDTs, i.e. *p,p'*-
161 DDE, *p,p'*-DDT, were measured. In all samples, only high chlorinated PCBs were measured.
162 However, in a few samples (those with the highest concentrations of PCBs), we have attempted
163 to measure lower-chlorinated PCB congeners. Yet, detection limits were higher and the lower
164 chlorinated PCB congeners measured (CB28, CB52, CB95 etc.) were below the limit of
165 quantification (<LOQ) and are thus not reported. Concentrations of analytes were expressed as
166 ng g $^{-1}$ dry weight (dw). Internal standards were purchased from Accustandard (New Haven, CT,
167 USA), while pesticide-grade solvents (Merck, Darmstadt, Germany) were used throughout the
168 entire process. Mean recoveries of internal standards were 52% \pm 13 for PCBs in all samples.
169 The same internal standard (CB 143) was used for other OCPs. For quality assurance, in each

batch of 10 samples, a procedural blank was prepared and analyzed. LOQs for different analytes were set at 3*SD of the procedural blank values. When analytes were not detected in blanks, the LOQ was calculated using a 10:1 signal to noise ratio.

2.3. Stable isotopes measurement

Composition of stable nitrogen and carbon isotopes was measured at the Center for Permafrost (University of Copenhagen, Denmark). We adapted the previously reported procedure by Eulaers et al. (2014a) for the measurement of SIs in feathers. Briefly, a representative homogenized subsample of 0.5 to 2.0 mg was wrapped into a tin combustion cup, and the ratios for stable carbon and nitrogen isotopes were measured by continuous flow using an elemental analyzer (CE 1110, Thermo Electron, Milan, Italy) coupled to a mass spectrometer (Finnigan MAT Delta PLUS, Thermo Scientific, Bremen, Germany). The ratios of SIs were expressed as

$$\delta X (\text{‰}) = \left(\frac{R_{\text{sample}}}{R_{\text{standard}}} - 1 \right)$$

with X representing the C or N SIs and R representing their corresponding ratios ($^{13}\text{C}/^{12}\text{C}$, $^{15}\text{N}/^{14}\text{N}$) in the sample or standard. Reference samples (Atropin) were included for the positive evaluation of analytical performance. The instrument was calibrated by employing pure gases of CO₂ and N₂ against the certified reference material of sucrose and (NH₄)₂SO₄ provided by the International Atomic Energy Agency (IAEA, Vienna, Austria). The SI ratios were calculated against the international standards Vienna PeeDee Belemnite (vPDB) and atmospheric N₂ (AIR) respectively. Analytical precision was maintained at 0.1‰ SD.

2.4. Statistical analysis

All the statistical computations were performed using SPSS (IBM 20) and R (version 3.2.3). Firstly, screening of the data was performed as suggested by Zuur et al. (2010) to avoid common statistical errors. Data was \log_{10} transformed after testing for normality using Q-Q plots and Shapiro-Wilk's tests (all $p < 0.05$). Only those compounds which were detected above the limit of quantification (>LOQ) in at least 50% of the samples of a species were treated for further statistical analysis. Missing values for these compounds were substituted with the proportion of detected samples*LOQ. The *null*-hypothesis was rejected at $\alpha=0.05$. Firstly, differences of PCB and OCP concentrations among species, families, omnivorous, scavengers, carnivorous and piscivorous trophic guilds, habitats (aquatic/terrestrial), feeding regime (diurnal or nocturnal) as

198 well their associations to dietary proxies ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$) were tested through analysis of variance
199 (ANOVA). Subsequent post-hoc Tukey's tests for honest significant differences (HSD) were
200 used for multiple comparisons. Further, above-mentioned variables were evaluated for their
201 capacity to explain the observed variation in levels of PCBs and OCPs using Akaike's
202 Information Criteria (AICc) as discussed previously (Johnson and Omland, 2004). A separate
203 AIC-based selection was run for each compound to evaluate the factors best governing the
204 observed variation in PCB and OCPs concentrations. Separate AIC based model was run for
205 each compound based on of the fact that they have different physicochemical properties and may
206 be influenced differently by different factors. Associations of PCBs and OCPs with dietary
207 proxies ($\delta^{13}\text{C}$, $\delta^{15}\text{N}$) were tested through linear regression. A separate ANOVA was performed to
208 determine the variation in concentrations among different feather types sampled from black kite.
209

210 **3. Results and Discussion**

211 *3.1. Variation in OC concentrations and profiles*

212 To the best of our knowledge, so far PCBs and OCPs have never been quantified in feathers of
213 predatory birds from Pakistan. The measured concentrations for different compounds of PCBs
214 and OCPs are summarized in table S2. Out of the 19 PCBs congeners targeted, 14 congeners,
215 i.e. CB 105, 118, 146, 153, 138, 187, 183, 128, 156, 180, 170, 199, 196/203, 194 were detected
216 above LOQ in $\geq 50\%$ samples for minimum one to maximum all the species. Among the studied
217 compounds CB 153, HCB, *p,p'*-DDE and *p,p'*-DDT were detected in all the studied species
218 whereas all other compounds were variably detected (Figure S1). Compounds such as CB 171,
219 174, 177, 206, 209, as well as OXC and CN were not detected above LOQ in $\geq 50\%$ of the
220 samples in any species (Figure S1), hence not further discussed. In general, the trend
221 $\Sigma\text{DDTs} > \text{HCB} > \text{PCBs} > \text{TN}$ was depicted in tail feathers of predatory birds from Pakistan. The
222 concentration ranges (minimum-maximum) recorded in this study were $0.11\text{-}2163\text{ ng g}^{-1}$ dry wt.
223 for DDTs, $0.02\text{-}34\text{ ng g}^{-1}$ dry wt. for HCB and $0.03\text{-}16\text{ ng g}^{-1}$ dry wt. for ΣPCBs and $0.01\text{-}0.13\text{ ng g}^{-1}$ dry wt.
224 for TN respectively. Previously, screening of OCs has only been carried out in
225 eggs of little egret (Sanpera et al., 2003) and cattle egret (Khan et al., 2014; Malik et al., 2011)
226 from Pakistan. In those studies, reported concentrations of OCs were higher probably due to the
227 higher lipid content in egg. In recent global literature, PCBs and OCPs in predatory birds have
228 been mostly reported in egg, muscle, liver, kidney and other non-keratinous tissues (Chen et al.,

229 2009; Jaspers et al., 2006; Kocagöz et al., 2014; Lavoie et al., 2010; Peng et al., 2015; Sun et
230 al., 2014; Zhang et al., 2011) whereas only few studies are available for comparison of PCBs
231 and OCPs in feathers. Compared to findings of the present study, \sum DDTs were found
232 comparable whereas \sum PCBs and HCB were approximately 5 to >50 fold higher in feathers from
233 different predatory bird species from south-west of Iran (Behrooz et al., 2009). Similarly,
234 feather concentrations of \sum DDTs and HCB were found comparable to our findings, whereas
235 \sum PCBs levels were relatively higher in different waterbird species from the Caspian Sea coast,
236 Northern Iran (Rajaei et al., 2011). Regarding the European scenario, compared to our study
237 Jaspers et al., (2007) reported relatively higher \sum PCBs, comparable HCBs and lower \sum DDTs
238 levels in tail feathers of multiple predatory species from Belgium. Further, Jaspers et al., (2009)
239 reported a comparable level of *p,p'*-DDE (1.07-139 ng g⁻¹ dry wt.), relatively lower level of
240 *p,p'*-DDT (0.38-11.8 ng g⁻¹ dry wt.) and fairly high range of \sum PCBs (2.92-236 ng g⁻¹ dry wt.) in
241 tail feathers of common magpie (*Pica pica*) from Belgium. Similarly, concentrations of *p,p'*-
242 DDE, HCB and TN but not \sum PCBs of this study were found comparable or slightly higher than
243 reported in tail feathers of white-tailed eagle (*Haliaeetus albicilla*) from western Greenland
244 (Jaspers et al., 2011).

245 The contribution of the detected compounds is illustrated in figure 2a, whereas profiles for PCBs
246 and DDTs are shown in figure S2a,b. Among the detected compounds, *p,p'*-DDE was found as
247 the predominant compound in predatory birds of the current study followed by *p,p'*-DDT and
248 congeners of PCBs, HCB and TN respectively, which is in line with previous studies (Chen et
249 al., 2009; Rajaei et al., 2011). Among PCBs, CB 118, 153, 138, 180, 170 and 194 were observed
250 as more prevalent congeners in tail feathers (figure S1). In the present study, PCB congeners
251 with six (*hexa*-CBs) and seven chlorines (*hepta*-CBs) dominated in terrestrial species whereas
252 those containing five chlorines (*penta*-CBs) were more prevalent in aquatic species (Figure S2a),
253 which is in agreement with previous findings (Yu et al., 2014; Jaspers et al., 2007). Previously,
254 Abbasi et al., (2016) reported that *p,p'*-DDE has been unanimously detected as predominant
255 metabolite of DDTs in Asian studies on birds. In contrast, *p,p'*-DDE has been found as a
256 predominant compound in feathers of European predatory birds (Jaspers et al., 2007; Eulaers et
257 al., 2013). In general, the elevated level of DDTs in this study corresponds to their wide scale use
258 as pesticide in Pakistan (Ali et al., 2014). Similar to our findings, CB 153, 180, 138 have been
259 reported as predominant congeners in feathers of predatory birds from different parts of the

260 world (Chen et al., 2009; Behrooz et al., 2009, Jaspers et al., 2007, 2011). Earlier, Dauwe et al.
261 (2005) suggested that elevated levels of lower-chlorinated PCB congeners in feathers may be
262 associated with differential elimination and distribution mechanisms.

263 *3.2. Intraspecific variation*

264 We aimed at elucidating intraspecific variation in OC exposure through linear regression of
265 concentrations versus the dietary proxies (SI values) and also by plotting the individuals of
266 species on a $\delta^{13}\text{C}$ / $\delta^{15}\text{N}$ biplot (Figure 3a). The distribution of species in the $\delta^{13}\text{C}$ / $\delta^{15}\text{N}$ biplot
267 reflects within and among species variations based on the differences in values of dietary
268 proxies (figure 3a). $\delta^{13}\text{C}$ values reflect dietary separation of carbon sources, whereas values of
269 $\delta^{15}\text{N}$ are used as a proxy for their position at trophic food chain (Yu et al., 2011). Aquatic
270 species i.e. grey heron and great cormorant, in the present study were found to be feeding at a
271 higher trophic position compared to terrestrial birds ($p<0.01$), which is in line with previous
272 studies (Hong et al., 2014; Jaspers et al., 2007). However, relatively scattered distribution
273 (figure 3a) of the individuals of aquatic birds depicted their wide dietary flexibility. Earlier,
274 based on stable isotope characterization, Sørmo et al., (2011) found that the diet of coastal
275 herring gull (*Larus argentatus*) was influenced by terrestrial sources. Moreover, Morkuné et al.,
276 (2011) reported that great cormorant switched its diet at various stages of life whereas grey
277 herons showed consistent dietary habits throughout their life span. We suspect that higher
278 trophic positions on $\delta^{13}\text{C}$ / $\delta^{15}\text{N}$ layout and relatively scattered distribution of aquatic birds in our
279 study were because of their more specialized dietary habits as well as varying exposure when
280 compared to terrestrial species. In contrast, individuals of Indian vulture and white-rumped
281 vultures were found with a relatively clustered distribution in the $\delta^{13}\text{C}$ / $\delta^{15}\text{N}$ biplot which might
282 be associated with more specialized dietary habits (Yu et al., 2011). We sampled these two
283 vultures from their isolated remote colonies where they mostly consumed the locally available
284 carrions, which restricts their choice for diverse food sources. Interestingly, Indian vultures
285 were observed to be feeding at a relatively higher trophic level as compared to white-rumped
286 vultures suggesting differences in dietary habits of these two species. Similarly, a scattered
287 distribution of black kites in the SI biplot reflects the availability of diverse food choices for
288 birds dwelling in human proximity and close to urban environments. Earlier, Barón et al.,
289 (2014) observed black kite as a versatile feeder ranging from human refusals, small insects,
290 invertebrates, up to small mammals, frogs and snakes in urban and township areas. The results

confirm our assumption of exploitation of diverse feeding sources by black kites. Similarly, Eurasian sparrowhawk, red-necked falcon and common kestrel were also found with relatively wide ranging distributions in the $\delta^{13}\text{C} / \delta^{15}\text{N}$ biplot indicating flexibility in food choices for these species as well (Chen et al., 2009; Elliot et al., 2009, Luzardo et al., 2014). Among two owl species, spotted owlet residing in urban and suburban localities depicted a relatively more scattered distribution in the $\delta^{13}\text{C} / \delta^{15}\text{N}$ biplot suggesting its dietary flexibility compared to tight clustering of little owl. Certain overlap among terrestrial species but not aquatic species is obvious from the $\delta^{13}\text{C}/\delta^{15}\text{N}$ biplot (figure 3a) suggesting their potential sympatric distribution (Zhang et al., 2011) as well as shared feeding sources (Elliott et al., 2009). Distribution of individuals birds of different predatory species on $\delta^{13}\text{C}/\delta^{15}\text{N}$ layout suggest that OC bioaccumulation is considerably influenced by habitat and dietary exposure in addition to different other factors. Regression analysis revealed a weak and non-significant (except few) but positive association between dietary proxies and concentration of compounds analyzed (R^2 ranged between 0.01 to 0.99). For $\delta^{15}\text{N}$, regression was significant in black kite (for PCBs; $R^2=0.44$ and HCB; $R^2=0.42$, both $p<0.01$) and great cormorant (for DDTs; $R^2=0.42$, $p=0.02$). Conversely, for $\delta^{13}\text{C}$ values regression was only significant in Eurasian sparrowhawk (for PCBs, $p<0.03$; and DDTs $p<0.04$). The regressions between dietary proxies ($\delta^{13}\text{C}$, $\delta^{15}\text{N}$) and OCs concentrations were found non-significant ($p>0.05$) for other studied species.

Locational differences of POPs accumulation were also done for three species for which samples were available for comparison between sites. We have only three species, i.e. black kite, Eurasian sparrowhawk and spotted owlet, for comparison between two different locations because all other species were sampled from only one site (table S1). For black kite, significant differences for $\delta^{13}\text{C}$ ($p=0.03$), $\delta^{15}\text{N}$ ($p=0.02$) and HCB ($p=0.02$) were observed between sites. ΣDDTs at Lahore and ΣPCBs at Rawalpindi were found slightly higher although not significantly different ($p>0.05$) between sites which is possibly associated with significant differences for the values of dietary proxies. None of the compounds nor stable isotope values differed significantly ($p>0.05$) between sites for spotted owlet or Eurasian sparrowhawk indicating similar exposure to pollutants at both sites. The above results for stable isotopes suggest that black kite, being a more urban dwelling species, may switch its feeding choices (Barón et al., 2014) based on availability at different sites hence reflect differential exposure to OCs. Conversely, spotted owlet and Eurasian sparrowhawk, which remain consistent between

322 sites due to lower availability of choices at suburban to forested sites hence reflect similar
323 exposure to OCs.

324 *3.3. Interspecific differences*

325 In the present study, highest median concentrations (minimum-maximum) of Σ PCBs at 7.9 ng g^{-1} dry wt.
326 ($0.4\text{-}15.4 \text{ ng g}^{-1}$ dry wt.) in red-necked falcon, Σ DDTs at 195.5 ng g^{-1} dry wt. (7.1-
327 1022.2 ng g^{-1} dry wt.) in common kestrel and HCB in Eurasian sparrowhawk at 0.7 ng g^{-1} dry
328 wt. ($0.1\text{-}34.4 \text{ ng g}^{-1}$ dry wt.) were recorded. All three species of the current study are terrestrial
329 predators that mainly feed upon small birds, rats, mouse, frogs, snakes and invertebrates with
330 some flexibility in their dietary choices (Behrooz et al., 2009). Further, these species mainly
331 reside in suburbs and agricultural lands around cities where they can get their prey easily.
332 Relatively higher concentrations of OCPs in common kestrel and Eurasian sparrowhawk in
333 particular is possibly attributed to their higher dietary exposure to agricultural used pesticides
334 (Behrooz et al., 2009). This was further corroborated through the significant regression between
335 origin of dietary carbon ($\delta^{13}\text{C}$) and PCBs ($R^2=0.45, p=0.03$) as well as with DDTs ($R^2=0.41;$
336 $p=0.04$) in Eurasian sparrowhawk. Regression was not significant for common kestrel possibly
337 because of limited movement and exposure of this species (Jaspers et al., 2007). However,
338 relatively higher concentrations of PCBs in red-necked falcon must be considered with caution
339 because of the low sample size. Moreover, it is observed that in winter Eurasian sparrowhawk
340 and red-necked falcon move towards towns and cities from the agricultural lands to overcome
341 winter harshness, which increases their exposure to the urban sources of OCs (Chen et al.,
342 2009). Based on their utility and disposal, PCBs in particular and HCB up to some extent are
343 originating from urban sources and hence bioaccumulate in the tissues of top predators during
344 their winter feeding exposure at temporary stopover sites. To the best of our knowledge, the
345 exposure to PCBs and OCPs in vultures has never been studied. In the present study, we
346 sampled both vulture species from their remotely located colonies from Sindh province (figure
347 1, table S1), where the exposure to urban as well as agricultural chemicals is expected to be
348 minimal, corresponding to the lower concentrations of PCBs and OCPs in vultures of the
349 current study. In contrast, the concentrations of PCBs and OCPs were found relatively higher in
350 black kite and spotted owl because of higher exposure of these urban/suburban dwelling
351 species to both agriculture and urban sources of OCs (Barón et al., 2014). The concentrations of
352 Σ DDTs but not Σ PCBs were found relatively higher in aquatic birds of our study suggesting

353 that the surplus use of OCPs and environmental leaching of PCBs is more bioavailable in water
354 reservoirs than terrestrial food chain ([Rajaei et al., 2011](#)). Unexpectedly, we have observed
355 relatively high concentrations of \sum DDTs in grey herons, which were sampled from a relatively
356 high altitude pristine location (figure 1, table S1), suggesting that these compounds may also
357 move towards high laying areas through long range transport from their origin ([Wania and](#)
358 [Mackay 1996](#)).

359 Previously, interspecific variation in contaminant levels was attributed to the combined
360 influence of several biological, ecological and spatial factors ([Eulaers et al., 2013, 2014; Peng et](#)
361 [al., 2015](#)). In the present study, we have evaluated the importance of various factors, such as
362 trophic level, taxonomic affiliation, habitat and feeding regime, as drivers of OC exposure, and
363 have investigated how values of $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ may specifically serve as dietary tracers that
364 govern difference in OC concentrations among species. Concentrations of compounds such as
365 \sum PCBs and \sum DDTs, but not HCB ($p=0.08$), differed significantly among species ($p<0.01$ for
366 both) as well as trophic guilds (all $p<0.03$). Multiple comparison (Tukey HSD) test revealed that
367 more specialized predators such as Eurasian sparrowhawk and red-necked falcon differed
368 significantly from omnivorous and piscivorous birds (Table 1). Among families, only \sum PCBs
369 ($p<0.01$), but not \sum DDTs ($p<0.88$) nor HCB ($p<0.82$), were significantly different. The
370 bioaccumulation trend of PCBs in falconidae family was found significantly different from all
371 other families. Conversely, the corresponding differences between habitats and feeding regimes
372 were non-significant for \sum PCBs ($p<0.27$; $p<0.91$), \sum DDTs ($p<0.45$; $p<0.62$) and HCB ($p<0.45$;
373 $p<0.62$) respectively (Table 1). Values for $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ are found significantly different
374 ($p<0.01$) for all the above mentioned variables, except for feeding regime (both $p=0.22$).
375 Further, associations between dietary habits of species and contaminants concentrations were
376 obtained by regressing the \log_{10} normalized concentrations for \sum PCBs, \sum DDTs and HCB with
377 $\delta^{15}\text{N}$ values (figure 3b, c, d). Regression slopes for species depicted that association of log
378 normalized \sum PCBs and \sum DDTs with $\delta^{15}\text{N}$ values were stronger when compared to HCBs. This
379 suggest that different OC compounds vary in their potential of bioaccumulation in birds. This
380 bioaccumulation differences could possibly be attributed to differences of exposure and
381 physicochemical properties of compounds ([Behrooz et al., 2009](#)). However, species feeding at
382 different trophic level ($\delta^{15}\text{N}$ values) depicted similar trends for each \sum PCBs, \sum DDTs and HCBs
383 accumulation except for few species. In case of common kestrel, regression slope was positive

for \sum PCBs while it is relatively straight or negative for \sum DDTs and HCB. Alternatively, regression slope for \sum PCBs was negative for Indian vulture followed by positive and straight lines for \sum DDTs and HCB respectively. Regression slopes for all other species somehow depicted similar bioaccumulation trend for \sum PCBs, \sum DDTs and HCB with varying degree of positive or negative trends. Although there is no clear differences, as depicted by slopes that bioaccumulation of PCBs and OCPs vary between terrestrial and aquatic species, however the bioaccumulation trend (slopes) for grey heron but not greater cormorant were somehow similar to most of the terrestrial species. This differences between two aquatic species could be attributed to sufficient terrestrial exposure ([Ito et al., 2013](#)) and potential influence of terrestrial feeding sources in grey heron ([Sørmo et al., 2011](#)) compared to great cormorant which strictly rely on aquatic food sources. We have also evaluated the importance of different variables in explaining the magnitude of exposure of PCBs and OCPs in feathers of predatory birds through their respective AIC values (table 3). In different models which are separately run for each of the compounds, variables with the lowest AIC value (shown bold in table 3) best explain the observed variation in concentrations of the different compounds. The models suggested that the concentrations of \sum PCBs, *p,p'*-DDE, *p,p'*-DDT and \sum DDTs are best explained by the variable species. But we have run separate models for each compounds assuming that physicochemical differences of these compounds may varyingly influenced by factors governing their bioaccumulation. Interestingly, most of the PCB congeners are best explained either by trophic guild, $\delta^{13}\text{C}$ values or taxonomic affiliation of species. Concentration differences of lower-chlorinated compounds were found to be more governed by trophic guilds. Earlier, [Behrooz et al., \(2009\)](#) suggested that the interspecific variations in OC concentrations are mainly due to differences in feeding habits of the species. Very few congeners, i.e. CB 128 and 187, were best predicted by habitat or feeding regime differences. For future studies, we recommend to further elucidate the factors best predicting the bioaccumulation of OCs by investigating large sample size for each species.

410 *3.4. Variation based on feather type*

411 To present date, very few studies have reported the differential accumulation pattern of OCs in
412 different feather types of predatory birds. Earlier, [Jaspers et al. \(2011\)](#) found significant
413 differences in levels of different organic contaminants among different feathers types of white-

tailed eagles from Greenland. Similarly, Eulaers et al. (2014b) also detected varying accumulation trends of OCs in different feather types of barn owl (*Tyto alba*) from Belgium. Based on the assumption that OCs bioaccumulate differently among feathers types, we tested the concentrations of PCBs and OCPs as well as values of $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ among body, tail, primary and secondary feathers of six individuals of black kite. The concentration pattern of detected compounds in different types of feathers of black kite is shown in figure S5 (a-d) whereas detection frequencies are shown in figure S1b. Detection frequencies for body and tail feathers were similar, but higher than wing (primary and secondary) feathers. The concentration trend of OCs in different feathers types was also found as $\Sigma\text{DDTs} > \Sigma\text{PCBs} > \text{HCB} > \text{TN}$. Test results for significant differences of OCs among different feather types are presented in table 2.

Analysis of variance revealed that concentrations of ΣPCBs , ΣDDTs , TN differed significantly ($p<0.01$ for all three compounds) among body, tail, primary and secondary feathers. Whereas differences were non-significant for HCB ($p=0.86$), $\delta^{13}\text{C}$ ($p=0.65$) and $\delta^{15}\text{N}$ ($p=0.64$) among feathers types. Among the different feather types, body feathers were found with highest mean concentrations of ΣPCBs , ΣDDTs , TN and $\delta^{13}\text{C}$ whereas HCB was highest in secondary feathers (table 2). Based on the higher detected concentrations and detection frequencies, we believe that body feathers could be used as a most useful tool for future biomonitoring studies in predatory birds as suggested earlier by Jaspers et al., (2011). However, we urge for a more elaborate investigation in the future with larger sample sizes to confirm this. Eulaers et al., (2014b) suggested that relatively elevated concentrations of OCs in tail and body compared to primary feathers are mainly associated with preening activity and moult pattern of the barn owl. Further, external contamination through preening has also been suggested to alter the level of OCs in feathers (Jaspers et al., 2008; 2013). The specific moulting pattern of different feathers types in black kite is currently unknown. However, we predict that a higher influence of preening activity on body and tail feathers of black kites because of their proximity to preen gland or beak (Eulaers et al., 2014b) may be associated with higher concentrations of OCs in these feather types. It has been suggested that moulting pattern and age (Jaspers et al., 2011) as well as length and growth of feathers (Bortolotti et al., 2010) are key factors to describe the variations in OCs levels among different types of feathers. Besides these, various confounding factors such as differences in moulting strategy (Jaeger et al., 2013) as well as feeding and migratory habits (García-Fernández et al., 2013) of species can influence the levels of OCs in

445 feathers. Although we have not evaluated any of the above mentioned factors in this study
446 because of lack of information during data collection, we suspect that a combined effect of just
447 mentioned factors may be responsible for differences in bioaccumulation patterns of OCs. In
448 future studies, we suggest to further elucidate the factors responsible for differential
449 accumulation of OCs among various feather types.

450

451 **4. Conclusions**

452 The present study is the first to report the levels of PCBs and OCPs in predatory birds of
453 Pakistan. Various contributing factors explaining the intra- and interspecific differences in PCBs
454 and OCPs levels in predatory birds were also evaluated. We concluded that PCBs and OCPs
455 could easily be quantified in predatory birds using feathers, particularly body feathers because of
456 their high detectability. Significant differences in concentrations of PCBs and some OCPs among
457 different feathers types emphasize the need for appropriate feather choice in future toxicological
458 studies. Compared to PCBs, concentrations of OCPs, particularly DDTs, were found higher in
459 predatory birds reflecting the large scale historical and current application of pesticides in
460 Pakistan. In general, PCB levels reported in predatory birds of Pakistan are found lower than
461 those of European studies whereas OCPs are relatively comparable. To get a broader picture, we
462 urge future research to investigate the significance of contributing factors influencing the levels
463 of OCs using multiple species of predatory birds from a wide geographical range.

464 **Conflict of interest**

465 The authors declare no conflicts with any person or organization.

466 **Supplementary information**

467 All the supporting material cited in this manuscript is available in the online supplementary files.

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474

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Table 1: Pairwise comparisons for PCBs, DDTs, HCB (ng g⁻¹ dry wt.) and stable isotope values (‰) in tail feathers among different groups of birds based on ANOVAs. Means are shown under each variable whereas differences through multiple comparisons are illustrated through different alphabetic characters. Groups with a single different letter differ significantly from the other. Sample sizes in each group are given in parentheses.

Test variable	Compounds/SIs	Test for significance			Mean concentrations and Significance levels (alphabets) among/between groups (Tukey-HSD)									
		df	F	P	BK (10)	WRV (12)	IV (9)	ESH (10)	CK (4)	RNF (2)	GH (6)	GC (4)	SO (10)	LO (6)
Species	δ13C	9,60	6.14	<0.01	-20.49 AB	-17.54 B	-17.53 B	-20.89 AB	-20.34 AB	-24.73 A	-24.00 A	-21.23 AB	-17.20 B	-20.64 AB
	δ15N	9,60	8.28	<0.01	9.01 AB	9.57 ABC	11.51 BCD	8.58 A	8.33 A	8.51 A	12.18 CD	13.60 D	10.67 ABC	9.97 ABC
	ΣPCBs	9,66	2.55	<0.01	2.05 B	0.45 B	0.57 B	3.56 AB	5.20 AB	8.41 A	4.10 AB	2.46 AB	2.04 B	2.74 AB
	ΣDDTs	9,66	2.61	<0.01	12.97 B	2.03 B	6.15 B	647.98 A	355.03 AB	43.4 B	55.69 B	13.15 B	132.49 B	25.97 B
	HCB	9,66	1.80	0.08	0.11	0.10	0.06	5.86	0.11	0.29	0.09	0.05	0.14	0.18
Families					Accipitridae (44)		Ardeidae (6)		Falconidae (6)		Phalacrocoracidae (4)		Strigidae (16)	
	δ13C	4,65	5.1	<0.01	-19.17 B		-24.00 A		-21.81 AB		-21.30 AB		-18.76 B	
	δ15N	4,65	9.19	<0.01	9.58 C		12.18 AB		8.39 C		13.60 A		10.35 BC	
	ΣPCBs	4,71	3.24	<0.01	1.55 B		4.10 AB		6.27 A		2.46 B		2.30 B	
	ΣDDTs	4,71	0.29	0.88	152.91		55.69		251.15		13.15		92.55	
Trophic guilds	HCB	4,71	0.37	0.82	1.41		0.09		0.17		0.05		0.15	
					Omnivorous (13)		Scavengers (21)		Carnivorous (32)		Piscivorous (10)			
	δ13C	3,66	9.95	<0.01	-20.49 AB		-17.53 C		-20.23 B		-23.08 A			
	δ15N	3,66	12.91	<0.01	9.01 B		10.4 B		9.26 B		12.65 A			
	ΣPCBs	3,72	4.45	<0.01	2.05 AB		0.28 B		3.44 A		3.45 A			
Habitat	ΣDDTs	3,72	2.92	0.03	12.97 B		3.80 B		295.86 A		38.67 B			
	HCB	3,72	1.09	0.35	0.11		0.09		1.94		0.08			
					Terrestrial (66)		Aquatic (10)							
	δ13C	1,68	12.66	<0.01	19.36		-23.08							
	δ15N	1,68	27.35	<0.01	9.61		12.66							
Feeding regime	ΣPCBs	1,74	1.19	0.27	2.17		3.45							
	ΣDDTs	1,74	0.56	0.45	147.22		38.68							
	HCB	1,74	0.37	0.54	1.00		0.08							
					Diurnal (60)		Nocturnal (16)							
	δ13C	1,68	1.5	0.22	-20.04		18.77							
	δ15N	1,68	0.44	0.50	9.93		10.35							
	ΣPCBs	1,74	0.01	0.91	2.34		2.31							
	ΣDDTs	1,74	0.24	0.62	143.70		92.55							
	HCB	1,74	0.58	0.44	1.07		0.16							

Table 2: Tests for strength of significance for PCBs and OCPs (ng g⁻¹ dry wt.) and stable isotope residues (‰) among different feather types of 6 individuals of black kites from Pakistan. Means are shown under each feather types whereas different alphabetic characters are used to illustrate significant differences through multiple comparison (Tukey-HSD) test.

Compounds/SIs	Test for significance			Mean concentrations and Significance levels (alphabets) among feather types (Tukey-HSD)			
	df	F	P	Body	Tail	Primary	Secondary
δ13C	3,20	0.56	0.65	-20.24	-21.55	-20.97	-22.32
δ15N	3,20	0.57	0.64	9.17	10.16	8.68	9.22
ΣPCBs	3,20	5.13	<0.01	4.21 B	2.57 AB	1.48 A	1.92 A
ΣDDTs	3,20	13.57	<0.01	27.57 B	12.17 A	7.68 A	9.97 A
HCB	3,20	0.24	0.86	0.13	0.14	0.14	0.16
TN	3,20	7.37	<0.01	0.05 B	0.03 AB	0.019 A	0.016 A

Table 3. Evaluation of factors governing OCs concentration differences based on Akaike's information criteria (AIC). Variables with lowest AIC (shown in bold) best explained the concentration of the respective compound. Significant differences (ANOVA) of compounds among/between tested variables are shown with bold p values.

	Species	Families	Trophic guilds	Food chain	Feeding regime	$\delta^{13}\text{C}$	$\delta^{15}\text{N}$
CB-105	AIC	116.41	114.42	116.93	115.58	115.87	118.00
	F, (p)	2.16 (0.08)	2.78 (0.04)	3.99 (0.02)	8.17 (0.00)	3.74 (0.06)	NC*
CB-118	AIC	204.85	203.05	198.85	202.92	213.28	210.78
	F, (p)	2.85 (0.01)	4.51 (0.00)	5.51 (0.00)	11.53 (0.00)	0.85 (0.35)	NC
CB-146	AIC	132.21	128.37	127.41	125.65	124.94	124.24
	F, (p)	0.88 (0.51)	0.82 (0.51)	0.60 (0.55)	0.00 (0.99)	1.56 (0.21)	NC
CB-153	AIC	226.48	240.69	223.07	245.41	251.33	249.54
	F, (p)	2.67 (0.01)	2.88 (0.02)	4.05 (0.01)	1.32 (0.25)	0.07 (0.79)	NC
CB-138	AIC	209.73	214.53	202.81	212.32	215.32	214.25
	F, (p)	1.56 (0.15)	1.38 (0.25)	2.66 (0.05)	1.10 (0.29)	0.36 (0.55)	NC
CB-187	AIC	114.26	108.47	108.06	106.07	107.79	107.69
	F, (p)	1.27 (0.29)	1.91 (0.14)	1.14 (0.32)	1.24 (0.27)	0.01 (0.89)	NC
CB-183	AIC	84.34	82.36	82.79	83.89	84.09	84.07
	F, (p)	2.07 (0.10)	2.85 (0.05)	1.34 (0.27)	0.72 (0.40)	0.21 (0.64)	NC
CB-128	AIC	154.67	149.27	148.62	147.42	145.59	146.78
	F, (p)	0.56 (0.77)	0.75 (0.56)	0.24 (0.78)	0.32 (0.57)	2.03 (0.15)	NC
CB-156	AIC	87.59	86.16	87.16	86.22	86.81	84.05
	F, (p)	3.41 (0.02)	2.45 (0.08)	0.76 (0.47)	0.01 (0.90)	0.75 (0.39)	NC
CB-180	AIC	218.61	215.15	213.21	216.7	216.9	214.6
	F, (p)	1.44 (0.18)	1.45 (0.22)	2.51 (0.06)	1.01 (0.31)	0.12 (0.72)	NC
CB-170	AIC	202.26	196.43	196.79	197.19	197.07	192.91
	F, (p)	1.40 (0.21)	1.58 (0.18)	1.99 (0.12)	0.72 (0.39)	0.00 (0.95)	NC
CB-199	AIC	66.40	64.51	63.31	63.31	66.57	66.66
	F, (p)	1.13 (0.36)	1.78 (0.19)	1.40 (0.25)	1.40 (0.25)	0.24 (0.62)	NC
CB-196/203	AIC	139.60	135.90	136.15	134.23	134.57	133.85
	F, (p)	1.21 (0.32)	1.38 (0.26)	1.55 (0.22)	0.84 (0.36)	0.21 (0.64)	NC
CB-194	AIC	149.63	149.22	150.14	153.12	153.52	150.73
	F, (p)	2.50 (0.03)	2.90 (0.04)	2.11 (0.10)	0.72 (0.40)	0.47 (0.49)	NC
ΣPCBs	AIC	217.74	273.26	238.12	279.91	283.48	282.77
	F, (p)	2.55 (0.01)	3.24 (0.01)	4.45 (0.00)	1.19 (0.27)	0.01 (0.91)	NC
TN	AIC	71.03	71.03	69.10	75.06	74.02	75.90
	F, (p)	4.91 (0.00)	4.91 (0.00)	7.59 (0.00)	4.40 (0.04)	7.88 (0.00)	NC
HCB	AIC	207.83	245.68	233.55	241.42	243.80	242.46
	F, (p)	1.80 (0.08)	0.37 (0.82)	1.09 (0.35)	0.37 (0.54)	0.58 (0.44)	NC
<i>p,p'</i> -DDE	AIC	272.43	315.55	277.44	317.85	317.24	318.66
	F, (p)	2.60 (0.01)	0.21 (0.92)	2.71 (0.05)	0.45 (0.50)	0.26 (0.60)	NC
<i>p,p'</i> -DDT	AIC	268.20	292.00	273.85	294.74	295.88	298.21
	F, (p)	2.67 (0.01)	1.51 (0.20)	3.15 (0.03)	1.10 (0.29)	0.10 (0.75)	NC
ΣDDTs	AIC	263.33	305.87	269.40	308.57	307.31	308.72
	F, (p)	2.61 (0.01)	0.29 (0.88)	2.92 (0.03)	0.56 (0.45)	0.24 (0.62)	NC

* NC (not calculated) are shown where ANOVA could not be quantified

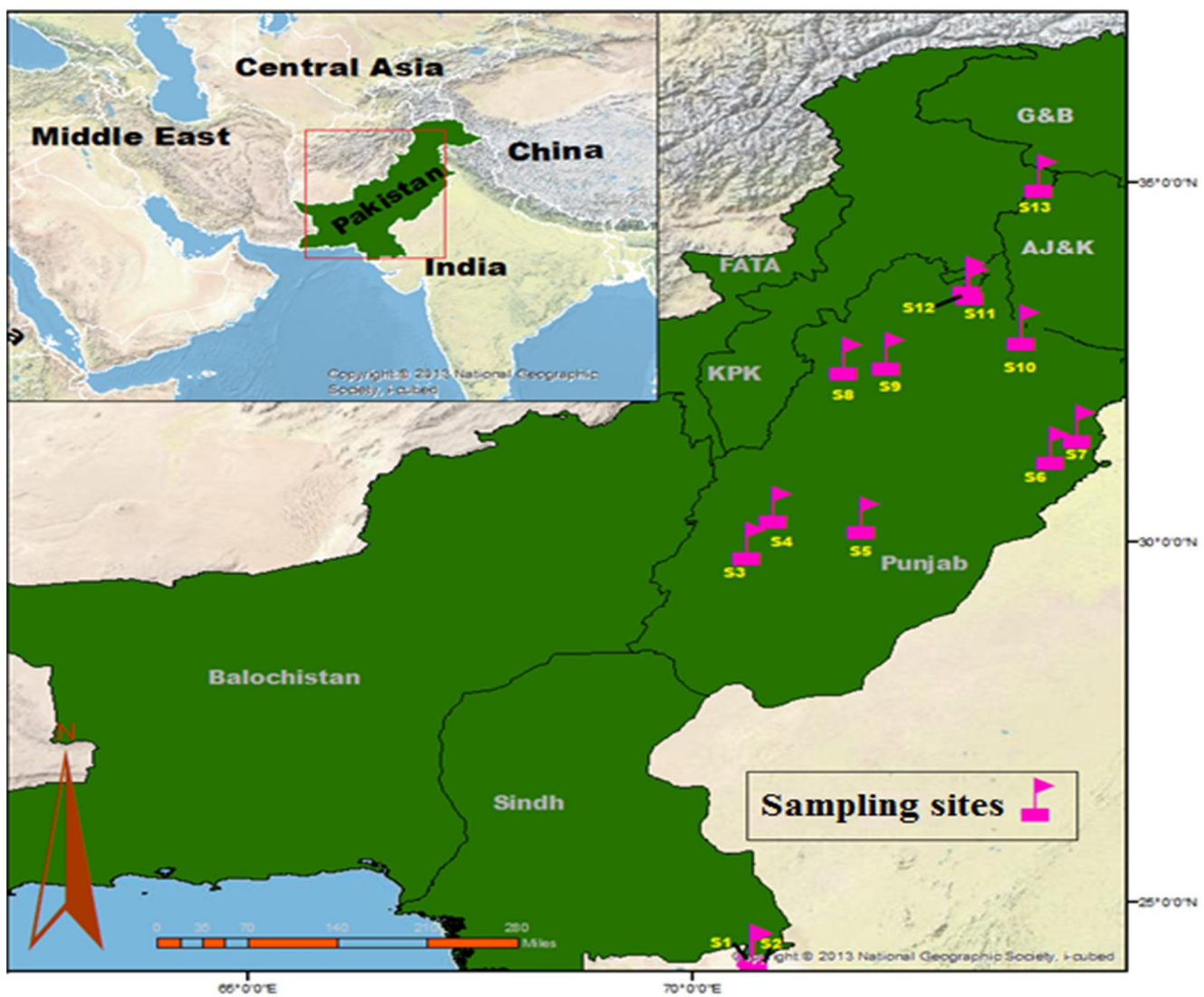


Figure 1. Map showing different sampling sites in Pakistan. The details of birds collected at different sampling sites are given in table S1 in the supplementary information.

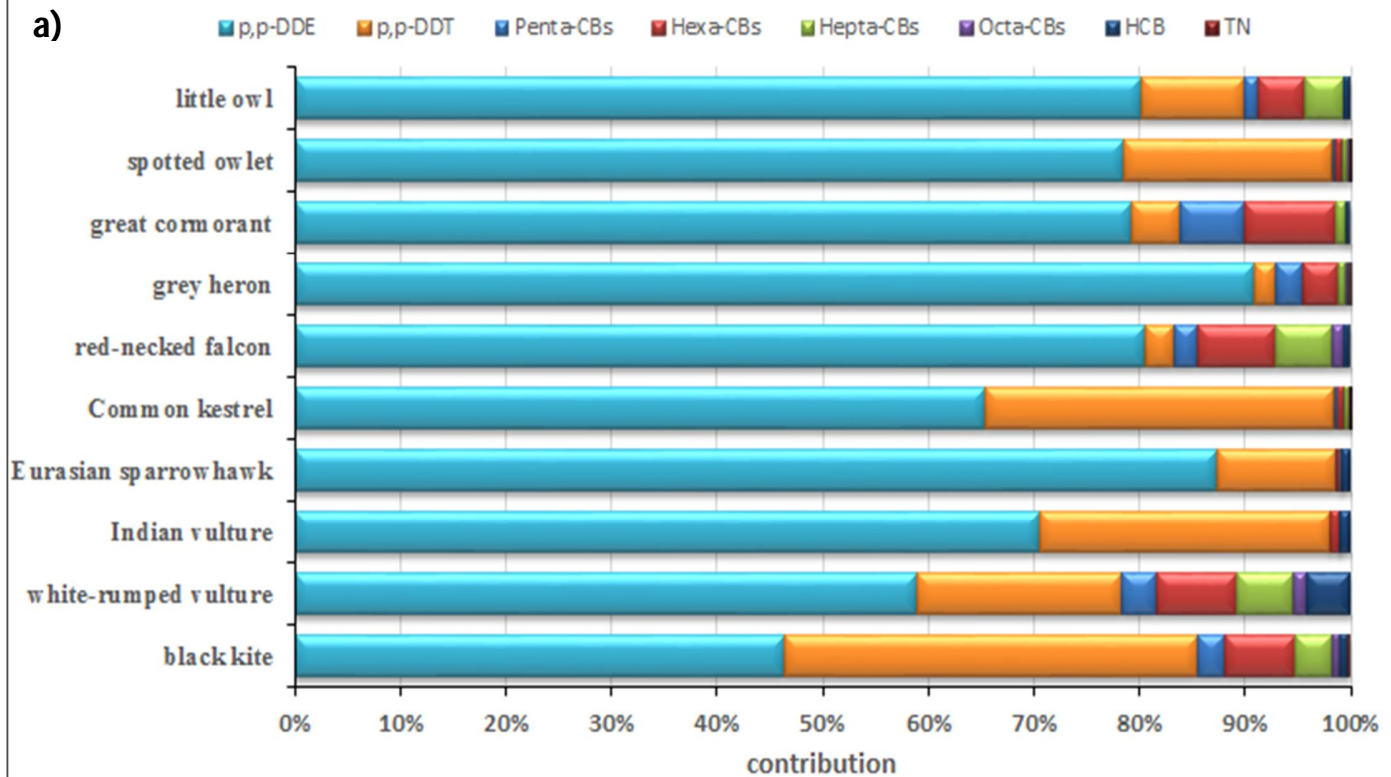
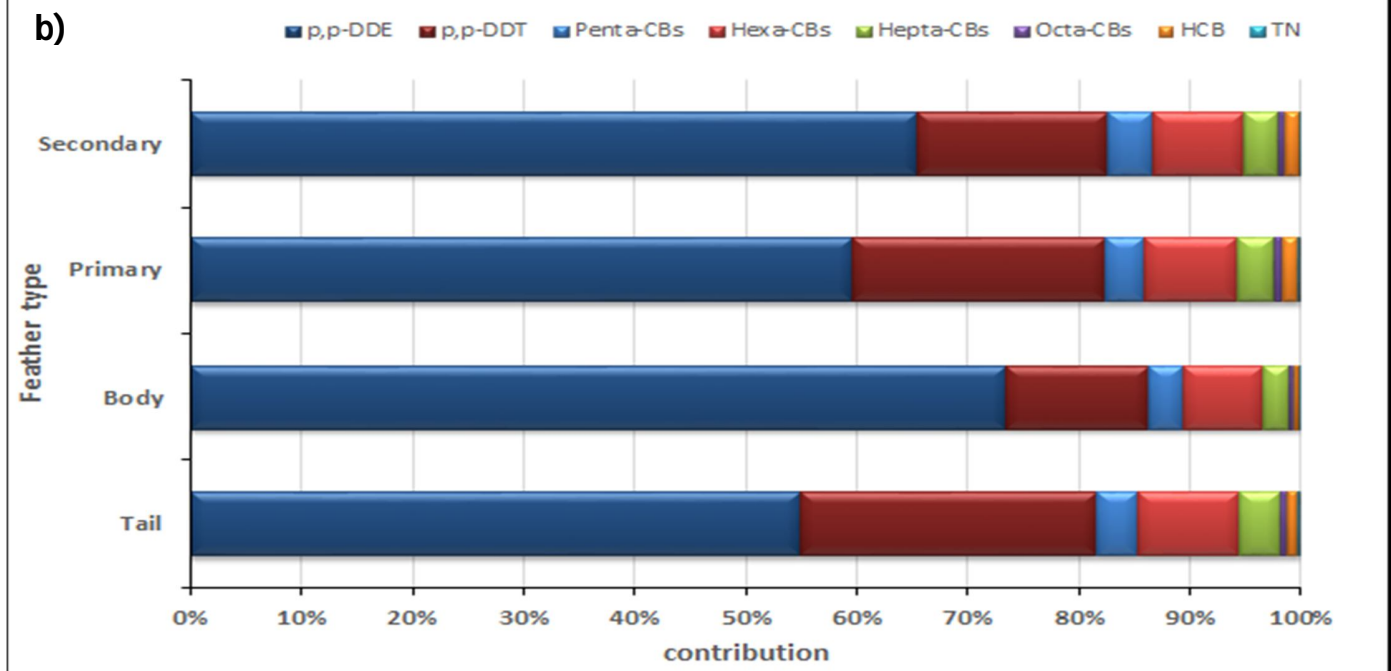
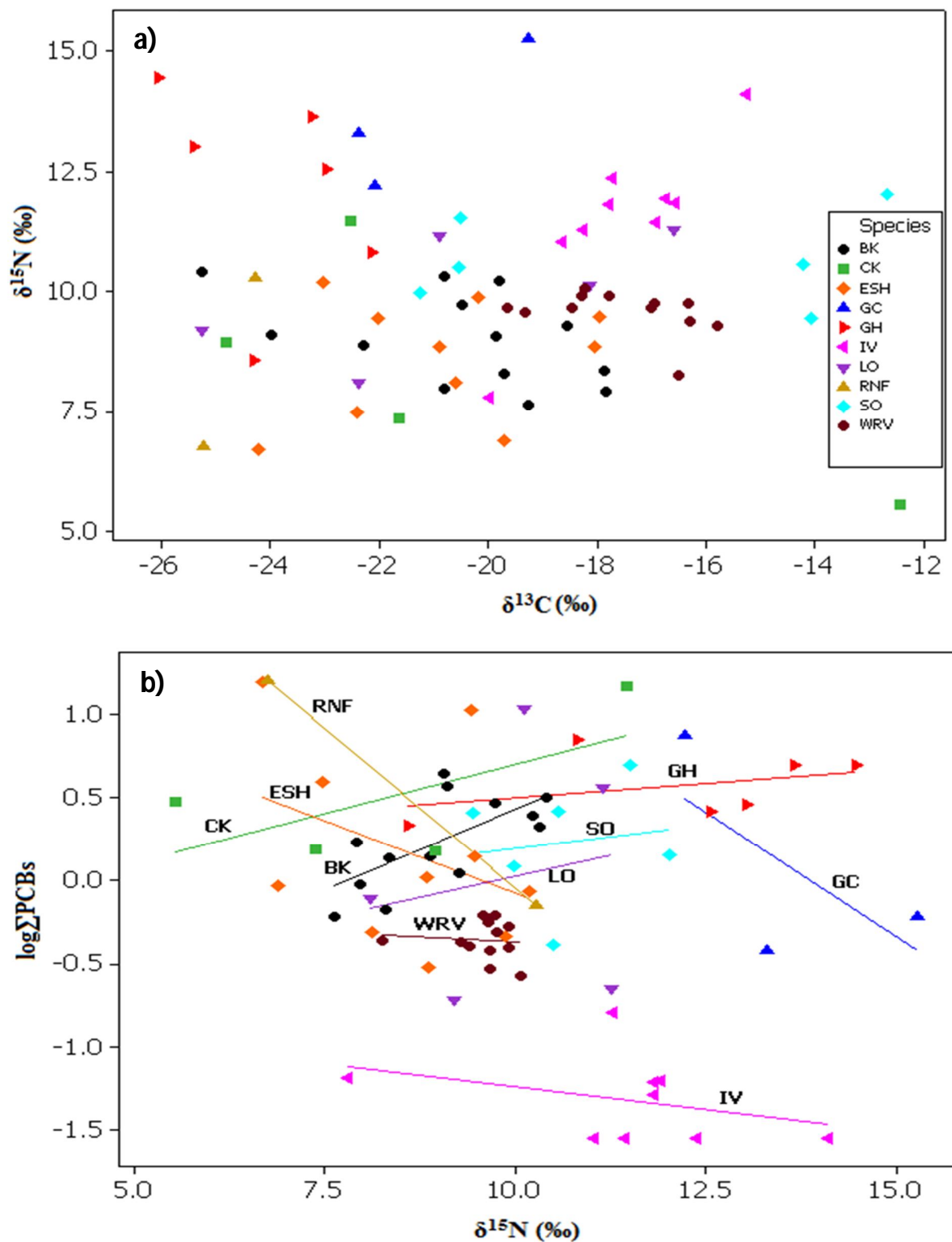
a)**b)**

Figure 2: Contribution profile (percentage) of PCBs and OCPs in (a) tail feathers of predatory birds and (b) different feather types of black kite from Pakistan.



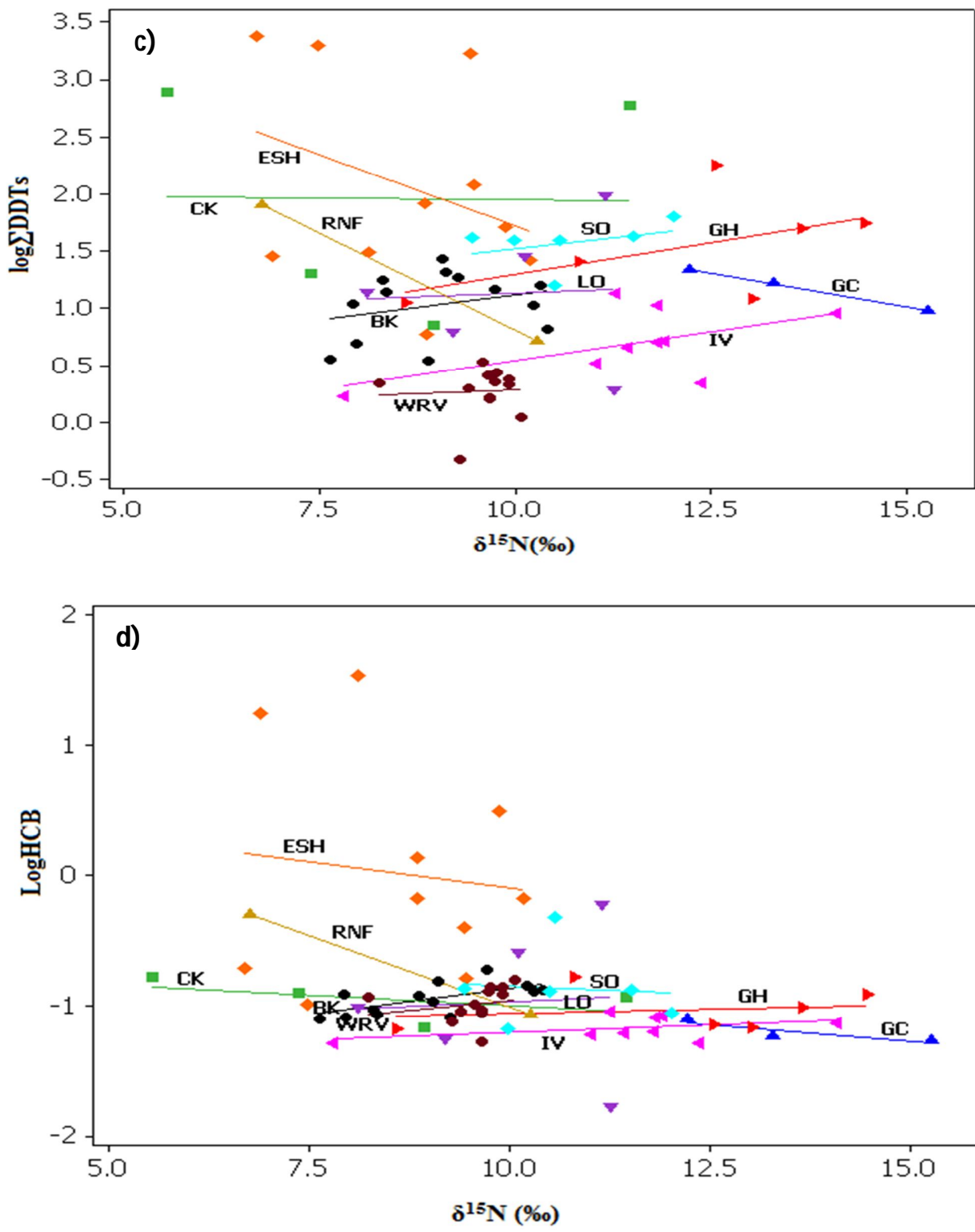


Figure 3. Scatter plots indicating interspecific differences in $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values (a), and species-specific regressions of \log_{10} -transformed concentrations of $\sum\text{PCBs}$ (b), $\sum\text{DDTs}$ (c) and HCB (d) on stable isotope values. Species abbreviations are as follows: BK=black kite, WRV=white-rumped vulture, IV=Indian vulture, ESH=Eurasian sparrowhawk, GH=grey heron, RNF=red-necked falcon, CK=common kestrel, GC=great cormorant; SO=spotted owlet, LO=little owl