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1	Accumulation of Anticoag	ulant Rodenticides in a Non-target Insectivore,
2	the European	hedgehog (Erinaceus europaeus)
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#### 42 Abstract

43 Studies on exposure of non-targets to anticoagulant rodenticides have largely focussed 44 on predatory birds and mammals; insectivores have rarely been studied. We investigated the 45 exposure of 120 European hedgehogs (Erinaceus europaeus) from throughout Britain to first-46 and second-generation anticoagulant rodenticides (FGARs and SGARs) using high 47 performance liquid chromatography coupled with fluorescence detection (HPLC) and liquid-48 chromatography mass spectrometry (LCMS). The proportion of hedgehogs with liver SGAR 49 concentrations detected by HPLC was 3-13% per compound, 23% overall. LCMS identified 50 much higher prevalence for difenacoum and bromadiolone, mainly because of greater ability 51 to detect low level contamination. The overall proportion of hedgehogs with LCMS-detected 52 residues was 57.5% (SGARs alone) and 66.7% (FGARs and SGARs combined); 27 (22.5%) 53 hedgehogs contained >1 rodenticide. Exposure of insectivores and predators to 54 anticoagulant rodenticides appears to be similar. The greater sensitivity of LCMS suggests 55 that hitherto exposure of non-targets is likely to have been under-estimated using HPLC 56 techniques.

57

58

59 Keywords: first- and second-generation anticoagulant rodenticide, insectivore,

60 brodifacoum, bromadiolone, difenacoum, flocoumafen, coumatetralyl, warfarin, non-

61 target

62

*Capsule:* Exposure of insectivorous hedgehogs to anticoagulant rodenticides in
 Britain is similar to predatory birds and mammals that specialise in eating small
 mammals, and hitherto exposure levels have been underestimated using HPLC
 techniques.

67

#### 68 **1. Introduction**

69

Globally, rodents destroy or spoil substantial amounts of food intended for 70 71 human or animal consumption (Singleton et al., 1999; Stenseth et al., 2003). Consequently, a range of methods is employed to reduce rodent density and 72 73 associated damage. This is most commonly done in developed countries using 74 anticoagulant rodenticides, vitamin K antagonists that prevent the synthesis of 75 functional prothombrin and related blood-clotting factors. Extensive use of first-76 generation anticoagulant rodenticides (FGARs) during the 1950s, however, led to the 77 evolution of genetic resistance in brown rats (Rattus norvegicus), with widespread 78 cross-resistance to other compounds (Cowan et al., 1995; Thijssen, 1995). As a 79 result, more potent second-generation anticoagulant rodenticides (SGARs) were 80 developed which have a greater affinity to binding sites, resulting in greater 81 accumulation, persistence and toxicity (Parmar et al., 1987; Huckle and Warburton, 82 1986). 83 Given their mode of action, both FGARs and SGARs are potentially harmful to all 84 vertebrates, and so users are expected to adopt measures that limit direct exposure 85 to non-target species. However, the degree to which these preventive measures are 86 adhered to, particularly by non-professionals, is unknown. For example, in Britain 87 some products are readily available to householders who may be less aware of the

risks of non-target poisoning and/or less likely to follow manufacturer's guidelines.

89 Non-target species may also be deliberately poisoned (Barnett et al., 2006).

Most studies investigating indirect exposure of non-target species to
anticoagulant rodenticides have focussed on the consumption of poisoned rodents by
predatory birds and mammals (Newton et al., 1990, 1999a; Berny et al., 1997;

93 McDonald et al., 1998; Shore et al., 1999, 2003a). However, invertebrates can be a 94 route of contamination for insectivorous vertebrates (Spurr and Drew, 1999) and, although exposure of insectivorous birds has been reported (Borst and Counotte, 95 96 2002; Dowding et al., 2006), exposure of insectivorous mammals has not been studied. Potential routes of uptake by invertebrates include: the consumption of 97 98 rodent faeces (Laas et al., 1985; Craddock, 2002; Eason et al., 2002); the 99 consumption of rodent carcasses; ingestion of soil-bound residues by e.g. 100 earthworms; and direct consumption of poison baits (Spurr and Drew, 1999; Dunlevy 101 et al., 2000; Craddock, 2002). Given that many ecological communities typically 102 contain larger numbers of insectivorous vertebrates relative to predators, the 103 contamination of invertebrates potentially poses the greater risk of non-target 104 poisoning in terms of species and individuals.

105 The European hedgehog (*Erinaceus europaeus*) is a medium-sized (0.8 - 1.2 kg) 106 insectivorous mammal distributed throughout Britain and across Western Europe 107 (Morris and Reeve, 2008). Hedgehogs are of particular interest in terms of exposure 108 to anticoagulant rodenticides, as they are reputed to have declined significantly in the 109 last few decades in Britain, and poisoning by industrial chemicals, including 110 rodenticides, may have been a contributory factor (Battersby and Tracking Mammals 111 Partnership, 2005). Our overall aim in this study was to investigate the scale and 112 severity of exposure of hedgehogs throughout Britain to some of the first-generation 113 (warfarin, coumatetralyl) and all of the second-generation (difenacoum, 114 bromadiolone, brodifacoum, flocoumafen) anticoagulant rodenticides that are 115 licensed for use in Britain; the indandione compounds were not determined using the 116 analytical techniques available to us in this study. The current study is the first to 117 assess anticoagulant rodenticide contamination in Britain of species at this trophic

118 level. Furthermore, we analysed tissue residues using both high performance liquid 119 chromatography coupled with fluorescence detection (hereafter HPLC) and liquid-120 chromatography mass spectrometry (LCMS). To date, characterisation of exposure 121 of non-target species has mostly used HPLC (for example, McDonald et al., 1998; 122 Shore et al., 2003a, 2006a; Walker et al., 2008) but LCMS is potentially a more 123 sensitive technique and, perhaps more importantly, enables compounds with similar 124 chemical structure to be differentiated with greater confidence since identification is 125 based upon mass rather than elution times. Our specific objectives were: to compare 126 and contrast the (i) frequency of occurrence and (ii) average residue magnitude of 127 FGARs and SGARs in hedgehogs by analysing liver concentrations using both HPLC 128 and LCMS techniques; (iii) to determine whether there were differences in levels of 129 contamination between males and females and between geographical regions; and 130 (iv) on the basis of these results, compare the extent of sub-lethal exposure of 131 hedgehogs in Britain with that of predatory birds and mammals, and assess whether 132 hedgehogs are at risk of acute toxicity from their exposure.

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#### 134 **2. Materials and Methods**

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During 2004-2006, 20 adult hedgehog carcasses were collected from wildlife rehabilitation hospitals from each of six (Scotland, Wales, Midlands and West, South-Western, South-Eastern, and Eastern) of the seven regions of Britain as defined by the Department for Environment, Food and Rural affairs when assessing rodenticide usage (Dawson et al., 2003); we were unable to obtain samples from the remaining region (Northern England). All 120 hedgehogs used in the study had either died following admission or were euthanased due to their injuries or illness.

Each carcass was weighed, sexed and stored at -20°C until dissection, when it was inspected for lesions, injuries or other abnormalities. These observations, along with information collected at admission, were used to determine the cause of death or reason for euthanasia. The whole liver, the primary organ for accumulation of rodenticides (Huckle and Warburton, 1986), was removed, weighed to two decimal places and stored in aluminium foil at -20°C until further analysis.

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150 2.1. Residue analyses

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Anticoagulant rodenticide residues were quantified using both HPLC and LCMS. The four main SGARs licensed for use in the UK (brodifacoum, bromadiolone, difenacoum and flocoumafen) were quantified using both techniques. The two most commonly applied FGARs in the UK, coumatetralyl and warfarin (Dawson and Garthwaite, 2004), were also analysed using LCMS only. All reagents were from Rathburn Chemical Co. Ltd, Walkerburn, Scotland and of a grade suitable for HPLC and LCMS analysis.

159 Extraction procedures for second-generation compounds followed Hunter (1985) 160 and Jones (1996). Samples were analysed in randomised batches of 15. Each liver 161 was defrosted at room temperature and a subsample of approximately 1g (mean wet 162 weight±SE=0.98±0.01g) ground to a homogenous paste using acid-washed furnace-163 cleaned sand and anhydrous sodium sulphate. A 30ml aliquot of extraction solvent (50:50 acetone/chloroform) was mixed thoroughly with the ground tissue, stood for 1 164 165 hour, then decanted and collected in a 100ml measuring cylinder through a funnel containing glass wool and anhydrous sodium sulphate. The ground tissue was 166 167 subsequently washed with 30ml aliguots of extraction solvent and washings were

added to the original extraction aliquot until a total volume of 100ml was collected.
The mixture was mixed by inversion and left to stand at room temperature for a
minimum of 12 hours. Subsequently the extract was divided into 50ml for analysis by
HPLC and 30ml was archived at 4°C in the dark for later analysis by LCMS. Both
samples were reduced to zero volume by evaporation of solvent in a fume cupboard
and the remaining 20ml was poured to waste.

174 The reduced extract was re-dissolved in 1ml of extract solvent and 4ml 175 acetronitrile and cleaned using an SPE Isolute C<sub>18</sub> (EC) 1g column (Internation 176 Sorbent Technology, Mid-Glamorgan, UK) connected to an SPE 500-mg NH<sub>2</sub> column 177 solvated with methanol. Columns were conditioned with 5ml methanol followed by 178 5ml acetronitrile. The re-dissolved extract was loaded onto the C<sub>18</sub> column and 179 washed with three 5ml aliquots of acetronitrile at <4ml/minute. The C<sub>18</sub> column was 180 then removed and 4ml ammoniacal methanol was washed through the NH<sub>2</sub> column 181 (flow <4ml/min). The resulting eluant was combined with 5ml methanol, reduced to 182 near dryness (to remove ammonia) and re-dissolved in 0.5ml methanol. Samples 183 were finally transferred to a chromatography vial via a 4mm syringe filter (Whatman 184 International Ltd, Kent, UK).

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186 2.2. High performance liquid chromatography

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High performance liquid chromatography (HP Series 1100, Agilent Technologies,
Bracknell, Berkshire, UK) was performed using a ODS Hypersil 200mm x 4.6mm
5µm column (Thermo electron corporation, Runcorn, Cheshire, UK) at 30°C. A 15µl
aliquot of cleaned-up extract was injected onto the column using 76:24
methanol:water (v/v) supplemented with 0.25% (v/v) acetic acid and 40mM

193 ammonium acetate, as the mobile phase pumped at 1.0ml/min isocratically. SGARs 194 were detected by fluorescence spectronomy (HP 1100 series fluorescence detector) 195 using three excitation wavelengths (313nm, 320nm and 350nm) simultaneously to 196 allow for correction of co-eluting peaks that interfered with the fluorescence of the 197 rodenticides. The emission for each excitation wavelength was measured at 380nm. 198 The excitation wavelength of 313nm gave the greatest emission signal at 380nm and 199 was thus used for quantification. The ratio the emission response elicited by the 200 320nm wavelength to that elicited by 313nm and the ratio elicited by 350nm to that 201 elicited by 313nm were both used to aid identification. A chromatographic peak was 202 identified as a specific SGAR if the ratios of the signals for each excitation 203 wavelength matched the ratios in the standards and if the absolute retention time of 204 the peak fell within the retention time window of the calibration standards.

205

### 206 2.3. Liquid chromatography mass spectrometry

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208 The archived extraction samples were cleaned using methods previously outlined and analysed by liquid-chromatography tandem mass spectrometry 209 210 conducted on a Zorbax Eclipse C18 3µm column (150 x 2mm). The analysis was 211 conducted using an isocratic mobile phase consisting of acetronitrile water containing 212 0.1% formic acid in the ratio 75:25 and at a flow rate of 200µl/min. The column was 213 maintained at 35°C; injection volume was set at 15µl. A Surveyor HPLC system 214 (Thermo Corporation, Hemel Hempstead, Hertfordshire, UK) was used to separate the sample and deliver it to an LCQ Duo, API ion trap mass spectrometer (Thermo 215 216 Corporation, Hemel Hempstead, Hertfordshire, UK).

Analyses were performed using electrospray ionisation in the negative mode. The capillary temperature was set at 270°C with an ionisation voltage of -36.0V. The sheath and auxiliary gasses used were helium and nitrogen maintained at 80psi and 20psi respectively. Sensitivity was increased using single ion monitoring, scanning for the molecular ion of each of the rodenticides. Selectivity and conformational analysis was undertaken using tandem mass spectrometry.

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224 2.4. Quality assurance

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226 Quantification of residues was carried out by comparison with rodenticide 227 standards (Chemservice, Greyhound Chromatography, Merseyside, UK) for all the 228 FGARS and SGARS that were quantified. For HPLC analysis, the linear calibration 229 range was 50-500ng/ml and the limit of detection (LoD) for peaks identified as 230 SGARs was determined from the linear regression of the multilevel calibration using 231 the equation  $Y=Y_0+3S_{v/x}$ , where Y is the LoD response,  $Y_0$  is the intercept and  $S_{v/x}$  is 232 the standard error of the regression line. The HPLC LoDs for bromadiolone, difenacoum, flocoumafen and brodifacoum based on the standards were 0.03, 0.01, 233 234 0.01 and 0.02µg respectively, which were analogous to previous analyses of polecat 235 (Mustela putorius) livers (Shore et al., 2003a). The LoDs for LCMS were obtained 236 using a similar method and were 0.002µg for all compounds. 237 For LCMS analysis, three concentrations (100, 50 and 10ng/ml) of the standards

for all the FGARs and SGARS were run alongside procedural blanks after every eight
samples to determine day-to-day quantitation. Calibration curves were obtained
using a range of concentrations (500, 400, 200, 100, 50, 20, 10, 5, 1 and 0.1ng/ml) of

these standards; the average areas of ten determinations of each standardconcentration were used to produce these curves.

For both HPLC and LCMS analysis, procedural blanks (reagents only) were 243 244 analysed alongside samples to detect possible contamination during sample 245 preparation. Chicken liver samples were each spiked with known concentrations of 246 each SGAR and were prepared, stored and analysed in the same way as unknown 247 samples to determine sample matrix recovery and percent recovery data. For HPLC 248 the mean (±SE%) recovery, determined from analyses of eight spiked samples, were 249 108±11.5%, 81.6±5.0%, 95.2±9.8% and 93.3±9.0% for difenacoum, bromadiolone, 250 flocoumafen and brodifacoum respectively. Corresponding figures for LCMS recovery 251 were 59.2±9.9%, 27.3±12.0%, 59.2±9.9% and 65.9±7.3%, determined from analyses 252 of four samples spiked for each SGAR. The apparently lower recovery associated 253 with LCMS than HPLC may have been an artefact reflecting poor stability of spiked 254 samples when archived. The bromadiolone and difenacoum concentrations in the 255 actual samples of hedgehog livers were not significantly lower when quantified by 256 LCMS than when measured by HPLC (see Results). Concentration data in tissue 257 samples were not recovery-corrected.

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259 2.5. Statistical analysis

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The numbers of samples with detectable and non-detectable rodenticide residues as determined by HPLC and LCMS were compared using Fisher's exact tests. Liver concentrations were not normally distributed and average residue concentrations are given as medians. Median liver concentrations in animals with detectable residues were compared using Mann-Whitney U tests. Wilcoxon matched

pairs tests were used to compare residue concentrations detected by the two
techniques within the same individual. Binary logistic regression was used to
examine the effect of region, batch number and sex on the presence/absence of
contamination; batch was included as a factor to confirm that batching samples for
analysis did not introduce any analytical biases. All analyses were conducting using
SPSS, Release 15.0 (Field, 2005).

272

273 **3. Results** 

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275 Reasons cited by wildlife hospitals for admission of the hedgehogs used in this 276 study were: injury (n=55); unknown (n=46); natural causes (n=18); and suspected 277 poisoning (n=1), although this diagnosis was not confirmed clinically or chemically. 278 No obvious signs of haemorrhage other than that associated with trauma were found 279 during *post-mortem* examinations (n=120).

280 Using HPLC, detectable liver concentrations of brodifacoum, bromadiolone, difenacoum and flocoumafen were found in four, 13, 16 and zero animals 281 282 respectively (Table 1); in total, SGARs were detected in 27 individuals (23% of the 283 animals analysed: Table 2). In contrast, SGARs were detected in 69 (57.5%) 284 hedgehogs when the analysis was conducted by LCMS (Table 2). FGARs (only 285 determined by LCMS) were detected in 27 (22.5%) animals (Table 2). Overall, 286 residues of at least one FGAR or SGAR were detected in two thirds of hedgehogs when samples were analysed by LCMS. Fifty-three (44%) individuals had liver 287 residues of one compound; 21 (18%), five (4%) and one (1%) animal contained 288 289 residues of two, three and four compounds respectively.

290 The greater frequency of detection of SGARs by LCMS than HPLC was largely 291 because more instances of difenacoum and bromadiolone contamination were 292 detected by LCMS (Table 2); the difference in frequency of detection between the analytical methods was significant for difenacoum (Fisher's Exact test, P<0.001) and 293 approached significance for bromadiolone (two-tailed Fisher's Exact test, P=0.10). 294 295 Much of this higher frequency of detection was due to the greater sensitivity of the 296 LCMS. Liver difenacoum and bromadiolone concentrations below 0.025µg/g wet 297 weight (ww) and 0.05µg/g ww. respectively, were not detected by HPLC, whereas 298 these concentrations comprised 25-50% of the LCMS detections for these 299 compounds (Fig. 1). Overall, detection of these low level difenacoum and 300 bromadiolone residues by LCMS accounted for an extra 30 hedgehogs (25% of the 301 sample) being identified as containing rodenticide.

302 The average magnitude of residues (Table 3), not just the frequency of 303 occurrence, also varied with analytical technique. When only hedgehogs with HPLC 304 and/or LCMS detectable residues were included in the statistical analysis, the 305 median liver bromadiolone concentration was lower when determined by LCMS than 306 by HPLC (Mann Whitney U test: U=61.0,  $n_1=23$ ,  $n_2=13$ , P<0.01; Fig. 2). This reflected 307 the presence of low-level bromadiolone concentrations (typically < 0.1  $\mu$ g/g ww; Fig. 308 1) that were detected by LCMS but not by HPLC (and so were not included in the 309 HPLC dataset of animals with detected residues). When the statistical analysis was 310 further restricted to a matched pair comparison of just animals with bromadiolone 311 residues detected by *both* analytical methods, there was no significant difference 312 between LCMS and HPLC measurements (Wilcoxon matched pairs test: n=10, Z=-313 0.663, P>0.05). This again suggested that differences between HPLC- and LCMSdetermined measurements were solely due to detection of low-level concentrations 314

by LCMS. However, this was not true for difenacoum. Median liver concentrations of difenacoum in animals with detectable residues did not differ with the method of determination (U=427.5,  $n_1$ =16,  $n_2$ =57, P>0.05), despite the presence of a relatively large number of low-level difenacoum residues in the LCMS sample (Fig. 1). This may reflect differential responses (involving enhancement or quenching of response) of the two techniques, as matched-pair analysis indicated that residues were higher in animals when measured by LCMS (n=9, Z=-2.429, P<0.05).

322 Analyses of potential differences in residue magnitude with sex and region were 323 based on LCMS data. Geographical region was not significantly associated with the 324 presence/absence of (i) FGARs (coumatetralyl and warfarin), (ii) bromadiolone and 325 difenacoum combined (the most commonly found SGARs), (iii) all four SGARs, or (iv) 326 all FGARs and SGARs combined (Table 4). Sex did, however, approach significance 327 in two of the four models (bromadiolone and difenacoum combined, P=0.052; all 328 SGARs, *P*=0.072; Table 4), with a greater frequency of occurrence of contamination 329 in males than females.

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### 331 **4. Discussion**

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The major proportion of hedgehog diet consists of invertebrates, particularly molluscs, beetles and earthworms (Wroot, 1984). Invertebrates have different bloodclotting mechanisms to vertebrates and so are less susceptible to anticoagulant rodenticides than birds and mammals (Shirer, 1992; Pain et al., 2000; Craddock, 2002; Johnston et al., 2005). However, ground-dwelling invertebrates can access and feed on rodenticides, including those placed in bait stations (Spurr and Drew, 1999; Dunlevy et al., 2000; Craddock, 2002), and retain ingested compound in their

340 bodies for four weeks or longer (Booth et al., 2001; Craddock, 2002). Additional 341 exposure of invertebrates to rodenticides may also arise through ingesting 342 contaminated soil (where baits have not been protected or have been displaced or 343 removed from bait stations), rodent food caches and rodent carcasses. Thus, 344 predation of contaminated invertebrates is likely to be a major pathway by which 345 hedgehogs are exposed to anticoagulant rodenticides. However, hedgehogs will 346 consume small mammal carcasses if they are available (Yalden, 1976) and may also 347 access spilt, cached or unprotected baits directly, and these may be alternative 348 secondary and primary exposure routes.

349 Whatever the route of exposure, it is clear from our results that contamination of 350 hedgehogs with anticoagulant rodenticides is commonplace. These compounds may 351 therefore similarly pose a risk to other species at the same trophic level, such as 352 insectivorous birds (Rammell et al., 1984; Empson and Miskelly, 1999; Robertson 353 and Colbourne, 2001). The frequencies with which we detected SGAR residues by 354 HPLC were towards the mid (brodifacoum, bromadiolone) or low (difenacoum) end of 355 the spectrum documented for predatory birds and mammals in Britain (Table 1), but 356 were comparable in some instances to prevalence rates in species considered to be 357 specialist predators of small mammals, such as the polecat (Shore et al., 2003a), 358 barn owl (Tyto alba) (Newton et al., 1999b) and tawny owl (Strix aluco) (Walker et al., 359 2008). Likewise, the magnitudes of residues were also broadly similar to those 360 measured in predatory birds and mammals in Britain (Table 3). Thus, hedgehogs in Britain appear to be at similar risk of exposure and effects from anticoagulant 361 362 rodenticides as non-target predatory birds and mammals.

Our data also suggest that exposure of hedgehogs is geographically widespread.
 The absence of any significant difference between the proportion of individuals with

365 residues and region indicates that the scale of exposure of hedgehogs does not vary 366 markedly across Britain, consistent with studies of polecats (Shore et al., 2003a), 367 even though the apparent use of rodenticides in arable regions varies geographically 368 (Dawson et al., 2003). In part, however, the likelihood of detecting correlated patterns 369 between prevalence rates in animals and regional patterns of use will be affected by 370 exactly which specific compounds are used. This is because compounds, and 371 particularly FGARs and SGARs, differ in their biological half-life and toxicity (Eason 372 et al., 2002). Furthermore, geographical variation in arable use of rodenticides is 373 unlikely to be of relevance to those animals that were from urban areas. There are no 374 published data for rodenticide use in urban areas in Britain and so it is not possible to 375 assess how urban use may relate to exposure of hedgehogs. Finally, our finding that 376 male hedgehogs tended to be more likely to accumulate rodenticides than females 377 may also have a spatial, albeit small scale, explanation. Males have a greater 378 ranging behaviour than females (Reeve, 1994) and this is likely to increase the 379 likelihood of individuals finding baits and contaminated forage.

380 The overall similarity between hedgehogs and specialist avian and mammalian 381 predators of small mammals was unexpected. This may simply indicate that 382 secondary exposure is more common than previously anticipated for food chains in 383 which small mammals are not a major component. However, this similarity may mask 384 other factors, such as differences in the likely exposure of non-target species in 385 urban and rural areas. We had no information on the exact location in which our 386 hedgehogs were found. Our reliance on analysing the carcasses of animals admitted 387 to wildlife hospitals may have biased the sample towards urban areas because their 388 relatively high human population density may mean that sick/injured hedgehogs are 389 more likely to be found. In contrast, most UK studies on secondary exposure in

390 predatory birds and mammals have analysed animals that are predominantly from 391 rural areas. It is not clear whether an urban-biased sample would tend to increase or 392 decrease the likelihood of detecting exposure. Rodenticides are widely used on 393 farms in rural Britain but are also commonly used throughout urban and suburban 394 landscapes by both professional practitioners and the general public. The density of 395 baits and contaminated prey relative to population numbers of non-target species in 396 rural and urban areas is completely unknown. Furthermore, it is possible that 397 hedgehogs may be particularly susceptible to exposure in urban areas where 398 untrained domestic users may be prone to unintentional misuse. Animals may also 399 be more likely to suffer traumatic injuries in human-dominated habitats through 400 collisions with motor vehicles or injuries arising from misadventure (Reeve and 401 Huijser, 1999). If such injuries occur independently of levels of rodenticide uptake, 402 such a sample would give a reliable indication of levels of sub-lethal contamination in 403 those areas, but if rodenticide uptake increases the likelihood of injury (Fournier-404 Chambrillon et al., 2004), then urban samples in particular may over-estimate 405 exposure rates. Comparison of exposure rates of hedgehogs or other species from 406 known urban and rural locations is merited.

The analysis of our sample of hedgehog tissues using LCMS as well as HPLC has shown that exposure, particularly low-level exposure, is markedly underestimated by HPLC. The proportion of hedgehogs exposed to SGARs increased by two- to three-fold when the analysis was conducted by LCMS. We postulate that current estimates of the exposure of predatory birds and mammals to SGARs have been similarly under-estimated where they have been determined using HPLC measurements.

414 Although exposure of hedgehogs to anticoagulants may be widespread, there is 415 no evidence from our study that this commonly causes lethal poisoning. The post 416 mortem examination of the animals in our study did not identify any instances of 417 haemorrhage that appeared consistent with rodenticide poisoning. Although there is 418 no precise liver concentration in hedgehogs or other species that is diagnostic of 419 lethal poisoning, SGAR residues in excess of 0.2µg/g ww are considered to be of 420 concern in barn owls (Newton et al., 1999a) and residues of >1µg/g ww are generally 421 considered to be very high. Irrespective of the measurement technique in our study, 422 the percentage of hedgehogs with summed SGAR residues above 0.2µg/g ww and 423 1µg/g ww was <11% and <5% respectively. The detection of liver residues exceeding 424 1µg/g ww suggests that lethal poisoning by rodenticides is likely to occur in some 425 hedgehogs, but the lack of haemorrhaging and relatively low magnitude of most 426 residues suggests that, for animals in our study, contamination with rodenticides was 427 generally not a contributory factor in their admission to wildlife hospitals. Overall, 428 however, poisoning of non-target wild animals by anticoagulant rodenticides is 429 difficult to monitor and studies such as ours may underestimate poisoning events 430 because animals with fatal doses may become lethargic some hours before death 431 and die in cryptic locations (Newton et al., 1999a). Furthermore, there is a general 432 lack of knowledge about whether sub-lethal exposure, as appears to be common in 433 hedgehogs, may be associated with any sub-lethal impacts or an increased 434 susceptibility to toxicity following repeated exposures.

435

## 436 **5. Conclusion**

This study has shown that the European hedgehog, an insectivorous species,
has similar rates of exposure (judged from the proportion of animals with HPLC-

439 detected liver concentrations and the size of those residues) to those of specialist 440 predators of small mammals. Given that hedgehogs only rarely eat rodents, these 441 results indicate that anticoagulant rodenticides are finding their way into ecosystems 442 via transfer pathways other than through consumption of contaminated rodents. 443 Furthermore, our data indicate that analysis of samples using LCMS can increase the 444 estimate of exposure by two- to three-fold, largely through the detection of low-level residues, and that the use of HPLC may have markedly under-estimated the true 445 446 scale of exposure of other non-target species to anticoagulant rodenticides. 447 448 **Acknowledgements** 449 450 We thank the RSPCA Westhatch, RSPCA Stapeley Grange, RSPCA East 451 Winch, RSPCA Mallydams Wood, The Gower Bird Hospital, St Tiggywinkles and 452 Hessilhead Wildlife Rescue Trust for supplying hedgehog carcasses and the 453 Dulverton Trust (C.V. Dowding and S. Harris) for financial support. 454 455 References 456 457 Barnett, E.A., Fletcher, M.R., Hunter, K., Sharp, E.A. 2006. Pesticide poisoning of 458 animals in 2005: investigations of suspected incidents in the United Kingdom.

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# 587 Figure legends

588

589 Fig. 1. Frequency distribution of bromadiolone and difenacoum liver concentrations in

590 hedgehogs detected by HPLC and LCMS.

591

592 Fig. 2. Median and interquartile ranges of liver concentrations of first- and second-

593 generation anticoagulant rodenticides in hedgehogs with detectable residues as

594 quantified using HPLC and LCMS. Sample sizes are given in Table 2.

595

597 Table 1

598 Percentage occurrence of the residues of the first-generation anticoagulant

599 rodenticide coumatetralyl (coum) and the second-generation anticoagulant

600 rodenticides brodifacoum (brod), bromadiolone (brom), difenacoum (difen) and

601 flocoumafen (floc) in the livers of predatory birds and mammals in British wildlife as

602 identified using high performance liquid chromatography. ND indicates residue not

603 detected; - indicates chemical was not investigated

604

Species	n	Coum	Brod	Brom	Difen	Floc	Total <sup>a</sup>	Ref <sup>b</sup>
Hedgehog ( <i>Erinaceus</i>	120	-	3.3	10.8	13.3	ND	22.5	1
europaeus)								
Polecat (Mustela putorius)	100	-	3.0	12.0	22.0	ND	36.0	2
Stoat (Mustela erminea)	40	15.0	2.5	6.7	-	-	22.5	3
Weasel (Mustela nivalis)	10	30.0	-	10.0	-	-	30.0	3
Red fox (Vulpes vulpes)	92	7.6	5.4	26.1	16.3	-	45.7	4
Barn owl ( <i>Tyto alba</i> )	717	-	3.9	11.0	16.7	1.1	26.1	5
Barn owl ( <i>Tyto alba</i> )	52	-	5.8	28.8	30.8	ND	42.3	6
Buzzard (Buteo buteo)	40	-	2.5	5.0	32.5	2.5	37.5	6
Tawny owl (S <i>trix aluco</i> )	172	-	4.7	11.6	5.8	ND	19.2	7
Red kite (Milvus milvus)	20	-	-	-	-	-	70.0	8
Kestrel (Falco tinnunculus)	36	-	-	-	-	-	67.0	8
Kestrel (Falco tinnunculus)	40	-	15.0	40.0	72.5	ND	84.6	9

605 606 607 <sup>a</sup> Total percentage of individuals positive for one or more chemicals. <sup>b</sup> Reference: 1 - present study; 2 - Shore et al. (2003a); 3- McDonald et al. (1998); 4 - Shore et al. (2003b); 5 - Newton et al. (1999b); 6 - Shore et al. (2006a);

7 - Walker et al. (2008), 8 - Shore et al. (2000); 9 - Shore et al. (2006b).

- Table 2
- Number and percentage (out of sample of 120) of hedgehogs with first- (FGAR) and
- second-generation anticoagulant rodenticides (SGAR) detected using high
- performance liquid chromatography (HPLC) and liquid-chromatography mass
- spectrometry (LCMS)

	Hedgehogs with residues detected by						
	HP	LC	LCMS				
	%	n	%	n			
Coumatetralyl (FGAR)			14.2	17			
Warfarin (FGAR)			8.3	10			
Brodifacoum (SGAR)	3.3	4	5.0	6			
Bromadiolone (SGAR)	10.8	13	19.2	23			
Difenacoum (SGAR)	13.3	16	47.5	57			
Flocoumafen (SGAR)	0	0	0.8	1			
Total SGARs only	22.5	27	57.5	69			
Total FGARs and SGARs	-		66.7	80			

Coumatetralyl and warfarin only determined using LCMS

- 617 Table 3
- 618 Mean  $\pm$  SE (*n*) concentration ( $\mu$ g/g ww) of second-generation anticoagulant
- 619 rodenticide residues in British wildlife identified using high performance liquid
- 620 chromatography. Figures are the concentrations only for those animals where
- 621 residue was detected
- 622

Species <sup>a</sup>	n	Brodifacoum	Bromadiolone	Difenacoum	Ref <sup>b</sup>
Hedgehog	120	0.05±<0.01 (4)	0.59±0.24 (13)	0.10±0.03 (16)	1
Polecat	50	0.06±0.01 (3)	0.12±0.03 (12)	0.30±0.07 (22)	2
Stoat	9	0.12	0.20±0.10 (3)	-	3
Weasel	3	-	0.25 (1)	-	3
Barn owl	88	0.02±<0.01 (9)	0.09±0.02 (23)	0.03±0.01 (35)	4
Kestrel	40	0.08±0.03 (6)	0.18±0.04 (16)	0.08±0.02 (29)	4
Red kite	8	0.35±0.22 (5)	0.11±0.01 (3)	0.20 (1)	5
Tawny owl	172	0.25±0.14 (8)	0.21±0.05 (20)	0.06±0.02 (10)	6

<sup>a</sup> For Latin names, see Table 1; <sup>b</sup> reference: 1 present study; 2 - Shore et al. (2003a); 3 - McDonald et al. (1998);
4 - Shore et al. (2006b); 5 - Carter and Burn (2000), 6 - RF Shore (unpubl. data).

625

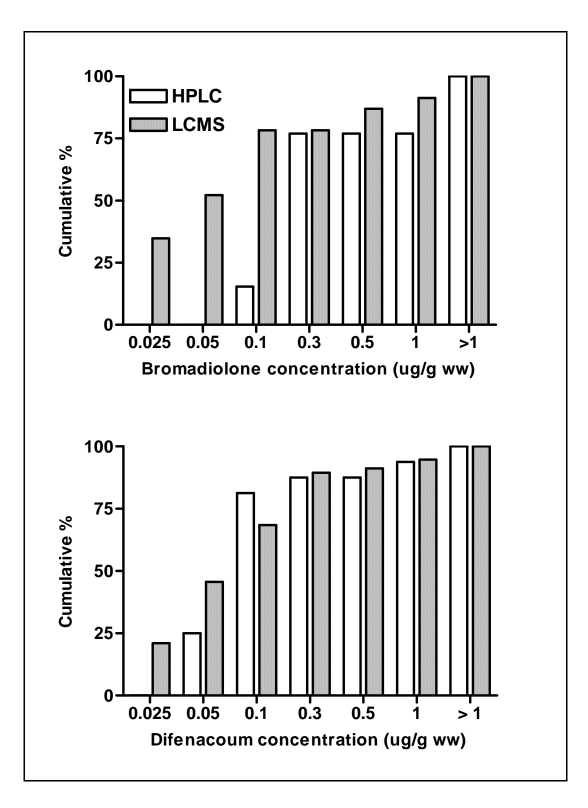
- 627 Table 4
- 628 Binary logistic regression models examining the relationship between region, batch
- and sex and the presence/absence of (a) first-generation anticoagulant rodenticides
- 630 (coumatetralyl and warfarin), (b) the second-generation anticoagulant rodenticides
- 631 bromadiolone and difenacoum, (c) all second-generation anticoagulant rodenticides
- 632 (brodifacoum, bromadiolone, difenacoum and flocoumafen) and (d) all first- and
- 633 second-generation anticoagulant rodenticides in hedgehogs from across Britain
- 634 (*n*=120)
- 635

Model	Variable	В	S.E.	Wald	d.f.	Р
а	Batch			3.426	8	0.905
	Region			6.885	5	0.226
	Sex	0.154	0.506	0.092	1	0.762
	Constant	-1.997	1.255	2.531	1	0.112
b	Batch			5.494	8	0.704
	Region			4.353	5	0.500
	Sex	0.831	0.428	3.775	1	0.052
	Constant	-1.130	0.951	1.412	1	0.235
С	Batch			2.821	8	0.945
	Region			3.474	5	0.627
	Sex	0.760	0.423	3.233	1	0.072
	Constant	-0.561	0.931	0.363	1	0.547
d	Batch			1.182	8	0.997
	Region			3.602	5	0.608
	Sex	0.730	0.450	2.635	1	0.105
	Constant	0.388	0.985	0.155	1	0.693

636 Male:female ratio for hedgehogs from different regions were: South-Eastern 12:8; South-Western

637 15:5; Eastern 11:9; Midlands and West 8:12; Wales 12:8; Scotland 7:13

639 Fig. 1



642 Fig. 2



