NSAIDs detected in Iberian avian scavengers and carrion after diclofenac registration for veterinary use in Spain

Marta Herrero-Villar, Roser Velarde, Pablo R. Camarero, Mark A. Taggart, Victor Bandeira, Carlos Fonseca, Ignasi Marco, Rafael Mateo

PII: S0269-7491(20)32452-0

DOI: https://doi.org/10.1016/j.envpol.2020.115157

Reference: ENPO 115157

To appear in: Environmental Pollution

Received Date: 3 April 2020

Revised Date: 17 June 2020

Accepted Date: 30 June 2020

Please cite this article as: Herrero-Villar, M., Velarde, R., Camarero, P.R., Taggart, M.A., Bandeira, V., Fonseca, C., Marco, I., Mateo, R., NSAIDs detected in Iberian avian scavengers and carrion after diclofenac registration for veterinary use in Spain, *Environmental Pollution* (2020), doi: https://doi.org/10.1016/j.envpol.2020.115157.

This is a PDF file of an article that has undergone enhancements after acceptance, such as the addition of a cover page and metadata, and formatting for readability, but it is not yet the definitive version of record. This version will undergo additional copyediting, typesetting and review before it is published in its final form, but we are providing this version to give early visibility of the article. Please note that, during the production process, errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

© 2020 Published by Elsevier Ltd.



CRediT author statement

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. † These authors contributed equally to this work.

Marta Herrero-Villar: Validation, Formal analysis, Investigation, Writing - Original Draft, Writing - Review & Editing, Roser Velarde: Investigation, Pablo R. Camarero: Validation, Investigation, Mark A. Taggart: Writing - Review & Editing, Victor Bandeira: Resources, Funding acquisition, Carlos Fonseca: Resources, Funding acquisition, Ignasi Marco: Resources, Supervision, Project administration, Funding acquisition, Rafael Mateo: Conceptualization, Methodology, Resources, Writing-Original Draft, Writing - Review & Editing, Supervision, Project administration, Funding acquisition.



Jonugal

1	NSAIDs detected in Iberian Avian Scavengers and Carrion after Diclofenac
2	Registration for Veterinary use in Spain
3	Marta Herrero-Villar ^{1*} , Roser Velarde ² , Pablo R. Camarero ¹ , Mark A. Taggart ³ , Victor
4	Bandeira ⁴ , Carlos Fonseca ⁴ , Ignasi Marco ^{2,5†} , Rafael Mateo ^{1†}
5	
6	¹ Instituto de Investigación en Recursos Cinegéticos (IREC), CSIC-UCLM-JCCM, 13005
7	Ciudad Real, Spain.
8	² Wildlife Ecology & Health group (WE&H) and Servei d'Ecopatologia de Fauna Salvatge
9	(SEFaS), Departament de Medicina i Cirurgia Animals, Universitat Autònoma de Barcelona,
10	08193 Bellaterra, Spain.
11	³ Environmental Research Institute, University of the Highlands and Islands, Thurso,
12	Scotland, KW14 7JD, UK.
13	⁴ Department of Biology & Centre for Environmental and Marine Studies (CESAM),
14	University of Aveiro, Campus Universitário de Santiago, 3810-193 Aveiro, Portugal.
15	⁵ Wildlife Conservation Medicine Research Group (WildCoM), Departament de Medicina i
16	Cirurgia Animals, Universitat Autònoma de Barcelona, 08193 Bellaterra, Spain.
17	† These authors contributed equally to this work.
18	
19	Corresponding author
20	Marta Herrero Villar*, PhD Student. From the Wildlife Toxicology research group in the
21	Institute for Game and Wildlife Research (IREC), CSIC-UCLM-JCCM.
22	Ronda de Toledo 12,
23	13005, Ciudad Real, Spain.
24	Phone: +34 926 29545 ext. 3353
25	Email contact: Marta.Herrero@uclm.es

26 Abstract

Despite the now well recognised impact of diclofenac on vultures across the Indian 27 subcontinent, this non-steroidal anti-inflammatory drug (NSAID) was registered in 2013 for 28 29 livestock treatment in Spain, Europe's main vulture stronghold. We assessed the risk of exposure to diclofenac and nine other NSAIDs in avian scavengers in the Iberian Peninsula 30 (Spain and Portugal) after the onset of diclofenac commercialization. We sampled 228 31 livestock carcasses from vulture feeding sites, primarily pig (n=156) and sheep (n=45). We 32 also sampled tissues of 389 avian scavenger carcasses (306 Eurasian griffon vultures, 15 33 cinereous vultures, 11 Egyptian vultures, 12 bearded vultures and 45 other facultative 34 scavengers). Samples were analysed by liquid chromatography with mass spectrometry 35 (LCMS). Seven livestock carcasses (3.07%) contained NSAID residues: flunixin (1.75%), 36 37 ketoprofen, diclofenac and meloxicam (0.44% each). NSAID residues were only detected in sheep (4.44%) and pig (3.21%) carcasses. Fourteen dead avian scavengers (3.60%) had 38 NSAID residues in kidney and liver, specifically flunixin (1.03%) and meloxicam (2.57%). 39 40 Flunixin was associated with visceral gout and/or kidney damage in three (0.98%) dead Eurasian griffons. To date, diclofenac poisoning has not been observed in Spain and Portugal, 41 however, flunixin would appear to pose an immediate and clear risk. This work supports the 42 need for well managed carrion disposal, alongside appropriate risk labelling on veterinary 43 NSAIDs and other pharmaceuticals potentially toxic to avian scavengers. 44

45

46 Capsule: NSAIDs were present in livestock carrion and wild avian scavengers in Spain, but
47 only flunixin was associated with visceral gout and/or kidney damage in three (0.98%)
48 Eurasian griffons.

49

50 **Keywords**: Veterinary pharmaceutical; Europe; vultures; flunixin; poisoning.

51

52 **1. Introduction**

During the late-1990s and early-2000s, South Asian Gyps vulture populations 53 collapsed (by up to 99.9%), almost leading to their extinction (Prakash et al., 2003). 54 Demographic studies showed alarming rates of adult mortality in India, Pakistan and Nepal of 55 white-rumped vulture (Gyps bengalensis), Indian vulture (Gyps indicus) and slender-billed 56 vulture (Gyps tenuirostris) (Gilbert et al., 2006; Prakash et al., 2012). Based on consistent 57 pathological findings in dead vultures (visceral gout with tubular nephrosis), dietary exposure 58 to a toxicant was considered to be a plausible cause. Oaks et al. (2004) hypothesized that 59 Asian vultures were being intoxicated with veterinary products used in livestock treatment 60 and further investigation identified diclofenac as the potential driver. This was corroborated 61 by a clear association between the presence of diclofenac residues in kidneys of dead vultures 62 and visceral gout (Oaks et al., 2004; Oaks & Watson, 2011). To confirm diclofenac toxicity, 63 experimental dosage studies were also conducted with non-releasable captive Gyps vultures, 64 and these established a lethal dose (LD₅₀) between 0.098-0.225 mg kg⁻¹ body weight (bw), 65 with death occurring after ingesting carrion containing 0.007-0.94 mg kg⁻¹ diclofenac (Oaks 66 et al., 2004; Swan et al., 2006). Based on this data, Green et al. (2004) further estimated that 67 just 0.13-0.75% of carcasses available to vultures in South Asia would need to contain a 68 lethal dose of diclofenac to cause rapid population declines. In fact, carcass monitoring 69 confirmed that 11.1-13.9% of carcasses available in India had detectable diclofenac residues, 70 with levels in livers ranging between 0.01 and 10.1 mg kg⁻¹ (Taggart et al., 2007, 2007b, 71 2009). Having identified this widespread problem in South Asia, Governments banned the 72 manufacture and importation of diclofenac for veterinary use in India, Pakistan and Nepal in 73 2006 (and in 2010 in Bangladesh), which by 2011 decreased diclofenac positive carcasses by 74 ~50% in India (Chaudhry et al., 2012; Prakash et al., 2012; Khan et al., 2013; Cuthbert et al., 75

76 2016). However, diclofenac was still being detected in carrion, and at lethal levels (Cuthbert77 et al., 2011, 2011b).

Identifying vulture-safe NSAIDs, that can also serve as alternatives to diclofenac for 78 79 veterinary use, is key to successfully reducing risks to vultures. To date, meloxicam is the only available NSAID that has clearly been identified as vulture-safe (Swarup et al., 2007; 80 Naidoo et al., 2008; Cuthbert et al., 2014). Meloxicam has been detected (below the limit of 81 quantification) in two eggs from captive-reared bearded vultures (Gypaetus barbatus) but no 82 adverse effects on reproduction have been observed (Zorrilla et al., 2018). Several other 83 NSAIDs have also undergone robust vulture safety testing, but all have shown varying 84 degrees of toxicity in *Gyps* vultures; these include ketoprofen (Naidoo et al., 2010, 2010b), 85 carprofen (Cuthbert et al., 2007; Fourie et al., 2015), flunixin (Fourie et al., 2015; Zorrilla et 86 al., 2015), nimesulide (Cuthbert et al., 2016) and aceclofenac, which is known to metabolize 87 to diclofenac in vivo when administered to cattle (Galligan et al., 2016). 88

This risk scenario is however not just limited to South Asia. Diclofenac is currently 89 90 authorized as a veterinary drug in certain African and European countries, and its toxicity has been demonstrated in native vulture species of these regions (Naidoo et al., 2009). In Europe, 91 diclofenac was first registered for veterinary use in Spain, Italy, Estonia, Latvia and the 92 Czech Republic in 2013. The Iberian Peninsula (Spain and Portugal) is particularly important 93 in this context, as it hosts ~95% of all European vultures. Spain holds around 31000 breeding 94 pairs of Eurasian griffon vultures (*Gyps fulvus*), >2500 pairs of cinereous vultures (*Aegypius*) 95 monachus), >1450 pairs of Egyptian vultures (Neophron percnopterus) and 125 pairs of 96 bearded vultures (Gypaetus barbatus) (Del Moral, 2009, 2017; Margalida et al., 2014; Del 97 Moral & Molina, 2018). Portugal hosts 500-1000 breeding pairs of Eurasian griffons, 5-8 98 pairs of cinereous vultures and 50-100 pairs of Egyptian vultures (BirdLife International, 99 2015). Due to concerns regarding risks to these important populations, the Spanish Agency of 100

Medicines and the Ministry of Agriculture and Environment performed a risk assessment (AEMPS & MAGRAMA, 2014) and estimated that residues of diclofenac in carrion may cause 15-39 deaths of Eurasian griffons in Spain per year, which was in stark contrast to another study that calculated 715-6389 deaths to be plausible (Green et al., 2016).

Eleven NSAIDs are used for livestock treatment in Spain (CIMAVET, 2020), and 105 most of these are now considered potentially toxic to *Gyps* vultures (Oaks et al., 2004; 106 Cuthbert et al., 2007; Naidoo et al., 2010, 2010b; Fourie et al., 2015; Zorrilla et al., 2015). 107 One of them, flunixin, has already been linked to wild vulture mortality in Spain (Zorrilla et 108 al., 2015). However, until now, there have been no published monitoring data regarding 109 NSAID residues in livestock carcasses and vultures in the Iberian Peninsula. The objectives 110 of this work have therefore been [1] to quantify the presence of diclofenac and other NSAIDs 111 in livestock carrion disposed of at supplementary feeding stations for vultures in Spain, [2] to 112 assess NSAID residues (and potential poisoning) in avian scavengers found dead or moribund 113 in Spain and Portugal between 2013 and 2019, and [3] to discuss the potential impact that 114 NSAIDs may have on Iberian populations of avian scavengers. 115

116

117 2. Material and methods

118

119 2.1. Carrion sampling

Carrion sampling was performed by trained environmental technicians and agents at supplementary feeding stations in Castile and Leon, Valencia, Aragon, La Rioja, Asturias, Galicia, Navarra and Extremadura (Fig. 1). Sampled carrion were 156 pig, 45 sheep, 4 goat, 4 horse, 6 cow and 13 unknown (the species were not recorded) carcasses. During the first sampling period (in 2016; performed under project TEC0004566), muscle, liver and kidney were collected from 125 pig carcasses. In the second period (2018-2019; performed under

- project D16ZO-046-MAF-AvianScaven), liver and kidney were collected from the remaining
 carcasses listed. Samples were collected into plastic zip-lock bags, frozen at -20 °C and sent
 to the Institute for Game and Wildlife Research (IREC) for NSAID analysis.
- 129

130 2.2. Avian scavenger sampling

We tested 389 avian scavengers from forensic cases admitted to wildlife rehabilitation 131 centres in Spain and Portugal (Fig. 1). This included four vulture species corresponding to 132 Eurasian griffons (n=306: 290 from Spain and 16 from Portugal), cinereous vultures (n=15: 133 13 from Spain and 2 from Portugal), bearded vultures (n=12 from Spain) and Egyptian 134 vultures (n=11; 10 from Spain and 1 from Portugal). We also included 45 facultative avian 135 scavengers from Spain corresponding to common buzzards (Buteo buteo, n=26), red kites 136 137 (Milvus milvus, n=15), black kites (Milvus migrans, n=3) and booted eagle (Hieraaetus pennatus, n=1). Although booted eagle does not show scavenging habits, this bird was 138 included in the analysis because it showed lesions of visceral gout. 139

The tissues collected and analysed at IREC were liver (n=384) and kidney (n=284). 140 Sample collection was performed during two mortality monitoring programs. Since 2004, 141 IREC has offered toxicological analyses of wildlife incidents to Spanish public 142 administrations. In cases of suspected poisoning, during post-mortem examinations, 143 veterinary staff from wildlife rehabilitation centres in Spain submitted liver and other samples 144 (or whole carcasses) for toxicological analysis. Between 2013 and 2019, liver samples from 145 avian scavengers (n=149) were collected through this analytical service and were tested for 146 NSAID presence, among other toxic substances. Since 2017, liver and kidney sampling of 147 148 avian scavengers (n=240) has also been carried out specifically for NSAID detection. These samples were taken during necropsies at wildlife rehabilitation centres (n=157), at the 149 Universitat Autònoma de Barcelona (n=56) and at IREC (n=8) in Spain, and at two 150

151 Portuguese wildlife rehabilitation centres (Centro de Estudos e Recuperação de Animais Selvagens (CERAS) and Centro de Recuperação e Investigação de Animais Selvagens 152 (RIAS) (n=19), whose samples were received by the University of Aveiro in Portugal. All 153 samples were collected into polypropylene tubes or zip-lock bags and held frozen at -20 °C 154 until analysis. Additionally, one Eurasian griffon which died in a wildlife rehabilitation centre 155 in 2015 after iatrogenic poisoning with flunixin was analysed, but this case was not included 156 in our statistical analysis and more details of the circumstances of this poisoning were given 157 in Soler et al. (2016). The flunixin tissue levels detected in this vulture were used as a 158 159 reference for confirmed poisoning.

160

161 2.3. NSAID Analysis

The veterinary NSAIDs registered in Spain (with number of commercial formulations 162 for use in livestock) are ketoprofen (23), meloxicam (19), flunixin meglumine (14), 163 acetylsalicylic acid (8), phenylbutazone (6), sodium salicylate (4), carprofen (4), metamizole 164 (4), tolfenamic acid (3), diclofenac (2) and suxibuzone (1) (CIMAVET, 2020). The selected 165 compounds tested were those initially covered by a method developed by Taggart et al. 166 (2009) to monitor carrion from India, which included carprofen, diclofenac, flunixin, 167 indomethacin, ketoprofen, meloxicam and naproxen. We also added tolfenamic acid, 168 suxibuzone and phenylbutazone to include most of the NSAIDs registered in Spain for 169 veterinary use in livestock according to the AEMPS database (CIMAVET, 2020). 170 Indomethacin and naproxen were not registered in Spain but were also monitored as they 171 were already covered by the method used. The only NSAIDs registered in Spain for 172 173 veterinary use in livestock not included in this work were metamizole, acetylsalicylic acid and sodium salicylate (CIMAVET, 2020). 174

Analytical standards (Table S1) were acquired from Sigma-Aldrich and Supelco: carprofen (33975 Supelco), diclofenac disodium salt (D6899 Sigma-Aldrich), flunixin meglumine (F0429 Sigma-Aldrich), indomethacin (I8280 Sigma-Aldrich), ketoprofen (34016 Supelco), meloxicam hydrated sodium salt (M3935 Sigma-Aldrich), naproxen (N8280), tolfenamic acid (T0535 Sigma-Aldrich), suxibuzone (S2400000 Sigma-Aldrich) and phenylbutazone (P8386 Sigma-Aldrich). Flunixin-d3 (34083 Supelco) was used as an internal standard.

The tissue extraction method was based on Taggart et al. (2009), with some 182 modifications. First, 0.5 g of tissue was weighed into a polypropylene tube to which 2 mL of 183 acetonitrile and 80 μ L of flunixin-d3 (at 80 ng μ L⁻¹ in acetonitrile) were added. This mix was 184 homogenized using an IKA-T8 homogenizer for 1 min. Between each sample, the 185 homogenizer was thoroughly cleaned using Extran MA 01 solution (Merck), Milli-Q water 186 and acetonitrile to avoid cross-contamination between samples. Once homogenized, the 187 sample was sonicated for 5 min and then centrifuged at 1000 rcf for 5 min. Next, 1 mL of the 188 supernatant was syringe-filtered through a 0.25 µm nylon filter into a 2 ml HPLC vial. The 189 extract obtained was analysed immediately, or when this was not possible, stored at -20 °C 190 until analysis (for no longer than 24 h). 191

Muscle, liver and kidney samples from the first 125 pig carcasses and the liver of the 192 first 10 forensic avian scavenger cases were analysed by liquid chromatography with 193 electrospray ionization mass spectrometry (LC-ESI-MS) using an Agilent 1100 LC coupled 194 to an Agilent 6110 single quadrupole MS following the method described by Taggart et al. 195 (2009). For subsequent samples, we used ultra-high-performance liquid chromatography 196 (UHPLC) with MS/MS time-of-flight mass spectrometry (LC-QTOF-MS; AB Sciex 197 TripleTOFTM4600 System). Chromatographic separation was carried out using a Poroshell-198 120EC-C18 column (2.1 x 150 mm, 2.7 µm). Chromatography conditions were as follows: 199

flow 0.5 mL min⁻¹; column temperature 40°C; gradient elution with (A) 0.1% formic acid in 200 Milli-Q and (B) 0.1% formic acid in acetonitrile. Initial conditions were 40% phase A and 201 60% phase B for 1 min, then a 5 min linear gradient to 35% A and 65% B, followed by 100% 202 B for 2 min, returning over 1 min to initial conditions. Injection volume was 5 µL and vials 203 were kept cool at 4 °C in the autosampler. The Q-TOF parameters were as follows: gas flow 204 (CUR) at 20 psi, source 1 gas (GS1) at 40 psi, source 2 gas (GS2) at 40 psi, maximum 205 temperature 400 °C (TEM), collision energy (CE) -35 V, propagation of collision energy 206 (CES) 15 V and fragmentation potential (DP) -100 V. The molecular weights for the 207 precursor ions and the three main fragmentation ions for the NSAIDs analysed in MRM 208 (multiple reaction monitoring) mode with positive ionization are shown in Table S1. 209 Quantification was performed using the most abundant fragment ion, with a fragmentation 210 voltage of 50 to 500 V and a capillary voltage of 4500 V. 211

Calibrations were performed using diluted working solutions made up from stock 212 solutions at 1 mg mL⁻¹ for each NSAID. From these, mixed working solutions were prepared 213 and kept at 4 °C until use. Mixed standards were made at concentrations of 50, 100, 200 and 214 400 ng mL⁻¹ in a final volume of 1 mL whilst including 0.25 ng mL⁻¹ of internal standard 215 (flunixin-d3). Blank and fortified samples were also made using chicken liver (tissue 216 surrogate) at NSAID levels of 50, 100 and 200 ng g⁻¹. These were processed daily in order to 217 estimate the accuracy and precision of the analytical technique (% recovery \pm RSD). We 218 obtained recovery rates between 87% (for tolfenamic acid) and 129% (for suxibuzone) and 219 RSD values ranged between 5.72 (for diclofenac) and 19.45 (for tolfenamic acid) (Table S2). 220 Regression coefficients (R^2) in fortified calibration spikes were between 0.879 (for naproxen) 221 and 0.989 (for meloxicam) (Table S2). Limits of quantification (LOOs) were established at 222 10 times the signal to noise ratio, and were between 0.0002 mg kg⁻¹ (for flunixin) and 0.02 223 $mg kg^{-1}$ (for naproxen) (Table S2). 224

225

226 2.4. Data analysis and interpretation

227	Detection frequency for each NSAID in carrion and avian scavengers was calculated
228	and compared (i.e., between regions and species) using Fisher's exact tests with IBM SPSS
229	Statistics 24. To analyse the risk of intoxication by NSAIDs from carrion ingestion, we used
230	median lethal dose (LD ₅₀) information, where available. For diclofenac, we used the LD ₅₀ of
231	0.098-0.225 mg kg ⁻¹ bw calculated for white-rumped vulture by Swan et al. (2006). For
232	flunixin, we used a lethal dose range of 1-4.5 mg kg ⁻¹ bw estimated for Rüppell's griffon
233	vulture (Gyps rueppellii) and cinereous vulture (Cuthbert et al., 2007), and for ketoprofen
234	1.5-5 mg kg ⁻¹ estimated in Cape griffon vulture (<i>Gyps coprotheres</i>) and white-backed vulture
235	(Naidoo et al., 2010b). Using the concentrations detected in carrion, we calculated the
236	estimated theoretical exposure (ETE) in Eurasian griffon with a mean body weight of 7.4 kg
237	and with an average daily intake of 1.2 kg of food (Donázar, 1993). These ETEs were used
238	with the LD ₅₀ values to estimate toxicity exposure ratios (TER=LD ₅₀ /ETE) for each NSAID
239	for Gyps. TERs were estimated using the minimum and maximum LD ₅₀ values noted above.
240	This ratio is widely used to evaluate the first-tier risk of exposure to a chemical substance
241	(such as a pesticide in birds) and must be <10 to represent an acute toxicity risk to wild birds
242	(EFSA, 2009). In the case of diclofenac, we also calculated the per-meal probability of death
243	in vultures feeding on the analysed carrion using the parameters of the dose-response curves
244	for Gyps species (Swan et al., 2006).

NSAID concentrations in vulture tissues have been interpreted based on previous studies, which associated residues in vultures with adverse effects and/or mortality. Diclofenac levels of 0.05-0.64 mg kg⁻¹ in kidney and flunixin levels of 2.7 mg kg⁻¹ in liver and 6.5 mg kg⁻¹ in kidney have been considered compatible with lethal poisoning by these NSAIDs in vultures (Oaks et al., 2004; Zorrilla et al., 2015). Likewise, the presence of visceral gout at post-mortem examination in birds with NSAID residues was considered
additional evidence of NSAID intoxication (Oaks et al., 2004; Zorrilla et al., 2015; Cuthbert
et al., 2016).

253

254 **3. Results**

255

256 *3.1. Detection of NSAIDs in carrion and risk assessment for vultures*

NSAID residues were detected in 3.07% (7/228) of all carrion tested (Table 1; Table 257 S3). We detected 5 NSAID positive samples in pigs originating from intensive production, 258 which represented 3.20% of the total pig carcasses analysed (n=156). Further, there were 2 259 positive sheep samples, representing 4.44% of the total sheep carcasses tested (n=45). Pig 260 samples were positive for flunixin (n=2, 1.28%), diclofenac (n=1, 0.64%), ketoprofen (n=1, 261 0.64%), and meloxicam (n=1, 0.64%). Sheep were positive for flunixin (n=2, 4.44%) (Table 262 1). None of the carcasses of goat (n=4), horse (n=4), cow (n=6) or 'unknown' species had 263 NSAID residues. However, these differences in prevalence between species were not 264 statistically significant. Positive carrion were detected in three regions: Castile and Leon (2 of 265 35, 5.71%), Aragon (3 of 112, 2.68%), and Valencia (2 of 18, 11.11%) (see Fig. 1 and Table 266 S4). There was a marginal significant difference between Aragon and Valencia (Fisher's test, 267 p=0.051). Prevalence is also shown in more detail by provinces (Fig. S1). 268

The estimated acute TER value was well above 10 for all samples, except for the one positive to diclofenac and one positive to flunixin (Table 1). The diclofenac positive pig muscle with 0.171 mg kg⁻¹ (ETE of 0.028 mg kg⁻¹ bw) resulted in a per-meal probability of death for vultures of 25.4% or 0.8% using the relevant dose-response curves and with an LD₅₀ of 0.098 or 0.225 mg kg⁻¹ bw, respectively. However, it must be acknowledged that diclofenac residues in this muscle tissue were detected only at an injection point (Fig. S2), so,

the real risk from this specific carcass was probably lower. The sheep liver with the highest
level of flunixin had a TER value ranging between 0.22-1, so the risk of poisoning here was
very high. The per-meal probability of death for vultures could not be calculated in this case
because there is no available dose-response curve for flunixin in *Gyps* vultures.

279

280 *3.3. Monitoring NSAIDs in dead avian scavengers including vultures*

We observed that 3.60% (14/389) of individuals had detectable NSAID residues in 281 liver and/or kidney (Table 2; Table S5). Eleven Eurasian griffons analysed (3.59%) were 282 positive for NSAIDs, specifically meloxicam (n=7, 2.29%) and flunixin (n=4, 1.30%). 283 Meloxicam was also detected in one Egyptian vulture (9.09%), one common buzzard (3.84%) 284 and one black kite (33.33%) (Table S5). Concentrations ranged between 0.023-20.35 mg kg⁻¹ 285 for flunixin and between 0.033-2.44 mg kg⁻¹ for meloxicam (Table 2). By region, the 286 prevalence was highest in Castile and Leon (1/2, 50%), followed by Cantabria (1/12, 8.33%), 287 Madrid (1/15, 6.67%), Aragon (4/85, 4.71%), Catalonia (6/133,4.51%) and Extremadura 288 (1/28, 3.57%) (Fig. 1). However, prevalence was not significantly different between regions 289 or between species. The situation in Castile and Leon may warrant further research given that 290 one out of two animals had NSAID residues. Prevalence is also included in more detail by 291 province (Fig. S1). 292

Post-mortem examinations showed that 10 out of 306 dead Eurasian griffons had degenerative lesions in kidney and/or liver (3.27%) and four of these presented extensive visceral gout (1.31%) (Fig. S2). Two of these cases (#2 and #4 in Table 2) also had elevated flunixin levels in tissues (20.35 mg kg⁻¹ in the kidney and 11.32 mg kg⁻¹ in the liver, and 4.91 mg kg⁻¹ in liver, respectively). The other two Eurasian griffons with visceral gout had no detectable NSAID residues in their tissues. A third Eurasian griffon (found dead under a cliff with lesions of traumatism) also had 0.33 mg kg⁻¹ of flunixin in liver and renal degeneration

(#1 in Table 2), while a fourth Eurasian griffon with 0.023 mg kg⁻¹ flunixin in liver had no
lesions, gout or kidney damage on necropsy (#3 in Table 2). In addition to these wild birds,
one Eurasian griffon that died in a wildlife rehabilitation centre was tested as it was suspected
to have died from iatrogenic flunixin poisoning (Soler et al., 2016). This bird had 2.83 mg kg⁻¹
in liver and 0.44 mg kg⁻¹ in muscle and visceral gout (#5 in Table 2); as such, these levels
were comparable with Eurasian griffons found dead in the field with this lesion.

306

307 4. Discussion

Residues of diclofenac and three other NSAIDs (flunixin, ketoprofen and meloxicam) have been detected in livestock carcasses supplied to supplementary feeding stations for avian scavengers in Spain. Diclofenac poisoning has not been detected in the avian scavengers tested, but flunixin poisoning has been confirmed in three wild Eurasian griffons in which the presence of the chemical was accompanied with visceral gout and/or renal damage.

314

315 *4.1. Risk assessment based on NSAID residues in carrion*

The first objective of the present study was to evaluate the risk of exposure to 316 diclofenac in avian scavengers in the Iberian Peninsula. Two commercial formulations of 317 diclofenac have been registered for veterinary use in livestock since 2013 in Spain 318 (CIMAVET, 2020). Diclofenac is not yet authorized in Portugal by the national authority 319 (Direção-Geral de Alimentação e Veterinária), despite a vote in favour of its use in the 320 Portuguese Parliament in January 2019. The carrion analyses performed here shows a 321 potential risk of exposure to diclofenac in Iberian avian scavengers because one pig carcass 322 was found positive. Therefore, the labelling of commercial diclofenac formulations, which 323

includes warnings to avoid disposal of carrion from treated animals for vulture feeding, is notbeing effective in all cases.

The pig carrion with diclofenac residues in muscle was possibly from an animal treated 326 327 more than 168 h before death, because residues in the muscle were limited to an area associated with the likely injection point (Fig. S2), and no residues were detected in liver or 328 kidney (Naidoo et al., 2018). In Spain, diclofenac dosage for pig is specified at 2.3 mg kg⁻¹ (1 329 mL per 20 kg bw of a solution with 46 mg mL⁻¹ of diclofenac) administered intramuscularly 330 in a three-day treatment pattern, with no more than 3 mL injected in a single point. Therefore, 331 a pig weighing 120 kg would need a daily dose of 276 mg, i.e., 138 mg in each of two 332 injection points every day, resulting in six points after a 3-day treatment (AEMPS, 2018). 333 The pharmacokinetics for diclofenac in pig indicate a 3.4 h elimination half-life and a 334 maximum plasma level of 4.7 μ g mL⁻¹ at 0.5 h (AEMPS, 2018). In addition, experimental 335 studies discussed in Green et al. (2006) describe a higher half-life in muscle (15 h) than in 336 kidney and liver (6-8 h). In the case of this diclofenac positive pig carcass, acute poisoning 337 could occur if scavengers consumed muscle from the injection sites, as has been observed for 338 carprofen (Naidoo et al., 2018). 339

Green et al. (2004) estimated that just 0.13-0.75% of carcasses needed to contain a 340 lethal level of diclofenac to explain (alone, without any other drivers) the rapid population 341 declines seen for *Gyps* vultures in South Asia. We found that 0.64% of pigs tested positive to 342 diclofenac, so the risk to Iberian avian scavengers exists. The relatively small number of 343 carrion samples tested, alongside the fact that the single positive was from a pig with 344 diclofenac residues at an injection site only, limits the possibility to perform a more robust 345 risk assessment. However, the scenario observed here, on the Iberian Peninsula, is far from 346 that seen in India where diclofenac residue prevalence prior to any legal ban was ~10% 347

nationally, with certain states monitored with 22.3% diclofenac positive carcasses (Taggart etal., 2007).

In addition to diclofenac, we detected other NSAIDs in pig and sheep carcasses, 350 351 specifically flunixin, ketoprofen and meloxicam. A risk to avian scavengers (according to TER calculations) was only noted in one sheep due to the high level of flunixin found. 352 According to data reported from 2004-2018 in the Spanish Residue Research National Plan 353 (PNIR: the focus of which is human food safety), two NSAIDs have been detected in samples 354 obtained from slaughterhouses (both in 2016), specifically diclofenac in a horse (5.88%, 355 1/17) and flunixin in a cow (25%, 1/4) (PNIR, 2016). Flunixin and ketoprofen are both 356 thought to be toxic to *Gyps* vultures, causing visceral gout and rapid death mortality, although 357 potentially at higher doses than for diclofenac (Cuthbert et al., 2007; Naidoo et al., 2010, 358 2010b; Zorrilla et al., 2015). Based on pharmacokinetic data, the two flunixin positive pigs 359 here probably died >48 h after treatment (Buur et al., 2006) and the ketoprofen positive pig 360 likely died >25 h after treatment (Mustonen et al., 2012). In terms of the two flunixin cases in 361 sheep, the animal with the highest level $(27.48 \text{ mg kg}^{-1})$ probably died quickly after treatment 362 (within 5 h) due to the elevated levels in liver, while the second (at 0.297 mg kg⁻¹ in liver) 363 likely died 10 to 15 h after treatment (Cheng et al., 1998). 364

365

366 *4.2. NSAID poisoning in Iberian avian scavengers*

We have not detected cases of diclofenac poisoning in avian scavengers from the Iberian Peninsula to date, despite the previous estimations of mortality performed by AEMPS & MAGRAMA (2014) and Green et al. (2016). In the specific case of porcine livestock, AEMPS & MAGRAMA (2014) assumes that vultures consume 38413 intensively reared pig carcasses per year, of which, 0.14-0.24% could contain diclofenac residues. Based on this, and proposing different diclofenac concentration scenarios in carrion (0.1, 0.4 and 0.8 mg kg⁻

373 ¹) and time intervals between last diclofenac injection and carrion intake (0-3, 3-12 and 12-24 h), AEMPS & MAGRAMA (2014) estimated that the number of vultures that could die per 374 year in Spain (from diclofenac in pig carcasses) would be between 4-7 individuals. This 375 markedly contrasts with the estimations of Green et al. (2016) that calculated 364-4609 376 annual deaths of Eurasian griffons due to pig carcasses. The main difference between these 377 studies is that Green et al. (2016) assumed that all carrion available (containing diclofenac 378 residues) had the potential to be toxic, given that experimental studies have indicated marked 379 interindividual variations. Mortality after exposure has been observed at doses as low as 380 0.007 mg kg⁻¹ bw (Oaks et al., 2004; Swan et al., 2006). Although the only carcass in our 381 study with diclofenac residues would likely not pose a high risk to vultures, we used our 382 0.64% diclofenac prevalence value to recalculate the proportion of carcasses that could 383 contain lethal levels for vultures in Spain and then refine the risk assessment. We can 384 estimate that the probability of dying in the first 8 h after last treatment would be 4.76% (8 385 h/168 h) for all diclofenac treated animals, so the percentage of carrion with potentially lethal 386 diclofenac levels would be $0.0476 \times 0.64 = 0.03\%$ (Table 3). Here we assume that probability 387 of death at a determined time is constant throughout the 168 h period after treatment, during 388 which diclofenac residues in tissues are above our limit of quantification. With this 389 percentage (0.03%) and the number of swine carcasses available (38413, AEMPS & 390 MAGRAMA, 2014), we can estimate the number of treated pigs with toxic levels (12) and 391 392 the number of meals available to vultures from these carrion (1600). Following the approach of Green et al. (2016) (with the proportion of vultures killed by feeding on a contaminated 393 pig treated 8 h before death), we can estimate that 78-600 vultures would die per year (with 394 LD_{50} of 0.098 and 0.225 mg kg⁻¹, respectively), which is between the ranges given in 395 previous estimations (Table 3). These estimates are based on LD_{50} data and dose-response 396

curves showing that some individuals can be especially sensitive to diclofenac, so somemortality may occur at doses much lower than the median value.

In contrast with diclofenac, flunixin poisoning has been detected in three Eurasian 399 griffons in this study, each showing visceral gout (Fig. S3) and/or kidney degeneration and 400 flunixin residues in liver between 0.33 and 11.32 mg kg⁻¹. These residue levels are 401 comparable with those detected in an iatrogenic flunixin poisoning in one Eurasian griffon, 402 with 2.83 mg kg⁻¹ of flunixin in liver and visceral gout (Soler et al., 2016); and, the case 403 described by Zorrilla et al. (2015) of another Eurasian griffon with 2.7 mg kg⁻¹ of flunixin in 404 liver and visceral gout. Flunixin poisoning has also been described in two Rüppell's griffon 405 vultures (Gyps rueppellii) and one white-backed vulture in captivity, with 0.016-0.039 mg kg⁻ 406 ¹ of flunixin in several tissues, who fed on flunixin contaminated beef with 31.35 mg kg⁻¹ 407 (Eleni et al., 2019). Flunixin has also been linked to possible iatrogenic poisoning in other 408 birds in captivity, including three cinereous vultures, one Rüppell's griffon and one white-409 backed vulture, at exposure doses of 1-4.5 mg kg⁻¹ (Cuthbert et al., 2007). Thus, our results 410 clearly confirm that vultures are dying due to flunixin in Spain, and the mortality observed 411 here of 3 out of 306 Eurasian griffons represents 0.98% of the studied cases (see Table S6). 412 With 30946 breeding pairs of Eurasian griffons in Spain, a productivity of 0.56 chicks/nest 413 and considering a stable population (natality~mortality), we would estimate an annual 414 mortality of 170 griffon vultures due to flunixin poisoning. 415

Meloxicam residues were detected in seven vultures with evidence of traumatism, electrocution, intoxication or suspected previous intoxication. None of the meloxicam positive birds had visceral gout. Further, meloxicam is not thought to be a risk to vultures as extensive vulture safety testing has taken place to demonstrate this (Swan et al., 2006b; Swarup et al., 2007; Naidoo et al., 2008; Mahmood et al., 2010).

Finally, it should be noted that while visceral gout is a frequent lesion observed in relation to NSAID poisoning in birds, confirmation must be attained using parallel chemical analysis of kidney or liver tissues. Beyond NSAID poisoning, gout can also be caused by metabolic disorders, dehydration, infectious etiology, renal damage or other nephrotoxic agents (Echols, 2016).

426

427 **5.** Conclusions

The Eurasian griffon population has increased in Spain from 24541 to 30946 breeding 428 pairs from 2008 to 2018 (+21.16%; Del Moral & Molina, 2018), so at the moment, there is no 429 evidence of a population level impact of diclofenac use in livestock on this species. 430 Nevertheless, monitoring efforts to study causes of mortality in Iberian avian scavengers 431 must continue because of the observed risk posed by the potential disposal of diclofenac 432 treated carrion in the field or in supplementary feeding stations. The presence of diclofenac in 433 one carcass indicates a failure in the formulation advisory systems in Spain (i.e., given on 434 product labels) which states that diclofenac treated animal carrion should "never reach the 435 trophic chain of wild animals". Likewise, the same recommendations should be applied to 436 formulations of flunixin and ketoprofen marketed in Spain, and to that of any veterinary 437 pharmaceutical known to be toxic to scavengers. For flunixin, levels capable of causing acute 438 toxicity in vultures were clearly identified and as such changes to labelling/advice are 439 certainly needed to protect these scavengers. In addition, this NSAID is not currently 440 registered in Europe for veterinary use in sheep (EMEA, 2000), so these results clearly 441 suggest that veterinary drugs have extra-label use. An effective risk assessment for veterinary 442 drugs must always consider the possibility that these may enter wildlife food webs through a 443 livestock carrion pathway. But also, farmers, veterinarians and wildlife technicians in charge 444 of managing supplementary feeding stations or the disposal of carrion in the field must be 445

aware of the risks that pharmaceutical treated livestock may represent for avian scavengers
(Mateo et al., 2015; Zorrilla et al., 2015; Casas-Díaz et al., 2016).

448

449 Acknowledgments

The authors gratefully acknowledge support from regional governments and other institutions 450 in Spain, specifically from Aragon, Catalonia, Extremadura (Acción por el Mundo Salvaje: 451 AMUS), Asturias, Andalusia, La Rioja, Castile and Leon, Madrid, Cantabria, Navarra, the 452 Balearic Islands, Valencia, Galicia, the Basque Country and Asturias. We would like to thank 453 the veterinarians who carried out post-mortem examinations, for their assistance in tissue 454 collection in the field and at wildlife rehabilitation centres. A special mention is given to 455 Rafael Molina and Olga Nicolas for performing a high number of the vulture necropsies and 456 submitting samples to IREC, specifically some key individuals within the intoxication case 457 group. We thank all the TRAGSATEC technicians and other staff from the regional 458 governments who co-operated and contributed in terms of providing tissue samples from 459 livestock carcasses from supplementary feeding stations. 460

461

462 Funding

The authors gratefully acknowledge funding from the Morris Animal Foundation within the 463 project D16ZO-046-MAF-AvianScaven, TRAGSATEC through contract Ref. TEC0004566 464 and CGL2013–40975-R project from I+D+I National Plan funded by the Spanish Ministry of 465 Economy and Competitiveness. We also acknowledge that the Portuguese samples were 466 received financial due FCT/MCTES CESAM 467 to support to (UIDP/50017/2020+UIDB/50017/2020), through national funds. 468

469

470 **CRediT** author statement

471	The manuscript was	written	through	contributions	of	all	authors.	All	authors	have	given
472	approval to the final	version c	of the man	nuscript.							

Marta Herrero-Villar: Validation, Formal analysis, Investigation, Writing - Original Draft,
Writing - Review & Editing, Roser Velarde: Investigation, Pablo R. Camarero: Validation,
Investigation, Mark A. Taggart: Writing - Review & Editing, Victor Bandeira: Resources,
Funding acquisition, Carlos Fonseca: Resources, Funding acquisition, Ignasi Marco:
Resources, Supervision, Project administration, Funding acquisition, Rafael Mateo:
Conceptualization, Methodology, Resources, Writing-Original Draft, Writing - Review &

479 Editing, Supervision, Project administration, Funding acquisition.

480

481 **Declaration of Competing Interest**

482 The authors declare no competing financial interests or personal relationships that could have

483 appeared to influence the present study.

484

485 **References**

- 486 Agencia Española de Medicamentos y Productos Sanitarios; AEMPS, 2018.
- 487 <u>https://cimavet.aemps.es/cimavet/pdfs/es/p/2760+ESP/P_2760+ESP.pdf</u> (Accessed
 488 Jan 8, 2020).
- 489 AEMPS & MAGRAMA, 2014. Análisis de riesgo de uso de medicamentos veterinarios con
- 490 diclofenaco sobre las poblaciones de buitres en España: Recomendaciones de
- 491 *actuación y escenarios potenciales de afección*. Agencia Española de Medicamentos y
- 492 Productos Sanitarios & Ministerio de Agricultura, Alimentación y Medio Ambiente,

493 España.

494 BirdLife International, 2015. European Red List of Birds. Luxembourg: Office for Official

495 Publications of the European Communities.

496 http://datazone.birdlife.org/userfiles/file/Species/erlob/supplementarypdfs/22695219_ gyps fulvus.pdf (Accessed May 23, 2020). 497 BirdLife International, 2015. European Red List of Birds. Luxembourg: Office for Official 498 499 Publications of the European Communities. http://datazone.birdlife.org/userfiles/file/Species/erlob/supplementarypdfs/22695231_ 500 aegypius_monachus.pdf (Accessed May 23, 2020). 501 BirdLife International, 2015. European Red List of Birds. Luxembourg: Office for Official 502 Publications of the European Communities. 503 504 http://datazone.birdlife.org/userfiles/file/Species/erlob/supplementarypdfs/22695180_ neophron_percnopterus.pdf (Accessed May 23, 2020). 505 Buur, J. L.; Baynes, R. E.; Smith, G.; Riviere, J. E., 2006. Pharmacokinetics of flunixin 506 meglumine in swine after intravenous dosing. J. Vet. Pharmacol. Ther. 29, 437-440; 507 DOI 10.1111/j.1365-2885.2006.00788.x. 508 Casas-Díaz, E.; Cristòfol, C.; Cuenca, R.; Agustí, S.; Carneiro, M.; Marco, I.; Lavín, S.; 509 Margalida, A. 2016. Determination of fluoroquinolone antibiotic residues in the 510 plasma of Eurasian griffon vultures (Gyps fulvus) in Spain. Sci. Total Environ. 557-511 558, 620-626; DOI 10.1016/j.scitotenv.2016.03.083. 512 Chaudhry, M. J. I.; Ogada, D. L.; Malik, R. N.; Virani, M. Z.; Giovanni, M. D., 2012. First 513 evidence that populations of the critically endangered long-billed vulture *Gyps indicus* 514 in Pakistan have increased following the ban of the toxic veterinary drug diclofenac in 515 south Asia. Bird Conserv. Int. 22, 389-397; DOI 10.1017/S0959270912000445. 516 Cheng, Z.; McKeller, Q.; Nolan, A., 1998. Pharmacokinetic studies of flunixin meglumine 517 and phenylbutazone in plasma, exudate and transudate in sheep. J. vet. Pharmacol. 518 Ther. 21, 315-321; DOI 10.1046/j.1365-2885.1998.00144.x. 519

- 520 Centro de Información online de Medicamentos Veterinarios de la AEMPS; CIMAVET,
- 521 2020. <u>https://cimavet.aemps.es/cimavet/publico/home.html</u> (Accessed June 8, 2020).
- 522 Cuthbert, R.; Parry-Jones, J.; Green, R. E.; Pain, D. J., 2007. NSAIDs and scavenging birds:
- 523 Potential impacts beyond Asia's critically endangered vultures. *Biol. Lett.* 3, 90-93;
- 524 DOI 10.1098/rsbl.2006.0554.
- 525 Cuthbert, R.; Taggart, M. A.; Prakash, V.; Saini, M.; Swarup, D.; Upreti, S.; Mateo, R.;
- 526 Chakraborty, S. S.; Deori, P.; Green, R. E., 2011. Effectiveness of action in India to
- 527 reduce exposure of *Gyps* vultures to the toxic veterinary drug diclofenac. *PLoS ONE*.
- 528 6, DOI 10.1371/journal.pone.0019069.
- 529 Cuthbert, R. J.; Prakash, V.; Saini, M.; Upreti, S.; Swarup, D.; Das, A.; Green, R. E.; Taggart,
- 530 M., 2011b. Are conservation actions reducing the threat to India's vulture
- 531 populations? *Curr. Sci.* 101, 1480-1484.
- 532 Cuthbert, R. J.; Taggart, M. A.; Prakash, V.; Chakraborty, S. S.; Deori, P.; Galligan, T.;
- 533 Kulkarni, M.; Ranade, S.; Saini, M.; Sharma, A. K.; Shringarpure, R.; Green, R. E.,
- 534 2014. Avian scavengers and the threat from veterinary pharmaceuticals. *Philos*.

535 Trans. R. Soc. Lond. B. Biol. Sci. 369 (1656), DOI 10.1098/rstb.2013.0574.

- 536 Cuthbert, R. J.; Taggart, M. A.; Saini, M. A.; Sharma, A.; Das, A.; Kulkarni, M. D.; Deori,
- 537 P.; Ranade, S.; Shringarpure, R. N.; Galligan, T. H.; Green, R. E., 2016. Continuing
- 538 mortality of vultures in India associated with illegal veterinary use of diclofenac and a
- potential threat from nimesulide. *Oryx*. 50, 104-112; DOI
- 540 10.1017/S003060531500037X.
- 541 Del Moral, J. C., 2009. Resultados generales, in: Moral, J. C. (Ed.), El Alimoche Común en
- 542 España: Población Reproductora en 2008 y Método de Censo. SEO/BirdLife, Madrid,
- 543 pp 14-24. <u>https://www.seo.org/wp-content/uploads/2012/04/31_alimoche.pdf</u>
- 544 (Accessed January 8, 2020).

545	Del Moral, J. C., 2018. Resultado generales, in: Moral, J. C. (Ed.), El Buitre Negro en
546	España: Población Reproductora en 2017 y Método de Censo. SEO/BirdLife, Madrid,
547	pp 12-28.
548	https://www.seo.org/boletin/seguimiento/censos/45%20buitre%20negro/html5forpc.ht
549	<u>ml?page=0</u> (Accessed January 8, 2020).
550	Del Moral, J. C.; Molina, B., 2019. Resultado generales, in: Moral, J. C.; Molina, B. (Eds.),
551	El Buitre Leonado en España: Población Reproductora en 2018 y Método de Censo.
552	SEO/BirdLife, Madrid, pp 17-34.
553	https://www.seo.org/boletin/seguimiento/censos/50%20buitre%20leonado/html5forpc
554	<u>.html?page=0&bbv=1&pcode=</u> (Accessed January 8, 2020).
555	Donázar, J. A., 1993. Los Buitres Ibéricos. Biología y Conservación. In: J. M. Reyero (Ed.),
556	Madrid, Spain, 285 pp.
557	Echols, M. S., 2006. Evaluating and Treating the Kidneys. In: Harrison, J. G., Lightfoot, T.,
558	(Eds.), Clinical Avian Medicine. Spix Publishing Palm Beach, Florida, USA, pp. 453-
559	487. DOI 10.1647/10826742(2006)20[285:CAM]2.0.CO;2.
560	Eleni, C.; Neri, B.; Giannetti, L.; Grifoni, G.; Meoli, R.; Stravino, F.; Friedrich, K. G.; Scholl,
561	F.; Di Cerbo, P.; Battisti, A., 2019. Death of captive-bred vultures caused by flunixin
562	poisoning in Italy. Environ. Toxicol. Pharmacol. 68, 91-93; DOI
563	10.1016/j.etap.2019.03.011.
564	European Food Safety Authority, 2009. Guidance Document on Risk Assessment for Birds
565	and Mammals on request from EFSA. EFSA Journal. 7, 1438; DOI
566	10.2903/j.efsa.2009.1438.
567	European Medicines Agency, 2000. Committee for veterinary medicinal products: flunixin
568	(extension to horses). EMEA, Summary Report 2. EMEA/MRL/744/00-FINAL.
569	https://www.ema.europa.eu/en/documents/mrl-report/flunixin-extension-horses-

23

	Journal Pre-proof
570	summary-report-2-committee-veterinary-medicinal-products_en.pdf (Accessed April
571	6, 2020).
572	Fourie, T.; Cromarty, D.; Duncan, N; Wolter, K.; Naidoo, V., 2015. The safety and
573	pharmacokinetics of carprofen, flunixin and phenylbutazone in the Cape vulture
574	(Gyps coprotheres) following oral exposure. PLoS ONE. 10, 1-12; DOI
575	10.1371/journal.pone.0141419.
576	Galligan, T. H.; Taggart, M. A.; Cuthbert, R. J.; Svobodova, D.; Chipangura, J.; Alderson,
577	D.; Prakash, V. M.; Naidoo, V., 2016. Metabolism of aceclofenac in cattle to vulture-
578	killing diclofenac. Conserv. Biol. 30, 1122-1127; DOI 10.1111/cobi.12711.
579	Gilbert, M.; Watson, R.T.; Virani, M. Z.; Oaks, J. L.; Ahmed, S.; Chaudry, M. J. I.; Arshad,
580	M.; Mahmood, S.; Ali, A.; Khan, A. A., 2006. Rapid population declines and
581	mortality clusters in three Oriental white-backed vulture Gyps bengalensis colonies in
582	Pakistan due to diclofenac poisoning. Oryx. 40, 388-399; DOI
583	10.1017/S0030605306001347.
584	Green, R. E.; Newton, I.; Shultz, S.; Cunningham, A. A.; Gilbert, M.; Pain, D. J.; Prakash,
585	V., 2004. Diclofenac poisoning as a cause of vulture population declines across the
586	Indian subcontinent. J Appl Ecol. 4, 793-800; DOI 10.1111/j.0021-
587	8901.2004.00954.x.
588	Green, E. E.; Taggart, M. A.; Das, D.; Pain, D. J.; Kumar, S. C.; Cunningham, A. A.;
589	Cuthbert, R., 2006. Collapse of Asian vulture populations: risk of mortality from
590	residues of the veterinary drug diclofenac in carcasses of treated cattle. J Appl Ecol.
591	43, 949-956; DOI 10.1111/j.1365-2664.2006.01225.x.
592	Green, R. E.; Donázar, J. A.; Sánchez-Zapata, J. A.; Margalida, A., 2016. Potential threat to
593	Eurasian griffon vultures in Spain from veterinary use of the drug diclofenac. J Appl
594	<i>Ecol.</i> 53, 993-1003; DOI 10.1111/1365-2664.12663.

- Khan, M. M. H., 2013. Population, breeding and threats to the white-rumped vulture *Gyps bengalensis* in Bangladesh. *Forktail*. 29, 52-56.
- 597 Mahmood, K. T.; Ashraf, M.; Ahmad, M. U., 2010. Eco-Friendly Meloxicam Replaces Eco-
- 598 Demaging Diclofenac Sodium in Veterinary Practice in South Asia A Review. J.
 599 *Pharm. Sci. Res.* 2, 672-685.
- Margalida, A.; Bogliani, G.; Bowden, C. G. R.; Donázar, J. A.; Genero, F.; Gilbert, M.;
- 601 Karesh, W. B.; Kock, R.; Lubroth, J.; Manteca, X.; Naidoo, V.; Neimanis, A.,
- 602 Sánchez-Zapata, J. A.; Taggart, M. A.; Vaarten, J.; Yon, L.; Kuiken, T.; Green, R. E.,
- 603 2014. One Health approach to use of veterinary pharmaceuticals. *Science*. 346, 1296-
- 604 1298; DOI 10.1126/science.1260260.
- Mateo, R.; Sánchez-Barbudo, I. S.; Camarero, P. R.; Martínez, J. M., 2015. Risk assessment
- 606 of bearded vulture (*Gypaetus barbatus*) exposure to topical antiparasitics used in
- 607 livestock within an ecotoxicovigilance framework. *Sci. Total Environ.* 536, 704-712;
- 608 DOI 10.1016/j.scitotenv.2015.07.109.
- Mustonen, K.; Niemi, A.; Raekallio, M.; Heionnen, M.; Peltoniemi, O. A.; Palviainen, M.;
- 610 Siven, M.; Peltoniemi, M.; Vainio, O., 2012. Enantiospecific ketoprofen
- 611 concentrations in plasma after oral and intramuscular administration in growing pigs.
- 612 *Acta Vet. Scand.* 54, DOI 10.1186/1751-0147-54-55.
- Naidoo, V.; Wolter, K.; Cromarty, A. D.; Bartels, P.; Bekker, L.; McGaw, L.; Taggart, M. A.;
- 614 Cuthbert, R.; Swan, G. E., 2008. The pharmacokinetics of meloxicam in vultures. *J*.
- 615 *Vet. Pharmacol. Ther.* 31, 128-34; DOI 10.1111/j.1365-2885.2007.00923.x.
- 616 Naidoo, V.; Wolter, K.; Cuthbert, R.; Duncan, N., 2009. Veterinary diclofenac threatens
- 617 Africa's endangered vulture species. *Regul. Toxicol. Pharmacol.* 53, 205-208; DOI
- 618 10.1016/j.yrtph.2009.01.010.

	urn		D			
U	սո	al			ιU	

- Naidoo, V.; Venter, L.; Wolter, K.; Taggart, M.; Cuthbert, R., 2010. The toxicokinetics of
- 620 ketoprofen in *Gyps coprotheres*: toxicity due to zero-order metabolism. *Arch. Toxicol.*
- 621 84, 761-766; DOI 10.1007/s00204-010-0521-0.
- Naidoo, V.; Wolter, K.; Cromarty, D.; Diekmann, M.; Duncan, N.; Meharg, A. A.; Taggart,
- 623 M. A.; Venter, L.; Cuthbert, R., 2010b. Toxicity of non-steroidal anti-inflammatory
- drugs to *Gyps* vultures: a new threat from ketoprofen. *Biol. Lett.* 6, 339-341; DOI
- 625 10.1098/rsbl.2009.0818.
- 626 Naidoo, V.; Taggart, M. A.; Duncan, N.; Wolter, K.; Chipangura, J.; Green, R. E.; Galligan,
- 627 T. H., 2018. The use of toxicokinetics and exposure studies to show that carprofen in
- 628 cattle tissue could lead to secondary toxicity and death in wild vultures. *Chemosphere*.
- 629 190, 80-89; DOI 10.1016/j.chemosphere.2017.08.167.
- 630 Oaks, J. L.; Gilbert, M.; Virani, M. Z.; Watson, R. T.; Meteyer, C. U.; Rideout, B. A.;
- 631 Shivaprasad, H. L.; Ahmed, S.; Chaudhry, M. J. I.; Arshad, M.; Mahmood, S.; Ali, A.;
- 632 Khan, A. A., 2004. Diclofenac residues as the cause of vulture population decline in

633 Pakistan. *Nature*. 427, 630-633; DOI 10.1038/nature02317.

- Oaks, J. L.; Watson, R. T., 2011. South Asian Vultures in Crisis: Environmental
- 635 Contamination with a Pharmaceutical. In: Elliot, J. E., Bishop, C. A., Morrisey, C. A.,
- 636 (Eds.), Wildlife Ecotoxicology. Springer New York, Oak Ridge, pp. 413-439. DOI
- 637 10.1007/978-0-387-89432-4_14.
- 638 PNIR, 2016. Plan Nacional de Investigación de Residuos.
- 639 https://www.mapa.gob.es/es/ganaderia/temas/sanidad-animal-higiene-
- 640 ganadera/higiene-de-la-produccion-primaria-ganadera/plan-nacional-de-investigacion-
- 641 <u>de-residuos-pnir/</u> (Accessed June 8, 2020).
- 642 Prakash, V.; Pain, D. J.; Cunningham, A. A.; Donald, P. F.; Prakash, N.; Verma, A.; Gargi,
- 643 R.; Sivakumar, S.; Rahmani, A. R., 2003. Catastrophic collapse of Indian white-

	Journal Pre-proof
644	backed Gyps bengalensis and long-billed Gyps indicus vulture populations. Biol.
645	Conserv. 109, 381-390; DOI 10.1016/S0006-3207(02)00164-7.
646	Prakash, V.; Bishwakarma, M. C.; Chaudhary, A.; Cuthbert, R.; Dave, R.; Kulkarni, M.;
647	Kumar, S.; Paudel, K., Ranade, S.; Shringarpure, R.; Green, R. E., 2012. The
648	Population decline of Gyps vultures in India and Nepal has slowed since veterinary
649	use of diclofenac was banned. PLoS ONE. 7, e49118; DOI
650	10.1371/journal.pone.0049118.
651	Soler, F.; Sánchez, S.; Pérez-López, M.; Mateo, R., 2016. Intoxicación por flunixin en buitre:
652	precaución en el laboratorio de diagnóstico toxicológico. Presented at Jornadas de
653	Toxicología Española e Iberoamericana, Sevilla, Spain, 2016; TV-P2, pp 53-54.
654	https://www.aetox.es/wp-content/uploads/2016/05/Libro-Abstracts-Jornadas-
655	Toxicolog-a-2016.pdf
656	Swan, G. E.; Cuthbert, R.; Quevedo, M.; Green, R. E.; Pain, D. J.; Bartels, P.; Cunningham,
657	A. A.; Duncan, N.; Meharg, A. A.; Oaks, J. L.; Parry-Jones, J.; Shultz, S.; Taggart, M.
658	A.; Verdoorn, G.; Wolter, K., 2006. Toxicity of diclofenac to Gyps vultures. Biol.
659	Lett. 2, 279-282; DOI 10.1098/rsbl.2005.0425.
660	Swan, G.; Naidoo, V.; Cuthbert, R.; Green, R. E.; Pain, D. J.; Swarup, D.; Prakash, V.;
661	Taggart, M.; Bekker, L.; Das, D.; Diekmann, J.; Diekmann, M.; Killian, E.; Meharg,
662	A.; Patra, R. C.; Saini, M.; Wolter, K., 2006b. Removing the threat of diclofenac to
663	critically endangered Asian vultures. PLoS Biol. 4, DOI
664	10.1371/journal.pbio.0040066.
665	Swarup, D.; Patra, R. C.; Prakash, V.; Cuthbert, R.; Das, D.; Avari, P.; Pain, D. J.; Green, R.
666	E.; Sharma, A. K.; Saini, M.; Das, D.; Taggart, M., 2007. Safety of meloxicam to
667	critically endangered Gyps vultures and other scavenging birds in India. Anim.
668	Coserv. 10, 192-198; DOI 10.1111/j.1469-1795.2006.00086.x.

	Dra	nro	$^{-1}$
JUUIIIAI		ρισ	U

669	Taggart, M. A.	; Cuthbert,	R.; Das,	D.; Sashikumar,	C.; Pain,	D. J.; Green.	R. E.;	Feltrer,	Y.;
		,		2, , , , , , , , , , , , , , , , , , ,	<i>c</i> , <i>i</i> , <i>m</i>	2.0., 0.000	,	,	-

- 670 Shultz, S.; Cunningham, A. A.; Meharg, A. A., 2007. Diclofenac disposition in Indian
- 671 cow and goat with reference to *Gyps* vulture population declines. *Environ. Pollut.*
- 672 147, 60-65; DOI 10.1016/j.envpol.2006.08.017.
- Taggart, M. A.; Senacha, K. R.; Green, R. E.; Jhala, Y. V.; Raghavan, B.; Rahmani, A. R.;
- 674 Cuthbert, R.; Pain, D. J.; Meharg, A. A., 2007b. Diclofenac residues in carcasses of
- domestic ungulates available to vultures in India. *Environ Int.* 33, 759-765; DOI
- 676 10.1016/j.envint.2007.02.010.
- Taggart, M. A.; Senacha, K. R.; Green, R. E.; Cuthbert, R.; Jhala, Y. V.; Meharg, A. A.;
- 678 Mateo, R.; Pain, D. J., 2009. Analysis of nine NSAIDs in ungulate tissues available to
- 679 critically endangered vultures in India. *Environ. Sci. Technol.* 43, 4561-4566; DOI
 680 10.1021/es9002026.
- Zorrilla, I.; Martinez, R.; Taggart, M. A.; Richards, N., 2015. Suspected flunixin poisoning of
 a wild Eurasian griffon vulture from Spain. *Conserv. Biol.* 29, 587-592; DOI
- 683 10.1111/cobi.12417.
- Zorrilla, I.; Richards, N. L.; Benítez, J. R.; Calvino, M.; Fernandez, I.; Rodriguez, F. 2018
- 685 Case study: Detection of two nonsteroidal anti-inflammatory drugs (NSAIDs) in the
- eggs of captive-reared bearded vultures at a breeding center in southern Spain. J
- 687 *Wildl. Rehabilitation.* 38, 15-27.

688 Figure legend

689

- 690 Fig. 1. Iberian Peninsula (Portugal and Spain) map with the distribution by regions of
- sampled carrions and avian scavengers and the percentage of samples with NSAID residues.
- 692

Journal Prevention

- 1 Table 1. Concentrations of NSAIDs detected in positive carrions in Spain alongside a first-tier risk assessment based on toxicity-to-exposure
- 2 ratios (TERs) in vultures.

Carrion	Species	Year	Region	NSAID	Concentration in carrion (mg/kg)		ETE (mg/kg)	LD ₅₀ or observed LD (mg kg ⁻¹)		Acute	<mark>e TER</mark>	
					Muscle	Liver	<mark>Kidney</mark>	-	min	max	<mark>min</mark>	max
<mark>1</mark>	<mark>Pig</mark>	<mark>2016</mark>	Castile and Leon	Diclofenac	<mark>0.171</mark>	<loq< th=""><th><loq< th=""><th><mark>0.028</mark></th><th>0.1^a</th><th>0.2^a</th><th><mark>3.57</mark></th><th><mark>7.14</mark></th></loq<></th></loq<>	<loq< th=""><th><mark>0.028</mark></th><th>0.1^a</th><th>0.2^a</th><th><mark>3.57</mark></th><th><mark>7.14</mark></th></loq<>	<mark>0.028</mark>	0.1 ^a	0.2 ^a	<mark>3.57</mark>	<mark>7.14</mark>
<mark>2</mark>	<mark>Pig</mark>	<mark>2018</mark>	<mark>Aragon</mark>	Ketoprofen	-	<loq< th=""><th>0.173</th><th><mark>0.028</mark></th><th>1.5^c</th><th><mark>5°</mark></th><th><mark>53.6</mark></th><th><mark>178.6</mark></th></loq<>	0.173	<mark>0.028</mark>	1.5 ^c	<mark>5°</mark>	<mark>53.6</mark>	<mark>178.6</mark>
<mark>3</mark>	Pig	<mark>2016</mark>	Valencia	<mark>Meloxicam</mark>	<loq< th=""><th><mark>0.023</mark></th><th><loq< th=""><th><mark>0.378</mark></th><th>-</th><th>-</th><th>-</th><th>-</th></loq<></th></loq<>	<mark>0.023</mark>	<loq< th=""><th><mark>0.378</mark></th><th>-</th><th>-</th><th>-</th><th>-</th></loq<>	<mark>0.378</mark>	-	-	-	-
<mark>4</mark>	Pig	<mark>2017</mark>	Valencia	<mark>Flunixin</mark>	<loq< th=""><th><mark>0.004</mark></th><th><loq< th=""><th><mark>0.001</mark></th><th>1^b</th><th><mark>4.5^b</mark></th><th>1000</th><th><mark>4500</mark></th></loq<></th></loq<>	<mark>0.004</mark>	<loq< th=""><th><mark>0.001</mark></th><th>1^b</th><th><mark>4.5^b</mark></th><th>1000</th><th><mark>4500</mark></th></loq<>	<mark>0.001</mark>	1 ^b	<mark>4.5^b</mark>	1000	<mark>4500</mark>
<mark>5</mark>	Pig	<mark>2016</mark>	Castile and Leon	<mark>Flunixin</mark>	<loq< th=""><th><loq< th=""><th><mark>0.008</mark></th><th><mark>0.001</mark></th><th></th><th></th><th>1000</th><th><mark>4500</mark></th></loq<></th></loq<>	<loq< th=""><th><mark>0.008</mark></th><th><mark>0.001</mark></th><th></th><th></th><th>1000</th><th><mark>4500</mark></th></loq<>	<mark>0.008</mark>	<mark>0.001</mark>			1000	<mark>4500</mark>
<mark>6</mark>	Sheep	<mark>2018</mark>	<mark>Aragon</mark>	<mark>Flunixin</mark>	, <mark>P</mark>	<mark>27.5</mark>	-	<mark>4.55</mark>			<mark>0.22</mark>	1
<mark>7</mark>	Sheep	<mark>2018</mark>	Aragon	Flunixin	<u> </u>	<mark>0.297</mark>	•	<mark>0.048</mark>			<mark>20.8</mark>	<mark>93.8</mark>

- 3 ETE: estimated theoretical exposure; LD₅₀: median lethal dose (LD₅₀ only available for diclofenac); TER: toxicity exposure ratio; LOQ: limit of
- 4 quantification.
- ^a Swan et al. 2006, ^b Cuthbert et al. 2007, ^c Naidoo et al. 2010b.
- 6

<mark>Bird</mark>	Species	Year	Region	Diagnosis	NSAID	Concentration (mg k		kg^{-1})
					_	Liver	Kidney	Muscle
<mark>1</mark>	<mark>Eurasian griffon</mark>	<mark>2010</mark>	<mark>Aragon</mark>	Traumatism/ Renal degeneration	<mark>Flunixin</mark>	<mark>0.33</mark>	-	-
2	<mark>Eurasian griffon</mark>	<mark>2015</mark>	Madrid	Indeterminate/ Visceral gout	<mark>Flunixin</mark>	<mark>4.91</mark>	-	-
<mark>3</mark>	<mark>Eurasian griffon</mark>	<mark>2017</mark>	Extremadura	Feather disease	<mark>Flunixin</mark>	<mark>0.023</mark>	-	-
<mark>4</mark>	Eurasian griffon	<mark>2018</mark>	Catalonia	Indeterminate/Visceral gout	<mark>Flunixin</mark>	<mark>11.32</mark>	<mark>20.35</mark>	-
<mark>5</mark>	Eurasian griffon	<mark>2015</mark>	Extremadura	Iatrogenic ^a	<mark>Flunixin</mark>	<mark>2.83</mark>	-	<mark>0.44</mark>
<mark>6</mark>	Eurasian griffon	<mark>2018</mark>	Catalonia	Traumatism	Meloxicam	<mark>0.641</mark>	<mark>0.264</mark>	-
<mark>7</mark>	Eurasian griffon	<mark>2018</mark>	Catalonia	Traumatism	Meloxicam	<mark>0.159</mark>	<mark>0.231</mark>	-
<mark>8</mark>	Eurasian griffon	<mark>2019</mark>	Aragon	Traumatism/Pb intoxication	Meloxicam	<mark>2.44</mark>	<loq< th=""><th>-</th></loq<>	-
<mark>9</mark>	Eurasian griffon	<mark>2019</mark>	Cantabria	Indeterminate	Meloxicam	<mark>1.84</mark>	-	-
<mark>10</mark>	Eurasian griffon	<mark>2019</mark>	Aragon	Suspected Pb intoxication	Meloxicam	<mark>1.06</mark>	-	-
<mark>11</mark>	Eurasian griffon	<mark>2019</mark>	Aragon	Suspected Pb intoxication	Meloxicam	<mark>0.887</mark>	<loq< th=""><th>-</th></loq<>	-
<mark>12</mark>	Eurasian griffon	<mark>2019</mark>	Extremadura	Traumatism	Meloxicam	-	<mark>0.829</mark>	-
<mark>13</mark>	Black kite	<mark>2019</mark>	Catalonia	Indeterminate	Meloxicam	<mark>0.033</mark>	<mark>0.046</mark>	-
<mark>14</mark>	Egyptian vulture	<mark>2018</mark>	Castile and Leon	Carbofuran intoxication	Meloxicam	<mark>0.141</mark>	-	-
<mark>15</mark>	Common buzzard	<mark>2018</mark>	Catalonia	Electrocution	Meloxicam	<mark>0.838</mark>	1.245	-

Table 2. Positive cases of NSAIDs with the region of origin, presumptive diagnosis, NSAID concentrations by tissue. 7

LOQ: limit of quantification. ^a Soler et al. (2016). 8

9

10

- 11 Table 3. Estimation of diclofenac treated pigs from intensive production with toxic levels for vultures, number of the corresponding toxic meals
- 12 for vultures and number of vultures killed by diclofenac poisoning.

Study	Treated	Weight of	Total weight of treated	Treated meals available to vultures	<mark>Vultures ki</mark>	lled per year	
	carcasses <mark>carcass (1</mark>		carcasses available to	(80% of mass consumed by vultures,	(F = D x proj	portion killed) ^a	
	available to (B)		<mark>vultures (kg)</mark>	regular meal of 1.2 kg per vulture)	Proportion killed		
	vultures (A)		$(C = A \times B)$	$(D = C \times 0.8/1.2)$	<mark>0.049</mark>	<mark>0.375</mark>	
Present study	12	200	2400	1600	78	600	
AEMPS	77 00	•			4	7	
Green et al.	<mark>55-</mark> 92	200	11122-18430	7415-12287	364-603	2781-4609	

^{a.}The number of dead vultures by feeding on meals containing a diclofenac concentration toxic for vultures was calculated following Green et al. (2016) with the proportion of killed vultures after feeding on a contaminated pig treated 8 h before death were 0.375 and 0.049. These proportions were obtained from the LD50 values of 0.098 and 0.225 mg kg⁻¹ calculated by Swan et al. (2006) from the experimental data of Oaks et al. (2004).

17

18

19



Highlights

- NSAID residues were detected in 3.07% of livestock carcasses in Spain.
- Diclofenac was detected in one livestock carcass (0.44%), but not in avian scavengers.
- Flunixin (1.03%) and meloxicam (2.57%) were found in tissues of avian scavengers. •
- Three Eurasian griffons (0.98%) may have died due to flunixin poisoning. ٠
- Flunixin can pose a risk to vultures and this must be addressed by product labelling.

Declaration of interests

 \boxtimes The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

Journal Prerk