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REVIEW

Multiple mortality events in bats: a global review

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ABSTRACT

- 1. Despite conservation concerns for many species of bats, factors causing mortality in bats have not been reviewed since 1970. Here, we review and qualitatively describe trends in the occurrence and apparent causes of multiple mortality events (MMEs) in bats around the world.
- 2. We compiled a database of MMEs, defined as cases in which ≥ 10 dead bats were counted or estimated at a specific location within a maximum timescale of a year, and more typically within a few days or a season. We tabulated 1180 MMEs within nine categories.
- **3.** Prior to 2000, intentional killing by humans caused the greatest proportion of MMEs in bats. In North America and Europe, people typically killed bats because they were perceived as nuisances. Intentional killing occurred in South America for vampire bat control, in Asia and Australia for fruit depredation control, and in Africa and Asia for human food. Biotic factors, accidents, and natural abiotic factors were also important historically. Chemical contaminants were confirmed causes of MMEs in North America, Europe, and in islands. Viral and bacterial diseases ranked low as causes of MMEs in bats.
- **4.** Two factors led to a major shift in causes of MMEs in bats at around 2000: the global increase of industrial wind-power facilities and the outbreak of white-nose syndrome in North America. Collisions with wind turbines and white-nose syndrome are now the leading causes of reported MMEs in bats.
- 5. Collectively, over half of all reported MMEs were of anthropogenic origin. The documented occurrence of MMEs in bats due to abiotic factors such as intense storms, flooding, heat waves, and drought is likely to increase in the future with climate change. Coupled with the chronic threats of roosting and foraging habitat loss, increasing mortality through MMEs is unlikely to be compensated for, given the need for high survival in the dynamics of bat populations.

INTRODUCTION

Bats number over 1300 species and occur in all continents except Antarctica (Fenton & Simmons 2014). Losses of roosting and foraging habitat and other stressors have led to widespread declines of bat populations (e.g. Mickleburgh et al. 1992, Hutson et al. 2001). Nevertheless, mortality in bats has not been reviewed since the work of Gillette and Kimbrough (1970). Many species of bats are highly gregarious and thus potentially vulnerable to 'die-offs', also referenced as multiple mortality events (MMEs). Additionally, bats are sources of zoonotic viral diseases (e.g. Calisher et al. 2006, Luis et al. 2013). Few viral disease-induced MMEs seem to have been documented in bats (e.g. Messenger et al. 2003, O'Shea et al. 2014), suggesting that many microparasites of bats are low in virulence or do not cause MMEs. However, documentation of die-offs due to any cause may be rare simply because bats are secretive, and thus MMEs due to disease may not be disproportionately uncommon. Here, we review and qualitatively describe trends in the occurrence of MMEs, including those caused by disease, in bats around the world.

METHODS AND SCOPE

We defined an MME as a case in which ≥ 10 dead bats were counted or estimated in a given locality within a maximum timespan of 1 year, and more typically within a few days or a season. We included accounts in which the authors qualitatively estimated the number of deaths (as many, dozens, hundreds, etc.). With few exceptions, we did not include a report unless it involved observations of carcasses. Because Mickleburgh et al. (2009) reviewed the consumption of bats for human food, we did not include observations of bats at marketplaces or as imports, unless the report indicated the number of bats killed over a given period of time at a specific location. We did not include records of bats killed by researchers. We compiled published literature and Internetaccessible reports that included observations of carcasses. We did not solicit unpublished material or personal communications. We searched Web of Science, Internet search engines (e.g. Google), regional mammal summaries, specialised outlets and newsletters (e.g. 'Bats' magazine), and other resources. Our search terms included the union of the terms 'bats' or 'Chiroptera' with mortality, die-offs, disease, epizootics, killing, mass mortality, multiple mortality, and so forth.

We identified nine categories of MMEs, according to the cause of death: 1) intentional killing by humans; 2) biotic factors other than disease (e.g. predation, biotoxins); 3) natural abiotic factors (e.g. weather, floods, fire, volcanism); 4) exposure to environmental contaminants, including pesticides; 5) accidents (e.g. entrapment, impalement, colli-

sions with objects other than wind turbines); 6) collisions with wind turbines; 7) infectious viral and bacterial diseases; 8) the fungal disease white-nose syndrome (WNS); and 9) unexplained causes of death. We also classified MMEs geographically, as occurring in: Africa, Asia, Australia, Europe (including the British Isles, Cyprus, Malta), North America, South America (including Trinidad, Curacao), or in islands (>100 km from continental mainland). Primary sources and detailed information about these MMEs are tabulated as online supporting information (Appendices S1–S9, with supporting literature cited listed separately as Appendix S10). We limit our analyses to descriptive summaries because the literature we compiled has many biases.

RESULTS AND DISCUSSION

We compiled 1180 accounts of MMEs in 152 species of bats in all regions, beginning in 1790 (Table 1 and Fig. 1). Cumulatively, collisions with wind turbines caused the highest number of MMEs (a number biased by regulatory reporting requirements in North America and Europe), followed by MMEs due to WNS (Table 1 and Fig. 2). MMEs attributed to infectious viral and bacterial diseases ranked lowest (Table 1 and Fig. 2).

A notable temporal shift in causes of MMEs in bats took place around 2000 (Fig. 3). From 1790 to 1999, 58% of reported MMEs globally were due to intentional killing by humans (39%) or biotic causes (19%; Fig. 3). From 2000 onwards, 70% of all MMEs were due to collisions with wind turbines (35%) and WNS (35%; Fig. 3). These two latter categories represent new and alarming challenges to bat populations.

Anthropogenic sources (human-caused categories and human-caused MMEs within other categories, e.g. accidents, contaminants) account for 54% of MMEs in all years (593 anthropogenic vs. 514 natural, unexplained cases excluded). WNS in North America was not considered anthropogenic. Comparisons with other mammals suggest that the high proportion of anthropogenic MMEs in bats is a cause for concern. In one analysis, from 1940-2012 'human perturbations' were listed as a cause for 20-25% of die-offs of all animal groups combined, but only 0-25% each decade for mammals with no increase in magnitude through time (Fey et al. 2015, bats not included). In another study, anthropogenic causes accounted for 52% of 2209 MMEs involving 10 or more deaths in 27 species of largeand medium-sized (>1 kg) mammals of North America, but more than half of that fraction was due to managed legal harvests (Collins & Kays 2011).

We present an overview of findings within the major categories of MMEs, highlighting regional differences where pertinent. Reports of MMEs in bats are biased regionally

Table 1. Summary of numbers of mass mortality events (MMEs) reported in bats, by category and region (see Appendices S1–S10 for details and references). The order of magnitude for maximum unadjusted numbers of carcasses documented for the largest MME within each category is given in parentheses, following the number of reports for each region and category

Category	Africa	Asia	Australia	Europe	Islands	North America	South America	Events (n)	Species (n)
Intentional killing (Appendix S1)	11 (104)	20 (104)	13 (104)	21 (104)	50 (10³)	58 (10 ⁴)	32 (10 ⁵)	205 (10 ⁵)	69
, , , ,	5 (10¹)	1 (10¹)	19 (10³)	16 (102)	16 (10%)	40 (10 ³)	10 (10¹)	107 (10 ³)	75
Biotic (Appendix S2)	2 (10.)	(' '	,	16 (10²)	16 (10²)	,	10 (10.)	,	
Abiotic (Appendix S3)	0	$6 (10^3)$	71 (10³)	0	$13 (10^3)$	24 (10 ⁵)	0	114 (10 ⁵)	23
Contaminants (Appendix S4)	0	0	1 (10¹)	27 (10 ⁴)	1 (10 ²)	14 (10³)	0	43 (104)	16
Accidental (Appendix S5)	1 (10¹)	0	8 (10¹)	34 (10 ²)	1 (10¹)	22 (104)	0	66 (10 ⁴)	37
Wind turbines (Appendix S6)	1 (10 ²)	0	2 (10 ¹)	59 (10 ²)	2 (10 ¹)	213 (10 ²)	4 (10 ¹)	281 (10 ²)	41
Viral or bacterial disease (Appendix S7)	1 (10 ²)	1 (10 ¹)	2 (10³)	2 (10³)	6 (10³)	13 (10³)	0	25 (10³)	14
White-nose syndrome (Appendix S8)	0	0	0	0	0	266 (10 ⁴)	0	266 (10 ⁴)	6
Unexplained (Appendix S9)	0	0	3 (10 ²)	30 (10 ³)	$2(10^3)$	38 (10 ⁵)	0	73 (10 ⁵)	20
Totals	19 (104)	28 (104)	119 (104)	189 (10 ⁴)	91 (10³)	688 (10 ⁴)	46 (10 ⁵)	1180	152

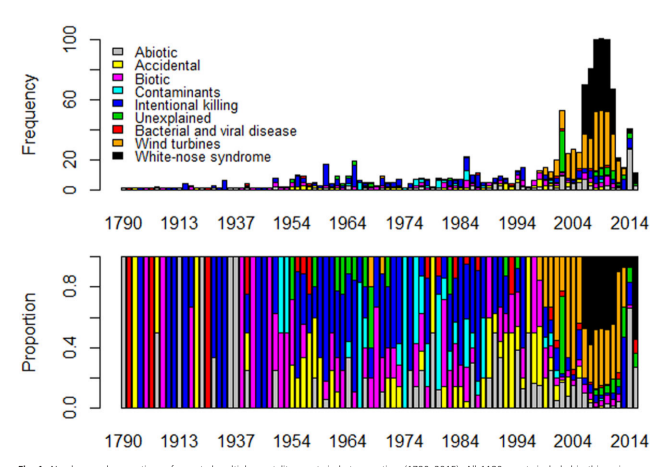


Fig. 1. Numbers and proportions of reported multiple mortality events in bats over time (1790–2015). All 1180 events included in this review are shown.

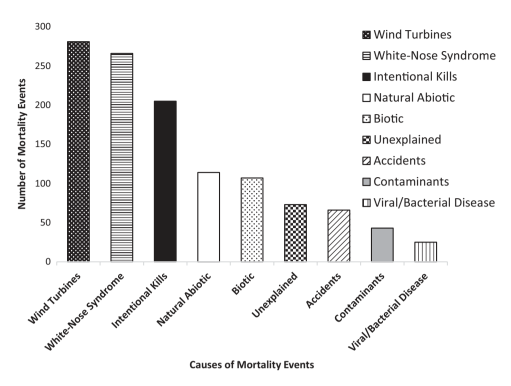


Fig. 2. Cumulative (1790–2015) frequencies of 1180 reported multiple mortality events in bats, by causal category.

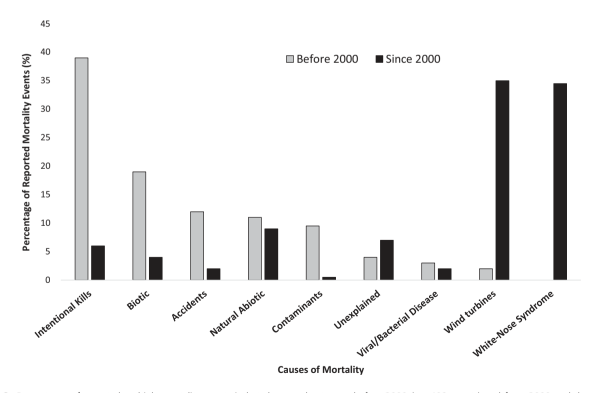


Fig. 3. Percentages of reported multiple mortality events in bats by causal category, before 2000 (n = 409 events) and from 2000 and thereafter (n = 771 events).

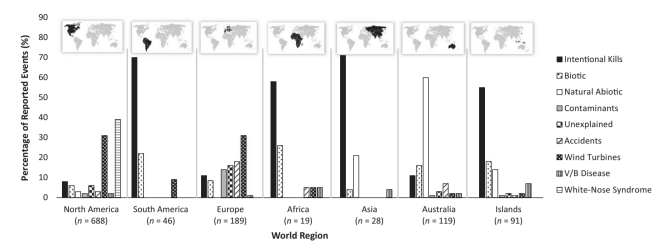


Fig. 4. Causes of multiple mortality events in bats, expressed as percentages of all events (n), for each geographic region.

towards North America, Europe, Australia, and islands; comparatively few events are recorded for Africa, Asia, and South America (Table 1 and Fig. 4). Similar biases are apparent in the reporting of mass mortality for all animals (Fey et al. 2015). We encourage greater reporting and documentation of MMEs in bats globally, perhaps in conjunction with other ongoing bat-monitoring programmes.

MMEs due to intentional killing

Prior to 2000, humans intentionally killing bats caused the greatest number of reported MMEs. These events occurred worldwide and are recorded as early as the 1800s; in total, we documented 205 cases involving millions of individuals of at least 69 species (Table 1 and Appendix S1). Intentional killing was widespread, and continues today, but education and protective legislation have undoubtedly helped reduce its occurrence. Reports of people killing bats for food are common in Asia, Africa, and in islands, but extremely rare in South America, and absent elsewhere (Appendix S1). Reports of people legally and illegally killing fruit bats (ostensibly for crop protection) are common in Australia, and to a lesser extent elsewhere (primarily in the Paleotropics). Killing of bats for crop protection is likely to be under-reported (e.g. Vincenot et al. 2015). In parts of the Americas within the range of common vampire bats Desmodus rotundus, state-sanctioned poisoning of bats and uncontrolled destruction of roosts aimed at reducing common vampire bat bites on livestock has affected thousands of caves and is likely to have killed millions of bats including many non-target species [see later and Appendix S1; dozens of other species share roosts with common vampire bats (Greenhall et al. 1983)]. In addition to the intentional killing of bats for food, crop protection, and common vampire bat control, people have attempted to exterminate colonies by acts of vandalism, including burning, shooting, bludgeoning, and fumigation with poisons (Appendix S1). We provide further summaries of MMEs in this category by region.

AFRICA

We found reports of people intentionally killing bats in Africa dated a century ago (Appendix S1). Lang and Chapin (1917a) noted "Night after night they [caverniculous insectivorous bats in central Africa] return to their accustomed roosts, which they do not abandon even when frequent raids made upon them by the natives have thinned out their numbers. Large, juicy lumps of fat, deposited in and about their abdominal cavity, stimulate the natives to kill all they can." Lang and Chapin (1917b) provide other accounts of people killing apparently large numbers of insectivorous bats from other types of roosts. Deaths of multiple pteropodids (family Pteropodidae) taken for food at specific locations are reported primarily for straw-coloured fruit bats *Eidolon helvum* (Appendix S1), but other species have been reported at markets (Mickleburgh et al. 2009).

AUSTRALIA

Starting in the 1800s, people decimated roosting camps and killed pteropodids for fruit crop protection (Appendix S1; Hall & Richards 2000). A limited number of these MMEs meet our documentation criteria (Appendix S1), however evidence suggests that millions of pteropodids were intentionally killed. Early methods included shooting, netting, trapping, explosives, fire and smoke; recent methods include electrified grids (Martin 2011). This culling was widespread and common, predicated on beliefs that local aggregations of bats consumed commercial fruit and could be controlled,

and that the bats sent 'scouts' to search for ripening crops. Research suggests that such beliefs are largely erroneous. Grey-headed flying foxes Pteropus poliocephalus, for example, exist in large open populations in which major eruptive movements follow the availability of native food (mostly nectar and pollen from eucalypts; see reports in Eby & Lunney 2002). Exclusion nets protect fruit crops but are too expensive for some growers. From 2000, legal protections designated for some species led to heated conflicts over pteropodid conservation and crop protection (Eby & Lunney 2002). Some states are phasing out legal killing, or regulating take by issuing permits. For example, the government of Queensland issued permits to kill 10580 bats of four species in 2013-2014 (Anonymous 2014a). Despite protections, illegal killing continues. We found little information about intentional killing of microchiropteran bats in Australia. In 1965, about 200 long-fingered bats (Miniopterus sp.) were found dead on the floor of a cave with traumatic injuries caused by vandals (Appendix S1).

ASIA

Commercial fruit growers entangled and killed over 1000 pteropodids (of more than one species) in Thailand (Appendix S1). Egyptian fruit bats Rousettus aegyptiacus in Israel were fumigated with ethylene dibromide and lindane, leading to mass mortality and declines in several non-target species of cave-dwelling insectivorous bats. Mortality through fumigation was reported from 1958 to 1985 (Appendix S1) and led to the 'complete extermination' of Geoffroy's myotis Myotis emarginatus and greater horseshoe bats Rhinolophus ferrumequinum at an Israeli nature reserve (Makin & Mendelssohn 1987). Market-hunters killed 10000 frugivorous, nectarivorous, and insectivorous bats every month in one cave in Thailand during the early 1980s. Large numbers of insectivorous as well as pteropodid bats were, and still are, killed for food markets in Laos, peninsular Malaysia, and Nagaland (India; Fig. 5; Appendix S1).

EUROPE

Stebbings (1988) summarised multiple instances of people intentionally killing bats at roosts in Europe from the 1920s through the 1980s. Many thousands of bats were killed by fires, shooting, bludgeoning, gassing, explosives, and direct application of insecticides as poisons. Nine species in three families were subject to this intentional killing in England, Scotland, Norway, Gibraltar, Malta, France, Germany, Cyprus, and the former Yugoslavia (Appendix S1; Stebbings 1988). People directly fumigated roosts with insecticides in Europe, causing MMEs such as the loss of 2000 individuals of *Pipistrellus* sp. at a building in Scotland during 1971 (Stebbings 1988).





Fig. 5. Intentional killing of bats for food. (a) Dead bats on the ground outside a cave in Nagaland, India, where villagers kill several thousand bats as an annual event extending back perhaps 150 years (see Appendix S1; photograph by Pilot Dovih). (b) *Pteropus* species for sale at a market in Manado, Northern Sulawesi, Indonesia (photograph by D.T.S. Hayman).

ISLANDS

In Borneo, hundreds of individuals of the nearly extinct Bulmer's fruit bat *Aproteles bulmerae* were shot in their cave roost for food on a single day, and about 4500 large flying foxes *Pteropus vampyrus natunae* were killed by hunters at a forest patch in 2003 (Appendix S1). Goodman (2006) reported recent food hunts for three species of insectivorous bats at caves in Madagascar, including about 2700 Commerson's leaf-nosed bats *Hipposideros commersoni* killed in a 3-month season (Appendix S1). Currently, a cull targeting tens of thousands of greater Mascarene flying foxes *Pteropus niger* is underway in Mauritius in the belief that the bats depredate fruit crops (Aldred 2015).

NORTH AMERICA

People intentionally killed at least 11 species of North American bats by burning, shooting, bludgeoning, poisoning, and other activities (Appendix S1). Reports of eradication extend over 100 years into the past. For example, "over two washtubfulls of the pesky critters" (presumed Brazilian free-tailed bats *Tadarida brasiliensis*) were destroyed at a Texas building in 1908 (Appendix S1).

Discovery of rabies in North American bats in the 1950s increased public fear of health risks, leading to attempted eradication of entire bat colonies, especially in buildings. Killing was widespread and became commercialised within the pest control industry. Additionally, fear or misunderstanding of bats also motivated killing at colonies, often referred to as 'vandalism'. Some MMEs from deliberate eradication are documented in the literature (Appendix S1). People used chemical control methods (primarily fumigation with dichlorodiphenyltrichloroethane - DDT) to poison colonies of multiple species at buildings in the USA, Canada, and Mexico (Appendix S1). Considering the large number of pest control companies operating in North America, the true number of MMEs from chemical eradication until the 1980s is probably orders of magnitude larger than reported in the literature. Killing bats with chemicals may continue in some parts of North America, but is more restricted and is considered less effective than physical exclusion.

SOUTH AMERICA

In the late 1950s, people used organochlorine pesticides to kill insectivorous bats considered nuisances in houses in Trinidad: 1592 Pallas's mastiff bats Molossus molossus and 339 black mastiff bats Molossus rufus were found dead after up to 15 houses were sprayed (Appendix S1). Killing of bats for vampire control, however, is the most heavily reported cause of MMEs in South America. Common vampire bats have been recognised as an economic and health problem in Latin America for at least a century. These bats consume blood from livestock, reducing animal health and productivity; more significantly, common vampire bats transmit rabies to cattle, which has been estimated to cause tens to hundreds of thousands of livestock deaths annually (Baer 1991). Recently, vampire-bat transmission of rabies to humans has also become recognised as a serious health problem in rural areas (e.g. Streicker et al. 2012). People use various methods to control common vampire bats in response to livestock and human health problems, including lighting or covering corrals, hanging spiny branches around cattle, capturing vampires in livestock corrals or roosts with nets and bat traps, clubbing and shooting bats at roosts, poisoning, setting fires in roosts, and offering bounties for vampire bat carcasses (Constantine 1970, Flores-Crespo & Arellano-Sota 1991). Some control methods at roosts have been indiscriminate, killing many bats of other species, whereas other methods specifically target common vampire

bats. More targeted state-sanctioned killing of common vampire bats now makes use of gels with oral anticoagulants applied to the bodies of common vampire bats captured at ranches; the bats return to roosts, spread the substance within the colony, and ingest the toxic compounds during grooming (including allogrooming; Arellano-Sota 1988, Flores-Crespo & Arellano-Sota 1991, Johnson et al. 2014). Bite wounds on cattle are also treated with anticoagulants that are then ingested by common vampire bats during feeding (Flores-Crespo & Arellano-Sota 1991).

People caused MMEs at common vampire bat roosts using methods such as flamethrowers, dynamite, fumigation with gases and poisons, electrocution, introduction of disease agents, shooting, netting, trapping, and application of anticoagulants (Greenhall & Schmidt 1988, Brown 1994; Appendix S1). Concerned biologists and conservationists have issued resolutions against destroying multi-species roosts for common vampire bat control (Anonymous 1968, 1998) and some governments have banned the destruction of bat roosts on state-owned reserves (e.g. Kikuti et al. 2011). Nonetheless, agencies and private individuals continue to destroy multi-species roosts and misapply anticoagulants to non-target species (e.g. Mayen 2003, Aguiar et al. 2010, Streicker et al. 2012, D. Streicker, personal observation). Recent quantitative analyses suggest that culling vampire bats may be ineffective for controlling bovine rabies (e.g. Streicker et al. 2012, Blackwood et al. 2013, Johnson et al. 2014).

Governments still try to control common vampire bats as part of national rabies control plans (D. Streicker, personal observation) and improving methods of control is an area of active research (Corrêa-Scheffer et al. 2014), but reports chronicling the current extent and effects of these practices on non-target species are limited. Earlier literature suggests a severe impact that may be ongoing. In Venezuela, people indiscriminately dynamited and discharged poisonous organophosphate gas into caves, killing an estimated 900000 bats of multiple species each year from 1964 to 1967 (see Constantine 1970 for references). In 1963–1968, authorities in the state of Rio Grande do Sul, Brazil, indiscriminately killed bats in 8240 caves (Constantine 1970). In an unusual approach in Colombia, a cave was fumigated with atomised Newcastle's disease virus (a Paramyxovirus of poultry), and an estimated 5000 common vampire bats were later found sick or dead, apparently from the virus (Constantine 1970).

MMEs due to biotic factors other than disease

MMEs caused by biotic factors other than disease principally involve predation; however, impalement on burrs of plants, paralysis from tick neurotoxins, and poisoning by a toxic algal bloom were also reported (Appendix S2). We tabulated 107 MMEs beginning in the 1890s, involving over 75 species of bats globally (Table 1 and Appendix S2). Bats have numerous predators, nearly all opportunistic, and the literature includes many anecdotal accounts of individual cases (Gillette & Kimbrough 1970). However, MMEs involving 10 or more bats taken as prey at specific locations have been reported for well over a century and have occurred globally. The predators known to cause MMEs in bats include invertebrates, snakes, birds, and mammalian mesocarnivores (Appendix S2). Birds are most widely cited as predators of bats globally, and may be a selective force for nocturnality (Mikula et al. in press). Two species of raptor specialise on bats, one in the Paleotropics (Macheiramphus alcinus) and one in the Neotropics (Falco rufigularis). These specialists hunt at dusk, particularly during emergence of bats, but rely on other prey as well. Predation by birds can be substantial globally, but estimates are unavailable. The author of the most extensive regional study estimated that avian predation in the British Isles accounted for over 200000 bat deaths per year (Speakman 1991). In some areas, predation by house cats also can be extensive: in one study it was estimated that cats kill 170000 bats annually in Britain (Appendix S2).

MMEs due to natural abiotic factors

Unseasonably cold weather, snow storms, flooding of roosts during heavy rainfall, overheating during unusual hot spells, burning or suffocation during landscape-level fires, and volcanic eruptions all cause MMEs in bats (Appendix S3). MMEs due to such abiotic factors have been known since the 1790s. Reports are unevenly spread globally: most observations are made in Australia, North America, and in islands. We documented 114 MMEs due to natural abiotic factors involving 23 species (Table 1), but found no published reports of such events in Africa, Europe, or South America.

ASIA

MMEs in pteropodid bats were reported at several locations in India as a result of unusually hot periods during 2010 and 2015 (Appendix S3).

AUSTRALIA

Deaths of many thousands of pteropodid bats due to extremely hot spells were reported in eastern Australia as recently as 2014 (Appendix S3). Ambient temperatures exceeded 48 °C at numerous locations during the 2014 event (Anonymous 2014b). Large numbers of flying foxes died during presumably lesser heat spells in 1790 and

during the early 1900s (Appendix S3), and many thousands of grey-headed and black flying foxes Pteropus alecto died during 2000–2007, presumably from excessive heat (Appendix S3; Welbergen et al. 2008). The main cause of these MMEs is hyperthermia. Welbergen et al. (2008) found that ambient temperatures of ≥42 °C were associated with lethal hyperthermia; up to 13% of the black flying foxes in one large, well-studied colony died from hyperthermia, the majority being dependent young. These MMEs took place during the summer when bats are in optimal body condition. MMEs from hyperthermia are expected to increase in number and severity with climate change (Welbergen et al. 2008). Droughts and resulting bush fires often accompany extreme heat, and have also killed 'many' flying foxes through smoke inhalation, heat exposure, or immolation (Hall & Richards 2000). Droughts also reduce food supplies and cause MMEs in grey-headed flying foxes through starvation (Appendix S3). At the other abiotic extreme, over 1000 flying foxes (species unspecified) died in northeastern New South Wales and southeastern Oueensland in 1990 and 1991 during wet, windy, cold winter weather when food supplies were low (Hall & Richards 2000).

ISLANDS

Major storms (hurricanes, typhoons, cyclones; Appendix S3) have caused MMEs in bats in islands. These MMEs primarily involved pteropodid bats, including Pacific flying foxes Pteropus tonganus in the Vava'u islands of Tonga in 2001, Pacific flying foxes and and Solomons flying foxes Pteropus rayneri in the Solomon Islands in 1986, Pacific flying foxes and Samoan flying foxes Pteropus samoensis in Tutuila, American Samoa in 1990 and 1991, Rodrigues flying foxes Pteropus rodricensis in Rodrigues, Mauritius in 1979, and greater Mascarene flying foxes in Réunion during 1960 and 1979 (Appendix S3). In many of these cases, some of the mortality following severe storms was delayed and attributable to lack of food caused by winds stripping vegetation, and to opportunistic hunting by people (e.g. Cheke & Dahl 1981, Appendix S3). Such MMEs are exacerbated by increased vulnerability to hunting due to lack of concealing vegetation, and by bats being forced to forage in daylight on the ground where they also are vulnerable to predation by dogs, cats, and pigs (Pierson & Rainey 1992, Stinson et al. 1992).

One report confirmed storms causing an MME of insectivorous bats in islands: flooding from Cyclone Hyacinthe in 1980 drowned over 3000 Mauritian little mastiff bats *Mormopterus acetabulosus* in a cave in Réunion (Cheke & Dahl 1981). Hurricanes in Puerto Rico in 1989 and 1994 preceded dramatic declines in populations of the red figeating bat *Stenoderma rufum*, but no carcasses were reported (Gannon & Willig 1994). Similarly, marked

decreases and changes in relative abundance among frugivorous species of bats followed the 1989 hurricane in Montserrat in the Caribbean (Pedersen et al. 2009).

Eruptions have caused MMEs in bats in volcanic islands. Direct mortality of multiple individuals of Seychelles flying foxes *Pteropus seychellensis* was observed in Grande Comore in the Comoros Islands in the Indian Ocean (Appendix S3). Volcanic activity may also have caused MMEs in pteropodid bats in some Pacific islands (Lemke 1992). Beginning in 1995, volcanic activity in Monserrat in the Caribbean "reduced the eastern and western flanks of the volcano to an ecological wasteland and have buried much of the southern half of the island under varying amounts of volcanic ash" (Pedersen et al. 2009). Volcanic action destroyed roosts in Monserrat, and frugivorous bat abundance decreased; although no mortality was directly observed and no species were lost, survivors showed signs of sublethal pathologies (Pedersen et al. 2009, 2012).

NORTH AMERICA

Flooding of caves during unusually rainy weather or spring runoff has caused MMEs in bats in North America. Floods in the Ohio River valley, USA, in 1964 led to declines in four species of bats at one hibernaculum in Kentucky, which dropped from about 6000 bats in February to about 500 in March after the flood; numerous carcasses were found trapped among flood debris and mud after the waters receded (Appendix S3). Similarly, about 6500 dead southeastern myotis Myotis austroriparius were observed awash in one Florida cave in 1989 following a summer downpour, an estimated 57000 individual southeastern myotis died in a second cave during record high water in 1990, and flooding in 1994 killed 85000 individual southeastern myotis in Snead's Cave, Florida (Appendix S3). An estimated 10000 grey bats Myotis grisescens were found dead after flooding in a Tennessee cave in 1970 (Appendix S3). Based on skeletal deposits, an estimated 300000 Indiana bats Myotis sodalis died from past flooding at Bat Cave, Kentucky, thought by some to be a major flood in 1937 (Appendix S3). MMEs in Indiana bats caused by flooding at other caves involved 3000 individuals in 1997, and hundreds at two caves in 1996 (Appendix S3).

Despite the thermal stability of caves and mines used by bats for hibernation, extremely cold weather has caused MMEs at hibernacula in North America. Subfreezing temperatures killed 200 individual Indiana bats at a hibernaculum in Indiana in 1977, and "large numbers" at a hibernaculum in Missouri (Appendix S3). Over 100 big brown bats *Eptesicus fuscus* were found dead in snow drifts formed by an early winter storm that blocked access to hibernacula in Minnesota in 1940 (Appendix S3). Hundreds of little brown bats *Myotis lucifugus* were found dead or

dying in the streets of a Wisconsin town in autumn 1936, and were thought to have become exhausted while migrating in association with a cold front (Appendix S3). In the Mojave Desert of California, over 40 Brazilian free-tailed bats attempting to drink in a normally ice-free pond were entrapped and frozen at the surface during an unusual cold snap in 1930 (Appendix S3).

MMEs due to exposure to environmental contaminants

Chemical contaminants caused MMEs in bats in Australia, Europe, New Zealand and North America, and at a global scale are suspected of causing far more mortality than has been clearly documented. We compiled 43 accounts of MMEs caused by chemical contaminants in 16 species of bats since 1952; organochlorine pesticides such as DDT were the most prevalent agents (Appendix S4). Worldwide use of many organochlorines markedly declined in recent decades due to legislative action and international treaties. Impacts of substitute chemicals on bats have not been adequately investigated (e.g. O'Shea & Johnston 2009).

Organochlorine insecticides and their metabolites can persist in the environment and may become concentrated in fatty tissues of insects and the bats that eat them. Organochlorines may reside in the bodies of bats with no overt effects until stored fats are metabolised, whereupon the chemicals increase in brains until they reach lethal, neurotoxic concentrations. Research has established threshold concentrations in bat brains that are diagnostic of lethality, and correlated concentrations measured in carcasses or guano also indicate lethal exposure (Clark & Shore 2001, O'Shea & Johnston 2009). Although circumstantial evidence and simple presence of chemical residues in carcasses are not truly diagnostic of this cause of death, some die-offs due to unexplained causes (Appendix S9) were very likely to have been caused by chemical contaminants. MMEs described below and in Appendix S4 pertain to poisoning through the food-web or other environmental exposures, and to unintended consequences of pesticide treatments of timber at roosts in buildings. Cases where chemicals were applied directly to bats or their roosts for 'pest control' are summarised under intentional killing (Appendix S1). We found no conclusive evidence that chemical contaminants directly caused MMEs through food-chain exposures in Africa, Asia, or South America, although organochlorines were applied much more recently in these regions.

AUSTRALIA

About 30% of the flying foxes (three species, Appendix S4) found dead in urban areas of Brisbane during the 1980s had histopathological lesions and concentrations in tissues indi-

cating exposure to lethal amounts of lead, likely as an atmospheric pollutant. Investigators hypothesised that insecticide exposure caused die-offs of long-fingered bats in the 1960s, but reported no corroborating diagnostic evidence (Appendix S4).

EUROPE

Treating of timbers in buildings with organochlorine compounds (DDT, dieldrin, lindane, pentachlorophenol) as protection against wood-boring insects, which took place from about 1950 to the 1980s, is thought to have been "the most important factor in killing bats and reducing breeding success" in Europe (Stebbings & Griffith 1986). First documented in the literature during the early 1970s (Appendix S4), timber treatment occurred on a massive scale; over 1500 private companies were directly involved in Britain alone during the 1980s (Mitchell-Jones et al. 1989). Many species of bats in Europe are highly dependent on woodenframe buildings for roosts. Organochlorine compounds were used in hundreds of thousands of such buildings, and remained available for uptake by roosting bats through contact, ingestion while grooming, and inhalation of vapours for years after application (e.g. Shore et al. 1990). Captive bats experimentally exposed to treated timbers died (reviewed in Clark & Shore 2001), and such uptake has been implicated in killing nine species of bats in the wild (Appendix S4). We are unaware of reports of food-web exposure to environmental contaminants causing MMEs in Europe. Researchers detected potentially toxic concentrations of lead in pipistrelles Pipistrellus sp. in England, but not definitively in association with MMEs (Appendix S4).

ISLANDS

The anti-coagulant rodenticide diphacinone caused the death of at least 115 New Zealand lesser short-tailed bats *Mystacina tuberculata* in North Island, New Zealand; presumably it was ingested from prey or while ground-foraging (Appendix S4).

NORTH AMERICA

Dieldrin (or the parent compound aldrin) clearly caused MMEs through food-web exposure (including juveniles nursing contaminated milk) in grey bats in the USA during the 1970s (Appendix S4). Grey bats roost in caves where carcasses can be easily found. Although the full extent of organochlorine-caused die-offs is unknown, lethal concentrations of dieldrin, DDT, endrin, heptachlor, and their metabolites have also been found in Indiana bats (Appendix S4) and possibly Brazilian free-tailed bats (Reidinger 1976). Similarly, exposure to DDT and its metabolites have been linked to major declines in populations of Brazilian

free-tailed bats in the southwestern USA (Geluso et al. 1976, Clark 2001), but these data were not obtained from carcasses found during MMEs. The role of other classes of insecticides in causing MMEs is more difficult to determine, but has been long suspected. For example, Brazilian free-tailed bats were found dead in Arizona agricultural fields where organophosphate insecticides were applied near a large colony that suffered a major contemporaneous decline (Clark & Shore 2001; Appendix S9). Carbamate poisoning was recently reported in bats in Idaho (Appendix S4).

Mining operations that utilise cyanide for gold extraction and then store contaminated waters in ponds on site attract bats to drink. MMEs in bats at these ponds due to cyanide poisoning have been recorded in South Carolina, Arizona, and Nevada (Appendix S4), and are likely to be more prevalent globally than documented in the literature.

MMEs due to accidents

Since at least 1906, accidental deaths of multiple bats have been reported from Africa, Australia, Europe, the Seychelles Islands, and North America (Appendix S5), and stem from both natural and anthropogenic sources. We tabulated 66 MMEs involving 37 species of bats. 'Natural' accidental mortality includes entrapment of bats in buildings (e.g. Dietz et al. 2009), entrapment of lasiurine bats (Genus Lasiurus; not normally cavernicolous) that cannot find their way out of large caves in North America, first noted in 1907 (Appendix S5), and odd accidents such as being crushed by falling trees (Appendix S5). Accidental falls from roost ceilings probably caused mortality of juvenile bats of several species in North America (Appendix S5). Such MMEs involved about 13000 young in one season in dense populations of Brazilian free-tailed bats in Texas caves, and as few as 36 per season for grey bats in Kentucky (Appendix S5). This mortality is strongest in the first days of life (Foster et al. 1978, Hermanson & Wilkins 1986), and fallen young are sometimes eaten by scavengers. Some young bats may fall from weakness due to disease, malnutrition, or pesticide poisoning, but ultimate causes are generally unknown.

Human-caused MMEs from accidents (Appendix S5) include electrocution of flying foxes on utility wires in Australia and in islands, incidental demolition of buildings with roosts (Europe), collisions with aircraft (Australia), and collisions with motor vehicles on roadways (Europe, North America). Collisions with motor vehicles may be a large, but widely unrecognised mortality factor for bats. Globally there are over 35 million km of roadways (Anonymous 2015); most reported MMEs on roads have been from studies in Europe that covered only about 150 km of roadway in total, yet revealed deaths in 26 species of bats (Appendix S5).

MMEs due to collisions with wind turbines

A recent and unexpected form of MMEs in bats is associated with the global expansion of industrial wind energy production. Multiple fatalities have been reported from wind turbines in North America, Europe, South America, Africa, and Australia, most during the past decade (Appendix S6). Wind turbines are increasing globally and MMEs in bats are likely to occur at most facilities, but the majority of available reports are limited to Europe and North America. We tabulated 281 MMEs involving 41 species; some carcass counts numbered in the hundreds (Table 1 and Appendix S6). This cause of MMEs is very recent (Figs 1 and 3), widespread, and growing rapidly. Numbers of deaths vary among sites for unknown reasons (Arnett et al. 2008). However, estimates that include bias corrections (see Appendix S6) range to thousands of bat deaths annually at some facilities (Appendix S6). Cumulative deaths of bats at turbines tabulated for Europe for the period 2003-2013 involved 5626 bats of 27 species in 18 countries (Rodrigues et al. 2014), only a fraction of the likely mortality. Most deaths of bats at wind turbines in temperate latitudes occur during late summer and autumn, and disproportionately affect migratory species that roost in trees (Cryan & Barclay 2009, Arnett & Baerwald 2013). In some regions, deaths of some species at wind turbines far exceed other known sources of mortality (Cryan 2011). Causes of susceptibility to wind turbines are not fully understood, but some bats seem attracted to them (Cryan et al. 2014).

MMEs attributable to or suggestive of viral or bacterial diseases

Despite the high number of viruses known to infect bats (e.g. Calisher et al. 2006, Luis et al. 2013), MMEs attributable to infectious viral or bacterial diseases are rarely reported. The absence of epizootics in bats has been noted as remarkable by field researchers over many years. In his review of Australian pteropodid ecology, Ratcliffe (1932) stated: "No reliable evidence of the occurrence of epidemics among the fruit-bat population was discovered." In studies of the southeastern myotis Myotis austroriparius, Rice (1957) remarked "Disease is apparently unimportant...During the course of this study, which involved observations on over a million bats in every known cave colony in Florida, I have never found a dead bat, and have seen only one which appeared diseased." In extensive research on cavernicolous bats in North America, Twente (1955) concluded "it would not seem probable that disease is an important limiting factor." Similarly, in ecological studies of Brazilian free-tailed bats in Texas, which form the largest aggregations of mammals on Earth, Davis et al. (1962) observed that "Better conditions for epizootic spread

of disease can hardly be imagined, yet we did not observe anything that looked like epizootic disease".

We found 25 MMEs in 14 species of bats (Table 1) attributable to or suggestive of disease other than WNS, many without confirmatory evidence of a specific agent or with unexplained components (Appendix S7). MMEs in this category involved maximum numbers in the thousands, included some of the earliest (1839) reported die-offs of bats, and occurred in Africa, Asia, Australia, Europe, North America, and in islands (Appendix S7). Nine MMEs involved pteropodid bats (six in islands). Brazilian free-tailed bats accounted for seven of the 13 cases in North America (Appendix S7). One event in North America provided supporting evidence (bacterial isolation and pathology) for a bacterial agent (Pasteurella multocida) killing about 100 individual big brown bats during a 4-week period at a roost in Wisconsin, USA (Blehert et al. 2014). With the exception of rabies, few of the other MMEs had strong confirmatory evidence of a causal organism; only three non-rabies reports suggested a specific agent. A Bunvavirus was implicated in deaths of wrinklelipped free-tailed bats Chaerephon plicatus in Cambodia, Lagos bat virus (a rabies-like lyssavirus) was found in about 10-15% of several hundred dead Wahlberg's epauletted fruit bats Epomophorus wahlbergi examined in South Africa (Appendix S7), and the recently identified Lloviu filovirus was found in carcasses of Schreiber's long-fingered bat Miniopterus schreibersii that were sampled during MMEs involving tens to hundreds of bats at two locations in Spain (Appendix S7). In this latter case, the finding of the filovirus is possibly incidental and the responsible cause remains unconfirmed (Olival & Hayman 2014). Multiple roosts in Spain, France, and Portugal were also subject to MMEs contemporaneously, but these populations were not extensively sampled diagnostically, and pathological and virological results were inconclusive (Appendix S9).

Rabies is enzootic in populations of Brazilian free-tailed bats, and given the huge size of some of these colonies, it is not unexpected that deaths of 10 or more bats from rabies have been reported within a season at roosts (Appendix S7). These deaths from rabies, however, are not of epizootic proportions (e.g. Davis et al. 2012 in Appendix S7). MMEs in Brazilian free-tailed bats during the 1950s at Carlsbad Caverns were equivocal: although > 10 individuals had died of rabies each year, others may have died of pesticide poisoning (Clark 2001), whereas inclement weather and abnormally cool conditions may have been associated with many of the deaths in 1955 and 1956, but not in 1957 (Appendix S7 and references therein). Constantine (1967) postulated that inclement, cool weather during periods of migratory stress contributed to or perhaps caused the MMEs at Carlsbad Caverns, as well as some unexplained MMEs in this highly migratory species elsewhere (Appendix S9). The Old World Schreiber's long-fingered bat is also

a migrant that gathers in large colonies, and similarly has been reported to suffer mortality that is unexplained or suspected to be due to pesticides, migratory stress, or inclement weather (Appendices S3 and S9).

MMEs suggested to be due to disease affected large numbers of pteropodid bats in islands (Appendix S7). Perhaps disease is more likely to kill pteropodids in immunologically naïve populations in islands, but corroborating evidence for any infectious agents remains slim. In two MMEs, disease was suspected because the die-off was contemporaneous with epidemics in humans (measles and dysentery). In a third case, subsequent authors suggested alternatively that invasive ants may have been a causal factor (Appendices S2 and S7). Two events involving mass abortions of pteropids in Australia (Appendix S7) may have been due to disease, but there was no confirmatory evidence, and other possible causes (malnutrition, weather, non-infectious agents) were not ruled out. However, as in other cases, availability and application of diagnostic tools for diseases of bats has been limited.

MMEs attributable to WNS

WNS is a fungal disease recognised in bats within the past decade in North America. WNS is the only epizootic disease known to cause widespread and high mortality in multiple species of bats over multiple years. First documented at a cave in New York, USA, during 2006, WNS has spread in subsequent winters to affect most species of hibernating bats in eastern North America (Turner et al. 2011). This ongoing epizootic has killed millions of bats and is affecting six or more species (Turner et al. 2011; Appendix S8); important populations are in serious decline due to WNS (Frick et al. 2010a, Thogmartin et al. 2013).

WNS is caused by the cold-growing fungus *Pseudo-gymnoascus destructans*, which severely infects the skin tissues of hibernating bats and catastrophically disrupts hibernation and physiology during winter (e.g. Lorch et al. 2011, Warnecke et al. 2012). The fungus also infects bats in Europe, where it is not known to cause MMEs (Puechmaille et al. 2011). Macroecological analysis supports a hypothesis that more severe WNS mortality in Europe may have occurred much earlier than in North America (Frick et al. 2015). WNS continues to spread across North America, and will probably continue to cause MMEs in additional species and regions.

Unexplained MMEs

Seventy-three MMEs in 20 species of bats in Australia, Europe, North America, and in islands lacked causal evidence (Table 1 and Appendix S9). Nearly all of these MMEs involved species that depend strongly on caves as roosts.

Two MMEs in Peter's ghost-faced bat *Mormoops megalophylla* during the 1950s in Mexico and cave myotis *Myotis velifer* in the USA were mysterious; thousands of mummified remains covered the walls and floors of caves (Appendix S9). Other cases occurred in the 1960s during peak usage and impacts of organochlorine insecticides; pesticide poisoning may have caused some of these MMEs, as previous authors speculated (Appendix S9). Forty-one unexplained MMEs involved migratory species (Brazilian free-tailed bats, Schreiber's long-fingered bat, mouse-eared myotis *Myotis myotis*); some researchers suggest migratory stress as a possible cause (Appendix S9).

Bat population dynamics, MMEs, and implications for the future

Recent studies indicate that population dynamics of bats may be particularly sensitive to mortality. These studies primarily involved temperate-zone species, but similar dynamics are seen in large tropical pteropodids (Hayman et al. 2012). A recent brief review noted the similarity in the relative importance of life history traits of bats to those of many large mammals: for the size of a bat population to remain stable or to increase, annual survival of adults must be relatively high: > 75-80% (O'Shea et al. 2011a). Population growth rates of bats show greater sensitivity and elasticity to adult survival than to reproduction or juvenile survival. This suggests that, compared with that of other small mammals, the demography of bats is adapted to a narrow range of mortality drivers, such as the natural biotic and abiotic factors, disease, and natural accidents documented in this review. These natural factors (with the recent exception of WNS) may operate more diffusely than some anthropogenic factors. The magnitude of mortality due to past intentional killing, recent incidence of WNS, and increases in levels of accidental mortality from collisions with wind turbines are likely to be additive; we doubt that bat populations can sustain such additive mortality for long. Furthermore, future climate change may increase the frequency of MMEs through severe weather events, such as extreme droughts, more frequent storms, and flooding (examples in Appendix S3). Recent researchers (see Herring et al. 2014) have concluded that climate change drove the high temperatures in Australia in 2013 and 2014 that killed hundreds of thousands of flying foxes (Appendix S3). Population growth can also be negatively impacted through the suppressed reproduction and reduced juvenile survival of bats seen during periods of drought (Adams 2010, Frick et al. 2010b, O'Shea et al. 2010, 2011b).

CONCLUSIONS

Many MMEs in bats have been reported over the years. Of the nine potential causes that we differentiated, intentional killing by people caused the greatest proportion of MMEs prior to 2000. People killed bats because they were considered sources of zoonotic disease, nuisances (e.g. bats that roosted in buildings), or, in Australia and Asia, competitors for fruit crops. People still kill and eat both insectivorous and pteropodid bats in Asia, Africa, and in some islands. Efforts to control bovine rabies transmitted by common vampire bats in South America and southern North America led to indiscriminate killing of non-target cavernicolous bat species that continues to the present. Prior to 2000, about 11% of the reported MMEs were attributed to natural abiotic factors. Projected extreme weather due to continuing climate change (e.g. severe storms, flooding, and drought) may increase the number of abiotic MMEs in the future.

Two new causes of MMEs have taken precedence since around 2000: death due to collisions with wind turbines globally, and the fungal disease causing WNS in eastern North America. Reports of MMEs due to these two causes will probably soon outnumber all prior reports from all categories combined. Among all categories, MMEs due to viral or bacterial diseases were most rarely reported. Unexplained MMEs were not very common. This supports the hypothesis that many microparasitic infections of bats do not result in MMEs.

We believe that the life history attributes of bats historically allowed populations to compensate more easily for natural causes of mortality. Intentional killing by humans and very recent increases in mortality from other anthropogenic sources has put markedly greater pressures on many populations of bats. Bats globally could benefit from policy, education, and conservation actions targeting human-caused mortality. Such actions are particularly important in the face of the new and increasing threats of the 21st century.

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SUPPORTING INFORMATION

Additional supporting information may be found in the online version of this article at the publisher's web-site:

Appendix S1. Reports of multiple bat deaths due to intentional killing by humans, including counts of dead bats taken for use as food.

Appendix S2. Reports of multiple bat deaths due to biotic factors other than infectious disease.

Appendix S3. Reports of multiple bat deaths due to natural abiotic factors.

Appendix S4. Reports of multiple bat deaths due to environmental contaminants, including pesticides.

Appendix S5. Reports of multiple bat deaths due to accidents other than collisions with wind turbines.

Appendix S6. Reports of multiple deaths due to fatal interactions with the blades of industrial wind turbines.

Appendix S7. Reports of multiple bat deaths due to or suggestive of infectious viral or bacterial disease.

Appendix S8. Reports of multiple bat deaths due to the fungal agent of white-nose syndrome, *Pseudogymnoascus destructans*.

Appendix S9. Reports of multiple bat deaths due to unexplained causes.

Appendix S10. Literature cited for Appendices S1-S9.