Proximity sensors reveal social information transfer in maternity colonies
of Common noctule bats
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#### 26 Summary

- Bats are a highly gregarious taxon suggesting that social information should be readily
   available for making decision. Social information transfer in maternity colonies might
   be a particularly efficient mechanism for naïve pups to acquire information on
   resources from informed adults. However, such behaviour is difficult to study in the
   wild, in particular in elusive and small-bodied animals such as bats.
- The goal of this study was to investigate the role of social information in acquiring
  access to two types of resources, which are crucial in the life of a juvenile bat: suitable
  roosting sites and fruitful feeding grounds. We hypothesized that fledging offspring
  will make use of social information by following informed members of the social
  groups to unknown roosts or foraging sites.
- In the present study we applied for the first time the newly developed miniaturized
  proximity sensor system 'BATS', a fully automated system for documenting
  associations among individual bats both while roosting and while on the wing. We
  quantified associations among juveniles and other group member while switching
  roosts and during foraging.
- 42 4. We found clear evidence for information transfer while switching roosts, mainly
  43 among juveniles and their genetically identified mothers. Anecdotal observations
  44 suggest intentional guidance behaviour by mothers, indicated by repeated commuting
  45 flights among the pup and the target roost. Infrequent, short meetings with colony
  46 members other than the mother indicate local enhancement at foraging sites, but no
  47 intentional information transfer.
- 5. Our study illustrates how advances in technology enable researchers to solve longstanding puzzles. Miniaturized proximity sensors facilitate the automated collection of
  continuous data sets and represent an ideal tool to gain novel insights into the
  sociobiology of elusive and small-bodied species.

52

## 53 Key-words

54 Evolved signals, foraging, information centre hypothesis, local enhancement, maternal care,

- 55 maternal guidance, Nyctalus noctula, roost switching
- 56

## 57 Introduction

58 The early development is a critical phase for an animal since it paves the way for later life by 59 affecting survival rate and overall fitness. It has been shown for several species that limited 60 access to food resources within the first weeks after birth has a negative impact on 61 reproductive success (Lindström 1999; Lummaa & Clutton-Brock 2002). However, a central 62 question is: how do offspring get access to resources in early life once they start to become 63 independent? Behaviours related to foraging might be genetically predefined, which is the 64 case in many invertebrates but also in vertebrate taxa (van Schaik et al. 2017). Over time, 65 'personal information' which is acquired by direct interaction with the environment (Dall et 66 al. 2005) or individual learning by trial and error augment an individual's capabilities (van 67 Schaik et al. 2017). If animals are born and raised in presence of their parents or other 68 conspecifics, information may be socially acquired (Galef Jr & Giraldeau 2001). Adopting 69 information from group members may be more efficient and less costly than individual 70 learning and contributes to an individual's behavioural flexibility under changing conditions 71 (Dall et al. 2005; van Schaik et al. 2017).

Information obtained within colonies or groups of conspecifics enables better decisions in various contexts such as predator avoidance, reduction of parasitism, habitat choice and foraging (Evans, Votier & Dall 2016). The mechanisms of acquiring social information may vary widely in complexity. Individuals may use 'inadvertent social information', which is generated by social cues of conspecifics, i.e., eating may inform about the location of food or

77 fleeing about the presence of a predator (Dall *et al.* 2005). Such public information is created 78 non-deliberately and in group-living animals it may be difficult to hide certain information, 79 e.g., on foraging success. This might particularly apply for breeding colonies, where parents 80 must return to their young and may inadvertently inform others on foraging success via, e.g., 81 time of arrival or fatness (Evans, Votier & Dall 2016). If information is provided deliberately, 82 'evolved signals' are used to actively exchange information (Dall et al. 2005). Black-capped 83 chickadees, e.g., broadcast alarm calls which contain information on the presence and even 84 the size of a predator (Templeton, Greene & Davis 2005) and honey bees use the waggle 85 dance to inform conspecifics on the location of a food source (Leadbeater & Chittka 2007). 86 The ready availability of information at communal roosts gave rise to the Information Center 87 Hypothesis (ICH), which states that such assemblages primarily evolved for the efficient exploitation of unevenly distributed food sources (Ward & Zahavi 1973). According to the 88 89 ICH colony members must assess the success of returning foragers, which are later followed 90 to food patches. However, information can also be generated right at the resource, a 91 mechanism termed 'local enhancement'. Black-browed albatrosses, e.g., indirectly detect food 92 patches by approaching aggregations of foraging predators over sea (Grünbaum & Veit 2003). 93 Both advertent and inadvertent information can be used by juveniles to find food. Juvenile 94 rats prefer feeding sites where adults are present and scent marks and trails of adults cause 95 juveniles to explore such sites (reviewed by Galef Jr and Giraldeau (2001)). Juveniles, since 96 they are naïve, seem to particularly rely on acquiring social information from more 97 experienced individuals of the group like their parents (termed 'vertical transmission') or 98 other adults ('oblique transmission') (van Schaik et al. 2017).

99 Bats are an ideal taxon to study the mechanisms of social information use in groups since the 100 vast majority of species is gregarious and long-lived (Wilkinson & Boughman 1999; Kerth 101 2008; Smith, Lacey & Hayes 2017). However, there is surprisingly little known on whether 102 and how juveniles benefit from social information provided by group members. An interesting

103 case of information transfer across generations is reported for greater sac-winged bats. Here, 104 vocal development of pups is influenced by imitation of territorial songs of harem males and 105 leads to a group signature which is independent of relatedness (Knörnschild et al. 2010; 106 Eckenweber & Knörnschild 2013). When it comes to learning where to roost and where to 107 forage, knowledge on juveniles becomes scarce and existing literature focusses on horizontal 108 information transfer among adult peers. Empirical studies in the wild demonstrated that 109 several insectivorous species are attracted to feeding buzzes by conspecifics which may be 110 used as a signal of foraging success (Gillam 2007; Dechmann et al. 2009; Cvikel et al. 2015). 111 Female greater spear-nosed bats coordinate their foraging bouts at the day roost by screech 112 calls (Wilkinson & Boughman 1998). However, neither local enhancement nor recruitment at 113 the roost has been demonstrated in foraging juveniles, so far. Spatial association among home ranges of mothers and offspring in at least three species and simultaneous feeding of mother-114 115 pup pairs in vampire bats suggest that vertical information transfer, possibly in form of 116 following behaviour, might provide juveniles with insights on where to forage, but this 117 mechanism has yet has to be demonstrated (Wilkinson 1995; Schnitzler, Moss & Denzinger 118 2003).

119 Similarly little is known on how juveniles learn about the location of suitable roosts and the 120 few existing studies only involved adults. In Common noctule bats local enhancement by 121 inadvertent acoustic cues significantly reduces the time required to locate a roost both in 122 captive and in wild experiments (Ruczyński, Kalko & Siemers 2007; Furmankiewicz et al. 123 2011). Kerth and Reckardt (2003) tracked nightly roost switching behaviour in Bechstein's 124 bats and assumed that recruitment of naïve by informed individuals already started at the day 125 roost, but could not unequivocally prove it. An exclusion experiment by Wilkinson (1992) 126 demonstrated that juvenile bats are able of relocating at a new roost with their mothers and the 127 author concluded that following behaviour is the only plausible explanation. One of the still 128 standing mysteries in bat ecology is how juveniles of temperate species locate swarming or

hibernation sites, which are often long distances from where they are born and reared. It has been hypothesized that mothers guide their offspring (Sachteleben 1991), but so far nobody was able to track such guidance behaviour. These examples emphasize that there are plenty of indications on following and guidance behaviour in juvenile bats, but so far technological limitations prevented the final proof.

134 While emerging technologies have revolutionized the field of bio-logging and in turn our 135 understanding of behaviour of wild animals during the past decades, studies on small-bodied 136 vertebrates still lag behind due to the scarcity of fully automated lightweight devices (Kays et 137 al. 2015). Proximity loggers represent a powerful tool for the study of information transfer 138 (Rutz et al. 2015; St Clair et al. 2015), but studies making use of such devices are generally 139 rare and the loggers used in the aforementioned studies are by far too heavy for tagging 140 medium-sized bats. Smaller tag versions of acceptable weight, however, show dramatically 141 reduced runtimes of less than 24 h (Levin et al. 2015). In the present study we used the newly 142 developed miniaturized proximity sensor system 'BATS', a fully automated system for 143 documenting associations among individuals at a tag weight of one to two gram and runtimes 144 of at least one to two weeks (Duda, Weigel & Koelpin 2018). Our developments enable us to 145 study interactions among tagged bats both while roosting and while on the wing. Here we 146 report on the first extensive study to apply our system and proximity sensors to free-ranging 147 bats, in general.

The goal of this study was to investigate the use of social information in acquiring access to two types of resources, which are crucial in the life of a juvenile bat: suitable roosting sites and fruitful feeding grounds. We hypothesized that fledging offspring will make use of social information by following either the mother or other informed members of the social groups to unknown roosts or foraging sites. If juveniles use social information when switching roosts, we expect that the successfully switching juvenile will be associated with at least one individual of the group shortly before and shortly after leaving the current roost and shortly

before and shortly after arriving at the new roost. If social information is used for finding foraging grounds, we would expect juveniles to be associated with at least one roosting partner shortly before and shortly after starting the bout, during several minutes after starting while commuting to the foraging ground, and possibly, but not necessarily when returning to the roost. The BATS-tracking system enabled us to classify and quantify the aforementioned events.

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#### 162 **2. Materials and methods**

163 2.1 Field site and study species

164 This study was conducted in "Königsheide Forst", a mixed forest in Berlin, Germany, from 165 June to August 2016 and 2017, respectively. The study site comprises ample of roosting 166 opportunities for bats such as natural tree holes and roughly 130 bat boxes. During this time 167 of the year females of the common noctule bat (Nyctalus noctula) form temporary groups, so 168 called maternity colonies, to jointly give birth and rear their young. Mothers give birth to one 169 or two offspring and individuals of a maternity colony frequently switch roosts, but usually 170 stay within the area of the nursing colony. Moving among roosts may involve a change in 171 group composition. However, strong, non-random inter-individual bonds have been observed 172 in captive studies as well as a certain degree of maternal care such as allogrooming of 173 offspring (Kleiman 1969; Kozhurina 1993).

The ideal opportunity to observe information transfer in maternity colonies should be the moment when the offspring start to fledge in order to track their behaviour during the first nights of independent flight. Therefore, we daily monitored the bat boxes, including checks after sunset when adults and already flying juvenile had emerged from the roost. We aimed at tagging the majority of a social group including juveniles, which have started fledging only recently or which not fledged at all, yet. We therefore prepared to capture on the following

day when around a third of the offspring were still inside the roost while the rest of colony(including already fledged youngsters) was foraging.

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183 2.2 Sample collection, molecular analysis and identification of mother pup pairs

184 In 2016 we captured a social group from a single bat box while in 2017 bats were caught from 185 two different bat boxes which were roughly 300m apart. Bats were kept in cotton cloth bags 186 until they were weighted, sexed and the forearm was measured using a calliper. If the 187 epiphyseal gaps were closed and the phalangeal-metacarpal joints were knobby, individuals 188 were considered adult (Brunet-Rossinni & Wilkinson 2009). We collected tissue samples with 189 a biopsy punch (Ø 4 mm, Stiefel Laboratorium GmbH, Offenbach, Germany) and preserved 190 them in 80% ethanol. In the lab we used the salt-chloroform procedure (Miller, Dykes & 191 Polesky 1988) modified by Heckel et al. (1999) for DNA isolation.

We used the DNA Analyser 4300 and the SAGA<sup>GT</sup> allele scoring software (both: LI-COR 192 193 Biosciences, Lincoln, NE, USA) to genotype a total of 75 individuals (n = 33 adult 194 females,  $n \equiv \exists 42$  juveniles) at 9 polymorphic microsatellite loci. We used the loci P11, P217, 195 P219 and P223 which were isolated from the focus species Nyctalus noctula (Mayer, 196 Schlötterer & Tautz 2000). Nleis3 and Nleis4 were isolated from the closely related Nyctalus 197 leisleri (Boston, Montgomery & Prodöhl 2009) and G6-Mluc, G31-Mluc, H23-Mluc and 198 H29-Mluc have originally been isolated from *Myotis myotis* (Castella & Ruedi 2000), but 199 were subsequently modified for cross-species utility in vespertilionid species (Jan et al. 2012). 200 To calculate allele frequencies all adult individuals from both years (n=33) were used. All 201 individuals were genotyped at least at eight loci, and genotypes were 99.7% complete. See 202 Table S1 in supporting information for allele numbers per locus, results of Hardy–Weinberg 203 tests, null allele frequencies, and non-exclusion probabilities for the nine microsatellite 204 markers.

205 Parentage analyses were performed with CERVUS v. 3.0 (Kalinowski, Taper & Marshall 206 2007) separately for the social groups caught in 2016 and 2017, respectively, since our 207 objective was to identify mother-pup pairs within year, not across years. The 2016 data set 208 comprised 20 juveniles and 13 adult females (candidate mothers), while 22 juveniles and 24 209 candidate mothers were used for 2017. Four of the 24 adult females in 2017 were recaptures 210 that were already caught in 2016 as juvenile (n=1) and adults (n=3). 211 Simulations were run with  $100 \ 000$  cycles, a proportion of 80% sampled candidate mothers, 212 an estimated genotyping error of 2%, and for two confidence levels (80% and 95%). One 213 mismatch per mother-offspring dyad was accepted to account for genotyping errors or 214 mutations. A mother could be assigned to 40 (2016: n=18; 2017: n=22) of the 42 analysed 215 juveniles. Thirty-three mother-pup pairs were assigned at 95 % confidence with no mismatch, 216 six at 95 % confidence with one mismatch and only one with 80 % confidence and one 217 mismatch.

218

219 2.3 Automated encounter detection among tagged bats

220 Our team developed a tracking system for direct encounter detection, which bases on wireless 221 sensor network technology for field strength related distance estimation between individuals. 222 The system is fully automated including remote data download and does not require 223 recapturing tagged animals thus reducing disturbance of the animals to a minimum. The 224 centrepiece of the tracking hardware is the animal-borne mobile node, in the following 225 referred to as 'proximity sensor'. Once deployed, a wake-up receiver on the proximity sensor 226 permanently scans its surroundings for signals of other proximity sensors, which are 227 constantly broadcasted every two seconds. This operation mode is independent of any further 228 infrastructure. Whenever one or more tracking sensors are within reception range of ca. 10 m 229 maximum distance (Ripperger et al. 2016), a so called 'meeting' is created. As soon as no 230 signal has been received by the respective meeting partner for five sending intervals

231 (corresponding to 10 seconds), the meeting is closed and stored to on-board memory along 232 with the ID of the meeting partner, a timestamp, total meeting duration, and a maximum 233 signal strength indicator (RSSI) of the meeting. The signal, which is broadcasted every 2 s, is 234 simultaneously used as an indicator of presence at a site of interest, e.g. a roost, when the 235 signal is received by a stationary node, in the following referred to as 'base stations'. We 236 positioned base stations near potential roosts to detect presence of individual tagged bats 237 inside a particular bat box or tree hole and we therefore termed a bat signal which are picked 238 up by base stations 'presence signal'. Base stations also provide remote data download, while 239 all downloaded data is locally stored and can be accessed by the user. In 2016 the system 240 could operate a maximum of 30 IDs at a time, while in 2017 the maximum number of 241 observable individuals has been increased to 60. In the following we give a brief overview of 242 the hardware components and the functionality of the system. For an elaborate, in-depth 243 description of the software see Cassens et al. (2017) and for hardware see Duda, Weigel and 244 Koelpin (2018).

245

246 2.3.1 Proximity sensors: We used a refined version of miniaturized proximity sensors, which 247 has been described and tested in free-ranging bats first in Ripperger et al. (2016). The 248 proximity sensor comprises a System-on-Chip (SoC) for communication control and on-board 249 data processing, a transceiver which enables communication in the 868MHz frequency band 250 with other proximity sensors or base stations and a wake-up-receiver which activates full 251 system functionality from an energy-saving low-power mode whenever communication 252 partners are in range. A lithium-polymer battery powers the mobile node. We built two 253 versions of the proximity sensor that differ in weight since adult females and offspring of 254 noctule bats varied considerably in body weight. The low-weight version for tagging offspring 255 was equipped with a 15 mAh battery and was housed in the fingertip of a nitrile lab glove. 256 The heavier version for adult females was equipped with either two 15 mAh battery of a

single 24mAh batteries and housed in a 3D-printed plastic case ensuring longer runtime. The
different proximity sensor versions resulted in a total weight of 1.1 to 1.9 g including battery
and housing.

260

261 2.3.2 Base stations and data access

The base station contains a receiver for the reception of presence signals and transmitted data. Presence signals and downloaded data are stored by a Raspberry Pi (Raspberry PI Foundation, Cambridge, UK) to a SD card along with the ID of the transmitting proximity sensor and the receiving base station, respectively, and a timestamp which is provided by a GPS unit. At the same time the Raspberry Pi hosts a WiFi hotspot allowing the user remote data access. The data is then stored in a MySQL database.

268

## 269 2.4 Tagging and data collection

On July 15<sup>th</sup> 2016 we tagged in total 26 individuals, 10 juveniles and 16 adult females, from a 270 single bat box. On July 18<sup>th</sup> 2017 we tagged in total 34 individuals, 19 juveniles and 15 adult 271 272 females from two bat boxes. This adds up to a total of 60 tracked bats, 29 of which were 273 juveniles and 31 were adult females. According to individual body weight we used different 274 versions of the proximity sensors. Bodyweight ranged from 17g to 25g for juveniles and 275 averaged at 21.26g + 2.04, while adult body weight ranged from 23.5g to 35g at an average 276 of 27.73g +/- 2.27. Individual tag-to-body weight ratios ranged from 4.4% to 7% for juveniles 277 and from 4.2% to 8% in adults, which is well within the recommendations for short-term 278 biologging studies in bats (Amelon et al. 2009; O'Mara, Wikelski & Dechmann 2014). 279 Proximity sensors were glued to the fur on the back of the bats using surgical cement (Perma-280 Type, Plainville, CT, USA) and drops off when the cement loses its tackiness. 281 Data collected during the first night after the tagging event was discarded to account for

282 potential behavioural changes right after tagging and actual data collection started the night

after in order to allow the bats to get used to the tag. In 2016 data collection lasted until July 284 28<sup>th</sup> (12 days) and in 2017 until August 8<sup>th</sup> (20 days). We installed three respectively five base 285 stations in 2016 and 2017 at day roosts to download data and to receive presence signals for 286 individual bats. Whenever bats switched to unknown roosts we used a handheld 868 MHz 287 panel antenna (HSP-868C, WiMo, Herxheim, Germany) connected to a base station to 288 localize the unknown roost and relocate base station.

289

290 2.5 Analysis of tracking data

291 We used the library RMySQL in R (James & DebRoy 2012) to access the data, which were 292 managed in HeidiSQL, a Windows client for MariaDB. In a first step we plotted and visually 293 explored presence signals and meetings received at base stations to define foraging bouts and 294 roost switching events, respectively, for all individual juveniles (see Fig. 2a-c for examples). 295 To evaluate potential information transfer we queried the meeting database for events, which 296 matched the timestamp of foraging bouts or roost switches, respectively. If information 297 transfer would play a role during foraging or roost switching, we would expect to find 298 meetings among offspring and other group members associated with these events. In detail, 299 we proceeded as follows.

300

301 2.5.1 Evaluation of information transfer during roost switching

We defined a roost switch as an event during which an individual changes its roost and potentially its roosting partners without prolonged absence times which may indicate foraging. A roost switch can be detected if a bat switches between two roosts which are both equipped with a base station receiving presence signals (Fig. 2b). If at least one roost is equipped with a base station, presence signals can be used to determine departure time or arrival, respectively. If the unmonitored roost is occupied by other tagged bats (indicated by reciprocal, stable meetings) we can at least unequivocally classify this event as a roost switch

309 (Fig. 2c). However, we cannot determine the time of arrival respectively departure at the 310 unmonitored roost because bats may leave jointly. If juveniles use social information when 311 switching roosts, we expect that the switching juvenile will be associated with at least one 312 individual of the group shortly before and shortly after leaving the current roost or arriving at 313 the new roost. To this end we define the moment of departing from or arriving at a monitored 314 roost, respectively, when the steady reception of signal beacons at a base station gets cut off 315 or starts. We subsequently queried the meeting database for meetings which are active or 316 which started within 60s before and within 60s after the moment of leaving or arriving at a 317 roost.

318

319 2.5.2 Evaluation of information transfer during foraging bouts

320 We defined a foraging bout as an event where an individual starts from a known roost, returns 321 to the same roost and does not visit other monitored roosts or roosts with tagged bats 322 (indicated by stable, lasting meetings) in between (Fig. 2a). We chose these strict rules to 323 ensure that the events we are looking at relate to foraging and do not overlap with roost 324 switching events. If social information would play a role in locating foraging grounds we 325 would expect a juvenile to associate with at least one roosting partner upon starting the bout, 326 during several minutes after departure while commuting to the foraging ground, and possibly, 327 but not necessarily when returning to the roost. As described above we equally defined the 328 start and the end of the foraging bout as the end and the start of the steady reception of the 329 presence signal, respectively. We then queried all meetings which were ongoing or started 330 within 60s before and within 60s after starting a foraging bout and returning, respectively. In 331 addition, we queried all meetings which originated during the entire foraging bout.

332

333 2.5.3 Statistical testing

334 We used a Mantel-Test to test whether social information used by juveniles is obtained by 335 their mothers in the first place or by any roosting partner. To this end we created a binary 336 matrix containing "1" for dyads which have been associated while roost switching and "0" for 337 dyads which have never been observed switching communally. Accordingly, foraging 338 associations were transformed into a binary martrix. For testing the effect of maternity we 339 created a second binary correlation matrix which listed the genetically determined identity of mother-pup pairs as "1", while all other dyads were marked "0". We tested the years 2016 and 340 341 2017 separately and ran Mantel tests in the library "ade4" version 1.7-11 in RStudio 1.1.453 342 using Monte-Carlo permutation tests with 9999 replicates (Dray & Dufour 2007; R 343 Developing Core Team 2015).

344

#### 345 **3. Results**

346 3.1 Genetic analyses

347 Mother and juvenile bats were caught in day roots at the time of weaning. In 24 determined

348 mother-pup pairs, both individuals were tagged with proximity sensors (2016: n=9, all

assigned at 95 % confidence with no mismatch; 2017: n=15, 12 pairs assigned at 95 %

confidence with no and three pairs at 95 % confidence with one mismatch). These 24 mother-

351 pup pairs generated the data for the following section.

352 3.2 Tracking results

In 2016 we received a total of 561,795 presence signals and 13,292 meetings from 23

individual bats and in 2017 we received 2,667,409 localization signals and 53,391 meetings

from 33 individual bats. One individual in 2016 and three individuals in 2017 did not get in

356 contact with base stations. These four individuals may have left the study area between

357 tagging and the following night.

358 3.2.1 Evaluation of joint roost switching events

359 To evaluate information transfer on roosts we screened the data set for joint departures from 360 and joint arrivals at roosts for all tagged juveniles. In 2016 we observed ten events of seven 361 individual juveniles being associated with another individual while switching among two 362 roosts. In all except one event the associated bats arrived together at a new roost, even though 363 successful switching took several approaches in two cases and temporary roosts may be used 364 in between (Table 1, Fig. 3). In six cases both roosts have been monitored by a base station, in 365 two cases the juveniles left a monitored roost and switched to a roost where other tagged bats 366 have been roosting and in the remaining two cases the juveniles switched from a monitored 367 roost to an unknown roost where no other tagged bats were present, except the one which 368 accompanied the juvenile during switching. In all 10 cases the juvenile was in company of its 369 identified mother and no other tagged bat.

In 2017 we observed six events where 5 individual juveniles switched roosts in company. Twice, the juvenile switched among two monitored roosts and four times among one monitored and an unmonitored site. Twice, the juvenile was associated with its identified mother, in four cases with another adult female.

Some juveniles switched directly among roosts. Such events took only seconds to minutes (see table xx). During other events stopover sites were used and several attempts of mothers re-associating with their young were necessary before both arrived at the new roost. Such unsuccessful tandem flights underline that the offspring was actively flying and not carried by the mother.

In both years significantly more mother-pup dyad have been observed switching roosts communally than expected by chance (Mantel tests, 9999 replicates; 2016: r = 0.88, p < 0.001; 2017: r = 0.21, p < 0.01).

382

383 3.2.2 Associations during foraging bouts

384 In total we detected 42 foraging bouts of juveniles, which matched our definition above, 385 conducted by 13 individuals (2016: four juveniles, eight bouts; 2017: nine juveniles, 34 386 bouts). Foraging bouts lasted on average 1:14:53 h with a standard deviation of 36:19 min. 387 During 6 of these 42 foraging bouts (14 %, n = 7 individual juveniles, all 2017) we detected in 388 total 28 short meetings, which lasted between 1 and 30 seconds (average: 7.4 s +/- 8.6). Two 389 of these meetings occurred within less than 90 s after two co-roosting individuals left a roost; 390 however, no further meetings have been documented during these foraging bouts. All 391 remaining meetings originated at least several minutes after emergence from the roost. Eight 392 times the meeting partner was another juvenile and twice an adult female. Only in one case 393 the meeting partner was the identified mother. Accordingly, meetings among mother-pup 394 dyads have not been observed more often than expected by chance (Mantel tests, 9999 395 replicates, r = 0.08, p > 0.05).

In 13 out of the 42 foraging bouts of juveniles (2016: n = two individuals; 2017: n = seven individuals) the identified mother was co-roosting before both started a bout. In all 13 cases the mother started its foraging bout considerably earlier than the juvenile (between 4:31 min and 1:26:02 h, average: 36:45 min (+/- 24:57)).

400

### 401 **Discussion**

The study of information transfer in free-ranging bats is particularly challenging due to their small body size and their elusive, nocturnal life. We tracked bats using novel, miniaturized proximity sensors and demonstrated that juveniles use social information of group members and for finding roosts mothers seem to intentionally guide their young. However, during foraging mothers did not guide their offspring, but meetings with other colony members may reflect local enhancement at feeding grounds.

408 To the best of our knowledge our study shows for the first time that recruitment to a new roost

409 starts already at the occupied roost. Furthermore, the repeated commuting flights we observed

410 in at least two cases until the juvenile arrives at the target roost represents anecdotal evidence 411 that at least in some cases deliberate, evolved signals rather than inadvertent social cues are 412 used. The existence of evolved signals and the strong bias towards information transfer 413 among mother-pup pairs suggests that the observed behaviour is best explained by kin 414 selection. Some studies have reported on the use of social information in bats for finding 415 suitable roosts, however, studies are scarce and the mechanisms are in parts poorly 416 understood, in particular when it comes to naïve juveniles. Studies on a range of 417 vespertilionid species including the focus species *N. noctula* have shown that conspecific calls 418 enhance roost finding efficiency in captive experiments as well as in the wild (Ruczyński, 419 Kalko & Siemers 2009; Schöner, Schöner & Kerth 2010; Furmankiewicz et al. 2011). These 420 studies demonstrate that bats may eavesdrop on vocalizations to localize an occupied roost 421 once within hearing distance. Since playbacks from varying contexts have been used we 422 conclude that the studied bats relied on inadvertently broadcasted public information. On the 423 contrary, Spix's disk winged bats deliberately produce signals to facilitate group cohesion, by 424 a remarkable call-and-response system among flying bats in search of a roost and bats 425 occupying a roost (Chaverri, Gillam & Vonhof 2010; Chaverri, Gillam & Kunz 2012). A 426 common theme of all abovementioned studies is that the mechanism of recruitment of 427 conspecifics is best explained by local enhancement, i.e. the social information is acquired at 428 the new roost, when searching bats are in hearing distance. Kerth and Reckardt (2003) were 429 first to present experimental evidence for information transfer about roosts in bats. The 430 authors presumed that naïve Bechstein's bats are recruited to a novel roost already at the 431 dayroost by experienced conspecifics, however, they could not unequivocally exclude local 432 enhancement at the target roost. Our study finally demonstrates that this inferred mechanism 433 does exist in roost-switching bats.

We classify the advertent information transfer from mothers to their young as a form ofmaternal care which has to the best of our knowledge not been observed in free-ranging bats,

436 so far. Mammalian offspring is usually strongly dependent on maternal care for food, 437 protection and warmth (Balshine 2012) and maternal investment in young is also wide-spread 438 in bats (Smith, Lacey & Hayes 2017). Besides weaning maternal care has been demonstrated 439 in form of post-weaning food provisioning (Wilkinson 1990; Geipel et al. 2013), grooming 440 (Kleiman 1969; Wilkinson 1986; Kozhurina 1993) or pup guarding (Bohn, Moss & 441 Wilkinson 2009). Carrying young in flight is also commonly observed and Jones (2000) 442 summarizes some reports where young are possibly carried to temporary roosts or feeding 443 grounds. However, this is the first study to document maternal guidance to roosts, which has 444 been hypothesized as a plausible explanation for young to reach swarming and hibernation 445 sites, but could not be confirmed, possibly due to the lack of appropriate tracking technology 446 (Sachteleben 1991; Burns & Broders 2015; Stumpf et al. 2017).

447 Previous work on bats indicated that roosts may act as information centres where bats may 448 obtain information on food by inadvertent cues (Ratcliffe & ter Hofstede 2005; O'Mara, 449 Dechmann & Page 2014). A considerable part of the diet of Common noctule bats consists of 450 insects, which fly in swarms and often over water (Gloor, Stutz & Ziswiler 1995). Such rich and patchy, but ephemeral foraging sites are required for the establishment of information 451 452 centres (Ward & Zahavi 1973) and juveniles in particular might benefit from rich food patches when collecting experience on where and how to forage. However, we did not 453 454 observe recruitment at the roost to feeding grounds in young noctules, which complies with 455 most foregoing studies that showed that 'ICH' operates in colonial roosts, but is rarely 456 demonstrated in breeding colonies (summarized by Evans, Votier and Dall (2016)). Our 457 observation that juveniles start foraging bouts considerably later than their mothers suggests 458 that juvenile noctules conduct opportunistic, explorative foraging flights. Rare and short 459 contacts to tagged colony members other than the mother during foraging bouts suggest that 460 local enhancement by eavesdropping on conspecifics while hunting may play a role as it has 461 been shown for several insectivorous bat species (Gillam 2007; Dechmann et al. 2009; Cvikel

*et al.* 2015). However, our data cannot unequivocally prove this theory since the exact contextof the meetings remains unknown.

464 Our observations raise the following question: Why is social information transfer among 465 mothers and their offspring context dependent? One possible explanation is that group 466 cohesion is crucial for energy-saving social warming and prolonged lactation periods in bats 467 require mothers to stay in contact with their young for 3 weeks to 2 months depending on the 468 species (reviewed by Kerth (2008)). Extended weaning, which was observed in captive 469 noctules for up to 2 months (Kleiman 1969), and the broad spectrum of insects they feed on 470 (Gloor, Stutz & Ziswiler 1995) may in turn enable juveniles to forage opportunistically and – 471 if available – make use of social information by local enhancement. In general, suitable roosts 472 of high quality may be harder to find opportunistically than insect prey and information on 473 roosts is likely to accumulate in adults, in particular in philopatric females. This should favour 474 information transfer on roosts since failing to relocate at an occupied roost might be more 475 costly than low foraging success, which might subsequently be balanced by extended 476 weaning. Adverse climatic conditions may have detrimental effects on single bats (Lindström 477 1999) and might therefore be a strong driver of the evolution of the observed guidance 478 behaviour, since local enhancement by vocalization at the roost (Furmankiewicz et al. 2011) 479 might not be functional for long-distance localization of roosting partners.

480

#### 481 **Conclusions**

Bats are facing ideal prerequisites for social information transfer, since they are long-lived and the vast majority of species is living in group. Regarding information use in offspring Wilkinson and Boughman (1999) speculated already 20 years ago that young bats almost certainly follow adults in situations other than foraging. However, this is also how long it took to unequivocally track mother-pup pairs switching among roosting sites. Our study shows that

487	the	current	revolution	in	tracking	technology	provides	powerful	tools	to	investigate
488	beha	avioural	ecology and	soc	iobiology	in free-rangi	ng small b	odied anim	als suc	ch as	s bats.

489

490 Authors' contribution
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491 SR and FM conceived the ideas and designed the sampling scheme; MH, ND, AK, BC and

492 RK developed and tested all tracking equipment. SR, LG, FM and HW collected the data; SR

493 and LG analysed the data; SR and FM led the writing of the manuscript. All authors

494 contributed critically to the drafts and gave final approval for publication.

495

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113-G0008/16).

501

### 502 Data accessibility

503 Data will be deposited on GFBio and made available upon acceptance.

504

## 505 Literature

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# 661 Supporting information

- 662 The following Supporting Information is available for this article online.
- Table S1: Results from allele frequency calculations with CERVUS v. 3.0 (Kalinovski et al.
- 664 2007).

## 665 **Tables**

Table 1: Summary of joint roost switching events of pups and associated partners. Switching

- 667 durations were only determined when both roosts were known and equipped with base
- stations, while NA represents unmonitored sites or uncertain switching mode.

		partnerID	start time	end time	duration	Switching
pupID	sex	(mother)	[date-time]	[date-time]	[hh:mm:ss]	mode
9307	f	9338 (m)	2016-07-21 02:59:35	NA	NA	NA
9311	f	9330 (m)	2016-07-16 23:26:22	2016-07-17 01:15:06	01:48:44	3*
9311	f	9330 (m)	2016-07-20 03:18:35	2016-07-20 03:19:31	00:00:56	1
9318	f	9330 (m)	2016-07-16 23:26:33	2016-07-16 23:28:31	00:01:58	1*
9318	f	9330 (m)	2016-07-20 04:14:25	2016-07-20 04:14:54	00:00:29	3
9319	f	9327 (m)	2016-07-16 22:48:01	NA	NA	NA
9312	f	9334 (m)	2016-07-17 01:29:14	NA	NA	1
9323	m	9340 (m)	2016-07-16 23:58:13	2016-07-17 01:37:06	01:38:53	2
9323	m	9340 (m)	2016-07-19 02:05:26	NA	NA	1
9325	m	9336 (m)	2016-07-17 02:44:31	2016-07-17 03:55:19	01:10:48	2
9376	f	9383	2017-07-20 04:23:43	2017-07-20 04:23:49	00:00:06	1
9376	f	9383	2017-07-22 02:25:53	NA	NA	1
9370	f	9368 (m)	2017-07-20 04:35:41	NA	NA	1
9373	m	9412	2017-07-22 02:38:13	2017-07-22 02:38:39	00:00:26	1
9380	m	9413	2017-07-20 04:17:52	NA	NA	1
9391	f	9385 (m)	NA	2017-07-22 02:22:25	NA	1

Switching mode: 1 quick, direct switch among two roosts; 2 switch included a joint stopover
at an unknown site indicated by stable meetings; 3 roost switch after commuting flights by
mother; \* see Fig. 3 for a schematic representation of these events

#### 673 **Figures**

674

Figure 1: Unpackaged proximity sensor (a) and tagged adult Common noctule bat (*Nyctalus noctula*) ready for take-off (b).

677

678 Figure 2: Visual representation of foraging bouts and roost switches based on presence signals 679 at bat boxes (base stations) and meeting data. (a) A foraging bout is characterized by an 680 interrupt of the presence signals of an individual bat which are received by a base station at a 681 specific roost. Usually, variation of the received signal strength indicator (RSSI) increases 682 when a bat is leaving a roost compared to when it is roosting (notice the pronounced spike 683 upon departure and return). (b) A roost switch among two monitored sites is displayed. The 684 presence signals interrupt at base station two while the strong variation in RSSI indicates that 685 the bat is flying. Presence signals are then received by base station 1. (c) A roost switch 686 occurs among an unmonitored to a monitored site. Roosting at the unmonitored site is 687 indicated by long-lasting stable meetings among three bat individuals. Meetings interrupt 688 when a bat individual leaves the unmonitored site followed by signal beacons being received 689 by the base station at the monitored site.

690

691 Figure 3: Schematic representation of a mother and its twins switching roosts: repeated 692 commutes indicate intentional behaviour of the mother. (a) A mother and its twins jointly 693 leave roost 1 and the mother successfully transfers to roost 2 with pup 1. The meeting to pup 2 694 aborts and pup 2 is flying back to roost 1. (b) Pup 2 moves solitarily from roost 1 to an 695 unknown location where it is joined by its mother after a few minutes. Both fly in company 696 towards roost 2, but pup 2 flies back to roost 1 while a meeting starts among the mother and 697 pup 1 at roost 2. (c) The mother joins pup 2 in an unknown location and they jointly switch to 698 roost 1. They jointly leave roost 1, but only the mother arrives at roost 2 starting a meeting

- 699 with pup 1, while the pup 2 flies back to roost 1. (d) The mother joins pup 2 in an unknown
- 100 location, around 00:50 am the meeting is interrupted for several minutes (possibly because at
- rol least one individual left), before the mother commutes twice between its two pups. Finally,
- around 01:15 am the mother successfully switches with pup 2 to roost 2 while a meeting is
- ongoing. The triad stays at roost 2 until shortly before 2 am.





Legend: O Presence signal at base station 1

 $\triangle$  Presence signal at base station 2

Presence at unknown roost indicated by stable meetings among three individuals

