Estimating the Density of Honeybee Colonies across Their Natural Range to Fill the Gap in Pollinator Decline Censuses

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Abstract: Although pollinator declines are a global biodiversity threat, the demography of the western honeybee (Apis mellifera) has not been considered by conservationists because it is biased by the activity of beekeepers. To fill this gap in pollinator decline censuses and to provide a broad picture of the current status of honeybees across their natural range, we used microsatellite genetic markers to estimate colony densities and genetic diversity at different locations in Europe, Africa, and central Asia that had different patterns of land use. Genetic diversity and colony densities were highest in South Africa and lowest in Northern Europe and were correlated with mean annual temperature. Confounding factors not related to climate, however, are also likely to influence genetic diversity and colony densities in honeybee populations. Land use showed a significantly negative influence over genetic diversity and the density of honeybee colonies over all sampling locations. In Europe honeybees sampled in nature reserves had genetic diversity and colony densities similar to those sampled in agricultural landscapes, which suggests that the former are not wild but may have come from managed hives. Other results also support this idea: putative wild bees were rare in our European samples, and the mean estimated density of honeybee colonies on the continent closely resembled the reported mean number of managed hives. Current densities of European honeybee populations are in the same range as those found in the adverse climatic conditions of the Kalahari and Saharan deserts, which suggests that beekeeping activities do not compensate for the loss of wild colonies. Our findings highlight the importance of reconsidering the conservation status of honeybees in Europe and of regarding beekeeping not only as a profitable business for producing honey, but also as an essential component of biodiversity conservation.
**Keywords:** Apis mellifera, apicultura, conservación de abejas, declive de polinizadores, función del ecosistema, uso de suelo

Estimación de la Densidad de Colonías de Abejas en su Rango Natural para Cerrar la Brecha en los Censos del Declive de Polinizadores

Resumen: Aunque la declinación de polinizadores es una amenaza global para la biodiversidad, la demografía de la abeja (Apis mellifera) no ha sido considerada por conservacionistas ya que está sesgada por la actividad de los apicultores. Para cerrar esta brecha en los censos del declive de polinizadores y proporcionar una visión amplia del estatus actual de las abejas en su rango natural, utilizamos marcadores genéticos de microsatélites para estimar las densidades de colonias y la diversidad genética en diferentes localidades en Europa, África y Asia central que tienen diferentes patrones de uso de suelo. La diversidad genética y las densidades de colonias fueron mayores en África del Sur y menores en el norte de Europa y se correlacionaron con la temperatura media anual. Sin embargo, es probable que factores de confusión no relacionados con el clima también afecten a la diversidad genética y las densidades de colonias en poblaciones de abejas. Otros resultados también soportan esta idea: abejas tentativamente silvestres fueron raras en nuestras muestras europeas y la densidad promedio de colonias de abejas estimada en el continente se aproxima bastante al número promedio de colmenas comerciales reportado. Las densidades actuales de abeja europea están en el mismo rango que las encontradas en las condiciones climáticas adversas de los desiertos Kalabari y Sabara, lo que sugiere que las actividades apícolas no compensan la pérdida de colonias silvestres. Nuestros resultados resaltan la importancia de revisar el estatus de conservación de las abejas en Europa y de considerar a la apicultura no solo como un negocio rentable para la producción de miel, sino también como un componente esencial de la conservación de la biodiversidad.

Palabras Clave: apicultura, Apis mellifera, conservación de abejas, declive de polinizadores, funcionamiento del ecosistema, uso de suelo

Introduction

Pollination is an important ecosystem service because the majority of wild plants are insect pollinated, as are most commercial crops (e.g., Buchmann & Nabhan 1996; Constanza et al. 1997; Klein et al. 2007). Pollinators, however, are under increasing threat from anthropogenic factors, including fragmentation and destruction of natural habitats through intensive land use, toxic pesticides used in agriculture, and invasive plant and animal species and their diseases (Allen-Wardell et al. 1998; Brown & Paxton 2009; but see also Ghazoul 2005). Without pollinators, ecosystem functioning, trophic cascades, and the survival and maintenance of genetic diversity of many wild plant populations would be at risk (Kearns et al. 1998), and economic yields of crops may suffer a drastic reduction (Gallai et al. 2009).

Even though pollinator declines are a global biodiversity threat, the demography of the western honeybee (Apis mellifera) has not been taken into account by conservationists because it is influenced by the activities of beekeepers and therefore is regarded as unnatural (e.g., Kremen et al. 2002). Nevertheless, honeybees are key generalist pollinators. They live in large perennial colonies that ensure high local pollination throughout the flowering season (Seeley 1985) and have a vast foraging area that enhances gene flow in plant communities (Roubik 1989; Free 1993). The western honeybee is endemic to Africa, Europe, and the Middle East (Ruttner 1988; Whitfield et al. 2007), but has been introduced worldwide for exploitation as a honey producer and crop pollinator (Huryn 1997; Moritz et al. 2005). Despite their commercial use, honeybees have suffered a progressive decline throughout the world over recent decades (Matheson et al. 1996). The recent occurrence of so-called colony collapse disorder (CCD) in the United States and a similar phenomena in Europe resulted in the death of hundreds of thousands of colonies (van Engelsdorp et al. 2008).

The global decline of managed honeybees has been discussed primarily in light of the important economic losses they represent (Southwick & Southwick 1992; Morse & Calderone 2000; Allsopp et al. 2008), and little attention has been paid to wild and feral honeybee populations (Moritz et al. 2005). Wild populations are important reservoirs of local adaptations that ultimately determine the survival of honeybees in the wild (Matheson et al. 1996). Moreover, a simultaneous decline of wild and managed honeybees may have dramatic consequences on pollination services to crops and natural
ecosystems (Gallai et al. 2009). On the basis of question-
able evidence for competition between honeybees and
other native pollinators (reviewed by Huryn 1997 and
Paini 2004, but see Roubik & Wolda 2001), many Eu-
ropean governments (e.g., Belgium, Netherlands, United
Kingdom) have implemented national policies that ex-
clude managed pollinators from protected areas (e.g., sec-
tions 14 and 16 of the Wildlife and Countryside Act 1981,
United Kingdom). In contrast, on the basis of the same
evidence, other countries such as Germany and Austria,
permit apiculture within designated protected areas. The
inconsistency in honeybee conservation policies across
Europe and the magnitude of recent colony losses jus-
tify the inclusion of A. mellifera in pollinator abun-
dance surveys.

The real magnitude of pollinator declines is not easily
determined and has been the subject of a controversial re-
view (Ghazoul 2005). Accurate measurements of popula-
tion densities are essential for any meaningful assessment
of decline (Paini 2004). For nonsocial pollinators this can
be done with direct counts of individuals and classical
abundance measures. For social bees, however, the num-
ber of colonies rather than the number of individuals
is the crucial parameter for conservation. The number
of colonies in the wild is, nevertheless, very difficult to
assess. Density estimates derived from direct counts of
feral and wild honeybee colonies are tedious and can be
unreliable because natural nests are hard to detect (Otis
Molecular DNA tools are a more effective mean to as-
ssess honeybee population density (Estoup et al. 1995;
Moritz et al. 2007). In particular, the use of tightly linked
microsatellite DNA markers to reconstruct queen geno-
types from samples of honeybee drones or workers is the
most powerful tool currently available (Shaibi et al. 2008)
that allows recognition of the offspring of closely related
queens (Jaffe et al. 2009). We used these genetic markers
to quantify the density of A. mellifera colonies in natural
and agricultural landscapes of Europe, Africa, and central
Asia and identified the factors influencing the current de-
ography of honeybees. In addition, to provide insights
into the demographic status of feral and wild honeybee
populations in Europe, we contrasted our estimates with
the number of managed colonies surrounding the sam-
ping locations where beekeeping occurs.

**Methods**

**Sampling**

We collected honeybee samples from 25 locations (10 in
Africa, 2 in central Asia, and 13 in Europe; see Support-
ing Information for coordinates and climatic data). Hon-
eybee drones gather at drone congregation areas (DCAs)
that virgin queens visit to mate with tens of partners
(Ruttner & Ruttner 1972; Moritz & Southwick 1995). The
DCAs attract drones from most colonies within recruit-
ment range (Baudry et al. 1998; Jaffe et al. 2009), so
genotyping drones collected from a DCA allows estimat-
ing the number of drone-contributing colonies within
the recruitment perimeter of the DCA (i.e., the density
of colonies). Honeybee drones can be lured by synthetic
queen pheromone into a trap kept approximately 20 m
above ground by a weather balloon (Williams 1987). We
captured drones from previously identified DCAs, by fly-
ing the pheromone trap between 12:00 and 17:00, under
sunny and windless conditions, during the swarming sea-
son.

At 12 locations where beekeeping is practiced, we al-
ternatively collected worker offspring from 10 managed
colonies headed by locally mated queens (Table 1). Hon-
eybee queens mate with many haploid drones (Moritz
& Southwick 1995). By genotyping workers of a single
queen, it is possible to deduce the genotype of the queen
and that of her mates (Estoup et al. 1994). To avoid sam-
ping workers that drifted from a foreign colony into the
sample hive, we collected young worker bees directly
from the combs upon opening of the hives. With this ap-
proach, failing to detect some fathers in a colony would
be equivalent to failing to sample some drones at a DCA.
Individuals were stored in 95% ethanol until genetic anal-
ysis.

**Genotyping**

We used different sets of microsatellite markers to deter-
mine individual genotypes (Table 1) and a Chelex proto-
col to extract DNA (Walsh et al. 1991). The microsatel-
rite target sequences were amplified by multiplex poly-
merase chain reactions (PCR) with fluorescently labeled
primers. Samples containing no DNA were included in
all plates as negative controls. We then resolved PCR
products in a MegaBACE 1000 capillary sequencer and
determined allele sizes with the Fragment Profiler soft-
ware (Amersham Biosciences, Germany).

**Genetic Diversity Measures and Reconstruction of Queen
Genotypes**

For each sample set, we constructed tables with the geno-
types of all drones (obtained either directly, by genotyping
drones caught in a DCA, or indirectly, by inferring
their genotype from worker offspring of a single queen).
Per locus allelic richness (AR) and gene diversity (ex-
pected heterozygosity, $H_e$) were calculated with FSTAT
(version 2.9.3.2; Goudet 2002). Because drones are pro-
duced parthenogenetically and only carry alleles from
their mother, genotyping drones allows for their assign-
ment to specific queens. To reconstruct the genotype
of individual drone-producing queens, we performed a
sibship reconstruction analysis employing COLONY (ver-
sion 1.3; Wang 2004).
### Table 1. Sample sizes, genetic diversity measures, nondetection errors, and number of estimated and managed honeybee colonies in 25 sampling locations spaced across Africa, central Asia, and Europe.

<table>
<thead>
<tr>
<th>Location</th>
<th>Sample</th>
<th>Allelic richness (SD)</th>
<th>Heterozygosity (SD)</th>
<th>NDE</th>
<th>Number of colonies median</th>
<th>Total</th>
<th>Managed</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Africa</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Al Kufrah, Libya</td>
<td>W</td>
<td>7.92 (3.58)</td>
<td>0.73 (0.04)</td>
<td>5.3×10⁻⁵</td>
<td>29 (1)</td>
<td>45</td>
<td>176</td>
<td>local beekeeper</td>
</tr>
<tr>
<td>Khartoum, Sudan</td>
<td>W</td>
<td>8.24 (3.50)</td>
<td>0.75 (0.07)</td>
<td>4.2×10⁻²</td>
<td>19 (4)</td>
<td>23</td>
<td>15 *</td>
<td>local beekeeper</td>
</tr>
<tr>
<td>El-Faw, Sudan</td>
<td>W</td>
<td>7.25 (1.89)</td>
<td>0.77 (0.04)</td>
<td>3.2×10⁻²</td>
<td>19 (4)</td>
<td>23</td>
<td>0 *</td>
<td>-</td>
</tr>
<tr>
<td>Leeuwfontein, South Africa</td>
<td>D</td>
<td>16.99 (3.31)</td>
<td>0.88 (0.01)</td>
<td>8.4×10⁻¹⁰</td>
<td>17 (0)</td>
<td>23</td>
<td>0 *</td>
<td>-</td>
</tr>
<tr>
<td>Ezenvelo, South Africa</td>
<td>D</td>
<td>18.02 (4.85)</td>
<td>0.91 (0.01)</td>
<td>1.5×10⁻¹⁰</td>
<td>24 (2)</td>
<td>37</td>
<td>0 *</td>
<td>-</td>
</tr>
<tr>
<td>Suikerbosrand, South Africa</td>
<td>D</td>
<td>16.68 (4.79)</td>
<td>0.77 (0.02)</td>
<td>2.0×10⁻⁷</td>
<td>23 (0)</td>
<td>34</td>
<td>0 *</td>
<td>-</td>
</tr>
<tr>
<td>Tswalu, South Africa</td>
<td>D</td>
<td>11.92 (1.37)</td>
<td>0.79 (0.06)</td>
<td>3.6×10⁻²</td>
<td>14.5 (2)</td>
<td>29</td>
<td>0 *</td>
<td>-</td>
</tr>
<tr>
<td>Pietermaritzburg, South Africa</td>
<td>D</td>
<td>12.36 (5.34)</td>
<td>0.82 (0.05)</td>
<td>1.9×10⁻⁵</td>
<td>25 (5)</td>
<td>52</td>
<td>0 *</td>
<td>-</td>
</tr>
<tr>
<td>Elsenburg, South Africa</td>
<td>D</td>
<td>10.22 (2.76)</td>
<td>0.81 (0.01)</td>
<td>4.8×10⁻⁴</td>
<td>17 (1)</td>
<td>29</td>
<td>57</td>
<td>ARC-Plant Protection Research Institute, Honeybee Research Section, South Africa</td>
</tr>
<tr>
<td>Jonkershoek, South Africa</td>
<td>D</td>
<td>11.81 (3.26)</td>
<td>0.84 (0.02)</td>
<td>3.8×10⁻⁵</td>
<td>26 (3)</td>
<td>40</td>
<td>0 *</td>
<td>-</td>
</tr>
<tr>
<td><strong>Central Asia</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dshuuku, Kyrgyzstan</td>
<td>W</td>
<td>7.44 (4.77)</td>
<td>0.71 (0.05)</td>
<td>3.9×10⁻⁵</td>
<td>14 (1)</td>
<td>21</td>
<td>60</td>
<td>local beekeeper</td>
</tr>
<tr>
<td>Tjurgen-Ak-Suu, Kyrgyzstan</td>
<td>W</td>
<td>6.48 (3.70)</td>
<td>0.72 (0.04)</td>
<td>2.5×10⁻⁷</td>
<td>12 (1)</td>
<td>23</td>
<td>40</td>
<td>local beekeeper</td>
</tr>
<tr>
<td><strong>Europe</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Caravaca de la Cruz, Spain</td>
<td>W</td>
<td>8.84 (5.28)</td>
<td>0.66 (0.09)</td>
<td>3.1×10⁻⁴</td>
<td>22 (0)</td>
<td>37</td>
<td>100</td>
<td>Veterinary Faculty, University of Murcia, Spain</td>
</tr>
<tr>
<td>Farindola, Italy</td>
<td>W</td>
<td>8.71 (3.10)</td>
<td>0.75 (0.07)</td>
<td>1.6×10⁻²</td>
<td>37 (8)</td>
<td>45</td>
<td>25 *</td>
<td>CRA-API, Italy</td>
</tr>
<tr>
<td>Intermesoli, Italy</td>
<td>W</td>
<td>9.97 (3.23)</td>
<td>0.84 (0.02)</td>
<td>2.0×10⁻²</td>
<td>33 (6)</td>
<td>39</td>
<td>45</td>
<td>CRA-API, Italy</td>
</tr>
<tr>
<td>Elba, Italy</td>
<td>W</td>
<td>11.06 (5.43)</td>
<td>0.74 (0.08)</td>
<td>1.9×10⁻²</td>
<td>34.5 (7)</td>
<td>42</td>
<td>50</td>
<td>CRA-API, Italy</td>
</tr>
<tr>
<td>Svinjicko, Croatia</td>
<td>D</td>
<td>9.94 (6.58)</td>
<td>0.71 (0.08)</td>
<td>1.3×10⁻⁴</td>
<td>17 (2)</td>
<td>35</td>
<td>50</td>
<td>Faculty of Agriculture, University of Zagreb, Croatia</td>
</tr>
<tr>
<td>Pokupsko, Croatia</td>
<td>D</td>
<td>9.58 (5.67)</td>
<td>0.63 (0.09)</td>
<td>1.9×10⁻³</td>
<td>15 (0)</td>
<td>22</td>
<td>300</td>
<td>Faculty of Agriculture, University of Zagreb, Croatia</td>
</tr>
<tr>
<td>Schwarzenau, Germany</td>
<td>W</td>
<td>4.35 (2.18)</td>
<td>0.67 (0.06)</td>
<td>8.9×10⁻⁵</td>
<td>10 (0)</td>
<td>19</td>
<td>n/a</td>
<td>Kreisimkerverein Halle e.V., Germany</td>
</tr>
<tr>
<td>Halle, Germany</td>
<td>D</td>
<td>8.58 (4.75)</td>
<td>0.70 (0.06)</td>
<td>3.8×10⁻⁴</td>
<td>17 (1)</td>
<td>30</td>
<td>32</td>
<td>Kreisimkerverein Halle e.V., Germany</td>
</tr>
<tr>
<td>Hochharz, Germany</td>
<td>W</td>
<td>6.64 (5.65)</td>
<td>0.68 (0.08)</td>
<td>1.6×10⁻⁴</td>
<td>11 (0)</td>
<td>20</td>
<td>n/a</td>
<td>-</td>
</tr>
<tr>
<td>Müritz, Germany</td>
<td>W</td>
<td>5.69 (4.25)</td>
<td>0.71 (0.05)</td>
<td>7.1×10⁻⁴</td>
<td>16 (0)</td>
<td>23</td>
<td>n/a</td>
<td>-</td>
</tr>
</tbody>
</table>

*continued*
Table 1. (continued)

<table>
<thead>
<tr>
<th>Location</th>
<th>Samplea</th>
<th>n0</th>
<th>Allelic richness (SD)</th>
<th>Heterozygosity (SD)</th>
<th>Number of colonies mediand</th>
<th>Totald</th>
<th>Managedf</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caher, Republic of Irelandg</td>
<td>D</td>
<td>96</td>
<td>2.89 (2.04)</td>
<td>0.32 (0.12)</td>
<td>2.7 × 10^{-2}</td>
<td>7 (0)</td>
<td>14</td>
<td>Bee Improvement and Bee Breeders' Association, Republic of Ireland</td>
</tr>
<tr>
<td>Belfast, Northern Irelandg</td>
<td>D</td>
<td>96</td>
<td>7.54 (3.87)</td>
<td>0.64 (0.12)</td>
<td>1.0 × 10^{-2}</td>
<td>10 (0)</td>
<td>16</td>
<td>Institute of Northern Ireland Beekeepers, U.K.</td>
</tr>
<tr>
<td>Gotland, Swedeng</td>
<td>D</td>
<td>23</td>
<td>1.76 (0.64)</td>
<td>0.38 (0.11)</td>
<td>1.4 × 10^{-2}</td>
<td>2 (0)</td>
<td>3</td>
<td>Swedish University of Agricultural Sciences, Sweden</td>
</tr>
</tbody>
</table>

aDrone genotypes were obtained either directly, by genotyping drone samples collected at DCAs (D), or indirectly, by genotyping worker offspring of mated queens (W).
bFinal sample sizes, given as the number of drone genotypes.
cNondetection errors, calculated as in Boomsma and Ratnieks (1996).
dNumber of colonies represented by at least a median number of drones, followed by the number of colonies remaining undetected because of an insufficient sample (in parentheses).
eSample size corrected number of reconstructed drone-producing queens.
fNumber of managed hives within a radius of 2.5 – 3 km from each sampling location (n/a: not available). This is given as an indicator for the occurrence of putative wild colonies (cases where we estimated more colonies than the managed ones are indicated with an asterisk [*]).
gSamples genotyped using sets of five, five and four tightly linked microsatellite markers. These three linkage groups are located on different chromosomes: loci HB-16-01, AC006, HB-16-02, HB-16-03, HB-16-05 on chromosome 16; loci HB-THE-01, HB-THE-02, HB-THE-03, HB-THE-04, HB-THE-05 on chromosome 13; and loci HB-SEX-01, HB-SEX-02, HB-SEX-03, UN351 on chromosome 3 (Shaibi et al. 2008).
hSamples genotyped using one set of 12 linked microsatellites located on chromosome 16: loci HB-002, HB-005, HB-007, HB-008, HB-010, HB-012, HB-15, HB-016, HB-017, HB-018, HB-019, and SV-240. In the Tswalu sample, the individuals analyzed by Moritz et al. (2008) were pooled with additional drones that had been collected at this site at the same time.
iBeekeeping activity is absent in the area and thus the samples represent wild honeybee populations.
jSamples previously genotyped by Moritz et al. (2007) using 10 microsatellite markers. Their genotypic data were included in our study because a different approach was used here to estimate colony densities (see Methods).
kSamples genotyped using five unlinked microsatellite markers (A7, A24, A79, A107 and A113) in combination with the three sets of tightly linked microsatellite markers (Shaibi et al. 2008).
lPerson or institution surveying the number of managed hives in the vicinity of our sampling locations.

The multilocus genotypes of the samples analyzed by Moritz et al. (2007) were directly introduced into COLONY to assign drones to colonies. In the samples analyzed with the linked markers, we first grouped all individuals sharing the same allelic combination at all loci within each linkage group. Individuals sharing a particular allelic combination were assigned to a single haplotype. Individuals that could not be assigned to a specific haplotype in at least one linkage group (because of low polymorphism or missamplifications at some loci) were excluded from all subsequent analyses. The haplotypes found in each linkage group were then used as individual alleles so that they could be introduced into COLONY. In the six samples in which only one set of linked markers was used for genotyping, we did not use COLONY because we did not have unlinked sets of markers. Instead, given that queens are diploid, we calculated a conservative estimate of the number of drone-producing queens by halving the sample-size-corrected number of haplotypes found in each population.

Nondetection and Nonsampling Errors

Two kinds of errors affected our estimated number of drone-producing queens. First, two identical genotypes could have been detected in two different individuals by chance (nondetection errors [NDEs]). The probability of this type of error is determined on the basis of the number of markers used and their level of polymorphism. We calculated this probability from the allele and haplotype frequencies in the samples analyzed with unlinked and linked markers, respectively, following Boomsma and Ratnieks (1996). Second, queens may have gone undetected because of an insufficient sample size (nonsampling errors). We assessed this type of error with a fitted Poisson distribution of the drones (obtained either
directly or indirectly through worker offspring) among the assigned queens (Jaffé et al. 2009). The final sample-size-corrected number of queens (estimated number of colonies) was obtained by adding the undetected queens (those with an expected frequency of zero) to the detected ones at each location. In the six samples in which we used only one set of linked markers for genotyping, the sample-size-corrected number of haplotypes found in each population was obtained from the distribution of drones among the assigned haplotypes.

**Census of Managed Hives and Detection of Putative Wild Colonies**

The number of managed hives within a radius of 2.5–3 km from each sampling location were exhaustively surveyed by beekeepers or members of beekeeping organizations. We asked these people to conduct the surveys because of their knowledge of local beekeeping operations. Sites were mapped through the use of free geographic information system software available online (Google Maps). In contrast to official hive censuses, our surveys included “registered” hives, from professional beekeepers, and “unofficial” hives, belonging to unregistered amateurs. Censuses were taken during or shortly after each sampling event and therefore represent the local situation at the time the samples were collected. We subtracted the number of estimated colonies from the number of managed colonies surveyed at each location to infer the putative number of wild (i.e., unmanaged) honeybee colonies.

**Density Estimation**

Honeybee colonies are headed by a single queen (Seeley 1985), and therefore the number of queens detected at a given location provides an estimate of colony density. To calculate this density from the number of queens detected, it is necessary to know the distance over which drones and queens fly. Drones can only fly up to 7 km (Ruttner & Ruttner 1972), and 15 km is the maximal mating distance (drone plus queen flight distance) ever recorded (Jensen et al. 2005). On average, however, drones fly between 600 and 1200 m (Taylor & Rowell 1988). Taylor and Rowell (1988) and Koeniger et al. (2005) report the number of drones recaptured at different distances from source hives. On the basis of these distributions, we calculated an average drone flight distance of 900 m. Therefore, in all subsequent density calculations, we used 900 m as the mean flight distance of drones and 2.5 km² as the mean mating area of drones (assuming the area is circular). Comparing the two approaches for estimating colony number (genotyping drones vs. worker offspring), Moritz et al. (2008) found that the population size estimated with worker offspring is 1.8 times that obtained from a drone sample, presumably due to the greater flight range of queens plus drones. For this reason we considered the queen’s mating area to be 1.8 times larger than that of drones (4.5 km²).

Using the total number of estimated colonies and the mean flight distance of drones to estimate the density of colonies, however, will inevitably overestimate colony density, given that distant colonies contributing drones to the DCA are considered within an average flight distance. Assuming a similar drone investment, distant colonies will contribute fewer drones than colonies located in the vicinity of a DCA, likely to be represented by many drones. For this reason we excluded those colonies represented by less than a median number of drones in all density calculations (the distribution of drones among colonies resembles a Poisson distribution; Baudry et al. 1998). For each location we therefore quantified the number of colonies represented by at least a median number of drones (Supporting Information). Dividing this number by the mean mating area of drones (for the drone samples) or queens (for the worker samples), we obtained an estimate of the local density of colonies at each sampling location.

**Statistical Analyses**

Except for gene diversity ($H_e$) and NDEs, all variables (allelic richness, colony density, temperature, and rainfall) were normally distributed. The $H_e$ and NDE were therefore arcsine and log, transformed, respectively. Homoscedasticity was not violated in any case. Linear regressions were then performed to detect colinearity of variables. Because temperature was positively correlated with colony density and gene diversity, we performed analyses of covariance (ANCOVA) with colony density, gene diversity, and allelic richness as dependent variables, pattern of land use (agricultural land or nature reserve) as a categorical predictor, and temperature as a covariate. We repeated the ANCOVA considering only the 13 samples from European locations. Finally, we used a t test to compare the mean density of honeybee colonies estimated in our 13 European locations with the mean density of managed hives of 35 European countries (calculated from Appendix I in De la Rúa et al. 2009).

**Results**

Genetic diversity measures were negatively correlated with NDE (AR-ln NDE regression’s $\beta = -0.62$, $t_{23} = -3.81$, $p < 0.001$; arcsine $H_e$-ln NDE regression’s $\beta = -0.50$, $t_{23} = -2.80$, $p = 0.01$). NDE, however, was not correlated with colony density (ln NDE-density regression’s $t_{23} = -1.62$, $p = 0.12$). Genetic diversity was highest in South Africa and lowest in Northern Europe (Table 1). Likewise, the highest colony densities were found in South Africa.
Figure 1. *Apis mellifera* colony densities at 10 locations in Africa, 2 locations in central Asia, and 13 locations in Europe. The diameter of the circles is proportional to the density at each location (given within the circle is the number of colonies per square kilometer), and color (yellow or green) indicates the pattern of land use (agricultural landscape or nature reserve respectively). The multicolor legend on the lower left relates to the Köppen climate classification map (adapted from Peel et al. [2007] and used here under the Creative Commons Attribution, Noncommercial and ShareAlike License) (A, tropical/megathermal climates; B, dry [arid and semi-arid] climates; C, temperate/mesothermal climates; D, continental/microthermal climates; E, polar climates [see Supporting Information for a detailed legend]).

Genetic diversity measures correlated positively with colony density (AR-density regression’s beta = 0.81, $t_{23} = 6.64, p < 0.001$; arcsine $H_e$-density regression’s beta = 0.70, $t_{23} = 4.68, p < 0.001$). Whereas average monthly rainfall was not correlated with any of our dependent variables (regression’s $r_{23} < 1, p > 0.4$ in all cases), mean annual temperature was an important predictor of gene diversity (temperature-arcsine $H_e$ regression’s beta = 0.54, $t_{23} = 3.04, p = 0.006$) and of colony density (temperature-density regression’s beta = 0.41, $t_{23} = 2.18, p = 0.04$; Fig. 2), but less so of allelic richness (temperature-AR regression’s beta = 0.39, $t_{23} = 2.04, p = 0.05$). For this reason, we used temperature as a continuous predictor in the subsequent analyses of covariance.

Land use had a significant negative influence on genetic diversity measures and the density of honeybee colonies, even when accounting for temperature variation (AR: $F_{1,22} = 10.60, p = 0.004$; $H_e$: $F_{1,22} = 12.99, p = 0.002$; density: $F_{1,22} = 8.36, p = 0.008$). Overall, allelic richness, gene diversity, and colony density were higher in nature reserves (mean [SD] = 11.88 [4.30], 0.93 [0.12], and 7.23 [2.67], respectively) than in agricultural landscapes (mean [SD] = 7.46 [2.67], 0.74 [0.17],...
and 4.57 [2.03], respectively). Within Europe, however, nature reserves showed similar genetic diversity and colony densities as agricultural landscapes (AR: $F_{1,10} = 0.01; H_e: F_{1,10} = 3.32$; density: $F_{1,10} = 0.02; p \geq 0.1$ in all cases). Putative wild (unmanaged) honeybees in Europe were only identified in Ireland and Italy, where we detected more colonies among genotypic samples than those kept by local beekeepers (Table 1). Finally, the mean density of honeybee colonies estimated across the 13 European sampling locations (Table 1) did not differ from the average density of managed hives reported in a recent survey (De la Rúa et al. 2009) of 33 European countries (estimated mean density [SD] = 4.83 [2.39], mean density of managed hives [SD] = 4.04 [2.82], $t_{44} = 0.89, p = 0.4$).

**Discussion**

Our estimates of the density of honeybee colonies were directly dependent on the number of genetic markers used and their level of polymorphism. Although our detection power was affected by the level of genetic diversity at each location, NDEs and colony densities were not correlated, which shows that NDEs had little impact on our estimates. Sampling biases caused by seasonality, on the other hand, represented a more important source of inaccuracy in our estimation of colony numbers (Hepburn & Radloff 1998; Jaffé et al. 2009). The samples from Europe and central Asia were collected in late spring through summer, near the drone production peak; and hence, we expect to have detected most of the reproductively active colonies at each sampling location.

In our selected African locations, where seasonality is not as clear-cut and pronounced as in temperate regions, avoiding the effects of seasonality was more difficult. Nevertheless, sampling was performed during summer and rainy months, when enough floral resources were available to colonies for growth and reproduction, which minimized the occurrence of colonies not investing in drone production. An additional source of potential bias in our density estimates is that queen and drone flight distances may differ among honeybee subspecies or sampling locations. Unfortunately, data are not available on the impact of environmental or genetic factors on the flight distance of drones or queens. Despite these limitations, the estimated number of colonies remaining undetected because of an insufficient sample was usually low (Table 1). Moreover, our local density estimates matched previously reported values (Otis 1991; Ratnieks et al. 1991; McNally & Schneider 1996; Baum et al. 2008; but see Oldroyd et al. 1997); and hence, we are confident they provide an informative picture of the reproductively active colonies found at our different sampling locations.

In our study genetic diversity and the density of honeybee populations were strongly influenced by the mean annual temperature. Brood production and reproductive swarming are determined by the availability of forage and the ability of bees to gather nectar and pollen (Hepburn & Radloff 1998). Rainfall is an important predictor of forage availability in dry, seasonal ecosystems, such as those of our African locations. In our global analysis, however, rainfall was not correlated either with genetic diversity or colony density. A possible explanation of this finding is that temperature is likely to be a more important predictor of colony densities in temperate climates, where...
Agricultural intensification and forestry reduce the diversities (Biesmeijer et al. 2006; Fitzpatrick et al. 2007). Most European countries have intensive land use, and many life-history traits. African subspecies disperse via long-distance migratory swarms, leave the nest in response to disturbance or disease (absconding) more readily, and have a faster generation time and smaller colonies than European honeybees (reviewed by Schneider et al. 2004). These traits promote population gene flow and high genetic diversity, boosting effective population sizes in Africa (Estoup et al. 1995; Jaffé et al. 2009). The impact of such subspecific differences on colony density estimates, however, is not clear. Whereas the density of wild honeybee colonies in Botswana is <5 colonies/km² (McNally & Schneider 1996; see also Hepburn & Radloff 1998 and references therein), estimates for feral Africanized colonies in the Neotropics span 6–20 colonies/km² (Otis 1991; Ratnieks et al. 1991). These figures overlap with the densities of European colonies we found (<1 to >8 colonies/km²; Fig. 1). Moreover, even though honeybees of African origin replaced European honeybees in the American tropics, an 11-year study in a Texan nature reserve shows that both can establish similar colony densities in temperate regions (Pinto et al. 2004).

Beekeeping practices can also have a profound influence over wild and feral honeybee populations (De la Rúa et al. 2009). Coexistence with managed bees exposes feral and wild populations to bee pathogens and to introgressive hybridization. Apiculture typically involves the concentration of colonies in apiaries, which facilitates the horizontal transfer of pathogens and parasites from colony to colony (Fries & Camazine 2001). This can have catastrophic consequences, as demonstrated by the drastic decline of feral U.S. bee populations after the introduction of parasitic Varroa mites in the 1980s (Kraus & Page 1995; Needham et al. 1988). Indeed, honeybee colonies in northern latitudes do not usually survive infestation by Varroa mites without treatment (Fries et al. 2006). Given the extent of the beekeeping industry in Europe, human-mediated disease dissemination is expected to cause important declines in European honeybee populations (De la Rúa et al. 2009).

In addition to hosting an extensive beekeeping industry, most European countries have intensive land use. The human-mediated destruction of natural habitats has led to a progressive reduction of suitable habitats for honeybees, with a negative impact on feral and wild populations (Biesmeijer et al. 2006; Fitzpatrick et al. 2007). Agricultural intensification and forestry reduce the diversity and abundance of native bees in the United States, which diminishes their pollination services by 3- to 6-fold (Kremen et al. 2002). In support of these findings, we found lower genetic diversity and colony densities in agricultural landscapes than in nature reserves. Restricting the analysis to Europe, however, erased the significant effect of land use, making agricultural landscapes and nature reserves indistinguishable in terms of genetic diversity or colony density. This may partially be due to the fact that variation in land-use patterns is much larger between than within continents. We nevertheless caution that other factors that we did not consider are also likely to affect the relationship between land use and the density of honeybee colonies (e.g., Winfree et al. 2007). More thorough comparisons, with larger sample sizes and that control for multiple factors, are needed to assess the extent to which land use is detrimental to honeybee populations. Only two of our sampling locations provided a test case to address the effect of land use on honeybee colony densities: the deforested farmland of Elsenburg and the neighboring nature reserve of Jonkershoek, 20 km apart, where no beekeeping takes place. Both share similar climatic conditions and a honeybee subspecies (A. m. capensis) that is tolerant to Varroa mites (Allsopp 2006). Habitat destruction thus seems to be the main factor responsible for the lower colony density observed in Elsenburg.

Another explanation for the lack of a difference in genetic diversity and colony densities between agricultural landscapes and nature reserves in Europe is that wild honeybee populations may be absent from nature reserves. For instance, we did not detect a higher number of colonies in most European sampling locations compared with those kept by local beekeepers. Only in the Italian nature reserve surrounding Farindola, did we detect a large excess of colonies compared with the number kept by local beekeepers (Table 1). Hence, we were more confident about the presence of wild bees in this location. At the European scale, our mean estimated density of honeybee colonies closely resembled the average density of managed hives reported by De la Rúa et al. (2009), which suggests that the European honeybee populations we studied here were mainly composed of managed bees. As disease dissemination and land use continue to threaten European honeybees (Brown & Paxton 2009), only populations managed by beekeepers are likely to survive. Although we cannot discount apiculture itself as involved in the decline of wild European honeybees, our data suggest that without beekeeping, honeybees in Europe would be close to extinction. Our estimates of honeybee population density across Europe were in the same range as those found in the adverse climatic conditions of the Kalahari and Saharan deserts (Fig. 1), which suggests that beekeeping activities do not compensate for the loss of wild honeybee colonies. Moreover, the current pollination services provided by honeybees might
not fulfill agricultural demand (Aizen & Harder 2009). Added to the parallel decline of other pollinators, honeybee losses could therefore seriously threaten pollination services and ecosystem functioning in Europe. Because honeybees are “rescue pollinators” and thus able to compensate for the decline of native pollinators in degraded habitats (Aizen & Feinsinger 1994; Dick 2001), we suggest that promoting beekeeping activities could compensate for habitat loss and ensure adequate pollination of wild plants and crops. In the light of severe declines in beekeeping activities in various European countries (COLOSS 2009), we recommend that beekeeping with local subspecies should be encouraged to preserve native genetic diversity and to reduce the transmission of pathogens throughout the continent (De la Rúa et al. 2009). A first step toward this could be to reconsider the conservation status of honeybees in Europe and to regard beekeeping not only as a profitable business for producing honey, but also as an essential component of biodiversity conservation.

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Supporting Information

The coordinates and climatic data for all sampling locations (Appendix S1), a schematic representation of our approach to estimate honeybee colony densities (Appendix S2), and the Köppen climate classification detailed legend (Appendix S3) are available as part of the on-line article. The author is responsible for the content and functionality of these materials. Queries (other than absence of the material) should be directed to the corresponding author.

Literature Cited


