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Analysis of honey environmental DNA indicates that the honey bee (Apis mellifera L.) trypanosome parasite Lotmaria passim is widespread in the apiaries of the North of Italy

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1	Short	Commun	ication

3	Analysis of honey environmental DNA indicates that the honey bee (Apis mellifera L.)
4	trypanosome parasite Lotmaria passim is widespread in the apiaries of the North of Italy
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16 Highlights

17

- 18 Axenic culture of *L. passim* from Italian isolates was established
- We analysed environmental DNA extracted from honey samples to detect the presence of L.

20 *passim*

• L. passim was present in 78% of the honey samples collected in the North of Italy

23 Abstract

24 Lotmaria passim is a trypanosomatid that infects honey bees. In this study, we established an axenic culture of L. passim from Italian isolates and then used its DNA as a control in subsequent 25 analyses that investigated environmental DNA (eDNA) to detect this trypasonosomatid. The source 26 of eDNA was honey, which has been already demonstrated to be useful to detect honey bee parasites. 27 DNA from a total of 164 honey samples collected in the North of Italy was amplified with three L. 28 passim specific PCR primers and 78% of the analysed samples gave positive results. These results 29 indicated a high prevalence rate of this trypanosomatid in the North of Italy, where it might be 30 considered another threat to honey bee health. 31

- 32
- **33 Keywords:** health/monitoring/parasite/PCR/Trypanosomatidae.

34 1. Introduction

35 Analysis of environmental DNA (eDNA), defined as DNA obtained directly from environmental-related matrixes or samples, is considered an efficient, non-invasive and easy-to-36 standardize methodology that has been proposed to facilitate the detection and monitoring of cryptic 37 or invasive organisms, that would be expensive or difficult to be sampled and identified with other 38 methods (Taberlet et al., 2012; Bohmann et al., 2014; Bass et al., 2015; Ribani et al., 2020). 39 40 Environmental DNA can extend and facilitate the possibility to evaluate the distribution of elusive organisms, including parasites, even over time and geographic areas (Thomsen and Willerslev, 2015). 41 Different analytical approaches based on eDNA have been tested according to the objective, the 42 43 methods of specimen sampling and the targeted organisms (Jerde et al., 2011; Jain et al., 2013; Wilcox et al., 2013). 44

Honey is a natural matrix produced by honey bees (Apis mellifera L.) from two main types of 45 secretions derived directly (i.e. nectar) or indirectly (i.e. honeydew) from plants. This product is 46 mainly made by sugars with other minor components, including DNA, a component that is usually 47 neglected even if it includes interesting information that can provide environment-derived 48 fingerprints (Utzeri et al., 2018a, 2018b; Bovo et al., 2018, 2020). That means that honey DNA 49 derives from all organisms that directly or indirectly are involved in its production or that are part of 50 51 the ecological niche in which it is produced, including parasites and pathogens of A. mellifera (Bakonyi et al., 2003; Lauro et al., 2003; McKee et al., 2003; D'Alessandro et al., 2007; Ribani et al., 52 2020). We recently demonstrated that this eDNA can be used for monitoring purposes of some main 53 54 health threats of honey bees, simplifying the possibility to obtain information on the incidence and distribution of honey bee pathogens and parasites (Bovo et al., 2018, 2020; Utzeri et al., 2019; Ribani 55 et al., 2020). 56

57 The trypanosomatid *Lotmaria passim* is a unicellular obligate parasite that infects honey bees. 58 *Lotmaria passim* has been only recently well characterized and distinguished from other 59 trypanosomatids, particularly from *Crithidia mellificae* (Schwarz et al., 2015). In-depth molecular 60 analyses have contributed to clarify that previous studies and published DNA sequences assigned to C. mellificae actually belonged to L. passim, that should be also considered the predominant and 61 widespread trypanosomatid of A. mellifera (Cepero et al., 2014; Ravoet et al., 2015; Schwarz et al., 62 2015). Based on these studies, a few molecular methods have been proposed to detect L. passim 63 (Arismendi et al., 2016; 2020; Stevanovic et al., 2016; Vejnovic et al., 2018). Few studies have, 64 however, reported detailed information about the distribution and prevalence of this trypanosomatid 65 in different parts of the world. Recent reports have evaluated the presence and prevalence of L. passim 66 in some European countries, USA and South America (Arismendi et al., 2016; Stevanovic et al., 2016; 67 Vargas et al., 2017; Castelli et al., 2019; Williams et al., 2019; Michalczyk et al., 2020). At present 68 69 there is no detailed information in Italy, apart from a preliminary investigation that we carried out 70 using honey eDNA (Ribani et al., 2020).

In this study, after having characterized an axenic culture of *L. passim* that was used as source of control DNA in PCR analyses, we took advantage from the possibility to use honey eDNA to obtain a more detailed and specific analysis of the distribution of *L. passim* in the North of Italy.

74

75 2. Materials and methods

76 2.1. Isolation and characterization of *Lotmaria passim*

77 Sixty A. mellifera workers were collected from an apiary in the province of Bologna (Italy), where in the past the presence of intestinal flagellates had been preliminarily microscopically 78 observed. Each honey bee was immobilized by chilling at -20 °C for 4-5 min, briefly washed in 99% 79 80 ethanol, and decapitated prior to dissection in a sterile environment. The intestine was removed with sterile tools, submerged in 0.5 mL of supplemented DS2 medium [Insectagro DS2 serum free/protein 81 free medium without L glutamine (CorningTM, NY, USA) plus 5% fetal bovine serum (Sigma Aldrich, 82 St. Louis, MO, USA) and 1% Antibiotic/Antimycotic solution (Sigma Aldrich, St.Louis, MO, USA)] 83 in a 1.5-mL microtube, gently macerated with a sterile pestle and incubated at 26 °C. Ten µL of each 84 culture was observed on wet mount slide, at light microscope with 400× objective, after 24-48 and 85

86 72 hours to verify the presence of free active flagellates. One out of 60 honey bees examined 87 individually was positive for flagellates. The established active culture was expanded in 88 supplemented DS2 medium and maintained by subculture steps every 4-10 days in fresh medium 89 (ratio 1:5). The procedure was a modification of the protocols reported by Runckel et al. (2011) and 90 Schwartz et al. (2015).

Morphological observations and image acquisition were performed on active cultures, both on wet and stained slides, at 400× and 1000× magnification through Leica DMLS light microscope (Leica, Wetzlar, Germany), equipped with digital camera Nikon DS-Fi2 with imaging software NIS Elements 4.10.01 (Nikon, Tokyo, Japan). The first type of observations was obtained from a drop of culture (10 μ L) under coverslip whereas the second type was derived from a thin film of the culture that was air-dried quickly and May-Grunwald Giemsa or Giemsa stained.

97 Cultures at peak density were centrifuged at 2900 rpm for 15 min. The obtained pellet was 98 washed twice in Phosphate Buffered Saline (PBS), resuspended in 200 μ L of PBS and cryopreserved 99 at -20 °C until DNA extraction.

100

101 2.2 Honey samples

A total of 164 honey samples, produced in 2018, were directly provided by beekeepers. The samples derived from apiaries located in all regions of the North of Italy (Liguria, n. 6; Piedmont, n. 10; Valle d'Aosta, n. 8; Lombardia, n. 10; Emilia-Romagna, n. 100; Trentino Alto Adige, n. 10; 105 Veneto, n. 10; and Friuli Venezia Giulia, n. 10). Geographic coordinates were used to locate the 106 production sites. Detailed information on the analysed samples is reported in Table S1.

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108 2.3. DNA extraction from axenic cultures and from honey samples

109 DNA was extracted from the cultivated flagellates, subsequently identified as being from L.

110 passim, using the Wizard Genomic DNA Purification kit (Promega Corporation, Madison, WI, USA)

111 following the Tissue Culture Cells manufacturer's protocol, starting from 300 µL of the axenic culture

112 containing about 1×10^6 cells. Isolated DNA from cell culture was resuspended in 30 µL of sterile 113 water and stored at -20 °C for further analyses. The same protocol was used to extract DNA from a 114 standard cell culture of *Crithidia mellificae*, purchased from ATCC (ATCC 30254).

DNA extraction from honey samples was based on the protocol described by Utzeri et al. 115 (2018c). Briefly, starting from a total of 50 g of honey divided in four 50 mL tubes, 40 mL of ultrapure 116 water was added to each tube, vortexed and incubated at 40 °C for 30 minutes. Then tubes were 117 118 centrifuged for 25 min at 5000 g at room temperature and the supernatant was discarded. The pellet was resuspended in 5 mL of ultrapure water and the content of the four tubes was merged in one and 119 then diluted with ultrapure water. After centrifugation for 25 min at 5000 g at room temperature, the 120 121 supernatant was discarded, and the pellet was resuspended in 0,5 mL of ultrapure water. The DNA 122 extraction protocol included the following steps: 1) 1 mL of CTAB extraction buffer [2% (w/v) cetyltrimethylammoniumbromide; 1.4 M NaCl; 100 mM Tris-HCl; 20 mM EDTA; pH 8] and 5 µL of 123 RNase A solution (10 mg/mL) were added to each honey pre-treated sample; 2) samples were 124 incubated for 10 min at 60 °C and, after the incubation, 30 µL of proteinase K (20 mg/mL) were 125 added; 3) subsequently, samples were incubated at 65° C for 90 min with gentle mixing; 4) then, 126 samples were cooled at room temperature and centrifuged for 10 min at 16,000 g; 5) 700 µL of the 127 resulting supernatant was transferred in a tube containing 500 µL of chloroform/isoamyl alcohol 128 129 (24:1), mixed by vortexing and centrifugated for 15 min (16,000 g at room temperature); 6) the supernatant was transferred in a new 1.5 mL tube and the DNA was precipitated with 500 µL of 130 isopropanol and washed with 500 µL of ethanol 70%; 7) the resulting DNA pellets were rehydrated 131 132 with 30 μ L of sterile H₂O and stored at -20°C until PCR analyses.

Extracted DNA was quality checked using the nanophotometer IMPLEN P300 (Implen GmbH,
Munchen, Germany) and visually inspected by 1% agarose gel electrophoresis in TBE 1X buffer after
staining with 1X GelRed Nucleic Acid Gel Stain (Biotium Inc., Hayward, CA, USA).

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137 2.4. Primer pairs, PCR analyses, sequencing and sequence data analyses

Primer pairs used in this study and PCR conditions are listed in Table S2. To assess the 138 possibility to successfully amplify the extracted DNA from honey, we first verified if amplification 139 could occur for honey bee DNA using primers designed on the mitochondrial DNA (mtDNA) region 140 of Apis mellifera to amplify a short fragment that had an amplification success rate from this source 141 of DNA equal to 100% (Utzeri et al., 2018c). Three primer pairs were then used to specifically 142 amplify L. passim DNA. One pair that targets the cytochrome b (CYTB) gene of L. passim was from 143 Stevanovic et al. (2016). Two other primer pairs were re-defined from the work of Arismendi et al. 144 (2016) to amplify two shorter DNA fragments from L. passim than those reported in the mentioned 145 study. This was needed to facilitate the amplification of the DNA from the degraded honey DNA. Of 146 147 the two new primer pairs derived from that work, the pair that amplified the 18S fragment included the same reverse primer of Arismendi et al. (2016) and a new forward primer whereas the pair that 148 amplified the GAPDH fragment included the same forward primer of Arismendi et al. (2016) and a 149 150 new reverse primer. The new primers were designed to have a 100% match with the corresponding L. passim sequence (accession number KM066244) and to have several mismatches and frameshifts 151 in the homologous gene regions of C. mellificae (e.g. accession number KJ713345). Primers were 152 (http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi). 153 selected using PRIMER3 Additional PCR primers derived from studies that specifically targeted *Crithidia spp.* were also used: 154 155 a pair of primers that specifically amplifies a short region of the C. mellificae GAPDH gene and a pair of primers that specifically targets a short fragment of the C. bombi TOPII locus (Bartolomé et 156 al., 2018; Table S2). 157

PCR were performed on a 2700 Thermal Cycler (Life Technologies, Carlsbad, CA, USA) in a total volume of 20 μ L using KAPA HiFi HotStart Mastermix (Roche, Basel, Switzerland) with the following PCR profile: initial denaturation step at 95 °C for 3 min, then 35 cycles of alternate temperatures (20 s at 98 °C, 15 s at the specific annealing temperature for the different primer pairs as indicated in Table S2, 30 s at 72 °C), followed by a final extension step at 72 °C for 1 min. Amplified DNA fragments were electrophoresed in 2.5 % agarose gels in TBE 1× buffer and stained with 1× GelRed Nucleic Acid Gel Stain (Biotium Inc., Hayward, CA, USA). Primer specificity was
tested using DNA extracted from axenic cultures of *L. passim* (reported above) and *C. mellificae*(ATCC 30254). All two primer pairs specific for one of the two targeted *Crithidia spp*. (Table S2)
amplified this *C. mellificae* DNA.

Amplified fragments from pure cell culture DNA and from 30 honey DNA samples (7 µL of 168 PCR product) were purified with 1 µL of ExoSAP-IT® (USB Corporation, Cleveland, OH, USA) for 169 15 min at 37 °C and then sequenced using the BrightDye® Terminator Cycle Sequencing Kit 170 (NIMAGEN, Nijmegen, the Netherlands). Sequencing reactions were purified using EDTA 0.125 M, 171 ethanol 100% and ethanol 70%, following a standard protocol, and then were loaded on an ABI3100 172 173 Avant Genetic Analyzer sequencer for detection of DNA sequences (Life Technologies, Carlsbad, 174 CA, USA). Obtained electropherograms were visually inspected using MEGA 7 (Kumar et al., 2016) **BLAST**n algorithm the online platform 175 and the was run on BLAST (http://www.ncbi.nlm.nih.gov/BLAST/) in order to compare and validate the assignment of the 176 obtained DNA sequences to the correct organism. 177

178

179 3. Results and discussion

We first established and characterized an axenic culture of a trypanosomatid, that according to 180 181 its features (Figure 1) was clearly consistent with L. passim, as described by Schwartz et al. (2015). However, while the morphology of the flagellated stage of C. mellificae and L. passim can be 182 generally useful to discriminate these two species within honey bees, cryptic species may be present 183 184 that cannot be distinguished from one another by cell morphology, supporting the need for a genetic confirmation (Schwartz et al., 2015), that was subsequently carried out by PCR analysis and 185 sequencing with primer pairs specifically designed and here tested for this purpose (Table S2). All 186 three primer pairs designed on L. passim (Table S2) produced the expected amplicons in the PCR 187 analyses of this newly established isolate, with 100% identity with the corresponding GenBank 188 entries (cytb: Accession no. MG494247, E-values = 8×10^{-115} ; 18S: MG182398, E-values = 9×10^{-74} ; 189

190 GAPDH: KX953207, E-values = 2×10^{-63}). The established axenic culture was then used to obtain 191 DNA useful for the validation of PCR tests designed to identify the presence of *L. passim* in Italian 192 colonies using its DNA footprint recovered from honey.

The same PCR primers were used to amplify the DNA extracted from honey produced in all regions of the North of Italy by 164 different beekeepers (Figure 2; Table S1). For all samples, honey bee DNA was always amplified confirming that the extracted DNA was not completely degraded and could be used for subsequent analyses (Ribani et al., 2020).

Positive samples for L. passim were then considered those honey samples from which at least 197 two out of three primer pairs obtained a clearly visible amplified fragment of the expected size, 198 199 according to the criteria applied for general pathogen diagnosis based on PCR detection assays (Sachse, 2004). A total of 128 honey samples (78%) gave positive results for this trypasonomatid 200 using the tested primer pairs. For 90% of the 128 positive samples (115 honey samples), all three 201 202 primer pairs produced the expected amplicons. Among the remaining 13 honey samples from which only two primer pairs tested returned an amplified fragment, the largest fragment of 247 bp was not 203 amplified in 10 of them, indicating that the lack of amplification could be due to a highly degraded 204 DNA that was isolated from these problematic samples. All sequenced fragment obtained from 205 amplification of honey DNA had the same sequence already reported from the DNA of the established 206 207 axenic cell culture. None of the other two primer pairs designed on Crithidia spp. sequences produced any amplified fragment from honey extracted DNA, further confirming that L. passim is the prevalent 208 trypasonomatid parasite in this part of Italy. Table 1 reports the percentage of positive honey samples 209 210 divided by administrative regions of the North of Italy. Even if the number of tested samples was not proportional to the geographic areas, the frequency of positive samples ranged from 33 to 100% in 211 Liguria and Friuli Venezia Giulia regions, respectively. In the region with the highest number of 212 samples (Emilia-Romagna, n. 100), 88% were positive. These results are in line with annual 213 frequencies reported for Serbia (38.9% to 83.3%) for period 2007-2015 (Stevanovic et al., 2016). 214

Other studies, that however were based on DNA assays applied on individual bees, detected

frequently *C. mellificae* and *C. bombi* on *Apis mellifera* (Graystock et al., 2015; Bartolomè et al., 2018) even if *L. passim* resulted the prevalent trypasonomatid in all parts of the world where the prevalence of this parasite has been evaluated at large scale (e.g. Castelli et al., 2019; William et al., 2019). The results we obtained for the North of Italy support the global and almost housekeeping presence of *L. passim* also in the Italian Peninsula.

221

222 4. Conclusions

The trypanosome *L. passim* has been suggested as an emerging potential contributor to honey bee health decline even if its role has not been completely clarified yet. In this study we first established an axenic culture of *L. passim*. Then, we demonstrated the usefulness of honey eDNA in monitoring studies and established the first prevalence map of *L. passim* in the North of Italy. The study indicated that this trypanosomatid was present in almost all apiaries from which honey samples were collected.

229

230 5. Declaration of interest

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

233

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241 References

- Arismendi, N., Bruna, A., Zapata, N., Vargas, M., 2016. PCR-specific detection of recently described
 Lotmaria passim (Trypanosomatidae) in Chilean apiaries. J. Invertebr. Pathol. 134, 1–5.
 https://doi.org/10.1016/j.jip.2015.12.008.
- Arismendi, N., Caro, S., Castro, M.P., Vargas, M., Riveros, G., Venegas, T., 2020. Impact of mixed
 infections of gut parasites *Lotmaria passim* and *Nosema ceranae* on the lifespan and immune-
- related biomarkers in *Apis mellifera*. Insects 11, 420. <u>https://doi.org/10.3390/insects11070420</u>.
- Bakonyi, T., Derakhshifar, I., Grabensteiner, E., Nowotny, N., 2003. Development and evaluation of
 PCR assays for the detection of *Paenibacillus larvae* in honey samples: comparison with
 isolation and biochemical characterization. Appl. Environ. Microb. 69, 1504-1510.
 https://doi.org/10.1128/AEM.69.3.1504-1510.2003.
- Bartolomé, C., Buendía, M., Benito, M., De la Rúa, P., Ornosa, C., Martín-Hernández, R., Higes, M., 252 Maside, X. 2018. A new multiplex PCR protocol to detect mixed trypanosomatid infections in 253 species of Apis and Bombus. J. Invertebr. Pathol. 154, 37-41. 254 https://doi.org/10.1016/j.jip.2018.03.015. 255
- Bass, D., Stentiford, G.D., Littlewood, D.T.J., Hartikainen, H., 2015. Diverse applications of
 environmental DNA methods in parasitology. Trends Parasitol, 31, 499-513.
 https://doi.org/10.1016/j.pt.2015.06.013.
- Bohmann, K., Evans, A., Gilbert, M.T.P., Carvalho, G.R., Creer, S., Knapp, M., Yu D.W., De Bruyn,
 M. 2014. Environmental DNA for wildlife biology and biodiversity monitoring. Trends Ecol.
 Evol, 29, 358-367. https://doi.org/10.1016/j.tree.2014.04.003.
- Bovo, S., Ribani, A., Utzeri, V.J., Schiavo, G., Bertolini, F., Fontanesi, L., 2018. Shotgun
 metagenomics of honey DNA: Evaluation of a methodological approach to describe a multikingdom honey bee derived environmental DNA signature. PLoS One 13, e0205575.
 <u>https://doi.org/10.1371/journal.pone.0205575</u>.

- Bovo, S., Utzeri, V.J., Ribani, A., Cabbri, R., Fontanesi, L., 2020. Shotgun sequencing of honey DNA
 can describe honey bee derived environmental signatures and the honey bee hologenome
 complexity. Sci. Rep. 10, 1-17. <u>https://doi.org/10.1038/s41598-020-66127-1</u>.
- 269 Castelli, L., Branchiccela, B., Invernizzi, C., Tomasco, I., Basualdo, M., Rodriguez, M., Zunino, P.,
- 270 Antúnez, K. 2019. Detection of *Lotmaria passim* in Africanized and European honey bees from
- 271 Uruguay, Argentina and Chile. J. Invertebr. Pathol, 160, 95-97.
 272 https://doi.org/10.1016/j.jip.2018.11.004.
- 273 Cepero, A., Ravoet, J., Gómez-Moracho, T., Bernal, J. L., Del Nozal, M. J., Bartolomé, C., Maside,
- 274 X., Meana, A., Gonzàlez-Porto, A. V., De Graaf, D. C., Martìn-Hernàndez, R., Higes, M. 2014
- Holistic screening of collapsing honey bee colonies in Spain: a case study. BMC Res. Notes 7,
- 276 649. <u>https://doi.org/10.1186/1756-0500-7-649</u>
- D'Alessandro, B., Antúnez, K., Piccini, C., Zunino, P., 2007. DNA extraction and PCR detection of 277 Paenibacillus larvae spores from naturally contaminated honey and bees using spore-decoating 278 freeze-thawing techniques. World J. Microbiol. Biotechnol. 23, 593-597. 279 and https://doi.org/10.1007/s11274-006-9261-y. 280
- Graystock, P., Goulson, D., Hughes, W. O. 2015. Parasites in bloom: flowers aid dispersal and
 transmission of pollinator parasites within and between bee species. Proc. Biol. Sci. 282,
 20151371. https://doi.org/10.1098/rspb.2015.1371.
- Jain, S.A., Jesus, F.T.D., Marchioro, G.M., Araújo, E.D.D. 2013. Extraction of DNA from honey and
 its amplification by PCR for botanical identification. Food Sci. Technol. (Campinas) 33, 753756. https://doi.org/10.1590/S0101-20612013000400022.
- Jerde, C. L., Mahon, A. R., Chadderton, W. L., Lodge, D. M., 2011. "Sight-unseen" detection of rare
 aquatic species using environmental DNA. Conserv. Lett. 4, 150-157.
 https://doi.org/10.1111/j.1755-263X.2010.00158.x.

- Langridge, D.F., McGhee, R.B., 1967. *Crithidia mellificae* n. sp. an acidophilic trypanosomatid of
 the honey bee *Apis mellifera*. J. Protozool, 14, 485-487. https://doi.org/10.1111/j.15507408.1967.tb02033.x
- Lauro, F.M., Favaretto, M., Covolo, L., Rassu, M., Bertoloni, G., 2003. Rapid detection of
 Paenibacillus larvae from honey and hive samples with a novel nested PCR protocol. Int J
 Food Microbiol, 81, 195-201. https://doi.org/10.1016/S0168-1605(02)00257-X.
- Michalczyk, M., Bancerz-Kisiel, A., Sokół, R. 2020. *Lotmaria passim* as third Parasite
 gastrointestinal tract of honey bees living in tree trunk. J. Apic. Sci., 64, 143-151.
 https://doi.org/10.2478/jas-2020-0012
- McKee, B.A., Djordjevic, S.P., Goodman, R.D., Hornitzky, M.A., 2003. The detection of
 Melissococcus pluton in honey bees (*Apis mellifera*) and their products using a hemi-nested
 PCR. Apidologie 34, 19-27. <u>https://doi.org/10.1051/apido:2002047</u>.
- 302 Ravoet, J., Schwarz, R.S., Descamps, T., Yañez, O., Tozkar, C.O., Martin-Hernandez, R., Bartolomé,
- C., De Smet, L., Higes, M., Wenseleers, T., Schmid-Hempel, R., Neumann, P., Kadowaki, T.,
 Evans, J.D., de Graaf, D., 2015. Differential diagnosis of the honey bee trypanosomatids *Crithidia mellificae* and *Lotmaria passim*. J. Invertebr. Pathol. 130, 21–27.
 http://dx.doi.org/10.1016/j.jip.2015.06.007.
- Ribani, A., Utzeri, V.J., Taurisano, V., Fontanesi, L. 2020. Honey as a source of environmental DNA
 for the detection and monitoring of honey bee pathogens and parasites. Vet. Sci. 7, 113.
 https://doi.org/10.3390/vetsci7030113.
- 310 Runckel, C., Flenniken, M.L., Engel, J.C., Ruby, J.G., Ganem, D., Andino, R., DeRisi, J.L., 2011.
- Temporal analysis of the honey bee microbiome reveals four novel viruses and seasonal prevalence of known viruses, *Nosema*, and *Crithidia*. PLoS One 6, e20656. https://doi.org/10.1371/journal.pone.0020656
- 314 Sachse, K., 2004. Specificity and performance of PCR detection assays for microbial pathogens. Mol
- Biotechnol, 26, 61-79. <u>https://doi.org/10.1385/MB:26:1:61</u>.

Schwarz, R.S., Bauchan, G., Murphy, C., Ravoet, J., de Graaf, D.C., Evans, J.D., 2015.
Characterization of two species of Trypanosomatidae from the honey bee *Apis mellifera*: *Crithidia mellificae* Langridge and McGhee, 1967 and *Lotmaria passim* n. gen., n. sp. J.
Eukaryot. Microbiol. 62, 567–583. http://dx.doi.org/10.1111/jeu.12209.

- 320 Stevanovic, J., Schwarz, R.S., Vejnovic, B., Evans, J.D., Irwin, R.E., Glavinic, U., Stanimirovic, Z.,
- 321 2016. Species-specific diagnostics of *Apis mellifera* trypanosomatids: A nine-year survey
- 322 (2007–2015) for trypanosomatids and microsporidians in Serbian honey bees. J. Invertebr.
- 323 Pathol. 139, 6-11. <u>https://doi.org/10.1016/j.jip.2016.07.001</u>.
- Taberlet, P., Coissac, E., Hajibabaei, M., Rieseberg, L.H., 2012. Environmental DNA. Towards next generation biodiversity assessment using DNA metabarcoding. Mol. Ecol. 21, 1789-1793.
- 326 <u>https://doi.org/10.1111/j.1365-294X.2012.05470.x</u>.
- Thomsen, P.F., Willerslev, E., 2015. Environmental DNA–An emerging tool in conservation for
 monitoring past and present biodiversity. Biol. Conserv. 183, 4-18.
 <u>https://doi.org/10.1016/j.biocon.2014.11.019</u>
- Utzeri, V.J., Ribani, A., Fontanesi, L., 2018a. Authentication of honey based on a DNA method to
 differentiate *Apis mellifera* subspecies: Application to Sicilian honey bee (*A. m. siciliana*) and
- 332 Iberian honey bee (*A. m. iberiensis*) honeys. Food Control 91, 294-301.
 333 <u>https://doi.org/10.1016/j.foodcont.2018.04.010</u>.
- Utzeri, V.J., Ribani, A., Schiavo, G., Bertolini, F., Bovo, S., Fontanesi, L., 2018b. Application of next
 generation semiconductor based sequencing to detect the botanical composition of monofloral,
- polyfloral and honeydew honey. Food Control 86, 342-349.
 https://doi.org/10.1016/j.foodcont.2017.11.033.
- Utzeri, V.J., Schiavo, G., Ribani, A., Tinarelli, S., Bertolini, F., Bovo, S., Fontanesi, L., 2018c.
 Entomological signatures in honey: an environmental DNA metabarcoding approach can
 disclose information on plant-sucking insects in agricultural and forest landscapes. Sci. Rep. 8,
- 341 1-13. <u>https://doi.org/10.1038/s41598-018-27933-w</u>.

342	Utzeri, V.J., Schiavo, G., Ribani, A., Bertolini, F., Bovo, S., Fontanesi, L., 2019. A next generation
343	sequencing approach for targeted Varroa destructor (Acari: Varroidae) mitochondrial DNA
344	analysis based on honey derived environmental DNA. J. Invertebr. Pathol. 161, 47-53.
345	https://doi.org/10.1016/j.jip.2019.01.005

- 346 Vargas, M., Arismendi, N., Riveros, G., Zapata, N., Bruna, A., Vidal, M., Rodríguez, M., Gerding,
- M., 2017. Viral and intestinal diseases detected in *Apis mellifera* in Central and Southern Chile.
 Chilean J. Agr. Res. 77, 243-249. http://dx.doi.org/10.4067/S0718-58392017000300243.

Vejnovic, B., Stevanovic, J., Schwarz, R.S., Aleksic, N., Mirilovic, M., Jovanovic, N.M.,
Stanimirovic, Z., 2018. Quantitative PCR assessment of *Lotmaria passim* in *Apis mellifera*colonies co-infected naturally with *Nosema ceranae*. J. Invertebr. Pathol, 151, 76-81.
<u>https://doi.org/10.1016/j.jip.2017.11.003</u>.

353 Wilcox, T.M., McKelvey, K.S., Young, M.K., Jane, S.F., Lowe, W.H., Whiteley, A.R., Schwartz,

M.K., 2013. Robust detection of rare species using environmental DNA: The importance of primer specificity. PLoS One 8, e59520. <u>https://doi.org/10.1371/journal.pone.0059520</u>.

- 356 Williams, M.K.F., Tripodi, A.D., Szalanski, A.L. 2019. Molecular survey for the honey bee (Apis
- 357 *mellifera* L.) trypanosome parasites *Crithidia mellificae* and *Lotmaria passim*. J. Apicul. Res.
- 358 58, 553-558. https://doi.org/10.1080/00218839.2019.1568956.

- **Table 1.** Frequency of honey samples produced in different regions of the North of Italy that were
- 361 positive for *Lotmaria passim*.

Region	No. of samples	Samples positive to <i>L. pas-</i> sim	% of positive samples
Valle D'Aosta	8	5	62.5%
Piedmont	10	6	60.0%
Liguria	6	2	33.3%
Lombardia	10	5	50.0%
Emilia-Romagna	100	88	88.0%
Trentino-Alto Adige	10	7	70.0%
Veneto	10	5	50.0%
Friuli-Venezia Giulia	10	10	100.0%
Total	164	128	78.0%

Figure 1. May Grunwald-Giemsa stained slides from culture: a) cells aggregates known as "rosettes";
b) morphological polymorphism; c, d, e) promastigote morphotype tear-drop shaped, with short
caudate (tail-like) posterior extension; f) transitional variants and spheroid stage; g, h) spheroids
forms in old cultures.

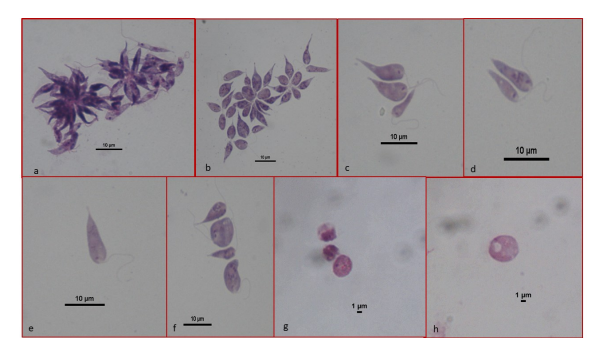
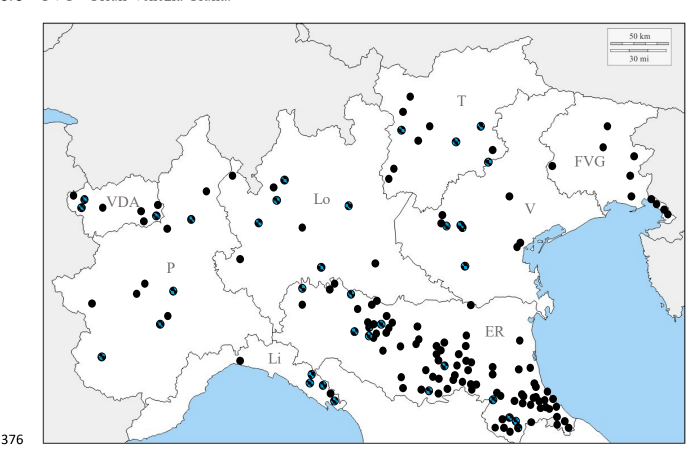


Figure 2. Geographic distribution of the analysed honey samples that were tested for the presence
of *L. passim* DNA. Full black dots: samples that were positive; black and blue striped dots: samples
that were negative. The initial of the region name are shown: VDA = Valle D'Aosta; P = Piedmont;
Lo = Lombardia; Li= Liguria; ER = Emilia-Romagna; V = Veneto; TAA = Trentino-Alto Adige;
FVG = Friuli-Venezia Giulia.



378 Supplementary Material

Table S1. List of honey samples used in this study: regions, localities and provinces of origin, year of production and presence/absence of *Lotmaria*

passim DNA.

Sample	Honey	Region	Locality	Municipality	Province	Year of pro- duction	Lotmaria pas- sim presence
1	Mixed flower from Alpes	Valle D'Aosta	La Thuile (Mont du Parc)	La Thuile	Aosta	2018	Absent
2	Silver fir honeydew	Valle D'Aosta	Les Combes	Introd	Aosta	2018	Present
3	Rhododendron	Valle D'Aosta	La Joux	La Salle	Aosta	2018	Absent
4	Mixed flower from Alpes	Valle D'Aosta	Barasson	St Oyen	Aosta	2018	Present
5	Mixed flower from Alpes	Valle D'Aosta	Fontaney	Challand Saint Victor	Aosta	2018	Present
6	Rhododendron	Valle D'Aosta	Niel	Gaby	Aosta	2018	Present
7	Rhododendron	Valle D'Aosta	Fontainemore	Fontainemore	Aosta	2018	Absent
8	Mixed flower	Valle D'Aosta	Breil	Perloz	Aosta	2018	Present
9	Mixed flower	Piedmont	Parco delle Vallere	Moncalieri	Torino	2018	Present
10	Acacia (Robinia)	Piedmont	Bra	Bra	Cuneo	2018	Absent
11	Honeydew (Forest)	Piedmont	Pocapaglia	Pocapaglia	Cuneo	2018	Present
12	Mixed flower	Piedmont	Mongrando	Mongrando	Biella	2018	Present
13	Acacia (Robinia)	Piedmont	Vinovo (Fraz. Tetti Rosa)	Vinovo	Torino	2018	Present
14	Rhododendron	Piedmont	Fraz. Chiotti	Castelmagno	Cuneo	2018	Absent
15	Chestnut	Piedmont	Ameno	Ameno	Novara	2018	Present
16	Cherry	Piedmont	Mondoni	San Germano Chisoni	Torino	2018	Present
17	Acacia (Robinia)	Piedmont	Vallone	Cossato	Biella	2018	Absent
18	Acacia (Robinia)	Piedmont	Roncaglia	Pralormo	Torino	2018	Absent
19	Mixed flower	Liguria	Beverino	Beverino	La Spezia	2018	Absent
20	Mixed flower	Liguria	Usurana	Calice al Cornoviglio	La Spezia	2018	Absent
21	Acacia (Robinia)	Liguria	Sesta a Godano	Sesta a Godano	La Spezia	2018	Absent

22	Chestnut	Liguria	Santa Maria	Calice al Cornoviglio	La Spezia	2018	Present
23	Acacia (Robinia)	Liguria	Ziona	Carro	La Spezia	2018	Absent
24	Ailanthus	Liguria	Genova	Genova	Genova	2018	Present
25	Acacia (Robinia)	Lombardia	Rengione	Missaglia	Lecco	2018	Absent
26	Linden/Lime tree (Tilia)	Lombardia	Onno	Oliveto Lario	Lecco	2018	Present
27	Mixed flower	Lombardia	Diga nel Parco Adda Nord	Vaprio d'Adda	Milano	2018	Present
28	Acacia (Robinia)	Lombardia	Oggiono	Oggiono	Lecco	2018	Absent
29	Acacia (Robinia)	Lombardia	Iseo	Iseo	Brescia	2018	Absent
30	Mixed flower	Lombardia	Casalmoro	Casalmoro	Mantova	2018	Present
31	Acacia (Robinia)	Lombardia	Missaglia	Missaglia	Lecco	2018	Absent
32	Mixed flower	Lombardia	Cascina Pirolo Casello	Pizzighettone	Cremona	2018	Absent
33	Mixed flower	Lombardia	Graglio (Val Veddasca)	Maccagno - Pino - Veddasca	Varese	2018	Present
34	Acero	Lombardia	Parco del Ticino	Vigevano	Pavia	2018	Present
35	Linden/Lime tree (Tilia)	Emilia-Roma- gna	Madonna dei Prati	Zola Predosa	Bologna	2018	Present
36	Acacia (Robinia)	Emilia-Roma- gna	Zola Vecchia	Zola Predosa	Bologna	2018	Present
37	Linden/Lime tree (Tilia)	Emilia-Roma- gna	Zola Vecchia	Zola Predosa	Bologna	2018	Present
38	Mixed flower	Emilia-Roma- gna	Osteria Nuova di Monte Co- lombo	Montescudo - Monte Colombo	Rimini	2018	Present
39	Acacia (Robinia)	Emilia-Roma- gna	Borgo Pedrosa	Montefiore Conca	Rimini	2018	Present
40	Acacia (Robinia)	Emilia-Roma- gna	Valle del Limenta	Castel di Casio	Bologna	2018	Absent
41	Mixed flower	Emilia-Roma- gna	Valle del Limenta	Castel di Casio	Bologna	2018	Present
42	Chestnut	Emilia-Roma- gna	Sant'Andrea	Caste del Rio	Bologna	2018	Present
43	Acacia (Robinia)	Emilia-Roma- gna	Sant'Andrea	Castel del Rio	Bologna	2018	Present
44	Linden/Lime tree (Tilia)	Emilia-Roma- gna	San Gabriele	Minerbio	Bologna	2018	Present
45	Sulla	Emilia-Roma- gna	Casoni di Romagna	Monterenzio	Bologna	2018	Present
46	Acacia (Robinia)	Emilia-Roma- gna	Neviano degli Arduini	Neviano degli Arduini	Parma	2018	Absent

47	Acacia (Robinia)	Emilia-Roma-	Pisignano	Cervia	Ravenna	2018	Present
48	Mixed flower	gna Emilia-Roma-	Pisignano	Cervia	Ravenna	2018	Present
49	Mixed flower	gna Emilia-Roma- gna	Fellicarolo	Fanano	Modena	2018	Present
50	Linden/Lime tree (Tilia)	Emilia-Roma- gna	Soliera	Soliera	Modena	2018	Present
51	Acacia (Robinia)	Emilia-Roma- gna	Mercato Saraceno	Mercato Saraceno	Forlì - Cesena	2018	Absent
52	Mixed flower	Emilia-Roma- gna	Bertinoro	Bertinoro	Forlì - Cesena	2018	Present
53	Mixed flower	Emilia-Roma- gna	Perticara	Novafeltria	Rimini	2018	Present
54	Linden/Lime tree (Tilia)	Emilia-Roma- gna	Forlì	Forlì	Forlì - Cesena	2018	Present
55	Mixed flower	Emilia-Roma- gna	Castiglione di Ravenna	Castiglione di Ravenna	Ravenna	2018	Present
56	Mixed flower	Emilia-Roma- gna	Forlimpopoli	Forlimpopoli	Forlì - Cesena	2018	Present
57	Mixed flower	Emilia-Roma- gna	da Cervia a Rontagnano So- gliano	Cervia - Rontagnano Sogliano	Ravenna	2018	Present
58	Acacia (Robinia)	Emilia-Roma- gna	Mercato Saraceno	Mercato Saraceno	Forlì - Cesena	2018	Absent
59	Mixed flower	Emilia-Roma- gna	Longiano	Longiano	Forlì - Cesena	2018	Present
60	Acacia (Robinia)	Emilia-Roma- gna	Sorrivoli	Roncofreddo	Forlì - Cesena	2018	Present
61	Chestnut	Emilia-Roma- gna	Bagno di Romagna	Bagno di Romagna	Forlì - Cesena	2018	Present
62	Linden/Lime tree (Tilia)	Emilia-Roma- gna	Bellaria	Rimini	Rimini	2018	Present
63	Mixed flower	Emilia-Roma- gna	San Mauro Mare	San Mauro Pascoli	Forlì - Cesena	2018	Present
64	Chestnut	Emilia-Roma- gna	Alfero	Alfero	Forlì - Cesena	2018	Present
65	Mixed flower	Emilia-Roma- gna	Gatteo	Gatteo	Forlì - Cesena	2018	Present
66	Mixed flower	Emilia-Roma- gna	Olmo	Gattatico	Reggio Emilia	2018	Present

67	Linden/Lime tree (Tilia)	Emilia-Roma-	Pianura	Gattatico	Reggio Emilia	2018	Absent
68	Honeydew (Forest)	gna Emilia-Roma-	Gattatico	Gattatico	Reggio Emilia	2018	Present
69	Acacia (Robinia)	gna Emilia-Roma-	Marazzano	Montescudo	Rimini	2018	Present
70	Acacia (Robinia)	gna Emilia-Roma- gna	Az. Masèra	Modigliana	Forlì - Cesena	2018	Present
71	Honeydew (Forest)	Emilia-Roma- gna	Celle - Pergola	Faenza	Ravenna	2018	Present
72	Mixed flower	Emilia-Roma- gna	Monte Trebbio	Modigliana	Forlì - Cesena	2018	Present
73	Linden/Lime tree (Tilia)	Emilia-Roma- gna	Parco della Contessa	Castel Bolognese	Ravenna	2018	Present
74	Mixed flower	Emilia-Roma- gna	Cà Corradini	Monterenzio	Bologna	2018	Present
75	Mixed flower	Emilia-Roma- gna	Antica Miniera di Bisano	Monterenzio	Bologna	2018	Present
76	Mixed flower	Emilia-Roma- gna	S. Maria Ripoetra	Sogliano al Rubicone	Forlì - Cesena	2018	Present
77	Acacia (Robinia)	Emilia-Roma- gna	San Polo d'Enza	San Polo d'Enza	Reggio Emilia	2018	Present
78	Bastard indago (<i>Amorpha fruti-</i> cosa)	Emilia-Roma- gna	Bodriazzo	Zibello	Parma	2018	Absent
79	Mixed flower	Emilia-Roma- gna	Castel San Pietro	Castel San Pietro	Bologna	2018	Present
80	Linden/Lime tree (Tilia)	Emilia-Roma- gna	Val di Zena	Pianoro	Bologna	2018	Present
81	Mixed flower	Emilia-Roma- gna	Gallo Bolognese	Castel San Pietro	Bologna	2018	Present
82	Mixed flower	Emilia-Roma- gna	Gallo Bolognese	Castel San Pietro	Bologna	2018	Present
83	Honeydew (Forest)	Emilia-Roma- gna	Montechiarugolo	Montechiarugolo	Parma	2018	Present
84	Mixed flower	Emilia-Roma- gna	Monticelli	Montechiarugolo	Parma	2018	Absent
85	Acacia (Robinia)	Emilia-Roma- gna	Lesignano de' Bagni	Lesignano de' Bagni	Parma	2018	Present
86	Chestnut	Emilia-Roma- gna	San Darmiano	Camugnano	Bologna	2018	Present

87	Mixed flower	Emilia-Roma-	San Darmiano	Camugnano	Bologna	2018	Present
88	Acacia (Robinia)	gna Emilia-Roma-	Montalbano	Zocca	Modena	2018	Present
89	Linden/Lime tree (<i>Tilia</i>)	gna Emilia-Roma-	Le Budrie	San Giovanni in Persiceto	Bologna	2018	Present
90	Chestnut	gna Emilia-Roma- gna	Montalbano	Zocca	Modena	2018	Present
91	Alfalfa	Emilia-Roma- gna	Balsemano	Villanova sull'Arda	Piacenza	2018	Present
92	Mixed flower	Emilia-Roma- gna	San Polo	Torrile	Parma	2018	Present
93	Honeydew (Forest)	Emilia-Roma- gna	Montecchio Emilia	Montecchio Emilia	Reggio Emilia	2018	Present
94	Ailanthus	Emilia-Roma- gna	San Giovanni	Cavriago	Reggio Emilia	2018	Present
95	Mixed flower	Emilia-Roma- gna	Bertinoro	Bertinoro	Forlì - Cesena	2018	Present
96	Bastard indago (<i>Amorpha fruti-</i> <i>cosa</i>)	Emilia-Roma- gna	Zona Fiume Po	Villanova sull'Arda	Piacenza	2018	Present
97	Honeydew (Forest)	Emilia-Roma- gna	Castelnuovo Rangone	Castelnuovo Rangone	Modena	2018	Present
98	Mixed flower	Emilia-Roma- gna	San Polo d'Enza	San Polo d'Enza	Reggio Emilia	2018	Present
99	Acacia (Robinia)	Emilia-Roma- gna	Campremoldo Sotto	Gragnano Trebbiene	Piacenza	2018	Absent
100	Mixed flower	Emilia-Roma- gna	Frassineta	Monghidoro	Bologna	2018	Present
101	Honeydew (Forest)	Emilia-Roma- gna	Montebabbio	Castellarano	Reggio Emilia	2018	Present
102	Linden/Lime tree (Tilia)	Emilia-Roma- gna	San Polo d'Enza	San Polo d'Enza	Reggio Emilia	2018	Present
103	Honeydew (Forest)	Emilia-Roma- gna	Monteveglio (Parco Abbazia)	Valsamoggia	Bologna	2018	Present
104	Coriander	Emilia-Roma- gna	Ca' de Fabbri	Minerbio	Bologna	2018	Present
105	Sulla	Emilia-Roma- gna	Montecalderaro	Castel S Pietro Terme	Bologna	2018	Present
106	Mixed flower	Emilia-Roma- gna	San Polo d'Enza	San Polo d'Enza	Reggio Emilia	2018	Present

107	Mixed flower	Emilia-Roma-	Covignano	Rimini	Rimini	2018	Present
108	Acacia (Robinia)	gna Emilia-Roma-	Saludecio	Saludecio	Rimini	2018	Present
109	Mixed flower	gna Emilia-Roma-	Lama Mocogno	Lama Mocogno	Modena	2018	Present
110	Mixed flower	gna Emilia-Roma-	Colline di Predappio	Predappio	Forlì - Cesena	2018	Present
111	Coriander	gna Emilia-Roma- gna	Bertinoro	Bertinoro	Forlì - Cesena	2018	Present
112	Mixed flower	Emilia-Roma- gna	Montiano	Montiano	Forlì - Cesena	2018	Present
113	Acacia (Robinia)	Emilia-Roma- gna	Vignale	Traversetolo	Parma	2018	Absent
114	Mixed flower	Emilia-Roma- gna	Cesata	Tredozio	Forlì - Cesena	2018	Absent
115	Alfalfa	Emilia-Roma- gna	Taglio Corelli	Alfonsine	Ravenna	2018	Present
116	Mixed flower	Emilia-Roma- gna	Montesasso	Mercato Saraceno	Forlì - Cesena	2018	Absent
117	Acacia (Robinia)	Emilia-Roma- gna	Sasso Marconi	Sasso Marconi	Bologna	2018	Absent
118	Chestnut	Emilia-Roma- gna	Castelnuovo di Vergato	Vergato	Bologna	2018	Present
119	Mixed flower	Emilia-Roma- gna	Monte Acuto Ragazza	Grizzana Morandi	Bologna	2018	Present
120	Mixed flower	Emilia-Roma- gna	Acqua Partita	Bagno di Romagna	Forlì - Cesena	2018	Present
121	Acacia (Robinia)	Emilia-Roma- gna	Selbagnone	Meldola	Forlì - Cesena	2018	Present
122	Mixed flower	Emilia-Roma- gna	Ozzano dell'Emilia	Ozzano dell'Emilia	Bologna	2018	Present
123	Honeydew (Forest)	Emilia-Roma- gna	Lungo il Torrente Tiepido	Castelnuovo Rangone	Modena	2018	Present
124	Acacia (Robinia)	Emilia-Roma- gna	Tibbio	Sarsina	Forlì - Cesena	2018	Present
125	Mixed flower	Emilia-Roma- gna	S. Andrea in Fiume	Cesena	Forlì - Cesena	2018	Present
126	Sulla	Emilia-Roma- gna	Castel del Rio	Castel del Rio	Bologna	2018	Present

127	Honeydew (Forest)	Emilia-Roma-	Castal San Pietro Terme	Castal San Pietro Terme	Bologna	2018	Present
128	Mixed flower	gna Emilia-Roma-	Boncellino	Bagnacavallo	Ravenna	2018	Present
129	Mixed flower	gna Emilia-Roma- gna	Carpineti	Carpineti	Reggio Emilia	2018	Present
130	Ailanthus	Emilia-Roma- gna	Colorno	Colorno	Parma	2018	Present
131	Acacia (Robinia)	Emilia-Roma- gna	Noceto	Noceto	Parma	2018	Present
132	Honeydew (Forest)	Emilia-Roma- gna	Colorno	Colorno	Parma	2018	Present
133	Honeydew (Forest)	Emilia-Roma- gna	Ravalle	Ferrara	Ferrara	2018	Present
134	Acacia (Robinia)	Emilia-Roma- gna	Celleri	Carpaneto	Piacenza	2018	Present
135	Rhododendron	Trentino-Alto Adige	Malga Vallina d'Amola	Giustino	Trento	2018	Absent
136	Rhododendron	Trentino-Alto Adige	Malga Bissina	Val Daone	Trento	2018	Present
137	Rhododendron	Trentino-Alto Adige	Malga Lavazzè	Rumo	Trento	2018	Present
138	Dandelion	Trentino-Alto Adige	Taio	Taio	Trento	2018	Present
139	Rhododendron	Trentino-Alto Adige	Tognola	Primiero di San Martino di Ca- strozza	Trento	2018	Present
140	Dandelion	Trentino-Alto Adige	Fraz. Piazzola	Rabbi	Trento	2018	Present
141	Raspberry	Trentino-Alto Adige	Bellamonte	Predazzo	Trento	2018	Absent
142	Rhododendron	Trentino-Alto Adige	Valfloriana	Valfloriana	Trento	2018	Absent
143	Mixed flower from Alpes	Trentino-Alto Adige	Malga Sadron	Croviana	Trento	2018	Present
144	Rhododendron	Trentino-Alto Adige	Malga Bissina	Daone	Trento	2018	Present
145	Acacia (Robinia)	Veneto	Castello	Arzignano	Vicenza	2018	Absent
146	Acacia (Robinia)	Veneto	Arzignano	Arzignano	Vicenza	2018	Absent
147	Mixed flower	Veneto	Montagnana	Montagnana	Padova	2018	Absent

148	Mixed flower	Veneto	Val d'Alpone	Bolca	Verona	2018	Present
149	Acacia (Robinia)	Veneto	Volpago del Montello	Volpago del Montello	Treviso	2018	Present
150	Acacia (Robinia)	Veneto	Chiampo	Chiampo	Vicenza	2018	Absent
151	Rhododendron	Veneto	Passo Valles	Falcade	Belluno	2018	Absent
152	Honeydew (Forest)	Veneto	Chiampo	Chiampo	Vicenza	2018	Present
153	Mixed flower	Veneto	Pianura Bassa Padovana	Correzzola	Padova	2018	Present
154	Mixed flower	Veneto	Pianura Bassa Padovana	Correzzola	Padova	2018	Present
155	Linden/Lime tree (Tilia)	Friuli-Venezia Giulia	Trebiciano	Trieste	Trieste	2018	Present
156	Bastard indago (Amorpha fruti- cosa)	Friuli-Venezia Giulia	Pianura Medio Friuli	Buttrio	Udine	2018	Present
157	Mixed flower	Friuli-Venezia Giulia	Poscolle	Cavazzo Carnico	Udine	2018	Present
158	Mixed flower	Friuli-Venezia Giulia	Piscianzi-Sottomonte	Trieste	Trieste	2018	Present
159	Morello Cherry (<i>Prunus maha-</i> <i>leb</i>)	Friuli-Venezia Giulia	Carso triestino Santa Croce	Trieste	Trieste	2018	Present
160	Erica carnea	Friuli-Venezia Giulia	Monte Corno	Trasaghis	Udine	2018	Present
161	Linden/Lime tree (Tilia)	Friuli-Venezia Giulia	San Martino	Terzo di Aquileia	Udine	2018	Present
162	Mixed flower	Friuli-Venezia Giulia	Caneva	Caneva	Pordenone	2018	Present
163	Linden/Lime tree (Tilia)	Friuli-Venezia Giulia	San Pietro al Natisone	San Pietro al Natisone	Udine	2018	Present
164	Mixed flower	Friuli-Venezia Giulia	Barcola	Trieste	Trieste	2018	Present

Table S2. PCR primer pairs used in this study.

Species	Original primer pair name	Primer sequences (5'-3'): forward and reverse	Size of the amplified DNA in bp	Amplified DNA regions	Ann. T. (°C)	References
Apis mellifera	Apis_trnL_group_F Apis_trnL_group_R	GGCAGAATAAGTGCATTG TTAATATGAATTAAGTGGGG	C 85, M 139, A 153	mtDNA COI- COII	51	Utzeri et al. (2018)
Lotmaria passim	LpCytb_F1 LpCytb_R	CGAAGTGCACATATATGCTTTAC GCCAAACACCAATAACTGGTACT	247	mtDNA cytb	59	Stevanovic et al. (2016)
Lotmaria passim	Lp_163_F Lp2R	CATTTGACTTGAATTAGCAAGC ACCACAAGAGTACGGAATGC	163	188	55	Arismendi et al. (2016) – Reverse – Forward: this study.
Lotmaria passim	Lp-gF Lp_140_R	TTGCGAAGAGCTCGCCTGAGGT GGTCGACTCGATCACGTACT	140	GAPDH	60	Arismendi et al. (2016) – Forward - Reverse: this study.
Crithidia mellifi- cae	CmGAPDH-F4 CmGAPDH-R1b	CGGCGTGGACTACGTGATT ACGACGTGGTGCTTGGAC	177	GAPDH	60	Bartolomè et al. (2018)
Crithidia bombi	CbTOP-F1 CbTOP-R1	CGAGGTGCGGCTCAACA GATGCAGCCATTCGGGCT	133	TOPII	62	Bartolomè et al. (2018)