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**1 DNA barcoding for identification of sand fly species (Diptera: Psychodidae) from  
2 leishmaniasis-endemic areas of Peru**

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29

30      **Abstract**

31      Phlebotomine sand flies are the only proven vectors of leishmaniases, a group of human and  
32      animal diseases. Accurate knowledge of sand fly species identification is essential in  
33      understanding the epidemiology of leishmaniasis and vector control in endemic areas. Classical  
34      identification of sand fly species based on morphological characteristics often remains difficult  
35      and requires taxonomic expertise. Here, we generated DNA barcodes of the cytochrome c  
36      oxidase subunit 1 (COI) gene using 159 adult specimens morphologically identified to be 19  
37      species of sand flies, belonging to 6 subgenera/species groups circulating in Peru, including the  
38      vector species. Neighbor-joining (NJ) analysis based on Kimura 2-Parameter genetic distances  
39      formed non-overlapping clusters for all species. The levels of intraspecific genetic divergence  
40      ranged from 0 to 5.96%, whereas interspecific genetic divergence among different species  
41      ranged from 8.39 to 19.08%. The generated COI barcodes could discriminate between all the  
42      sand fly taxa. Besides its success in separating known species, we found that DNA barcoding is  
43      useful in revealing population differentiation and cryptic diversity, and thus promises to be a  
44      valuable tool for epidemiological studies of leishmaniasis.

45

46      *Keywords:* DNA barcode, sand flies, leishmaniasis, vector, Peru

47

48     **1. Introduction**

49       Phlebotomine sand flies (Diptera: Psychodidae, Phlebotominae) are of significant public  
50       health importance in many parts of the world as vectors of the causative agents of leishmaniasis,  
51       bartonellosis and sand fly fever (Maroli et al., 2013). Leishmaniasis remains one of the world's  
52       most neglected diseases; endemic in 98 countries, putting 350 million people at risk, with some  
53       two million new cases every year (WHO, 2010; Alvar et al., 2012). The disease is caused by  
54       several species of cell-infecting parasites of the genus *Leishmania*, and presents three major  
55       distinct clinical manifestations: cutaneous, mucocutaneous and visceral forms (Bates, 2007).  
56       Sand flies are the only proven vectors of leishmaniases and approximately 800 species have been  
57       recorded in five major genera; *Phlebotomus* and *Sergentomyia* in the Old World, and *Lutzomyia*,  
58       *Brumptomyia* and *Warileya* in the New World (Munstermann, 2004). However, only species  
59       belonging to the genera *Phlebotomus* and *Lutzomyia* are the putative vectors of *Leishmania*  
60       (Munstermann, 2004; Bates, 2007; Kato et al., 2010). Therefore, correct sand fly species  
61       identification is very important to design strategies for surveillance and control of leishmaniasis  
62       in given endemic areas.

63       In Peru, three sand fly species, *Lutzomyia* (*Lu.*) *peruensis*, *Lu. ayacuchensis* and *Lu.*  
64       *verrucarum* have been incriminated as the vectors of *Leishmania* (*Viannia*) *peruviana*, the  
65       aetiological agent of Andean-type cutaneous leishmaniasis in the endemic areas (Davies et al.,  
66       1993; Perez et al., 1991, 1994, 2007; Caceres et al., 2004; Kato et al., 2008, 2011). Although  
67       these sand fly species role in disease transmission has been well studied, detailed molecular  
68       taxonomic knowledge of each species is far from complete. Presently, sand flies are typically  
69       identified based on morphological features; mainly by internal structures such as the cibarium,  
70       pharynx, spermatheca of females and terminal genitalia of males (Young and Duncan, 1994).

71 However, this morphological classification is laborious, time-consuming and often complicated  
72 by phenotypic plasticity and cryptic species complexes, as well as demanding considerable skill  
73 and taxonomic expertise. To overcome these technical limitations, in the last three decades  
74 molecular approaches have been increasingly employed to explore the taxonomy, population  
75 structure and phylogeny in insect vectors including sand flies (Mukhopadhyay et al., 2000; Uribe  
76 Soto et al., 2001; Depaquit et al., 2002; Beati et al., 2004; Khalid et al., 2010; Fujita et al., 2012;  
77 Yamamoto et al., 2013; Nzelu et al., 2014; Gomez et al., 2014; Kato et al., 2015).

78 Recently, DNA barcoding has been widely shown to be an effective molecular tool for  
79 species identification of many insects (Hebert et al., 2003a, b). Among the mitochondrial genes,  
80 cytochrome c oxidase I (COI) is reported to be the most conserved gene and hence has distinct  
81 advantages for taxonomic studies (Knowlton and Weigt, 1998). Studies have demonstrated the  
82 use of DNA barcoding as a valid molecular tool for identifying dipterans; mosquitoes (Cywinska  
83 et al., 2006), black flies (Pramual et al., 2011) and sand flies (Azpurua et al., 2010; Kumar et al.,  
84 2012; Contreras Gutiérrez et al., 2014; Nzelu et al., 2014), and revealing sand fly cryptic species  
85 (Cohnstaedt et al., 2011; Scarpassa et al., 2012; Kumar et al., 2012). Construction of DNA  
86 barcodes for each species of sand fly would provide an efficient tool for their identification. The  
87 aim of the present study was to explore the utility of the DNA barcode approach for  
88 identification of sand fly species in Peru, where leishmaniasis is endemic, by analyzing  
89 morphologically identified adult specimens.

90

91      **2. Materials and methods**

92      *2.1 Sand fly collection*

93      Sand flies were caught using CDC light traps and Shannon traps peridomiciliary,  
94      intradomiciliary and extradomiciliary at 12 Departments; Piura, Lambayeque, Cajamarca, La  
95      Libertad, San Martin, Huánuco, Lima, Huancavelica, Ayacucho, Apurímac, Cusco and Puno in  
96      Peru between 2001 and 2013 (Fig. 1, Table 1). All the flies were captured throughout the night  
97      between 18:00 p.m. and 6:00 a.m. with CDC light traps and between 18:00 p.m. and 22:00 p.m.  
98      with Shannon traps. The sand flies were morphologically identified mainly based on  
99      measurements of wing veins, the ratio of palpus length to length of antenna and the color of the  
100     thorax (Young and Duncan, 1994), and then fixed in absolute ethanol and stored at room  
101     temperature.

102

103      *2.2 DNA extraction*

104      Individual ethanol-fixed sand flies were lysed in 50 µl of DNA extraction buffer [150 mM  
105     NaCl, 10 mM Tris-HCl (pH8.0), 10 mM EDTA and 0.1% sodium dodecyl sulfate (SDS)] with 100  
106     µg/ml proteinase K. The samples were incubated at 37 °C overnight. Afterwards, 25 µl of  
107     distilled water was added, heated at 95 °C for 5 min, and then, 0.5 µl portions were directly used  
108     as templates for PCR amplification.

109

110      *2.3 Polymerase chain reaction and sequence analysis*

111      The mitochondrial COI was amplified using the primers LCO 1490 (5'-  
112     GGTCAACAAATCATAAAGATATTGG3') and HCO 2198

113 (5'TAAACTTCAGGGTGACCAAAAAATCA-3') (Folmer et al., 1994). The reaction was carried  
114 out in a volume of 15 µl using a pair of primers (0.4 µM each), Ampdirect Plus (Shimadzu  
115 Biotech, Tsukuba, Japan), and BIOTaq HS DNA polymerase (Bioline, London, UK). After an  
116 initial denaturation at 95 °C for 5 min, amplification was performed with 35 cycles consisting of  
117 denaturation at 95 °C for 1 min, annealing at 49 °C for 30 sec, extension at 72 °C for 2 min,  
118 followed by a final extension at 72 °C for 10 min. PCR products were purified using a FastGene  
119 Gel/PCR Extraction kit (NIPPON Genetics) and sequenced with a forward primer by the dideoxy  
120 chain termination method using a BigDye Terminator v3.1 Cycle Sequencing Kit (Applied  
121 Biosystems, Foster City, CA). To confirm the authenticity of the COI sequences obtained, some  
122 individuals from each sand fly species (in total 40 specimens) were subjected to PCR using a high  
123 fidelity DNA polymerase (KOD-Plus-ver.2; TOYOBO, Tokyo, Japan), and the sequences were  
124 determined as described above.

125

#### 126 2.4 Data analysis

127 The COI sequences were aligned by CLUSTAL W software (Thompson et al., 1997)  
128 incorporated into MEGA (Molecular Evolutionary Genetics Analysis) version 5.2 (Tamura et al.,  
129 2011). The nucleotide compositions and sequence divergences within and between species were  
130 calculated using the distance model Kimura 2-Parameter (K2P) (Kimura, 1980). This model  
131 provides the best estimate of divergence when genetic distances are low, as in recently diverged  
132 taxa (Nei and Kumar, 2000). The ‘best match’ method in the SpeciesIdentifier module of  
133 TaxonDNA software v1.7.8 (Meier et al., 2006) was used to test the frequency of successful  
134 identification. A Neighbor-joining (NJ) tree of K2P distances was created to provide a graphic  
135 representation of the clustering pattern among different species (Saitou and Nei, 1987). Branch

136 support for NJ was calculated using the bootstrapping method with 1,000 replicates in MEGA  
137 5.2 (Tamura et al., 2011). All sequences and other specimen information are available in the  
138 dataset project ‘DBSFP’ (Process IDs: DBSFP001-15 to DBSFP159-15) on the Barcode of Life  
139 Data System (BOLD) at <http://www.boldsystems.org>. Sequence data are also available in the  
140 DDBJ/EMBL/GenBank databases (<http://www.ncbi.nlm.nih.gov/genbank/>) under the accession  
141 numbers AB984357–AB984520 (Table 1).

142

143    **3. Results**

144    *3.1 Sequence analysis*

145    A total of 159 COI sequences from 19 morphologically identified species of phlebotomine  
146    sand flies belonging to 6 subgenera/species groups were obtained. Each species was analyzed  
147    using 1 to 17 specimens (Table 1, 2). The COI sequences obtained varied from 650 to 672 bp by  
148    direct sequencing. No deletions, insertions or stop codons were found indicating that all  
149    sequences constituted functional mitochondrial products (Funk and Omland, 2003). As these 159  
150    COI sequences contained no indels, alignments were straight forward. The COI sequences from  
151    19 species of sand flies had high adenosine and thymine (AT)-rich content bias (65.9%), similar  
152    to that observed for other insects (Simon et al., 1994), with an average nucleotide composition of  
153    A=28.2%, T=37.7% , C=18.0%, and G=16.2%.

154    Intraspecific genetic distances based on Kimura 2-Parameter values for the sand fly species  
155    ranged from 0 to 5.96% (Table 2). Maximum intraspecific genetic diversity was observed in 7 of  
156    the 19 species, especially within the three vector species of *Leishmania (Viannia) peruviana*;  
157    *Lutzomyia (Lu.) peruensis* (5.96%), *Lu. verrucarum* (3.48%) and *Lu. ayacuchensis* (2.76%)  
158    (Table 2). The high level of intraspecific divergence was related to geographical distribution in  
159    some species. The mean intraspecific K2P distance ranged from 0.05% (*Lu. caballeroi*) to 3.36%  
160    (*Lu. peruensis*) (Table 2). Interspecific K2P genetic distances for the sand fly species included in  
161    the study ranged from 8.39 to 19.08%. The overall nucleotide diversity recorded among the 19  
162    species under the Subfamily Phlebotominae included in the study was found to be 13.9%. Low  
163    levels of minimum interspecific divergence typically occurred in closely related species, such as  
164    *Lu. tejadai* and *Lu. quillabamba* (8.39%), *Lu. bicornuta* and *Lu. battistinii* (8.88%), and *Lu.*  
165    *castanea* and *Lu. caballeroi* (10.15%). Collectively, the distances between species belonging to

166 the subgenus *Lutzomyia* (*Lu. bicornuta* and *Lu. battistinii*), and the *Verrucarum* group (*Lu.*  
167 *verrucarum* and *Lu. nuneztovari*) were 8.88 and 12.27%, respectively, whereas the distances  
168 between species belonging to the subgenus *Pifanomyia* (*Lu. maranonensis*, *Lu. robusta* and *Lu.*  
169 *nevesi*) ranged from 11.46 to 15.43%. On the other hand, the diversity within the subgenus  
170 *Helcocyrtomyia* (*Lu. munaypata*, *Lu. quillabamba*, *Lu. pescei*, *Lu. noguchii*, *Lu. castanea*, *Lu.*  
171 *caballeroi*, *Lu. tejadai*, *Lu. ayacuchensis* and *Lu. peruvensis*) ranged from 8.39 to 16.5%. Using  
172 SpeciesIdentifier, the overall percentage of correct species identification was 100% as  
173 recognized by traditional taxonomy based on morphological features. Generally, the COI  
174 divergence within species did not exceed the genetic distance between species. As a result, there  
175 was a distinction between intraspecific and interspecific genetic divergences, which is critical for  
176 DNA barcoding success in taxa discrimination (Fig. 2).

177

178 3.2 *Neighbor-joining tree*

179 The Neighbor-joining tree showed shallow intraspecific and deep interspecific divergences  
180 (Fig. 3). Specimens of the same species were grouped closely together, regardless of the  
181 collection site. However, in exceptional cases, obvious geographic differences in sequences  
182 within the same species were found in three species, namely, *Lu. peruvensis*, *Lu. verrucarum* and  
183 *Lu. maranonensis*. Each of these species was monophyletic, but the individuals were divided into  
184 two clades corresponding to geographic origin (Fig. 3), namely, *Lu. peruvensis*: Cusco (southeast)  
185 clade, and Lima (central) and La Libertad (northwest) clade; *Lu. verrucarum*: Lima (central)  
186 clade, and Piura (northwest) clade; *Lu. maranonensis*: Querocotillo (Cajamarca-northwest) clade,  
187 and San José de Lourdes (Cajamarca-northwest) clade. Additionally, the *Lu. robusta* cluster was

188 monophyletic, but the individuals were divided into two clades: Cajamarca I (northwest) clade,  
189 and Cajamarca II (northwest) clade (Fig. 3).

190

191     **4. Discussion**

192     Adequate knowledge of the ecological and medical importance of sand flies relies on correct  
193     identification of these minute and fragile insects. The gold standard for any robust taxonomic  
194     system is its ability to deliver reliable and accurate species identifications (Hebert et al., 2003a).  
195     In the present study, we were able to recover and align the targeted COI fragment from all sand  
196     fly species examined. The strongly biased AT-richness noted within the DNA barcode sequences  
197     was consistent with results from other insects including different sand fly genera (Cywinska et  
198     al., 2006; Pramual et al., 2011; Kumar et al., 2012; Contreras Gutiérrez et al., 2014). The range  
199     of intraspecific variation was between 0 and 5.96%, while the interspecific variability for species  
200     differentiation ranged between 8.39 and 19.08%. The range of intraspecific genetic divergence in  
201     this study, 0 to 5.96%, was very similar to values reported for New World sand flies in Colombia  
202     (0 to 6%; Contreras Gutiérrez et al., 2014), whereas the 5.96% maximum intraspecific  
203     divergence was higher than that earlier reported for Old World sand flies in India (2%; Kumar et  
204     al., 2012). High intraspecific sequence divergence was found among the three vector species of  
205     *Leishmania (Viannia) peruviana* in Peru (Davies et al., 1993; Perez et al., 1991, 1994, 2007;  
206     Caceres et al., 2004; Kato et al., 2008, 2011); *Lu. (Helcocyrtomyia) peruensis* (5.96% maximum  
207     intraspecific divergence), *Lu. (Verrucarum) verrucarum* (3.48%), and *Lu. (Helcocyrtomyia)*  
208     *ayacuchensis* (2.76%) (Table 2). Overall, there were no overlaps between the intra- and inter-  
209     specific divergence levels, despite the high values of intraspecific divergences observed in some  
210     species, supporting the utility of mtDNA COI in discriminating species (Hebert et al., 2003b).

211     Inclusion of DNA sequence data in phylogenetic analysis of individuals allows for quick  
212     determination of monophyletic groups and recognition of hidden or potential species. In addition  
213     to delineation of species, DNA barcoding can reveal cryptic diversity (Hebert et al., 2004;

214 Pramual et al., 2011; Kumar et al., 2012). The Neighbor-joining (NJ) tree analysis distinctively  
215 clustered the COI sequences obtained from individual *Lutzomyia* sand flies as previously  
216 determined by morphological features; specimens of single species formed barcode clusters with  
217 tight cohesion that were clearly distinct from those of allied species. However, in this study four  
218 taxa, *Lu. peruvensis*, *Lu. verrucarum*, *Lu. maranonensis* and *Lu. robusta* were found to be  
219 complexes of at least 2 genetically distinct groups. The NJ tree revealed two deep-divergence  
220 reciprocally monophyletic clades in *Lu. peruvensis*, the main vector of *L. (V.) peruviana* (Perez et  
221 al., 1991, 2007; Kato et al., 2008, 2011) that clustered by geographic collection area (Fig. 3), and  
222 indicated that the *Lu. peruvensis* group from Cusco (southeast) was genetically distinct from those  
223 from Lima (central) and La Libertad (northwest). A similar scenario was previously observed in  
224 *Lu. peruvensis* based on the analysis of *cyt b* gene, among specimens from two regions, Ancash  
225 and Lima (central), and La Libertad (northwest) (Yamamoto et al., 2013). Additionally, we also  
226 observed a high genetic distance between the Lima (central) and Piura (northwest) groups of *Lu.*  
227 *verrucarum*. Therefore, these observed genetic distances among *Lu. peruvensis* and *Lu.*  
228 *verrucarum* groups indicated that the groups in each taxa were highly isolated, with little or no  
229 genetic exchange between the groups.

230 Phlebotomine sand flies are known to have limited sustained flight and tend to remain  
231 associated with their emergence sites, typically not dispersing more than a half kilometer  
232 (Alexander, 1987; Killick-Kendrick, 1990; Morrison et al., 1993). Such putatively low flight  
233 capacity and dispersal range of sand flies coupled with the heterogeneous terrain of the Andes  
234 may cause the isolation of these populations throughout their distributions, thereby producing  
235 genetically distinct but morphologically similar individuals. Consequently, genetic  
236 differentiation observed within some species, may be ascribable to local adaptation or genetic

237 drift. Moreover, the observation of genetic differentiation among the vector species (*Lu.*  
238 *peruensis*, *Lu. verrucarum*, and *Lu. ayacuchensis*) may have serious implications for  
239 epidemiological studies and vector control strategies. Indeed it has been shown that insecticide  
240 susceptibility and vector capacity can vary within evolutionary lineages, even within the same  
241 species (Lanzaro et al., 1993; O' Loughlin et al., 2008; Hassan et al., 2012). Collectively, the fact  
242 that some species showed higher conspecific divergence does not compromise the use of COI  
243 sequences for their identification; rather, it allows delineation of the regional lineages that  
244 constitute them (Hebert et al., 2003b). Nonetheless, further investigations with more specimens  
245 from various localities will be required to provide additional information on the levels of  
246 divergence within these vector species and other prevalent sand fly species in Peru. Such studies  
247 may also contribute to more effective leishmaniasis control in endemic areas.

248 In conclusion, this study provided the COI barcodes for several Peruvian sand flies and  
249 showed their effectiveness in discriminating species recognized through prior conventional  
250 taxonomic work. Aside from its success in discriminating known species, DNA barcoding was  
251 found to be useful in revealing population differentiation. A better understanding of the  
252 implication of the genetic differentiation among the vector species will require further molecular  
253 (genetic information from multiple loci), ecological and morphological studies.

254

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261

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- 409

410    **Figure Legends**

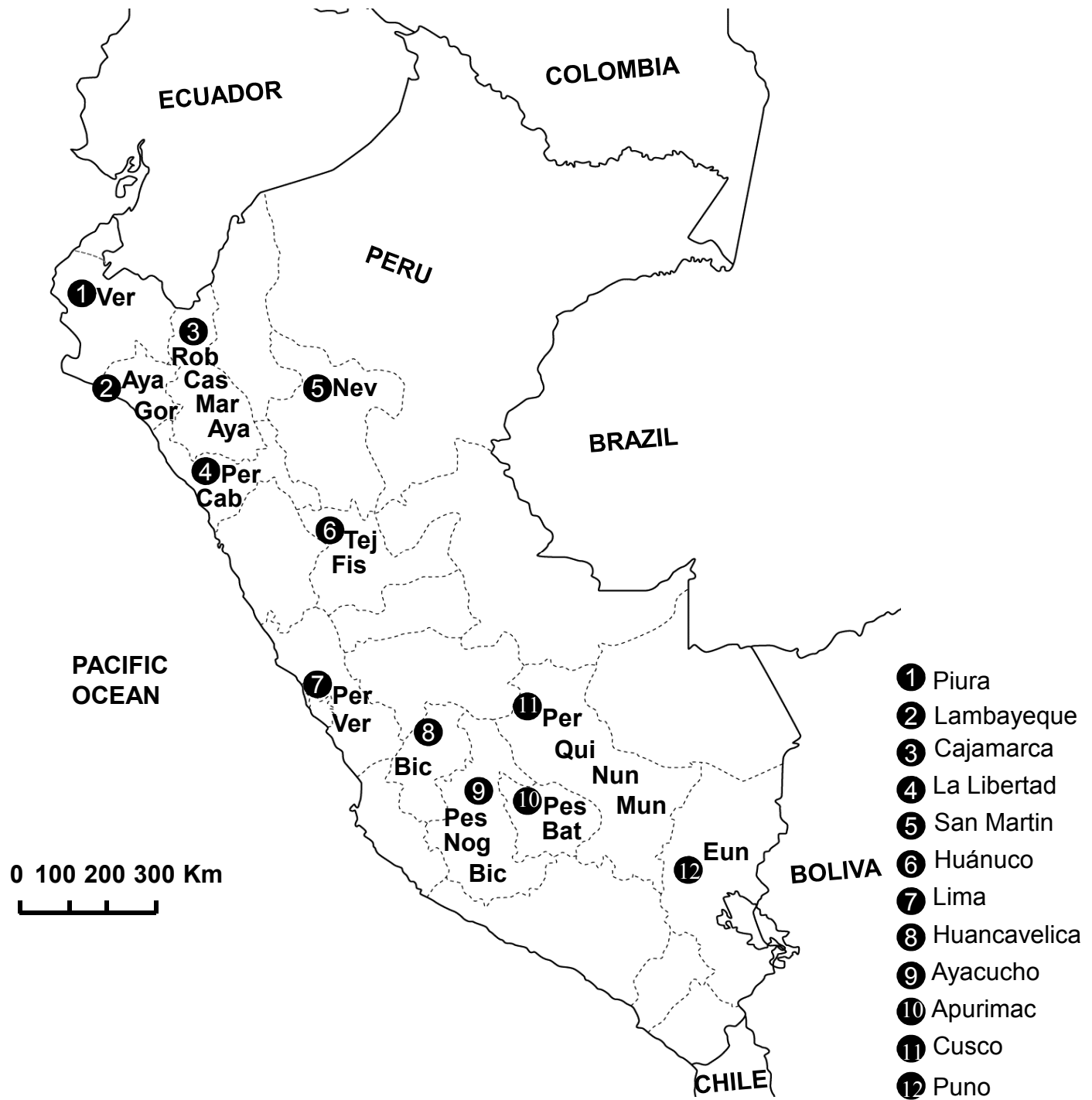
411    **Fig. 1.** Map of Peru showing the geographic locations where sand flies were collected along with  
412    the respective species analyzed in this study. Bat: *Lu. battistinii*; Bic: *Lu. bicornuta*; Ver: *Lu.*  
413    *verrucarum*; Nun: *Lu. nuneztovari*; Mar: *Lu. maranonensis*; Nev: *Lu. nevesi*; Rob: *Lu. robusta*;  
414    Tej: *Lu. tejadai*; Qui: *Lu. quillabamba*; Cab: *Lu. caballeroi*; Cas: *Lu. castanea*; Pes: *Lu. pescei*;  
415    Aya: *Lu. ayacuchensis*; Per: *Lu. peruensis*; Mun: *Lu. munaypata*; Nog: *Lu. noguchii*; Fis: *Lu.*  
416    *fischeri*; Gor: *Lu. gorbitzi*; Eun: *Warileya euniceae*.

417

418    **Fig. 2.** Distribution of intraspecific (grey) and interspecific (black) genetic distances for the  
419    cytochrome c oxidase subunit I (COI) sequences of sand fly species from Peru.

420

421    **Fig. 3.** Neighbor-joining tree based on Kimura 2-parameter genetic distances of mitochondrial  
422    COI sequences of *Lutzomyia* sand fly species from Peru. The bar scale represents 0.01%  
423    divergences. Bootstrap values are shown above or below branches. Specimen IDs (in  
424    parentheses).



**Fig. 1**

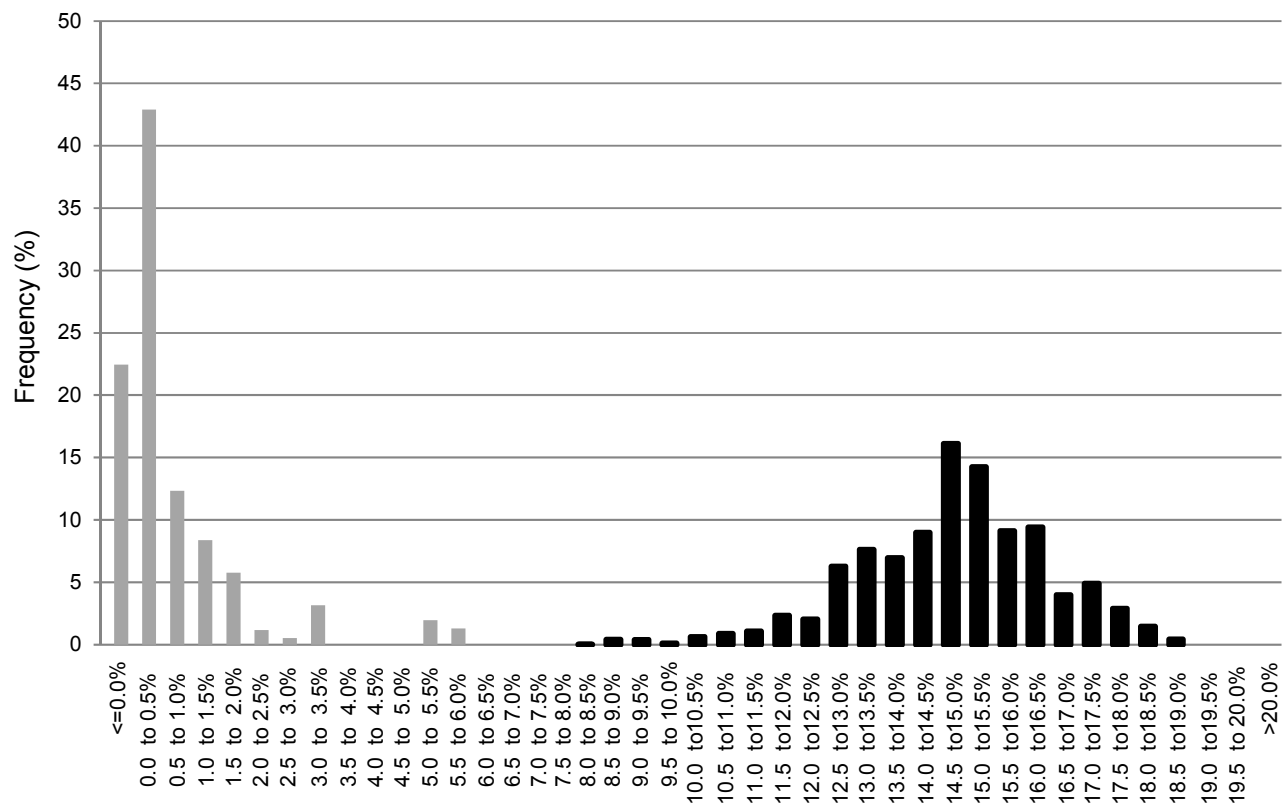


Fig. 2

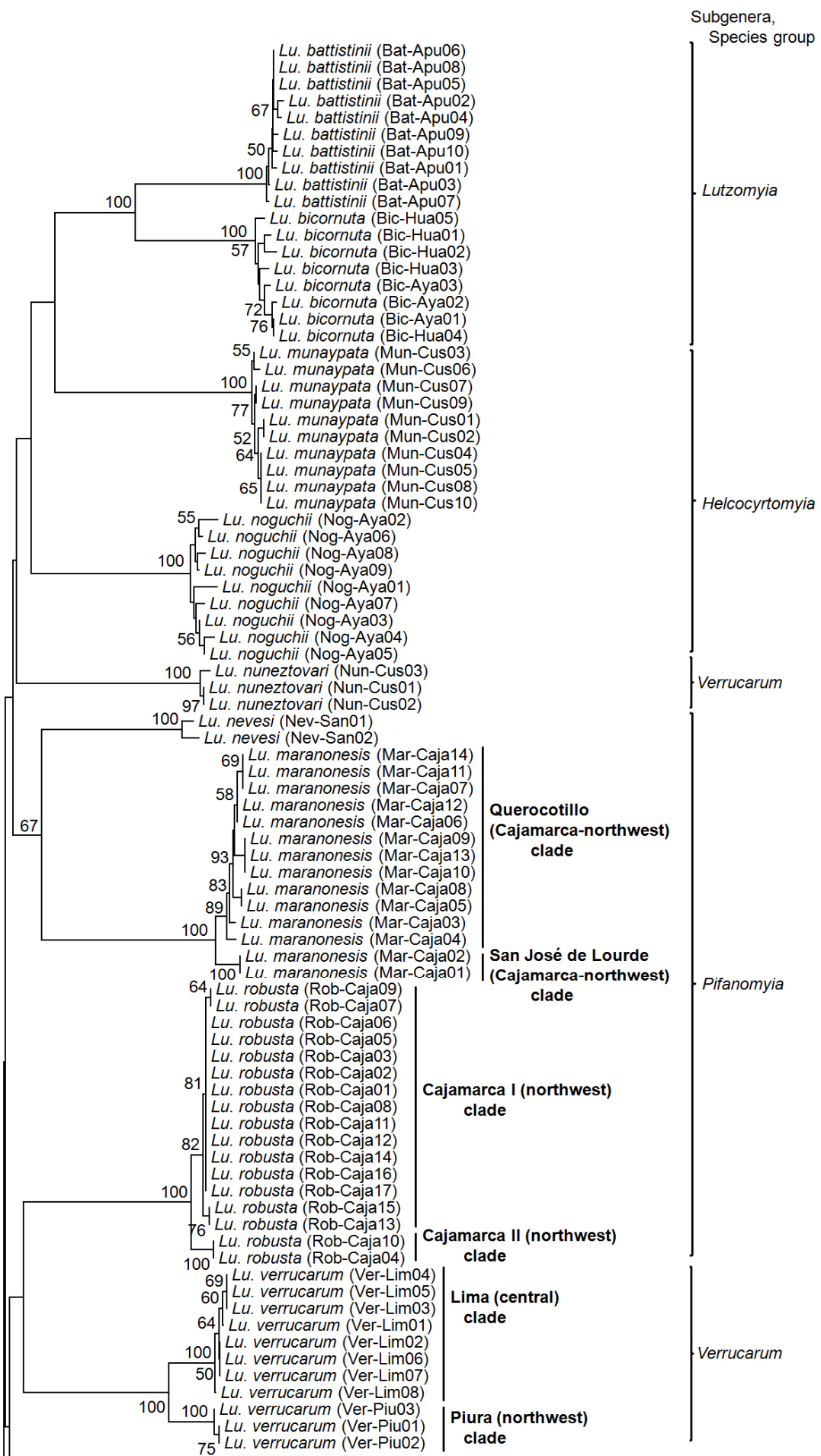


Fig. 3

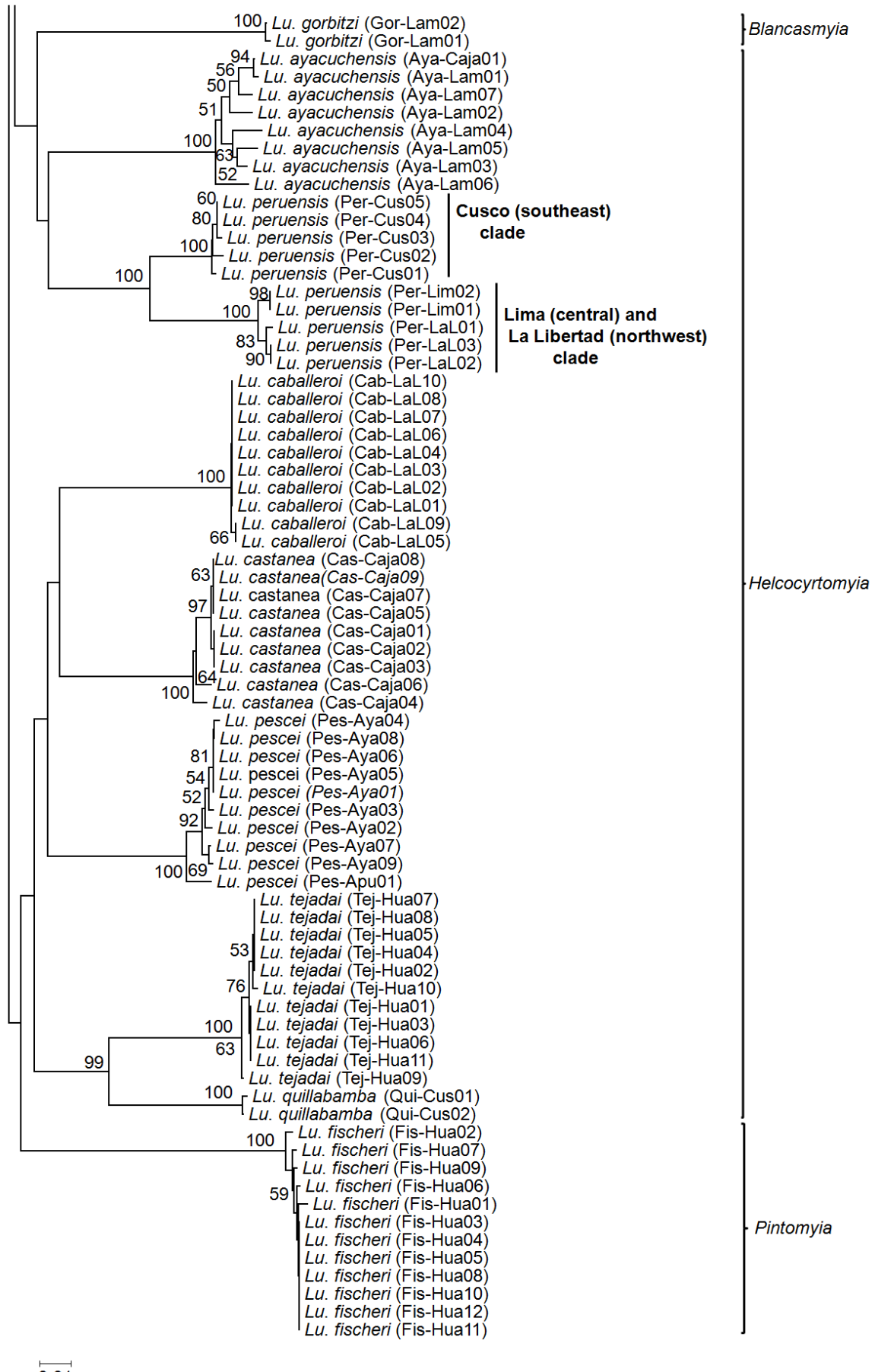


Fig. 3 (Continued)

**Table 1**

Details of sand fly specimens used for DNA barcoding and analysis in the study along with the BOLD Process IDs and GenBank accession numbers.

Sand fly species	No. of samples	Location		Trapping method	Year of capture	BOLD Process IDs	GenBank accession nos.
		District	Department				
<i>Lu. battistinii</i>	10	Abancay	Apurimac	CDC	2012	DBSFP001-15- DBSFP010-15	AB984357-AB984366
<i>Lu. bicornuta</i>	3	Luricocha	Ayacucho	CDC	2009	DBSFP011-15- DBSFP013-15	AB984367-AB984369
<i>Lu. bicornuta</i>	5	San Marcos de Rocchac	Huancavelica	CDC	2011	DBSFP014-15- DBSFP018-15	AB984370-AB984374
<i>Lu. munaypata</i>	10	Maranura	Cusco	CDC	2007	DBSFP019-15- DBSFP028-15	AB984375-AB984384
<i>Lu. quillabamba</i>	2	Calca	Cusco	CDC	2011	DBSFP029-15- DBSFP030-15	AB984385-AB984386
<i>Lu. tejadai</i>	11	Huánuco	Huánuco	CDC	2007, 2012	DBSFP031-15- DBSFP041-15	AB984387-AB984397
<i>Lu. pescei</i>	9	Luricocha	Ayacucho	CDC	2009	DBSFP042-15- DBSFP050-15	AB984398-AB984406
<i>Lu. pescei</i>	1	Chacoche	Apurimac	CDC	2012	DBSFP051-15	AB984407
<i>Lu. caballeroi</i>	10	La Cuesta	La Libertad	CDC	2007	DBSFP052-15- DBSFP061-15	AB984408-AB984417
<i>Lu. castanea</i>	5	San José de Lourdes	Cajamarca	ShT	2001	DBSFP062-15- DBSFP066-15	AB984418-AB984422
<i>Lu. castanea</i>	4	San Ignacio	Cajamarca	CDC	2007	DBSFP067-15- DBSFP070-15	AB984423-AB984426
<i>Lu. fischeri</i>	12	Ambo	Huánuco	CDC	2013	DBSFP071-15- DBSFP082-15	AB984428-AB984439
<i>Lu. gorbitzi</i>	2	Motupe	Lambayeque	CDC	2013	DBSFP083-15- DBSFP084-15	AB984440-AB984441
<i>Lu. ayacuchensis</i>	7	Motupe	Lambayeque	CDC	2013	DBSFP085-15- DBSFP091-15	AB984442-AB984448
<i>Lu. ayacuchensis</i>	1	San José de Lourdes	Cajamarca	CDC	2013	DBSFP092-15	AB984449
<i>Lu. peruensis</i>	2	Ambar	Lima	CDC	2007	DBSFP093-15- DBSFP094-15	AB984450-AB984451
<i>Lu. peruensis</i>	3	La Cuesta	La Libertad	CDC	2007	DBSFP095-15- DBSFP097-15	AB984452-AB984454
<i>Lu. peruensis</i>	1	Cusipata	Cusco	CDC	2011	DBSFP098-15	AB984455
<i>Lu. peruensis</i>	1	Urubamba	Cusco	CDC	2012	DBSFP099-15	AB984456
<i>Lu. peruensis</i>	3	Ollantaytambo	Cusco	CDC	2012	DBSFP100-15- DBSFP102-15	AB984457-AB984459
<i>Lu. verrucarum</i>	3	Huancabamba	Piura	CDC	2004	DBSFP103-15- DBSFP105-15	AB984460-AB984462
<i>Lu. verrucarum</i>	2	San Antonio de Chaclla	Lima	CDC	2013	DBSFP106-15- DBSFP107-15	AB984463-AB984464
<i>Lu. verrucarum</i>	6	Santa Eulalia	Lima	CDC	2013	DBSFP108-15- DBSFP113-15	AB984465-AB984470
<i>Lu. nuneztovari</i>	3	Maranura	Cusco	CDC	2007	DBSFP114-15- DBSFP116-15	AB984471-AB984473
<i>Lu. noguchii</i>	9	Pullo	Ayacucho	CDC	2007	DBSFP117-15- DBSFP125-15	AB984474-AB984482
<i>Lu. nevesi</i>	2	Tarapoto	San Martin	CDC	2007	DBSFP126-15- DBSFP127-15	AB984483-AB984484
<i>Lu. maranonesis</i>	2	San José de Lourdes	Cajamarca	CDC	2001	DBSFP128-15- DBSFP129-15	AB984485-AB984486
<i>Lu. maranonesis</i>	12	Querocotillo	Cajamarca	CDC	2006	DBSFP130-15- DBSFP141-15	AB984487-AB984498
<i>Lu. robusta</i>	2	San José de Lourdes	Cajamarca	ShT	2001	DBSFP142-15- DBSFP143-15	AB984501-AB984502
<i>Lu. robusta</i>	10	San José de Lourdes	Cajamarca	CDC	2001	DBSFP144-15- DBSFP153-15	AB984503-AB984512
<i>Lu. robusta</i>	5	San Juan de Cutervo	Cajamarca	CDC	2007	DBSFP154-15- DBSFP158-15	AB984513-AB984517
<i>Wa. euniceae</i>	1	Alto Inambari	Puno	ShT	2013	DBSFP159-15	AB984520

*Lu:* *Lutzomyia*; *Wa:* *Warileya*; CDC: CDC light trap; ShT: Shannon trap; BOLD: Barcode of Life Data System.

**Table 2**

Intraspecific Kimura 2-Parameter (K2P) pairwise distances of sand fly species from Peru.

Subgenera/Species group	Sand fly species	No. of Specimens	Minimum (%)	Mean (%)	Maximum (%)
<i>Lutzomyia</i>	<i>Lu. battistinii</i>	10	0	0.27	0.46
	<i>Lu. bicornuta</i>	8	0	0.76	1.25
<i>Verrucarum</i>	<i>Lu. verrucarum</i>	11	0	1.59	3.48
	<i>Lu. nuneztovari</i>	3	0	0.3	0.46
<i>Pifanomyia</i>	<i>Lu. maranonesis</i>	14	0	0.81	1.87
	<i>Lu. nevesi</i>	2		0.93	0.93
	<i>Lu. robusta</i>	17	0	0.37	1.55
<i>Helcocyrtomyia</i>	<i>Lu. tejadai</i>	11	0	0.16	0.46
	<i>Lu. quillabamba</i>	2		0.15	0.15
	<i>Lu. caballeroi</i>	10	0	0.05	0.15
	<i>Lu. castanea</i>	9	0	0.5	1.24
	<i>Lu. pescei</i>	10	0	0.58	1.69
	<i>Lu. ayacuchensis</i>	8	0.2	1.85	2.76
	<i>Lu. peruensis</i>	10	0	3.36	5.96
	<i>Lu. munaypata</i>	10	0	0.28	0.46
	<i>Lu. noguchii</i>	9	0.2	0.92	1.72
<i>Pintomyia</i>	<i>Lu. fischeri</i>	12	0	0.2	0.75
<i>Blancasmyia</i>	<i>Lu. gorbitzi</i>	2		0.15	0.15
—	<i>Wa. euniceae</i>	1		N/A	

N/A: not applicable.