Leishmania donovani Transmission Cycle Associated with Human Infection, Phlebotomus alexandri Sand Flies, and Hare Blood Meals, Israel¹

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Cutaneous leishmaniasis caused by Leishmania major or L. tropica and visceral leishmaniasis caused by L. infantum have been reported in Israel. We collected Phlebotomus spp. sand flies in the Negev desert of southern Israel to identify circulating Leishmania spp. Of 22,636 trapped sand flies, 80% were P. alexandri. We sequenced Leishmania-specific internal transcribed spacer 1 fragments and K26 genes. Of 5,019 Phlebotomus female sand flies, 2.5% were Leishmania DNA-positive; 92% of infections were L. donovani. Phylogenetic analyses showed separate clustering of L. donovani and L. infantum. P. alexandri flies positive for L. donovani harbored blood meals from European hares. Leishmania DNA isolated from a patient with cutaneous leishmaniasis who lived in the survey area was identical to L. donovani from P. alexandri flies. We report circulation of L. donovani, a cause of visceral leishmaniasis, in southern Israel. Prompt diagnosis and Leishmania spp. identification are critical to prevent leishmaniasis progression.

Zoonotic leishmaniasis is endemic to Israel. *Leishmania tropica, L. major,* and *L. infantum* infect humans in different areas of Israel and circulate through distinct zoonotic transmission cycles (1). Cutaneous

Author affiliations: Ministry of Health Central Laboratories for Public Health, Jerusalem, Israel (L. Studentsky, L. Orshan, I. Ben Avi, D. Diaz, S. Elbaz, M. Davidovich-Cohen); The Hebrew University of Jerusalem, Rehovot, Israel (L. Studentsky, G. Baneth); Ministry of Health National Laboratory for Public Health, Tel Aviv, Israel (F. Akad); Soroka University Medical Center, Beer-Sheba, Israel (O. Sagi); Israeli Ministry of Environmental Protection, Jerusalem (G. Zagron); Maccabi Healthcare Services, Tel Aviv (L. Valinsky) leishmaniasis (CL) is caused by *L. major*, which is transmitted by *Phlebotomus papatasi* sand flies, and *L. tropica*, which is transmitted by *P. sergenti* and *P. arabicus* sand flies. Canine leishmaniasis and human visceral leishmaniasis (VL) are caused by *L. infantum* in Israel, and the putative vectors are *P. perfiliewi*, *P. syriacus*, and *P. tobbi* sand flies (1–9). Reservoirs for *L. major* are sand rats (*Psammomys obesus*), gerbils (*Gerbillus dasyurus*), jirds (*Meriones crassus* and *M. tristrami*), and possibly also voles (10–13), whereas rock hyraxes (*Procavia capensis*) are considered the animal reservoir for *L. tropica* in Israel (14). Domestic dogs (*Canis lupus familiaris*), jackals (*C. aureus*), foxes (*Vulpes vulpes*), and wolves (*C. lupus*) are recognized reservoir hosts for *L. infantum* (15).

A substantial increase in CL incidence has been recorded since 2002, and endemic transmission has occurred in areas of Israel where it was previously unknown (6,16,17). Although not life-threatening, CL is a considerable public health problem in Israel; CL is diagnosed in hundreds of new patients annually. During 2001-2018, CL incidence rates increased 7-fold, from 0.4 to 2.9/100,000 population; a peak was observed in 2012, when the mean annual incidence increased to 4.4/100,000 population (18,19). Our study combines results from sand fly surveys, Phlebotomus spp. blood meal analysis, and human patient clinical data from the mountainous area of central Negev in southern Israel during the summer months of 2018-2020. We found a fourth leishmaniasis transmission cycle associated with human illness.

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Materials and Methods

Study Area and Sand Fly Trapping

We conducted our study in the mountainous desert area of central Negev in southern Israel (Figure 1). In this region, elevations range from 50 to 1,037 m above sea level, large differences occur between peak daytime and nighttime temperatures, and annual average precipitation is 30–150 mm (20,21). We collected sand flies outdoors in August 2018, September 2019, and August 2020 by using modified traps from the



Figure 1. Locations of Phlebotomus spp. sand fly collection sites within the central Negev region of Israel in study of Leishmania donovani transmission cycle associated with human infection, Phlebotomus alexandri sand flies, and hare blood meals. We collected sand flies outdoors in August 2018, September 2019, and August 2020 by using modified traps from the US Centers for Disease Control and Prevention. The traps operated without light and were powered by 2 AA (1.2V) rechargeable batteries and baited with ≈1 kg dry ice. Traps were placed in an updraft vertical position overnight; openings were ≈10 cm above the ground, and collection cups hung above the motor and fan. Different colored circles indicate sites where specific Leishmania spp. infections were identified in trapped Phlebotomus sand flies. Empty circles indicate sites where sand flies were negative for Leishmania spp. Inset shows location of the survey area in Israel (red box).

US Centers for Disease Control and Prevention. The traps operated without light and were powered by 2 AA (1.2V) rechargeable batteries and baited with \approx 1 kg dry ice. We placed traps in an updraft vertical position overnight; openings were \approx 10 cm above the ground, and collection cups hung above the motor and fan (22,23).

Identification and Sample Preparation

We transferred live sand fly catches to the laboratory, which we then chilled and processed. We counted dead sand flies and sorted by sex, identifying all male flies at the species level by using specific morphologic keys for genitalia (24,25). We kept all engorged females and \leq 10–15 unfed females from each trap individually. If the number of female sand flies in the trap was >15, we pooled those flies with others in groups of 20 specimens each. We noted the blood meal size and freshness for each engorged female (26). We stored all female fly specimens in collection microtubes at –80°C until DNA extraction.

Molecular Analysis by Real-Time PCR, HRM Assay, and Sequencing

We extracted total DNA from sand fly samples by using the QIAsymphony DSP DNA Mini Kit and QIAsymphony SP robot (QIAGEN, https://www. qiagen.com). We homogenized the samples for 5 min in 50 μ L lysis buffer and stainless steel beads by using a TissueLyser II instrument (QIAGEN). The lysis buffer contained DNase- and proteinase-free RNaseA (ThermoFisher Scientific, https://www.thermofisher.com), proteinase K, and ATL tissue lysis buffer (QIAGEN). After homogenization, we added 200 μ L lysis buffer to each samples and incubated at 56°C for 2 h. We performed centrifugation and transferred the samples directly to the robot. We extracted DNA in accordance with the manufacturer's instructions and eluted the DNA in 100 μ L of elution buffer.

We performed all real-time PCR reactions by using a Roche LightCycler 96 (Roche, https://www.roche. com) and AccuMelt HRM SuperMix (Quantabio, https://www.quantabio.com). We analyzed all female sand flies for *Leishmania* spp. infection and single and engorged female flies to determine *Phlebotomus* sand fly species and blood meal source. We performed high-resolution melting (HRM) assays at the final step of each real-time PCR. We performed amplicon dissociation analysis by capturing fluorescence signals in 0.1°C/s increments and holding for 60 s in each range of the melting curve (60°C-85°C for sand fly species and blood meal detection assays or ≤95°C for *Leishmania* PCR). Sanger sequencing was performed at the Center for Genomic Technologies at Hebrew University of Jerusalem.

We screened all female sand flies for *Leishmania* DNA and identified parasite species by amplifying an internal transcribed spacer (ITS) 1 rRNA fragment with ITS1–219 PCR primers (Appendix Table 1, https://wwwnc.cdc.gov/EID/article/29/5/22-1657-App1.pdf) and by using the HRM assay (27). For PCR controls, we extracted DNA from parasite promastigote cultures of international reference strains: *L. major* (MHOM/PS/1967/JerichoII), *L. tropica* (MHOM/IL/1990/P283), *L. infantum* (MHOM/SD/62/2S), *L. donovani* (MHOM/SD/1962/1S-CLD2), and *L. aethiopica* (MHOM/ET/1972/L102). High purity water for molecular biology (Bio-Lab, http://www.biolabchemicals.com) was used as a negative control.

We included DNA isolated from skin lesions from 4 patients who had leishmaniasis diagnosed at the parasitology laboratory at Soroka Medical Center, Beer-Sheba, Israel; leishmaniasis was caused by *L. donovani/L. infantum* complex in those patients (Table 1). Leishmaniasis was diagnosed at the hospital by using multiplex real-time PCR with 5 probes for the ITS region of *Leishmania* sp. (28,29). We analyzed the samples further at the Ministry of Health by using real-time PCR-HRM amplification of the ITS1 fragment and ITS region and then sequencing.

We amplified the entire 1,020-bp ribosomal ITS region from *Leishmania*-positive field, clinical, and control samples by using PCR primers LITSR and LITSV (Appendix Table 1). If the entire ITS region was not successfully amplified with LITSR and LITSV primers, we used an internal pair of primers, L5.8S and L5.8SR, to amplify ITS1 and ITS2 separately (30). We performed ITS1 amplicon sequencing for 12 of the positive samples, and the entire ITS region was sequenced from 6 unfed and 3 engorged females, 3 pooled *Phlebotomus* spp. samples, all 4 human samples, and 4 *Leishmania*-positive controls. We amplified the repeat region of the *L. donovani* and *L. infantum HASPB* (known as K26) gene for additional

separation of *L. donovani* complex–positive samples by using primers K26F and K26R (*31*).

To identify *Phlebotomus* spp., we amplified a 368–393-bp fragment of the cytochrome b gene by using a universal primer set designed for this study (cytb-F and cytb-R; Appendix Table 1). The specificity of the designed primers was tested against DNA sequences from hematophagous arthropods, including sand flies, mosquitoes, and ticks. Male sand flies identified at the species level by using morphologic characteristics were used as positive controls and molecular biology grade water was used as a negative control. We analyzed all individual samples, and 1 third of samples from each melting curve pattern were sequenced.

We identified blood meal sources in *Phlebotomus* sand fly specimens by amplifying a 500-bp segment of host 12S and 16S mitochondrial rRNA genes by using modified vertebrate universal primers N12–16F and N12–16R (32). We included negative (water) and positive (100 ng of human DNA) controls in each PCR. We sequenced 50 samples that represented all HRM curve patterns and all female sand flies containing blood meals that had a melting curve of a rare host (<5 samples).

We used DNA from *Leishmania* reference strains, male sand fly specimens identified by morphologic characteristics, and human blood as templates for realtime PCR and HRM curve standardization. Each species produced a unique melting curve that was easily distinguishable from other species and consistent with observed nucleotide differences (Appendix Figures 1–3). We compared normalized HRM curves of field samples with the positive control included in each PCR, which enabled species determination (27). We validated species identification by sequencing 1 third of the samples; complete matches were observed for speciation by HRM curve analysis and DNA sequencing.

We aligned and corrected nucleotide sequences by using BioNumerics version 8.0 software (Applied Maths, https://www.applied-maths.com) and

Table 1. Human clinical samples from Soroka Medical Center in study of <i>Leishmania donovani</i> transmission cycle associated with							
human infection, <i>Phlebotomus alexandri</i> sand flies, and hare blood meals, Israel*							
		Age,					
Patient no.	Diagnosis	y/sex	Clinical description	Residence	Infecting <i>Leishmania</i> sp.†		
1	VL	47/M	Splenomegaly	Northern	L. infantum (100% identity with L. infantum, GenBank		
				Israel	accession no. KU680954)		
2	VL	4/F	Splenomegaly,	Hebron	L. infantum (99.19% identity with L. infantum, GenBank		
			hepatomegaly		accession no. MN503527)		
3	CL	69/F	Skin ulcer	Negev	L. infantum (99.87% identity with L. infantum, GenBank		
					accession no. KU680954)		
4	CL	51/M	Skin ulcer	Arava	L. donovani (99.75% identity with L. donovani, GenBank		
					accession no. LC459330)		

*CL, cutaneous leishmaniasis; VL, visceral leishmaniasis

+Leishmania spp. were identified by PCR high resolution melting curves and Sanger sequencing.

			_		
		MW534746 Leishmania infantum MHOM/SD/62/2S / Promastigote culture / Israel			
		MZ366761 Leishmania infantum (VL478) / Human / Israel			
		MZ366762 Leishmania infantum (CL409) / Human / Israel			
		MZ366763 Leishmania infantum (VL489) / Human / Israel			
		MW587846 Leishmania infantum (SF25609) / Phlebotomus / Israel			
		MW587845 Leishmania infantum (SF21413) / Phlebotomus / Israel			
		MZ366768 Leishmania infantum (SF1) / Phlebotomus / Israel			
	94	GU591397 Leishmania infantum / Jackal / Israel		Leishmania	
[KM677141 Leishmania infantum MCAN/IL/2011/TR4 / Dog / Israel		infantum	
		KM677143 Leishmania infantum MCAN/IL/2011/TA2 / Dog / Israel			
		KM677133 Leishmania infantum MCAN/IL/2011/NT6 / Promastigote culture / Israel			
		KM677128 Leishmania infantum / Promastigote culture / Israel			
		MK311268 Leishmania infantum / Human / West Bank			
		KU975157 Leishmania infantum MHOM/CN/2009/KSF-4 / Promastigote culture / China			
72		MT302167 Leishmania infantum / Human / Iran			
		MN245034 <i>Leishmania infantum /</i> Cat / Brazil	_		
		MG564280 Leishmania infantum / Dog / Egypt	٦		
		MW534748 Leishmania donovani MHOM/SD/1962/1S-CLD2 / Promastigote culture / Israel			
		MZ366765 Leishmania donovani (CL659) / Human / Israel			
		MW587836 Leishmania donovani (SF2222) / Phlebotomus / Israel			
		MW587838 Leishmania donovani (SF2262) / Phlebotomus / Israel			
		MW587839 Leishmania donovani (SF2096) / Phlebotomus / Israel		Leishmania	
		MZ366764 Leishmania donovani (SF2103) / Phlebotomus / Israel		donovani	
	89	MZ366766 Leishmania donovani (SF7321) / Phlebotomus alexandri / Israel			
		MZ366767 Leishmania donovani (SF11658) / Phlebotomus alexandri / Israel			
		KY658229 Leishmania infantum / Promastigote culture / Morocco			
		KY982638 Leishmania donovani / Human / Thailand			
		AJ634361 Leishmania infantum MHOM/SD/62/3S / Human / Sudan			
		AJ000291 Leishmania donovani MHOM/SD/75/LV139 / Promastigote culture / Sudan			
		KU949582 <i>Leishmania major</i> / Dog / Israel	-		
	99	MN604098 Leishmania major / Human / Israel		Leishmania	
		MZ366759 Leishmania major MHOM/PS/1967/Jericholl / Promastigote culture / Israel		major	
		GQ920673 Leishmania aethiopica / Promastigote culture / USA	_		
	100	HG512946 Leishmania aethiopica MHOM/ET/70/L96 / Promastigote culture / Belgium		Leishmania	
		EU683619 Leishmania aethiopica MHOM/ET/1972/L102 / Promastigote culture / Israel		aethiopica	
		FJ595949 Leishmania tropica / Rock hyrax / Israel	٦		
		MZ366760 Leishmania tropica MHOM/IL/1990/P283 / Promastigote culture / Israel			
		KY524300 Leishmania tropica / Dog / Israel			
		GU591396 Leishmania tropica / Jackal / Israel		Leishmania	
		FN677345 Leishmania tropica MHOM/PS/01/ISL590 / Human / West Bank		tropica	
	95	MW111286 Leishmania tropica / Phlebotomus sergenti / West Bank			
		GU591395 Leishmania tropica / Fox / Israel			
		EU683617 Leishmania tropica ISER/IL/1998/LRC-L758 / Promastigote culture / Israel			
		MT966013 Leishmania tropica / Phlebotomus sergenti / West Bank	J		

Figure 2. Phylogenetic analysis of *Leishmania* internal transcribed spacer 1 rRNA fragments in study of *Leishmania donovani* transmission cycle associated with human infection, *Phlebotomus alexandri* sand flies, and hare blood meals, Israel. *Leishmania*-specific internal transcribed spacer 1 rRNA fragments (201 bp) were amplified by PCR from *P. alexandri* sand flies, pooled female *Phlebotomus* spp. flies, and patient samples and then sequenced. Tree was constructed by using the maximum-likelihood method and Tamura 3-parameter model, estimated by using the Aikaike information criterion (*33*). Dendogram includes sequences from *L. donovani* and *L. infantum* isolated from sand flies and clinical samples in this study compared with *Leishmania* spp. reference controls and GenBank sequences from Israel and other countries. Tree shows substantial separate clustering of *L. infantum* (boostrap 94%) and *L. donovani* (bootstrap 89%) sequences. Empty circles are *Leishmania* international reference strains, black triangles are the 10 sequences from our study deposited in GenBank, and black circles are additional *L. infantum*—positive sand flies samples from Israel. Available GenBank sequences for *L. major, L. tropica, L. infantum*, and *L. donovani* from Israel and other countries are also included. GenBank accession numbers, *Leishmania* spp., isolate source, and country are indicated. Only bootstrap values >70% are shown. Not to scale.

compared sequences against the GenBank database by using BLASTN (http://blast.ncbi.nlm.nih.gov). We identified *Leishmania* spp., blood meal sources, and sand fly species on the basis of >98% identity with sequences obtained during the BLAST search. We submitted sequences of the ITS1 fragments and entire ITS and *K*26 regions obtained in this study to GenBank (Appendix Table 2).

We constructed phylogenetic trees on the basis of marker gene sequences in this study and relevant



Figure 3. Phylogenetic analysis of entire *Leishmania* internal transcribed spacer region in study of *Leishmania donovani* transmission cycle associated with human infection, *Phlebotomus alexandri* sand flies, and hare blood meals, Israel. *Leishmania*-specific internal transcribed spacer region (988 bp) was amplified by PCR from *P. alexandri* sand flies, pooled female *Phlebotomus* spp. flies, and patient samples and then sequenced. Tree was constructed by using by using the maximum-likelihood method and Tamura 3-parameter model of all relevant *Leishmania* spp. and *Trypanosoma cruzi* as an outgroup. Sand fly and clinical samples from this study (black triangles), *L. infantum* isolates from Israel (black circles), *Leishmania* international reference strains (empty circles), and available GenBank *Leishmania* sequences are shown. GenBank accession numbers, isolate source, and country of origin are shown for each sequence. Only bootstrap values >70% are shown next to branches. Not to scale.



Figure 4. Phylogenetic analysis of *Leishmania K26* gene in study of *Leishmania donovani* transmission cycle associated with human infection, *Phlebotomus alexandri* sand flies, and hare blood meals, Israel. *Leishmania*-specific *K26* gene fragment (348 bp) was amplified by PCR from *P. alexandri* flies, pooled female *Phlebotomus* spp. flies, and patient samples and then sequenced. Tree was constructed by using the maximum-likelihood method and Hasegawa-Kishino-Yano model. *K26* phylogenetic analysis shows separation between *L. infantum* and *L. donovani*. Sand fly and clinical samples from this study (black triangles), *L. infantum* isolates from Israel (black circles), *Leishmania* international reference strains (empty circles), and available GenBank *Leishmania* sequences are shown. GenBank accession number, isolate source, and country of origin are shown for each sequence. Only bootstrap values >70% are shown next to branches. Not to scale.

sequences of other *Leishmania* spp. deposited in Gen-Bank. We used MEGA X software (33) to infer phylogenetic trees after nucleotide sequence alignment was performed by using ClustalW software (http:// www.clustal.org) and maximum-likelihood and neighbor-joining algorithms. We used 1,000 bootstrap replicates to determine percentages of replicate trees. We constructed a phylogenetic tree composed of 45 analyzed partial sequences of the ITS1 locus, including sequences of *Leishmania* spp. from Israel and other countries deposited in GenBank and *Trypanosoma* *cruzi* as an outgroup (Figure 2). We constructed a second tree that included 30 nearly complete ITS sequences of all relevant *Leishmania* spp. and *T. cruzi* as an outgroup (Figure 3) and an additional phylogenetic tree that included 20 *K26* gene sequences of *L. donovani* complex–positive samples (Figure 4).

Results

We collected 22,636 *Phlebotomus* spp. sand fly specimens (15,720 female and 6,916 male; sex ratio 2.3) during 7 trapping nights by using 118 traps placed

_	20	18	20	19	20	20		
Phlebotomus spp.	F	Μ	F	М	F	Μ	Total no. (%)	
P. alexandri	129	1,452	70	126	431	4,039	6,247 (80.0)	
P. kazeruni	38	837	22	11	78	142	1,128 (14.5)	
P. sergenti	5	109	26	9	34	56	239 (3.1)	
P. papatasi	25	97	8	8	8	27	173 (2.2)	
P. syriacus	0	0	3	0	17	3	23 (0.3)	
Not identified*	8,344	0	265	0	6,217	0	14,826	
Total	8,541	2,495	394	154	6,785	4,267	22,636	
*Sand flies were pooled	'Sand flies were pooled before molecular testing in batches of 20 speciments each.							

 Table 2. Number of female and male Phlebotomus spp. collected during 2018–2020 in central Negev in study of Leishmania donovani transmission cycle associated with human infection, Phlebotomus alexandri sand flies, and hare blood meals, Israel

at 94 sites. After identifying all male and 894 (6%) female flies, we found the catches consisted of 5 species. The most abundant sand fly species were *P. alexandri* (80%), *P. kazeruni* (14.4%), *P. sergenti* (3.1%), *P. papatasi* (2.2%), and *P. syriacus* (0.3%) (Table 2).

Among the 4,140 unfed female sand flies tested in 210 pools, we found 41 pools were positive for *Leish*mania spp. In addition, 6/688 single female flies and 4/206 engorged female flies were positive for Leishmania spp. Of the 51 Leishmania-positive samples, the HRM curves of 47 (36 pools, 6 single females, and 3 engorged female flies) were similar to the HRM curve of the L. donovani control (Table 3; Appendix Figure 1). The HRM curves for 2 pooled fly samples from 2018 were identical to the HRM curve of the L. tropica control. One pooled fly sample and 1 engorged female fly collected in 2020 had an HRM curve identical to the L. major control. The ITS1-PCR sequences of 20 samples (11 pools and all 9 single and engorged females) that had HRM curves similar to the L. donovani control HRM curve were also 100% identical to the L. donovani control sequence.

Leishmaniasis was diagnosed in 4 human patients (Table 1). The ITS1 HRM curve and sequence from patient 4 with CL were similar to the *L. donovani* control and 47 *L. donovani*–positive *Phlebotomus* spp. samples. The ITS1 HRM and sequences from patients 1 and 2 with VL and patient 3 with CL were similar to the L. infantum control. Alignment of ITS1 sequences from L. infantum and L. donovani controls, 3 representative sand fly samples showing HRM identical to L. donovani, and the 4 patient samples showed clustering into 2 distinct groups. The first group comprised the L. donovani control, 3 sand fly samples, and patient 4. The second group comprised the L. infantum control and samples from patients 1, 2, and 3. The difference between the groups was at position 71-74 in ITS1; the L. donovani group had a 4-nt (ATAT) insertion that was missing in the *L. infantum*-positive samples. A comparison of ITS1 sequences with those in GenBank showed 100% query coverage and 99.65%-100% identity with GenBank sequences for L. infantum and L. donovani from various countries (data not shown).

We aligned DNA sequences from the entire ITS region obtained from *Leishmania*–positive *P. alexandri* samples from our study and the *L. infantum* and *L. donovani* controls (Appendix Figure 4). We found 2 additional regions containing polymorphic sites that distinguished between *L. infantum* and *L. donovani*: a 2-nt (GG) deletion at position 724–725 and 1-nt (G) insertion at position 817 in the *L. donovani* sequence.

We constructed a phylogenetic tree of ITS1 rRNA fragments of *Leishmania* sequences obtained from *P. alexandri* flies, pooled female *Phlebotomus* spp. flies, and

Table 3. Number of Leishmania spl transmission cycle associated with	 detected in sand fly samples human infection, <i>Phlebotomus</i> 	s by PCR during 2018–20 s alexandri sand flies, and	20 in study of <i>Leishma</i> hare blood meals, Isra	ania donovani ael*
Year	Total no. tested	L. donovani	L. tropica	L. major
2018				
Females (no. pools)	2,938 (148)	24	2	0
Single unfed females	108	0	0	0
Single engorged females	89	0	0	0
2019				
Females (no. pools)	262 (15)	4	0	0
Single unfed females	121	3	0	0
Single engorged females	8	0	0	0
2020				
Females (no. pools)	940 (47)	10	0	1
Single unfed females	459	3	0	0
Single engorged females	109	3	0	1
Total no	5 019	47	2	2

*All engorged females and ≤10–15 unfed females from each trap were maintained and analyzed individually. If the number of female sand flies in the trap was >15, they were pooled in batches of 20 specimens each.

Total no.

meals, Israel					
Blood meal source	P. alexandri	P. kazeruni	P. papatasi	P. sergenti	Total no.
Lepus europaeus (hare)	107	8	5	6	126
Equus hemionus (onager)	31	1	0	1	33
Gazella dorcas (gazelle)	10	0	4	2	16
Canis lupus familiaris (dog)	1	0	3	0	4
Homo sapiens (human)	1	0	0	0	1
Psammomys obesus (fat sand rat)	0	0	1	0	1
Vulpes vulpes (fox)	1	0	0	0	1

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Table 4. Number of female Phlebotomus spp. sand flies collected in the central Negev region engorged with different blood meals in study of Leishmania donovani transmission cycle associated with human infection. Phlebotomus alexandri sand flies, and hare blood

patient samples from this study. We compared those sequences with Leishmania spp. controls and GenBank sequences from Israel and other countries. The tree showed substantial separate clustering of *L. infantum* (boostrap 94%) and L. donovani (bootstrap 89%) sequences (Figure 2). Phylogenetic analysis of the entire ITS region showed separate clustering of *L. infantum* (bootstrap 84%) and L. donovani (bootstrap 95%) sequences (Figure 3). K26 phylogenetic analysis also showed separation between *L. infantum* and *L. donovani* (Figure 4).

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We identified the blood meal source for 182/206 (88%) engorged female sand flies that represented the 4 most abundant sand fly species within the study area. We observed 7 types of HRM curves. We compared blood meal sequences with GenBank sequences and determined similarities between HRM curves. We identified European brown hare (Lepus europaeus) blood in 126 (69.2%) flies, onager (Equus hemionus) blood in 33 (18.3%) flies, gazelle (Gazella dorcas) blood in 16 (8.8%) flies, and domestic dog (C. lupus familiaris) blood in 4 (2.2%) flies; 1 female sand fly each contained blood from either a fat sand rat (Psammomys obesus), fox (V. vulpes), or human (Table 4). Hare blood was the dominant blood meal found in all 4 Phlebotomus spp. flies: P. papatasi, 38%; P. sergenti, 67%; P. alexandri, 71%; and P. kazeruni, 89%. The 12S-16S hare blood meal sequences were 99.8% similar to L. europaeus hares and only 95.3% similar to *L. capensis* hares.

Of the 47 *Phlebotomus* spp. sand fly samples with HRM curve patterns and sequences similar to the L. donovani control, 9 were single P. alexandri female sand flies, 3 of which were engorged with hare blood. Of the 2 Phlebotomus spp. samples positive for L. major, 1 was in a single engorged P. papatasi female sand fly that had an unsuccessful blood meal identification. The 2 identified *L. tropica* samples were from pooled female sand flies.

Discussion

We found a fourth transmission cycle of leishmaniasis in the central Negev region of southern Israel.

On the basis of molecular analysis of the ITS region and K26 gene and phylogenetic analysis, we concluded that the parasite found in patient 4, who lives in the survey area, and in *P. alexandri* sand flies was L. donovani sensu stricto. We found that L. infantum was the cause of illness in the other 3 patients with leishmaniasis. A case report describing patient 4 was published in 2016; the authors concluded that the infecting parasite was likely L. infantum because of prevailing knowledge of endemic Leishmania transmission in Israel (29). An earlier study reported another patient from the Arava region of central Negev, close to where patient 4 lives, who had symptoms of both CL and VL (34). The cause of infection was identified as L. donovani; the authors noted that this infection was unusual because L. donovani was not known to circulate in Israel. The earlier study substantiates our findings of L. donovani in both sand flies and another human patient within the same geographic area. L. infantum was identified as the causative agent in the other 3 patients in our study and was also described in canines in Israel (1).

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The high abundance of *P. alexandri* sand flies within the study area and the association with *L. don*ovani infections suggest that the P. alexandri sand fly is the putative vector of L. donovani in Israel. P. alexandri flies have been associated with L. donovani sensu latu transmission in other parts of the Old World. Natural infection by L. donovani was found in field-collected P. alexandri sand fly specimens in China, and inoculation of hamsters with those parasites caused VL (35). Another study reported the susceptibility of *P. alexandri* to artificial infection with *L. donovani* isolated from human patients in China (36).

We found blood meals from European brown hares in $\approx 70\%$ of engorged female *Phlebotomus* sand flies in our study. The high feeding rates on hares, presence of L. donovani in female P. alexandri sand flies engorged with hare blood and illnesses reported in humans infected with L. donovani suggest a zoonotic L. donovani transmission cycle in Israel. Those data suggest that the hare could be a potential reservoir and P. alexandri flies could be the putative vector for L. donovani. The role of hares as a reservoir host for L. donovani requires further investigation; however, a related hare species, Lepus granatensis, was reported as a potential sylvatic reservoir for L. infantum in a leishmaniasis outbreak in Madrid, Spain (37,38). Furthermore, studies in Greece and Italy detected L. donovani complex infection in L. europaeus hares (39,40), providing support for hares as a potential reservoir for L. donovani in Israel. Dogs were identified as reservoirs for L. donovani in India, Sudan, and Ethiopia, and different rodent species have been identified as possible reservoirs of *Leishmania* spp. from the *L*. donovani complex (41-48). However, no L. donovani infections in canines and rodents have been reported in Israel; infections in sand fly blood meals found in our study do not implicate those hosts in the local life cvcle of L. donovani.

In conclusion, we found circulation of L. donovani in the Negev region of southern Israel that was associated with cutaneous lesions in humans. We determined that P. alexandri was the putative sand fly vector and that hares were the main reservoir host of L. donovani. We found 2 distinct Leishmania spp. in the L. donovani complex in Israel. Previously, the few reported human cases of CL resulting from L. donovani infections were attributed to either L. infantum or nonautochthonous infections. Analysis of patient samples in our study indicates that, in addition to L. major and L. tropica (the known agents causing CL), L. donovani is also a cause of autochthonous CL in Israel. Our results suggest that CL in Israel can be caused by L. donovani, a primary cause of VL. Therefore, prompt diagnosis, identification of the Leishmania sp., and treatment with drugs intended for visceral leishmaniasis, such as pentavalent antimonials or liposomal amphotericin B (49), are critical to prevent disease progression and death among patients with leishmaniasis.

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etymologia revisited

Dermatophilus congolensis

[dur"mə-tof'i-ləs con-go-len'sis]

From the Greek *derma* (skin) + *philos* (loving), *Dermatophilus congolensis* is a Gram-positive, aerobic actinomycete, and facultatively anaerobic bacteria (Figure 1). D. congolensis infects the epidermis and produces exudative dermatitis termed dermatophilosis that was previously known as rain rot, rain scald, streptotrichosis, and mycotic dermatitis.

In 1915, René Van Saceghem (Figure 2), a Belgian military veterinarian stationed at a veterinary laboratory in the former Belgian Congo (thus, the species name *congolensis*), reported *D. congolensis* from exudative dermatitis in cattle. Local breeders and veterinarians had observed the disease since 1910, but the causal agent was not identified.

Dermatophilosis affects animals, mainly cattle, and more rarely humans. Outbreaks of *D. congolensis* infection have severe economic implications in the livestock and leather industries.

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Leishmania donovani Transmission Cycle Associated with Human Infection, Phlebotomus alexandri Sand Flies, and Hare Blood Meals, Israel

Appendix

Appendix Table 1. Real-time PCR primers and protocols used for the detection and identification of Leishmania species, blood	b
meal sources, and sand fly species*	

Target gene	Primer	Amplicon, bp	Reference ⁺ or PCR protocol
ITS1	ITS1–219F: AGCTGGATCATTTTCCGATG	265	(27)
	ITS1–219R: ATCGCGACACGTTATGTGAG		
ITS	LITSR: CTGGATCATTTMCGATG	1020	(30)
	LITSV: ACACTCAGGTCTGTAAAC		
ITS1	LITSR: CTGGATCATTTMCGATG	320	
	L5.8S: TGATACCACTTATCGCACTT		
ITS2	L5.8SR: AAGTGCGATAAGTGGTA	700	
	LITSV: ACACTCAGGTCTGTAAAC		
K26	K26F: ACGAAGGACTCCRCAAAG	350	(31)
	K26R: TTCCCATCGTTTTGCTG		
cytb‡	cytb-F: GGAGGAGTAATYGCHYTTGTWATATC	368–393	95°C, 5 min; then 45 cycles of 95°C, 5 s;
	cytb-R: AAGATATTTACCYGCTTCKTTATGTT		38°C, 2 s; 60°C, 45 s; 95°C, 60 s; 40°C,
			60 s; 65°C, 1 s; 85°C, 1 s; 37°C, 30 s
12S, 16S	N12–16F: ACAYACCGCCCGTCACCCTC	500 bp	(32)
	N12-16R AACCAGCTATCACMAGGCTCG		

*ITS, internal transcribed spacer, entire region; ITS1, internal transcribed spacer 1; ITS2, internal transcribed spacer 2; 12S, 16S, mitochondrial rRNA gene; cytb, cytochrome b gene.

†References are from the main text.

‡Primers designed for our study.

Appendix Table 2. Details of the samples from this study submitted to GenBank*

Accession no.†	Leishmania sp.	Isolate	Source	Locus	Size, bp
MW587834	Leishmania donovani	SF2101	Phlebotomus	ITS1	200
MW587835	Leishmania donovani	SF2103	Phlebotomus	ITS1	228
MW587836	Leishmania donovani	SF2222	Phlebotomus	ITS1	287
MW587837	Leishmania donovani	SF2226	Phlebotomus	ITS1	170
MW587838	Leishmania donovani	SF2262	Phlebotomus	ITS1	290
MW587839	Leishmania donovani	SF2096	Phlebotomus	ITS1	287
MW587841	Leishmania donovani	CL659 (#4)	Human	ITS1	232
MW587842	Leishmania infantum	CL409 (#3)	Human	ITS1	227
MW587843	Leishmania infantum	VL478 (#1)	Human	ITS1	246
MW587844	Leishmania infantum	VL489 (#2)	Human	ITS1	245
MW587845	Leishmania infantum	SF21413	Phlebotomus	ITS1	312
MW587846	Leishmania infantum	SF25609	Phlebotomus	ITS1	198
MW534746	Leishmania infantum	MHOM/SD/62/2S	Promastigote culture	ITS	1,024
MW534748	Leishmania donovani	MHOM/SD/1962/1S-CLD2	Promastigote culture	ITS	980
MZ366759	Leishmania major	MHOM/PS/1967/Jericholl	Promastigote culture	ITS	815
MZ366760	Leishmania tropica	MHOM/IL/1990/P283	Promastigote culture	ITS	783
MZ366761	Leishmania infantum	VL478 (#1)	Human	ITS	782
MZ366762	Leishmania infantum	CL409 (#3)	Human	ITS	783
MZ366763	Leishmania infantum	VL489 (#2)	Human	ITS	782
MZ366764	Leishmania donovani	SF2103	Phlebotomus	ITS	782

Accession no.†	<i>Leishmania</i> sp.	Isolate	Source	Locus	Size, bp
MZ366765	Leishmania donovani	CL659 (#4)	Human	ITS	787
MZ366766	Leishmania donovani	SF7321	P. alexandri	ITS	786
MZ366767	Leishmania donovani	SF11658	P. alexandri	ITS	785
ON796535	Leishmania donovani	Khartoum	ATCC	ITS	967
ON796536	Leishmania donovani	SF21301	Phlebotomus	ITS	985
ON796537	Leishmania infantum	MHOM/TN/80/IPT-1	ATCC	ITS	985
ON796538	Leishmania infantum	CL109	Human	ITS	1,029
ON796539	Leishmania infantum	CL110	Human	ITS	1,018
ON796540	Leishmania infantum	SF21413	Phlebotomus	ITS	712
ON858810	Leishmania infantum	CL110	Human	k26	766
ON858811	Leishmania infantum	VL478	Human	k26	620
ON858812	Leishmania donovani	CL659	Human	k26	360
ON858813	Leishmania donovani	SF14779	Phlebotomus	k26	361
ON858814	Leishmania donovani	SF7274	Phlebotomus	k26	367
ON858815	Leishmania donovani	SF7321	P. alexandri	k26	370
ON858816	Leishmania donovani	SF11658	P. alexandri	k26	365
ON858817	Leishmania donovani	SF19824	Phlebotomus	k26	371
ON858818	Leishmania donovani	SF21301	Phlebotomus	k26	362
ON858819	Leishmania infantum	SF21413	Phlebotomus	k26	504
ON858820	Leishmania infantum	SF25609	Phlebotomus	k26	478
ON858821	Leishmania donovani	Khartoum	ATCC	k26	278
ON858822	Leishmania donovani	MHOM/SD/1962/1S-CLD2	Promastigote culture	k26	278
ON858823	Leishmania infantum	MHOM/TN/80/IPT-1	ATCC	k26	619
ON858824	Leishmania infantum	MHOM/SD/62/2S	Promastigote culture	k26	616

*ITS, internal transcribed spacer, entire region; ITS1, internal transcribed spacer 1.

†GenBank accession no.



Appendix Figure 1. Normalized high resolution melting curves of 370-bp cytochrome b PCR amplicons from different *Leishmania* spp. isolated from sand flies trapped in the central Negev region, Israel. Normalized fluorescence is plotted against temperature.



Appendix Figure 2. Normalized high resolution melting curves of *Leishmania*-specific internal transcribed spacer 1 rRNA fragment PCR amplicons isolated from different *Phlebotomus* spp. sand flies trapped in the central Negev region, Israel.



Appendix Figure 3. Normalized high resolution melting curves of blood meal PCR amplicons from different animal sources found in engorged sand fly females trapped in the central Negev region, Israel.

	ITS1 poly (TA)	ITS2 poly (G)	ITS2 poly (GA)	ITS2 poly (G)
L. donovani / P. alexandri	() TATATATAT <mark>ATAT</mark> GTAG	5G () GGGG- –TCG	AGGGAGAGAGGCT) AATGGGGGG <mark>G</mark> AGGT ()
L. donovani MHOM/SD/1962/1S-CLD2	() TATATATAT <mark>ATAT</mark> GTAG	3G () GGGG– –TCG	AGGGAGAGAGGCT) AATGGGGGG <mark>G</mark> AGGT ()
L. infantum MHOM/SD/62/2S	() TATATATAT GTAG 62 7	66 () 6666 <mark>66</mark> 7C6 9 720	AGGGAGAGAGGCT (738) AATGGGGGGG-AGGT () 808 821

Appendix Figure 4. Alignment of sequences from the entire internal transcribed spacer region of the *18S* gene from *Leishmania donovani* found in *Phlebotomus alexandri* sand flies in the central Negev, Israel, and *L. donovani* and *L. infantum* international reference strains. Nucleotides marked in blue represent insertions, and discontinuous lines represent absence of nucleotides in sequences. Numbers correspond to nucleotide sequence position in the region. ITS1, internal transcribed spacer 1; ITS2, internal transcribed spacer 2.