Transmission of *Leishmania tropica* was studied in 2 adjacent foci in Israel where vector populations differ. Only *Phlebotomus sergenti* was found infected with *L. tropica* in the southern focus; *P. arabicus* was the main vector in the northern focus. Rock hyraxes (*Procavia capensis*) were incriminated as reservoir hosts in both foci. *L. tropica* strains from the northern focus isolated from sand flies, cutaneous leishmaniasis cases, and rock hyraxes (*Procavia capensis*) were antigenically similar to *L. major*, and strains from the southern focus were typically *L. tropica*. Laboratory studies showed that *P. arabicus* is a competent vector of *L. tropica*, and *P. sergenti* is essentially refractory to *L. tropica* from the northern focus. Susceptibility of *P. arabicus* may be mediated by O glycoproteins on the luminal surface of its midgut. The 2 foci differ with respect to parasites and vectors, but increasing peridomestic rock hyrax populations are probably responsible for emergence of cutaneous leishmaniasis in both foci.

*Leishmaniases* are parasitic diseases with a wide range of clinical symptoms and currently threaten 350 million persons in 88 countries (1). In Israel and its vicinity, *Leishmania major* and *L. tropica* cause cutaneous leishmaniasis (CL), and *L. infantum* can result in visceral leishmaniasis (2). Until recently, relatively little information was available on the epidemiology of CL caused by *L. tropica* in this region. Outbreaks were not investigated, and cases were usually grouped together with CL cases caused by *L. major* (3). However, in recent years, new foci of CL caused by *L. tropica* are emerging in different parts of the country, such as the Galilee region of northern Israel and the Judean Desert east of Jerusalem that warrant thorough investigations (4,5). Clinically, lesions caused by *L. tropica* last longer and are more difficult to treat than those caused by *L. major* (6). Although *L. tropica* can be anthropotonic, foci in Israel appear to be zoonotic, with rock hyraxes (*Procavia capensis*) serving as probable reservoir hosts (4).

*Leishmania* development in sand flies is facilitated by interaction with midgut molecules of the vector. Laboratory studies showed that sand flies are composed of 2 groups. Species such as *Phlebotomus (Phlebotomus)* papatasi, the vector of *L. major* and *P. (Paraphlebotomus) sergenti*, the main vector of *L. tropica*, show specificity for *Leishmania* they transmit in nature (7,8). Conversely, species such as *Lutzomyia longipalpis*, the vector of *L. infantum* in South America, and many others are permissive and support development of several *Leishmania* spp. (8,9).

Studies performed with *L. major* and *P. papatasi* showed that attachment in the midgut is mediated by O glycosylated terminal epitopes on the luminal surface of its midgut. The 2 foci differ with respect to parasites and vectors, but increasing peridomestic rock hyrax populations are probably responsible for emergence of cutaneous leishmaniasis in both foci.
and P. (Paraphlebotomus) sergenti that transmit *L. tropica* in 2 adjacent foci in the Galilee region of northern Israel.

*L. tropica* is genetically heterogeneous, and strains are readily distinguishable by antigenic, biochemical, and molecular techniques (14–16). We report findings of extensive studies in 2 adjacent CL foci that demonstrate conclusively that both vector species and parasite strains from the northern focus are different from those in the southern focus, a mere 10 km away (Figure 1).

**Materials and Methods**

**Study Area**

Studies were conducted in 2 adjacent foci in the Galilee region of northern Israel (Figure 1). The northern focus comprises several villages situated on generally south-facing slopes ≈5 km north of Lake Kinneret in the eastern lower Galilee of northern Israel (32°55′N, 35°36′W). The area investigated encompasses the villages of Amnun (at sea level), Karkom (100–150 m above sea level), and Korazim (150 m above sea level), which have ≈1,200 inhabitants living in ≈300 single-family houses surrounded by gardens and built on basalt rock. Many boulders from the cleared land have been piled into large heaps separating individual plots and surrounding the villages. These boulder mounds are inhabited by numerous rock hyraxes (*P. capensis*)

The southern focus includes the city of Tiberias (32°47′N, 35°32′W; population = 38,952). Studies were conducted in the outskirts of urban neighborhoods, where boulder mounds were inhabited by large populations of rock hyraxes. These neighborhoods are built on north- to northeast-facing slopes.

**Collection, Dissection, and Identification of Sand Flies**

Sand flies were trapped by using CDC light traps (John W. Hock, Gainesville, FL, USA) in September 2002 and 2004. Dead flies were stored in 70% alcohol and identified by using several keys (17–19). Live female flies were immobilized on ice, rinsed briefly in 96% ethanol, and dissected in 0.9% sterile saline. Guts were microscopically examined for parasites. Heads and genitalia were used for identification. Guts containing promastigotes were aseptically placed in glass vials (2.5 mL) containing blood agar made from defibrinated rabbit blood overlaid with a 1:1 mixture of RPMI 1640 medium and Schneider *Drosophila* cell culture medium supplemented with 10% fetal calf serum (Sigma, Saint Louis, MO, USA, and Gibco-BRL, Gaithersburg, MD, USA), 10,000 IU penicillin (Biotika, L’upea, Slovakia), 100 µg/mL amikacin (Bristol-Myers Squibb, Princeton, NJ, USA), and 1,500 µg/mL 5-fluorocytosine (Sigma). Some data on *Leishmania* isolates from the northern focus were obtained from our previously published results.

**Collection of Animals**

Rock hyraxes were trapped by using raccoon traps (http://www.havahart.com) baited with fresh leaves and anesthetized with ketamine (10 mg/kg given intramuscularly). Samples of blood and skin were obtained for parasite culture and blotted onto filter paper for PCR analysis. Animals were released at the site of capture. Skin biopsy specimens were homogenized and placed in blood agar culture medium in flat tubes (Nuncelon; Nunc Nalgene International, Rochester, NY, USA). Rats (*Rattus rattus*) were trapped by using steel mesh traps (Tomahawk Live Trap Co., Tomahawk, WI, USA) placed in sewers and rock crevices. Spiny mice (*Acomys cahirinus*) were captured by using Sherman traps (H.B. Sherman Traps, Tallahassee, FL, USA). Rodents were anaesthetized with ketamine/xylazine (150 mg/kg and 15 mg/kg, respectively, given intraperitoneally). Blood from the tip of the tail was blotted on filter paper. Ear biopsy specimens were treated as described for hyraxes. Cultures were checked at 4–7-day intervals for 1 month.
DNA Extraction
DNA from wild-caught sand flies kept frozen or preserved in 100% ethanol was extracted as previously described (20). DNA from filter paper disks was extracted by using the phenol-chloroform method (21).

Detection and Identification of Leishmania infections by PCR
The ribosomal internal transcribed spacer region 1 (ITS1) was amplified with Leishmania-specific primers. ITS1 PCR products showing a Leishmania-specific band on agarose gels were digested with HaeIII for species identification (22). Restriction fragments were subjected to electrophoresis on agarose gels and compared with DNA of L. infantum (Li-L699), L. major (Lm-L777), and L. tropica (Lt-L590).

Antigenic Characterization of Parasite Isolates
Initial screening of isolates was performed by using gel diffusion of glycoconjugates secreted into culture media (excreted factor) and several antileishmanial serum samples (23). Leishmania-specific monoclonal antibodies (MAbs) were used in indirect immunofluorescent antibody (IFA) assays to determine surface antigenic characteristics of parasites (14). Briefly, promastigotes from primary cultures of new isolates and controls of L. infantum (Li-L699), L. major (Lm-L777), and L. tropica (Lt-L590) were placed in wells of fluorescent antibody slides (Bellco Glass Inc., Vineland, NJ, USA), dried, and fixed in cold acetone. Slides were blocked with 5% fetal bovine serum in phosphate-buffered saline (PBS) for 1 hour at room temperature. Mouse MAbs specific for L. major (T1), L. tropica (T11, T14, and T15), L. tropica/L. major (T3), and L. infantum/L. donovani (D2) were applied for 1 hour at 37°C. Goat anti-mouse immunoglobulin G conjugated with fluorescein isothiocyanate was applied for 40 minutes at 37°C in the dark. The preparations were washed 3 times with PBS plus 5% Tween 20 between incubations. Slides were mounted in 3% DABCO (Sigma) in PBS/glycerol and viewed with an Axiovert microscope (Zeiss, Göttingen, Germany).

Experimental Infection of Sand Flies
Laboratory colonies of P. sergenti and P. arabicus females were established from gravid females caught in the northern focus. The colonies were maintained at 23°C–25°C, 100% humidity, and 14:10 light:dark photoperiod. Adults had access to cotton wool soaked in 50% honey. Females were allowed to feed twice a week on mice anaesthetized with a ketamine/xylazine mixture (150 mg/kg and 15 mg/kg). Fed females were placed in plaster of paris–lined oviposition containers, and larvae were maintained on a decaying rabbit feces/rabbit chow mixture (24). Sand flies were infect-ed by membrane feeding on heat-inactivated rabbit blood containing 5×10^5 promastigotes/mL. Fed females were maintained at 23°C and dissected on day 9 after feeding, when infections were mature. Guts were checked microscopically for Leishmania promastigotes. Infection intensity was scored as light (<50 promastigotes/gut), moderate (50–500 promastigotes/gut), and heavy (>500 promastigotes/gut). L. tropica strains from the northern (IARA/IL/2001/L810, Amnumfly1) and southern (MHOM/IL/2001/L-836, Tiberias) foci were used for comparing susceptibility of sand flies to local strains. Promastigotes from the same culture and sand flies from the same batch were used in individual experiments. For every combination, the experiment was repeated twice. Statistical tests were performed by using Statgraphics version 4.2 software (StatPoint, Englewood Cliffs, NJ, USA).

Glycosylation of Sand Fly Midguts
Midguts were dissected from 5- to 10-day-old P. sergenti and P. arabicus females. Midgut proteins were separated by sodium dodecyl sulfate–polyacrylamide gel electrophoresis on 10% gels under reducing conditions in a Mini-Protean III apparatus (Bio-Rad Laboratories, Hercules, CA, USA) at 200 V. Gels were stained with Coomassie brilliant blue R-250 or transferred to nitrocellulose membranes by using a Semiphor unit (Hoefer Scientific Instruments, San Francisco, CA, USA). Western blotting was performed for 90 minutes at 1.5 mA/cm². Membranes were incubated with 20 mmol/L Tris, 150 mmol/L NaCl, 0.05% Tween (TBS-Tw) with 5% bovine serum albumin for 2 hours and then with Helix pomatia agglutinin (HPA) biotinylated lectin, which recognizes N-acetyl-D-galactosamine (GalNAc), a typical carbohydrate in O-glycans. In the control groups, HPA reactions were competitively inhibited by preincubation with 250 mmol/L GalNAc for 30 minutes. After repeated washing in TBS-Tw, blots were incubated for 1 hour with streptavidin peroxidase in TBS-Tw. The peroxidase reaction was developed with the substrate 4-chloro-1-naphthol. All chemicals for lectin blotting were obtained from Sigma.

Random Amplified Polymorphic DNA Analysis
Twenty wild-caught sand flies morphologically identified as P. sergenti, 10 from the northern focus and 10 from the southern focus, were included in the analysis. Two flies from Tulek, Turkey, were included as an outgroup. DNA from thoraxes was extracted by using the High Pure PCR template preparation kit (Roche, Paris, France). Five decamer random primers (OPD5, OPE4, OP11, OP114, and OP118; Operon Technologies Inc, Alameda, CA, USA) were used. The reaction mixture contained 12.5 μL master mixture (75 mmol/L Tris·HCl, pH 8.8, 20 mmol/L (NH₄)₂SO₄, 0.001% Tween 20, 800 μM deoxynucleotide
DNA samples for RAPD analysis were used for ITS2 sequencing. One specimen from each study area was included as previously described (25).

Results

Sand Fly Species

A total of 1,491 sand flies (7 species, 4 subgenera) from the northern focus and 876 sand flies (7 species, 4 subgenera) from the southern focus were identified. Phlebotomine fauna in the southern focus were relatively species poor with P. (Paraphlebotomus) sergenti comprising >90% of the flies. The most striking difference in the species composition between the foci was the absence of P. (Adlerius) arabicus and P. (Adlerius) simici from the southern focus, both of which were prominent species in the south-facing slopes of the northern focus (Table 1) (26).

Leishmania Infections in Sand Flies

To detect infections and obtain parasite isolates, sand fly females were dissected in sterile saline and guts were examined microscopically. Four (6.6%) of 61 P. arabicus and 1 (0.8%) of 125 P. sergenti from the northern focus had promastigotes in their guts. Infection intensity in P. sergenti from the northern focus was low, but all P. arabicus had heavy, mature infections. A total of 213 flies from the southern focus were dissected; 196 were P. sergenti, and 19 (9.7%) had promastigotes. Eleven of these females had heavy infections, and 8 had moderate-to-light infections. All infected females were caught at 1 sublocality in the southern focus, where the local infection rate was 19.6%. Promastigote cultures were established from 4 P. arabicus and 1 P. sergenti captured in the northern focus and from 18 P. sergenti females captured in the southern focus (Table 2). None of the other sand fly species were infected.

Sand flies that were not dissected fresh were kept frozen and were subjected to ITS1 PCR for detection of Leishmania. Nine (18%) of 50 P. sergenti females from the southern focus were positive for Leishmania ribosomal DNA. HaeIII digestion of the ITS1 PCR products confirmed that all P. sergenti had L. tropica (Figure 2).

Identification of Infections in Mammals

Rodents collected in the northern focus were tested for L. tropica infection by ITS1 PCR. Dried blood and skin samples from 28 rats (R. rattus) and 46 spiny mice (A. cahirinus) were negative for Leishmania DNA. Eight of 73 rock hyraxes from the northern focus and 6 of 46 rock hyraxes from the southern focus were positive for L. tropica DNA by ITS-1 amplification and reverse-line blotting using sequence-specific probes (data not shown). Of the positive animals, 11 were adults (9 females and 2 males) and 1 was a juvenile male. Parasites from 1 rock hyrax captured in the northern focus were cultured and identified by ITS1 PCR and digestion with HaeIII (Figure 3).

Antigenic Characterization of Leishmania Isolates

IFA assays with species-specific MAbs were used to characterize different isolates. L. tropica isolates from the northern focus were antigenically distinct from all other isolates, including those from the southern focus (Table 3).

Susceptibility of P. arabicus and P. sergenti to L. tropica

In laboratory experiments, L. tropica parasites from the northern focus infected only P. arabicus, and parasites from the southern focus infected both P. arabicus and P. sergenti. Susceptibility of P. arabicus for infection with L. tropica strains from both northern and southern foci was high (94% and 97%, respectively). In contrast, P. sergenti was not permissive for L. tropica strains from the northern focus (1 of 64 flies). Susceptibility of P. sergenti for

| Table 1. Phlebotomus sand fly species in the Galilee foci, northern Israel* |
|----------------------------------|------------------|------------------|------------------|
| Species                          | Northern focus   | Southern focus   |
|                                  | No. (%) females  | No. (%) males    | No. (%) females  | No. (%) males    |
| P. (Adlerius) arabicus           | 62 (15)          | 234 (22)         | –                | –                |
| P. (Adlerius) simici             | 35 (9)           | 118 (11)         | –                | –                |
| P. (Paraphlebotomus) sergenti    | 131 (32)         | 317 (29)         | 267 (91)         | 532 (92)         |
| P. (Larroussius) tobbi           | 167 (40)         | 337 (31)         | 11 (4)           | 23 (4)           |
| P. (Larroussius) syriacus        | –                | 12 (1)           | 8 (3)            | 16 (2)           |
| P. (Larroussius) perfluswii      | 10 (2)           | 31 (3)           | 1 (<1)           | 2 (<1)           |
| P. (Phlebotomus) papatasi        | 9 (2)            | 28 (3)           | 5 (2)            | 7 (1)            |
| Total                            | 414              | 1,077            | 292              | 580              |

*Species comprising <1% of the fauna (1 P. [Paraphlebotomus] alexandri and 3 P. [Adlerius] halopennis) found in the southern focus were not included.
infection with \textit{L. tropica} from the southern focus strain was lower (66\%) than that of \textit{P. arabicus} (Figure 4).

**Glycosylation of Luminal Midgut Proteins**

Incubation of \textit{P. sergenti} midgut lysates with HPA showed no reaction, indicating a lack of O-glycosylated proteins (Figure 5). In contrast, an abundant glycoprotein (37–43 kDa) was strongly labeled by HPA in \textit{P. arabicus} midgut lysates. Controls of \textit{P. arabicus} midgut lysates incubated with HPA blocked by preincubation with GalNAc showed no reaction, which confirmed the specificity of the lectin reactions in experimental blots (Figure 5). Labeling of midguts with fluorescein-conjugated HPA confirmed the presence of GalNAc-containing glycoproteins in the midguts of \textit{P. arabicus}. Intensity of labeling in \textit{P. sergenti} midguts was weaker, which reflected a nonspecific background reaction (Figure 5).

**Comparison of \textit{P. sergenti} Populations by RAPD and ITS2 Sequencing**

Flies from both foci shared the same banding pattern and differed from Turkish \textit{P. sergenti} (Figure 6). ITS 2 sequences of \textit{P. sergenti} from both foci were identical with each other and nearly identical (99\%) with the ITS 2 sequence of a \textit{P. sergenti} specimen from the West Bank (GenBank accession no. AF462325) (data not shown).

**Discussion**

We have identified 2 emerging foci of CL in which rock hyraxes serve as reservoir hosts of the causative agent \textit{L. tropica}. Despite their geographic proximity, the 2 foci show fundamental differences with regard to transmission cycles. Parasites and vector species in the southern focus are typical of most Asian zoonotic \textit{L. tropica} foci, but the northern focus is characterized by antigenically distinct parasites that are transmitted by a newly incriminated sand fly vector.

\textit{L. tropica} is widely distributed in eastern and northern Africa, the Middle East, and large parts of Asia. A recent study using 21 microsatellite loci showed that \textit{L. tropica} is a genetically heterogeneous species composed of >80 genotypes. The genetic makeup of this complex suggests a probable African origin, with isolates from the northern focus more related to African isolates than to other strains from the Middle East (16).

The major surface molecule of \textit{Leishmania} promastigotes is LPG, which has been shown to mediate attachment of parasites to the midgut of the sand fly (8). LPG of \textit{L. tropica} from the northern focus is characterized by abundant terminal \(\beta\)-galactose residues on side chains. Conversely, \(\beta\)-galactose residues on LPG side chains of other \textit{L. tropica} isolates are mostly capped with glucose (27). Differences in sugar moieties may have a role in infection of \textit{P. sergenti} (Figure 4). Although \(\beta\)-galactose residues are present in \textit{L. major} LPG, strains of \textit{L. tropica} from the northern focus were not infective to \textit{P. papatasi}, the natural vector of \textit{L. major} (M. Svobodova, unpub. data) (4).

\textit{P. sergenti} is probably a species complex, and its component populations show several molecular and morphologic differences (25). RAPD-PCR is a powerful tool for estimating genetic variability and was successfully used to compare genetic variation within and between 5 sympatric \textit{Phlebotomus} species in Spain (28). Using the same primer sets, we did not find any differences between \textit{P. sergenti} flies from the 2 foci (Figure 6). We deduce that populations from both foci are probably freely interbreeding.

![Figure 2](https://example.com/figure2.png)
\textit{P. sergenti} is of Palearctic origin; flies migrated into North Africa during the Miocene era (29). Thus, \textit{L. tropica} and \textit{P. sergenti} apparently originated in different continents and their geographic overlap probably arose at a later time. \textit{P. sergenti}, \textit{P. (Larroussius) guggisbergi}, \textit{P. (Paraphlebotomus) saevus}, and perhaps \textit{P. arabicus} are vectors in Africa (30,31). Since \textit{L. tropica} variants from both foci develop in \textit{P. arabicus}, but only the variant from the southern focus completes development in \textit{P. sergenti}, we postulate that \textit{L. tropica} was initially transmitted by \textit{P. arabicus} or another permissive vector such as \textit{P. (Adlerius) halepensis} (9). The more common transmission cycle is a later adaptation to \textit{P. sergenti}, a dominant, widely distributed phlebotomine species.

Refractoriness of \textit{P. sergenti} to variants of \textit{L. tropica} from the northern focus is probably due to the lack of HPA-binding proteins on the luminal surface of midgut epithelium. HPA-binding epitopes are present in permissive vectors such as \textit{P. arabicus} (Figure 4), \textit{P. halepensis} (P. Volf, unpub. data), and \textit{Lu. longipalpis} (32). These findings support infections with multiple species of \textit{Leishmania} (9,33).

The absence of \textit{P. arabicus} from the north-facing slopes of the southern foci contrasts dramatically with its predominance in the south-facing slopes of the northern focus. Although a satisfactory explanation for this fact is lacking, such phenomena are not unusual. For example, species richness of insects was much higher in the drier and warmer south-facing slopes of a narrow canyon (100–400 m wide) in Mount Carmel, Israel, than in the north-facing slope of the same canyon (34). \textit{P. arabicus} is

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**Figure 3.** A) PCR of \textit{Leishmania} internal transcribed spacer region 1 (ITS1) of cultured \textit{Leishmania} promastigotes isolated from rock hyrax. B) \textit{HaeIII} digestion of restriction fragment length polymorphisms of ITS1 PCR products shown in A. Lane MW, molecular mass marker; lane 1, \textit{L. infantum} (Li-L699); lane 2, \textit{L. major} (Lm-L777); lane 3, \textit{L. tropica} (Lt-L590); lane 4, rock hyrax.

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**Figure 4.** Artificial infection of laboratory-reared \textit{Phlebotomus arabicus} and \textit{P. sergenti} with \textit{Leishmania tropica} isolates from 2 foci in Galilee, Israel. Note the high susceptibility of \textit{P. arabicus} for both strains and refractoriness of \textit{P. sergenti} for the northern strain. Nf, northern focus; Sf, southern focus.

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**Table 3.** Characterization of \textit{Leishmania tropica} isolates from the Galilee foci, northern Israel*

| Focus/source          | Monoclonal antibody specificity | Excreted factor serotype |
|----------------------|---------------------------------|--------------------------|
|                      | \textit{L. major} T1 | \textit{L. major}/\textit{L. tropica} T3 | \textit{L. tropica} T11 |                   |
| Northern             |                                |                          |                          |                   |
| \textit{Phlebotomus arabicus} | 5+                             | 5+                       | ±                        | A4                |
| Girl with CL†        | 4+                             | 5+                       | –                        | A4                |
| Rock hyrax           | 5+                             | 5+                       | ±                        | A4                |
| Southern             |                                |                          |                          |                   |
| \textit{P. sergenti} | –                              | 4+                       | 3+                       | A9B2              |
| Man with CL†         | –                              | 2+                       | 4+                       | A9B2              |
| Reference strains    |                                |                          |                          |                   |
| \textit{L. major}    | 5+                             | 5+                       | –                        | A1                |
| \textit{L. tropica}  | ±                              | 3+                       | 3+                       | A9                |

*Characterization was performed by using excreted factor serotyping (23) and species-specific monoclonal antibodies (14). CL, cutaneous leishmaniasis. Values indicate relative intensity of fluorescence under UV light. \textit{L. tropica} isolates from the northern focus were antigenically similar to \textit{L. major} and distinct from other \textit{L. tropica} strains.

†Specimens were isolated by skin scraping for diagnostic purposes at the Department of Dermatology at Hadassah Hospital, Jerusalem.
widely distributed in Africa and the Arabian peninsula (17), and the Galilee focus forms the northern limit of its distribution. Since P. arabicus originates in warmer regions, finding it in warmer, drier, south-facing slopes and not in cooler, shadier north-facing slopes of the hills in Galilee is not surprising (Table 1).

Rock hyraxes in both foci were found infected with L. tropica, and 1 isolate was obtained from an adult male in the northern focus. Although rock hyraxes were suspected reservoir hosts of L. tropica in Africa (35,36) and have been previously implicated in the northern focus (4), this is the first report of a rock hyrax isolate that was identified as L. tropica and shown to be identical to those obtained from humans and sand flies in the same focus (Table 3).

Rock hyrax populations in many parts of Israel are expanding rapidly and encroaching upon human habitation. They were extremely common in both foci studied, as well as in other L. tropica foci in the region (D. Meir and A. Warburg, unpub. data; [4,5]). In the Galilee foci, rock hyraxes inhabit crevices within boulder mounds that were created when land was cleared for the construction of houses. These artificial caves also afford suitable breeding sites for sand flies. Rock hyraxes are susceptible to L. tropica, and infected rock hyraxes are infective to feeding P. arabicus and P. sergenti. Sand flies are attract-
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