



LeishMan: a European consortium for the harmonisation of diagnosis, parasite typing and treatment of patients with leishmaniasis

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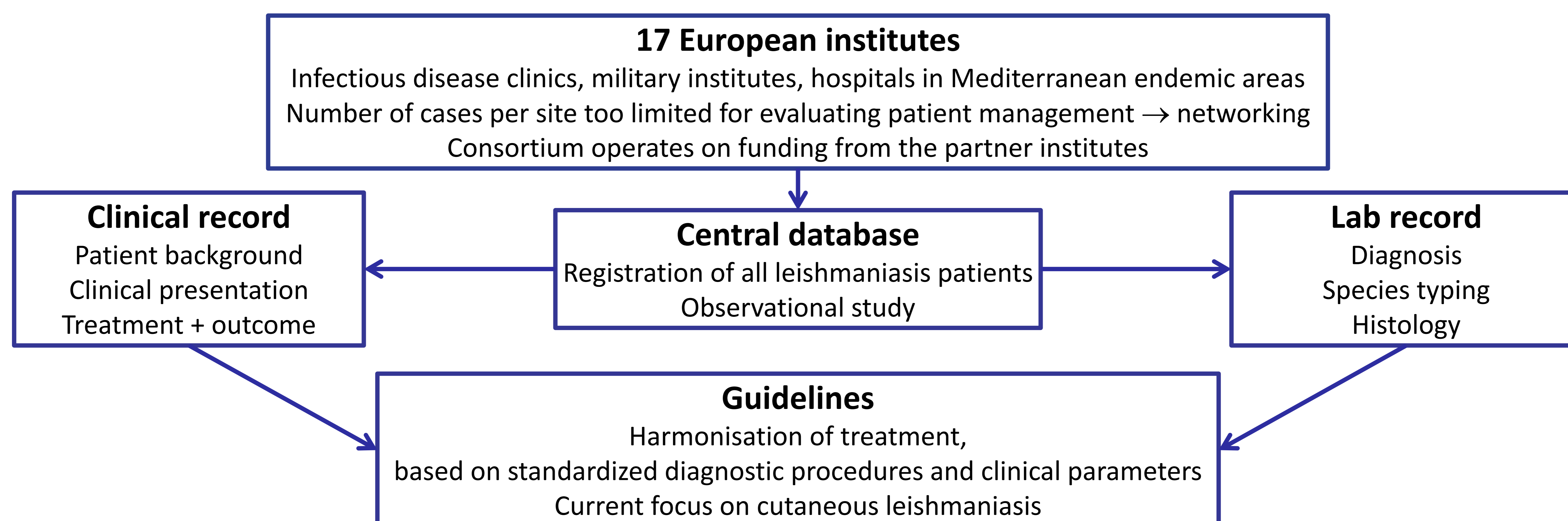
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Current output / related abstracts

Meetings: twice a year

Publications:

- 1) Blum et al. (2012) Local or systemic treatment for New World cutaneous leishmaniasis? Re-evaluating the evidence for the risk of mucosal leishmaniasis. *International Health* 4; 153–163
- 2) Leishman: Developing evidence based clinical recommendations for the treatment of cutaneous leishmaniasis in Europe (Abstract 616)
- 3) Systematic comparison of five genotypic markers for species discrimination in *Leishmania* (Abstract 324)

Joining the network

- 1) The LeishMan network is open for additional members who can actively contribute to the database in terms of patient and diagnostic data.
- 2) European institutes dealing with travel medicine, as well as those seeing patients in endemic areas of southern Europe are welcomed.
- 3) Institutes wanting to join should be able to perform their own molecular diagnosis.
- 4) Interested parties can contact the coordinator Dr. Johannes Blum (johannes.blum@unibas.ch)

Contacts

Network coordinator: Dr. Johannes Blum (johannes.blum@unibas.ch)

Members present @ WorldLeish5:

Names in red on the network map

Full member list: see www.leishman.eu

