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Seroprevalence of five neglected parasitic diseases among immigrants accessing five infectious and tropical diseases units in Italy: a cross-sectional study

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112	Abstract
113	Objective: This multicentre cross-sectional study aims to estimate the prevalence of five
114	neglected tropical diseases (Chagas disease, filariasis, schistosomiasis, strongyloidiasis,
115	toxocariasis) among immigrants accessing health care facilities in five Italian cities
116	(Bologna, Brescia, Florence, Rome, Verona).
117	Methods: Individuals underwent a different set of serological tests, according to country of
118	origin and presence of eosinophilia. Seropositive patients were treated and further
119	followed up.
120	Results: A total of 930 adult immigrants were enrolled: 477 men (51.3%), 445 women
121	(47.9%), 8 transgender (0.8%); median age was 37.81 years (range 18-80). Most of them
122	were coming from the African continent (405/930, 43.5%), the rest from East Europe,
123	South America and Asia. A portion of 9.6% (89/930) were diagnosed with at least one of
124	the infections under study. Seroprevalence of each specific infection varied from 3.9%
125	(7/180) for Chagas diseases to 9.7% (11/113) for toxocariasis. Seropositive people were
126	more likely to be 35 to 40 years-old male and to come from South East Asia, Sub-Saharan
127	Africa or South America.
128	Conclusions: The results of our study confirm that neglected tropical diseases represent a
129	substantial health problem among immigrants and highlight the need for addressing this
130	emerging public health issue.
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133	Introduction
134	The World Health Organization (WHO) defines the neglected tropical diseases (NTDs) as
135	a diverse group of infections mainly affecting poor populations, increasing poverty, and
136	having a low priority in the political and scientific agenda [1].

137	Human migration is a key factor in the appearance/re-appearance of NTDs in non or
138	former endemic contexts [2] [3]. We live in an era of unprecedented human mobility, with
139	approximately 232 million international migrants and 740 million internal migrants
140	worldwide [4]: Eurostat estimated that a total of 3.4 million people migrated to one of the
141	European Community countries in 2013; half of them came from non-member countries
142	[5]. In 2014, immigrants accounted for the eight per cent of the total Italian resident
143	population [6]; another 326,000 undocumented immigrants and refugees were present in
144	Italy. [7].
145	Immigrants are generally young and in good health conditions [8], nevertheless, the
146	prevalence of some infectious diseases may be significant among immigrants, as a result
147	of the wide diffusion of these conditions in their countries of origin [9] and the further
148	exposure during migration [10] [3]. Many infections, including several NTDs, may be
149	asymptomatic and hence remain undiagnosed [11]. As a consequence, seropositive
150	individuals can develop chronic forms (e.g. Chagas disease, schistosomiasis), fatal
151	complications (e. g. Chagas disease, schistosomiasis or strongyloidiasis) and can
152	potentially transmit the disease [11].
153	The research on the burden of communicable diseases among immigrants in Western
154	countries mainly focuses on HIV, tuberculosis and viral hepatitis [8] [12]. NTDs were rarely
155	addressed, possibly because they are often asymptomatic and have a relatively low
156	transmission in the absence of environmental and biological reservoirs/vectors. Moreover
157	in most of the European countries there is no systematic mandatory regulation regarding
158	NTDs reporting and surveillance [13]. Given the considerable immigration flows to Italy
159	and the scarce information on the relevance of the NTDs, seldom considered at hospital
160	level, this study aims to estimate the seroprevalence of five NTDs (Chagas disease,
161	filariasis, schistosomiasis, strongyloidiasis and toxocariasis) among immigrants attending

162 hospitals and relative outpatient clinics in five Italian cities: Bologna, Brescia, Florence, 163 Rome, and Verona.

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Methods

Study population, data collection and patient management

A cross-sectional survey was performed in five Italian infectious and tropical diseases units located in five different hospitals (Bologna University Teaching Hospital; Florence University Teaching Hospital; Hospital Sacro Cuore - Negrar, Verona; Spedali Civili General Hospital, Brescia; L. Spallanzani University Teaching Hospital, Rome) and in one outpatient clinic for undocumented immigrants (Brescia Local Health Authority outpatient clinic). In the hospital setting, patients were usually referred from primary care, Emergency Departments or other secondary care services; they were either inpatients or individuals with chronic infections followed-up in specialised outpatient clinics. Individuals who attended any of the above mentioned centres for any reason in the study period (November 2012 to November 2014) and who were born in an endemic country (see online appendix Annex 1 for details), older than 18 years and with sufficient knowledge of Italian or a timely access to a linguistic mediator, were eligible. In each clinical centre one or two investigators were responsible for offering participation to the project to each eligible patient seeking care during any of their routine clinical activities. After signing the informed consent, enrolled patients underwent a different set of serological tests according to the criteria reported in online appendix Table 1. The choice of the infections was based on the most common areas of origin of immigrants in Italy, the potential severity of the disease if not treated, the availability and quality of diagnostic tools, the amenability to treatment and the potential for spreading in the community. The definition of endemic country for a certain infection was based on the WHO geographical

188	classification of NTDs [14] (see online appendix Annex 1 for details). For filariasis, only
189	endemic countries for lymphatic filariasis, onchocerciasis and loiasis were included.
190	Serology for toxocariasis and filiariasis was limited to individuals with eosinophilia These
191	two diseases usually present with a raised eosinophil count [15]. However, in order to
192	increase case detection, eosinophil cut-off level was set at 300/ μ L instead of 450-500/ μ L,
193	as routinely suggested [15], for its good positive predictive value for helminthiases [16].
194	Concurrently, clinical and socio-demographic information, including country of origin, list of
195	visited countries, time since arrival, and educational level were collected. The investigators
196	offered treatment and follow-up to seropositive patients, while they supplied seronegative
197	individuals with the results of their tests. The centres elaborated operational guidelines for
198	the management of each disease which were made available on the study website.
199	Study protocol was approved by the ethics committee of the coordinating site (Bologna
200	University Teaching Hospital) under the resolution number 124/2012/O/Oss and by those
201	of all other participating units.

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Microbiological diagnosis procedures

A sample of 12 ml of venous blood was collected from each participant. Blood samples were centrally tested at the Service of Epidemiology and Laboratory for Tropical Diseases of the Hospital Sacro Cuore - Don Calabria, Negrar in order to reduce inter-laboratory variability. Serum samples were tested for specific antibodies using commercial immunoenzymatic assays according to manufacturer's instructions. The qualitative presence of antibodies for Trypanosoma cruzi (etiological agent of Chagas disease) was tested employing two enzyme-linked immunosorbent assays (ELISA), one based on recombinant antigens ("BioELISA Chagas", Biokit, Lliça d'Almunt, Spain), the other based on crude antigens ("BioELISA Chagas III", BiosChile, Santiago, Chile). For the other infections a single ELISA was used ("Filariasis ELISA kit", Bordier Affinity Products SA,

214	Crissier, Switzerland, for filariasis; "Schistosoma mansoni ELISA kit", Bordier Affinity
215	Products SA, Crissier, Switzerland for schistosomiasis; "Strongyloidiasis ELISA kit" based
216	on Strongyloides ratti antigens, Bordier Affinity Products SA, Crissier, Switzerland for
217	stronglyloides; "DRG Toxocara canis ELISA", DRG Instruments GmbH, Marburg,
218	Germany, for toxocariasis).
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220	Statistical analysis
221	Categorical variables were described through frequencies and the median and the ranges
222	were used to describe age. Countries of origin were subsequently grouped into 11 regions
223	following the Geosentinel classification [17]. This choice relies on the fact that Geosentinel
224	system splits the globe into a higher number of regions (eleven) than WHO (six), with more
225	precise identification of risk areas.
226	Prevalence point estimates and their 95% confidence intervals were obtained. Chi-square
227	tests were performed to assess differences between groups. Data were managed and
228	analysed using STATA 14.1.
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231	Results
232	Description of the study population
233	From November 2012 to November 2014 a total of 930 individuals were enrolled across
234	the six centres. Two thirds of them were outpatients. Due to refused consent or scarce
235	knowledge of Italian/lack of linguistic mediator, 4.9% (48/978) of the individuals were not
	enrolled.
236	
236 237	Socio-demographic information of the enrolled population is summarized in online
	Socio-demographic information of the enrolled population is summarized in online appendix Table 2. The male-to-female ratio was 1:1, and the median age was 37.8 years

240	years. Individuals coming from the African region represented 43.5% of the total (405/930)
241	other frequent areas of origin were Eastern Europe (197/930, 21.2%), South and Centra
242	America (177/930, 19.0%) and Asia (142/930, 15.3%). More than a half of patients
243	declared a medium or high level of education (high school diploma or degree). The socio-
244	demographic profile of the individuals varied slightly across the six centres. In the clinic for
245	undocumented immigrants in Brescia, enrolled subjects were younger than the total
246	population (median age of 35.2 versus 37.8 years, age range of 18-64 versus 18-80) and
247	their time since arrival was slightly shorter (50.9%, 56/110, of them arrived in the last four
248	years versus 31.3%, 290/930, in the total). Differences across the centres in terms of
249	origin might mirror the differing immigrant flows to the Italian cities: despite the high
250	presence of African immigrants in the whole sample, individuals enrolled in the Roman
251	hospital were mainly South Americans and the ones enrolled in Bologna mainly came from
252	Eastern Europe.
253	A white blood cell count was available for 583 individuals: among them, 19.4% (113) had
254	eosinophilia.

Seroprevalence of the neglected infectious diseases

Among the 930 enrolled individuals, 96 new infections were detected: 42 cases of strongyloidiasis, 31 of schistosomiasis, 11 of toxocariasis, 7 of Chagas disease, and 5 of filariasis. Eighty-nine patients were diagnosed with one or more NTDs, which leads to an overall seroprevalence of 9.6% (95%Cl 7.8-11.6) in the study population. Seven individuals had two infections simultaneously. Across the centres the prevalence varied between 6.3%, 7/110, (in Brescia clinic for undocumented immigrants) and 15.3%, 30/193,

(in Verona).

> Seropositive individuals were mostly men (M:F=2:1) with a median age of 38.8 years (range 21-78). The seroprevalence was twice as high in men as in women (p-value<0.05)

266 for all infections except for Chagas disease. The Geosentinel region with the highest NTDs 267 prevalence was South East Asia, followed by Sub-Saharan Africa and South America. 268 Among the 189 patients who were known to be HIV positive, 14 (7.4%) were also 269 seropositive for at least one of the NTDs under study (8 cases of strongyloidiasis, 4 cases 270 of schistosomiasis, 2 cases of toxocariasis and 1 case of Chagas disease). 271 Global seroprevalence, women to men ratio and regions with highest prevalence are 272 shown in Table 1. Detailed prevalence estimated by infections and Geosentinel regions 273 are listed in online appendix Table 3.

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Discussion

Approximately one out of 10 individuals in our study was seropositive for at least one of the infections. This figure represents a considerable burden given the potential consequences of these conditions. In particular, strongyloidiasis and Chagas disease can lead to chronic infections, which might represent a serious threat for the individual and the health systems [18]. Strongyloidiasis is responsible of the hyperinfection syndrome, a rare life-threatening complication that mainly affects immunosuppressed individuals, thus early detection and treatment are particularly relevant [19]. Similarly, the potential transmissibility of Chagas disease outside endemic areas, through blood transfusions, organ or tissue transplants, or mother-to-child, highlights the importance of its early detection [20]. Furthermore, most of these infections are treatable with affordable and generally well-tolerated therapies, especially when compared to the severity of the untreated consequences [21]. Interestingly, we noticed a higher prevalence of the five neglected infections among patients coming from South East Asia, Sub-Saharan Africa and South America, suggesting that immigrants coming from these areas are most at risk. However, this broad geographic subdivision may mask differences at country level which we were not able to account for,

292	given the limited sample size. The prevalence of all infections was twice as high in men as
293	in women (p-value<0.05) but Chagas disease. A potential higher environmental and
294	working exposure risk for intestinal parasites and other vector-borne infections among
295	male immigrants can contribute to explain this finding [22]. Age and time since arrival in
296	Italy were not associated with the presence of infections.
297	These results are in line with the findings of similar studies carried out in Spain [8] [23],
298	except for Chagas disease prevalence, which was much higher in the Spanish samples.
299	This difference may be explained by the larger proportion of enrolled Latin American
300	subjects in the Spanish studies, as a consequence of a different migration pattern and the
301	availability of widespread screening programs. A cross sectional study carried out in
302	Australia [11] among recent immigrants and two others conducted in the United States in
303	refugees [9] [24] reported a higher prevalence of intestinal parasitic infections than in our
304	study. These figures may be explained by the different population under study and by our
305	diagnostic approach based on antibody detection. The mentioned studies mainly enrolled
306	refugees and recent immigrants and used microscopic examination of the stools,
307	identifying also parasites for which no serological test is available.
308	As already reported by others [25], we noticed a significant seroprevalence of NTDs in the
309	subgroup of patients with a known HIV infection. In this subgroup of patients treatment
310	should be strongly recommended, because of the risk of severe complications especially
311	in the case of strongyloidiasis and Chagas disease [25].
312	In our study, the proportion of individuals with an increased eosinophil count was in line
313	with other studies [26] [15], despite a possible overestimation due to the lower cut-off level
314	for an abnormal eosinophil count. Eosinophilia generally occurs in approximately 10% of
315	individuals returning from the tropics [15], and a prevalence up to 30% is often reported
316	among immigrant populations [26]. In this last group helminthic infections are the
317	commonest identifiable cause of eosinophilia, accounting for 14% to 64% of the total

318	cases [15]. Among those who were screened for strongyloidiasis, positive patients were
319	more likely to have a high eosinophil count (data not shown). This result confirms what
320	was previously found and emphasizes the importance of investigating the presence of S.
321	stercoralis among immigrants, particularly in presence of raised eosinophil levels [15].
322	Whilst for Chagas disease and schistosomiasis serology is deemed to be a valid screening
323	tool [8], there is no standard method for the detection of other intestinal parasites [27].
324	When stool microscopy had been used for screening purpose, the prevalence was found
325	to be relatively low [8] [23]. Amongst intestinal parasites we focused on strongyloidiasis,
326	because of its potentially fatal complications in immunosuppressed populations and to its
327	long-term persistence in the host [28].
328	The main limitation of the study is its generalizability. The enrolment took place at hospital
329	level and not at the community level; this means that our results may not be entirely valid
330	for the general immigrant population. Individuals who access the health services may be
331	those who have been in the host country for a longer time; they may differ from the general
332	population in terms of socio-demographic characteristics and, therefore, may have a
333	different risk profile for the infections. Moreover, no formal sample size calculation was
334	performed; some prevalence estimates on specific infections and subpopulations showed
335	a great uncertainty due to the small sample size of these groups.
336	Additionally, a selections bias cannot be ruled out. Enrolment in the study completely
337	relied on the staff dedicated to the project. It is likely that a proportion of patients eligible
338	for the study had been missed; however, they should not have been systematically
339	different from those enrolled because the physicians responsible for the enrolment
340	covered different care settings within their units. Moreover, not all the patients who were
341	asked to take part in the study gave their consent. The proportion of patients who did not
342	participate despite their eligibility was less than five percent in all the centres suggesting
343	that this source of bias is not substantial.

344	Another limitation of the study lies in the exclusive use of serology to estimate the
345	prevalence of selected NTDs in immigrants. As a matter of facts, these tests may not
346	distinguish between prior infection and active disease since antibodies may persist for
347	many months to years after successful treatment in most of the NTDs evaluated and these
348	tests are prone to cross-reactions with other parasite antigens [29]. Additional tests on
349	stool or urine samples would have certainly increased the diagnostic sensitivity but were
350	deemed not be feasible, given the logistic arrangement of the study that relied on a
351	centralised laboratory. Since the therapies for the infections under study are mostly short-
352	term and well-tolerated, we opted for treating all seropositive patients, except those
353	affected by T. cruzi infection, who underwent disease staging before treatment according
354	to current agreements [30].
355	Despite these limitations, our findings highlight the importance of tackling the NTDs
356	challenge in a non-endemic setting. The call for systematic detection and appropriate
357	management is even more urgent because it has been reported that health professionals
358	rarely consider these diseases. As a consequence, NTDs are highly likely to be
359	underdiagnosed at present, or diagnosed too late or inefficiently managed [31]. As
360	previously suggested in the European context, screening protocols seem to be a sensible
361	option [32]. A presumptive anthelminthic therapy for immigrants coming from areas at high
362	risk had been previously demonstrated to be cost effective in certain setting [27]. However,
363	this approach is not free from drawbacks, including toxicity, under-treatment of certain
364	infections [33] and risk of focusing on a single medical intervention while neglecting a
365	proper follow-up and a more comprehensive approach to migrants' health.
366	The importance of diagnosing and treating these infections is crucial among
367	immunosuppressed patients (for example those receiving chemotherapy, chronic steroid
368	or immunosuppressive treatment) and donors/recipients of solid organ and hematopoietic
369	stem cell transplantation, as well as blood transfusion [34]. Indeed, the rising success and

370	adop	tion of transplantation reasonably increases the proportion of the immigrant					
371	popu	lation who will become donor/recipient of organ transplantation and					
372	blood	blood/hemoderivates in destination countries. Many of the pathogens that cause NTDs car					
373	be e	ither reactivated during immunosuppression or transmitted via organ graft or blood					
374	trans	fusions [34], making a screening approach in these contexts life-saving.					
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378	Wed	declare that we have no conflicts of interests.					
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384	Refe	rences					
385							
386	[1]	WHO. First WHO report on neglected tropical diseases: working to overcome the					
387		global impact of neglected tropical diseases [Internet]. World Health Organization.					
388		2010. Available from:					
389		http://apps.who.int/iris/bitstream/10665/44440/1/9789241564090_eng.pdf?ua=1					
390	[2]	Aagaard-Hansen J, Nombela N, Alvar J. Population movement: A key factor in the					
391		epidemiology of neglected tropical diseases. <i>Trop Med Int Heal</i> . 2010;15:1281–8.					
392	[3]	Smith Darr J, Conn DB. Importation and Transmission of Parasitic and Other					
393		Infectious Diseases Associated with International Adoptees and Refugees					
394		Immigrating into the United States of America. <i>Biomed Res Int.</i> 2015;2015:763715.					
395	[4]	International Organization for Migration. World Migration Report: Migrants and					
396		Cities: New Partnerships to Manage Mobility [Internet]. 2015. 204 p. Available from:					
397		http://publications.iom.int/system/files/wmr2015_en.pdf					
398	[5]	Eurostat F. Migration and Migrant Population Statistics. <i>Eurostat</i> [Internet].					
399		2014;2012:1–13. Available from:					
400		http://epp.eurostat.ec.europa.eu/statistics_explained/index.php/Migration_and_migr					
401		ant_population_statistics#Foreign_and_foreign-born_population					

402	[6]	Perego G, Soddu F. XXIV Rapporto Immigrazione 2014 [Internet]. 2014. Available
403		from:
404		http://www.caritasitaliana.it/materiali//Pubblicazioni/libri_2015/Rapporto_Immigrazio
405		ne/sintesi_rapportoimmigrazione2014_quadroregionale.pdf
406	[7]	Blangiardo C. Stima delle presenze irregolari. Vari anni. ISMU. Vol. 2010. 2012.
407	[8]	Monge-Maillo B, Lòpez-Vélez R, Norman FF, Ferrere-Gonzàlez F, Martìnez-Pérez
408		A, Pérez-Molina JA. Screening of imported infectious diseases among
409		asymptomatic Sub-Saharan African and Latin American immigrants: A public health
410		challenge. Am J Trop Med Hyg. 2015;92:848–56.
411	[9]	Lifson AR, Thai D, O'Fallon A, Mills WA, Hang K. Prevalence of tuberculosis,
412		hepatitis B virus, and intestinal parasitic infections among refugees to Minnesota.
413		Public Health Rep. 2002;117:69–77.
414	[10]	Zammarchi L, Bartalesi F, Bartoloni A. Tuberculosis in tropical areas and
415		immigrants. Mediterr J Hematol Infect Dis. 2014;6:e2014043.
416	[11]	Caruana SR, Kelly HA, Ngeow JYY, Ryan NJ, Bennett CM, Chea L, et al.
417		Undiagnosed and potentially lethal parasite infections among immigrants and
418		refugees in Australia. J Travel Med. 2006;13:233-9.
419	[12]	Hargreaves S, Seedat F, Car J, Escombe R, Hasan S, Eliahoo J, et al. Screening
420		for latent TB, HIV, and hepatitis B/C in new migrants in a high prevalence area of
421		London, UK: a cross-sectional study. BMC Infect Dis. 2014;14:657.
422	[13]	ECDC. Annual epidemiological reports 2016.pdf [Internet]. 2016. Available from:
423		http://ecdc.europa.eu/en/publications/surveillance_reports/annual_epidemiological_i
424		eport/Pages/epi_index.aspx
425	[14]	WHO. Neglected tropical diseases [Internet]. Available from:
426		http://www.who.int/neglected_diseases/resources/en/
427	[15]	Checkley AM, Chiodini PL, Dockrell DH, Bates I, Thwaites GE, Booth HL, et al.
428		Eosinophilia in returning travellers and migrants from the tropics: UK
429		recommendations for investigation and initial management. J Infect. 2010;60:1-20.
430	[16]	Bisoffi Z, A C, F B. Eosinofilia in patologia di importazione: criteri diagnostici e
431		terapeutici. G Ital di Mal Infett. 1995;1:293-8.
432	[17]	Harvey K, Esposito DH, Han P, Kozarsky P, Freedman DO, Plier DA, et al.
433		Surveillance for travel-related disease-GeoSentinel Surveillance System, United
434		States, 1997-2011. Morb Mortal Wkly report Surveill Summ. 2013;62:1–23.
435	[18]	WHO. Investing To Overcome the Global Impact of Neglected Tropical Diseases,

436		Third WHO report on neglected tropical diseases [Internet]. 2015. Available from:
437		http://www.who.int/neglected_diseases
438	[19]	Buonfrate D, Requena-Mendez A, Angheben A, Munoz J, Gobbi F, Van Den Ende
439		J, et al. Severe strongyloidiasis: a systematic review of case reports. BMC Infect
440		Dis. 2013;13:78.
441	[20]	Angheben A, Anselmi M, Gobbi F, Marocco S, Monteiro G, Buonfrate D, et al.
442		Chagas disease in Italy: Breaking an epidemiological silence. Eurosurveillance.
443		2011;16.
444	[21]	El Moghazy W, Kashkoush S, O'hali W, Abdallah K. Long-term outcome after liver
445		transplantation for hepatic schistosomiasis: a single-center experience over 15
446		years. Liver Transplant Off Publ Am Assoc Study Liver Dis Int Liver Transplant
447		Soc. 2015;21:96–100.
448	[22]	Abu-Madi MA, Behnke JM, Boughattas S, Al-Thani A, Doiphode SH, Deshmukh A.
449		Helminth infections among long-term-residents and settled immigrants in Qatar in
450		the decade from 2005 to 2014: temporal trends and varying prevalence among
451		subjects from different regional origins. Parasit Vectors. 2016;9:153.
452	[23]	Bocanegra C, Salvador F, Sulleiro E, Sànchez-Montalvà A, Pahissa A, Molina I.
453		Screening for imported diseases in an immigrant population: Experience from a
454		teaching hospital in Barcelona, Spain. Am J Trop Med Hyg. 2014;91:1277–81.
455	[24]	Geltman PL, Cochran J, Hedgecock C. Intestinal parasites among African refugees
456		resettled in Massachusetts and the impact of an overseas pre-departure treatment
457		program. Am J Trop Med Hyg. 2003;69:657-62.
458	[25]	Salvador F, Molina I, Sulleiro E, Burgos J, Curran A, Van den Eynde E, et al.
459		Tropical diseases screening in immigrant patients with human immunodeficiency
460		virus infection in Spain. Am J Trop Med Hyg. 2013;88:1196–202.
461	[26]	Pardo J, Carranza C, Muro A, Angel-Moreno A, Martín AM, Martín T, et al.
462		Helminth-related eosinophilia in African immigrants, Gran Canaria. Emerg Infect
463		Dis. 2006;12:1587–9.
464	[27]	Muennig P, Pallin D, Sell RL, Chan MS. The cost effectiveness of strategies for the
465		treatment of intestinal parasites in immigrants. N Engl J Med. 1999;340:773–9.
466	[28]	Rapoport AB, McCormick D, Cohen PA. Screening for Schistosoma mansoni and
467		Strongyloides stercoralis Infection Among Brazilian Immigrants in the United States
468		Open forum Infect Dis. 2015;2:ofv003.
469	[29]	Conway DJ. Atkins NS. Lillywhite JE. Bailey JW. Robinson RD. Lindo JF. et al.

470		Immunodiagnosis of Strongyloides stercoralis infection: a method for increasing the
471		specificity of the indirect ELISA. Trans R Soc Trop Med Hyg. 1993;87:173-6.
472	[30]	Bern C. Chagas' Disease. N Engl J Med. 2015;373:456–66.
473	[31]	Zammarchi L, Strohmeyer M, Bartalesi F, Bruno E, Muñoz J, Buonfrate D, et al.
474		Epidemiology and Management of Cysticercosis and Taenia solium Taeniasis in
475		Europe, Systematic Review 1990-2011. PLoS One. 2013;8.
476	[32]	Karki T, Napoli C, Riccardo F, Fabiani M, Grazia Dente M, Carballo M, et al.
477		Screening for Infectious Diseases among Newly Arrived Migrants in EU/EEA
478		Countries???Varying Practices but Consensus on the Utility of Screening. Int J
479		Environ Res Public Health. 2014;11:11004–14.
480	[33]	Muennig P, Pallin D, Challah C, Khan K. The cost-effectiveness of ivermectin vs.
481		albendazole in the presumptive treatment of strongyloidiasis in immigrants to the
482		United States. Epidemiol Infect. 2004;132:1055–63.
483	[34]	Machado CM, Martins TC, Colturato I, Leite MS, Simione AJ, De Souza MP, et al.
484		Epidemiology of neglected tropical diseases in transplant recipients. Review of the
485		literature and experience of a Brazilian HSCT center. Rev Inst Med Trop Sao Paulo
486		2009;51:309–24.
487		

L. C. Alice	Numerator and denominator	Overall prevalence (95% CI)	Women to men ratio	Geosentinel Regions with highest prevalence		
Infection				Geosentinel Region	Numerator and denominator	Region-specific prevalence (95% CI)
				North East Asia	2/31	6.45% (1.48-23.93)
Strongyloidiasis	42/939	4.51% (3.35-6.05)	1:2	Sub-Saharan Africa	20/330	6.06% (3.93-9.22)
				South Central Asia	5/90	5.55% (2.29-12.85)
	31/519	5.97% (4.22-8.37)	1:2.8	Sub-Saharan Africa	25/323	7.73% (5.27-11.22)
Schistosomiasis				South America	3/46	6.52% (2.02-19.05)
				South Central Asia	1/34	2.94% (0.37-19.77)
Chagas disease	7/180	3.88% (1.85-7.98)	2.5:1	South America*	7/172	4.06% (1.93-8.34)
			(2)	South East Asia	2/6	33.33% (4.18-85.13)
Toxocariasis	11/113	9.73% (5.42-16.86)	1:1.8	South America	4/19	21.05% (7.33-47.32)
				Eastern Europe	3/24	6.89% (3.73-34.48)
Filariasis	5/54	9.25% (7.83-11.63)	1:1.5	Sub-Saharan Africa*	5/34	14.70% (5.96-31.91)

^{*} Only one region is reported because the infection was not found in people coming from other areas