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http://dx.doi.org/10.1017/S0031182020000645

| **Title** | Human African trypanosomiasis: the current situation in endemic regions and the risks for non-endemic regions from imported cases |
|-----------|-------------------------------------------------------------------------------------------------------------------------|
| **Authors** | Gao, J-M, Qian, Z-Y, Hide, G, Lai, D-H, Lun, Z-R and Wu, Z-D |
| **Type** | Article |
| **URL** | This version is available at: http://usir.salford.ac.uk/id/eprint/57024/ |
| **Published Date** | 2020 |

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Human African trypanosomiasis: the current situation in endemic regions and the risks for non-endemic regions from imported cases

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Summary

Human African trypanosomiasis (HAT) is caused by *Trypanosoma brucei gambiense* and *T. b. rhodesiense* and caused devastating epidemics during the 20th century. Due to effective control programs implemented in the last two decades, the number of reported cases has fallen to a historically low level. Although fewer than 977 cases were reported in 2018 in endemic countries, HAT is still a public health problem in endemic regions until it is completely eliminated. In addition, almost 150 confirmed HAT cases were reported in non-endemic countries in the last three decades. The majority of non-endemic HAT cases were reported in Europe, United States and South Africa, due to historical alliances, economic links or geographic proximity to disease endemic countries. Furthermore, with the implementation of the “Belt and Road” project, sporadic imported HAT cases have been reported in China as a warning sign of tropical diseases prevention. In this paper, we explore and interpret the data on HAT incidence and find no positive correlation between the number of HAT cases from endemic and non-endemic countries. This data will provide useful information for better understanding the imported cases of HAT globally in the post-elimination phase.

**Key words:** Human African trypanosomiasis; *Trypanosoma brucei gambiense*; *T. b. rhodesiense*; sub-Saharan Africa; non-endemic disease countries
Introduction

Human African trypanosomiasis (HAT), also known as sleeping sickness, is a neglected tropical disease, which is found in 36 African endemic countries (Simarro et al., 2010). It is caused by subspecies of the parasitic protozoan *Trypanosoma brucei* and transmitted by tsetse flies. Two subspecies are able to infect humans: *T. b. gambiae* causes a chronic form of HAT in West and Central Africa, while *T. b. rhodesiense* is the pathogenic agent for the acute form of the disease in Eastern Africa (Barrett et al., 2003; Stich et al., 2002). In the last decade of the 20th century, the number of reported cases of HAT reached alarming levels with up to 50,000 new cases reported each year (Hide, 1999). Therefore, surveillance and control activities against HAT were reinforced at the beginning of the 21st century. These effective measures enabled a steady decrease in HAT cases (Franco et al., 2014a; WHO, 2013a). As a consequence, HAT was included in the WHO NTD roadmap of diseases targeted for elimination by 2020. The WHO definition of elimination for HAT is that the incidence is less than 1 new case per 10000 population in at least 90% of foci and with fewer than 2000 cases reported globally. According to this definition, the criteria for elimination was achieved in 2017.

It is well-known that the prevalence of the pathogen, *T. brucei* is restricted to sub-Saharan Africa, due to the distribution limitations of its transmission vector, the tsetse flies of the genus *Glossina* (Migchelsen et al., 2011). However, human population movements, including travelling, business connections and migration may put people with little awareness of HAT in danger. This is evidenced by reports of HAT-positivity in some travelers, military personnel and immigrants. During the period 2000 to 2010, more than 100 cases of HAT were reported in 19 non-disease endemic countries (Simarro et al., 2012). According to the UN World Tourism...
Organization, the number of international tourist arrivals increased from 25 million in 1950 to 1035 million in 2012 (Norman et al., 2015). The strongest growth in international tourist arrivals was recorded in Asia, followed by the Pacific Region, Africa and the Americas. According to the Chinese government, the number of Chinese people living overseas has reached 60 million. Furthermore, approximately 122 million Chinese people traveled overseas in 2016. Additionally, a large number (up to 52.67 million) non-Chinese travelers crossed China’s borders in 2014. Sino-overseas cooperation, triggered by the “Belt and Road Initiative”, has promoted the international movement of a huge population of both Chinese and non-Sino peoples and this will greatly increase the importation risk of tropical diseases that were not endemic in China.

In this article, we critically analyze the incidence of human African trypanosomiasis (HAT) in endemic countries and non-endemic countries during the period between 1990 and 2018.

Search strategy and selection criteria

We searched PubMed for the terms “Trypanosoma brucei”, “Trypanosoma brucei gambiense”, “Trypanosoma brucei rhodesiense”, “Human African trypanosomiasis” to identify papers published between Jan 1, 1990 and Dec 31, 2018 on Human African trypanosomiasis cases in non-endemic countries. Additionally, we reviewed relevant articles cited in those references and included them as primary sources when appropriate.
Incidence of human African trypanosomiasis in endemic countries from 1990-2018

Human African trypanosomiasis was first described around the 14th century, but it can probably be considered as endemic in African regions since the appearance of humans. In the 20th century, because of the socio-economic impact caused by HAT, effective measures were taken to control the disease. This resulted in it reaching a very low, generalized, transmission by the mid-1960s with a minimum of 4,435 cases declared in Africa in 1964 (WHO, 2013b). However, post-1960s, disease control interventions were constrained due to conflicts, insecurity and other social instability factors which largely resulted from the inevitable (and justifiable) emergence of independent African states from their former European colonizers. Unintentionally, this led to a resurgence of the disease in the 1980s and 1990s (Franco et al., 2014a,b). As a consequence, the number of sleeping sickness cases reached alarming levels such that by 1998 the newly reported cases were up to 37991 and the total number of existing cases of HAT was estimated at 300,000 cases (WHO, 1998). At that time, the status of HAT triggered attention from the health authorities in endemic countries alongside bilateral and multilateral agencies (Franco et al., 2014a, b). The World Health Organization (WHO) also responded to this situation by coordinating international partners working in HAT control, raising awareness and political will, and bringing new resources from the public and private sectors to support national control programs. Control programs were reinforced again and as a result, in the past 20 years, important advances have been made in controlling this disease. A substantial, steady reduction in the number of cases has been observed, reaching the historical lowest figure in 2018 (Figure 1).

Among the two species which cause human African trypanosomiasis, more than
90% of HAT cases were caused by *T. b. gambiense* (Figure 2A and 2B). It is worth noting that the cases in the Democratic Republic of the Congo (DRC) consistently accounted for the vast majority of *gambiense*-HAT cases. Especially in 1997 and 1998, increases in new cases of *gambiense*-HAT (25094 and 26318) in the DRC, led to a sharp increase in total number of *gambiense*-HAT cases. In addition, new cases of *gambiense*-HAT in Angola also sharply increased during this period due to civil war, as 6726, 8275 and 6610 new cases were reported in 1996, 1997 and 1998 respectively (Figure 2A, Table S1). However, the total number of reported cases of rhodesiense HAT has shown a gradual descending trend over nearly three decades. We found that almost all of new cases were reported in Uganda and United Republic of Tanzania, and the number of cases in both countries accounted for over 80% of all cases from 1990 to 2007 (Figure 2B, Table S2). By 2008, the number of new reported cases dropped below 200 and less than 100 new cases were found in 2011.

According to reports from WHO, less than 1500 new cases of HAT (including 1420 new cases of *T. b. gambiense* (WHO, 2019a), representing about 98% of the total HAT cases, and 27 new cases of *T. b. rhodesiense* (WHO, 2019b), representing less than 2% of the total HAT cases) were recorded in 2017 (Table S1 and S2). As always, the majority of *gambiense*-HAT cases were distributed in the Democratic Republic of the Congo with up to 1110 cases (78% of the total *gambiense*-HAT) (Figure 3). Furthermore, 140 and 76 new cases of *gambiense*-HAT were reported in Guinea and the Central African Republic respectively. For the rest, less than 50 new cases of *gambiense*-HAT, were reported in Angola (18 cases), Cameroon (5 cases), Chad (28 cases), Congo (15 cases), Côte d’Ivoire (3 cases), Equatorial Guinea (4 cases), Gabon (9 cases) and South Sudan (12 cases) respectively. To our knowledge, there are no reported *gambiense*-HAT cases in other West and Central Africa countries.
However, during 2017, there were only 27 reported new cases of *rhodesiense*-HAT in Eastern African endemic areas, these were distributed in Uganda (13 cases), Malawi (7 cases), United Republic of Tanzania (3 cases), Zambia (3 cases) and Zimbabwe (1 case) respectively.

Based on the latest reports from WHO (WHO, 2019a,b), the case numbers of HAT were down to their lowest-ever levels in 2018 and the total number of new cases was less than 1000, including 953 cases of *T. b. gambiense*, and 24 new cases of *T. b. rhodesiense*. It was exciting that the number of reported *gambiense*-HAT cases from the Democratic Republic of the Congo was down to 660 and the HAT cases from every other country numbered fewer than 100 (Figure 3B). Surprisingly, the case number in Angola increased from 18 (2017) to 79 (2018). For *T. b. rhodesiense*, new cases were only reported in Malawi (15 cases), Uganda (4 cases) and Zambia (5 cases) respectively in 2018.

**Cases of human African trypanosomiasis reported in non-disease endemic countries (DECs)**

The human African trypanosomiasis cases in non-endemic countries from 1990 to 2010 have been collected in previous reports (Migchelsen *et al.*, 2011; Simarro *et al.*, 2012). Therefore, we will concentrate on cases in non-endemic countries reported between 2011 and 2018. A total of 15 cases of HAT were reported in 11 non-endemic countries (Figure 4; Table 1), including 9 cases of *rhodesiense*-HAT and 4 cases of *gambiense*-HAT and another 2 indeterminate cases (Table 1). Rhodesiense- HAT was mainly diagnosed in tourists after short visits to endemic countries, and the majority of them were in the first stage of disease. However, the majority of *gambiense*-HAT patients had been living in endemic countries for an extended period for various
reasons (eg. migrants or expatriates, etc). Therefore, by the time they were diagnosed, the disease had developed into the second stage. Due to historical or economic links with endemic countries and frequent travel to endemic countries, citizens from Europe have more chances of becoming infected with trypanosomes. More than half of the cases (8 cases, 53.3% of total) were diagnosed in European countries (including Belgium, 1 case; France, 1 case; Germany, 2 cases; Netherlands, 1 case; Spain, 1 case; and UK, 2 cases). In addition, another 2 cases were reported in North America (including one in Canada and one in USA) and South America (Argentina) respectively. The remaining 4 cases were identified in Asia. Interestingly, with the exception of 1 case from Israel, the other 3 HAT cases were all diagnosed in China, which is now starting to appear as a non-DEC with increasing numbers of imported HAT cases during the period of 2011 to 2018.

Furthermore, the number of reported cases diagnosed in non-endemic countries between 2011 to 2018 were compared with previous records (Figure 5). It is reported that many more HAT cases were diagnosed in non-endemic countries from 2001 to 2010 including 77 cases of *rhodesiense*-HAT and 31 cases of *gambiense*-HAT, as detailed in supplementary table 3. As shown in Figure 6, during this period of time, many imported HAT cases were diagnosed in South Africa (26 cases, accounting for 24.1% of the total 108 cases). This is probably due to its proximity to endemic countries with famous protected areas and game reserves. In addition, due to historical or economic links with endemic countries, most of the HAT cases were diagnosed in North American and European countries, 21 and 53 cases, account for 19.4% and 49.1%, respectively. Among them, the United States and United Kingdom contributed significant numbers of imported HAT cases (US, 18 cases; and the UK, 12 cases), followed by Netherlands (11), Italy (9), France (6) and Germany (5).
Only 21 cases of HAT (15 cases of *rhodesiense*-HAT and 6 cases of *gambiense*-HAT) were reported in non-endemic countries between 1990 to 2000 (Figure 5, Table S4). Among them, most cases were mainly diagnosed in the USA (6) and France (5), with the rest being diagnosed in the Netherlands (1), Germany (1), Italy (1), Mexico (1), New Zealand (1), South Africa (1), Switzerland (2) and the UK (2) (Figure 6). There was a resurgence of HAT in the 1990s, when the number of sleeping sickness cases reached the highest ever recorded in 1998. Why there is no positive correlation between number of HAT cases in non-endemic countries and endemic countries is puzzling? It may be that some HAT cases were not published in the literature or perhaps there have been some misdiagnosed cases in the past. However, from the beginning of this century, pharmacy services in non-endemic countries have had to request drugs from WHO to treat HAT cases and, at the same time, provide epidemiological, parasitological, biological, and clinical data. Therefore, this information has enabled WHO to maintain HAT surveillance system and a comprehensive database for non-endemic countries.

The sources of imported HAT cases in non-endemic regions

During the period from 1990 to 2018, more than 101 reported *rhodesiense*-HAT cases were exported from eastern and southern Africa (As shown in Table 2). It is supposed that the infection has been contracted in protected areas such as national parks (NP), wildlife reserves, and game reserves (Selby et al. 2013; Simarro et al., 2012). The country exporting the majority of rhodesiense HAT cases is the United Republic of Tanzania, with up to 56.4% (57 cases). Other exporting countries are Malawi (15%, 15 cases), Zambia (9.9%, 10 cases), Zimbabwe (6%, 6 cases), Rwanda (5%, 5 cases), Kenya (3%, 3 cases) and Uganda (2%, 2 cases). However, the number
of rhodesiense HAT endemic cases in a country does not positively correlate with the number of cases exported. For example, from 1990 to 2000, there were no new cases reported in Rwanda, nevertheless it exported more than 5 cases to non-endemic areas from this country; while the maximum number of endemic cases was reported in the Uganda, but no cases were exported from this country. In addition, only 8 cases were reported in Zimbabwe from 2001 to 2010, but as many as 5 cases were exported during this period. Furthermore, the number of exported cases from Kenya is the same as the number of domestic cases from 2011 to 2018. Perhaps the factors underlying the export of cases are related more to the awareness of travelers and the messaging around risk in specific countries than biological factors such as the epidemiology of exposure.

However, the majority of HAT cases were gambiense-HAT that were reported in western and central Africa, whereas there were only 40 cases of exported gambiense-HAT during 1990 to 2018 (Table 3). The countries of origin for exported gambiense-HAT are mainly the Gabon, Angola and DRC, each accounting for 25% (10 cases), 20% (8 cases) and 20% (8 cases), followed by Cameroon (10%, 4 cases), Equatorial Guinea, Nigeria, Central African Republic and Uganda (5%, 2 cases each), Sudan (2.5%, 1 case) and one uncertain case (2.5%) which may have originated from Nigeria or Gabon. To our surprise, as many as 7 and 3 cases were exported from Gabon and Cameroon respectively, despite numbers of indigenous cases being as low as 305 and 174 in these two countries from 2001 to 2010. In addition, although only 6 indigenous cases were reported in Nigeria, 1 case of exported HAT has been reported between 2011 and 2018. These seems to be a general lack of association between incidence and exported infections across many endemic countries.
HAT carriers may not be the main infective source for non-endemic countries

In an analysis of the dynamic change trend of HAT cases diagnosed in non-endemic countries and endemic countries, we found that there is no positive correlation ($P = 0.62$) between the two groups (Figure 7). It is possible that the number of people traveling to eastern Africa, endemic areas of the *rhodesiense*-HAT, is higher than that in other regions. This suggestion is supported by the most recent tourism data (2018) that shows higher numbers of incoming tourists in East African DEC countries (e.g. Malawi, 870k; Zambia, 1072k; Tanzania, 1506k) compared with West African DEC countries (e.g. DRC, 351k (2016 data); Congo 158k) (UNWTO, 2020). However, more than 70% of the cases reported from non-endemic countries were infected during trips to east African countries, in particular to popular tourist countries, such as Tanzania, Malawi or Zambia. By comparison, the overall proportion of *rhodesiense*-HAT cases reported from endemic countries accounts for only 10 percent of the total number of cases.

Interestingly, inspection of the data presented in Figure 7 shows that peaks of reported non-DEC infections occur a few years after the corresponding peaks in DECs (see for example the endemic disease peak in 1997-1998 with the peak of reported cases in non DECs 2001-2002, four years later). This suggests that there might be a time delay between the local disease status in endemic countries and the recording of reported cases in non-DECs. Analysis of correlation between the number of cases in DECs with those reported from non-DECs shows a highly significant correlation ($P = 0.0011$) when the comparative data sets are offset by 4 years (i.e. DEC data from 1990 to 2014 compared year by year with non-DEC data from 1994 to 2018). Significant
correlations are additionally found when comparisons are offset by 3 to 7 years. Such a delay between the endemic peak could be attributed to the relatively long incubation period in *T. b. gambiense* infections, a delay in diagnosis or reporting of cases in the recipient non-DEC or a generic delay in reporting back to WHO. Comparable significant correlations were found when this analysis was carried out separately on *T. b. gambiense* data (e.g. $P = 0.0009$; 4 year offset data) but no correlation at all was seen when analyzing *T. b rhodesiense* non-DEC case data (e.g. $P = 0.83$; 4 year offset data). This broad analytical approach has many flaws but suggests that more detailed case information and better stratification of the data might tell us more about the role of HAT carriers or the reporting processes in non-DEC cases.

In addition to human reservoirs, many livestock and wildlife species can act as animal reservoirs, from which *T. b. gambiense* and *T. b. rhodesiense* maybe spill over to humans (Büscher *et al*., 2017; Cecchi *et al*., 2014, 2015; Cordon-Obras *et al*., 2015; Jamonneau *et al*., 2004; Kageruka, 1989; Mehlitz, 1997; Molyneux, 1973; Njiooku *et al*., 2006; N’Djetchi *et al*., 2017; Simarro *et al*., 2010; Yesufu, 1971). It has been demonstrated that human-derived *T. b. gambiense* strains that were cyclically transmitted by tsetse flies between animals for more than a year remained transmissible to humans (Van Hoof, 1947). Compared to latent infections in humans, our current knowledge of *T. b. gambiense* and *T. b. rhodesiense* infections in animals is very limited and fragmented. Therefore, it is not known whether cryptic reservoirs in animals will compromise the sustainable elimination of human Africa trypanosomiasis.

**The risk of mechanical transmission of *Trypanosoma* spp.**

It is known that tsetse flies play the most important role, as cyclical vectors, in
the transmission of African human and livestock trypanosomes (*Trypanosoma congolense*, *T. vivax*, and *T. brucei*) (Desquesnes and Dia, 2003a,b; Hoare, 1972). However, previous research has demonstrated that many species of *Trypanosoma* can be mechanically transmitted by haematophagous insects or sucking flies (Desquesnes and Dia, 2003a,b). Importantly, *T. brucei rhodesiense* was reported to be mechanically transmitted by *Musca sorbens*, which could potentially pose a threat to human populations and livestock in non-endemic areas (Büscher et al., 2018; Simarro et al., 2010).

**China as a new destination for imported cases**

Until recently, there have been no reported cases of trypanosomiasis in China. As China further develops international connections through economic development and tourism in African countries, the risk of the emergence of non-endemic cases in China will increase. Such a development may require a greater awareness, in the Chinese health systems, of the possible emergence of cases. Strikingly, there have been three cases of imported human African trypanosomiasis in recent years, including 2 cases of *T. b. gambiense*, discovered in Chinese workers and presumably infected in Gabon, Western Africa; whereas another case of *T. b. rhodesiense* was found in a Chinese tourist visiting game parks in Tanzania, Eastern Africa (Zhou et al., 2018). According to the Chinese government, approximately 60 million Chinese people are now living overseas and in 2016, more than 122 million Chinese people travelled overseas (IOMMA, 2017). Additionally, the number of visitors entering China numbered up to 52.67 million in 2014 (CNTA, 2016). Therefore, economic globalization and population migration will increase the importation risk of tropical diseases that were not previously endemic in China and, of course, other countries too. This situation
may be mitigated as African countries are now moving into the elimination phase of sleeping sickness but the situation needs to be monitored in the possible event of any post-elimination resurgence of disease.

Conclusions

Sleeping sickness, also known as human African trypanosomiasis (HAT), is a neglected disease that impacts 70 million people living in 1.55 million km$^2$ in sub-Saharan Africa (Aksoy et al., 2017). The successful control activities carried out over the last less 20 years have substantially reduced the burden of disease. According to the latest report from WHO (WHO, 2019a,b), new cases of HAT were less than 1000 in 2018 (98% of total HAT is $T. b. gambiae$, while $T. b. rhodesiense$ is less than 2%). However, in some rural areas where trypanosomiasis is prevalent, the overall performance of peripheral health systems is often weak and characterized by unskilled staff, low attendance and low coverage.

Furthermore, political instability and insecurity are inescapable challenges in an elimination programme. Gambiense-HAT does not spread easily and humans are the only significant reservoir. The successful elimination of $gambiense$-HAT requires a turmoil-free socio-political context. However, the elimination of $rhodesiense$-HAT poses challenges because of the vast animal reservoirs.

Therefore, the strengthening of health systems to implement the activities included in the elimination strategies is essential. Past and current experiences show that HAT elimination is difficult and will require more effort, time, and money than initially anticipated. History teaches us that falling case numbers can result in a decline in donor and control agency interest, opening the door to swift and severe recrudescence (Büscher et al., 2017). Advocacy is needed to sensitize donors on the
importance of switching from a control to an elimination mindset.

It is known that the majority of HAT cases reported by endemic countries correspond to the gambiense form (98%), the opposite is true for imported cases in non-endemic countries, where 72% of cases are caused by \textit{T. b. rhodesiense} and 28% by \textit{T. b. gambiense}. However, the proportion of gambiense- to rhodesiense-HAT cases diagnosed in non-endemic countries is lower than one would probably expect. Moreover, it is difficult to establish the number of migrants and refugees traveling to non-endemic countries from HAT transmission areas, and even more difficult to ascertain how many of them are affected by HAT. Therefore, it is a potential threat to non-endemic countries because global mobility means that \textit{Trypanosoma brucei} can be spread in many ways.

**Acknowledgments.** We thank Fangli Lv at Zhongshan School of Medicine, Sun Yat-Sen University for her reviewing and editing the English language; and thank Zhishen Xu at School of Life Sciences, Sun Yat-Sen University for comments on data analysis.

**Financial support.** Authors’ laboratories are supported by grants from the National Science and Technology Major Project (2018ZX10101002-001 and 2018ZX10101002) and the National Natural Science Foundation (31720103918 and 31672276 to ZRL).

**Conflict of interest.** We declare no competing interests.

**Ethical standards.** Not applicable.
References

Aksoy S, Buscher P, Lehane M, Solano P and Van Den Abbeele J (2017) Human African trypanosomiasis control: Achievements and challenges. PLoS Negl Trop Dis 11(4), e0005454.

Barrett MP, Burchmore RJ, Stich A, Lazzari JO, Frasch AC, Cazzulo JJ and Krishna S (2003) The trypanosomiases. Lancet 362, 1469-1480.

Büscher P, Cecchi G, Jamonneau V and Priotto G (2017) Human African trypanosomiasis. Lancet 390, 2397-2409.

Büscher P, Bart JM, Boelaert M, Bucheton B, Cecchi G, Chitnis N, Courtin D, Figueiredo LM, Franco JR, Grébaut P, Hasker E, Ilboudo H, Jamonneau V, Koffi M, Lejon V, MacLeod A, Masumu J, Matovu E, Mattioli R, Noyes H, Picado A, Rock KS, Rotureau B, Simo G, Thévenon S, Trindade S, Truc P and Van Reet N (2018) Do Cryptic Reservoirs Threaten Gambiense-Sleeping Sickness Elimination? Trends Parasitol 34(3), 197-207.

Cecchi G, Paone M, Feldmann U, Vreysen MJ, Diall O and Mattioli RC (2014)
Assembling a geospatial database of tsetse-transmitted animal trypanosomosis for Africa. Parasit Vectors 7, 39.

Cecchi G, Paone M, Argilés Herrero R, Vreysen MJ and Mattioli RC (2015)
Developing a continental atlas of the distribution and trypanosomal infection of tsetse flies (Glossina species). Parasit Vectors 8, 284.

China National Tourism Administration (CNTA) (2017) Big data on the overseas travelling in China in 2016. http://www.cnta.gov.cn/xxfb/hydt/201701/t20170124_813152.shtml. Accessed on 3 September 2017 (in Chinese).

Clerinx J, Vlieghe E, Asselman V, Van de Casteele S, Maes MB and Lejon V (2012)
Human African Trypanosomiasis in a Belgian Traveller Returning from the Masai Mara Area, Kenya, February 2012. *Euro Surveill* 17(10).

Cordon-Obras C, Rodriguez YF, Fernandez-Martinez A, Cano J, Ndong-Mabale N, Ncogo-Ada P, Ndongo-Asumu P, Aparicio P, Navarro M, Benito A and Bart JM (2015) Molecular evidence of a *Trypanosoma brucei gambiense* sylvatic cycle in the human African trypanosomiasis foci of Equatorial Guinea. *Front Microbiol* 6, 765.

Cottle LE, Peters JR, Hall A, Bailey JW, Noyes HA, Rimington JE, Beeching NJ, Squire SB and Beadsworth MB (2012) Multiorgan Dysfunction Caused by Travel-Associated African Trypanosomiasis. *Emerg Infect Dis* 18, 287-289.

De Kyvon MA, Maakaroun-Vermesse Z, Lanotte P, Priotto G, Perez-Simarro P, Guennoc AM, Toffol BD, Paris L, Bernard L, Goudeau A, Chandenier J and Desoubeaux G (2016) Congenital Trypanosomiasis in Child Born in France to African Mother. *Emerg Infect Dis* 22, 935-937.

Desquesnes M and Dia ML (2003a) Mechanical transmission of *Trypanosoma congolense* in cattle by the African tabanid *Atylotus agrestis*. *Exp Parasitol* 105(3-4), 226-231.

Desquesnes M and Dia ML (2003b) *Trypanosoma vivax*: mechanical transmission in cattle by one of the most common African tabanids, *Atylotus agrestis*. *Exp Parasitol* 103, 35-43.

Franco JR, Simarro PP, Diarra A, Ruiz-Postigo JA and Jannin JG (2014a) The journey towards elimination of gambiense human African trypanosomiasis: not far, nor easy. *Parasitology* 141(6), 748-760.

Franco JR, Simarro PP, Diarra A and Jannin JG (2014b) Epidemiology of human African trypanosomiasis. *Clin Epidemiol* 6, 257-275.
Gomez-Junyent J, Pinazo MJ, Fernández S, Mas J, Chaguaceda C, Pellicé M, Gascón J and Muñoz J (2017) Human African Trypanosomiasis in a Spanish Traveler Returning from Tanzania. *PLoS Negl Trop Dis* 11, e0005324.

Hide G (1999) History of sleeping sickness in East Africa. *Clin Microbiol Rev* 12(1), 112-125.

Hoare CA (1972) The Trypanosomes of Mammals. A Zoological Monograph. Blackwell Scientific Publications, Oxford, UK: pp. 749.

Hu X, Sun Y, Xia X, Xu L, Min B, Zhou D, Wang L, Wang H, Niu Z, He M, Wang J, Deng A and Yang J (2016) Photo Quiz: A Man with Lymphadenopathy and Lethargy. *J Clin Microbiol* 54, 2631.

International Organization for Migration Migration Activities (IOMMA) (2017) https://www.iom.int/countries/china. Accessed on 3 September 2017 (in Chinese).

Jamonneau V, Ravel S, Koffi M, Kaba D, Zeze DG, Ndri L, Sane B, Coulibaly B, Cuny G and Solano P (2004) Mixed infections of trypanosomes in tsetse and pigs and their epidemiological significance in a sleeping sickness focus of Côte d'Ivoire. *Parasitology* 129, 693-702.

Kageruka P (1989) Réservoir animal de *Trypanosoma* (Trypanozoon) *brucei gambiae* en Afrique Centrale. *Ann Soc Belg Méd Trop* 69, 155-163.

Liu Q, Chen XL, Chen MX, Xie HG, Liu Q, Chen ZY, Lin YY, Zheng H, Chen JX, Zhang Y and Zhou XN (2018) *Trypanosoma Brucii Rhodesiense* Infection in a Chinese Traveler Returning from the Serengeti National Park in Tanzania. *Infect Dis Poverty* 7, 50.

Luintel A, Lowe P, Cooper A, MacLeod A, Büscher P, Brooks T and Brown M (2017) Case of Nigeria-Acquired Human African Trypanosomiasis in United Kingdom,
Mehlitz D (1977) The behaviour in the blood incubation infectivity test of four Trypanozoon strains isolated from pigs in Liberia. *Trans R Soc Trop Med Hyg* 71, 86.

Meltzer E, Leshem E, Steinlauf S, Michaeli S, Sidi Y and Schwartz E (2012) Human African Trypanosomiasis in a Traveler: Diagnostic Pitfalls. *Am J Trop Med Hyg* 87, 264-266.

Migchelsen SJ, Buscher P, Hoepelman AI, Schallig HD and Adams ER (2011) Human African trypanosomiasis: a review of non-endemic cases in the past 20 years. *Int J Infect Dis* 15, e517-e524.

Molyneux DH (1973) Animal reservoirs and Gambian trypanosomiasis. *Ann Soc Belg Med Trop* 53, 605-618.

N’Djetchi MK, Ilboudo H, Koffi M, Kaboré J, Kaboré JW, Kaba D, Courtin F, Coulibaly B, Fauret P, Kouakou L, Ravel S, Deborggraeve S, Solano P, De Meeûs T, Bucheton B and Jamonneau V (2017) The study of trypanosome species circulating in domestic animals in two human African trypanosomiasis foci in Côte d’Ivoire identifies pigs and cattle as potential reservoirs of *Trypanosoma brucei gambiense*. *PLoS Negl Trop Dis* 11, e0005993.

Njiokou F, Laveissière C, Simo G, Nkinin S, Grébaut P, Cuny G and Herder S (2006) Wild fauna as a probable animal reservoir for *Trypanosoma brucei gambiense* in Cameroon. *Infect Genet Evol* 6, 147-153.

Norman FF, Monge-Maillo B, Martínez-Pérez Á, Perez-Molina JA and López-Vélez R (2015) Parasitic infections in travelers and immigrants: part I protozoa. *Future Microbiol* 10, 69-86.

Pale CA and Vigna L (2013) Images in Clinical Medicine. African Trypanosomiasis in 2016. *Emerg Infect Dis* 23, 1225-1227.
Argentina. N Engl J Med 369, 763.

Pasternak J, Wey SB, Silveira PA and Camargo TZ (2013) An African Visitor in Brazil. Einstein (Sao Paulo) 11, 261-262.

Richter J, Göbels S, Göbel T, Westenfeld R, Müller-Stöver I and Häussinger D (2012) A Returning Traveller with Fever, Facial Swelling, and Skin Lesions. Bmj 344, e2092.

Selby R, Bardosh K, Picozzi K, Waiswa C and Welburn SC (2013) Cattle movements and trypanosomes: restocking efforts and the spread of Trypanosoma brucei rhodesiense sleeping sickness in post-conflict Uganda. Parasit Vectors 6, 281.

Simarro PP, Cecchi G, Paone M, Franco JR, Diarra A, Ruiz JA, Févre EM, Courtin F, Mattioli RC and Jannin JG (2010) The atlas of human African trypanosomiasis: a contribution to global mapping of neglected tropical diseases. Int J Health Geogr 9, 57.

Simarro PP, Franco JR, Cecchi G, Paone M, Diarra A, Ruiz Postigo JA and Jannin JG (2012) Human African Trypanosomiasis in Non-Endemic Countries (2000–2010). Journal of Travel Medicine 19 (1), 44-53.

Stich A, Abel PM and Krishna S (2002) Human African trypanosomiasis. BMJ 325, 203-206.

Streit JA and Matsumoto E (2016) African Trypanosomiasis. N Engl J Med 375, 2380.

United Nations World Tourism Organization. www.e-unwto.org/toc/unwtotfb/current.

Van Hoof LMJJ (1947) Observations on trypanosomiasis in the Belgian Congo. Trans R Soc Trop Med Hyg 40,728-761.

Wang XY, Ruan QL, Zhang WH, Gu JF, Qian YY, Chen MX, Liu Q, Lu Q and Zhang WH (2018) Human African Trypanosomiasis in Emigrant Returning to China from Gabon, 2017. Emerg Infect Dis 24, 400-404.
Wijsman CA, Hanssen JLJ, Scheper H, Visser LG and van Lieshout L (2018) A Case of Delayed Diagnosis of East-African Trypanosomiasis in a Dutch Traveller. J Travel Med 25, 10.

Wolf T, Wichelhaus T, Gottig S, Kleine C, Brodt HR and Just-Nuebling G (2012) Trypanosoma Brucei Rhodesiense Infection in a German Traveller Returning from the Masai Mara Area, Kenya, January 2012. Euro Surveill 17(10), 20114.

World Health Organization (1998) Control and Surveillance of African Trypanosomiasis. Report of a WHO Expert Committee. WHO Technical Report Series 881. Geneva, Switzerland: World Health Organization.

World Health Organization (2013a) Control and surveillance of human African trypanosomiasis. Geneva: World Health Organization.

World Health Organization (2013b) WHO Report on Global Surveillance of Epidemic-Prone Infectious Diseases. Geneva, Switzerland: World Health Organization; 2000. Available from: http://whqlibdoc.who.int/hq/2000/WHO_CDS_CSR_ISR_2000.1.pdf. Accessed December 21.

World Health Organization (2019a) Number of new reported cases (T.b. gambiense) Data by country. https://apps.who.int/gho/data/node.main.A1636?lang=en.

World Health Organization (2019b) Number of new reported cases (T.b. rhodesiense) Data by country. https://apps.who.int/gho/data/node.main.A1637?lang=en.

Yesufu HM (1971) Experimental transmission of Trypanosoma gambiense in domestic animals. Ann Trop Med Parasitol 65, 341-347.

Zhou XN, Qian MB, Priotto G, Franco JR and Guo JG (2018) Tackling imported tropical diseases in China. Emerg Microbes Infect 7(1), 12.
Table 1. Cases of human African trypanosomiasis (HAT) recorded in non-endemic countries from 2011 to 2018.

| Cases | Refs | Time  | Place of diagnosis | Place of infection | Sex/age | Activity  | Stage | Chancre | Species          |
|-------|------|-------|--------------------|--------------------|---------|-----------|-------|---------|------------------|
| 1     | (Wijsman et al., 2018) | 2018* | NA                 | Netherlands        | W56     | Traveller | 1     | Yes     | T. b. (unknown)  |
| 2     | (Wang et al., 2018)    | 2017  | August             | China              | M60     | worker    | 2     | NA      | T. b. gambiense   |
| 3     | (Lu et al., 2018)      | 2017  | August             | China              | W41     | Traveller | 1     | Yes     | T. b. rhodesiense |
| 4     | (Streit and Matsumoto, 2016) | 2016* | December           | USA                | W60     | Traveller | 1     | NA      | T. b. rhodesiense |
| 5     | (Launet et al., 2017)  | 2016  | August             | UK                 | W58     | NA        | 2     | NA      | T. b. gambiense   |
| 6     | (De Kuyser et al., 2016) | 2016* | May                | France             | M14month| Delivery  | 2     | NA      | T. b. gambiense   |
| 7     | (Gomez-Juynent et al., 2017) | 2015  | August             | Spain              | W49     | Traveller | 1     | Yes     | T. b. rhodesiense |
| 8     | (Hu et al., 2016)      | 2014  | September          | China              | M45     | Sailor    | 2     | Yes     | T. b. gambiense   |
| 9     | (Pale et al., 2013)    | 2013* | August             | Argentina          | M65     | Traveller | NA    | NA      | T. b. rhodesiense |
| 10    | (Pasternak et al., 2013) | 2013* | January            | Canadian           | M       | Traveller | 1     | NA      | T. b. rhodesiense |
| 11    | (Clerinx et al., 2012) | 2012  | February           | Belgium            | M       | Traveller | NA    | Yes     | T. b. rhodesiense |
| 12    | (Wolf et al., 2012)    | 2012  | January            | Germany            | M62     | Traveller | 1     | NA      | T. b. rhodesiense |
| 13    | (Meltzer et al., 2012) | 2012* | NA                 | Israeli            | W31     | Traveller | 1     | NA      | T. b. rhodesiense |
| 14    | (Richter et al., 2012) | 2012* | March              | Germany            | M58     | Traveller | NA    | NA      | T. b. (unknown)   |
| 15    | (Cottle et al., 2012)  | 2012* | February           | UK                 | W49     | Traveller | 1     | NA      | T. b. rhodesiense |

*: Time refers to the published date when the diagnostic date is unavailable.
NA: Information is unavailable on the case report.

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Table 2. Summary of *rhodesiense*-HAT cases in eastern African countries but reported in endemic countries or non-endemic countries.

|                      | 1990-2000 |          | 2001-2010 |          | 2011-2018 |          | Total in non-DECs |
|----------------------|-----------|----------|-----------|----------|-----------|----------|-------------------|
|                      | DECs | Non-DECs | DECs | Non-DECs | DECs | Non-DECs | DECs | Non-DECs |
| Kenya                | 164  | 1        | 23   | 0        | 2     | 2        | 3     |          |
| Malawi               | 736  | 1        | 465  | 14       | 180  | 0        | 15    |          |
| Rwanda               | 0    | 5        | 0    | 0        | 0    | 0        | 5     |          |
| Tanzania             | 3424 | 6        | 1294 | 49       | 16   | 2        | 57    |          |
| Tanzania or Kenya    | **   | **       | **   | **       | **   | **       | 1     | 1        |
| Tanzania, Kenya or Rwanda | **   | 1       | **   | **       | **   | **       | 1     |          |
| Uganda               | 5458 | 0        | 2660 | 2        | 319  | 0        | 2     |          |
| Zambia               | 41   | 1        | 81   | 7        | 42   | 2        | 10    |          |
| Zimbabwe             | 10   | 0        | 8    | 5        | 22   | 1        | 6     |          |
| Zimbabwe or Botswana | **   | **       | **   | **       | **   | **       | 1     | 1        |
| Total in non-DECs    | **   | 15       | **   | 77       | **   | 9        | 101   |          |

DECs=Disease endemic countries; Non-DECs= Non-disease endemic countries.
Table 3. Summary of *gambiense*-HAT cases in west and central African countries but reported in endemic countries or non-endemic countries.

|                | 1990-2000 |          | 2001-2010 |          | 2011-2018 |          | Total in non-DECs |
|----------------|-----------|----------|-----------|----------|-----------|----------|------------------|
|                | DECs      | Non-DECs | DECs      | Non-DECs | DECs      | Non-DECs |                  |
| Angola         | 43017     | 1        | 18048     | 7        | 402       | 0        | 8                |
| Cameroon       | 360       | 1        | 174       | 3        | 52        | 0        | 4                |
| Central African Republic | 6320 | 0        | 6990      | 2        | 1090      | 0        | 2                |
| DRC            | 176073    | 1        | 99451     | 6        | 25635     | 1        | 8                |
| Eq. Guinea     | 515       | 0        | 165       | 2        | 13        | 0        | 2                |
| Gabon          | 451       | 1        | 305       | 7        | 81        | 2        | 10               |
| Nigeria        | 65        | 1        | 107       | 0        | 6         | 1        | 2                |
| Nigeria or Gabon | **      | 1        | **        | **       | **        | **       | 1                |
| Sudan          | 6073      | 0        | 14149     | 1        | 843       | 0        | 1                |
| Uganda         | 14790     | 0        | 2928      | 2        | 90        | 0        | 2                |
| Total in non-DECs | **      | 6        | **        | 30       | **        | 4        | 40               |

DECs=Disease endemic countries; Non-DECs= Non-disease endemic countries.
Figure legends.

**Fig. 1.** Total number of reported cases of HAT in endemic countries from 1990 - 2018.

**Note:** Data taken from the World Health Organization.

**Fig. 2.** (A) Number of reported cases of *gambiense*-HAT per year from 1990 -2018 in endemic countries with a breakdown for each country. (B) Number of reported cases of *rhodesiense*-HAT per year from 1990 -2018 in endemic countries with a breakdown for each country. Data taken from the World Health Organization (WHO).
**Fig. 3.** Distribution of human African trypanosomiasis in endemic countries, 2017 and 2018. Data taken from the World Health Organization (WHO).
Fig. 4. HAT cases that were diagnosed and confirmed in non-endemic countries from 2011–2018. (A) (B) Dots represent the place of diagnosis; (C) Triangles indicate the probable place of infection. Numbers refer to the designations in Table 1.
**Fig. 5.** Number of HAT cases diagnosed in non-endemic countries in different periods.

**Fig. 6.** HAT cases reported in non-endemic countries in different time periods.

**Fig. 7.** The dynamic trend changes in HAT cases diagnosed in non-DECs and DECs.

Bar graphs indicate the annual figures. The line graphs indicate the distribution of 3 year averages.