Effects of Political Instability in Venezuela on Malaria Resurgence at Ecuador–Peru Border, 2018

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Mass migration from Venezuela has increased malaria resurgence risk across South America. During 2018, migrants from Venezuela constituted 96% of imported malaria cases along the Ecuador–Peru border. *Plasmodium vivax* predominated (96%). Autochthonous malaria cases emerged in areas previously malaria-free. Heightened malaria control and a response to this humanitarian crisis are imperative.

_Malaria_ is a vectorborne parasitic infection caused by *Plasmodium* spp. and transmitted by *Anopheles* mosquitoes, characterized by fever and hemolysis with chronic and fatal potential (7). Despite substantial strides toward elimination in the Americas, malaria remains a major concern; ≈975,700 cases occurred and 138 million persons were at risk in 2017 (2). Most malaria cases in South America occur in the Amazon region, and _P. vivax_ is more common than _P. falciparum_ (3).

*P. vivax* and _P. falciparum_ malaria were historically endemic to the Ecuador–Peru coastal border region. During 1990–2012, a total of 62,000 malaria cases were reported from El Oro Province, Ecuador, and 85,605 from Tumbes Region, Peru (4). Through vector control and active case surveillance and response, malaria was eliminated from El Oro Province in 2011 and Tumbes Region in 2012 (4). However, malaria cases elsewhere in Ecuador increased from 378 in 2013 (5) to 1,279 in 2017 (6). Peru and other countries in the region also reported increased malaria in 2017, indicating a major risk for reintroduction to elimination areas (2). In 2017, Venezuela alone accounted for more than half of all malaria cases in the Americas (2).

The public health sector in Venezuela is struggling with infectious disease epidemics, including malaria (7), despite a historically successful malaria control program (3). The worsening social and economic crisis has led to large-scale migration from and within Venezuela. The shortage of antimalarial drugs and lax in-country control efforts have exacerbated the situation, affecting countries throughout South America (8). Many people from Venezuela are migrating through Colombia and Ecuador to reach Peru and the southern cone of South America, stopping at various locations along the way (Figure). We report a series of imported malaria cases in migrants from Venezuela and the first autochthonous cases of malaria in the Ecuador–Peru border region since local elimination.
During February–November 2018, seven malaria cases (6 P. vivax, 1 P. falciparum) were detected in adults in El Oro Province and reported to the Ecuadorian Ministry of Health (Appendix, https://wwwnc.cdc.gov/EID/article/25/4/18-1355-App1.pdf). Five cases occurred in recent migrants from Venezuela, and 1 was imported from Peru. The most recent case (no. 7), reported in November 2018, was autochthonous. Plasmodium spp. infection was confirmed at the national reference laboratory in Guayaquil, Ecuador. Active surveillance within 1 km of each case-patient’s residence revealed no acute cases, and collateral thick blood smears were negative. Entomologic teams documented Aedes aegypti and Culex spp. mosquitoes in the homes but no Anopheles mosquitoes. The residences all had basic infrastructure and no history of malaria since local elimination in 2011.

During May–October 2018, a total of 20 P. vivax malaria cases were detected in adults in Tumbes Region and reported to the Peruvian Ministry of Health (Appendix). Seventeen cases occurred in Venezuelan migrants now living in the province, and 3 were autochthonous cases in persons residing in Tumbes. An epidemiologic investigation revealed that the autochthonous case-patients had no history of travel outside of Tumbes Region.

We cannot definitively state whether the migrants from Venezuela were exposed to malaria in Venezuela or during transit. Regardless, this population represents a highly vulnerable group with complex treatment issues. Malaria should be considered in the differential diagnosis for febrile patients from Venezuela and for local populations in nearby parts of South America. The transience of the migrant population presents treatment follow-up issues. The incubation period for P. vivax malaria is 12–18 days and, for P. falciparum malaria, 9–14 days. Case-patients (Appendix) often exhibited inadequately or untreated malaria. Imported cases are the likely source of the locally transmitted cases in Tumbes Region and El Oro Province because the primary mosquito vectors (An. albimanus and An. punctimacula) remain abundant in this area (9). Another concern is relapse of dormant P. vivax hypnozoites, which can occur up to...
several years after initial infection \((I)\). Issues with primaquine (i.e., \(CYP2D6\)-poor metabolizers or hemolysis risk in patients with glucose-6-phosphate dehydrogenase deficiency) complicate treatment of dormant hypnozoites that cause relapse \((I)\). A new treatment, tafenoquine, which still causes hemolysis in glucose-6-phosphate dehydrogenase deficiency, was recently approved in the United States as a single dose for prevention of \(P.\text{ vivax}\) malaria relapse \((10)\), although this medication might not reach at-risk groups in South America. Ecuador and Peru currently follow the Pan American Health Organization guidelines regarding primaquine use (https://www.paho.org/hq/dmdocuments/2011/TreatmentGuidelines-2nd-ed-2010-eng.pdf).

Local ministries of health responded quickly to these cases and implemented case surveillance. However, reductions in resources after elimination of local malaria transmission in 2011–2012 severely limited malaria control efforts in Ecuador and Peru. Imported cases of malaria at the Ecuador–Peru border region pose a serious threat of continued resurgence in local transmission. We urge international solutions for Venezuela’s humanitarian crisis and augmentation of infectious disease surveillance and control along migration routes and in surrounding regions.

**About the Author**

Dr. Jaramillo-Ochoa is an epidemiologist working for the Ministry of Health for the Health District of the city of Machala, El Oro Province, Ecuador. His primary research interests include the epidemiology of vectorborne diseases and monitoring and evaluating vector-control interventions.

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**Rickettsia parkeri and Candidatus Rickettsia andeanae in Ticks of the Amblyomma maculatum Group, Mexico**

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**Anopheles and Culex human biting rates in southern Ecuador. Malar J. 2017;16:479. http://dx.doi.org/10.1186/s12936-017-2121-4**

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We report Rickettsia parkeri and Candidatus Rickettsia andeanae in ticks of the Amblyomma maculatum group collected from dogs in Sonora, Mexico. Molecular characterization of these bacteria was accomplished by DNA amplification and sequence analysis of portions of the rickettsial genes gltA, htrA, ompA, and ompB.

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Appendix

Clinical and Demographic Details for Malaria Cases from El Oro Province, Ecuador, and the Tumbes Region, Peru, in 2018

Case Descriptions: El Oro Province, Ecuador

Six cases of malaria were reported in El Oro Province from February to November 2018. Details are presented below and in Appendix Table 1.

Case 1: A 27-year-old woman from Venezuela, at 8 weeks’ gestation, entered Ecuador in February 2018 with fever, chills, diaphoresis, and headache. She endorsed a prior history of incomplete treatment for *Plasmodium vivax* and *P. falciparum* on multiple occasions in Venezuela (Ciudad Bolívar). Thick blood smear revealed *P. vivax*. She received treatment with chloroquine, and subsequent thick blood smears (on days 7, 14, 21, 28) were negative. She continued with chloroquine preventive treatment until delivery.

Case 2: A 22-year-old man from Venezuela entered Ecuador in June 2018, with no prior history of malaria. His symptoms began 2 days after entry into Ecuador, and included fever, chills, diaphoresis, and headache. He was initially seen at an outpatient clinic and was prescribed amoxicillin and paracetamol. Six days later, he presented to a public hospital in the city of Machala, El Oro Province, Ecuador, with arthralgias, productive cough, and jaundice. While a cough is not usually thought of in malaria presentation, and in diagnosis, this has been reported in the literature (1). Diagnostic tests for dengue and leptospirosis were negative. Thick blood smear revealed *P. vivax*. He completed unspecified antimalarial treatment and follow-up thick blood smears (days 7, 14, 21, 28) were negative.
Case 3: A 41-year-old woman from Venezuela entered Ecuador in May 2018 with a past medical history of malaria on at least 16 occasions. She reported incomplete treatment due to lack of antimalarial medications in Venezuela and did not finish antimalarial treatment when *P. vivax* malaria was initially diagnosed in the city of Guayaquil, Ecuador (185 km north of Machala). She arrived in late July to the city of Machala, where *P. vivax* malaria was diagnosed again. She received treatment with chloroquine and primaquine for 7 days, and follow-up thick blood smears were negative. According to the Ecuadorean Ministry of Health team, she was treated an additional 14 days with primaquine to ensure killing of hypnozoites.

Case 4: A 51-year-old man entered Ecuador from the Peruvian Amazon (Canton Andoas) in August 2018 with fever, chills, and headache. He reported a history of malaria on 36 occasions. On August 22, *P. falciparum* malaria was diagnosed in Machala via thick blood smear. He was treated according to Pan American Health Organization guidelines (https://www.paho.org/hq/dmdocuments/2011/TreatmentGuidelines-2nd-ed-2010-eng.pdf).

Case 5: A 33-year-old man entered Ecuador in August 2018, having left Venezuela 10 days earlier. He had worked in the malaria-endemic Amazon region of Venezuela and had a history of malaria on at least 6 occasions. He reported treatment only with chloroquine in Venezuela. Symptoms (fever, chills, headache) began about 1 month after his arrival in Ecuador, and thick blood smear on September 17 showed *P. vivax*. He received unspecified antimalarial treatment. Follow-up thick blood smears (days 7,14,21, and 28) were negative.

Case 6: A 32-year-old man was diagnosed with *P. vivax* malaria in Venezuela (Bolivar state) 5 months prior, where he received treatment with chloroquine for 3 days and primaquine for 7 days. He left Venezuela in July 2018 and entered Ecuador 6 days later. His symptoms began in November of 2018, and included fever, chills, and headache. At that time, he presented twice to a health center in Machala, Ecuador, with unremarkable workup, including negative thick blood smears, and was treated symptomatically with paracetamol. One week later, he returned to the health center with jaundice, hepatosplenomegaly, and altered liver function test results. Thick blood smear
on November 14 showed *P. vivax*. He was treated with chloroquine and primaquine. Two of 4 follow-up thick blood smears (days 7, 14, 21, and 28) were negative. It was thought this was a relapse of *P. vivax* from liver hypnozoites; however, autochthonous transmission cannot be excluded.

Case 7: A 66-year-old man residing in the border city of Huaquillas in El Oro Province is thought to have the first autochthonous case of malaria in the province since 2011. The patient self-treated symptoms with paracetamol and sought medical care after 4 days with no improvement. A thick blood smear on November 28 revealed *P. vivax*, and the patient was treated with primaquine. Local control measures were implemented by the Ecuador Ministry of Health. No additional cases were found in the 1-km area around the patient’s home.

**Tumbes Region, Peru**

Twenty cases of malaria were reported from May to November 2018 in the Tumbes Region, Peru (Appendix Table 2). Additional clinical details are unavailable at this time.

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### Appendix Table 1. Summary of malaria cases from El Oro Province, Ecuador, in 2018*

| Case no. | Age, y/sex | Origin | Pregnant | Month of entry to Ecuador, 2018 | Symptoms | Month of diagnosis, 2018 | History of malaria | Diagnosis | Follow-up |
|----------|------------|--------|----------|---------------------------------|----------|--------------------------|--------------------|-----------|----------|
| 1        | 27/F       | Venezuela | Yes      | Feb                             | Fever, chills, diaphoresis, headache | February       | Yes                  | P. vivax | Negative |
| 2        | 22/M       | Venezuela | NA       | Jun                             | Fever, chills, diaphoresis, headache, arthralgias, productive cough, and jaundice | June           | No                   | P. vivax | Negative |
| 3        | 41/F       | Venezuela | No       | May                             | No details | July                   | Yes                | P. vivax | Negative |
| 4        | 51/M       | Peru     | NA       | Aug                             | Fever, chills and headache | August        | Yes                  | *P. falciparum* | Negative |
| 5        | 33/M       | Venezuela | NA       | Aug                             | Fever, headache and chills | Septembe r    | Yes                  | P. vivax | Negative |
| 6        | 32/M       | Venezuela | NA       | Jul                             | Fever, headache and chills, jaundice, hepatosplenomegaly | November      | Yes                  | P. vivax | Negative |
| 7        | 66/M       | Huaquillas | NA | Local resident | No details | November | Yes | P. vivax | Negative |

*All cases were diagnosed by thick blood smear and received complete treatment. NA, not applicable.

### Appendix Table 2. Summary of malaria cases from Tumbes Province, Peru, 2018*

| Case no. | Age, y/sex | Origin | Symptoms | Date of diagnosis, 2018 | Diagnosis |
|----------|------------|--------|----------|--------------------------|-----------|
| 1        | 24/M       | Venezuela | Fever, chills, diaphoresis, headache | May        | P. vivax               |
| 2        | 59/M       | Venezuela | Fever, chills, diaphoresis, headache | May        | P. vivax               |
| 3        | 30/F       | Venezuela | Fever, chills, diaphoresis, headache | Jun        | P. vivax               |
| 4        | 53/F       | Tumbes   | Fever, chills, diaphoresis, headache | Jun        | P. vivax               |
| 5        | 18/M       | Venezuela | Fever, chills, diaphoresis, headache | Jul        | P. vivax               |
| 6        | 20/F       | Venezuela | Fever, chills, diaphoresis, headache | Jul        | P. vivax               |
| 7        | 26/F       | Venezuela | Fever, chills, diaphoresis, headache | Jul        | P. vivax               |
| 8        | 22/M       | Venezuela | Fever, chills, diaphoresis, headache | Jul        | P. vivax               |
| 9        | 25/F       | Venezuela | Fever, chills, diaphoresis, headache | Jul        | P. vivax               |
| 10       | 35/M       | Venezuela | Fever, chills, diaphoresis, headache | Jul        | P. vivax               |
| 11       | 22/M       | Venezuela | Fever, chills, diaphoresis, headache | Jul        | P. vivax               |
| 12       | 48/F       | Tumbes   | Fever, chills, diaphoresis, headache | Jul        | P. vivax               |
| 13       | 39/M       | Venezuela | Fever, chills, diaphoresis, headache | Aug        | P. vivax               |
| 14       | 21/M       | Venezuela | Fever, chills, diaphoresis, headache | Aug        | P. vivax               |
| 15       | 23/M       | Venezuela | Fever, chills, diaphoresis, headache | Aug        | P. vivax               |
| 16       | 18/M       | Venezuela | Fever, chills, diaphoresis, headache | Aug        | P. vivax               |
| 17       | 39/M       | Venezuela | Fever, chills, diaphoresis, headache | Oct        | P. vivax               |
| 18       | 22/M       | Venezuela | Fever, chills, diaphoresis, headache | Nov        | P. vivax               |
| 19       | 41/M       | Venezuela | Fever, chills, diaphoresis, headache | Nov        | P. vivax               |
| 20       | 35/F       | Tumbes   | Fever, chills, diaphoresis, headache | Nov        | P. vivax               |

*Additional details unavailable.