**Management of Travel-Related Infectious Diseases in the Emergency Department**

Laura Throckmorton\(^1\) • Jonathan Hancher\(^2\)

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**Abstract**  
**Purpose of Review** Emergency physicians generally have limited exposure to internationally acquired illnesses. However, travelers can present quite ill, and delays in recognition and treatment can lead to increased morbidity and mortality. This paper aims to summarize typical presentations of common international diseases and provide the emergency physician with a practical approach based on current guidelines.

**Recent Findings** In the treatment of traveler’s diarrhea, azithromycin has become the treatment of choice due to the growing antibiotic resistance. Intravenous artesunate was approved in 2019 under investigational new drug protocol for the treatment of severe malaria, and artemisinin-based combination therapies (ACTs) have become the first-line treatment for most cases of uncomplicated malaria. Since the 2015 outbreak, Zika has become a concern to many travelers, but the current treatment is supportive.

**Summary** Clinicians should be aware of a few noteworthy updates in the treatment of internationally acquired illnesses, but more importantly, they must recognize warning signs of severe illness and treat promptly. Future research on workup and disposition could help emergency physicians identify which patients need admission in well-appearing febrile travelers.

**Keywords** International infectious diseases • Traveler’s diarrhea • Febrile traveler • Emergency department • Malaria • Global Health

**Introduction**

While variable by practice setting, most emergency physicians in the USA have little regular exposure to internationally acquired illnesses. Therefore, their ability to recognize these illnesses can be limited. While basic practice guidelines are available, confirmatory diagnostic testing in the emergency department is limited, and some treatments might not be readily available. This article aims to assist the emergency physician identify internationally acquired illnesses based on travel history, signs, and symptoms and to help guide management based upon current guidelines and their practical applicability in the emergency department.

**Prevention**

While the emergency physician is unlikely to be providing pre-travel care in the USA, many EM physicians travel internationally and therefore need to be aware of primary prevention strategies for themselves and patients they treat while abroad. Country-specific guidelines are published by the Centers for Disease Control (CDC), but general considerations include vaccination, prophylactic antibiotics, and protection from insect bites or infected water sources.

**Mosquito-Borne Illnesses**

- Bite prevention: Travelers to areas with malaria, Zika virus, and other mosquito-borne illnesses can minimize their
risk by wearing long sleeves and pants, using bed netting, minimizing outdoor activities around dusk, and avoiding travel during the rainy season [1].

- Chemoprophylaxis: Medication selection should be guided by local resistance as well as patient preferences in timing of travel, side effects, and medical history. Table 1 outlines the medications currently approved in the USA for malaria prophylaxis and specific considerations for each medication.

- Standby emergency treatment (SBET): While chemoprophylaxis is the most efficacious in malaria prevention, SBET has also become an option for travelers who are traveling to low-risk areas who do not want to take chemoprophylaxis for the duration of their travel. Patients using this method should begin taking prescribed antimalarials if they develop a fever and seek a medical evaluation as soon as possible.

### Vaccinations

- Hepatitis A [3]: Recommended before most international travel at least 4 weeks prior to departure
  - Second dose after 6 months for long-term immunity

- Typhoid fever [4]: Recommended before most international travel and can be administered via oral capsules or injection
  - Particularly important due to rising antibiotic resistance of *S. typhi*

### General Approach to the Febrile Traveler

When treating a febrile patient with a history of travel, important considerations include travel location, timing, sick contacts, weather, activities partaken while abroad, pregnancy, and medical comorbidities. Many infectious diseases acquired

| Atovaquone-proguanil | Dosing | Pregnancy | Special Considerations |
|-----------------------|--------|-----------|------------------------|
| - Daily dosing         | - 1–2 days before through 7 days after | - Contraindicated | - Side effects uncommon |
| Chloroquine            | - Weekly dosing                        | - Safe in all trimesters | - More expensive |
|                        | - 1–2 weeks before through 4 weeks after| - Likely safe but second line | - Many areas with resistance |
| Doxycycline            | - Daily dosing                         | - - Photosensitivity | |
|                        | - 1–2 days before through 4 weeks after | - Inexpensive | |
| Mefloquine             | - Weekly dosing                        | - Safe in all trimesters | - Can also prevent rickettsia and leptospirosis |
|                        | - 2 weeks before to 4 weeks after       | - - Some areas of resistance | |
| Primaquine             | - Daily dosing                         | - Contraindicated | - Not recommended for psychiatric conditions, seizures, cardiac conduction abnormalities |
|                        | - 1–2 days before through 7 days after  | - Used in areas of > 90% prevalence of *P. vivax* | |
| Tafenoquione (newly FDA approved in 2018) | - Daily dosing for 3 days before travel and transitions to weekly through 1 week after return | - Contraindicated | - Contraindicated in G6PD deficiency |

Source: CDC [2]
abroad can also take several weeks to manifest. Practitioners should remember to test for common local illness, particularly influenza during flu season, and cover for a broad range of illnesses in toxic-appearing patients.

Workup is dependent on location and timing of travel to help determine which diseases are most likely. For cases in which many pathogens seem possible, we would recommend lab workup to include complete blood count with differential, basic metabolic panel; liver function tests; coagulation screen; blood smear; CK level; pregnancy test; influenza swab; blood cultures; urinalysis; urine culture; and chest x-ray. While many findings are nonspecific, these tests will likely identify patients with warning signs of severe illness and might help guide the practitioner toward the most likely pathogen.

Many internationally acquired illnesses cannot be definitively diagnosed in the emergency department given the similar features of many diseases and the delay in diagnostic results. In determining disposition, clinicians should consider warning signs of severe disease for the most likely cause of illness, and if present, admit. Many febrile illnesses can be managed in the outpatient setting, but patients should be given strict return precautions and close follow-up. If the patient has a travel history which places him or her at risk of dengue, NSAIDs should be avoided until it has been definitively ruled out due to the risk of progression to dengue hemorrhagic fever.

Malaria

Epidemiology and Transmission

Malaria is the most common febrile illness among travelers to endemic areas and is caused by the mosquito-borne parasites of the *Plasmodium* genus, primarily *P. falciparum*, *P. vivax*, *P. ovale*, and *P. malariae*. Cases in the USA have been increasing over the last few decades to over 2000 cases reported to the CDC in 2016. The majority of cases diagnosed in the USA are among travelers returning from Africa, particularly West Africa. *P. falciparum* has been identified in nearly 70% of infections with *P. vivax* being the second most common. Mortality in the USA is < 0.5% [6•].

Signs and Symptoms

Symptoms of malaria include fever, headache, chills, diaphoresis, myalgias, diarrhea, vomiting, and cough. The onset of symptoms is dependent on the *Plasmodium* species with *P. falciparum* typically causing the most severe symptoms. In confirmed cases in 2016, over 90% of those with *P. falciparum* reported onset of symptoms within 1 month of returning to the USA [6•]. However, nearly half of cases of *P. vivax* or *P. ovale* had onset of symptoms more than 1 month after returning to the USA likely due to reactivation of dormant liver parasites [6•]. Febrile seizures can occur in children but should be considered a warning sign of cerebral malaria in any age group.

Severe malaria definitions vary between the CDC and the World Health Organization (WHO), but diagnosis can be made with any of the following signs and symptoms [7, 8, 55]:

- Seizures, altered mental status, or other neurologic manifestations
- Acute kidney injury
- Hemoglobin < 7 g/dL
- ARDS
- Hypoglycemia (< 40 mg/dL)
- Acidosis
- Liver failure or severe jaundice
- Hemodynamic instability
- > 5–10% parasitemia

Of confirmed US malaria cases reported in 2016, approximately 15% were classified as severe disease, and seven people died [6•].

Management

The diagnosis of malaria is typically by blood smear but can also be done by polymerase chain reaction. Additional laboratory abnormalities can include anemia, thrombocytopenia, elevated transaminases, mild coagulopathy, and elevated BUN and creatinine. Lumbar puncture has limited utility in cerebral malaria as results can be normal or show only mild elevations in total protein and cell counts with mildly depressed glucose [9]. If there is any concern for cerebral malaria, the patient should be treated empirically as mortality is high even with treatment.

Recommendations for treatment of malaria are dependent on the presence of any severe features, local resistance, and patient comorbidities. Access to antimalarials in the ED is likely to heavily influence treatment as even many large tertiary referral centers do not have most antimalarial drugs stocked. If the patient took prophylaxis while abroad, a different antimalarial should be selected for improved efficacy and reduced toxicity. The CDC does have a Malaria Hotline (770-488-7788) for treatment assistance with a staff member on call 24/7.

Based on CDC and WHO recommendations, we would recommend the following treatment for confirmed or suspected cases of malaria:

- Uncomplicated malaria [2, 8, 10]
  - Artemether-lumefantrine: The only artemisinin-based combination therapy (ACT) approved in the USA

WHO recommends ACTs as the first-line therapy due to highest cure rate.
Alternative first-line medications in quinine susceptible areas

- Chloroquine
- Hydroxychloroquine

Alternative first-line medications in quinine-resistant areas

- Atovaquone-proguanil
- Mefloquine
- Quinine + tetracycline, doxycycline, or clindamycin

• Pregnancy [2, 8]

• Artemether-lumefantrine: Approved in 2018 as first-line treatment in second and third trimesters

Second-line drug in first trimester due to limited safety data [11]
- Quinine + clindamycin
- Mefloquine

• Severe malaria [2, 8]

• First line: Intravenous antimalarials.

Artesunate: First-line therapy for severe malaria but only became available in the USA in 2019 under investigational drug protocol

Not accessible in the ED; must be shipped from CDC

Quinidine: Production in the USA discontinued in 2017 [12]

- Second line: Artemether-lumefantrine (oral).

Interim treatment until IV Artesunate can be obtained from the CDC

If unable to swallow pill, NG tube should be placed in ED

- Third line: atovaquone-proguanil or quinine.

- Intravenous clindamycin and doxycycline have been used in the past, but they are not recommended for the initial treatment of severe malaria as the onset of action is greater than 24 h [2].

Anyone with confirmed *P. falciparum* or species not yet known should be admitted to the hospital [10]. Patients with signs of severe malaria likely need admission to an intensive care unit. Those with no previous history of malaria, immunocompromised patients, children less than five, and pregnant women are at the highest risk for developing severe disease or rapid deterioration, and admission should be strongly considered [2, 13, 14].

Dengue

Epidemiology and Transmission

Dengue is a febrile illness caused by a mosquito-borne flavivirus. It is endemic throughout the tropics and is estimated to cause symptoms in only one quarter of infections. According to the WHO, dengue is the second most common febrile illness in travelers returning from low- or middle-income countries [15]. There are four known serotypes (DEN1–4), and subsequent infection with a different serotype places the individual at higher risk of developing severe dengue.

Signs and Symptoms

Symptoms generally start 4–7 days following mosquito bite, and the disease consists of three phases [15, 16]:

1. Febrile phase
   - 3–7 days
   - High fevers (40°C), severe headache, pain behind the eyes, myalgias, arthralgias, vomiting, lymphadenopathy, rash

2. Critical phase
   - 24–48 h
   - Defervescence, capillary leak, hypovolemia, potential development of severe dengue (dengue hemorrhagic fever [DHF] or dengue shock syndrome [DSS])

3. Convalescent phase
   - Fatigue lasting days to weeks

Severe Dengue

Severe dengue (DHF/DSS) is rare and primarily seen in cases of secondary infection with a different serotype. Therefore, severe dengue is particularly uncommon in travelers and should only be expected in those with frequent travel to endemic areas. Nonetheless, severe dengue can be fatal, so awareness of certain features can help distinguish DHF and DSS from other severe febrile illnesses. These findings include pleural effusions, ascites, elevated hemoglobin in the setting of thrombocytopenia, low ESR, hepatomegaly, shock with narrow pulse pressure, petechiae/positive tourniquet test, mucosal bleeding, and DIC [15–17].
**Management**

Within the first 7 days of illness, diagnosis is made by RT PCR or NS1 antigen testing. IgM antibodies can be detected 4–7 days after onset of symptoms but can cross-react with other flaviviruses including Zika, West Nile, Yellow Fever, and Japanese Encephalitis [17]. Other laboratory findings include hemoconcentration, thrombocytopenia, leukopenia, and elevated liver enzymes [15, 16].

The treatment of dengue, including severe dengue, consists of supportive care, particularly fluid resuscitation, fever control, and management of bleeding. It is important to note that NSAIDS should not be used in the treatment of fever due to the increased risk of bleeding. Platelet transfusions are recommended for platelet counts < 10,000 in the setting of active bleeding. Prophylactic transfusions are not recommended [18].

Anyone developing warning signs of severe dengue including severe abdominal pain, persistent vomiting, hemorrhagic, or severe fatigue within the critical phase should be admitted and watched closely [15]. Admission should also be considered in the settings of pregnancy, extremes of age, or significant comorbidities. Still, most cases of dengue can be managed in the outpatient setting. Patients should be educated on symptoms of severe dengue and told to return due to the risk of rapid progression [15].

**Leptospirosis**

**Epidemiology and Transmission**

Leptospirosis is an aerobic spirochete transmitted by contact with infected animal urine through abrasions, mucous membranes, or ingestion of contaminated food or water. Those affected typically have a history of freshwater exposure such as wading through flood waters or participating in water sports. In the USA, only 100–150 cases are identified each year, of which 50% are in Puerto Rico [19]. Cases have also been identified in Hawaii, the Pacific Coast, and the South. Internationally, leptospirosis can be acquired in most tropical regions with the highest risk in Southeast Asia [20].

**Signs and Symptoms**

Leptospirosis should be considered in patients with rapid onset fevers, myalgias, and headache with recent freshwater exposure or return from Southeast Asia. Incubation period is generally 5 to 14 days following exposure [21].

Leptospirosis consists of two phases [20]:

1. Phase one: Abrupt onset fevers, myalgias, and headache 1–3 weeks following exposure and lasting 2 to 9 days, concluding with cessation of fevers

- Conjunctival suffusion: Conjunctival injection without inflammatory exudates
  - May occur in 15–80% of cases and is highly specific [22–24]
- Other symptoms: Vomiting, diarrhea, hepatosplenomegaly, lymphadenopathy, pharyngitis, rash

2. Phase two, the “immune phase”: Recurrent fever and potential development of complications:

- Pulmonary hemorrhage, ARDS, uveitis, optic neuritis, myocarditis, rhabdomyolysis, and Weil’s disease (jaundice and nonoliguric renal failure)

**Management**

Diagnosis can be made from blood culture during phase one and urine culture during phase two [19, 25]. Additional pathogen-specific testing is hospital dependent. Routine lab findings are nonspecific but can include thrombocytopenia, hypokalemia, hyponatremia, elevated amylase, transaminitis, and hyperbilirubinemia [20, 21]. An elevated creatinine kinase can be useful in distinguishing leptospirosis from other diseases as it is elevated in up to 50% of patients [56]. CSF can show lymphocytic or neutrophilic pleocytosis, mildly elevated protein, and normal glucose. CSF culture is generally positive in the first 10 days of illness [20]. Chest x-ray should also be obtained for any respiratory symptoms due to risk of pulmonary hemorrhage and ARDS [20].

Most cases of leptospirosis are mild and can be managed outpatient with doxycycline (100 mg) or azithromycin (500 mg) [19, 25]. Patients with pulmonary involvement, CNS infection, jaundice, renal failure, or age over 60 are at highest risk of death [25, 26] and should be admitted and given IV doxycycline (100 mg), penicillin (1.5 million IU), or a third-generation cephalosporin [19, 25]. Until Rickettsia is ruled out, doxycycline is generally recommended as the initial treatment. In patients with severe disease, corticosteroids can be considered, but recent studies have conflicting data on their benefit [27, 28].

**Enteric (Typhoid) Fever**

**Epidemiology and Transmission**

Enteric fever is a broader term encompassing both typhoid fever caused by Salmonella enterica serotype Typhi and Paratyphoid fever caused by Salmonella enterica serotypes Paratyphi A, B, or C. While S. typhi is more common,
S. paratyphi is becoming more prevalent particularly in South Asia and is not covered by the typhoid vaccines. Enteric fever is contracted through ingestion of contaminated food or water, and the highest risk is from visits to areas of poor sanitation. The CDC estimates 400 cases per year in the USA with over 70% of cases occurring in travelers returning from India, Bangladesh, or Pakistan [29•].

**Signs and Symptoms**

Enteric fever classically occurs in three stages:

1. Week 1: Fever, chills, bacteremia
2. Week 2: “Rose spots” and abdominal pain develop
3. Week 3: Hepatosplenomegaly, intestinal bleeding, and potential for ileocecal perforation

   - In those hospitalized with enteric fever, incidence of perforation can be as high as 10% [30, 31].

Enteric fever should be considered in patients who have traveled to an endemic area within the preceding 3 weeks and who are presenting with gastrointestinal symptoms accompanied by 3 or more days of fever. While most patients will complain of abdominal pain, diarrhea is not always seen, and patients can instead present with constipation. Other common symptoms include headache, cough, arthralgias, and myalgias [32].

**Management**

If enteric fever is suspected, workup should include complete blood count with differential, complete metabolic profile; coagulation screen; EKG; blood cultures; and stool culture. Findings that can help point to enteric fever include anemia, leukopenia with left shift (adults) or leukocytosis (children), elevated LFTs, high fever (> 40 °C), and bradycardia [32].

Definitive diagnosis is made by blood culture, but this test has low sensitivity and will not provide a diagnosis in the ED. Therefore, patients with suspected enteric fever should be treated empirically based on clinical suspicion [57].

- Uncomplicated enteric fever: Oral azithromycin 1 g (followed by 500 mg daily ×5–7 days)
  - Second line: Fluoroquinolone if acquired in a region with low resistance
- Severe illness with systemic symptoms requiring hospitalization: Ceftriaxone 2 g or cefotaxime 1–2 g
  - Patients with altered mentation or signs of shock should also be given dexamethasone 3 mg/kg as this has been shown to dramatically reduce mortality [33, 34].
  - Recent travel to Pakistan: Carbapenem or azithromycin
  - Ongoing outbreak of multidrug-resistant strain of S. typhi since 2016 [35]

**Traveler’s Diarrhea**

**Epidemiology and Transmission**

Traveler’s diarrhea is the most common illness seen in individuals traveling from developed to resource-limited regions [36], occurring in up to 40% of travelers [37, 38•]. Transmission is fecal-oral, most often by food and water in regions with suboptimal sanitation and hygienic practices [39]. The highest risk regions include South and Southeast Asia, Africa (excluding South Africa), South America, Central America, and Mexico. Food from street vendors and staying in “all-inclusive” lodgings are specific risk factors for developing the illness [40].

Most episodes occur between 4 and 14 days after arrival to a resource-limited region [41]. Acute illness is most frequently caused by bacteria but can also be caused by parasites or viruses. Worldwide, the most common cause is enterotoxigenic Escherichia coli (ETEC), followed by Salmonella, Campylobacter jejuni, and Shigella [39]. Practitioners should consider geographic variation, as Campylobacter species are more common than ETEC in Southeast Asia [42].

**Signs and Symptoms**

Classic traveler’s diarrhea is defined as three or more unformed stools in 24 h plus at least one of the following: nausea, vomiting, abdominal pain, fever, or blood in stool. Symptoms are variable and partially dependent on the causative agent [39].

ETEC classically consists of malaise, anorexia, abdominal cramps followed by sudden onset watery diarrhea with very frequent stools, typically without blood or purulence. Patients may have a fever, nausea, or vomiting. Campylobacter or Shigella cause inflammatory diarrhea, which can present with similar symptoms but may progress to fever, tenesmus, or bloody diarrhea [39].

**Management**

Testing should include a basic metabolic panel to assess for dehydration or metabolic derangement. The determination of
microbiologic agent is typically unnecessary as ETEC cannot be distinguished from nonpathogenic *E. coli* on routine stool cultures [43]. Whether or not to pursue further testing should be based on clinical judgment and will likely include a shared decision-making conversation with the patient. Stool testing in the ED is reasonable when patients present with severe diarrhea, bloody or mucoid stools, antibiotics in preceding 8–12 weeks, systemic illness, or symptom lasting longer than 10–14 days. Recommended tests include:

- Stool culture to evaluate for *Campylobacter* or *Shigella*.
- Hospital dependent tests for ETEC or Shiga toxin.
- Stool O&P for *Giardia lamblia*, *Cyclosporidium*, *Isospora*, and other parasites.
- If recent antibiotic history, test for *Clostridioides* (*Clostridium*) *difficile*
- If the patient appears systemically ill, send blood cultures to evaluate for bacteremia, most commonly seen from *Salmonella* species (Typhi).

Consider admission for those patients with laboratory evidence of severe dehydration, acute kidney injury, need for electrolyte replacement and inability to tolerate orals, or systemic illness.

The treatment is typically symptomatic and supportive, as the vast majority of episodes is self-limited and resolves within three to 5 days. Although antibiotic stewardship is important, it is very reasonable to fill a 3-day antibiotic prescription prior to travel and start the antibiotic within 1–2 days of symptoms [39, 43]. Management includes the following:

- Fluid replacement, by mouth or intravenously.
- Antimotility agents such as loperamide or diphenoxylate can be helpful with those with frequent diarrhea.
- Antibiotics can reduce symptoms from several days to one or 2 days and are recommended for severe illness characterized by bloody or mucoid stool, or if symptoms substantially interfere with the purpose of travel [39, 43]. Options include:
  - Azithromycin 1000 mg once or 500 mg daily for 3 days
  - Fluoroquinolones: Levofloxacin 500 mg for one to 3 days or ciprofloxacin 750 mg once or 500 mg twice daily for 3 days

Poorly absorbed but remain alternatives for patients in whom fluoroquinolones or azithromycin are not appropriate.

### Chikungunya

#### Epidemiology and Transmission

Chikungunya is an arthropod-borne *Alphavirus* primarily transmitted by mosquito bites [44]. Interestingly, the name is derived from an African dialect meaning “stooched walk” due to the disease hallmark: debilitating joint and back pains. Chikungunya often occurs in outbreaks during the rainy season and has been seen globally in Africa, Asia, Europe, the Pacific Islands, and in the Americas [45].

Viremia occurs within a few days of infection, and the virus has a propensity to invade synovium, tenosynovium, and muscles [45]. The virus may linger in the joints for up to 2 weeks. Chronic arthritis develops in up to 60% of infected individuals [46].

#### Signs and Symptoms

Typically, symptoms start with fever and malaise lasting 3–5 days which are followed by polyarthralgias and dermatologic symptoms lasting 7–10 days [45]. However, arthralgias can persist for weeks, months, or even years. The rash is typically macular but can be patchy, diffuse, or pruritic. It often starts on the limbs and trunk and may involve the face. Arthralgias are the hallmark of chikungunya. The joint pain is typically unilateral and symmetric, affecting distal joints. The axial skeleton is involved in 34–52% of cases [46, 47]. Severe complications (respiratory failure, myocarditis, renal failure, hemorrhage, acute hepatitis, meningoencephalitis, acute flaccid paralysis, seizures) and death can occur, more often in elderly patients with medical comorbidities [44].

#### Management

Diagnosis can be made with chikungunya viral RNA RT-PCR within the first week (sensitivity 100% and specificity 98%). If RT-PCR is negative or if the patient has had symptoms for 8 or more days, diagnosis is made by virus serology via ELISA or IFA. If testing for chikungunya, one should test for dengue virus and Zika virus as well [58]. Given the severe arthralgias associated with chikungunya, it may be prudent to rule out septic arthritis if significant effusion or asymmetry is present. Joint fluid analysis will be consistent with inflammatory arthritis [47].

Indications for admission include significant comorbidities or inability to ambulate. Treatment is primarily supportive.
with rest, fluids, and acetaminophen [59]. NSAIDs should not be used until dengue has been excluded [45].

**Zika**

**Epidemiology and Transmission**

Zika virus is an arthropod-borne flavivirus transmitted by mosquitoes. Transmission can also occur via the maternal-fetal route, sexual intercourse (vaginal, anal, oral), or direct exposure to blood [48]. Outbreaks have occurred in Africa, Southeast Asia, the Pacific Islands, the Americas, and the Caribbean, most notably during the 2016 Olympic Games in Brazil [49, 50].

**Signs and Symptoms**

Approximately 20–25% of infected individuals have symptoms of infection, which are typically mild and last 2 to 7 days [51]. These include fever, pruritic rash, arthralgias, conjunctivitis, myalgias, headache, dysesthesia, and generalized weakness. Less commonly symptoms include abdominal pain, nausea, diarrhea, or mucous membrane ulcerations [52]. Zika has been implicated in serious neurologic complications such as congenital microcephaly, Guillain-Barré syndrome, myelitis, and meningoencephalitis [53].

**Management**

Zika can be diagnosed by PCR and serology testing. However, cross-reactivity with other flaviviruses is common, and testing is generally not recommended from the ED [54]. If labs are sent, CBC may show thrombocytopenia. Treatment is supportive with fluids and acetaminophen. Generally, patients will be discharged, and we recommend education surrounding transmission and pregnancy prevention. If the patient is suffering from a severe neurologic complication, we recommend brain/spine imaging, lumbar puncture, and admission.

**Conclusions**

The differential diagnosis in a febrile international traveler is broad, and in many cases, definitive diagnosis will not be made in the ED. In addition, while the CDC and WHO have useful guidelines for diagnosis and management, these tests and treatment might not be available in the emergency department. Therefore, one must be aware of alternative treatments particularly for severely ill patients. In an otherwise undifferentiated febrile traveler, basic labs can be drawn to narrow the differential and screen for risk of development of severe disease. Recognition of development of severe disease is critical as these patients have high risk of complications and death.

**Compliance with Ethical Standards**

**Conflict of Interest** The authors declare that they have no conflict of interest.

**Human and Animal Rights and Informed Consent** This article does not contain any studies with human or animal subjects performed by any of the authors.

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