Rickettsia prowazekii Infections

Etiology

*Rickettsia prowazekii* is a pleomorphic, obligate intracellular, Gram negative coccobacillus in the family Rickettsiaceae and order Rickettsiales of the α-Proteobacteria. It belongs to the typhus group of the *Rickettsia*. The disease associated with this organism is usually called epidemic typhus, although some authors use the term "sylvatic typhus" for human illnesses associated with flying squirrels. Strains of *R. prowazekii* may differ in virulence; however, a genetic analysis suggested that the strains found in squirrels are similar to other *R. prowazekii* and do not belong to a different subspecies.

Species Affected

In the U.S., the southern flying squirrel (*Glaucomys volans*) serves as a reservoir host for *R. prowazekii*. There was no evidence of infection in nearby eastern gray squirrels (*Sciurus carolinensis*) during some investigations. Antibodies were also absent from white-footed mice (*Peromyscus leucopus*), eastern ground squirrels (*Tamias striatus*), opossums (*Didelphis marsupialis*) and raccoons (*Procyon lotor*). One study found no evidence for *R. prowazekii* in northern flying squirrels (*Glaucomys sabrinus*) in California. Guinea pigs and voles of the genus *Microtus* can be infected experimentally by intraperitoneal inoculation, a route that bypasses normal immune defenses.

There is currently little or no evidence that *R. prowazekii* infects other animals. Some older studies (i.e., from the 1960s or earlier) reported finding antibodies to this organism in various domesticated and wild animals, including wild rodents. However, most commonly used serological tests cross-react with other species of *Rickettsia*, and the significance of these antibodies is unclear. In a few reports from the 1950s and 1960s, *R. prowazekii* was isolated from goats, sheep and a donkey, and from ticks collected from various livestock including ruminants and camels. Other groups were unable to isolate it from various domesticated or wild species, and one study found antibodies to other rickettsia but not *R. prowazekii*. Experimental infections, most dating before 1975, demonstrated limited or no replication in domesticated animals. Infections in lambs seemed to be limited and transient, with little replication of the agent. In one study, organisms were recovered after 5-6 days from the lung, spleen, liver and/or brain (but not lymph nodes or blood) of some lambs inoculated by intravenous or intraperitoneal routes, but only from the injection site of lambs inoculated subcutaneously. Attempts to demonstrate organisms in the blood of a limited number of experimentally infected dogs, donkeys, goat kids, calves and young camels failed. However, in some cases, animals were tested on only a few days and/or blood was not collected until 6-18 days after inoculation. Persistent seroconversion was reported in a donkey. Other animals also converted, but sometimes for only short
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periods before antibodies declined, with responses that were no stronger (and sometimes weaker) than to dead *R. prowazekii*. One group reported that rabbits could be infected experimentally with a very large dose of *R. prowazekii*, while another group could not detect organisms in the blood of rabbits given a lower dose.

**Zoonotic potential**

*R. prowazekii* affects people. In most parts of the world, humans are the only reservoir host for this organism.

**Geographic Distribution**

Human infections with *R. prowazekii* occur worldwide; however, cases in most countries now typically result from imported or recrudescent infections, rather than from local maintenance of the organism. Foci of endemic disease still exist in some resource-poor parts of Asia, sub-Saharan Africa, and Latin America. The situation in North Africa is unclear. At one time, endemic typhus seemed to have disappeared from this area; however, sporadic cases have since been reported in residents of the highlands of Algeria and visitors to that region.

The only known animal reservoir for *R. prowazekii*, the southern flying squirrel, occurs in eastern North America, including the U.S. and southern Canada. Zoonotic human cases have, to date, been reported only from the U.S.

**Transmission**

*R. prowazekii* is transmitted between people by the human body louse (*Pediculus humanus corporis*). Person-to-person transmission does not seem to occur when lice are absent. Lice become infected when they feed on the blood of an infected person, and excrete *R. prowazekii* in their feces after 2 to 6 days. Lice defecate when they feed, and organisms from louse feces or crushed lice can enter the body through the bite wound, other breaks in the skin or mucous membranes. *R. prowazekii* is also infectious by inhalation. Infected lice die prematurely within 2 weeks. However, *R. prowazekii* can survive in louse feces and dead lice for several weeks: in one study, this organism was still detected after 100 days. It can be transmitted between life stages of the louse (transstadial transmission), but does not pass to a new generation in eggs. Other vectors might be able to transmit *R. prowazekii*, although their role, if any, is thought to be minimal. Human head lice (*Pediculus humanus capitis*) have been infected in the laboratory, and can shed this organism. It has also been detected in various species of ticks. Transmission is theoretically possible in blood transfusions, during the period when rickettsia occur in the blood.

*R. prowazekii* can become latent in people, especially those who are not treated with antibiotics. There is no indication that latent carriers can pass the organism to lice while the infection is quiescent; however, *R. prowazekii* can become reactivated years after infection, probably when stress, concurrent illness or other factors cause immunity to wane. At this time, the organism can reappear in the blood and be transmitted to lice. Where *R. prowazekii* persists in people is not known, but studies in mice suggest it might be harbored in adipose tissues. Transplantation of organs containing latent rickettsia is a theoretical route of exposure.

*R. prowazekii* is thought to spread between flying squirrels via squirrel lice (*Neohaematopinus scuiropteri*). How squirrels transmit the disease to humans is still unclear. *N. scuiropteri* does not feed on humans, but squirrel fleas (*Orchopeas howardi*), other mammalian fleas and some mites can be infected in the laboratory and might act as mechanical vectors. Another possibility is contact with organisms in infected, dried squirrel louse feces (e.g., via inhalation), or squirrel tissues. Zoonotic infections could be transmitted between people if human body lice were present: zoonotic strains of *R. prowazekii* are capable of infecting these lice. Infections acquired from squirrels can become latent in humans, and later re-emerge.

**Disinfection**

*R. prowazekii* is expected to be susceptible to 1% sodium hypochlorite, 70% ethanol, 2% peracetic acid, 3-6% hydrogen peroxide, iodine, glutaraldehyde and formaldehyde, based on the effectiveness of these agents against other Gram negative prokaryotic organisms. *R. prowazekii* can also be inactivated by moist heat (121°C [250°F] for a minimum of 15 minutes) and dry heat (170°C [338°F] for a minimum of an hour).

**Infections in Animals**

Little has been published about the effects of *R. prowazekii* on southern flying squirrels; however, exposure seems to be common in some squirrel populations in the eastern U.S. Infections usually peak in the fall and early winter, when squirrel populations are concentrated in nests. One study reported that southern flying squirrels inoculated intraperitoneally with the highest doses of *R. prowazekii* died. No clinical signs were mentioned in squirrels that received more moderate doses and had rickettsia in the blood.

Experimental infections of domesticated animals including dogs, lambs, goat kids, calves, donkeys and young camels, did not result in clinical signs.

**Infections in Humans**

**Incubation Period**

The incubation period is 1-2 weeks, with most infections becoming evident after 10-14 days. Recrudescence cases can occur years after the initial infection.

**Clinical Signs**

The symptoms often begin suddenly, with nonspecific initial signs that may include headache, chills, fever, myalgia (which is often severe) and malaise. Some, but not
all, patients develop a rash after a few days. Small pink macules usually appear first on the upper trunk or axillae, and can spread to almost the entire body, typically sparing the face, palms and soles. As the disease progresses, the rash can become dark and maculopapular, petechial or even purpuric. However, some patients only have a transient rash. Petechiae may occasionally appear on the conjunctiva and/or soft palate. Rashes are more difficult to detect on darker skin. Other frequently reported signs include arthralgia, splenomegaly, and abdominal pain. Nausea, vomiting, and mild thrombocytopenia. Coughing is common, and some patients develop pulmonary complications including secondary bacterial pneumonia. Conjunctivitis, diarrhea and hematuria have been reported. CNS signs of varying severity (e.g., confusion, drowsiness, seizures, coma, hearing loss) can be seen in some patients. There may also be other serious complications, including shock, renal dysfunction, myocarditis and multiple organ dysfunction. Ischemia from vasculitis can result in gangrene, usually symmetrical, of the distal fingers and toes. More extensive involvement of the extremities is possible but uncommon. The acute, febrile stage of epidemic typhus lasts approximately 2 weeks in untreated patients, but full recovery may be slow, sometimes taking up to 2-3 months. A significant number of untreated cases are fatal. Residual neurological defects are reported to be rare in patients who survive.

Brill–Zinsser disease (recrudescent typhus) resembles acute epidemic typhus, but tends to be milder, with a lower risk of death and a shorter course. Epidemic typhus acquired from flying squirrels has also been milder and shorter in many cases, although severe illnesses have been reported. Rash was barely visible or absent in some people with the zoonotic form.

### Diagnostic Tests

Epidemic typhus is often diagnosed by serology. A fourfold rise in titer is diagnostic. Titters usually become detectable during the second week; thus, this test is most useful for retrospective confirmation. A number of serological tests have been described. They include indirect fluorescence antibody (IFA) tests, various agglutination tests (e.g., plate microagglutination, latex agglutination) and enzyme immunoassays such as ELISAs and dot blot assays. Complement fixation tests were also used in the past, and may still be employed in some countries. *R. prowazekii* can cross-react with *R. typhi*, the agent of murine typhus, in some tests. Differential reactions in immunoblotting, and cross-absorption assays, performed at reference laboratories, can distinguish these reactions. Patients with Brill–Zinsser disease usually have elevations in specific IgG but not IgM.

Organisms can be identified directly in tissue samples, including skin biopsies of the rash, with immunohistochemical or immunofluorescent staining, or polymerase chain reaction (PCR) assays. Blood can also be collected for PCR, which can distinguish *R. prowazekii* from other rickettsia, including *R. typhi*. Culture and identification of *R. prowazekii* is not usually available outside reference laboratories, as rickettsia are both fastidious and dangerous to laboratory personnel. If necessary, this organism can be cultured with a shell-vial assay using L929 cells. Inoculation into guinea pigs or embryonated eggs was commonly used in the past.

Test availability may be limited at diagnostic laboratories in areas where epidemic typhus is uncommon.

### Treatment

Epidemic typhus can be treated with tetracyclines. Chloramphenicol may also be used, but it is reported to be less effective, and also carries a risk of serious drug-related side effects.

### Prevention

Outside North America, epidemic typhus can be prevented by avoiding human lice. Bathing removes lice from the body, and insecticide treatment may also be recommended. Lice on clothing can be killed by laundering at 50°C (122°F). Because body lice are obligate parasites of humans and usually die within 5 days in the environment, another possibility is to keep infested clothing isolated for a week. In North America, flying squirrels should not be allowed to take up residence in attics, porches, walls or other areas of human residences. Close contact with these animals should be avoided.

There are no commercial vaccines for epidemic typhus. Experimental vaccines have sometimes been produced by the military (e.g., in the U.S.) and might be available for high-risk situations. The relevant authorities should be consulted for current information about vaccines.

### Morbidity and Mortality

In most parts of the world, epidemic typhus now occurs only as sporadic cases, originating from recrudescent or imported infections. Occasional zoonotic cases are also described in North America, typically between November and February.

Outbreaks usually occur in populations living in unsanitary, crowded conditions where lice are difficult to control. They are often associated with wars, famines, floods, and other disasters, and can occur in refugee camps. These outbreaks can be explosive. Historical outbreaks sometimes caused thousands or even millions of deaths, and serious outbreaks can still be seen in some countries. In 1997, epidemic typhus is thought to have affected more than 40,000 people in Burundi. Most epidemics occur during the colder months, when closer contact and heavier clothing facilitate lice infestations and person-to-person transmission. Underreporting may be an issue in some remote areas, particularly when the illness is mild.

The severity of a clinical case can vary with general health, age and the form of the disease. Children under the age of 10 years tend to have relatively mild cases compared to adults, while elderly or debilitated patients are
particular susceptible to serious complications. Case fatality rates up to 60% or more have been reported in untreated primary infections; however, the highest mortality rates were described in highly vulnerable populations, such as the elderly, or people who were severely undernourished. Some sources estimate that, overall, the case fatality rate is probably 10-30% in untreated cases. It is reported to be < 5% when treated.

Recrudescent typhus and zoonotic typhus are often less severe than primary, human louse-transmitted typhus, with an estimated mortality rate of 1% in recrudescent disease. Recrudescent typhus (Brill-Zinsser disease) seems to be uncommon in patients who were treated with antibiotics effective against R. prowazekii. No fatalities have been reported, to date, in zoonotic typhus. Nevertheless, there have been some severe cases that required intensive care.

Internet Resources

- Centers for Disease Control and Prevention (CDC)
- Rickettsial (Spotted & Typhus Fevers) & Related Infections
- European Centre for Disease Prevention and Control (ECDC). Epidemic Louse-borne Typhus
- PHAC. Pathogen Safety Data Sheets
- The Merck Manual
- World Health Organization

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References

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