## Virginia Department of Health Q Fever: Overview for Healthcare Providers

Organism	Coxiella burnetii: obligate intracellular bacterium, gram-negative coccobacillus which can persist in
	spore-like form in the environment for years
Reporting to Public	Suspected or confirmed cases require <u>immediate</u> notification to the local health department (LHD).
Health	See <a href="http://www.vdh.virginia.gov/local-health-districts/">http://www.vdh.virginia.gov/local-health-districts/</a>
Infectious Dose	1–10 organisms
Occurrence	Occurs worldwide.
	• In the United States, Q fever is rare, but is likely under-recognized; 156 cases were reported in
	2015. In Virginia, an average of 2.0 cases per year were reported during 2011–2015.
Natural Reservoir	Primarily sheep, cattle, and goats, but many other species (including cats, dogs, some wild
	mammals, and birds) can be infected.
	Tick vectors might be important for maintaining animal and bird reservoirs, but are not commonly
	associated with transmission to humans.
Route of Infection	<ul> <li>Most commonly by inhalation of contaminated airborne particles from birth products, excreta, or tissue. Also by inhalation of contaminated airborne particles from wool or bedding of infected animals.</li> </ul>
	Laboratory exposure through infective aerosols, droplets, or parenteral inoculation.
	Other reported routes: ingestion of unpasteurized dairy products from infected animals;
	transmission by blood or bone marrow transfusion; sexual transmission; laboratory transmission;
	possibly tick-borne transmission.
Communicability	Person-to-person transmission is extremely rare, but has occurred (e.g., during autopsy, delivery of
	baby).
Risk factors	Working with animals (e.g., livestock farms, meat processing plants, slaughterhouses, veterinary)
	clinics, animal research facilities), attending birth by infected animals, or living near livestock
	Consuming unpasteurized dairy products
	Handling infective laboratory specimens
	Chronic Q fever is more likely in those with valvular disease, blood vessel anomalies,
	immunosuppression, or infections during pregnancy.
Case-fatality Rate	Acute Q fever: low (< 2%) in untreated infections; negligible in treated infections
	• Chronic Q fever endocarditis: 25% –65% if untreated, <10% with appropriate treatment
Incubation Period	<ul> <li>Acute Q fever: Depends on dose, but typically 2 –3 weeks (range 3–39 days)</li> </ul>
	Chronic Q fever: months to years
Clinical Description	Severity varies and approximately half of infections are asymptomatic.
	Acute Q fever: nonspecific febrile illness usually accompanied by rigors, myalgia, malaise, and
	retrobulbar headache. GI symptoms (e.g., diarrhea, vomiting) might occur, particularly in children.
	Severe disease can include acute hepatitis, pneumonia and meningoencephalitis. Fever usually
	lasts 5-14 days but may continue for as long as 2 months. Placentitis and miscarriage possible in
	pregnancy.
	• Chronic Q fever (occurs in <5% of acute cases): endocarditis, hepatitis, osteomyelitis, post-Q fever fatigue syndrome
Differential Diagnosis	Variable depending on affected system
Radiography	Chest x-ray might be normal or have nonspecific abnormalities, including segmental or lobar
	consolidation (unilateral or bilateral), involving upper or lower lobes, or feature multiple or single
	opacities; pleural effusions are present in ~35% of cases.

Q Fever: Overview for Healthcare Providers VDH/OEPI/DSI

	Endocarditis may cause relatively small valvular vegetative lesions that are more easily visualized  with transpoort and a shape of the conditions are also as a shape of the condi
C	with transesophageal echocardiography than with a transthoracic echocardiogram.
Specimen Collection	A serologic diagnosis of acute Q fever is made by testing paired sera (acute-phase serum collected
and Laboratory	as soon as possible after symptom onset and convalescent-phase serum collected 2–3 weeks after
Testing	acute phase) to detect a 4-fold change or greater in phase II IgG antibody titer.
	• A serologic diagnosis of chronic Q fever is made by detecting elevated phase I IgG antibody >1:800
	(and is typically higher than phase II IgG) and an identifiable nidus of infection (e.g., endocarditis).
	PCR (whole blood, serum) and immunohistochemistry staining (tissue) can also be used to
	facilitate diagnosis of acute or chronic Q fever.
	• Culture is not recommended for routine diagnosis because it is difficult, time consuming, and requires Biosafety Level (BSL) 3 precautions.
	• If Q fever is suspected, notify the <u>LHD</u> immediately to discuss the case and laboratory testing.
	Specimens may be sent to Division of Consolidated Laboratory Services (DCLS) after VDH approves
	testing. For questions about specimen collection, the DCLS Emergency Officer can be reached 24/7
	at 804-335-4617.
Treatment	• Doxycycline is the preferred treatment for Q fever in non-pregnant adults and children ≥8 years of
	age. For details, refer to CDC's Diagnosis and Management of Q Fever — United States, 2013
	(http://www.cdc.gov/mmwr/pdf/rr/rr6203.pdf) and consult with package inserts.
	Treatment is most effective at preventing severe complications if it is started within 3 days of
	symptom onset.
	Treatment of acute Q fever should be initiated when Q fever is suspected and should not be
	withheld while laboratory testing is pending. Serologic monitoring is recommended for patients
	with acute Q fever, especially those at high risk of developing chronic Q fever.
	Treatment of chronic Q fever should be initiated after diagnostic confirmation.
	Treatment of asymptomatic patients or those who have already recovered from their infection is
	not usually recommended, but it might be considered for patients at high risk for developing
	chronic Q fever.
Post-Exposure	• PEP following a known exposure to naturally-occurring <i>C. burnetii</i> is not recommended; however,
Prophylaxis	self-monitoring for symptoms and periodic serologic testing might be recommended.
	• PEP (e.g., doxycycline) following an intentional release of <i>C. burnetii</i> can be considered for those at
	high risk for exposure. PEP is considered effective only if administered within 8-12 days of
	exposure.
Vaccine	In the United States, a vaccine is not commercially available.
Infection Control	Use Standard Precautions for all patients.
	• If splashes of infected material are anticipated (e.g., during infant delivery), a face mask and eye
	protection (goggles or face shield) are recommended. C. burnetii is easily aerosolized. If aerosol-
	generating procedures are performed or if exposure to infected birth fluids is anticipated,
	additional precautions are recommended, including use of a fit-tested N-95 respirator and goggles
	or a face shield for eye protection. Use of an airborne infection isolation room, or at least a
	private room, during such procedures is preferable.

Q Fever: Overview for Healthcare Providers VDH/OEPI/DSI